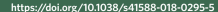
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A primer on deep learning in genomics

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Deep learning methods are a class of machine learning techniques capable of identifying highly complex patterns in large datasets. Here, we provide a perspective and primer on deep learning applications for genome analysis. We discuss successful applications in the fields of regulatory genomics, variant calling and pathogenicity scores. We include general guidance for how to effectively use deep learning methods as well as a practical guide to tools and resources. This primer is accompanied by an interactive online tutorial.

eep learning has made impressive recent advances in applications ranging from computer vision to natural-language processing. This primer discusses the main categories of deep learning methods and provides suggestions for how to effectively use deep learning in genomics. The primer is intended for bioinformaticians who are interested in applying deep learning approaches, and for genomicists and general biomedical researchers who seek a high-level understanding of this rapidly evolving field. Computer scientists may also use the primer as an introduction to the exciting applications of deep learning in genomics. However, we do not provide a survey of deep learning in the biomedical field, which has been broadly covered in recent reviews¹⁻⁵. This paper is accompanied by an interactive tutorial that we have created for interested readers to build a convolutional neural network to discover DNA-binding motifs (see URLs).

Deep learning as a class of machine learning methods

Machine learning techniques have been extensively used in genomics research^{3,6}. Machine learning tasks fall within two major categories: supervised and unsupervised. In supervised learning, the goal is predicting the label (classification) or response (regression) of each data point by using a provided set of labeled training examples. In unsupervised learning, such as clustering and principal component analysis, the goal is learning inherent patterns within the data themselves.

The ultimate goal in many machine learning tasks is to optimize model performance not on the available data (training performance) but instead on independent datasets (generalization performance). With this goal, data are randomly split into at least three subsets: training, validation and test sets. The training set is used for learning the model parameters (detailed discussion on parameter optimization in ref. 1), the validation set is used to select the best model, and the test set is kept aside to estimate the generalization performance (Fig. 1). Machine learning must reach an appropriate balance between model flexibility and the amount of training data. An overly simple model will underfit and fail to let the data 'speak'. An overly flexible model will overfit to spurious patterns in the training data and will not generalize.

Large neural networks, a main form of deep learning, are a class of machine learning algorithms that can make predictions and perform dimensionality reduction. The key difference between deep learning and standard machine learning methods used in genomics—e.g., support vector machine and logistic regression—is that deep learning models have a higher capacity and are much more flexible. Typical deep learning models have millions of trainable parameters. However, this flexibility is a double-edged sword. With appropriately curated training data, deep learning can automatically learn features and patterns with less expert handcrafting. It also requires greater care to train on and to interpret the underlying biology. Box 1 summarizes the main messages of this primer on how to effectively use deep learning in genomics.

Setting up deep learning

Deep learning is an umbrella term that refers to the recent advances in neural networks and the corresponding training platforms (e.g., TensorFlow and PyTorch). The starting point of a neural network is an artificial neuron, which takes as input a vector of real values and computes the weighted average of these values followed by a nonlinear transformation, which can be a simple threshold. The weights are the parameters of the model that are learned during training. The power of neural networks stems from individual neurons being highly modular and composable, despite their simplicity. The output of one neuron can be directly fed as input into other neurons. By composing neurons together, a neural network is created.

The input into a neural network is typically a matrix of real values. In genomics, the input might be a DNA sequence, in which the nucleotides A, C, T and G are encoded as [1,0,0,0], [0,1,0,0], [0,0,1,0] and [0,0,0,1]. Neurons that directly read in the data input are called the first, or input, layer. Layer two consists of neurons that read in the outputs of layer one, and so on for deeper layers, which are also referred to as hidden layers. The output of the neural network is the prediction of interest, e.g., whether the input DNA is an enhancer. Box 2 describes key terms and concepts in deep learning.

There are three common families of architectures for connecting neurons into a network: feed-forward, convolutional and recurrent. Feed-forward is the simplest architecture. Every neuron of layer i is connected only to neurons of layer i+1, and all the connection edges can have different weights. Feed-forward architecture is suitable for generic prediction problems when there are no special relations among the input data features.

