AI in Health: State of the Art, Challenges, and Future Directions

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Summary

Introduction: Artificial intelligence (AI) technologies continue to attract interest from a broad range of disciplines in recent years, including health. The increase in computer hardware and software applications in medicine, as well as digitalization of health-related data together fuel progress in the development and use of AI in medicine. This progress provides new opportunities and challenges, as well as directions for the future of AI in health.

Objective: The goals of this survey are to review the current state of AI in health, along with opportunities, challenges, and practical implications. This review highlights recent developments over the past five years and directions for the future.

Methods: Publications over the past five years reporting the use of AI in health in clinical and biomedical informatics journals, as well as computer science conferences, were selected according to Google Scholar citations. Publications were then categorized into five different classes, according to the type of data analyzed.

Results: The major data types identified were multi-omics, clinical, behavioral, environmental, and pharmaceutical research (R&D) data. The current state of AI related to each data type is described, followed by associated challenges and practical implications that have emerged over the last several years. Opportunities and future directions based on these advances are discussed.

Conclusion: Technologies have enabled the development of AI-assisted approaches to healthcare. However, there remain challenges. Work is currently underway to address multi-modal data integration, balancing quantitative algorithm performance and qualitative model inter-pretability, protection of model security, federated learning, and model bias.

Keywords
AI; health; deep learning; machine learning; natural language processing; federated learning

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1 Introduction

Artificial Intelligence (AI) refers to a set of technologies that allow machines and computers to simulate human intelligence. AI technologies have been developed to analyze a diverse array of health data, including patient data from multi-omic approaches, as well as clinical, behavioral, environmental, and drug data, and data encompassed in the biomedical literature.

Because of the potential to automate many tasks currently requiring human intervention, AI has attracted considerable interest from a variety of fields. AI methodologies are now commonly used to aid in computer vision, speech recognition, and natural language processing (NLP). In healthcare, the rapid development of computer hardware and software applications over recent years has facilitated digitization of health data, providing new opportunities [1] for the development of computational models and opportunities to use AI systems to extract insights from data.

AI technologies can simulate human intelligence at a variety of levels. Both machine learning (ML) and deep learning (DL) are subsets of AI. ML allows systems to learn from data at the most basic level. DL is a type of ML which uses more complex structures to build models. Conventional AI approaches (such as expert systems), according to Obemeyer and Emanuel [2], can “take general principles about medicine and apply them to new patients” in a manner similar to medical students in their first year of residency. ML abstracts rules from the data, similar to what a physician might experience during his residency [2].

2 Materials and Methods

This review includes works published over the past 3 to 5 years, according to the number of citations on Google Scholar. From this pool, five major types of data used in AI for health were identified. These data types include multi-omics data, clinical data, behavioral/wellness data, environmental data, as well as research and development data. The current state of AI related to each data type is discussed, followed by associated challenges and practical implications that have emerged over the last two years. Opportunities and future directions based on these data types are discussed.
3 AI for Common Biomedical Data Types

3.1 Multi-omics Data

Multi-omics data [4] refers to the biological process where different “-omics” data, such as genomics, proteomics, transcriptomics, epigenomics, and microbiomics are jointly collected and analyzed. In comparison to conventional single omics approaches, multi-omics offer a comprehensive understanding of biological processes. Separate omics data sources can often characterize the same or closely related biological processes. In ML, this is referred to as a multi-view setting [5], where each omic is regarded as a separate view. To integrate these inputs, either data-based integration or model-based integration is required.

**Data-based Integration.** Concatenation of the data from all of views, with or without transformation, can result in a single model. This integrative approach has been used successfully to combine data from single-nucleotide polymorphisms (SNPs) and messenger ribonucleic acid (mRNA) gene expression into a single matrix and explore the relationship between SNPs and mRNA to predict a quantitative phenotype (e.g., drug cytotoxicity) using a Bayesian integrative model [6].

