Local Search is a Remarkably Strong Baseline for Neural Architecture Search

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Abstract. Neural Architecture Search (NAS), i.e., the automation of neural network design, has gained much popularity in recent years with increasingly complex search algorithms being proposed. Yet, solid comparisons with simple baselines are often missing. At the same time, recent retrospective studies have found many new algorithms to be no better than random search (RS). In this work we consider, for the first time, a simple Local Search (LS) algorithm for NAS. We particularly consider a multi-objective NAS formulation, with network accuracy and network complexity as two objectives, as understanding the trade-off between these two objectives is arguably the most interesting aspect of NAS. The proposed LS algorithm is compared with RS and two evolutionary algorithms (EAs), as these are often heralded as being ideal for multi-objective optimization. To promote reproducibility, we create and release two benchmark datasets containing 200K saved network evaluations for two established image classification tasks, CIFAR-10 and CIFAR-100. Our benchmarks are designed to be complementary to existing benchmarks, especially in that they are better suited for multi-objective search. We additionally consider a version of the problem with a much larger architecture space. While we find and show that the considered algorithms explore the search space in fundamentally different ways, we also find that LS substantially outperforms RS and even performs nearly as good as state-of-the-art EAs. We believe that this provides strong evidence that LS is truly a competitive baseline for NAS against which new NAS algorithms should be benchmarked.

Keywords: neural architecture search · local search · evolutionary algorithm · random search · multi-objective NAS · NAS baseline

1 Introduction

Deep learning has achieved excellent results for machine learning tasks in heterogeneous fields, including image recognition [21], natural language processing [19], games [25,29], and genomics [24]. While the availability of large amounts of data [14] and hardware advancements (e.g., tensor parallel processing units [12]) have been key enabling factors for deep learning, the success of the field is also
due to the ingenious design of competent neural network architectures (consider, e.g., the invention of residual connections [10]).

Proper and innovative architecture design requires experience, intuition, and expensive trial-and-error. This has sparked research on techniques to automate this task, i.e., the field of Neural Architecture Search (NAS) has emerged [8, 30]. NAS research is quickly gaining popularity: 2019 alone counts almost 50 publications. Most NAS proposals present new, typically increasingly complex, NAS algorithms. However, recent work questions whether the need for many of these algorithms is actually justified: [23] and [32] showed that several NAS algorithms are no better than Random Search (RS) when validated properly.

While RS can be considered perhaps the simplest form of a baseline, it is also by far the worst form of heuristic search in many cases, with the worst scalability. For this reason, it is often not even considered in many modern heuristic design studies. Yet, from RS to more complex search heuristics such as EAs, is still quite a leap. In the gap for instance lie almost equally simple and classical, yet far less random in nature, Local Search (LS) techniques. Still LS has, to the best of our knowledge, not yet been applied to NAS. In this paper, we do this for the first time, and in doing so, we make three contributions:

1. We propose a simple and parameter-less LS algorithm for multi-objective (accuracy vs. architecture complexity) NAS.

2. To enable quick and easy benchmarking, we release two datasets containing cached performances for over 200,000 Convolutional Neural Network (CNN) architectures for CIFAR-10 and CIFAR-100. Our datasets are designed to be complementary to similar existing proposals [6, 33]. We remark that we designed these datasets to be able to compare NAS approaches, and not necessarily to obtain State-of-the-Art (SotA) neural networks.

3. We provide evidence on the validity of LS being a superior baseline for NAS by conducting several experiments. In particular, we consider NAS in different search spaces; We compare LS with RS as well as with a classic and a SotA Evolutionary Algorithm (EA); We include an experiment on CIFAR-100 where the possible architectures are less constrained, leading to a large search space containing over 104 Billion possible neural architectures.

