GeneCAI: Genetic Evolution for Acquiring Compact AI

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ABSTRACT
In the contemporary big data realm, Deep Neural Networks (DNNs) are evolving towards more complex architectures to achieve higher inference accuracy. Model compression techniques can be leveraged to efficiently deploy such compute-intensive architectures on resource-limited mobile devices. Such methods comprise various hyperparameters that require per-layer customization to ensure high accuracy. Choosing such hyperparameters is cumbersome as the pertinent search space grows exponentially with model layers. This paper introduces GeneCAI, a novel optimization method that automatically learns how to tune per-layer compression hyperparameters. We devise a bijective translation scheme that encodes compressed DNNs to the genotype space. Each genotype’s optimality is measured using a multi-objective score based on accuracy and number of floating point operations. We develop customized genetic operations to iteratively evolve the non-dominated solutions towards the optimal Pareto front, thus, capturing the optimal trade-off between model accuracy and complexity. GeneCAI optimization method is highly scalable and can achieve a near-linear performance boost on distributed multi-GPU platforms. Our extensive evaluations demonstrate that GeneCAI outperforms existing rule-based and reinforcement learning methods in DNN compression by finding models that lie on a better accuracy/complexity Pareto curve.

CCS CONCEPTS
• Computing methodologies → Neural networks; Genetic algorithms: Search methodologies; • Hardware → Hardware-software codesign; • Theory of computation → Bio-inspired optimization;

KEYWORDS
Deep Learning, Genetic Algorithms, Multi-objective Optimization, Computer aided/automated design, Parallel Optimization

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1 INTRODUCTION
With the growing range of applications for Deep Neural Networks (DNNs), the demand for higher accuracy has led to a continuous increase in the complexity of state-of-the-art models. Such high execution cost hinders the deployment of DNNs in real-time applications on commodity hardware. Fortunately, modern neural networks have been shown to incur high redundancies that can be eliminated without compromising inference accuracy [15]. Effective identification and removal of such redundancies has fueled research in two interlinked domains: (i) Developing model compression techniques, e.g., pruning [9, 11, 17, 21, 24], quantization [46], and coding [7]. (ii) Devising automated policies that learn how to configure compression techniques to jointly achieve accuracy and compactness [10, 16, 33, 34, 39, 42]. In this paper, we focus on the latter.

The effectiveness of contemporary compression techniques relies on careful tuning of several hyperparameters across DNN layers. These hyperparameters directly control the trade-off between accuracy and execution cost on a constrained device. The question to be answered is how to find an optimal hyperparameter configuration that results in a high compression rate while minimally affecting inference accuracy. Figure 1 shows how an intelligent hyperparameter selection policy can better estimate the geometry of the optimal Pareto front for the same compression technique (Pruning).

Finding the optimal set of hyperparameters is quite challenging as the space of possibilities grows exponentially with number of DNN layers. Such large search-space renders manual or computer-ized greedy algorithms sub-optimal or infeasible for hyperparameter tuning. Existing work in automated compression suggests the use of heuristic methods [9, 11, 17] or Reinforcement Learning (RL) [10, 41, 42]. To tackle the high-dimensionality of the search space, heuristics and RL algorithms specify the hyperparameters one layer at a time, thus overlooking the layer inter-dependencies.

In this paper, we take an alternative approach, called GeneCAI, that simultaneously tunes the compression hyperparameters for all DNN layers using genetic algorithms (GA). We devise a customized translator that encapsulates the hyperparameter configuration of all DNN layers as one fixed-length vector $x \in \mathbb{R}^d$, called an individual. Using this translation scheme, each hyperparameter vector translates to a unique compressed DNN. In this setting, the search for compression hyperparameters directly translates to optimizing an objective function $f(x) : \mathbb{R}^d \to \mathbb{R}$ over $x$ where $f(\cdot)$ is an arbitrary measure of quality. We suggest a unified formulation for $f(\cdot)$, called the score, which assesses the quality of individuals by combining inference accuracy and a desired execution cost, e.g., runtime.

GeneCAI iteratively evolves the initial population towards superior individuals. To promote high-quality initialization, we propose a context-aware boundary characterization that significantly contributes to convergence and search results. After initialization, each evolution iteration comprises three consecutive genetic operations, namely, selection, crossover, and mutation. We balance GeneCAI exploration and exploitation by devising a diversity-guided GA. In particular, we define and measure diversity at the genotype level and adapt our crossover and mutation operations accordingly.
maintaining a diversified population, GeneCAI ensures a fast and stable search and prevents premature convergence.

In high-level, GeneCAI offers several benefits: (I) it is highly cost-effective as it takes a post-processing approach where a pre-trained DNN is compressed while no training is required amid GA iterations. (II) GeneCAI exploits parallelism to reduce optimization time by concurrent sample evaluations; We develop an accompanying API that manages the infrastructure for multi-GPU execution of GeneCAI search. (III) GeneCAI makes no assumptions about the objective function; by adjusting the cost, one can apply GeneCAI to various DNN compression tasks and hardware constraints. (IV) GeneCAI encoding and genetic operations accommodate optimization of both discrete and continuous-valued hyperparameters.

To demonstrate GeneCAI generalizability, we evaluate our methodology on four compression techniques and several architectures. Our evaluations unveil the full potential of GA in learning to effectively combine multiple methods and push the limits of network compression. For VGG-16 on ImageNet, GeneCAI pushes the state-of-the-art FLOPs reduction from $5\times$ to $7.1\times$ with higher accuracy. For expert-designed efficient MobileNets, GeneCAI obtains on average $1.2\%$ higher top-1 accuracy than the MobileNet Pareto curve [12]. We further show that GeneCAI computational flow is highly scalable and enjoys a linear reduction in search time with an increased number of distributed computing resources.

