Automated fault detection without seismic processing

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Automated fault detection without seismic processing

Mauricio Araya-Polo1, Taylor Dahlke1,2, Charlie Frogner3, Chiyuan Zhang3, Tomaso Poggio3, and Detlef Hohl1

Abstract

For hydrocarbon exploration, large volumes of data are acquired and used in physical modeling-based workflows to identify geologic features of interest such as fault networks, salt bodies, or, in general, elements of petroleum systems. The adjoint modeling step, which transforms the data into the model space, and subsequent interpretation can be very expensive, both in terms of computing resources and domain-expert time. We propose and implement a unique approach that bypasses these demanding steps, directly assisting interpretation. We do this by training a deep neural network to learn a mapping relationship between the data space and the final output (particularly, spatial points indicating fault presence). The key to obtaining accurate predictions is the use of the Wasserstein loss function, which properly handles the structured output — in our case, by exploiting fault surface continuity. The promising results shown here for synthetic data demonstrate a new way of using seismic data and suggest more direct methods to identify key elements in the subsurface.

Introduction

Acquisition technology advances and exploration of complex areas are pushing the amount of data to be analyzed into the “big data” category. Current exploration workflows consist of many partially automated steps in which domain experts (geologists, geophysicists, rock physicists, etc.) command highly tuned applications and then curate the resulting data in search of valuable information. The data explosion is stressing these workflows to a point at which every year more of the data remains unused. Generally, the exploration process can be broken down into two main elements: advanced tools and manpower. Tools have progressed, and the addition of high-performance computing has helped to reduce turnaround times for seismic imaging (Rubio et al., 2009; Rastogi, 2011). Even in the extreme case in which execution time for processing tools would take nearly zero time, the problem of manpower remains; there is no sensible way in which domain experts can analyze and interpret all incoming data. The best solution must trade domain-expert time for computing time. Therefore, some of that domain knowledge needs to be formalized and implemented within existing and future tools. One way to achieve this is by taking advantage of algorithms that learn, for instance, from legacy data that have been properly vetted. Using machine learning, we can take advantage of new algorithms and software ecosystems, as well as specialized hardware. In this contribution, we will focus our attention on one such application of machine learning.

Seismic imaging is the primary tool used to build high-resolution models of the subsurface. In practice, it is typically part of an iterative workflow that alternates between imaging steps and model update steps. This iterative refinement is expensive in terms of human costs (picking out features/adjusting the velocity model) and computational costs (many repeated wave propagations and attributes computations). Most prior work focused on identifying features in already migrated images (Hale, 2012; Hale, 2013; Guillen, 2015; Addison, 2016; Bougher and Hermann, 2016). The literature is filled with refinements to this workflow, but ultimately, it remains largely the same.

Recent work (Zhang et al., 2014; Frogner et al., 2015; Dahlke et al., 2016) demonstrates a new approach that builds and uses a deep neural network (DNN) statistical model to transform raw-input seismic data directly to the final mapping of faults in 2D. In this case, fault locations were chosen as the output because of their relevance to optimizing production in existing fields. DNNs are built on the premise that they can be used to replicate any function (in theory, even a nonlinear one like acoustic wave propagation). We show that DNNs can be used to identify fault structure in 3D volumes with reasonable accuracy. The greater promise is that as computational tools improve, we can use even more complex neural networks to improve accuracy. The costs are all computational, mostly in the form of training incurred only once up front. Once the neural network is trained, predictions can be produced in a fraction of the training time and of the time needed for traditional physics modeling with partial differential equations. This means that nearly instantaneous earth models could be created as acquisition progresses. An interpreter aboard an acquisition vessel could identify hot spots as they travel over them and then modify acquisition to improve imaging over that same area on another pass. This is more cost effective than initiating a new acquisition campaign.

In this paper, we start by explaining the basics of the problem followed by discussions of DNNs and the Wasserstein loss function. Next, we introduce the general workflow used to train our DNN. After that, we discuss our performance results and some of our findings regarding network architectural parameters. Last, we will explain the directions for future work.

Problem overview

The key geologic feature we target in this work is a fault network. Shifted rock layers can trap liquid hydrocarbons, which will form reservoirs (Figure 1). The 3D structure of a fault network can be highly complex (Figure 2), representing a challenge for structured output methods of prediction.

Deep neural networks (DNN)

Using learning algorithms is an appealing alternative to classic seismic processing. Among that class of algorithms, we have
selected DNN for our experiments since there is increasing community agreement (Grzeszczuk et al., 1998; De et al., 2011; Lin and Tegmark, 2016) in favor of this kind of algorithm as a good surrogate for physics-based processes.