In a convolutional neural network (CNN), a neuron is scanned across the input matrix, and at each position of the input, the CNN

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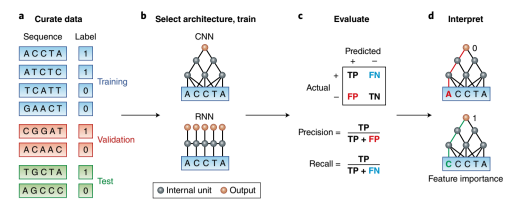


Fig. 1 Deep learning workflow in genomics. **a**, A dataset should be randomly split into training, validation and test sets. The positive and negative examples should be balanced for potential confounders (for example, sequence content and location) so that the predictor learns salient features rather than confounders. **b**, The appropriate architecture is selected and trained on the basis of domain knowledge. For example, CNNs capture translation invariance, and RNNs capture more flexible spatial interactions. **c**, True positive (TP), false positive (FP), false negative (FN) and true negative (TN) rates are evaluated. When there are more negative than positive examples, precision and recall are often considered. **d**, The learned model is interpreted by computing how changing each nucleotide in the input affects the prediction. The interactive tutorial illustrates the four steps of this workflow (see URLs).

computes the local weighted sum and produces an output value. This procedure is very similar to taking the position weight matrix of a motif and scanning it across the DNA sequence. CNNs are useful in settings in which some spatially invariant patterns in the input are expected.

Recurrent neural networks (RNN) are designed for sequential or time-series data⁷. At each point in the sequence, a neural network, which could be feed-forward or convolutional, is applied to generate an internal signal, which is also fed to the next step of the RNN. Hidden layers of the RNN can be viewed as memory states that retain information from the sequence previously observed and are updated at each time step.

In addition, there are neural network architectures used for unsupervised learning. The most common are autoencoders that perform nonlinear dimensionality reduction¹⁰, in contrast to principal component analysis, which is linear. In an autoencoder, the output is set to be the input, and the network is encouraged to find a low-dimensional space that compresses the original information and reconstructs the inputs.

Training a neural network starts with a labeled dataset of (X_i, Y_i) , where each X_i is the ith input, and Y_i is its output label. Each training point X_i is fed into the network, and the network's output is evaluated against the true label Y_i to produce a loss $L(Y_i, Y_i)$. Loss is the sum of the errors made for each example. The lower the loss, the better the model. Commonly used loss functions include squared error and cross-entropy. Squared error measures the difference between predicted and actual output values, which is especially relevant when the output is a continuous value. Cross-entropy measures the difference between two probability distributions over the same set of underlying events or classes, as is appropriate when the output is categorical?

To train the network, the derivative of $-L(Y'_i, Y_i)$ is computed with respect to the parameters of the network, which are the collection of neuron weights. By updating the weights in a small step in the direction of the derivative, the network's prediction loss can be decreased, and its accuracy can be increased. The derivative can be efficiently computed for each training point via the chain rule from calculus; this process is commonly called back-propagation. In parallel, the network's accuracy is also evaluated on the validation data, which are not used to update the weights. A good practice to avoid overfitting is to stop the training updates when the validation accuracy begins to decrease. More sophisticated techniques such as L_2 regularization on the weights and dropout are also effective in

controlling overfitting. L_2 regularization encourages simpler, and thus more generalizable, models by penalizing models with large weights⁷. Dropout is a training process that randomly ignores nodes to mitigate overfitting¹¹.

How to use deep learning effectively

The most important step in building an effective deep learning model is to first curate an appropriate training dataset and to select a suitable evaluation metric. The training set should be constructed to ensure that confounding biases that may artificially inflate performance are not introduced. For example, known pathogenic genetic variants may cluster in certain regions of the genome, i.e., exons or promoters, whereas known neutral variants may be more broadly distributed throughout the genome. A neural network naively applied to these unbalanced data may appear to perform well, but in reality it would probably learn to identify regions of the genome enriched in pathogenic variants without actually being able to distinguish neutral from pathogenic variants within these important genomic regions. Thus, it is important to design training datasets that are appropriately balanced for confounders that would detrimentally affect performance when applied to real-world-use cases¹².

Furthermore, genomics data are often highly imbalanced¹³. For example, there are many more variants that are not disease causing than are disease causing, or only a small fraction of the population may develop a particular disease to be predicted. Therefore it is often more meaningful to assess precision and recall, measures of classifier performance that account for the imbalance of classes in a dataset rather than simple accuracy¹⁴.

Successful application of deep learning also requires domain knowledge, as with all other machine learning methods. For example, in support-vector-machine classification and logistic regression, domain knowledge is used to construct features from data, and in Bayesian models, expertise is incorporated into the prior distributions. In deep learning, domain knowledge is built into the design of the network architecture. The performance of the network crucially depends on understanding the assumptions and limitations behind different architectures.