In this work, we consider NAS in a multi-objective setting. Single-objective NAS, i.e., searching only for good accuracy-networks, has already been amply investigated in [23, 32], where it was found that very different search algorithms, including RS, actually perform very similar. Multi-objective NAS on the other hand has been studied less (see Sec. 2), and is arguably a more interesting setting, where sophisticated search algorithms may potentially be needed. Alongside RS, we consider EAs, since they are a natural fit for multi-objective optimization, and are often used for NAS.

3 https://github.com/D-X-Y/Awesome-NAS
4 Benchmark datasets: https://github.com/ArkadiyD/MacroNASBenchmark
Source code for reproducibility: https://github.com/tdenottelander/MacroNAS
2 Related work

NAS can be broadly categorized by the level at which the search is performed, i.e., macro-level and micro-level [8] (different nomenclatures exist [30]). Macro-level search aims at composing (wiring) atomic cells of different type. A cell is a computational graph (a sub-net) composed of different operations/layers (convolutions, activation functions, etc.). In micro-level search, the position of cell types within the architecture is pre-decided (often a same cell type is repeated in multiple places), while it is the cells’ internal wiring that is optimized. Both levels can also be searched at the same time [15,17], potentially along with hyper-parameter settings [17,31]. In the remainder of the paper, we will refer to “architecture” and “network” (for brevity, “net”) interchangeably.

Search algorithms considered for NAS vary wildly. Reinforcement Learning (RL), Bayesian optimization approaches, and EAs are among the most popular [8]; the latter being typically employed in multi-objective scenarios. In this paper, we mostly focus on performing multi-objective NAS at macro-level. Therefore, in the following we report on recent works in this direction, and focus on whether comparisons against a baseline (i.e. RS) were made.

The work in both [4, 13] builds upon NSGA-II [5] to obtain a front of nets that trade off accuracy and compactness. Although nets with SotA performance are ultimately obtained, the search algorithm is not compared with any baseline. In [7], a multi-objective EA is proposed that includes Lamarckian mechanisms: instead of training nets from scratch, child nets inherit the weights of their parents and are only fine-tuned. Although the method is shown to outperform RS, it is unclear whether this is due to a difference with the way the EA searches, or solely because of weight inheritance. In [15], NSGA-Net is presented, which evolves fixed-length architectures at both macro- and micro-level. The authors show, through experimental analysis, that NSGA-Net is superior to RS. Summarizing, most of the literature work considers only RS as a baseline (sometimes in not entirely equal conditions), or no baseline at all. Importantly, we found no work investigating and comparing with an LS algorithm.

Since we release benchmark datasets of saved net evaluations for macro-level NAS, we also refer to important related works of this nature. NAS-Bench-101 [33] contains cached performance indicators for 423,000 nets on CIFAR-10, while NAS-Bench-201 [6] does the same also for CIFAR-100 and ImageNet, but for far less architectures (15,625). Both datasets are built for micro-level NAS. We did not consider NAS-Bench-201 because the limited number of possible architectures can limit the analysis of the search behavior for long run-times. We built new benchmark datasets for three reasons. First, we intended to provide the community with a macro-level search benchmark dataset for better reproducible NAS: we are not aware of other works proposing this. Second, NAS-Bench-101 includes infeasible solutions (our benchmark datasets do not), and choosing how to handle them substantially changes the search behavior of an algorithm. Lastly, for NAS-Bench-101 it has already been shown that RS is a competitive baseline in a single-objective setting [33]. We will show however that, in fact, in the
search space of NAS-Bench-101 it is hard to do any better than RS, even when considering a multi-objective setting (see Sec. 5.1).

3 Objectives, Encodings, Search algorithms

We now present the objectives to optimize, the encoding schemes we use, and the algorithms we consider in the experimental analysis. For a detailed description of the benchmark datasets we introduce, we refer the reader to the Appendix.