DNNs consist of “layers” of weighted nodes that are activated by inputs from previous layers. These networks are trained with examples in which the correct/true output (label) is known for a given input; the weight parameters in the nodes of the network evolve due to the minimization of the error between prediction and true value. This causes the network to become an increasingly better predictor of the training examples and, ultimately, of any example (assuming proper training) of a data class that is similar in nature to the training data.

This topology of information flow between nodes is inspired by the structure of neurons in the brain (Figure 3). The universal approximation theorem shows that neural networks can be used to approximate an arbitrary continuous function up to our desired accuracy (Hornik et al., 1989). For this reason, there is much promise in using this approach to make predictions of functions that are very nonlinear (like fault probability from time-series data). While the theoretical limits of this approach are generous, to find the neural network that best approximates our function is not trivial. Our ability to build effective neural networks is limited by computing constraints. More complex networks are more expensive to train, and the generation of accurate training examples can be computationally expensive due to large-scale forward modeling. Ultimately, our predictions are only as good as the complexity and refinement of our neural network coupled with the relevance and quality of the features we choose as inputs.

**Workflow**

The first step in our workflow (Figure 4) is to collect the training examples. Real data examples are impractical and limiting since the labels are assigned to fault locations by few domain experts. This means that the neural network’s best result would be bound by human performance and data quality. Instead, we generate realistic 3D velocity models synthetically, with the fault labeling generated concurrently for an unbiased ground truth. Next, we use an acoustic approximation to the wave equation to generate wavefields and record them as time-series signals with predefined acquisition geometry. This step is conducted on thousands of random velocity-model realizations, giving us many instances of labeled fault locations and the corresponding seismic traces for the entire data set. We reserve a portion of this data set unseen from the algorithm (holdout set) so that we can test it after we have trained our predictor. We set ranges on a relatively small number of parameters as bounds on the random model generator. These parameters include the number of layers in a model, the number of faults, the range of velocity, and the dip and strike angles for each possible fault. We believe that the randomized models produced in this manner are realistic enough to demonstrate the efficacy of neural network predictions (see Figure 5).

**Labeling**

Since the positions of the faults are known a priori, the labeling process is straightforward. However, complexity is added to this process when the labeling targets a subsampled grid for computational efficiency. Addition of a threshold parameter to define the labeling of the coarse grid is needed. This threshold value sets how many fine-sampled voxels inside a coarse voxel must be fault-labeled for that coarse voxel to be considered as having a fault or not. There are a variety of ways to choose that threshold.
so that the subsampled labeling appropriately matches the underlying fine-sampled labeling, with ours being chosen on a trial-and-error basis prior to any training.

**Structured output learning and the Wasserstein loss**

Our problem is naturally formulated as predicting a (subsampling) 3D “voxel map” of binary fault/nonfault indicators. This is similar to image segmentation problems in computer vision. A common way of handling this is to introduce a Markov random field (MRF) on the output voxels that captures the interactions and run an inference algorithm to get the jointly optimal predictions. More generally, the inference step can be incorporated into the objective function for learning. Commonly used formulations include structured support vector machines and conditional random fields (Nowozin and Lampert, 2011).

However, while the couplings for pixel labels in an image can be modeled naturally via neighborhood similarity or input-pixel-based similarity, the prior structure in our model is of much higher complexity. More specifically, the faults usually extend as smooth surfaces. This property cannot be characterized via factors that involve only a few nearby output variables. On the other hand, inference on MRFs with general high-order factors is computationally expensive. As a result, we choose to perform independent predictions for each output region and incorporate our prior knowledge via a novel loss function called the Wasserstein loss (Frogner et al., 2015).

Formally, let \( K = D \times D \times D \) be the number of output voxels. We normalize the ground-truth binary output vector \( y \in \{0, 1\}^K \) to \( y = \frac{y}{\|y\|} \), so that it represents a probability distribution over the 3D grid. Moreover, we model our predictor as a DNN with a softmax layer at the top so that it also produces a probability distribution \( \hat{y} \) on the output space. For our application, the output space is the 3D grid of voxels, and the natural ground metric is the Euclidean distance between the voxels. \( T \) in the loss term is a joint probability distribution that marginalizes to the ground truth and the prediction. Intuitively, \( T \) defines a transportation plan that maps probability mass from the prediction to the ground-truth, and \( \langle T, M \rangle \) measures the cost of this plan according to the ground metric. The loss is then defined by the cost of the optimal feasible transport plan. For the cases of Figure 7, the Wasserstein loss for the left plot will be smaller than the central plot due to the larger cost of this plan according to the ground metric.

