TensorFuzz: Debugging Neural Networks with Coverage-Guided Fuzzing

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Abstract

Machine learning models are notoriously difficult to interpret and debug. This is particularly true of neural networks. In this work, we introduce automated software testing techniques for neural networks that are well-suited to discovering errors which occur only for rare inputs. Specifically, we develop coverage-guided fuzzing (CGF) methods for neural networks. In CGF, random mutations of inputs to a neural network are guided by a coverage metric toward the goal of satisfying user-specified constraints. We describe how fast approximate nearest neighbor algorithms can provide this coverage metric. We then discuss the application of CGF to the following goals: finding numerical errors in trained neural networks, generating disagreements between neural networks and quantized versions of those networks, and surfacing undesirable behavior in character level language models. Finally, we release an open source library called TensorFuzz that implements the described techniques.

1 Introduction

Neural networks are gradually becoming used in more contexts that affect human lives, including for medical diagnosis [13], in autonomous vehicles [2, 7, 19], as input into corporate and judicial decision making processes [4, 37], in air traffic control [23], and in power grid control [40].

Neural networks have the potential to transform these applications, saving lives and offering benefits to more people than would be feasible using human labor. However, before this can be achieved, it is essential to ensure that neural networks are reliable when used in these contexts.

Machine learning models are notoriously difficult to debug or interpret [27] for a variety of reasons, ranging from the conceptual difficulty of specifying what the user wishes to know about the model in formal terms to statistical and computational difficulties in obtaining answers to formally specified questions. This property has arguably contributed to the recent ‘reproducibility crisis’ in machine learning [9, 16, 24, 28, 30, 33, 34] – it’s tricky to make robust experimental conclusions about techniques that are hard to debug. Neural networks can be particularly difficult to debug because even relatively straightforward formal questions about them can be computationally expensive to answer and because software implementations of neural networks can deviate significantly from theoretical models. For example, ReluPlex [23] can formally verify some properties of neural networks but is too computationally expensive to scale to model sizes used in practice. Moreover, ReluPlex works by analyzing the description of a ReLU network as a piecewise linear function, using a theoretical model in which all of the matrix multiplication operations are truly linear. In practice, matrix multiplication on a digital computer is not linear due to floating point arithmetic, and machine learning algorithms can learn to exploit this property to perform significantly nonlinear computation [10]. This is not to criticize ReluPlex but to illustrate the need for additional testing methodologies that interact directly with software as it actually exists in order to correctly test even software that deviates from theoretical models.

Preprint. Work in progress.
In this work, we leverage an existing technique from traditional software engineering—coverage-guided fuzzing (CGF) [39, 46]—and adapt it to be applicable to testing neural networks. In particular, this work makes the following contributions:

- We introduce the notion of CGF for neural networks and describe how fast approximate nearest neighbors algorithms can be used to check for coverage in a general way.
- We open source a software library for CGF called TensorFuzz.\(^1\)
- We use TensorFuzz to find numerical issues in trained neural networks, disagreements between neural networks and their quantized versions, and undesirable behaviors in character level language models.

Figure 1: Coarse descriptions of the main fuzzing loop. Left: A diagram of the fuzzing procedure, indicating the flow of data. Right: A description of the main loop of the fuzzing procedure in algorithmic form.

2 Background

This section gives background on coverage-guided fuzzing for traditional software, then discusses existing methods for testing neural networks before finally touching on how CGF relates to these methods.

2.1 Coverage-guided fuzzing

Coverage-guided fuzzing is used to find many serious bugs in real software [1]. Two of the most popular coverage-guided fuzzers for normal computer programs are AFL [46] and libFuzzer [39]. These have been expanded in various ways in order to make them faster or to increase the extent to which certain parts of the code can be targeted [5, 6].