Similarly, Mankoo et al. [7] developed an integrative approach using a multivariate Cox least absolute shrinkage and selection operator (LASSO) to predict remission rates and survival in ovarian cancer by integrating copy number alteration, methylation, microRNA (miRNA) and gene expression data. This group performed a survival analysis with a selected set of variables using Cox regression based on a variable selection via LASSO [7]. Shen et al. [8] proposed the iCluster framework for subtyping glioblastoma with three omics data types: copy number, mRNA expression, and DNA methylation data. The iCluster framework assumes all the omics data share a common set of latent variables during joint dimension reduction and data integration.

**Model-based Integration.** In this approach, a separate model based on each data view is built, followed by the aggregation of the model outputs. For example, the analysis tool for heritable and environmental network associations (ATHENA) [9-11] performed genomic analyses by integrating different omics data such as copy number alterations, methylation, miRNA and gene expression to identify associations with clinical outcomes such as ovarian cancer survival. In the integration process, base models and neural networks were first constructed based on each type of omic data, followed by integrative model building [6]. Wang et al. [12] proposed a network fusion approach for cancer subtyping, which begins by constructing patient similarity matrices. These matrices are based on mRNA expression, DNA methylation, and miRNA expression data. Matrix building is followed by an iterative nonlinear procedure to integrate the three base similarity matrices into a unified matrix, with the goal of identifying patient subtypes. Dr ghici and Potter [13] proposed an ensemble approach to help predict drug resistance in HIV protease mutants. This approach builds a base of predictive models with structural features from an HIV protease–drug inhibitor complex and DNA sequence variants, and then performs majority voting according to the predictions of the base models.

Challenges, opportunities, and practical implications of AI in using multi-omics data. Despite the promising results that have been achieved so far, there are still many challenges to developing effective AI approaches for multi-omic data analysis.

* Because multi-omic data are highly heterogeneous, simple concatenation of raw data or model outputs from each view will miss the opportunity to explore the potential connections and relationships across entities in different views. Network-based approaches, which treat entities as nodes and their relationships as edges in the network, hold great promise for integrative analysis of multi-omic data [14]. Conventional network analysis algorithms, such as label propagation [15, 16], focus more on the edges/connections within the network. The recently proposed Graph Neural Network (GNN) [17], which considers both the node features and edge connections, would be of great interest in this context.

* Different from conventional weighted networks, edges (e.g., gene regulations and protein interactions) are usually rich contexts associated in a network constructed from multi-omic data. The incorporation of such contexts may complicate the analysis on the networks. Some typical network properties, such as edge weight non-negativity or transitivity, could be violated. Moreover, conventional network analysis assumes the network is pairwise, i.e., each edge only connects a pair of nodes in the network. However, in many scenarios we are also interested in investigating higher order interactions among different entities, for which case pairwise network analysis is not enough [18]. Therefore, there is huge potential to develop novel AI methodologies for analyzing multi-omics networks.

3.2 Clinical Data

AI technologies have also been used extensively in analyzing clinical data, including medical images, electronic health records (EHRs), and physiological signals.

3.2.1 Medical Images

Conventional ML approaches for analyzing medical images are often based on feature engineering, where features or descriptors of the medical images are extracted and then fed into the learning models for different tasks such as segmentation or classification. Due to advances that have revolutionized DL methodologies, an ever-increasing number of DL models have been incorporated into the medical image analysis pipeline. For example, Gulshan et al. [19] trained the Inception-V3 model [20], which is a deep learning model for natural image analysis, on a set of 128,175 renal fundus photographs for the identification of diabetic retinopathy. The authors demonstrated that, in two validation sets of 9,963 images and 1,748 images, the algorithm had 90.3% and 87.0% sensitivity, and 98.1% and 98.5% specificity, respectively. Esteva et al. [21] applied the same model to a set of skin images to enable discrimination between benign and malignant lesions. They designed a transfer-learning mechanism which pretrains the convolu-
tional layers of the Inception-V3 model with trained weights from ImageNet, and then retrains the final, softmax layer using a local skin image data set, fine-tuning the model parameters across all layers. Using 127,463 training images and 1,942 testing images, they demonstrated that the model can discriminate between benign and malignant lesions at a level of accuracy similar that of dermatologists. Interestingly, Kermany et al. [22] also adopted the same model and transfer learning strategy on two-dimensional optical coherence tomography images by freezing the parameters on the convolution layers after pretraining, without any fine tuning. With 108,312 training images and 1,000 testing images, the authors found that the model demonstrated an area under the receiving operating characteristic curve (AUC) of 99.9%. These three works demonstrate the power of end-to-end deep learning models for medical image classification through superior quantitative performance. In clinical decision support, numbers are not enough, as clinicians also need to know how the decision is made and decisions must be supported by evidence.