3.1 Objectives

The first objective $f_1$ is to maximize accuracy (fraction of correct net predictions). Particularly, we set $f_1$ to the validation accuracy ($\text{acc}_{\text{val}}$), i.e., the accuracy of the net for a set of data that is not used for training the net, to assess whether the net generalizes. Because the validation set is still used in the NAS optimization loop, we consider the ability of nets with good $\text{acc}_{\text{val}}$ to generalize to a second set of completely held-out data, i.e., the test set, in Sec. 5.4.

The second objective $f_2$ is to maximize a net’s efficiency, i.e., to minimize the Mega Multiply–ACcumulate operations (MMACs, 1 MMAC $\approx$ 2 FLOPs), which is the number of GPU computations a net requires to make a prediction. MMACs is a popular complexity metric [26] and is deterministic (in contrast to actual time taken). Because large differences in scale of the objectives can influence search mechanisms (e.g., the crowding distance [5]), we set MMACs to be in the same range as $\text{acc}_{\text{val}}$, i.e., we normalize MMACs to lie in $[0, 1]$. The min/max MMAC values for the normalization are trivial to determine as it suffices to consider the smallest/largest achievable nets. For the sake of a consistent optimization direction in both objectives, the secondary objective $f_2$ becomes maximizing $1 - \text{normalize}(\text{MMACs})$. Since no MMACs were reported for NAS-Bench-101, for the experiment in Sec. 5.1 we use $1 - \text{normalize}(\#\text{parameters})$.

3.2 Encodings

We use a direct encoding that represents feed-forward CNNs. The encoding is an array of discrete variables $[x_1, \ldots, x_\ell]$, where each $x_i$ takes values (cell types) in $\Omega_i$, a position-dependent alphabet.

Fig. 1: Best $\text{acc}_{\text{val}}$ net on CIFAR-10 (left) and cell types (right). For our benchmarks, 3 types are possible for normal cells (types 1, 2 and 5) and reduction cells (positions 5, 10, and 15) are fixed to type 1. For the large-scale experiment, 5 types are available for all cells. For details on cell types see the Appendix.
In our benchmark datasets, the number of searchable cells (length of the encoding) is $\ell = 14$. In particular, only normal cells, i.e., cells that maintain the height and width of the input they receive \cite{34}, can be optimized. Conversely, cells that reduce spatial dimensions (reduction cells) are fixed. The alphabet $\Omega_i$ is identical for each position $i$ and contains 3 options. One option is the identity, i.e., a placeholder cell that forwards information as is. The other two options are commonly used convolutional blocks from literature (described in the Appendix). Fig. 1 shows an example of an encoding. Even though the encoding has a fixed length, it can represent variable-length architectures by means of identity cells. Note that the mapping from encoding to (computationally unique) architecture is redundant. The size of the search space (total number of encodings) is $3^{14} = 4,782,969$, which maps to 208,537 architectures. The latter number is sufficiently small to allow us to evaluate all architectures and save their performances to create the benchmark datasets.

For the large-scale experiment, we increase the number of variables $\ell$ to 17 by including the optimization of reduction cells, and also increasing the alphabet size $|\Omega_i|$ to 5, $\forall i$. Here, the position $i$ influences what $\Omega_i$ is used, as options for normal cells are different from options for reduction cells (see Fig. 1). The total search space of $5^{17} = 762,939,453,125$ maps to 104,086,030,125 architectures.

The encoding for NAS-Bench-101 \cite{33} (micro-level) is different. Here, both the type of operations in a (repeated) cell, and their connections, can be optimized. The encoding is a binary string of length 21 that represents connections, concatenated with a ternary string of length 5 that represents operations.

3.3 Search algorithms

The first algorithm we consider is NSGA-II \cite{5}. It is by far the most popular multi-objective EA in general. We use most commonly adopted parameter settings of NSGA-II: 2-point crossover, single-variable mutation with probability $p_c = 1/\ell$, tournament size 2 and population size $n_{\text{pop}} = 100$.