In coverage-guided fuzzing, a fuzzing process maintains an input corpus containing inputs to the program under consideration. Random changes are made to those inputs according to some mutation procedure, and mutated inputs are kept in the corpus when they exercise new “coverage”. What is coverage? It depends on the type of fuzzer and on the goals at hand. One common measure is the set of parts of the code that have been executed. By this measure, if a new input causes the code to branch a different way at an if-statement than it has previously, coverage has increased.

CGF has been highly successful at identifying defects in traditional software, so it is natural to ask whether it could be applied to neural networks. Traditional coverage metrics track which lines of code have been executed and which branches have been taken. In their most basic forms, neural networks are implemented as a sequence of matrix multiplications followed by elementwise operations. The underlying software implementation of these operations may contain many branching statements but many of these are based on the size of the matrix and thus the architecture of the neural network.

\(^1\) To be released upon publication.
so the branching behavior is mostly independent of specific values of the neural network’s input. A neural network run on several different inputs will thus often execute the same lines of code and take the same branches, yet produce interesting variations in behavior due to changes in input and output values. Executing an existing CGF tool such as AFL thus may not find interesting behaviors of the neural network. In this work, we elect to use fast approximate nearest neighbor algorithms to determine if two sets of neural network ‘activations’ are meaningfully different from each other. This provides a coverage metric producing useful results for neural networks, even when the underlying software implementation of the neural network does not make use of many data-dependent branches.

2.2 Testing of Neural Networks

Methods for testing and computing test coverage of traditional computer programs cannot be straightforwardly applied to neural networks. We can’t just naively compute branch coverage, for example, because of reasons discussed above. Thus, we have to think about how to write down useful coverage metrics for neural networks. Though this work is the first (as far as we know) to explore the idea of CGF for neural networks, it’s not the first to address the issues of testing and test coverage for neural networks. A variety of proposals (many of which focus on adversarial examples [43]) have been made for ways to test neural networks and to measure their test coverage. We survey these proposals here:

Pei et al. [35] introduce the metric of neuron coverage for a neural network with rectified linear units (ReLus) as the activation functions. A test suite is said to achieve full coverage under this metric if for every hidden unit in the neural network, there is some input for which that hidden unit has positive value. They then cross reference multiple neural networks using gradient based optimization to find misbehavior.

Ma et al. [29] generalize neuron coverage in two ways. In k-multisection coverage, they take – for each neuron – the range of values seen during training, divide it into k chunks, and measure whether each of the k chunks has been “touched”. In neuron boundary coverage, they measure whether each activation has been made to go above and below a certain bound. They then evaluate how well these metrics are satisfied by the test set.

Sun et al. [41] introduce a family of metrics inspired by Modified Condition / Decision Coverage [14]. We describe their ss-coverage proposal as an example: Given a neural network arranged into layers, a pair of neurons \( (n_1, n_2) \) in adjacent layers is said to be ss-covered by a pair \( (x, y) \) of inputs if the following 3 things are true: \( n_1 \) has a different sign for each of \( x, y \), \( n_2 \) has a different sign for each of \( x, y \), and all other elements of the layer containing \( n_1 \) have the same sign for \( x, y \).

Tian et al. [44] applies the neuron coverage metric to deep neural networks that are part of self-driving car software. They perform natural image transformations such as blurring, shearing, and transformation and use the idea of metamorphic testing [8] to find errors.

Wicker et al. [45] perform black box testing of image classifiers using image-specific operations. Concurrent with this work, Sun et al. [42] leverages a complementary approach called concolic execution. Whereas our approach is analogous to AFL or libFuzzer, their approach is analogous to CUTE [38].

2.3 Opportunities for improvement

It is heartening that so much progress has been made recently on the problem of testing neural networks. However, the success of e.g. AFL and libFuzzer in spite of the existence of more sophisticated techniques suggests that there is a role for an analogous tool that works on neural networks. Ideally we would implement CGF for neural networks using the coverage metrics from above. However, all of these metrics, though perhaps appropriate in the context originally proposed, lack certain desirable qualities. We describe below why this is true for the most relevant metrics.