Recently, De Fauw et al. [23] proposed a novel two-stage deep learning architecture for diagnosis and patient referral (e.g., urgent, semi-urgent, routine, and observation only) of retinal disease. In the first stage, a deep segmentation network (3D Unet [24]) was developed to create a “detailed device-independent tissue segmentation map” from 3D Optical Coherence Tomography (OCT) images. Then a deep classification convolutional neural network (CNN) was constructed in the second stage to analyze the segmentation map and suggestions on diagnosis and patient referrals. After training the systems on only 14,884 scans, the approach was applied to patient triage and referral in an ophthalmology clinic. Compared with the conventional single-stage end-to-end framework, this two-stage approach derived a “device-independent segmentation of OCT scans” which serves as “intermediate representations that are readily viewable by a clinical expert” [23] and thus provides evidence for the second stage of disease diagnosis or patient referral. This facilitates the integration of the system into clinical workflows.

Challenges, opportunities, and practical implications of AI in using medical images. According to a recent report in *The Lancet*, a dermatologist may review over 200,000 images of skin lesions over decades of work, compared to mere days that it could take for a computer to analyze the same images using AI-assisted techniques [25]. ML approaches have also been used to successfully analyze raw images in cardiovascular imaging studies. By expanding the size and variety of cardiovascular imaging databases, new DL approaches can be developed, according to Heglin and colleagues [26].

Challenges remain regarding the use of AI in medical imaging. Analysis of medical images relies heavily on deep learning architectures that were designed and trained on natural images, such as the inception-V3 model discussed above. Medical images are also used to further fine-tune models. This enhances the model’s ability to recognize image patterns in the training data but may not be generalizable to new image patterns. Moreover, there are few dedicated DL model architectures for medical image analysis. An associated challenge is that training a brand-new model architecture typically needs a large number of images [26], which may not be easy to obtain in medical applications.

In addition to the model challenges, there are also data challenges. For example, differences in images from patients with different ethnicities (e.g., light vs. dark skins) may introduce disparities in the model’s decisions implicitly [27]. For example, if a skin lesion classification model is trained on a set of images composed of many more light skins than dark skins, it tends to perform better to classify light skins than dark ones.

### 3.2.2 Electronic Health Records

EHRs are systematic collections of longitudinal patient health information [28]. There are two types of information contained in patient EHRs: 1) structured information, which refers to the fields that contain data using existing lexicons, such as demographics, diagnosis, laboratory tests, medications, and procedures; and 2) unstructured information, which is typically free text documents such as clinical notes from physicians and nurses. In recent years, efforts have been devoted to developing AI methodologies for EHR analysis.

Conventional machine learning models for analyzing the structured information in EHRs are mostly vector based [29, 30], where patient records within a certain time window are collapsed into vectors composed of the summary statistics of the values of the features in different dimensions. One major limitation of this approach is that the temporality among the clinical events within EHRs is lost. To explore such temporality, Wang et al. [31] proposed to represent patient EHRs as longitudinal matrices with one dimension corresponding to the features and the other dimension corresponding to the time. Matrix factorization [31] or CNN type of approaches [32] were then developed to analyze such matrices. One big challenge for such matrix representation is the ultra-high sparsity. To handle such challenge, sequence modeling approaches, such as Recurrent Neural Networks (RNN) [33] have been used to analyze structured EHR data. Choi et al. [34] leveraged RNN to predict the onset risk of Congestive Heart Failure (CHF). To further enhance the model interpretability, they developed the Reverse Time Attention Model (RETAIn) [35] for modeling EHR sequences, so that the most recent clinical visits received the highest level of attention. Bekhet et al. [36] tested the generalizability of RETAIn on CHF onset risk prediction with a larger patient cohort. One limitation of RNN-based models is that they are not good at capturing long-term dependencies for the events in sequences. To solve this problem, Xiao et al. [37] leveraged TopicRNN [38], which combines RNN and global topic modeling to predict CHF patient readmission risk using EHR sequences, where each global topic corresponds to a specific distribution of the events in the EHR sequence.

