Surrogate-Assisted Evolutionary Deep Learning Using an End-to-End Random Forest-based Performance Predictor

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Abstract—Convolutional neural networks (CNNs) have shown remarkable performance in various real-world applications. Unfortunately, the promising performance of CNNs can be achieved only when their architectures are optimally constructed. The architectures of state-of-the-art CNNs are typically hand-crafted with extensive expertise in both CNNs and the investigated data, which consequently hampers the widespread adoption of CNNs for less experienced users. Evolutionary deep learning (EDL) is able to automatically design the best CNN architectures without much expertise. However, existing EDL algorithms generally evaluate the fitness of a new architecture by training from scratch, resulting in the prohibitive computational cost even operated on high-performance computers. In this paper, an endto-end offline performance predictor based on the random forest is proposed to accelerate the fitness evaluation in EDL. The proposed performance predictor shows promising performance in term of the classification accuracy and the consumed computational resources when compared with 18 state-of-the-art peer competitors by integrating it into an existing EDL algorithm as a case study. The proposed performance predictor is also compared with the other two representatives of existing performance predictors. The experimental results show the proposed performance predictor not only significantly speeds up the fitness evaluations, but also achieves the best prediction among the peer performance predictors.

Index Terms—evolutionary deep learning, performance predictor, surrogate model, random forest, convolutional neural network.

I. INTRODUCTION

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C ONVOLUTIONAL Neural Networks (CNNs) [1], [2], as the most dominant deep learning approaches [3], have been demonstrating their promising performance in addressing various real-world applications, such as image classification [4], speech understanding [5], and natural language processing [6], to name a few. It is well known that the performance of CNNs highly relies upon their architectures, and a new architecture must be redesigned if the addressed data has been changed. Unfortunately, designing architecture with the best performance for the investigated data requires extensive expertise in both the CNNs and the data domain [7], which is not necessarily held by the interested users. Designing the best CNN architecture for the given data can be seen as an optimization problem, which can be mathematically formulated by (1):

$$\underset{A_{\lambda}}{\operatorname{arg\,max}} \ \mathcal{L}(A_{\lambda}, \mathcal{D}_{train}, \mathcal{D}_{test}) \ s.t. \ \lambda \in \Lambda$$
(1)

where λ refers to the parameters related to the architectures of CNNs, such as the number of convolutional layers and the configurations of pooling layers; Λ refers to the parameter space, A_{λ} denotes the CNN algorithm A adopting the architecture parameters λ , and \mathcal{L} means the performance measure of A_{λ} on the test data D_{test} after it has been trained on the training data \mathcal{D}_{train} . Generally, λ is with discrete values, e.g., the configurations of both convolutional and pooling layers must be integers, which results in the architecture optimization not being well-addressed by the exact optimization algorithms. To address this challenge, researchers have developed algorithms (e.g., [8]-[14]) to design the best CNN architecture by effectively solving the problems represented in (1). Among these algorithms, the evolutionary algorithm-based ones, which are called the evolutionary deep learning algorithms (EDLs), are much preferred because EDLs generally consume less computational resource than other CNN architecture design algorithms [11]-[14], while still can achieve the promising results. For example, on the CIFAR10 dataset [15], the evolutionary algorithm-based Automatic Evolving CNN (AE-CNN) algorithm finds the best CNN architectures by consuming 22 days using three Graphics Processing Units (GPUs), and achieves the classification error rate of 4.7%. However, the reinforcement learning-based neural architecture search method [9] consumes 28 days using 800 GPUs, while achieves the classification error rate of 6.01%which is a worse result than AE-CNN.

Evolutionary algorithm is a type of population-based heuristic computational paradigms [16]–[18], and has been widely used in addressing various complex optimization problems [19]-[21], mainly because of its gradient-free and insensitiveness to local minimum [22]-[24]. In EDLs, each individual is transformed to a CNN with the corresponding architecture through the mapping from genotype to phenotype, and then the weights of the CNN are initialized and iteratively trained based on \mathcal{D}_{train} commonly with stochastic gradient descent (SGD) [25], [26] given a number of epochs. During each epoch, the CNN also needs to be trained with multiple iterations in which the number is determined by the adopted batch size and the size of \mathcal{D}_{train} . Such a single fitness evaluation typically takes several hours to days. As a result, the potential promising architecture may not be found through EDLs within a limited computational budget due to the time-consuming fitness evaluation. This will become even more severe in the case of a large population and with a large number of generations, which is a more widely used configuration of evolutionary algorithms targeting at finding the better solutions [16], [18], [27]. Therefore, speeding up the fitness evaluation of CNN is essential in the design of EDL.

A common way to accelerate the fitness evaluation of CNN is to use a performance predictor [28]. In 2014, Swersky et al. proposed the Freeze-thaw Bayesian optimization algorithm [29] by using a Gaussian process regression [16], which built the model based on the training performance in the first t epochs to predict performance at the T (T > t) epoch. Followed by the same routine, the learning curve extrapolation methods in [30], [31] employed a set of functions from exponential families and the Bayesian neural network, respectively, to build the model based on the training performance in the first t epochs. Unfortunately, these mentioned performance predictors are typically based on expensive Markov chain Monte Carlo sampling procedures and manually designed curve functions, which are with high computational complexity. In 2017, Deng et al. proposed the Peephole algorithm [32] that predicted the performance of a CNN based on its architecture information, where a large number of different CNNs were trained, and the performance of all sampled networks in each epoch was collected as the training data, and then a long-short term memory neural network [33] is used to train the model. In addition, the train-less accuracy predictor (TAP) [34] and the accelerating neural network architecture search (ANNAS) [35] algorithms also employed similar methods to predict the performance of the given CNNs.

In fact, the optimization problem in EDLs is a type of computationally expensive problems, which is usually solved by surrogate-assisted evolutionary algorithms (SAEAs) [36], using cheap approximated regression and classification models (like Gaussian process model [37], radial basis function network [38], etc.) to replace the expensive fitness evaluation [39]. SAEAs have shown effectiveness and efficiency in various real-world optimization applications [40]. Generally, the models are trained from a small number of expensive fitness evaluations (i.e., the samples with evaluated objective values), and then the trained models are used as the fitness predictors in the evolutionary search to accelerate the optimization process [41].

According to [42], SAEAs can be divided into online and offline algorithms, which depend on whether expensive fitness evaluations are used to enrich the training data during the optimization process. Although new training data in online algorithms can significantly improve the quality of the model and optimization performance, offline algorithms are more practical than online algorithms due to the hardness and high cost in obtaining new data [40]. Without sampling new training data, offline algorithms can be very fast, and its data collection and optimization search are completely separated. So far, the base algorithms commonly applied in offline SAEAs are data pre-processing [43], data mining [42], and ensemble learning [44], since they can effectively and robustly address the problems having limited training data.