Sun et al. [41] claim that the neuron coverage metric is too easy to satisfy. In particular, they show that 25 randomly selected images from the MNIST test set yield close to 100% neuron coverage for an MNIST classifier. This metric is also specialized to work on rectified linear units (reLus) which limits its generality.
Neuron boundary coverage [29] is nice in that it doesn’t rely on using reLus, but it also still treats neurons independently. This causes it to suffer from the same problem as neuron coverage: it’s easy to exercise all of the coverage with few examples.

The metrics from Sun et al. [41] are an improvement on neuron coverage and may be useful in the context of more formal methods, but for our desired application, they have several disadvantages. They still treat reLus as a special case, they require special modification to work with convolutional neural networks, and they do not offer an obvious generalization that supports attention [3] or residual networks [15]. They also rely on neural networks being arranged in hierarchical layers, which is often not true for modern deep learning architectures.

What we would really like is a coverage metric that is simple, cheap to compute, and is easily applied to all sorts of neural network architectures. Thus, we propose storing the activations (or some subset of them) associated with each input, and checking whether coverage has increased on a given input by using an approximate nearest neighbors algorithm to see whether there are any other sets of activations within a pre-specified distance. We discuss this idea in more detail in Section 3.2.

3 The TensorFuzz library

Drawing inspiration from the fuzzers described in the previous section, we have implemented a tool that we call TensorFuzz. It works in a way that is analogous to those tools, but that is different in ways that make it more suitable for neural network testing. Instead of an arbitrary computer program written in C or C++, it feeds inputs to an arbitrary TensorFlow graph. Instead of measuring coverage by looking at basic blocks or changes in control flow, it measures coverage by (roughly speaking) looking at the “activations” of the computation graph. In 3.1, we discuss the overall architecture of the fuzzer, including the flow of data and the basic building blocks. In 3.2, we discuss building blocks in more detail. In particular, we describe how the corpus is sampled from, how mutations are performed, and how coverage and objective functions are evaluated. We pay extra attention to describing special challenges associated with defining a coverage metric for neural networks and explain how these challenges can be (at least partially) dealt with using approximate nearest neighbors.

3.1 The basic fuzzing procedure

The overall structure of the fuzzing procedure is very similar to the structure of coverage-guided fuzzers for normal computer programs. The main difference is that instead of interacting with an arbitrary computer program that we have instrumented, we interact with a TensorFlow graph that we can feed inputs to and get outputs from.

The fuzzer starts with a seed corpus containing at least one set of inputs for the computation graph. Unlike in traditional CGF, we don’t just feed in big arrays of bytes. Instead, we restrict the inputs to those that are in some sense valid neural network inputs. If the inputs are images, we restrict our inputs to have the correct size and shape, and to lie in the same interval as the input pixels of the dataset under consideration. If the inputs are sequences of characters, we only allow characters that are in the vocabulary extracted from the training set.

Given this seed corpus, fuzzing proceeds as follows: Until instructed to stop, the fuzzer chooses elements from the input corpus according to some component we will call the Input Chooser. For the purpose of this section, it’s ok to imagine the Input Chooser as choosing uniformly at random, though we will describe more complicated strategies in 3.2.

Given an input, the Mutator component will perform some sort of modification to that input. The modification can be as simple as just flipping the sign of an input pixel in an image, and it can also be restricted to follow some kind of constraint on the total modification made to a corpus element over time - see 3.2 for more on this.

Finally, the mutated inputs can be fed to the neural network. In TensorFuzz, two things are extracted from the neural network: a set of coverage arrays, from which the actual coverage will be computed, and a set of metadata arrays, from which the result of the objective function will be computed.

Once the coverage is computed, the mutated input will be added to the corpus if it exercises new coverage, and it will be added to the list of test cases if it causes the objective function the be satisfied.
See Figure 1 for complementary overviews of this procedure.