Analyzing the unstructured information in EHR has been a long-standing topic in medical informatics. The conventional NLP approaches have been mostly rule-based or regular-expression-based. These methods typically need rigorous definitions of rules or regular expressions before the analysis. One challenge of these approaches is that it is impossible to enumerate all possible rules/regular expressions. In recent years, because
of the huge success of AI methods in NLP, more and more data-driven methodologies are developed for clinical NLP. For example, Kaur et al. [39] developed a NLP algorithm that can automatically identify patients who meet asthma predictive index (API) criteria from patient EHRs. Luo et al. [40] proposed to represent high-order semantic features from clinical texts as graphs and developed a subgraph-augmented nonnegative tensor factorization approach to analyze them. They also proposed a segmented CNN [41] and RNN [42] to process short clinical notes and achieved state-of-the-art performance on relation classification. Filannino and Uzuner [43] performed a survey on the shared tasks for clinical NLP and identified data-driven approaches for tackling those tasks. Soysal et al. [44] developed a clinical language annotation, modeling, and processing (CLAMP) toolkit for customized clinical NLP applications.

3.2.3 Physiologic Data

Physiologic data refer to the signals from processes such as electrocardiograms (EKGs) and electroencephalograms (EEGs). These signals are usually categorized as continuous, in terms of time and value. Conventional signal processing methods usually transform those continuous-time signals into vectors through some transformations (e.g., Fourier or wavelet transform [46-48]), and then build analysis algorithms on top of these vectors. Recently, deep-learning based technologies have been used to analyze raw signals. For example, Hannun et al. [49] proposed a 34-layer CNN model to map EKG signals to a series of rhythm classes to detect heart arrhythmia. Schwab et al. [50] proposed to tackle the same problem with RNN techniques. Schirrmeister et al. [51] proposed to leverage CNN modeling to encode and visualize EEG signals. To leverage more available data, Liang et al. [52] developed a transfer learning strategy that leverages EEG data sources for seizure prediction using CNN models.

3.3 Behavioral Data

In addition to multi-omics and clinical data, behavioral data is also linked to health status. While the use of behavior data in health applications poses some specific challenges, due to the way such data is collected and housed, there are some research teams that investigate the relationship between behavior data and health.

Social Media. The use of social media, such as Facebook, Twitter, LinkedIn, and Instagram may differ according to health status. For example, Sinnenberg et al. [55] identified associations between Twitter posts and the risk of cardiovascular disease. From a set of 4.9 million tweets, this group found that users with cardiovascular disease can be characterized by the tone, style, and perspective of their tweets, as well as some basic demographics. Ra et al. [56] found “a significant association between higher frequency of modern digital media use and increase in symptoms of ADHD (attention-deficit/ hyperactivity disorder) over a 24-month period” in adolescents between the ages of 15 and 16, as compared to baseline. Researchers have examined social media analytics and mental health, and they identified markers in social media activity associated with worsening psychotic symptoms [57], schizophrenia [58], risk of suicidal ideation [59], and depression [60].