2 BACKGROUND AND RELATED WORK

Neural Network Compression. Existing model compression techniques can be roughly categorized in two classes: post-training compression and compression-aware training. The post-training approach is applied to pre-trained DNNs, without modifying the standard training loss or the back-propagation routines [9, 11, 17, 18, 21, 23, 24, 27, 32, 38]. Alternatively, in compression-aware training [1, 5], the training loss or back-propagation algorithm is altered to create models that are inherently compressible.

Each compression method has its own benefits and tradeoffs; in either case, the compression hyperparameters still need to be specified. Most of the existing work in DNN compression literature rely on hand-crafted or heuristic methods for per-layer configuration of the compression hyperparameters, which can be cumbersome and result in sub-optimal solutions. Different from heuristic methods that compress one layer at a time, GeneCAI performs whole-network compression where all layers are simultaneously optimized to account for inter-layer correlations. We target post-training compression and apply fine-tuning only once. This allows us to customize DNNs for various hardware platforms with little search/training overhead.

Automated Architecture Search. Modifying the DNN architecture graph in search for higher training/inference efficiency [15] has gained much research traction recently. Several works focus on designing automated methodologies for achieving such compact and accurate neural networks. GA has been applied to Neural Architecture Search (NAS) [14, 22, 26, 29, 37, 40] where the goal is to build a neural network architecture from scratch. Different from NAS, this paper focuses on learning hyperparameters for DNN customization, which acts on pre-trained DNNs and does not require training-from-scratch. This property allows our algorithm to enjoy a fast search-space exploration using parallel computing resources.

Automated DNN Compression. Authors of [13] develop a pruning scheme that selects the pruned filters using GA rather than magnitude-based or gradient-based approaches [21, 28]. Nevertheless, the amount of pruning applied per layer is still unknown. GeneCAI addresses this remaining challenge by learning the compression hyperparameters. As a result, GeneCAI can be applied to generalized compression techniques and not just network pruning. Reinforcement Learning (RL) [10, 39] is another promising tool for DNN compression. Although effective in finding near-optimal solutions, RL applies compression one layer at a time to overcome the curse of dimensionality. Thus, the RL agent typically needs many learning episodes to identify the inherent inter-layer correlations.

GA is shown to produce competitive results with RL on the challenging tasks of neural architecture search and robotic control while enjoying higher scalability and significantly lower training overhead [29, 31, 40]. Inspired by this, we develop an optimization method based on GAs that achieves better/similar final solutions as prior art with much lower runtime and computational overhead. GeneCAI optimizes all hyperparameters at once with a high scalability by exploiting parallelism specifically in distributed settings.

3 PROBLEM FORMULATION

DNN compression, in high-level, is a transformation $T(M, \vec{x})$ that converts a pre-trained model $M$ to a compressed model $\hat{M}_d$ with lower computational complexity. In this process, the adjustable hyperparameter vector $\vec{x} \in \mathbb{R}^d$ controls the complexity and accuracy of the output model. A desirable compressed network $\hat{M}_d$ should satisfy two properties: (i) the generalization capability of the compressed model should resemble the original network and (ii) the execution cost of $\hat{M}_d$ on the target hardware platform should be as low as possible. These properties outline a multi-objective black-box optimization problem over the hyperparameter vector $\vec{x}$, dubbed an individual. We assume we have access to a scoring oracle, $f(\cdot)$, that assesses each individual based on the classification accuracy on a validation set and computational complexity of its corresponding compressed model. The scoring scheme is designed to capture the trade-off between model accuracy $A(\hat{M}_d)$ and hardware cost $C(\hat{M}_d)$. GeneCAI’s customized score translates the original bi-objective problem of DNN compression to a single-objective optimization:

$$\max_{\vec{x} \in \mathbb{R}^d} f(A(\hat{M}_d), C(\hat{M}_d)),$$  (1)

For simplicity, we show $f(A(\hat{M}_d), C(\hat{M}_d))$ as $f(\vec{x})$ in the rest of the paper. Since full knowledge about $f(\cdot)$ and/or its first derivatives cannot be assumed, often empirical evaluations and optimization is the only viable strategy. Brute-force evaluation, in general, is infeasible as the search-space grows exponentially with $d$. Instead, we propose an empirical zero-th order optimization algorithm based on GA, dubbed GeneCAI, for finding the maximum value in Eq. (1), i.e., $f^*$. We denote the population at the $t^{th}$ step of the GA by $\mathcal{P}^t = \{\vec{x}^n\}_{n=1}^N$ where $N$ is the population size. GeneCAI aims at finding a near-optimal population after $T$ iterations, $\mathcal{T} = \{\mathcal{P}^t\}_{t=1}^T$, where the individuals are in vicinity of the optimal solution:

$$f(\vec{x}^n_T) \approx f^* \quad \forall n \in \{1, \ldots, N\}$$  (2)

In particular, by adopting a guided search through genetic operations, GeneCAI iteratively samples better DNNs with higher $f(\vec{x})$. 

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4 GENECAI APPROACH

We provide a generic solution to effectively compress a pre-trained DNN while maximally preserving model accuracy. GENECAI automation policy acts on a pool of hyperparameters and explores the corresponding search-space using customized genetic operations. An overview of GENECAI genetic solution is shown in Figure 2 and Algorithm 1 presents the corresponding pseudo code. Below, we summarize the key steps in our optimization framework:

I. First, a pre-processing step characterizes the search-space boundaries within which the optimal solution can reside. These boundaries are specified based upon task-enforced constraints on inference accuracy. Using the acquired boundaries, an initial population of hyperparameter vectors \( \mathbf{x}_0 = \{ \mathbf{x}_1^0, \ldots, \mathbf{x}_N^0 \} \) are sampled (Sec. 4.4).