Fundamentally better algorithmic design is foundational to EA research. This means building better problem-specific EAs or ones capable of doing better at large classes of interesting problems. In the latter direction lie linkage learning EAs that attempt to automatically detect and exploit building blocks of non-trivial sizes \cite{18,28}. To the best of our knowledge, such algorithms have not yet been tried for NAS. Studying if they have merit here as well is interesting. Thus, we consider the Multi-Objective Gene-pool Optimal Mixing Evolutionary Algorithm (MO-GOMEA). MO-GOMEA mostly differs from a classic genetic algorithm in that every generation it computes a model of linkage, i.e. the estimated strength of interdependency between variables, and uses this model to propagate potential building blocks during variation \cite{28}. If the NAS search space exhibits any linkage, this algorithm can likely exploit it. Also, MO-GOMEA uses clusters to partition the population in objective space, so it is generally able to improve upon specific parts of the front by recombining within niches. We use one of the latest implementations for discrete optimization \cite{16}, which includes the Interleaved Multi-start Scheme (IMS). The IMS allows to avoid the manual
setting of population size parameter by evolving populations of increasing sizes in interleaved fashion. For the initial population size, we use $n_{\text{pop}} = 8$ (default).

We further consider RS as a standard baseline. RS generates new nets repeatedly by sampling each cell uniformly at random from the alphabets $\Omega_i$.

Finally, we consider a simple random restart LS algorithm:

1. Initialize a random architecture (as in RS);
2. Sample a scalarization coefficient $\alpha \sim U(0, 1)$;
3. Consider all variables in random order, as follows:
   3.1. For each variable $x_i$, evaluate the net obtained by setting $x_i$ to each option in $\Omega_i$. Keep the best according to $\alpha \times f_1 + (1 - \alpha) \times f_2$;
4. Repeat step 1 until the computation budget (evaluations / time) is exhausted.

In other words, we perform one round of first-improvement LS at the level of variables, but best-improvement LS at the level of options for one variable. Fig. 2 illustrates one iteration of our LS algorithm (from now on, we simply refer to it as LS).

4 Experimental setup

All search algorithms store an archive $\mathcal{A}$ of best-found nets according to (strict) Pareto domination. Formally, it is: $x \succ y$ (“$x$ dominates $y$”) iff $f_i(x) \geq f_i(y) \land f(x) \neq f(y)$, for each objective $f_i$. The archive thus satisfies $\nexists x, y \in \mathcal{A} : x \succ y$.

We use no size limitation for the archive, and update the archive each time a new net is discovered (the time cost is insignificant compared to evaluating a net’s performance, see below).

To compare algorithms, we consider the improvement of the hypervolume enclosed by $\mathcal{A}$, using the origin as reference point. We only count the evaluations of unique architectures as evaluating a net is typically costly (minutes to hours). Caching is used to not re-evaluate a same architecture twice.

We handle the net with identity cells for all variables differently from the others. This trivial net can be considered fundamentally uninteresting to search for, but it still influences multi-objective metrics. We therefore set NSGA-II and MO-GOMEA to include one instance of it in the population at initialization. Similarly, we include it in the archive of RS and LS from the beginning.

For experiments concerning the benchmark datasets, i.e., where performances for different architectures are pre-stored, we perform 30 runs for each algorithm. For the large-scale search space, where pre-evaluating all nets is simply unfeasible and a single evaluation takes on average 6.5 minutes on Nvidia RTX 2080 Ti GPU, we perform 6 runs for each algorithm.
5 Results

First, we compare the algorithms on NAS-Bench-101 and on our benchmarks. Next, we analyze the performance on our CIFAR-100 benchmark, and on the large-scale version for this task, in more detail. Lastly, we discuss what the observed differences mean when considering the nets’ ability to generalize.