Our goal in this paper is to present an effective and efficient end-to-end performance predictor (in short named E2EPP) based on random forest [45]. The adoption of random forest lies in its effectiveness to a limited number of training samples, applicability to discrete variables, and robustness to its parameter settings [46], [47]. The proposed E2EPP is able to directly know the performance of a CNN once it looks at the new CNN architectures, which is naturally able to speed up the evolutionary process of EDLs. The contributions of this paper are summarized below:

- The training data of the random forest are a set of data pairs, and each pair is composed of the CNN architecture and its performance. However, the architecture is modelled by describing languages, e.g., a paragraph of words, which cannot be directly used as the input to the random forest. Therefore, we propose an effective encoding method that is capable of extracting the features of CNN architectures as the numerical values, which are further used as the samples for training the random forest. As a case study, AE-CNN [48], as the representative of the state-of-the-art EDLs, is adopted to investigate the details of encoding.
- 2) The trained random forest is based on a set of samples of CNN architectures (and their performance) which do not necessarily cover all the architecture space. If the performance prediction is adopted from the tree giving the best performance, the obtained result may be biased. To this end, we employ a selective ensemble strategy to choose diverse trees for combination in the local area of every generation.
- 3) Considering the two goals in term of using the performance predictor: the efficiency in speeding up the evolutionary process and the effectiveness in finding the best CNN architecture, we perform extensive experiments to verify these aspects.

The remainder of this paper is organized as follows. The background and related work are introduced in Section II. This is followed by the algorithm details of the proposed random forest-based performance predictor in Section III. In order to validate the effectiveness and the efficiency of the proposed performance predictor, the experiment designs and the experimental results are presented in Sections IV and V, respectively. Finally, the conclusion and future works are described in Section VI.

II. LITERATURE REVIEW

In this section, the EDL, AE-CNN and random forest are first introduced as the base algorithms in Subsection II-A, which is helpful to know the details of the proposed performance predictor. Then, the related work to performance predictors is presented in Subsection II-B, for better justifying the effectiveness and efficiency of the proposed work.

A. Background

1) EDL: A general framework of EDLs [11]–[14] is composed of the steps shown below:

- Step 1: Randomly initialize a population with the predefined size based on the corresponding genotype-to-phenotype mapping strategy.
- Step 2: Map each genotyped individual to the corresponding CNN, and train each CNN to obtain the classification accuracy on the validation dataset as its fitness value.
- Step 3: Use tournament selection to select the parent individuals based on the fitness, and then generate the new offspring achieving the same predefined size through the crossover and mutation operators.
- Step 4: Perform the environmental selection on the combined population to select a new population surviving into the next generation.
- Step 5: Go to step 3 if the termination condition is not satisfied. Otherwise, choose the individual with the best fitness and terminate the evolutionary process.

Indeed, the whole process of an EDL follows the procedure of an evolutionary algorithm, i.e., population initialization (Step 1), fitness evaluation (Step 2), offspring generation (Step 3) and environmental selection (Step 4). Generally, an evolutionary algorithm involves four parameters: the maximal generation number for the whole evolutionary process, the population size for the population initialization and the environmental selection, and the crossover rate as well as the mutation rate for the offspring generation. One major factor of the promising performance of evolutionary algorithms is the environmental selection to keep individuals with better fitness in the new population than the previous population. However, as described in Step 2, the fitness of each individual in EDL is evaluated through the training of the corresponding CNN. For a better understanding of the fitness evaluation, the framework of training a CNN is provided in Algorithm 1.

Generally, the training of a CNN is achieved by using SGD on a given training dataset through hundreds of epochs (Steps 5-11). The objective of the training is to maximize the classification accuracy by adjusting millions of the weights (Step 8) in CNN. For a moderate CNN on a commonly used benchmark dataset, e.g., CIFAR10 and CIFAR100 [15], the training process would take from several hours to days performing even on the high-performance GPUs. Since evolutionary algorithms belong to a class of population-based algorithms, their required numbers of evaluations are larger than that of SGD, which make the computational cost of EDL even higher. For example, on the CIFAR10 dataset, the large-scale evolutionary algorithm [12] consumed 11 days on 250 GPUs, and the hierarchical evolutionary algorithm [13] spent 1.5 days

Algorithm 1: Fitness Evaluation of A CNN in EDL

Input: The individual p, training dataset \mathcal{D}_{train} , validation dataset \mathcal{D}_{valid} , training epochs T, batch size b, learning rate γ , objective function \mathcal{L} .

Output: The fitness.

- 1 $net \leftarrow$ Decode p into the corresponding CNN architecture;
- 2 Add the classification layer to the end of *net*;

3 Randomly initialize the weights w in net;

4 $t \leftarrow 0;$

5 while t < T do

for each batch data in
$$\mathcal{D}_{train}$$
 do

$$\nabla w \leftarrow \text{Compute the gradient by } \partial \mathcal{L} / \partial w;$$

 $w \leftarrow w - \gamma \nabla w;$

9 er

6

7

8

10

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t \leftarrow t+1;
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11 **end**

12 Calculate the classification accuracy of net on \mathcal{D}_{valid} ;

13 **Return** the fitness.

on 200 GPUs. Such a large number of GPUs may not be available to many researchers. To address such a challenge, it is of great importance to reduce the computational cost for EDLs by developing performance predictors, which could replace most of those expensive fitness evaluations, to speed up EDLs relying on limited computational resources.

2) AE-CNN: As the performance predictor is part of EDL in this research, an EDL method should be provided before the performance predictor is detailed. In this work, the AE-CNN algorithm developed by the authors is selected as the representative EDL, which is mainly based on the reasons that: 1) it shows promising performance among existing EDLs [48], and 2) the source code of AE-CNN is available to the public. Noting that the proposed performance predictor is applicable to any existing EDLs [8]–[14].

The AE-CNN algorithm [48] is an automatic EDL algorithm based on the building blocks of the state-of-the-art ResNet [49] and DenseNet [50], which has shown the promising performance on the CIFAR10 and CIFAR100 datasets [15] compared to state-of-the-art CNNs manually designed and other semi-automatic and automatic EDL algorithms [48]. The AE-CNN algorithm used in this work is to prepare samples for training the random forest, and then the trained random forest is used as the performance predictor in AE-CNN to verify the effectiveness and efficiency. In this subsection, we mainly introduce the genotype-to-phenotype mapping of AE-CNN for the completeness of the presentation, which enables the ability of evolutionary algorithms to model CNN architectures. These CNN architectures are the input to the random forest (the details of how to feed the architecture information to the random forest will be discussed in Section III).

The CNN architectures generated by AE-CNN are composed of the DenseNet Blocks (DBs), ResNet Blocks (RBs) and Pooling Blocks (PBs). Each DB or RB is composed of multiple DenseNet Units (DUs) and ResNet Units (RUs), respectively, while a PB consists of only one pooling layer. Each DU, RU, or PB differs in terms of the parameter settings. The parameters of a DU are the sizes of input and output (denoted by in and out, respectively), while those of an RU are the same as those of a DU, in addition to an increasing factor (denoted by k). The parameter of a PB is only the pooling type (i.e., the maximal or *mean* pooling type) because its other parameters are all set to fixed values. Because a DB/RB is composed of multiple DUs/RUs, the parameters of a DB/RB are the corresponding parameters of a DU/RU and the amount (denoted by *amount*) of DUs/RUs in a DB/RB. In addition, as the DBs, RBs and PBs compose a CNN with an order, an extra parameter (denoted by *id*) is also used to represent the corresponding position in the CNN. Obviously, a CNN generated by AE-CNN is composed of sequential blocks which may be the DUs, RUs or PBs. When the block is an RB, the parameters are the *id*, *amount*, *in* and out; when the block is a DB, the parameters are *id*, *amount*, k, in and out; when the block is a PB, the parameter is the pooling type denoted by type. Noting that the *in* of the current RB/DB should be equal to the out of its previous RB/DB for the reason of constructing a valid CNN.