II. At each iteration, all individuals \( \mathbf{x} \) are translated to their compressed DNNs \( \hat{M}_C \) (Sec. 4.1). The optimization scores \( f(\mathbf{x}) \) are then evaluated in parallel (Sec. 4.2).

III. Based on the new evaluations, GENECAI genetic operations, i.e., selection, crossover, and mutation (Sec. 4.3), are performed to update the population towards a new, more competent, generation.

By iteratively applying steps II and III above, the non-dominated Pareto front for compressing a desired pre-trained DNN is obtained. In the following, we elaborate on each design component in detail.

Algorithm 1 GeneCAI Search Algorithm

Inputs: fitness oracle \( f(\cdot) \), population size \( (N) \), iterations \( (T) \), crossover parameters \( (P_{cross}, P_{swap}) \), mutation parameter \( P_{tweak} \).

Output: Population after \( T \) iterations, \( \mathbb{P}^T = \{ \mathbf{x}_n^T \}_{n=1}^N \).

1: \( \mathbb{P}^0 = \{ \mathbf{x}_n^0 \}_{n=1}^N \)\hspace{1cm} \( \triangleright \) Directed Initialization
2: \( \text{div}^{\text{thr}} = \frac{\text{div}(\mathbb{P}^0)}{2} \)
3: for \( t = 0 \ldots T - 1 \) do
4: Calculate scores: \( f_n = f(\mathbf{x}_n^t) \) \( \triangleright \) Evaluation
5: \( \mathbb{G}^t = \text{sample}(\mathbb{G}^t | \{ f_n \}_{n=1}^N) \) \( \triangleright \) Selection
6: Derive list of parents: \( \mathbb{G}^t \rightarrow \hat{\mathbb{G}}^t = \{ (p_1^t, p_2^t) \}_{n=1}^N \)
7: for \( (p_1, p_2) \) in \( \hat{\mathbb{G}}^t \) do
8: \( \mathbb{C}^t \text{.append}(\text{crossover}(p_1, p_2 | P_{cross}, P_{swap})) \) \( \triangleright \) Crossover
9: if \( \text{div}(\mathbb{C}^t) \leq \text{div}^{\text{thr}} \) then
10: \( \text{Adjust} P_{\text{mutate}} \)
11: \( \mathbb{P}^{t+1} = \text{Mutate}(\mathbb{C}^t | P_{\text{mutate}, P_{tweak}}) \)

4.1 Genetic Encoding

An initial step for application of GeneCAI is defining the pertinent search-space for the black-box optimization. To this end, we are in need of a global encoding scheme \( T(M, \mathbf{x}) \) that translates a vector of compression hyperparameters \( \mathbf{x} \) to its corresponding compressed architecture \( \hat{M}_C \). To ensure an effective search, our proposed translation possesses the following characteristics:

- **Distinctness.** Our encoding is bijective: given an individual \( \mathbf{x} \) and model \( M \), the compressed \( \hat{M}_C \) is uniquely determined.
- **Continuity.** A small change in an individual translates to a similarly small alternation in the corresponding compressed DNN. This is particularly important to ensure search convergence.
- **Scalability.** GeneCAI encoding offers a compact, low dimensional representation, enabling effective search. The search-space dimensionality scales linearly with the number of DNN layers and is invariant to number of DNN parameters/operations.

We direct our focus on **four** compression tasks, namely, structured [11] and non-structured [6] Pruning, Singular Value Decomposition [45], and Tucker-2 approximation [18]. Figure 3 shows a high-level view of our encoding scheme for a 4-layer neural network. For pruning, we allocate 1 hyperparameter per layer: \( \mathbf{x} \in \mathbb{R}^4 \). For low-rank approximation, we allocate 2 hyperparameters for Tucker-2 and 1 hyperparameter for SVD, per layer, resulting in \( \mathbf{x} \in \mathbb{R}^6 \). In the following, we explain each method’s hyperparameters and their interpretation as a compressed model in detail.

**Pruning.** Pruning reduces model size by setting a percentage of low priority parameters/channels to zero [6, 11, 21]. We allocate one continuous value \( p \in [0, 1] \), per layer, to represent the ratio of non-zero values. For the example of ResNet-50, each sample will thus be a vector \( \mathbf{x} \in \mathbb{R}^{10} \). We consider two contemporary DNN pruning methods, namely, structured and non-structured pruning. Structured pruning aims at removing a portion of feature-map channels while non-structured pruning removes a subset of DNN layer weights.

Having defined the vector of hyperparameters \( \mathbf{x} \) for pruning, we implement the transformation \( T(M, \mathbf{x}) \) following common practice in prior art. For structured pruning, we use the sum of absolute gradients of model loss with respect to ReLU feature-map channels for pruning priority [28]. For a ReLU layer with \( c \) feature-map channels and pruning rate \( p \), the \( \{p \times c\} \) channels with lowest priorities (lower absolute gradients) are removed. For non-structured pruning, we use the absolute value of weight parameters to prioritize them [6]. Given a weight \( W \in \mathbb{R}^{k \times k \times c \times f} \) and pruning rate \( p \), the \( \{p \times k \times k \times c \times f\} \) elements with lowest absolute values are pruned.

Figure 3: Vectorized representation of pruning for **Pruning** and **Decomposition**. Here, CONV and FC denote convolutional and fully-connected layers, respectively.
SVD. We apply SVD on weight parameters ($W$) of fully-connected layers ($W \in \mathbb{R}^{k \times f}$) and point-wise convolutions ($W \in \mathbb{R}^{1 \times 1 \times k \times f}$).