5.1 Preliminary experiments

We begin by analyzing the results for NAS-Bench-101 (Fig. 3 left). For this benchmark, differences among the algorithms are negligible. This can also be seen in Fig. 4, where fronts obtained by the algorithms are similar at different points in time. These results are in line with the ones reported for the single-objective experiments conducted in \cite{33}: differences among fundamentally different algorithms are relatively small.

Regarding the macro-level search spaces, i.e., on our benchmarks on CIFAR-10 and CIFAR-100 (Fig. 3 middle and right, respectively), a notable difference is found between RS and the other algorithms. There are however also small differences between the EAs and LS. In the zoomed-in views it can be seen that, for the first 100 evaluations, NSGA-II performs as good as RS: this is because the initial population ($n_{pop} = 100$) is evaluated. Meanwhile, MO-GOMEA already performed optimization on the small initial populations instantiated by its IMS.

The hypervolume obtained for CIFAR-100 is not as good as the one obtained for CIFAR-10. This is not surprising because CIFAR-100 is considered a harder classification task. On our benchmarks, the best $acc_{val}$ for CIFAR-10 is 0.925, the one for CIFAR-100 is 0.705. Furthermore, note that the fact that the hypervolumes for NAS-Bench-101 reach better values, is mainly because $f_2$ is different here (based on number of net parameters instead of MMACs). Still, NAS-Bench-101 contains nets with slightly larger $acc_{val}$ (yet easily found by all algorithms), up to 0.951. Since CIFAR-100 is the harder task, we focus on this task in the next sections.

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Fig. 3: Convergence graphs on NAS-Bench-101 (left) and on our benchmarks for CIFAR-10 (middle) and CIFAR-100 (right). Medians (solid lines) and 25/75th percentiles (bands) of 30 runs are shown. Note that only the horizontal axes are the same, and are in logarithmic scale in the zoomed-in views (insets).
5.2 Benchmark: CIFAR-100

Fig. 3 (right) shows that, particularly for the EAs, the hypervolume increases rather quickly in the beginning, and not notably afterwards. Covering the entire Pareto front is however hard (e.g., only 21 out of 30 runs of MO-GOMEA cover it within 100,000 evals). Overall, MO-GOMEA performs best, followed by LS until \( \sim 700 \) evaluations. At this point, differences between MO-GOMEA, NSGA-II and LS are not significant at the 99% confidence level (Friedman test p-value \( = 0.05 > 0.01 \)). We remark that what these differences truly mean is discussed in terms of generalization in Sec. 5.4. RS is clearly the inferior approach.

To obtain a better understanding of the search the algorithms perform, we present the fronts discovered for CIFAR-100 at different time steps in Fig. 6 and Fig. 11 (left) (we found very similar results for the benchmark on CIFAR-10). The plots for 10 and 100 evaluations confirm the fact that MO-GOMEA and LS are the quickest at approaching the front. Note that the archive at 10 evaluations (Fig. 6 left) of LS is already obtained during the very first iteration of LS.

Compared to MO-GOMEA, LS is especially quick at finding the less-densely populated areas in the objective space (low MMACs). This is likely because making networks with less MMACs is straightforward, as more layers can be set to identity, which is effectively searched by the LS (with the proper objective weighting) that tries every option in each cell. Moreover, the EAs need to rely on the encodings present in the initial population, where an identity cell is sampled in position \( i \) only with chance \( 1/|\Omega_i| \). As shown in Fig. 3 evaluating small architectures is necessary to cover the part of the Pareto front containing efficient nets, as the number of nets there is smaller.

It is clear that RS is inferior to the other algorithms, as it struggles to find efficient nets that occur only sparsely in the search space. This issue is due to the fact that RS does not build up from efficient nets and thus solely relies on the way it samples: sampling a net with many identity cell is rare. For example, the probability of sampling the full-identity net is \( 1/|\Omega_i| \) (recall that this net is pre-inserted in the archive because uninteresting).
5.3 Large-scale NAS: CIFAR-100

Fig. 10 (right; consider only $acc_{val}$, i.e., solid lines) shows how the hypervolume improves over time for the large-scale experiment on CIFAR-100. MO-GOMEA and LS obtain a good hypervolume similarly quickly, as observed for the benchmark version. NSGA-II converges more slowly than MO-GOMEA and LS, but it is capable of outperforming them in terms of hypervolume towards the end. Nonetheless, both LS and MO-GOMEA do appear to improve again near the end of our total evaluation budget.