As an illustration, suppose we want to build a model to predict whether a DNA sequence is an active enhancer⁸. Biological knowledge indicates that regulatory elements may be effective even after small spatial translations, thus suggesting that CNN might model them effectively. Moreover, the regulatory motifs are known to

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Box 1 | Deep learning in genomics: key elements and guidance

- Large training datasets (typically thousands of examples), curated to remove confounders, are required.
- The main architectures—feed-forward, convolutional and recurrent—correspond to different assumptions about data.
- Most genomics data do not require very deep networks.
- Researchers must be wary of high accuracy due to data imbalance or bias that makes classification too easy.
- A good practice is to compare against simpler machine learning models on the same dataset.
- Deep learning can achieve high accuracy, but the interpretation of results is more challenging than for standard statistical models.

have a tendency to be relatively short (<20 nt) thus suggesting that convolution filters should also be small (<20 nt). Finally, if enhancers are being modeled, most of the activity is known to have a tendency to be clustered in regions from several hundred base pairs to two kilobases. Consequently, a reasonable design would be to limit inputs to the network to <2 kb. Any choice of the network architecture places an implicit prior over the model, and, as with any other machine learning, mismatch between the model's prior and the underlying biology can lead to poor performance.

In computer vision, researchers have observed that performance can improve with very deep networks (more than 100 layers). In most genomics applications, fewer than five layers are sufficient^{13,15}. Even relatively shallow networks can still have millions of parameters, and the most important factor that determines the success of a model is the availability of a large corpus of labeled training data. Most successful biological applications of deep learning have at least several thousand labeled examples¹. It is always good practice to train simpler machine learning models—linear regression, least absolute shrinkage and selection operator (LASSO), gradient boosting or random forest—in parallel on the same dataset to compare against the deep learning models. Simpler models have fewer parameters and can work better when fewer training datasets are available¹⁶.

Interpreting deep learning models

In many genomic applications, researchers are more interested in the biological mechanisms revealed by the predictive model rather than the prediction accuracy itself^{13,17}. For example, the main motivation for building accurate deep learning models to predict chromatin patterns is a hope to learn new gene-regulation grammar by interpreting the trained model. Although deep learning can achieve state-of-the-art accuracy, it is more challenging to interpret than the more standard statistical models.

The simplest method to interpret a neural network is analogous to in silico mutagenesis. Given a particular data point X, each feature of X (for example, mutating each nucleotide) can be systematically varied while the rest of the features are fixed, and how the network's output changes can be tracked. This approach is easy to implement but can be computationally expensive—the network recomputes the output for every mutation of X. A computationally tractable approximation to mutagenesis is to take the derivative of the network output with respect to each feature of X. This derivative can be computed in one back-propagation pass, and it conveys the sensitivity of the output to small perturbations in input features. Features with large positive or negative derivatives may be more influential to the outcome.

In strict terms, the derivative is a valid measure of influence for only infinitesimally small perturbations to the input, whereas in practice, researchers are interested in larger changes (for example,

Box 2 | Vocabulary and concepts

Artificial neuron

A simple mathematical function that takes a vector as input and outputs a transformed weighted sum of the vector. The weights are the neuron's parameters.

Deep learning

Neural networks with multiple layers of artificial neurons. The output of one layer is fed as input into the next layer to achieve greater flexibility.

Cross-validation

A common machine learning strategy wherein the dataset is split multiple times into training and validation sets. The average validation performance across the multiple splits is used to select the final model.

Back-propagation

A common method to train neural networks by updating its parameters (i.e., weights) by using the derivative of the network's performance with respect to the parameters.

Feed-forward neural network

The most flexible class of neural networks, wherein each neuron can have arbitrary weights.

Convolutional neural network

A class of neural networks in which groups of artificial neurons are scanned across the input to identify translation-invariant patterns.

Recurrent neural network

A class of neural networks with cycles that can process inputs of varying lengths.

Autoencoder

A class of neural networks that performs nonlinear dimensionality reduction.

Feature importance

A saliency score for each input feature (for example, each nucleotide) that measures the extent to which changes in that feature affect the prediction.

mutation of A to C). Several variations of the derivative-based interpretation methods have been developed to partially address this limitation—for example, integrated gradient¹⁸ and DeepLift¹⁹—and this goal is still currently an active area of research. Other interpretation methods, such as LIME²⁰ select a small number of features to explain why a prediction is made.