3) Random Forest: Compared with other learning algorithms, such as neural networks, the random forest has the advantages of directly accepting the discrete data as input, almost having no extra parameters to tune and not relying on a large amount of training data [50]. These advantages are the exact reason for employing the random forest in this work. Random forest [45] is an ensemble learning method by operating a set of decision trees for classification and regression tasks [46]. During the training of random forest, each decision tree randomly selects a part of the whole feature set and then learns the mapping from the selected features to the corresponding target. During the predicting process, each decision tree selects the same features as it selects in the training phase, and then output the corresponding prediction. The random forest uses the average output of the selected decision trees as its final output. Randomly selecting a part of the whole feature set is also the famous random subspace method [47], which has been shown to be simple yet effective.

B. Related Work

The existing performance predictors can be classified into two different categories: performance predictors based on the learning curve and end-to-end performance predictors, both of which are based on the training-predicting computational paradigm. The representatives of these two types are the Freezethaw Bayesian Optimization (FBO) algorithm [29] and the Peephole algorithm [32].

The FBO algorithm uses a regression model based on Bayesian Optimization to predict the performance of a CNN. The regression model is built based on the data sampled from the learning curve regarding the first t epochs, i.e., a series of data pairs in different training epochs and the corresponding performance at these epochs. After the regression model that has been optimized by the Bayesian Optimization method, it is used to predict the performance of the same CNN at the T-th epoch where T > t. The Peephole algorithm uses a number of CNN architectures and their corresponding performances that have been achieved by training the CNNs with T epochs, as the training samples to train a long-short time memory neural network [33]. The trained neural network directly predicts the performance of a new CNN based its architecture, which is called the end-to-end mechanism because the end in the input is the raw data while the end in the output is the classification accuracy. Because the architecture cannot be directly used as the input data of the neural network, the Peephole algorithm employs the word vector technique to map the CNN architecture to a numerical value.

The major advantage of the FBO algorithm remains in the trained-CNN-free nature, i.e., it does not need any trained CNNs in advance. Because training a CNN is time-consuming, varying from several days to weeks, the FBO algorithm is efficient. However, it will not be effective when the learning curve is not smooth because the curve fitting works under the assumption that the curve is smooth. In recent deep learning applications, the learning curve is usually not smooth because a schedule of learning rates is usually used. Once the learning rate is changed, the learning curve will have a non-smooth segment. Another limitation of the FBO algorithm is the nonend-to-end nature (i.e., in predicting the performance of each CNN, a part of the training data regarding this CNN must be collected for training the predictor), which requires much more labour work when it is used. Owing to the end-to-end nature, the Peephole algorithm is more convenient for use. However, the major drawback of Peephole remains in the requirement of a large number of training samples, which results in added computational complexity of collecting train samples exceeding the EDL without using performance predictors. For example, Peephole used over 8,000 fully trained CNN architectures as the training data. However, EDLs generally achieve promising performance by evaluating only hundreds of individuals. If we have enough computational resources to evaluate the 8,000 CNN architectures, we will not need to develop the performance predictor. Such a limitation is largely caused by its adopted regression mode, i.e., the neural network-based algorithm, which typically highly relies on a large amount of labelled training data.

The proposed performance predictor has the advantages of the end-to-end manner and relying upon only limited training data, which address both limitations of the existing performance predictors discussed above.

III. PROPOSED ALGORITHM

As mentioned, the main challenge of EDL is the high computational cost of evaluating a single CNN architecture. Inspired by offline SAEAs, we use random forest [46] as the fitness predictor to replace the expensive fitness evaluation. For a better understanding, Fig. 1 shows the framework of the proposed performance predictor as well as the associated EDL. The random forest-based performance predictor is learned from a number of different CNN architectures that have been trained for a classification task with their accuracy. Compared with the CNN training process, the random forest construction and prediction is computationally cheaper and can be repeatedly

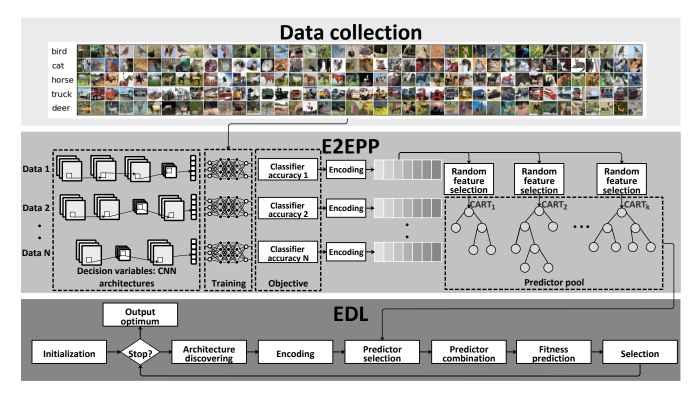


Fig. 1. Main frame of the proposed algorithm.

used throughout the evolutionary optimization. Thus, the high computational cost in EDL can be relieved.

As shown in Fig. 1 that is composed of three blocks, i.e., the data collection, E2EPP and EDL. The proposed E2EPP performance predictor is part of EDL. Firstly, a set of training data is collected for training the random forest-based predictor, where the collection is achieved by performing the corresponding EDL without using E2EPP. Each data sample is composed of the CNN architecture and the corresponding classification accuracy that is obtained by training the CNN from scratch. Secondly, those architectures are encoded into discrete code (shown in Subsection III-A) for building the random forest-based predictor pool with a large number (say K) regression trees (denoted as CARTs) [51] (shown in Subsection III-B). During each generation of the EDL, the newly generated CNN architecture is encoded as the input to the random forest, and then its performance is predicted by using the adaptive combination of CARTs from the predictor pool (shown in Subsection III-C). When the EDL terminates, the CNN architecture that has the best prediction performance is output. Noting that there is no further CNN training during the optimization process.

Since the CNN architecture is presented as discrete variables, random forest [52], which is suited for discrete regression tasks and also relies on only limited labelled data [53], is adopted as the fitness predictor in the proposed algorithm. Also, as a selective ensemble of CARTs considering both global and local landscape, the random forest-based predictor is adaptively updated over the generations of the evolutionary search, where the population is distributed in different local regions and a fixed random forest cannot guarantee accuracy in those local regions. In the following subsections, we will first introduce the details of encoding of the CNN architectures to the proper format for the random forest, training of the random forest, and fitness prediction using the trained random forest in Subsections III-A, III-B and III-C, respectively. Then, the strength and the weakness of the proposed performance predictor are summarized in Subsection III-D.