Fig. 7 shows the way the algorithms discover solutions over time, and Fig. 9 shows the density of architectures obtained at 2500 evaluations. Notably, MO-GOMEA discovers nets with the largest $acc_{val}$ (bottom-right points, consistent for all 6 runs). This may be because of two reasons. First, MO-GOMEA exploits linkage, which may be useful to recombine complex building blocks and obtain better performing nets. Second, MO-GOMEA restricts recombination to solutions that are similar.

Regarding RS, from Figs. 8, 9 it can be seen that RS mostly samples complex nets. Yet, as RS lacks any mechanism to refine and exploit solutions, it cannot match MO-GOMEA’s capability of discovering the most accurate nets. The main issue of RS is, as aforementioned, that it samples efficient nets too rarely.
NSGA-II exhibits a very interesting behavior: it searches progressively from more complex to simpler architectures (Figs. 7, 8). This happens because in the beginning it is more likely to generate complex nets (as RS does), while later on NSGA-II’s crowding distance operator steers the search towards the areas where points are most scattered, i.e., where efficient nets are located. Moreover, it is likely that its non-linkage informed operators are less likely to generate the best performing nets.

LS is found to be slightly inferior to the EAs in terms of final hypervolume (Fig. 10 right, solid lines), and it does not ultimately obtain the best front (Figs. 7 and 11 middle-right). Statistical testing (Friedman followed by post-hoc pairwise Mann-Whitney-U, 99% confidence interval) indicates that differences in hypervolume at the end of the runs (i.e., at 2500 evaluations) are significant. However, what we believe is most important is that LS is remarkably quick at obtaining fairly good nets (Fig. 7), with rather uniform spread in terms of trade-offs (Fig. 8). Moreover, LS has no issues similar to RS, in fact, it outperforms it markedly.

**Fig. 8:** MMACs of evaluated nets throughout one example run, smoothed for readability (moving avg. filter of size 75).

**Fig. 9:** Fronts obtained at 2500 evaluations by the search algorithms in one example run on the large-scale experiment on CIFAR-100, with a visual description of the nets. The colors of cells correspond to the legend in Figure 1. Heatmaps display the distribution of nets evaluated by the algorithms.

### 5.4 Generalization

Because the validation is used during the search, a risk exists to implicitly start overfitting to it. Hence, a second held-out set is used, i.e., given (a front of) best-acc_{val} nets, their acc_{test} is measured.

As can be seen from Figs. 10 (left) and 11 (left-most two), on our benchmark RS clearly remains the worst performing algorithm when considering generalization to the test set. However, MO-GOMEA and NSGA-II lose the lead to LS, between 100 and 18,000 evaluations. This is a consequence of the aforementioned phenomenon: despite the fact that nets are not trained on the set of data on which acc_{val} is measured, acc_{val} is used to select what nets are best, hence the EAs’ better search translates to more overfitting to this set. LS is slightly less
powerful at searching (worse hypervolume in terms of $acc_{\text{val}}$), yet the nets generalize well to the test set. Crucially, there are no significant differences among LS and the EAs at 100,000 evaluations (Friedman $p$-value = 0.55 > 0.01).

The findings for the enlarged search space reflect the ones found for the benchmark dataset: the difference in performance between LS and the EAs becomes smaller when considering $acc_{\text{test}}$ (see Fig. 10 right). Fig. 11 (right-most two) displays this especially well, as the most accurate nets discovered by MO-GOMEA turn out to be less accurate when tested again. Again, no statistical differences are observed at the end of the runs between LS and the EAs (Friedman test $p$-value = 0.85 > 0.01).