Video and Conversational Data. Use of video and conversational data has gained the attention of many, both inside and outside of fields such as healthcare. Tencent, the Chinese tech giant, claims to have developed a vision system that can spot Parkinson’s Disease in 3 minutes [61]. Recently, a clinical trial involving extensive interviews between patients and trained medical staff using linguistic markers as screening tools for mild cognitive impairment (MCI) detection has shown promise [62, 63]. Tang et al. [64] built a conversational agent based on transcripts from these clinical trials using reinforcement learning techniques [65]. This agent was trained to maximize the diagnosis accuracy of MCI with a minimum number of conversational events, and the agent performed significantly better than supervised learning models.
Data with clinical data can provide a unique person’s EHR. Therefore, linking behavioral we can only confirm the disease from the from his/her posts on social media. However, truth labels. For example, we can judge data is the difficulty of obtaining the ground direction.

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One challenge for analyzing behavioral data is the difficulty of obtaining the ground truth labels. For example, we can judge whether a person is likely to have depression from his/her posts on social media. However, we can only confirm the disease from the person’s EHR. Therefore, linking behavioral data with clinical data can provide a unique opportunity to impact health, from both an individual and a population standpoint.

In addition to patient behavior, it is also interesting to analyze clinician behavioral data for the purpose of better quality of care delivery. Yeung et al. [73] proposed the concept of “bedside computer vision,” which utilizes computer vision technol-

3.4 Environmental Data

Environmental factors are important in a number of diseases, including cardiovascular disease [74], chronic obstructive pulmonary disease (COPD) [75], Parkinson’s Disease [76], psychiatric disorders [77], and cancer [78]. AI technologies have been used to explore environmental data to better understand disease mechanisms and improve care quality. For example, Song et al. [79] explored the effect of environment on hand, foot, and mouth disease through time-series analyses. Stingone et al. [80] studied the association between air pollution exposures and children’s cognitive skills in the United States using ML models. Park et al. [81] leveraged advanced ML models to construct environmental risk scores and applied them to metal mixtures, oxidative, and cardiovascular disease. Hahn et al. [82] developed multifactor dimensionality reduction software for detecting gene–gene and gene–environment interactions.

Challenges, opportunities, and practical implications of AI in using environmental data.

From the above summary, we can see that behavioral data are heterogeneous. Different types of behavioral data characterize a person from different aspects, thus the integrative analysis of behavioral data can provide us a more holistic view. Insel [72] proposed the concept of digital phenotyping, which “involves collecting sensor, keyboard and voice and speech data from smartphones to measure behavior, cognition and mood.” There will be many opportunities on this direction.

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3.5 Pharmaceutical Research and Development Data

Medications play important roles in healthcare. Data collected in various stages of drug development often contain insights about disease mechanisms and treatments. AI methodologies have been adopted to extract insights from those data. Drug data are presented below according to the information source (i.e., PubChem, clinical trials, and spontaneous reports).

Chemical Compounds. PubChem [83] is a website which lists information related to small molecules and their bioactivities. Many researchers use the molecular structures contained in PubChem as a vocabulary and then adopt a footprint (zero-one) or bag-of-words representation for the analysis of specific compounds. For example, Zhang et al. [84, 85] used footprint-based representations to calculate drug similarities and combined them with patient or disease similarities to achieve personalized treatment recommendations. Recently, graph convolutional networks (GCN) [86] have been applied in molecular structure design and analyses, where each molecule is treated as a graph, with the atoms as graph nodes. Duvenaud et al. [87] designed a GCN structure to extract features (referred to as neural fingerprints) from the molecules, with good prediction capability, parsimony, and interpretability using this approach. According to Kearns et al. [88], molecular graph convolutions “represent a new paradigm in ligand-based virtual screening.”

Clinical Trials. Clinical trials are a key step in drug development. The participants in clinical trials are usually selected with strict inclusion and exclusion criteria. Clinical trial data provide a wealth of information for each pharmaceutical company. Recently, AI approaches have been used in clinical trial design and data mining. For example, Chekrour et al. [89] adopted feedforward feature selection and gradient boosting in cross-trial prediction of treatment outcomes in depression. Kohannim et al. [90] investigated the usage of a support vector machine to boost the power of clinical trials and reduce the clinical trial sample size.