For SVD, the compression parameter is the decomposition rank which takes an integer value in $\{1, \ldots, R\}$ where $R = \min(c, f)$. For efficiency, we discretize the space of approximation ranks in each layer into 64 values and encode them as follows:

$$rank \in \{1 \leftarrow \frac{R}{64}, \ldots, 64 \leftarrow R\}, \quad R = \min(c, f)$$  \hspace{1cm} (3)

Tucker-2. Tucker decomposition is a generalized Higher Order SVD (HOSVD) for arbitrary-shaped tensors. We apply this method on 4-way weight tensors in convolutional layers, $W \in \mathbb{R}^{b \times k \times c \times f}$. We focus on Tucker-2 which only decomposes the tensor along $c$ and $f$ directions, i.e., output and input channels. For Tucker-2 decomposition, the compression hyperparameter for each convolution layer is a tuple of integer-valued approximation ranks ($r_1, r_2$) where $r_1 \in \{1, \ldots, c\}$ and $r_2 \in \{1, \ldots, f\}$. To increase search efficiency, we quantize the space of decomposition ranks to 8 bins per-way as follows:

$$r_1 \in \{1 \leftarrow \frac{c}{8}, \ldots, 8 \leftarrow c\} \quad r_2 \in \{1 \leftarrow \frac{f}{8}, \ldots, 8 \leftarrow f\}$$  \hspace{1cm} (4)

4.2 Objective Evaluation

We develop a customized scoring mechanism to assess the quality of individuals in each iteration of GeneCAI algorithm. The multi-objective score of GeneCAI simultaneously reflects the compressed DNN’s accuracy and computational complexity. The objective of DNN compression then translates to maximizing this score. We formalize DNN compression as a constrained optimization as follows:

$$\max_{\vec{x} \in \mathbb{R}^d} \Delta C(M, \vec{x})$$  \hspace{1cm} subject to $A(\hat{M}_x) > acc_{thr}$ \hspace{1cm} (5)

where $\Delta C(M, \vec{x})$ represents the normalized difference in hardware cost, e.g., number of floating-point operations (FLOPs), between the uncompressed network, $M$, and the compressed model, $\hat{M}_x$. The function $A(\cdot)$ denotes the inference accuracy and $acc_{thr}$ is a task-enforced threshold on the post-compression accuracy. Having an accuracy constraint is crucial since the optimization algorithm will converge to a model size of zero otherwise. To solve the constrained optimization problem in Eq. (5), we formulate it as the following primal unconstrained optimization using penalty methods [2]:

$$\max_{\vec{x} \in \mathbb{R}^d} \Delta C(M, \vec{x}) - \log \left(\frac{\text{PEN}_{\text{acc}}(\vec{x})}{\text{PEN}_{\text{thr}}(\vec{x})}\right)$$  \hspace{1cm} (6)

where the term $\log \left(\frac{\text{PEN}_{\text{acc}}(\vec{x})}{\text{PEN}_{\text{thr}}(\vec{x})}\right)$ is the exterior penalty function [4] that enforces a constraint on the accuracy, i.e., $A(\hat{M}_x) > acc_{thr}$. The function $\text{PEN}_{\text{acc}}(\vec{x})$ measures the accuracy degradation as follows:

$$\text{PEN}_{\text{acc}}(\vec{x}) = \begin{cases} A(M) - A(\hat{M}_x) & A(\hat{M}_x) \geq acc_{thr} \\ A(M) - A(\hat{M}_x) + e^{acc_{thr} - A(\hat{M}_x)} & A(\hat{M}_x) < acc_{thr} \end{cases}$$  \hspace{1cm} (7)

Figure 4 visualizes the accuracy penalty. To prevent undesirable drop of accuracy, we greatly diminish the score of individuals that cause lower accuracies than the set constraint, $acc_{thr}$. The log penalty term can thus be estimated as follows:

$$\log \left(\frac{\text{PEN}_{\text{acc}}(\vec{x})}{\text{PEN}_{\text{thr}}(\vec{x})}\right) = \begin{cases} \log(A(M) - A(\hat{M}_x)) & A(\hat{M}_x) \geq acc_{thr} \\ \log(acc_{thr} - A(\hat{M}_x)) & A(\hat{M}_x) < acc_{thr} \end{cases}$$  \hspace{1cm} (8)

For accuracy values satisfying the threshold, this term enforces the accuracy maximization objective and applying the logarithm smoothens the accuracy variations by damping sudden changes. For accuracy values below $acc_{thr}$, a linear penalty is applied to prevent further accuracy loss. To prevent numerical instability, we define our score function as the exponential of the primal optimization of Eq. (6) which is formalized in Eq. (9):

$$f(\vec{x}) = e^{\Delta C(M, \vec{x}) / \text{PEN}_{\text{acc}}(\vec{x})}$$  \hspace{1cm} (9)

Note that maximizing the score function of Eq. (9) is equivalent to maximizing its logarithm value in Eq. (6). The developed score function encourages reduction in hardware cost (in the numerator) while penalizing the decrease in model accuracy caused by compression (in the denominator). Our proposed scoring mechanism is globally applicable to various compression tasks and can be modified to reflect a variety of hardware costs, e.g., power, memory footprint, and runtime. To ensure efficiency, inference accuracies are measured on a small held-out portion of the training samples. GeneCAI scoring function successfully models the ultimate goal of high compression with minimal accuracy degradation.

4.3 Optimization with Genetic Operations

GeneCAI genetic optimization is a metaheuristic approach inspired by natural evolution and the notion of “survival of the fittest” and can be leveraged to explore large search-spaces. Our GA works on a population of individuals. The core idea is to encourage creation of superior individuals and elimination of the inferior ones. To this end, an iterative process evolves the previous generation into a new, more competent population by performing a set of bio-inspired actions, i.e., selection, cross-over, and mutation.