A. Encoding

The encoding operates on the trained CNNs whose architectures are randomly initialized. To collect this data, we use AE-CNN to randomly generate a set of valid CNN architectures. A valid CNN architecture means that this architecture can be trained with a predefined training routine without any exceptions such as the out-of-memory errors causing a zero classification accuracy. Based on the description shown in Subsection II-A2, the collected training data are summarized as below:

- 1) RBs and DBs: Each generated CNN architecture is composed of four RBs and four DBs at most, the number of output channels of each block varies between [32, 512] that is set based on the conventions of state-of-the-art CNNs [50], [54].
- PBs: Each generated CNN architecture contains four pooling layers at most. There are two types of PBs: MAX and MEAN.

Generally, we encode a CNN architecture into a chromosome with $3N_b + 2N_p$ discrete variables, when the maximal number of RBs and DBs is N_b and the maximal number of PBs is N_p . For the first $3N_b$ variables, each RB or DB is encoded

Input: The CNN architecture A , the maximal number			
B_n of DBs and RBs, the maximal number N_p			
of PBs.			
Output: The encoded architecture information.			
1 $b_list \leftarrow \emptyset;$			
2 $p_list \leftarrow \emptyset;$			
$l \leftarrow Calculate$ the number of blocks in \mathcal{A} ;			
4 for $for i \leftarrow 1$ to l do			
5 $block \leftarrow \text{Get the } i\text{-th block of } \mathcal{A};$			
6 if block is a RB then			
7 Put 1 into $b_list;$			
8 Put the values of the <i>out</i> and <i>amount</i> of <i>block</i>			
into b_{list} ;			
9 else if block is a DB then			
10 Put the values of the k , out and amount of			
$block$ into b_{list} ;			
11 else			
12 if block is a maximal pooling layer then			
13 Put 1 to p_list ;			
14 else			
15 Put 0 to p_list ;			
16 end			
17 Put i to p_list ;			
18 end			
19 end			
20 Put zero to b_list until $ b_list = 3N_b$;			
21 Put zero to p_list until $ p_list = 2N_p$;			
22 Return $b_list \cup p_list$.			

into a triplet as [type, out, amount], where the block type for RBs is set to 1, and that for DBs are set to 12, 20 and 40 when k is equal to 12, 20 and 40, respectively. Noting that the parameter of in for each RB/DB is not encoded because it can be calculated by the out of its previous RB/DB, and the smaller number of decision variables could result in a better performance for regression model when the training data is limited [46]. For the following $2N_p$ variables, each pooling layer is encoded into a pair as [pooling type, layer position], the maximal and mean pooling types are presented by 1 and 0, respectively. If a CNN architecture has b RBs and DBs, and p PBs, then its 3b + 1-th to $3N_b$ -th variables are set to zeros, and its $3N_b + 2p + 1$ -th to $3N_b + 2N_p$ -th variables are set to zeros as well. Therefore, the performance predictor by using random forest is based on the input data with $3N_b + 2N_p$ discrete decision variables, and the output is a continuous value within the range of [0, 1]. Algorithm 2 shows the details of encoding a CNN architecture into the data that can be directly used by the random forest, and $|\cdot|$ is a countable operator.

B. Training of the Random Forest

A large number of CARTs are generated in the predictor pool. Each CART is trained by the whole training data with a random subset of features (i.e. discrete variables), where each discrete variable is assigned a probability of 0.5 in order to maximize the diversity of the predictor pool [55]. Each node of a CART presents a rectangle region in the decision space. The mean squared error of the output of those samples in that region (node) determines whether this node needs splitting or not (i.e., whether the mean squared error decrease is smaller than a set threshold T_s [51], and if so, this node is a leaf node). When K CARTs are obtained, the predictor pool is ready for the optimizer. The details of training the CARTs are shown in Algorithm 3.

Al	gorithm 3: Performance Predictor Training
Ι	nput: The K CARTs, the encoded training data
	\mathcal{D}_{train} , the feature number m .
0	Dutput: The K trained CARTs and their selected
	feature ids.
1 I	$\leftarrow \emptyset;$
2 f	or $i \leftarrow 1$ to K do
3	$CART \leftarrow$ Select the <i>i</i> -th CART from CARTs;
4	$v \in \mathbb{R}^m \leftarrow$ Randomly generated a vector from
	[0,1];
5	$I_i \leftarrow$ Collect the position of the elements whose
	values are greater than 0.5 in v ;
6	Train $CART$ on the features whose ids are in I_i ;
7	$I \leftarrow I \cup I_i;$
8 e	nd
0	Dutput: The K trained CARTs and the corresponding
	selected feature ids I.

C. Performance Prediction

Since the computational cost of training a single CNN is very high, it is impractical to obtain any new training data for the predictor pool during the optimization process. Thus, those predictors cannot be updated or validated, which is the main challenge of offline SAEAs [40]. To deal with the lack of training data, a large number of surrogate models are employed as ensemble members in a recent offline SAEA [44]. The results indicate that a selective ensemble surrogate can effectively improve the robustness of the obtained solution when the new training data is unavailable. Also, those ensemble members are adaptively combined in each generation to provide local information on the current population. Motivated by the combination strategy that was originally proposed in our previous work [44] for addressing decision variables having continuous values, we employ it to update the random forestbased predictor in every generation in the proposed algorithm, i.e., we choose Q CARTs from the predictor pool and then use their average prediction as the fitness value. Although the decision variables of the proposed algorithm are discrete, it is experimentally found that the combination strategy still works well.

In each generation, all of the K trained CARTs re-estimated the performance on the CNN \mathcal{A}^b that has the best-predicted fitness value; and then Q out of the K CARTs are uniformly selected from the K ordered CARTs based on their prediction values on \mathcal{A}^b . The Q CARTs are combined as the ensemble performance predictor to evaluate both parent and offspring population. Such selection is based on the performance diversity of CARTs around the current best CNN architecture \mathcal{A}^b . After that, the generated CNN architectures \mathcal{A} are evaluated by using the ensemble predictor of Q CARTs. Thus, the adaptive predictor can balance the global tendency and local information in the fitness landscape, where the combination of K CARTs predicts the global average landscape and that of Q diverse CARTs in a small area refines the local landscape. The details of the prediction process in a generation are shown in Algorithm 4.

D. Strength and Weakness of E2EPP

As have been introduced in Subsection II-B, the limitations of the existing performance predictors are the non-end-to-end nature, the strict assumption on the smoothness of the learning curve, and the availability of the large training samples. The proposed method has been carefully designed to address these limitations.

The proposed algorithm is end-to-end and does not rely on the learning curve no matter whether it is smooth or not. Firstly, the end-to-end nature is more convenient in practice because we do not need to prepare the training data in predicting the performance of each CNN. Secondly, because the proposed algorithm does not need to fit the learning curve, the predicted performance is better than the existing approaches based on the learning curve. This is theoretically evidenced by the universal approximation theorem [56] that the learning curvebased approaches can achieve promising performance only when the learning curve is smooth. However, in practice, the learning curve is not always smooth.

On the other hand, the proposed algorithm does not require a large number of training samples. Most existing approaches employing deep learning techniques succeed subject to the availability of a large amount of training data. In the CNN performance prediction, such training data is collected by training a lot of CNNs from scratch. However, the training is time-consuming even if performing on GPUs, while the main goal of designing performance predictors is to save the time of training CNNs. If we put enough time to collect a sufficient amount of training data, the performance predictor design will lose its original purpose. In the proposed method, we use the random forest as the base operator to learn the mapping between the CNN architecture and its performance. The reason is that random forest can achieve a promising performance on limited amount of training data, which has been theoretically proven and practically investigated [45], [46].