Spontaneous Reports. The FDA Adverse Event Reporting System (FAERS) [91] collects information on adverse events related to specific drugs. For the last decade, FAERS has been the major resource for conducting pharmacovigilance research. Sakaeda et
One limitation of clinical trials is that they
Although graph convolution approaches
above. We list a few of them below.
pharmaceutical R&D data as summarized
research, challenges still exist for analyzing
R&D data. Despite existing promising
names and outcomes to standard vocabu-
et al. [95] mapped drug
graphs obtained from side-effect profiles in
drug-drug interactions using drug similarity
oped a label propagation algorithm to predict
et al. [94] devel-
(tions using FAERS. Zhang
odding molecular, genomic, cellular, clinical,
avidity of de-novo
design, their interpretability remains
Specifically, in addition to
more efficient discovery of novel drug
cules, understanding associated
mechanisms of action is important. To
achieve this goal, we should incorporate
the domain knowledge from biology
and chemistry into the model building
process.

One limitation of clinical trials is that they
have very rigorous inclusion and exclusion
criteria for patient recruitment. The goal
is to eliminate the potential effect of con-
foundering factors. However, this will also
make the recruited patients “ideal” because
of the rigorous recruiting constraints,
and different from real world patients.
Similarly, FAERS data is composed of
a set of adverse drug reaction reports with
limited information. To make the insights
mined from clinical trial and FAERS data
more practical and useful, it is crucial to
link them with real world patient data
from EHRs or claims. FDA has released
a new strategic framework to advance the
use of real-world evidence to support the
development of drugs and biologics [96].
This will bring in lots of opportunities to
develop AI methodologies for the integrative
analysis of pharmaceutical R&D and
real-world clinical data.

3.6 Biomedical Literature Data
Published reports in the biomedical literature
are another important source of data for AI
in health applications. AI technologies and
NLP can be used to extract useful information
from the literature to inform health research. Many studies focus on biomedical
literature mining; for an early survey, refer
to Cohen and Hersh [97]. Recently, due to
the revolution of modern machine learning
approaches, such as deep learning, especially
in NLP; many advanced AI algorithms have
been developed in biomedical literature
mining and achieved state-of-the-art performance. There are two fundamental
problems on literature mining: (i) named entity recognition and normalization, which is the problem of identifying interested named entities
e.g., diseases, genes, genetic variants) in
the text and normalizing them (e.g., whether
two different textual descriptions correspond
to the same disease). For example, Leaman
et al. [98] developed DNorm, which is a
machine learning approach for disease name
normalization based on pairwise learning-
to-rank. The authors showed that comparing
with traditional lexical normalization and
matching approaches such as MetaMap [99]
and Lucene [100], DNorm can achieve an
improvement of 0.121 on micro-averaged F
measures. Recently researchers have also
shown that doing joint named entity recogni-
tion and normalization together can further
boost the performance of both tasks [101,
102]; (ii) relation classification, which is
the problem of identifying the relationships
among named entities once they have been
located in the literature. To deal with this
problem, Singhal et al. [103] developed a
rank aggregation approach to mine gen-
otype-phenotype relationships from bio-
medical literatures, and they demonstrated
a 28% performance improvement in terms
of F1 measures on benchmarks. Peng and
Lu [104] developed a multichannel depen-
dency-based CNN approach for extracting
protein-protein interactions from biomedical
literature searches and achieved a 24.4%
relative improvement in F1 measures over
the state-of-the-art methods.

Challenges, opportunities, and practical
implications of AI in using existing litera-
ture. In reality, a practical literature mining
engine would involve both components
we mentioned above, either explicitly or
implicitly. As an example, Zhang et al.
[105] developed a multi-view ensemble
learning pipeline to integrate the textual
features extracted from PubMed articles
with models to classify clinically action-
able genetic mutations found in specific
patients. However, because both tasks are
challenging, and the developed algorithms
are error-prone, the error could accumulate
across different stages in the pipeline and
may result in bad system performance.
Therefore, there is great potential on inte-
grated end-to-end learning of the model
parameters in different modules.