Figure 5 illustrates the process of evolving a new population from the previous one. After the current population is evaluated (see Eq. (9)), individuals are assigned scores representing their quality, i.e., fitness. The selection step then performs a sampling (with replacement) based on individuals’ fitness scores. Cross-over and mutation aim to explore the proximity of the selected individuals by injecting small random patterns. To promote a diverse non-dominated front of solutions, we tune the hyperparameters associated with mutation based on a diversity measure. This, in turn, maintains the balance between exploration and exploitation.
GeneCAI genetic operations are devised to be compatible with the translation method. In particular, our designed operations satisfy the “closure” property: the output vector of mutation/cross-over translates to a valid compressed DNN. Note that selection, and cross-over are unified across different compression techniques while the mutation operation needs to be customized per compression method.

**Selection.** GeneCAI selection stage attempts to choose high-quality individuals based on the ordering of Pareto dominance to generate the next population. GeneCAI adopts a fitness-proportionate selection mechanism. Let us denote the population at the beginning of $t$-th iteration by $\mathcal{P} = \{x_n\}_{n=1}^N$ with fitness scores $\{f_n\}_{n=1}^N$ obtained from Eq. (9). To select superior samples, we construct the following probability distribution over the population using the fitness scores:

$$p_n = \frac{f_n - f_{\min}}{\sum_{n=1}^N (f_n - f_{\min})},$$  

where $f_{\min}$ is the minimum score. Subtraction of the minimum score ensures that the probability of selecting the weakest individual is zero and it is always eliminated. The superior individuals, $S'_t$, are selected by performing a non-uniform random sampling of the old population where the probability of selecting an individual is derived from Eq. (10). The total number of individuals in the population remains unchanged across iterations. Superior individuals can be selected multiple times for the new population (sampling with replacement) while ones with low scores are rarely chosen. The combination of GeneCAI evaluation and selection stages preserves high-quality individuals and eliminates weak ones.

**Crossover.** We design a crossover operation which creates new individuals, i.e., new compression configurations, $C'_t$ by inheriting and combining compression ratios from a pair of parent individuals. The crossover is performed by randomly swapping corresponding elements of parent genotypes. Given the selected population $S_t$, we sort the individuals in descending order based on their fitness score. To form the cross-over pairs, we pick the fittest individual available as the first parent. We then choose the individual which has highest distance with the first parent as the second parent. Such distance-based selection of pairs is motivated by increasing diversity among the newly generated offspring, and promoting exploration. For two individuals $\bar{x}_1, \bar{x}_2 \in \mathbb{R}^d$, the distance is calculated as follows:

$$\text{dist}(\bar{x}_1, \bar{x}_2) = \frac{1}{d \times K} \sum_{i=1}^d (\bar{x}_1[i] - \bar{x}_2[i])^2$$  

where $K$ is a constant equal to the maximum allowed value for the corresponding compression hyperparameter, e.g., $K = 64$ for SVD decomposition. We use two parameters to control the crossover operation: $p_{\text{cross}}$ determines the probability of applying crossover between the parents, and $p_{\text{swap}}$ is the per-element swapping probability. The proposed crossover allows superior individuals to exchange learned patterns and transfer knowledge across the population.

**Mutation.** Mutation randomly tweaks each individual in the crossovered population $C'_t$ to create a new population $\mathcal{P}^{t+1}$ for the next iteration of the GA. We devise two variations of mutation for continuous and discrete-valued individuals.

- **Continuous-valued Individuals.** Each element of a continuous-valued individual is mutated by adding a random value drawn from a zero-mean Normal distribution $\mathcal{N}(0, 0.2)$. We then clip the values to ensure they remain in the allowed interval, i.e., $[0, 1]$.

- **Discrete-valued Individuals.** Discrete individuals are mutated by randomly incrementing/decrementing vector elements such that the value remains in the valid discrete range in Sec. 4.1.

Mutation allows exploration of the neighborhood of candidate points in the search-space. Similar to cross-over, we define two control parameters: $p_{\text{mutate}}$ is the probability that the individual gets mutated and $p_{\text{tweak}}$ determines the per-element tweaking probability. To maintain the balance between exploration and exploitation and avoid premature convergence, we tune the mutation parameters based on the diversity of individuals in $C'_t$. We use the dispersion of individuals in the population and define diversity as follows:

$$\text{div}(C'_t) = \frac{1}{N} \sum_{n=1}^N \left( \sum_{i=1}^d (x'_n[i] - x'_n[i])^2 \right)$$

where $x'_n \in \mathbb{R}^d$ is the mean of all individuals in $C'_t$. As can be seen, the diversity function is closely tied to the variance. The diversity can therefore be adjusted by controlling the per-element variance of the individuals in the crossovered population $\text{Var}[x_i]$. Let us denote an arbitrary individual after crossover by $\bar{y} \in C'_t$, which transforms to $y \in \mathbb{R}^{d+1}$ after mutation. The $i$-th element of $\bar{y}$ is thus sampled from a random variable $Y_i$ with the following probability distribution:

$$P(Y_i = \bar{y}_i) = \begin{cases} \frac{p_M}{1 - p_M} & \bar{y}_i = \bar{x}_i[i] + \eta \\ 1 - \frac{p_M}{1 - p_M} & \bar{y}_i = \bar{x}_i[i] \end{cases}$$

Here, $\sigma^2_\eta$ is the variance of the added perturbation. Summing up the vector values in Eq. (14) provides the new population diversity:

$$\text{div}(C^{t+1}) = \text{div}(C'_t) + d \times p_M \sigma^2_\eta$$

For a desired threshold on population diversity, we can therefore determine the mutation parameters using Eq. (15). Since tweaking multiple elements of the individual vector can result in drastic changes in the corresponding compressed DNN’s architecture and accuracy, we restrict $p_{\text{tweak}}$ to a small value ($p_{\text{tweak}} = 0.05$) and merely adjust $p_{\text{mutate}}$ for diversity control. In our experiments, we set the diversity threshold to be half the diversity for the randomly initialized population. Such adaptive tuning of mutation allows for a diversity-guided search and ensures a fast and stable convergence.