### 3.2 Details of the fuzzing procedure

In this section we describe in more detail the components of the fuzzer.

**Input Chooser:** At any given time, the fuzzer must choose which inputs from the existing corpus to mutate. The optimal choice will of course be problem dependent, and traditional CGFs rely on a variety of heuristics to make this determination. For the applications we tested, making a uniform random selection worked acceptably well, but we ultimately settled on the following heuristic, which we found to be faster: 

\[ p(c_k, t) = \frac{e^{t_k - t}}{\sum_k e^{t_k - t}}, \]

where \( p(c_k, t) \) gives the probability of choosing corpus element \( c_k \) at time \( t \) and \( t_k \) is the time when element \( c_k \) was added to the corpus. The intuition behind this is that recently sampled inputs are more likely to yield useful new coverage when mutated, but that this advantage decays as time progresses.

**Mutator:** Once the Input Chooser has chosen an element of the corpus to mutate, the mutations need to be applied. In this work, we had to implement mutations for both image inputs and text inputs. For image inputs, we implemented two different types of mutation. The first is to just add white noise of a user-configurable variance to the input. The second is to add white noise, but to constrain the difference between the mutated element and the original element from which it is descended to have a user-configurable \( L_\infty \) norm. This type of constrained mutation can be useful if we want to find inputs that satisfy some objective function, but are still plausibly of the same “class” as the original input that was used as a seed. In both types of image mutation we clip the image after mutation so that it lies in the same range as the inputs used to train the neural network being fuzzed.

For text inputs, since we can’t simply add uniform noise to the string, we mutate according to the following policy: we uniformly at random perform one of these operations: delete a character at a random location, add a random character at a random location, or substitute a random character at a random location.

**Objective Function:** Generally we will be running the fuzzer with some goal in mind. That is, we will want the neural network to reach some particular state - maybe a state that we regard as erroneous. The objective function is used to assess whether that state has been reached. When the mutated inputs are fed into the computation graph, both coverage arrays and metadata arrays are returned as output. The objective function is applied to the metadata arrays, and flags inputs that caused the objective to be satisfied.

**Coverage Analyzer:** The coverage analyzer is in charge of reading arrays from the TensorFlow runtime, turning them into python objects representing coverage, and checking whether that coverage is new. The algorithm by which new coverage is checked is central to the proper functioning of the fuzzer.

The characteristics of a desirable coverage checker are: we want it to check if the neural network is in a ‘state’ that it hasn’t been in before, so that we can find misbehaviors that might not be caught by the test set. We want this check to be fast (so we probably want it to be simple), so that we can find many of those misbehaviors quickly. We want it to work for many different types of computation graph without special engineering, so that practitioners can use our tooling without having to make special adaptations. We want exercising all of the coverage to be hard, otherwise we won’t actually cover very much of the possible behaviors. Finally, we want getting new coverage to help us make incremental progress, so that continued fuzzing yields continued gains.

As alluded to in Section 2.3, none the coverage metrics discussed in Section 2.2 quite meet all these desiderata, but we can design from first principles a coverage metric that at least comes closer to meeting them.

A naive, brute force solution is to read out the whole activation vector and treat new activation vectors as new coverage. However, such a coverage metric would not provide useful guidance, because most inputs would trivially yield new coverage. It is better to detect whether an activation vector is close to one that was observed previously. One way to achieve this is to use an approximate nearest neighbors algorithm, (as used for Neural Turing Machine [12] memory by Rae et al. [36]). When we get a new
activation vector, we can look up its nearest neighbor, then check how far away the nearest neighbor is in Euclidean distance and add the input to the corpus if the distance is greater than some amount $L$.

This is essentially what we do. Currently, we use an open source library called FLANN [31] to compute the approximate nearest neighbors. In general, you may not need to see all the activations. The set of activations to use is presently a matter for empirical tuning. We find it is often possible to obtain good results by tracking only the logits, or the layer before the logits.