For a CNN, it is also possible to visualize each convolution filter as a heat map or position weight matrix–style logo image. These visualizations are useful to obtain a sense of what local features the network might be learning. A caveat is that multiple convolution filters might be learning partially redundant features, and how the local features interact is less clear, because such interaction depends on the higher layers of the network.

The interpretation methods discussed here should not be confused with causal models that attempt to pinpoint cause–effect relationships. Interpreting a prediction model can identify salient features and generate hypotheses, but inferring actual causality requires experimental perturbation.

Applications in genomics

A growing number of publications are presenting deep learning approaches and tools to study the genome (Fig. 2). Functional genomics is a leading application domain of deep learning²¹. Examples include predicting the sequence specificity of DNA- and RNA-binding proteins and of enhancer and cis-regulatory regions, methylation status, gene expression and control of splicing. These tools are based on data generated by DNase I sequencing, assay for

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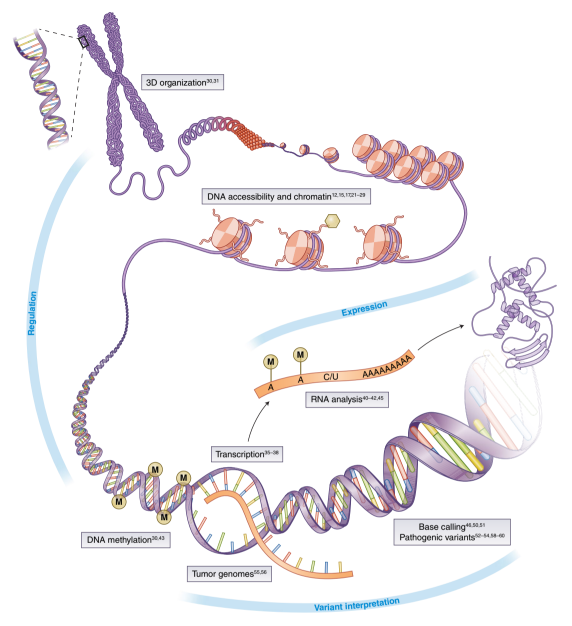


Fig. 2 | Applications of deep learning in genomics. The boxes highlight several application domains and references discussed in the text. Image adapted with permission from ref. 65, Springer Nature.

transposase-accessible chromatin using sequencing (ATAC-seq), chromatin immunoprecipitation (ChIP) sequencing, ChIP-on-chip, RNA immunoprecipitation sequencing, transcription-factor datasets and chromatin state^{15,17,22-24}. Similarly, the identification of transcription start sites as well as cis-regulatory/enhancer elements²⁵⁻²⁹ can be executed with the addition of features from the Encyclopedia of DNA Elements (ENCODE) project, as well as transcription-start-site sequencing and RNA-seq signals. The methylation state of DNA, which also influences the expression of genes, has been inferred from three-dimensional genome topology (on the basis of Hi-C) and DNA sequence patterns³⁰. Nucleotide sequence and DNase I assay signals predict Hi-C contacts at 1-kb resolution³¹.

Deep learning has been especially successful when applied to regulatory genomics, by using architectures directly adapted from modern computer vision and natural-language-processing applications³². Most approaches use CNN or RNN, which are well suited for the tasks of modeling regulatory elements, though genomics-specific modifications to the deep learning architecture

can be useful. For example, Shrikumar et al.³³ have addressed the phenomenon in which, in double-stranded DNA, the same pattern may appear identically on one strand and its reverse complement, owing to complementary base-pairing. Conventional deep learning models that do not explicitly model this property can produce different predictions on forward- and reverse-complement versions of the same DNA sequence.

Different tools are able to extract transcriptome patterns from large sets of gene expression data with the goal of estimating how much RNA is produced from a DNA template in a particular cell or condition^{34–37}. Deep learning can build predictive models of gene expression from genotype data³⁸ and can be used for studying the splicing-code model³⁹ as well as for the identification of long noncoding RNAs^{40–42}. Finally, deep learning has been used for the interpretation of regulatory control in single cells; for example, the detection of DNA methylation in single cells^{30,43}, and for the identification of subgroups of cells by improving the representation of single-cell RNA-seq data^{44,45}.

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Box 3 | Deep learning resources

All the resources, except for those listed in the 'specific for genomics' block, are relevant for deep learning in general. Because the field is developing rapidly, this information is likely to be considerably different in the future.