Unfortunately, the weakness of the proposed algorithm is the unknown of the minimal number of training samples for achieving promising performance, while the minimal number is case by case for different tasks. In practice, we need to use an incremental strategy to sample training samples until the desired performance is achieved.

IV. EXPERIMENT DESIGN

In order to verify the effectiveness and efficiency of the proposed performance predictor, a series of experiments are carefully designed and performed. Although the proposed performance predictor aims at speeding up the fitness evaluation of EDL, the ultimate goal of the performance predictor is

Algorithm 4: Performance Predicting

Input:	The K trained CARTs, the selected features ids
	<i>I</i> of each CART, the current best CNN \mathcal{A}^b , the
	number of most diverse prediction Q , the
	generated architectures \mathcal{A} to be evaluated.

Output: The fitness values of A.

1 $Y \leftarrow \emptyset$:

- 2 $\mathcal{A}^b_{encoded} \leftarrow$ Encode \mathcal{A}^b based on the details shown in Subsection III-A;
- **3** for $i \leftarrow 1$ to K do
- $CART \leftarrow$ Select the *i*-th CART from CARTs; 4
- $x \leftarrow \text{From } \mathcal{A}^b_{encoded}$ select the elements whose ids 5 are in I_i ;
- $y \leftarrow \text{Use } CART$ predict the classification accuracy 6 on x;
- $Y \leftarrow Y \cup x;$ 7

8 end

- 9 $Y \leftarrow$ Order the elements in Y;
- 10 $I_{selected}^{CART} \leftarrow$ Uniformly select Q CARTs based on Y; 11 for $i \leftarrow 1$ to $|\mathcal{A}|$ do
- $F_i \leftarrow$ the mean prediction of Q selected CARTs 12 $(I_{selected}^{CART})$ on $|\mathcal{A}_i|$;

13 end

14 Return F.

to find the best CNN architecture that achieves a promising classification performance on the image data at hand. Therefore, two experiments are performed in this paper: 1) investigating the classification performance of the proposed performance predictor with AE-CNN, and 2) inspecting the efficiency of the proposed performance predictor. In this section, the selected peer competitors and benchmark datasets, as well as the parameter settings for these two types of experiments, are detailed.

A. Peer Competitors

In comparing the classification performance, the chosen peer competitors are divided into three different categories: the state-of-the-art CNNs whose architectures are manually designed, the state-of-the-art CNN architecture designs based on non-evolutionary algorithms (mainly based on reinforcement learning), and the state-of-the-art EDL algorithms. The first category covers DenseNet [50], ResNet [54], Maxout [57], VGG [58], Network in Network [59], Highway Network [60], All-CNN [61], FractaNet [62]. Considering the promising performance of ResNet, we use its two different versions: ResNet with the depths of 101 and 1,202, respectively. For the convenience of the discussion, they are denoted as ResNet (depth=101) and ResNet (depth=1,202), respectively. The second category consists of NAS [9], MetaQNN [8], EAS [9], and Block-QNN-S [10]. The third category includes Genetic CNN [11], Large-scale Evolution [12], Hierarchical Evolution [13], and CGP-CNN [14]. Considering the proposed performance predictor is introduced in a case study of AE-CNN, AE-CNN combined with E2EPP (denoted as AE-CNN+E2EPP) is chosen to perform this experiment.



Fig. 2. Examples of CIFAR10 and CIFAR100 shown in Figs. 2a and 2b, respectively. Each row represents two samples from the same category and the words in the left refer to the corresponding category name.

For the second experiment, the selected peer competitors are the existing performance predictors [30], [32] discussed in Subsection II-B.

B. Benchmark Datasets

Benchmark datasets are only required by the first experiment. The CIFAR10 and CIFAR100 datasets are selected as the benchmark datasets. Both benchmark datasets are selected mainly because of their wide adoption by the state-of-the-art CNNs and CNN architecture design algorithms.

CIFAR10 is a 10-category natural object classification dataset, consisting of a training dataset with 50,000 images and a test dataset with 10,000 images, and each image has the dimension of 32×32 . In the training dataset, each category has roughly the equal number of samples, while in the test dataset, each category has the exact same number of samples.

CIFAR100 is just like CIFAR10, except that it is 100category. Because CIFAR100 has the same number of images in the training images and test images as those of CIFAR10, each category in CIFAR100 only has one-tenth of images as that in CIFAR10. Due to the smaller number of training data in each category and a larger number of classification categories than CIFAR10, CIFAR100 is a more complex dataset for classification than CIFAR10.

For each image of both benchmark datasets, the object to be classified commonly occupies a small area of the entire image, and also the size, the area, and the position of each object differ to each other in other images, even when they are from the same category. An example of these two benchmark datasets are shown in Fig. 2 for a glance, where each row denotes the objects from the same category, and the label leading in each row denotes the ground-truth of the corresponding object.

C. Parameter Settings

In this subsection, the parameter settings for generating the training data from AE-CNN are detailed at first, and then those of the proposed performance predictor are introduced next. Noting that these parameter settings are applied to both experiments.

Based on the conventions of the deep learning community, the SGD algorithm is used to train the CNN architectures whose weights are initialized with the commonly used Xavier method [63]; the batch size is set to 128; the weight decay is set to 5×10^{-4} ; each CNN is trained 350 epochs, and the learning rate is set to 0.01 for the first epoch and the 151-th to the 249-th epoch, 0.1 for the second to the 150-th epoch, and 0.001 for the remaining epochs. The classification accuracy is evaluated on the "validation dataset1". Because the used benchmark datasets do not have the corresponding validation dataset, 20% images are randomly selected from the corresponding training dataset as the validation dataset based on the conventions. All the CNNs are trained on three GPUs with the same model of NVIDIA 1080TI. In addition, we set both maximal numbers of RB and DB to 4 because of the limited GPU memory. Because each pooling operation halves the input size one time, and the input sizes of CIFAR10 and CIFAR100 are both 32×32 , the maximal number of pooling layers is set to 4. Furthermore, based on the conventions of DenseNet, k is selected from $\{12, 20, 40\}$, and the maximal numbers of DUs in a DB are 10 when k = 12 and k = 20, and 5 when k = 40. The maximal number of RUs in an RB is set to 10. Based on our configuration for generating the training data, N_b and N_p are set equal to 8 and 4, respectively, and the decision variable length of each training data for the random forest is 32 $(8 \times 3 + 4 \times 2)$. To build the performance predictor pool, we generate 1000 CARTs. The threshold of stopping node splitting is set as $1e - 4 \times \sigma^2$ (σ^2 is the variance of the training data) when growing each CART. In each generation, 100 CARTs are selected to combine the random forest-based predictor. All the parameter settings are summarized in Table I.