One potential optimization to note: you don’t actually need to know the nearest neighbor, you just need to know whether there exists a neighbor within some range. For this, you can use a distance-sensitive Bloom filter [25], but this will come at the expense of sometimes counting coverage as ‘not new’ when it is new. If your fuzzing application is one in which this is acceptable, this could speed things up for you. However, we leave the testing of this idea to future work.

3.3 Batching and nondeterminism

There are two other issues that are somewhat unique to TensorFlow graphs that merit a few words. First, nearly all TensorFlow graphs that exist in practice have been engineered to take advantage of hardware parallelism offered by e.g. modern GPUs. For this reason, it could be wasteful to only fetch the coverage and metadata for one mutated input at a time. Instead, we perform the mutations as a batch and feed a batch of inputs to the computation graph, then we check the coverage and objective function on a batch of output arrays. Second, computation graphs will often give nondeterministic outputs - both because of instructions that execute nondeterministically (e.g. large accumulations on GPUs) and because of fundamentally random operations built into the graph. For now, we have chosen to deal with this in the most naive way possible: if we can feed the same input in twice and get different coverage, we simply include the input twice in the corpus.

4 Experimental results

We briefly present a few different applications of the CGF technique to establish that it is useful in general settings.

4.1 CGF can efficiently find numerical errors in trained neural networks

Since neural networks use floating point math [11], they are susceptible to numerical issues, both during training and at evaluation time. These issues are notoriously hard to debug, partly because they may only be triggered by a small set of rarely encountered inputs. This is one case where CGF can help. We focus on finding inputs that result in not-a-number (NaN) values.

Numerical errors are important to find: Numerical errors, especially those resulting in NaNs, could cause dangerous behavior of important systems if these errors are first encountered ‘in the wild’. CGF can be used to identify a large number of errors before deployment, and reduce the risk of errors occurring in a harmful setting.

CGF can quickly find numerical errors: With CGF, we should be able to simply add check numerics ops to the metadata and run our fuzzer. To test this hypothesis, we trained a fully connected neural network to classify MNIST [26] digits. We used a poorly implemented cross entropy loss on purpose so that there would be some chance of numerical errors. We trained the model for 35000 steps with a mini-batch size of 100, at which point it had a validation accuracy of 98%. We then checked that there were no elements in the MNIST dataset that cause a numerical error. Nevertheless, TensorFuzz found NaNs quickly across multiple random initializations. See Figure 2 for more details.

Gradient-based search techniques might not help find numerical errors: One potential objection to CGF techniques is that gradient-based search techniques might be more efficient than more

\[2\text{ In fact, they’ve often been engineered to run on multiple GPUs, which suggests an obvious optimization to TensorFuzz: unlike stochastic gradient descent, in which parallelism is limited by stale gradients [32], fuzzing performance can theoretically scale linearly with hardware, assuming that the corpus does not need to be kept perfectly synchronized.}\]
Figure 2: We trained an MNIST classifier with some unsafe numerical operations. We then ran the fuzzer 10 times on random seeds from the MNIST dataset. The fuzzer found a non-finite element every run. Random search never found a non-finite element. Left: the accumulated corpus size of the fuzzer while running, for 10 runs. Right: an example satisfying image found by the fuzzer.

Random search is prohibitively inefficient for finding some numerical errors: To establish that random search is not sufficient and that coverage-guidance is necessary for efficiency, we compared to random search. We implemented a baseline random search algorithm and ran it for 100,000 samples from the corpus with 10 different random initializations. The baseline was not able to find a non-finite element in any of these trials.

4.2 CGF surfaces disagreements between models and their quantized versions

Quantization [18] is a process by which neural network weights are stored and neural network computations performed using numerical representations that consist of fewer bits of computer memory. Quantization is a popular approach to reducing the computational cost or size of neural networks, and is widely used for e.g. running inference on cell phones as in Android Neural Networks API or TFLite and in the context of custom machine learning hardware – e.g. Google’s Tensor Processing Unit [20] or NVidia’s TensorRT.