Resource type	Name	URL	Comment
Cloud platform	Amazon EC2	https://aws.amazon.com/ec2/	Most popular cloud platform
	Microsoft Azure	https://azure.microsoft.com/	Second-largest cloud platform
Plug-and-play cloud GPU services	FloydHub	https://www.floydhub.com/	All startups in the GPU service space; pay-by-the-hour model on top of basic monthly subscriptions
	PaperSpace	https://www.paperspace.com/	
	Valohai	https://valohai.com/	
	Google CloudML	https://cloud.google.com/ml-engine/	Can run your own models on Google's hardware, including tensor processor units
	Google Colaboratory	https://colab.research.google.com/	Notebook environment with free GPUs (during 12 h)
Design services for deep learning models	Fabrik	https://github.com/Cloud-CV/Fabrik/	Model export to Keras code; no training
	IBM Data Cloud	https://datascience.ibm.com/	Model export to Keras, PyTorch, TensorFlow or Caffe
	DeepCognition	http://deepcognition.ai/	Training and evaluation included
Prebuilt images with CUDA support	Docker Hub	https://hub.docker.com/r/nvidia/cuda/	Docker images from NVIDIA with CUDA/cuDNN GPU support
	Amazon Deep Learning AMIs	https://aws.amazon.com/machine-learning/amis/	Amazon Machine Images (AMIs) with GPU support
Software libraries (general)	Keras	https://keras.io/	More high-level than TensorFlow (below) but can be integrated with it in many ways
	TensorFlow	https://www.tensorflow.org/	Developed by Google; most popular deep learning framework
	PyTorch	http://pytorch.org/	Developed by Facebook
Software libraries	DragoNN	https://kundajelab.github.io/dragonn/	Tutorials included
(specific for genomics)	Kipoi	http://kipoi.org/	Model zoo for deep learning in genomics
Educational resources	fast.ai	http://www.fast.ai/	E.g., Deep Learning for Coders 1 and 2
	Coursera	https://www.coursera.org/specializations/deep-learning/	Deep-learning-specialization course package
	Textbook	http://neuralnetworksanddeeplearning.com/	Free online textbook with example code
	Fast.ai tips on configuring a deep learning environment	https://github.com/reshamas/fastai_deeplearn_part1/blob/master/README.md#platforms-for-using-fastai-gpurequired/	Instructions for configuring deep learning frameworks for a variety of platforms; from the fast.ai course but general; the details of these procedures change quickly
	Setting up TensorFlow with GPU on Google Cloud Engine	https://medium.com/google-cloud/jupyter-tensorflow-nvidia-gpu-docker-google-compute-engine-4a146f085f17/	Recipe for Docker-based setup of Google Cloud instance with TensorFlow, GPU support and Jupiter Notebooks

Predicting phenotypes from genetic data is also a major area of interest of deep learning. A first step in performing these types of predictions is to specify what genetic variants are present in an individual genome. This problem has been addressed by DeepVariant, which applies a CNN to make variant calls from short-read sequencing. The method treats DNA alignments as an image with a performance that appears to exceed that of standard variant callers⁴⁶. Other methods for variant calling using more traditional DNA representations have also been developed^{47–49}. Long-read-sequencing technologies also use deep learning for base calling^{50,51}.

Prioritizing variants on the basis of the likelihood that they are pathogenic is important. Several methods that predict the pathogenicity of coding variants have been proposed^{52–54}. These approaches are essentially aggregators that combine prior non–deep learning

predictors as well as features known to be useful for the prediction of variant pathogenicity. A more challenging problem, the prediction of functional consequences of noncoding variants, has seen some success via DeepSEA¹³, a CNN trained to predict functional genomics features that can also be adapted to predict variant effects. In cancer genomics, deep learning can extract the high-level features between combinatorial somatic mutations and cancer types⁵⁵ and learn prognostic information from multicancer datasets⁵⁶. Preliminary phenotype prediction in agricultural applications appears promising⁵⁷. However, a demonstration of the prediction of common complex human disease phenotypes is only now emerging: Zhou et al. have recently reported the extension of DeepSEA to the study of regulatory variants in autism spectrum disorder⁵⁸. The same team has published ExPecto, the ab initio prediction of gene

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expression levels and variant effects from sequences from more than 200 tissues and cell types⁵⁹. Also recently, Sundaram et al. have trained a DNN by using hundreds of common variants from population sequencing of nonhuman primate species to identify pathogenic variants in rare human diseases⁶⁰.

Resources

Implementation of deep learning approaches requires familiarity with new tools and resources. These include access to computing power and general and genomics-specific software libraries (Box 3).