On the other hand, in contrast with the
various biomedical data we introduced in
previous sections, biomedical literature
serves as the knowledge source derived from
biological or clinical research. Injecting
mined knowledge from such sources into
the biomedical data modeling processes can
make the developed models more reliable
and generalizable. Tools such as PubMed
Phrases [106], PubMed Labs [107], and
 LitVar [108] have recently been developed to
facilitate research exploration of biomedical
literature, which provides an unprecedented
opportunity for the integration of knowledge
and data driven insights from biomedical
research.

4 AI in Health: Future Directions
4.1 Integrative Analysis
As Francis Collins envisioned in his vision
about the precision medicine initiative
[109], the next generation of scientists
will “develop creative new approaches for
detecting, measuring, and analyzing a wide
range of biomedical information — includ-
ing molecular, genomic, cellular, clinical,
behavioral, physiological, and environmental parameters.” Data from different modalities can describe a health problem from different aspects, and by integrative mining of those heterogeneous data, holistic and comprehensive insights into health can be obtained.

Recent years have seen an increase in research and initiatives related to AI in health, integrating different aspects of clinical data [110], linking biorepositories with clinical data [111-113], and forging connections between pharmaceutical research and development with clinical data [84]. More importantly, combining knowledge and data is the key to developing successful AI algorithms for health. In contrast to other computer fields such as vision and speech analysis, where large data sets can be obtained, patient data is often limited and can vary widely. In addition, real-world health problems are typically complex. To help offset this problem, the expertise from clinicians and biologists is necessary to inform the model’s learning process so that the model does not overfit the data.

4.2 Model Transparency
Traditional AI technologies, such as rule-based systems, are highly interpretable. Recent AI technologies, such as deep learning models, can achieve good quantitative performance, but are largely treated as black boxes. There are lots of debates recently on whether model interpretability is needed. For example, in a recent interview [114], Geoff Hinton, a pioneer in DL, argued that policymakers should not insist on ensuring people understand exactly how a given AI algorithm works, because “people can’t explain how they work, for most of the things they do.” Poursabzi-Sangdeh et al. [115] conducted a controlled randomized experiment to examine how important model interpretability is to users. Surprisingly, the results showed that there was no significant difference on users’ trust of black-box and transparent models. Moreover, “increased transparency hampered people’s ability to detect when a model has made a sizeable mistake.” Holm [116] defended black-box models by drawing the analogy with human decision-making process, where decisions are largely subjective (“outcomes of their own ‘deep learning’”). That’s why today “neuroscience struggles with the same interpretability challenge as computer science.”

According to the authors of the present article, there are certain areas where model interpretability may not be that important, especially in applications where AI algorithms have already demonstrated the capability to produce accurate results in a reliable and generalizable manner. However, this is not the case for health, at least in the current stage of the computational technology for healthcare analytics. For example, it has been shown that deep learning models can only achieve similar performance as logistic regression in hospital readmission tasks using EHRs [117] or claims [118]. Even for medical image analysis where deep learning models have achieved state-of-the-art performance, it is still difficult to justify the model generalization ability. That is, if the model works well on the medical image data set from one radiology center, it is not easy to justify it can still work well for another radiology center. Moreover, in most healthcare settings, final decision makers will still be human clinicians, and AI algorithms are just assisting them. Therefore, it is important to provide specific rationales for the propositions of those AI algorithms, to make the clinician feel more comfortable. Moreover, to enhance the clinical utility of AI algorithms, they should be integrated into regular clinical workflows [119].

On the other hand, the state-of-the-art performance of AI algorithms in many health applications are far from perfect. We should still encourage the exploration of black-box models to see if better performance can be achieved. In this case, post-hoc explanation techniques [45] would be helpful to interpret how the model works. One example of such techniques is knowledge distillation [120], which employed a student-teacher scheme to learn a simpler/interpretable model whose performance can approximate the performance of the complicated black-box model, from which the dark knowledge is “distilled out.”