**4.4 Directed Initialization**

A na{"i}ve random initialization of samples in the first iteration can result in a slow and sub-optimal convergence. To address this, we utilize boundary characterization as a pre-processing step to enable a targeted sample initialization. The general behavior of inference accuracy for a compressed DNN $M_{\lambda}$ is monotonic: as the compression rates increase, the accuracy drops. As such, for any task-enforced threshold on accuracy, we can characterize the boundaries of $X_i$ on a per-layer basis. Boundary Characterization allows for a directed search that eliminates unnecessary exploration of outlier subspaces, i.e., regions that are unlikely to contain the optimization solution. Figure 6 visualizes the hyperparameter search-space and the outlier regions for pruning a two-layer neural network. The horizontal plane corresponds to different configurations of per-layer pruning rates and the vertical axis represents their quality (score $f(.)$).
The optimal solution to the search problem is a configuration with the highest score. The outlier regions in Figure 6 are therefore the flat sectors of the space. By filtering out the non-optimal outlier regions, GeneCAI sampling can find the solution within the region bounded by the initialization margins. We find a threshold vector $\bar{\theta}$ where each element constrains a single hyperparameter that corresponds to a certain layer in the compressed DNN. In the example of Figure 6, $\bar{\theta}$ has two elements, each presented by a dashed line ($[\bar{\theta}[1]$ and $[\bar{\theta}[2]$). Below we describe how the boundaries $\bar{\theta}[i]$ are obtained given an accuracy threshold $acc_{thr}$ for each compression task.

**Pruning.** We obtain a threshold vector $\bar{\theta} \in \mathbb{R}^d'$ with the $i^{th}$ element $\bar{\theta}[i]$ specifying the maximum pruning rate of the $j^{th}$ layer such that the inference accuracy of the compressed model $M_2$ does not violate the accuracy threshold $acc_{thr}$:

$$\bar{\theta}[i] = max\{p\} \text{ s.t. } \bar{x}[j] = \left\{ \begin{array}{ll} p & j = i \wedge A(\hat{M}_2) > acc_{thr} \\ 0 & j \neq i \end{array} \right. (16)$$

**Decomposition.** For SVD and Tucker-2 decompositions, the threshold vector $\bar{\theta}$ represents per-layer minimum ranks satisfying $acc_{thr}$:

$$\bar{\theta}[i] = min\{r\} \text{ s.t. } \bar{x}[j] = \left\{ \begin{array}{ll} r & j = i \wedge A(\hat{M}_2) > acc_{thr} \\ K_j & j \neq i \end{array} \right. (17)$$

where $K_j \in \{64, 8\}$ is the maximum allowed encoded rank for the $j^{th}$ hyperparameter (see equations 3&4).

Note that real-world neural networks have many more layers and the pertinent search-space is of much higher dimensionality than the 2-dimensional plane shown Figure 6. For a $d$-dimensional search-space, the proposed boundary characterization scheme reduces the effective (continuous) search volume from $[0, 1]^d$ to $\prod_{i=1}^d [\bar{\theta}[i]]$ for pruning. For decomposition, the (discrete) search-space size is reduced from $\prod_{j=1}^n K_j$ to $\prod_{i=1}^d (K_i - \bar{\theta}[i])$. Such approach significantly improves search convergence time and solution quality.

After performing boundary characterization, we randomly sample the initial population from the space enclosed by the threshold vector $\bar{\theta}$. For pruning, the $i^{th}$ element $\bar{x}[i]$ in each sample vector is drawn from a Normal distribution $N(\bar{\theta}[i]/2, \bar{\theta}[i]/2)$. For decomposition, the $i^{th}$ element $\bar{x}[i]$ is randomly selected from $[\bar{\theta}[i], \bar{\theta}[i] + 1, \ldots, K_i]$.  

**5 EXPERIMENTS**

We provide extensive evaluations on CIFAR-10 [20] and ImageNet [30] benchmarks. The evaluated network architectures include AlexNet, VGG, ResNet family, and MobileNets. All models are implemented using Pytorch library. The networks are trained from scratch following the parameter setup and training schedule adopted by the original papers [8, 12, 19, 35, 36]. For CIFAR-10, we use a VGG-variant as used in [17]. We compress the models using GeneCAI and fine-tune them for 20 and 60 epochs on CIFAR-10 and ImageNet, respectively. The genetic parameters are set as $P_{\text{tweak}} = 0.05, P_{\text{cross}} = 0.8, P_{\text{swap}} = 0.2 [40]$. $P_{\text{mutate}}$ is adaptively determined based on the population diversity, as discussed in Sec. 4.3. All results are from a single run of the GA.

**5.1 Quantitative Results on CIFAR-10**

We apply GeneCAI to pre-trained CIFAR-10 architectures and compare our results with prior art in Table 1. Specifically, we conduct experiments with non-structured pruning ($P_n$), structured pruning ($P_s$), SVD and Tucker decomposition ($D$), and combination of multiple compression methods ($D + P_s$). We set the population size to 100 for ResNet-56 and ResNet-50, 200 for ResNet-110, and 50 for VGG. We randomly select 1,000 images from the training data to use as validation set for score computation. The threshold ($acc_{thr}$) is set to 90% for ResNet-X and 60% for VGG. For all networks, we let GeneCAI run for 50 iterations.