Cloud platforms. Recent advances in deep learning have, to a considerable degree, been driven by developments in graphical processing unit (GPU) technology. GPUs can significantly increase training speed because the way in which neural networks are trained lends itself well to their architecture, thus allowing for fast vector and matrix multiplications, owing to the large number of processing units and high memory bandwidth in GPUs. Some academic highperformance computing clusters offer access to GPUs, but because progress in GPU technology is rapid, those processors are at risk of becoming outdated. The major cloud-computing platforms, such as Amazon EC2, Google Cloud Engine, Microsoft Azure and IBM Cloud, can be more flexible in offering on-demand GPU access. However, even these cloud platforms require some degree of configuration from the user, such as installing and compiling an appropriate version of CUDA (a popular parallel computing platform and programming toolkit) for general GPU programming, upon which many deep learning frameworks depend for GPU acceleration.

For users wishing to avoid semimanual setup procedures, there are specialized platforms offering 'plug-and-play' GPU access, for instance FloydHub, Valohai, Paperspace and Google CloudML. Typically, these platforms provide configuration-free access to GPUs on Amazon or Google cloud infrastructures for a modest markup compared with running 'raw' cloud instances directly. Perhaps the simplest alternative at the time of writing of this paper is Google Colaboratory, a Python notebook environment that provides free use of a K80 GPU during 12 consecutive hours. The tutorial accompanying this paper (see URLs) is built on this platform. In some cases, using lightweight virtual machines such as Docker or Singularity containers is convenient for packaging software and dependencies so that they can be directly run on a cloud instance with minimal setup. All these solutions, however, still require users to write the code for the models.

Some emerging platforms are offering the possibility to design deep learning models without coding. For example, IBM Data Cloud has recently introduced a neural network designer in which users can build a model from building blocks and export the finished model to runnable code. The open-source Fabrik framework offers a similar capability. DeepCognition offers a platform in which users can design, train and evaluate models in the same framework.

Software libraries. Many software libraries are commonly used for deep learning. Whereas earlier models were often written in frameworks developed at universities, such as Theano, Torch and Caffe, recent years have seen a strong showing of open-source Python libraries developed in corporate laboratories, for example TensorFlow (Google) and PyTorch (Facebook). Some frameworks were initially developed independently but subsequently received explicit backing from a company, for example, Keras (Google) and MXNet (Amazon). Keras has gained its popularity from being designed to be at a higher level of abstraction than most other frameworks; consequently, it allows for model construction in larger conceptual blocks such as network layers without requiring users to keep track of all the details.

Because of the relatively recent entry of deep learning into genomics, there are few genomics-specific software libraries (Box 3).

One example is DragoNN, a toolkit to teach and learn about deep learning for genomics, which includes tutorials in Python notebook format, a command-line interface, tools for running models on cloud CPU or GPU, and a model-interpretation module. Kipoi is a repository of ready-to-use trained models in genomics. It currently contains 2,031 different models ('model zoo'), covering canonical predictive tasks in transcriptional and post-transcriptional gene regulation⁶¹. Kipoi uses Conda virtual environments to install the correct dependencies for each model.

Conclusions and perspectives

Although deep learning has demonstrated impressive potential in genomics, a number of outstanding issues remain⁶². First is the challenge of how to design deep learning systems that best augment and complement human experience in making medical decisions (for example, genome interpretation). Second is the challenge of how to avoid biases in training sets and how to interpret predictions. Interpretation and robustness are two important directions of method development⁶³. Finally, there is a need for iterative experimentation, in which deep learning predictions can be validated by functional laboratory tests or by formal clinical assessment.

The most successful applications of deep learning in genomics to date has been in supervised learning, i.e., making predictions⁸. It is important to not confuse high prediction accuracy with the ultimate objective of extracting biomedical insights from data and making robust assessment in diverse settings that might be different from the training data. Beyond making predictions, deep learning can potentially become a powerful tool for synthetic biology by learning to automatically generate new DNA sequences and new proteins with desirable properties⁶⁴. Such generative models are an exciting new frontier of research.

URLs. Interactive tutorial to build a convolutional neural network to discover DNA-binding motifs, https://colab.research.google.com/drive/17E4h5aAOioh5DiTo7MZg4hpL6Z 0FyWr.

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Author contributions

All authors conceived and designed the project. J.Z., M.H., P.M., A. Torkamani and A. Telenti wrote the manuscript. J.Z. and A.A. wrote the online tutorial.

Competing interests

M.H. is an employee of Peltarion.

Additional information

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