Another related issue about model transparency is ownership. As Shah et al. has envisioned in their perspective [121], there is a worrying trend towards proprietary algorithms which are opaque, and the developers are “reluctant to transparently report” model details. This may raise the potential risk of harm when these models are applied in clinical practice [122]. In this case “regulatory and professional bodies should ensure the advanced algorithms meet accepted standards of clinical benefit, just as they do for clinical therapeutics and predictive biomarkers”, as Parikh et al. said in their discussion about predictive analytics in medicine [123].

4.3 Model Security
Conventionally we usually talk about the importance of protecting the security and privacy of health data, especially the data related to individual patients. With an increase in the number of AI models in health, we should also be aware of the potential security risk of those models. One example is adversarial attack, which refers to the process of constructing data that can confuse machine learning models and results in suboptimal or even incorrect decisions. For example, Sitawarin et al. [124] demonstrated that pollution on transportation signs can easily fool autonomous driving systems. Sun et al. [125] showed that slight modifications of lab values in a patient’s EHR can completely alter the mortality prediction made by what is otherwise a well-trained predictor. Finlayson et al. [126] provide a more detailed discussion on the potential concerns about the “incentives for more sophisticated adversarial attacks” in healthcare. From the authors of the present article’s perspective, it is important for (i) medical professionals to be aware of this potential risk; (ii) AI researchers to develop effective defense mechanisms in view of medical adversarial attacks; and (iii) policy makers to take into consideration the potential model security risk when they make new regulatory frameworks.

4.4 Federated Learning
Health data are widely distributed in and among health-related institutions, and each institution may be associated with a different set of stakeholders. In many cases, these
data are sensitive and cannot be aggregated. From a model-training perspective, it is desirable to have more and diverse data to inform model training.

Federated learning can assist with this challenge. According to Konečný et al. [127], “Federated Learning is a ML setting where the goal is to train a high-quality centralized model using training data distributed over a large number of clients”. These clients often have unreliable and relatively slow network connections. Developing federated health AI technologies is both important and highly demanding. Lee et al. [128] developed a privacy-preserving federated patient similarity learning approach and evaluated it on MIMIC III data [129]. They confirmed that in a federated setting, proper homomorphic encryption of patient information can indeed preserve the quality of patient similarity measures.

In addition to clinical data, there are more and more patient-generated data nowadays. For example, these data can be continuously generated from wearable devices or mobile phones. In this case, patients could be reluctant to share their data on some public cloud to train a predictive model for their future health status. With federated learning, the model will be stored in the cloud. Each user can download the current version of the model and improve it locally with his/her data. The model changes will be summarized as a focused update which will be sent back to the cloud with encrypted communication. Then the focused updates from different users will be averaged to improve the model. During the entire process, all data will remain on local devices and no individual update is stored in the cloud. Therefore, the model will be continuously updated in a secure way.

4.5 Data Bias

All AI models need training data samples. Typically, the size of the training sample obtained from patients is not large enough to capture all variations across patients and complexities of their health problems. Frequently, the model trained from patients at one hospital does not apply to patients in another hospital. We usually refer to this challenge as the bias carried in the data, and such data bias remains one of the major challenges to AI in health. As pointed out by Khullar [130], such bias can also worsen health disparities.

One way to reduce bias is to collect large and diverse patient data sets. Examples of such efforts include the OHDSI project [131] we introduced in Section 2.2, as well as the national clinical research network PCORnet created by the Patient-Centered Outcomes Research Institute (PCORI) [132] which currently includes 13 clinical data research Networks (CDRNs) collecting longitudinal patient data from a range of health systems across the United States. These efforts serve as a foundation for collecting large-scale, diverse data sets needed for robust, generalizable AI models. Researchers can also reduce bias during the model building process [133] using methods such as counterfactual Gaussian Process which is developed to perform both risk prediction and conduct “what-if” reasoning for individualized treatment planning.

5 Conclusion

The interest, applicability, and promise of AI in health is evidenced in recent literature. This review emphasizes some of the important aspects for future consideration and research. The work underway to overcome challenges in AI in health shows promise, and this progress will facilitate the expanding role that AI is likely to continue to play in health, from both an individual and population standpoint.

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