**Non-structured Pruning ($P_n$ in Table 1).** We perform non-structured pruning on ResNet-50 and report the ratio of non-zero model parameters. $acc_{thr}$, is set to 93% and we do not perform any fine-tuning on the compressed model. As shown, GeneCAI achieves 1.34× lower parameters compared to state-of-the-art RL method [10]. Note that the lower FLOPs and comparable accuracy of [25] are due to training the model from scratch whereas GeneCAI and [10] do not include any fine-tuning.

**Structured Pruning ($P_s$ in Table 1).** We implement structured pruning by adding masks after ReLU activation layers. We base our comparisons on FLOPs per inference relative to the uncompressed baseline. With similar or higher accuracy, GeneCAI compressed models achieve 1.35× lower FLOPs than prior art (on average).

**Decomposition and Pruning ($D + P_s$ in Table 1).** To unveil the full optimization potential of our method, we allow GeneCAI to learn and combine multiple compression techniques, namely, pruning, SVD, and Tucker. The $D + P_s$ experiments are conducted by first decomposing the network and then applying structured pruning. As shown in Table 1, GeneCAI pushes the limits of compression by 2.45× on average with less than 1% drop in accuracy compared to state-of-the-art works. We also report FLOPs reduction by only applying decomposition (SVD and Tucker), shown by $D$ in Table 1.

**5.2 Quantitative Results on ImageNet**

Table 1 summarizes GeneCAI results for ImageNet dataset. Number of samples is 100 for ResNet-50, 50 for VGG-16, and 20 for AlexNet. We let GeneCAI run for 50 iterations with $acc_{thr} = 10\%$ for all models. This target accuracy is compensated by fine-tuning.

**Non-structured Pruning ($P_n$ in Table 1).** We perform non-structured pruning on AlexNet and report the ratio of non-zero model parameters. GeneCAI achieves higher accuracy with 3% higher parameter size compared to a Bayesian Optimization approach [3].

**Structured Pruning ($P_s$ in Table 1).** On ResNet-50, GeneCAI compresses the model to 1.5× less FLOPs on average while achieving similar/higher test accuracy compared to best prior works. On VGG-16, GeneCAI outperforms all heuristic methods and gives competing results with [25] and the state-of-the-art RL method [10]. Note that [25] does not propose a hyperparameter optimization algorithm and merely focuses on the training of already-compressed DNNs. As such, their approach is orthogonal to GeneCAI and can be combined with our method to further improve final accuracy.

**Decomposition and Pruning ($D + P_s$ in Table 1).** Using a combination of decomposition and pruning, GeneCAI achieves 1.85× lower
Table 2: GeneCAI pruning for MobileNetV1 & V2 on ImageNet.

| Dataset      | Model | Top1 (%) | Top5 (%) | FLOPs    |
|--------------|-------|----------|----------|----------|
| CIFAR-10     | Baseline | 93.7     | 100      | 220 M    |
|               | AMC [10] | 93.5     | 40.0     | 220 M    |
|               | RF [25]  | 93.4     | 20.0     | 220 M    |
|               | GeneCAI (P) | 93.6   | 30.0     | 220 M    |
| ImageNet     | Baseline | 60.7     | 80.0     | 220 M    |
|               | AMC [10] | 54.8     | 5.0      | 220 M    |
|               | GeneCAI (P) | 56.1   | 78.2     | 220 M    |

Table 3: GeneCAI compressed MobileNets Speedup on embedded CPU and GPU for structured pruning on ImageNet.

| Model         | Theoretical Speedup | Real Speedup |
|---------------|---------------------|--------------|
| ResNet-50     | 1.7x                | 1.6x         |
| ResNet-100    | 2.0x                | 1.7x         |
| MobileNetV1   | 2.7x                | 2.5x         |
| MobileNetV2   | 4.0x                | 3.4x         |

Table 4: Search runtime of GeneCAI for pruning on various benchmarks.

| Dataset   | Arch. | b | N_{iters} | Search Time (minutes) |
|-----------|-------|---|-----------|-----------------------|
| ImageNet  | AlexNet | 50 | 50 | 10 | 3 | 3 | 3 |
|           | VGG-16 | 50 | 50 | 112 | 57 | 38 | 28 |
|           | ResNet-50 | 100 | 50 | 145 | 73 | 49 | 36 |
|           | MobileNetV1 | 50 | 100 | 97 | 48 | 32 | 24 |
|           | MobileNetV2 | 50 | 100 | 116 | 57 | 38 | 28 |
|           | Cifar-10 | 50 | 50 | 3 | 2 | 1 | 1 |
|           | ResNet-50 | 100 | 50 | 35 | 19 | 13 | 11 |
|           | ResNet-56 | 100 | 50 | 7 | 5 | 3 | 3 |
|           | ResNet-110 | 200 | 50 | 55 | 30 | 22 | 16 |

5.3 Compressing Compact Networks

To further demonstrate the effectiveness of GeneCAI optimization, we apply compression to MobileNet architectures trained on ImageNet dataset. These networks are specifically designed for embedded applications with a strict efficiency constraint. As such, MobileNets inherently have very low complexity/weights which renders their compression quite challenging. We apply pruning to MobileNetV1 and MobileNetV2 with a population size of 50, and let GeneCAI run for 100 iterations. We compare the compression rate and accuracy achieved by GeneCAI with the FLOPs-accuracy Pareto curve of the original MobileNet architectures [12, 35], the state-of-the-art AutoML approach [10, 41], and compression-aware training methodologies, i.e., US-Nets [43, 44].