In performing the CNN architecture design by using AE-CNN and the proposed E2EPP method, we set both the maximal generation number and the population size to 20, the probabilities for the crossover and mutation are set to 0.9 and 0.1, respectively, as suggested in AE-CNN. Noting that before E2EPP is used for each generated CNN architecture, the CNN architecture is evaluated in the GPUs for one epoch so that the generated CNN can be normally evaluated by E2EPP, i.e., the employed GPUs can carry the CNN architecture and will not lead to an out-of-memory error. This is also for keeping a similar data distribution between the training data and the test data for E2EPP. After the evolutionary process, the individual with the best classification accuracy is chosen and then fully trained with the same training routine as collecting

¹We are aware of different definitions on the validation (data)set from literature. Based on the convention of the machine learning community, the validation set is used only for controlling overfitting and model selection, and the selected model cannot be updated and improved via the validation set. In evolutionary deep learning, after the individuals (models) are selected based on their performance of the "validation set", they are updated and possibly improved by genetic operators during evolution. So we use the quotation marks to highlight its different meaning here from its traditional meaning.

the training data for five independent runs, and the best result is reported, which is followed the conventions from deep learning community.

TABLE I A summary of the parameter settings.

Parameter name	Parameter value
batch size	128
weight decay	5×10^{-4}
training epochs	350
learning rate	0.01 for 1, and 151-249 epochs;
	0.1 for 2–150 epochs; 0.001 for
	251-350 epochs
k of the DB	$\{12, 20, 40\}$
maximal number of DUs in a DB	10 when $k = 12$ and $k = 20; 5$
	when $k = 40$
maximal number of RUs in a RB	10
maximal number of DBs (N_p)	4
maximal number of RBs (N_b)	8
generation number	20
population size	20
crossover probability	0.9
mutation probability	0.1
number of CARTs	1000
threshold of stopping node splitting	$1e - 4 \times \rho^2$ (ρ^2 is the variance of
	the training data)
number of selected CARTs	100

V. EXPERIMENTAL RESULTS

The results of the designed experiments are presented and analyzed in this section. Specifically, the classification performance of AE-CNN+E2EPP, in terms of the classification accuracy and the consumed GPU days, are elaborated in Subsection V-A. In order to extensively investigate the proposed E2EPP performance predictor, its efficiency and effectiveness are individually tested, and the corresponding experimental results are shown in Subsections V-B and V-C, respectively. In addition, the comparision to Radial Basis Network (RBN) has also been done to show the promising performance of random forest employed in the proposed E2EPP performance predictor.

A. Overall Results

Table II presents the experimental results in terms of the classification accuracy and consumed GPU days of the compared algorithms. Specifically, Table II is divided into five rows: the second to fourth rows denote the state-of-the-art CNNs manually designed, the state-of-the-art CNN architecture design algorithms based on non-evolutionary algorithms, and the state-of-the-art CNN architecture design algorithms based on evolutionary algorithms (i.e., the EDL algorithms), respectively; the last row shows the result of the proposed performance predictor used in AE-CNN (i.e., AE-CNN+E2EPP). In addition, the first column shows the names of the compared algorithms; the second and the third columns show the classification accuracy on CIFAR10 and CIFAR100, and the fourth column shows the consumed GPU days for achieving the corresponding classification accuracy by the corresponding architecture design algorithms. These architecture design algorithms generally report their consumed numbers of GPUs and days. For the convenience of the comparison, we unify them by using "GPU

days" as an indicator expressing the utilized computational resource. For example, one GPU day means a GPU is fully consumed in one day to achieve the classification accuracy. All the results of the peer competitors in this table are extracted from their seminal papers, in addition to the GPU days that we convert it by multiplying the number of GPUs with the number of days taken. The symbol "–" means the corresponding paper does not provide the corresponding result publicly available. Noting that the GPU days of AE-CNN+E2EPP is mainly caused by collecting the training data but not performing E2EPP.

For the classification accuracy obtained by the state-of-the-art CNN manually designed, on the CIFAR10 benchmark dataset, AE-CNN+E2EPP achieves, on the average, the classification accuracy of 1.5% higher than ResNet (depth=101), VGG and All-CNN, 2.04% higher than ResNet (depth=1, 202) and Highway Network, and even 3.78% higher than Maxout and Network in Network, respectively, while slightly lower than DesnNet (0.04%) and FractalNet (0.06%). On the CIFAR100 benchmark dataset, AE-CNN+E2EPP, whose classification accuracy is 77.98%, outperforms all other peer competitors that achieve the classification accuracy varying from 77.70% (by FractalNet) to 61.40% (by Maxout). AE-CNN+E2EPP demonstrates the promising performance among peer competitors in this category.

Compared with non-evolutionary algorithms, AE-CNN+E2EPP achieves the better classification accuracy than NAS and MetaQQ, but a little worse than EAS and Block-QNN-S on CIFAR10; on CIFAR100, the classification accuracy of AE-CNN+E2EPP is significantly better than that of MetaQQ while slightly worse than that of Block-QNN-S. In addition, NAS, MetaQNN and Block-QNN-S achieve their classification accuracies by consuming 22, 400, 100 and 90 GPU days, respectively, while AE-CNN+E2EPP only consumes 8.5 GPU days. In addition, AE-CNN+E2EPP also consumes fewer GPU days than that of EAS.

Compared with other EDLs, AE-CNN+E2EPP shows lower classification accuracy than that of Hierarchical Evolution, while higher than those of Genetic CNN, Large-scale Evolution and CGP-CNN on CIFAR10. On CIFAR100, AE-CNN+E2EPP shows the highest classification accuracy than others. Furthermore, AE-CNN+E2EPP also consumes the fewest GPU days than other peer competitors in this category.

In fact, EAS, Block-QNN-S, Hierarchical Evolution and CGP-CNN are all semi-automatic CNN architecture design algorithms, i.e., they design the CNN architectures based on both the expertise of manually designed CNNs and the automatic ways of EDLs. To this end, it is reasonable that CNN+E2EPP is inferior to them in terms of the classification accuracy. In summary, AE-CNN+E2EPP wins 26 times out of the 30 classification accuracy comparisons, and also consumes the least GPU days among all architecture design peer competitors.

B. Efficiency of E2EPP

The experimental results presented in Table II have indicated that AE-CNN can achieve promising classification accuracy when the proposed E2EPP is used, which is the ultimate goal of designing the performance predictor. In this subsection,

	Peer Competitors	CIFAR10	CIFAR100	GPU Days
	DenseNet [50]	94.76	75.58	_
	ResNet (depth=101) [54]	93.57	74.84	_
	ResNet (depth=1,202) [54]	92.07	72.18	-
state of the out CNNs	Maxout [57]	90.70	61.40	_
state-of-the-art CNNs manually designed	VGG [58]	93.34	7.95	_
	Network in Network [59]	91.19	64.32	-
	Highway Network [60]	92.28	67.61	-
	All-CNN [61]	92.75	66.29	-
	FractalNet [62]	94.78	77.70	_
	NAS [9]	93.91	-	22,400
CNN architecture design algorithms	MetaQNN [8]	93.08	27.14	100
based on non-evolutionary methods	EAS [9]	95.77	-	10
	Block-QNN-S [10]	95.62	79.35	90
	Genetic CNN [11]	92.90	70.95	17
CNN architecture design algorithms	Large-scale Evolution [12]	94.60	77.00	2,750
based on evolutionary methods	Hierarchical Evolution [13]	96.37	-	300
based on evolutionary methods	CGP-CNN [14]	94.02	-	27
	AE-CNN + E2EPP	94.70	77.98	8.5

TABLE II THE COMPARISON OF AE-CNN+E2EPP AND THE PEER COMPETITORS IN TERMS OF THE CLASSIFICATION ACCURACY (%) AND THE CONSUMED GPU DAYS ON THE CIFAR10 AND CIFAR100 BENCHMARK DATASETS.

we will specifically investigate the efficiency of E2EPP by disabling and enabling it in AE-CNN. To do a fair comparison, AE-CNN is performed with the same parameter settings as those of AE-CNN+E2EPP on both CIFAR10 and CIFAR100. We record the consumed GPU days of AE-CNN when it has a new generated CNN architecture whose fitness value satisfies the condition formulated by (2)

$$\left|\frac{f_1 - f_2}{f_2}\right| \le 0.01 \tag{2}$$

where f_1 denotes the fitness of the new CNN architecture generated by AE-CNN, and f_2 denotes the fitness of AE-CNN+E2EPP shown in Table II.