Errors resulting from quantization are important to find: Of course, quantization is not very useful if it reduces the accuracy of the model too dramatically. Given a quantized model, it would thus be nice to check how much quantization reduced the accuracy.

Few errors can be found just by checking existing data: As a baseline experiment, we trained an MNIST classifier (this time without intentionally introducing numerical issues) using 32-bit floating point numbers. We then truncated all weights and activations to 16-bits. We then compared the predictions of the 32-bit and the 16-bit model on the MNIST test set and found 0 disagreements.

CGF can quickly find many errors in small regions around the data: We then ran the fuzzer with mutations restricted to lie in a radius 0.4 infinity norm ball surrounding the seed images, using the activations of only the 32-bit model as coverage. We restrict to inputs near seed images because these inputs nearly all have unambiguous class semantics. It is less interesting if two versions of the model disagree on out-of-domain garbage data with no true class. With these settings, the fuzzer was able to generate disagreements for 70% of the examples we tried. Thus, CGF allowed us to find real errors that could have occurred at test time. See Figure 3 for more details.

Random search fails to find new errors given the same number of mutations as CGF: As in Section 4.1 we tried a baseline random search method in order to demonstrate that the coverage-
Figure 3: We trained an MNIST classifier with 32-bit floats and then truncated the associated TensorFlow graph to 16-bit floats. Both the original and the truncated graph made the same predictions on all 10000 elements of the MNIST test set, but the fuzzer was able to find disagreements within an infinity-norm ball of radius 0.4 around 70% of the test images that we tried to fuzz. Left: the accumulated corpus size of the fuzzer while running, for 10 runs. Lines that go all the way to the right correspond to failed fuzzing runs. Right: an image found by the fuzzer that is classified differently by the 32-bit and 16-bit neural networks.

guidance specifically was useful in this context. The random search baseline was not able to find any disagreements that didn’t already exist when given the same number of mutations as the fuzzer.

4.3 CGF surfaces undesirable behavior in character level language models

Given a trained machine learning model, we may be able to characterize certain of its behaviors as undesirable, even though incorporating this characterization into the training loss might be inconvenient or even intractable. As an example, we train a character level language model of the type described in Karpathy [21]. In particular, we modify the code from https://github.com/sherjilozair/char-rnn-tensorflow to train a 2 layer LSTM [17] on the Tiny Shakespeare [22] dataset.

We then consider the application of sampling from this trained language model given a priming string. One can imagine such a thing being done in an auto-complete application, for example. For illustrative purposes, we identify two desiderata that we can approximately enforce via the fuzzer: First, the model should not repeat the same word too many times in a row. Second, the model should not output words from the blacklist.

We fuzz the model using for coverage the hidden state of the LSTM after the priming string has been consumed. We sample from the model according to a softmax over the logits, using a fixed random seed that we reset at every sampling. We use the mutation function described in Section 3.2.

We ran an instance of TensorFuzz and an instance of random search for 24 hours each. TensorFuzz was able to generate repeat words, but so was random search. However, TensorFuzz was able to generate six out of ten words from our blacklist while random search only generated one.

5 Conclusion

We have introduced the concept of coverage-guided fuzzing for neural networks and described how to build a useful coverage checker in this context. We have demonstrated the practical applicability of TensorFuzz by finding numerical errors, finding disagreements between neural networks and their quantized versions, and surfacing undesirable behavior in RNNs. Finally, we are releasing along with this paper the implementation of TensorFuzz, so that other people can both build on our work and also use our fuzzer to find real issues.
Acknowledgments

We thank Dave Anderson for helpful comments on an early version of the draft. We thank Rishabh Singh, Alexey Kurakin, and Martín Abadi for general input. We also thank Kostya Serebryany for helpful explanations of libFuzzer.
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