Table 2: GeneCAI pruning for MobileNetV1 & V2 on ImageNet.

| Policy       | Top1 (%) | Top5 (%) | FLOPs |
|--------------|----------|----------|-------|
| Baseline     | 70.6     | 89.5     | 569 M |
| MobileNetV1 (0.75x) [12] | 68.4 | 88.2 | 325 M |
| US-Nets [43] | 69.5 | 325 M |
| GeneCAI      | 70.8     | 89.3     | 223 M |
| MobileNetV1 (0.5x) [12] | 63.7 | 149 M |
| US-Nets [43] | 63.5 | 136 M |
| GeneCAI      | 64.1     | 85.4     | 136 M |
| MobileNetV2 (0.75x) [35] | 69.8 | 88.3 | 220 M |
| US-Nets [43] | 70.0 | 222 M |
| GeneCAI      | 70.1     | 89.5     | 220 M |

5.4 Search Overhead and Scalability

The core computational load in GeneCAI algorithm corresponds to the evaluation of a batch of b samples. For each sample x_i in the batch, the evaluation phase comprises transforming the sample to its corresponding compressed DNN \hat{M}_x, measuring inference accuracy on the validation data, and emulating the execution cost.

Since samples in a batch are independent, evaluation can be parallelized on multiple GPU devices to achieve faster search convergence. Table 4 summarizes the runtime of GeneCAI algorithm for several benchmarks and datasets. Runtimes are measured on a machine with an Intel Xeon E5 CPU and four NVIDIA Titan Xp GPUs. The results show high scalability: runtime drops almost linearly with the number of GPUs. The state-of-the-art RL algorithm reports ~ 1 hour to compress CIFAR-10 architectures [10]. For their most complex benchmark, i.e., ResNet-56, GeneCAI achieves a search time of only ~ 12 minutes on a single GPU and ~ 3 minutes on four GPUs.

Table 3: GeneCAI compressed MobileNets Speedup on embedded MobileNets for structured pruning on ImageNet.

| Model         | Theoretical Speedup | Real Speedup |
|---------------|---------------------|--------------|
| MobileNetV1   | 1.7x                | 1.6x         |
| MobileNetV2   | 2.0x                | 1.7x         |
|               | 2.7x                | 2.5x         |
|               | 4.0x                | 3.4x         |

MobileNetV1 1.4x 1.4x 1.4x
5.5 Analysis and Discussion

To illustrate GeneCAI methodology, we consider VGG architecture trained on CIFAR-10 compressed with structured (filter) pruning for $acc_{\text{thr}} = 60\%$. The initial samples obtained from our directed initialization scheme are shown in Figure 7-a, where each column corresponds to an individual and each row represents a model layer. After applying GeneCAI for $50$ iterations, the population pattern of Figure 7-b is learned. Upon convergence, individuals have similarly high scores. GeneCAI successfully learns expert-designed rules: first and last layers of the network (first and last rows in Figure 7-b) are given high densities to preserve accuracy.

![Initial Population](image1.png)

**Figure 7:** (a) Initialized population for structured pruning. (b) Population upon convergence.

GeneCAI performs whole-network compression by encoding all layers’ hyperparameters in one genotype. As such, GeneCAI effectively learns which hyperparameter configuration least affects model accuracy and most reduces the overall FLOPs. To show this capability, we present the per-layer FLOPs for VGG-16 network trained on ImageNet in Figure 8. The bars show the per-layer FLOPs percentage in the original model; the curve shows the per-layer percentage of pruned FLOPs in the compressed DNN. Different from prior art [17], GeneCAI prunes the first convolutions more and relaxes pruning for later layers as they have a minor role in FLOPs.

5.6 Ablation Study

In this section, we study the effect of various GeneCAI parameters on algorithm convergence and final FLOPs/accuracy. For brevity, we only focus on the task of structured pruning for VGG on CIFAR-10. We show the trend lines as well as a fraction of individuals (black dots) across GeneCAI iterations. **Figure 9-a** shows the evolution of FLOPs ratio for two initialization policies, one with uniformly random samples and one with our proposed initialization scheme discussed in Sec. 4.4. As seen, naive initialization greatly harms the convergence rate and final FLOPs.

**Effect of Initialization.** Figure 9-a shows the evolution of FLOPs ratio for two initialization policies, one with uniformly random samples and one with our proposed initialization scheme discussed in Sec. 4.4. As seen, naive initialization greatly harms the convergence rate and final FLOPs.

**Effect of Population Size.** Figure 9-b presents the effect of population size on GeneCAI convergence. A higher number of individuals results in a smoother convergence and lower final FLOPs. This effect saturates for a large enough population. In our experiments, we observed that the population size should be larger for searching vector spaces with higher dimensionality.

**Effect of Mutation and Cross-over Parameters.** We conduct two experiments, one with $(P_{\text{mutate}}, P_{\text{cross}}) = (0.8, 0.2)$ and the other with $(0.4, 0.1)$ and compare the convergence in Figure 9-c. Higher $(P_{\text{mutate}}, P_{\text{cross}})$ allows the GA to explore more possibilities, leading to faster convergence while smaller probabilities result in a more stable evolution. Thus, GeneCAI dynamically tunes the GA parameters to ensure balanced exploration and exploitation (see Sec. 4.3).

6 CONCLUSION

This paper introduces GeneCAI, a method to automate DNN compression using genetic algorithms. Our algorithm learns how the compression hyperparameters should be set across layers to achieve a better performance than models designed by human experts. The core idea behind GeneCAI is to translate compression hyperparameters into a vector of genes and explore the corresponding search-space using genetic operations. This approach allows GeneCAI to be generic and applicable to any combination of post-processing DNN compression methods. We showcase GeneCAI effectiveness on four DNN compression methods, namely, structured and non-structured pruning, SVD, and Tucker decomposition. Experiments show that GeneCAI achieves superior results compared to state-of-the-art heuristic and reinforcement-learning based algorithms on a variety of benchmarks. The proposed method is highly efficient as it does not require gradient computation and is entirely parallelizable.

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