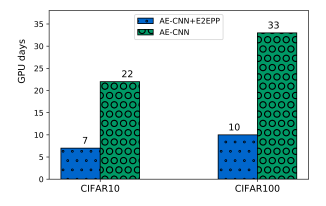


Fig. 3. The consumed GPU days of AE-CNN and AE-CNN+E2EPP when achieving the same classification accuracy on CIFAR10 and CIFAR100.

Fig. 3 shows the comparisons of the consumed GPU days when AE-CNN satisfies the condition of (2) on CIFAR10 and CIFAR100. The numbers above each bar display the consumed GPU days by the corresponding algorithm. The bars filled with the symbol of "·" denote the results of AE-CNN+E2EPP, while the bars filled with the symbol of "o" denote that of AE-CNN. Clearly, it can be seen from Fig. 3 that E2EPP has significantly improved the efficiency of AE-CNN for finding the best CNN architectures. Particularly, E2EPP saves 214% and 230% fewer GPU days on CIFAR10 and CIFAR100, respectively, when it is used in AE-CNN. Therefore, the goal of designing such a proposed performance predictor, in terms of speeding up the fitness evaluation, has been achieved.

C. Effectiveness of E2EPP

In order to check the effectiveness of E2EPP, the FBO [30] and Peephole [32] algorithms, which are the representatives of existing performance predictors as detailed in Subsection II-B, are selected to do the quantitative and qualitative comparisons against E2EPP. For the quantitative comparison, we choose the Mean Square Error (MSE), Kendall's Tau (KTau) [64] and the Coefficient of Determination (CoD) [65] as the indicators suggested in [32]. Specifically, KTau measures the correlation between the ranks of the prediction and their true ranks. A higher KTau value implies the higher correlation and the KTau value varies between [-1, 1]. The CoD measures the closeness degree of the prediction to its true value, and its value ranges from $-\infty$ to 1 where a larger value means the closer degree. For the qualitative comparison, we scatter the true values with the corresponding prediction on a 2-dimension space where the horizontal axis denotes the true values while the vertical axis denotes the prediction. The parameter settings of FBO and Peephole are set based on the suggestions in [32]. In addition, the experiment on each benchmark dataset is performed 10 times.

TABLE III THE MEAN SQUARE ERROR (MSE), KENDALL'S TAU (KTAU), COEFFICIENT OF DETERMINATION (COD) AND THE USED TIME OF FBO, PEEPHOLE, AND E2EPP ALGORITHMS ON THE **CIFAR10** DATASET.

	MSE	KTau	CoD	Time (seconds)
FBO	0.0077	0.2170	-4.346	3.702
Peephole	0.0019	0.5324	0.3765	359.296
E2EPP	0.0006	0.6604	0.5184	3.813

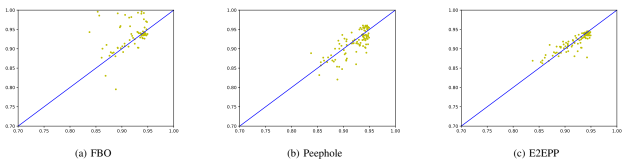


Fig. 4. The prediction and the true values of EBO, Peephole and E2EPP on the CIFAR10 dataset. In each subfigure, the horizon axis denotes the true values, while the vertical axis denotes the corresponding prediction.

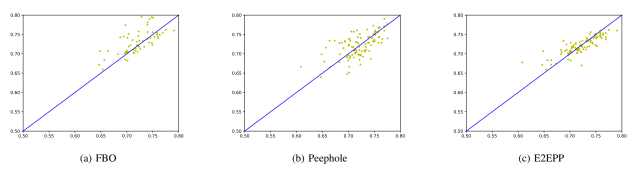


Fig. 5. The prediction and the true values of EBO, Peephole and E2EPP on the CIFAR100 dataset. In each subfigure, the horizon axis denotes the true values, while the vertical axis denotes the corresponding prediction.

TABLE IV THE MEAN SQUARE ERROR (MSE), KENDALL'S TAU (KTAU), COEFFICIENT OF DETERMINATION (COD) AND THE USED TIME OF FBO, PEEPHOLE, AND E2EPP ALGORITHMS ON THE **CIFAR100** DATASET.

	MSE	KTau	CoD	Time (seconds)
FBO	0.0078	0.1923	-2.631	3.725
Peephole	0.0024	0.5412	0.3238	458.652
E2EPP	0.0015	0.6501	0.3872	3.514

The quantitative comparison results on CIFAR10 and CI-FAR100 are shown in Tables III and IV, respectively. In addition, we also add the consumed time (seconds) measuring how long E2EPP, FBO and Peephole take to finish the experiment. Clearly as can be seen from Table III, FBO and Peephole obtain the MSE of 0.0077, and 0.0019, respectively, while E2EPP obtains the MSE of 0.0006 that is an order of magnitude less than FBO and Peephole can. This means that E2EPP could achieve a better prediction than FBO and Peephole. For the KTau indicator, FBO, Peephole and E2EPP achieve the values of 0.2170, 0.5324 and 0.6604, respectively, which imply that the performance predicted by E2EPP has the best correlation. For the CoD indicator, E2EPP again wins the highest value of 0.5184 (-4.346 for FBO and 0.3765 for Peephole). Based on the definition of CoD, the predicted value of E2EPP is the closest one to the ground truth value among the comparisons. In addition, E2EPP uses 3.813 seconds for the prediction, which is a little worse than FBO that uses 3.702 seconds. However, it is still much better than Peephole that

uses 359.296 seconds. The similar comparison results can also be investigated from Table IV. Specifically, E2EPP obtains the MSE of 0.0015 that is about half of that from Peephole and 1/5 of that from FBO, i.e., E2EPP has the minimal prediction error than the others. Furthermore, FBO obtains the KTau indicator value of 0.1923, while Peephole and E2EPP obtain the values of 0.5412 and 0.6501, respectively. For the CoD indicator, E2EPP achieves the best one, i.e., 0.3872, while FBO and Peephole acquire -2.631 and 0.3238, respectively. This shows the best prediction performance of E2EPP on the CIFAR100 dataset. In addition, E2EPP consumes 3.514 seconds to finish the prediction, which is the best among FBO (3.725 seconds) and Peephole (458.652 seconds).