![Fig. 10: Convergence graphs regarding $acc_{\text{val}}$ (solid lines) and $acc_{\text{test}}$ (dashed lines) on CIFAR-100 on the benchmark (left) and the enlarged search space (right). Medians (lines) and 25/75th percentiles (bands) of respectively 30 and 6 runs are shown for all algorithms. Note the different ranges for the vertical axes and the logarithmic scale for the horizontal axis in the zoomed-in view.](image1)

![Fig. 11: Validation and test archives for 10,000 evaluations on benchmark NAS (resp. first and second from the left) and 2500 evaluations on large-scale NAS (resp. second and first from the right). Archives are from one example run.](image2)

6 Discussion

While several works compare algorithms in terms of time, we preferred a comparison in terms of number of evaluations. Using a time budget is reasonable
for practical applications, but has its limitations: particularly, time-based experiments are influenced by hyper-parameter choices (e.g., network training budgets), and can put some algorithms at disadvantage (e.g., see [20] for RS).

In our macro-level comparisons, we systematically found that RS performed much worse than the other algorithms. This boiled down to the fact that RS has a very low probability of sampling efficient nets. This simple result is important when considering RS as a baseline to benchmark against: if the sampling process of RS is not well-aligned with the desired objectives, RS can hardly beat an algorithm that includes optimization, even if as simple as LS. Note that on NAS-Bench-101, where the encoding and sampling process is such that small nets are not extremely rare, RS performed similar to the other algorithms. Carefully setting RS to sample solutions in a way that is well-aligned with the objectives may be non-trivial, and in fact harder than using LS.

We showed that the fact that different EAs include different search mechanisms, such as linkage learning and cluster-restricted mating for MO-GOMEA, and non-dominated sorting and crowding distance for NSGA-II, does lead to notable differences in what nets are found at what point in time. This is important to consider in practical applications where the evaluation budget is limited. LS was found to be competitive in this respect, as it quickly obtained a rather uniform scattering of nets with different trade-offs. Moreover, LS proved capable of ultimately competing very well when generalization is accounted for: the capability of SotA EAs to optimize at a very refined level with respect to $\text{acc}_{\text{val}}$ does not always translate to better $\text{acc}_{\text{test}}$ due to implicit overfitting.

7 Conclusion

With several experiments, we have shown that a simple, parameter-less local search algorithm can be a competitive baseline for NAS. We found that the proposed local search algorithm competes with state-of-the-art EAs, even up to thousands of evaluated architectures, in a multi-objective setting. By a greedy single-variable search mechanism and a random scalarization of the objectives, LS discovers nets with rather uniform trade-offs between accuracy and complexity in few evaluations. Performance gaps with EAs evident at search time are often lost when generalization is assessed. Importantly, we found that local search outperforms random search consistently and markedly. We therefore recommend to adopt local search as a baseline for NAS.

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Appendix

In this section we describe in detail the procedure we used to create the benchmark datasets (for CIFAR-10 and CIFAR-100), and the changes introduced for the large-scale NAS experiment (of Sec. 5.3).

Design of the benchmark dataset

We consider feed-forward CNNs, consisting of 17 sequentially connected cells. Similarly to [33], each net includes two auxiliary components: the stem convolution prior to the first cell, and a classifier after the last cell, which maps the output of the last cell to a prediction score for each class. The stem convolution is implemented by a $3 \times 3$ convolutional layer which converts the input image to 32 feature maps. The classifier is implemented by a $1 \times 1$ convolutional layer, followed by global average pooling and the final linear layer. Fig. 1 shows an example of our NAS setting.