The qualitative comparisons on CIFAR10 and CIFAR100 are scattered in Figs. 4 and 5, respectively, where the x axis of each point (denoted as (x, y)) represents the true value and y axis denotes the corresponding prediction. For the convenience of the comparison, we also plot the line y = x in each figure. The points standing above y = x mean the predictions are higher than the true values, while the points standing below y = x means the predictions are lower than the true values. The more points closer to y = x, the better prediction of the corresponding performance predictor. As can be clearly seen from Figs. 4a and 4b, FBO has only half of the total points that are close to the line of y = x, and Peephole achieves better results by having more points close to the line of y = x. However, it is clearly observed from Fig. 4c showing the results of E2EPP, almost all of the points are close to the line y = x. This means that the predicted values of E2EPP have minimal prediction errors, which is consistent with the quantitative analysis shown in Table III. From Figs. 5a, 5b and 5c that show the results performed on CIFAR100 by FBO, Peephole and E2EPP, respectively, it can be seen that FBO and Peephole achieve the similar results that have a wide spread of points around the line of y = x, while the results of E2EPP have the closer distance to the line y = x. This also shows the same conclusion from the quantitative analysis observed in Table IV.

D. Comparison to Radial Basis Network (RBN)

In order to demonstrate the promising performance of the random forest employed by the proposed E2EPP performance predictor, the comparison to RBN is performed, and the results are shown and analyzed in this subsection.

Specifically, an RBN is a three-layer neural network, except that the activation function of each unit in the hidden layer is a Gaussian function, and the output layer is a linear weight summation of the output of the hidden layer. The RBN is often used for regression tasks, which is similar to the function of random forest used in the proposed E2EPP performance predictor. Another reason for choosing RBN here for the comparison is that RBN is the base of our previous work [44] that motivates the design of E2EPP in this paper. For a fair comparison, we only replace the random forest by an RBN, while the other parts are kept the same as that used for the experiment shown in Subsection V-A. For the convenience of the discussions, we denote the RBN used by the AE-CNN algorithm as AE-CNN+RBN.

In the comparison, we only show the classification accuracies on CIFAR10 and CIFAR100, but not showing the consumed GPU days since the performance of both AE-CNN+E2EPP and AE-CNN+RBN does not rely on GPUs. The number of GPU days shown in Table II of AE-CNN+E2EPP is used for measuring the validity of the generated CNN architectures during the evolutionary process. In addition, the number of hidden units of the RBN is set to the same as the input size based on the conventions of practice [44]. The comparison results are shown in Table V.

TABLE V THE COMPARISON OF AE-CNN+E2EPP AND AE-CNN+RBN IN TERMS OF THE CLASSIFICATION ACCURACY (%) ON THE CIFAR10 AND CIFAR100 BENCHMARK DATASETS.

	CIFAR10	CIFAR100
AE-CNN + E2EPP	94.70	77.98
AE-CNN + RBN	82.33	70.20

It can be seen from Table V that AE-CNN+E2EPP achieves the classification accuracies of 94.70% and 77.98% on CI-FAR10 and CIFAR100, respectively, while AE-CNN+RBN achieves those of 82.33% and 70.20%, respectively. This implies that the random forest employed in the proposed E2EPP performance predictor is better than RBN. In fact, it is easy to find the proof of this conclusion, when the RBN is used, the mean and standard deviations of the Gaussian functions are calculated based on the training data. However, the training data in E2EPP is encoded with some numerical numbers that do not refer to the real meaning. In addition, the RBN has been well-known to be suitable for continuous data, while the training data in E2EPP is discrete. In summary, AE-CNN+E2EPP outperforms AE-CNN+RBN on the CIFAR10 and CIFAR100 benchmark datasets.

As can be seen above, we have performed four groups of experiments that are carefully designed to verify the promising performance of the proposed E2EPP performance predictor. Particularly, the first is for investigating whether E2EPP could enable EDL to achieve the competitive performance in designing the CNN architectures. The second validates whether E2EPP could accelerate EDL or not. The third quantitatively and qualitatively investigates whether E2EPP could outperform the existing approaches regarding the performance prediction, and the fourth investigates whether the random forest employed in E2EPP could perform better than the RBN that is another commonly used regression model. In addition to the third one, all the others are performed in the case of AE-CNN. All the experimental results demonstrate the effectiveness and efficiency of the proposed E2EPP performance predictor. From these experimental results, we can recognize that the performance predictor is indeed able to speed up EDL, and the end-to-end nature is convenient to use in practice by avoiding manual interventions during the performing of EDL. In addition, the random forest, as an ensemble approach, achieves the promising performance but relying on only limited training samples, which is a good characteristic especially in EDL where sufficient training data is not easy to obtain. Furthermore, the random forest could directly use the discrete number as its input, which is also another good characteristic in predicting the performance of CNNs when the architecture information is used as the input data. This implies that the ensemble methods depend on discrete input data may be a proper choice in designing the performance predictors, which is not only suitable to EDL but also other deep learning model selection methods.

VI. CONCLUSIONS AND FUTURE WORK

The goal of this paper was to develop an effective and efficient end-to-end performance predictor, named E2EPP, to speed up evolutionary deep learning algorithms in designing the best CNN architectures for given tasks. This goal has been achieved by proposing a random forest-based offline surrogate model. In order to enable the random forest to accept the CNN architectures as its input data, we also develop an encoding schema that is able to uniquely map the description of a CNN architecture into a numerical decision variable. In addition, we took the advantage of random forest in terms of the ensemble characteristic that each tree in the random forest randomly selects a subspace of the whole features as its training data. Furthermore, in order to improve the diversity of used off-line model, only a part of the decision trees in the random forest is selected based on their prediction on the best individual to do the prediction, which in turn enhances the accuracy of the predicted values. The proposed performance predictor is firstly examined by integrating it into an existing evolutionary deep learning algorithm, which is called AE-CNN+E2EPP, on CIFAR10 and CIFAR100. Through the comparisons with eight CNNs whose architecture are manually crafted, and another eight CNN architecture designs, AE-CNN+E2EPP shows the very promising results on both classification accuracy and consumed GPU days. In addition, the efficiency of the E2EPP is also examined on AE-CNN by enabling and disabling it. The results show that E2EPP could save 2/3 of computational time when achieving the same classification accuracy. Further, the E2EPP is compared with the other two state-of-the-art performance predictors, namely FBO and Peephole, on three measurements. E2EPP shows the best results on these three measurements. Especially, E2EPP uses only 1/100 computational time of that Peephole uses. It also shows that E2EPP achieves the best prediction through the qualitative comparison among these three performance predictors on both CIFAR10 and CIFAR100 datasets.

The proposed performance predictor is based on the firsttrain-then-to-predict paradigm, which requires a set of trained samples in advance. Theoretically, the more trained samples, the better prediction results. However, collecting more trained samples implies consuming more computational resource, which leads to long processing time. Achieving a better performance prediction results conflicts to the goal of designing the performance predictors. Therefore, finding a good balance between these two objectives may be an efficient way to use the performance predictor, which is left for future work. In addition, we will also integrate E2EPP into other CNN architecture design algorithms to solve some specific tasks, such as recognizing breast cancer on the BreakHis dataset [66].

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