Similarly to [33], we pre-set certain positions to ensure a feasible memory consumption for all nets. Specifically, cells at positions 5, 10 and 15 (reduction cells) reduce the spatial input dimensionality and increase the number of channels: for an input of size $D \times H \times W$ ($D$ is number of feature maps, $H$ and $W$ are the spatial dimensions) they return an output of size $2D \times H/2 \times W/2$. This is implemented by means of a $1 \times 1$ convolutional layer, followed by max-pooling. The remaining 14 positions are subjected to search with cells that preserve spatial dimensionality (i.e., normal cells).

As options for the searchable cells, we use differently parameterized inverted bottleneck convolutional blocks (MBConv) [22], which are widely used in NAS and known for their efficiency [3,9,27]. We allow a total of 3 cell types: MBConv with expansion factor 3 and kernel size 3; MBConv with expansion factor 6 and kernel size 5; and the identity cell. The latter option performs no operation and effectively acts as a placeholder to represent smaller nets, i.e., nets with fewer cells. If two nets are identical except for the position of identity cells in-between reduction cells (i.e. from positions 1 to 4, 6 to 9, and 11 to 14), they are computationally equivalent (an example is shown in Fig. 12).

To evaluate all nets in a feasible time, we relied on an approach that approximates training from scratch: the one-shot NAS [1,9]. The approach trains
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A supernet, i.e., a massive net that contains all possible nets of the search space as sub-nets. We follow the training procedure described in [9]: for each batch of training samples, one path, i.e., a sequence of cells that represents one of the possible architectures (see Fig. 13), is sampled from the supernet, and weights in the cells that belong to that path are trained. We use standard training settings: cross-entropy loss and stochastic gradient descent. We use stochastic weight averaging (SWA) [11] for robustness. For both CIFAR-10/100, we use a stratified data split as done in [33]: 40K samples for training, 10K for validation, 10K for testing. Further details on the training process are summarized in Table 1.

To evaluate a single architecture, we follow the approach also described in [9]: 1) Take the weights from the trained supernet corresponding to the architecture evaluated; 2) Recalculate parameters of all batch normalization layers using a subset of the training data (we use a fixed random stratified subset of 10K samples); 3) Calculate validation and test accuracies.

Fig. 13: Graphical description of the supernet with a randomly sampled path highlighted. In our case, the choices correspond to the cells of Fig. 12.

Large-scale experiment

In the large-scale experiment, we make both normal and reduction cells searchable. The options for normal cells are now 5, and consist of MBConv blocks with all four possible combinations of the aforementioned expansion factors and kernels sizes ($\{3, 6\} \times \{3, 5\}$), and the identity cell. We now further allow for the search of reduction cells, again with 5 options. These options are the same MB-Conv blocks but with stride 2 and number of channels multiplication, and the same simple reduction cell as used in our benchmarks. Recall that the positions of reduction cells are fixed (at indices 5, 10 and 15).

For this experiment we adopt the one-shot NAS approach again to speed-up the search, but include a refinement step, to better approximate the accuracy that nets would achieve if trained from scratch [1]. In particular, to evaluate an architecture, we finetune the weights obtained from the supernet for 20 epochs before calculating validation and test accuracies. Further hyperparameter settings are summarized in Table 1.
| Setup                  | Benchmarks creation | Large-scale experiment |
|------------------------|---------------------|------------------------|
| Hyperparameter         | Supernet training   | Supernet training      |
| Optimizer              | SGD with momentum 0.9, weight decay $10^{-4}$ | SGD with momentum 0.9, weight decay $10^{-4}$ | SGD with momentum 0.9, weight decay $10^{-4}$ |
| Initial learning rate  | 0.1                 | 0.1                    | 0.01                   |
| Learning rate schedule | Cyclic cosine annealing | Cyclic cosine annealing | Cosine annealing        |
| Using SWA              | ✔️                   | ✔️                     | ✗                      |
| Number of epochs       | 370                 | 500                    | 20                     |
| Batch size             | 128                 | 128                    | 256                    |

Table 1: Training hyperparameters