\[
\ell_w(\hat{y}, y) = \min_{T \in \mathcal{H}(y)} \langle T, M \rangle,
\]

\[
\Pi(\hat{y}, y) = \left\{ T \in \mathbb{R}^{K \times K} : T1 = \hat{y}, T^T1 = y \right\},
\]

where \( \langle T, M \rangle = \text{tr}(T^T M) \) is the inner product, for a given ground metric matrix \( M_{k,k'} = d(k, k') \) for some ground metric \( d(y) \) on the output space. For our application, the output space is the 3D grid of voxels, and the natural ground metric is the Euclidean distance between the voxels. \( T \) in the loss term is a joint probability distribution that marginalizes to the ground truth and the prediction. Intuitively, \( T \) defines a transportation plan that maps probability mass from the prediction to the ground-truth, and \( \langle T, M \rangle \) measures the cost of this plan according to the ground metric. The loss is then defined by the cost of the optimal feasible transport plan. For the cases of Figure 7, the Wasserstein loss for the left plot will be smaller than the central plot due to the larger cost of this plan according to the ground metric.

**Feature generation and extraction**

Raw seismic data is far too unrefined and redundant to be immediately useful as inputs to our neural network. We improved prediction performance by extracting input features carefully, taking advantage of techniques from signal processing. The amount of collected features is large and grows by orders of magnitude when more realistic models are used. For instance, for all cases used in this work, the acquisition configuration is: nine shots per model, with 400 receivers in a 20 × 20 grid, and 1500 time samples per receiver. This configuration (smaller than realistic acquisitions) produces 5,400,000 raw features in total per model. If we were to use a more realistic acquisition, such as 100 shots with 1600 receivers and 2000 time samples, we would end up with up to 320,000,000 input attributes. These numbers only grow with denser acquisition geometries.
Choosing the architectural parameters of a neural network is not a deterministic process. A neural network that learns from this amount of features requires a large number of training epochs to train.

We use transformations of the data space coupled with feature-reduction methods for removing the features that are the least useful for prediction. These steps reduce the feature space to a more reasonable magnitude (thousands of input features rather than millions).

Implementation and results

In terms of computing, the training process takes a few hours per data set and minutes for testing. Putting this in perspective, the training process for one data set takes the same time that the migration of a single shot takes for a dense acquisition with anisotropic assumption. Once the network is trained, it can be reused many times with minimal cost, and given a certain generalization level, it can be exposed to many new data sets. This can be far less expensive than processing and interpreting one complete seismic acquisition, which happens every time the underlying model is changed. Our proprietary workflow is almost completely implemented in the MIT Julia language in which the heavy computing is offloaded to GPGPUs through NVIDIA’s CuDNN.

We generate thousands of random velocity models with up to four faults in them, of varying strike, dip angle, and position. Our models had between three and six layers each, with velocities varying from 2000 to 4000 [m/s], with layer velocity increasing with depth. These models were 140 × 180 × 180 grid points at the sampling used for wave propagation (using the acquisition geometry described earlier) but were subsampled to 20 × 20 × 20 and 32 × 32 × 32 for labeling purposes. The raw data collected was reduced aggressively to a feature set that can fit in an NVIDIA K80 GPGPU memory.

With the generated features and labels, a variety of fully connected deep neural networks are trained. The network architecture main parameters varied from two to 20 hidden layers and 256 to 2048 units per layer. For all cases presented in Table 1, we used the Wasserstein loss function for training. The output of the networks is a subsampled 3D voxel grid, with each voxel’s value indicating the likelihood of a fault being present within the voxel. Ground-truth labels on the same grid were binary valued, indicating presence or not of a fault in each voxel. The final predictions were generated by taking the likelihood values map output and applying a threshold to it, such that likelihood values above the threshold would be considered having a fault, while those below would not. As a result, a lower threshold will label more voxels as faults, while a high threshold labels less.

We use two different quantitative metrics of performance in this work: intersection over union (IoU) and area under the ROC curve (AUC).

The IoU value is a ratio of the number of voxels that are in the intersection of the ground truth and prediction, divided by the number of voxels in the union of the ground truth and prediction. This gives us an idea of how clustered or scattered a prediction is; the values range from 0 to 1, where higher values are better (example in Figure 6). Two predictions could theoretically have the same AUC value (see Table 1) but

![Figure 6. Comparison of (a) the Wasserstein- and (b) non-Wasserstein-based predictions, IoU metric (described in the article). Red areas show false positives, green shows true positives (correct predictions), and yellow shows false negative. (c) 2D slice of a 3D model. The predictions have very different IoU, where green means better.](image)

![Figure 7. ROC curves for representative DNN architectures. The closer to the top-left corner the better.](image)

<table>
<thead>
<tr>
<th>AUC</th>
<th>IoU</th>
<th>Hidden layers</th>
<th>Nodes per layer</th>
<th>Faults per model</th>
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<td>4</td>
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<td>4</td>
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</table>

Table 1. Results obtained on several representative sets of simulated test data. The first two columns report performance metrics; the other columns describe the parameter of the experiments. All results obtained using Wasserstein loss function, with 16,000 training models and 4000 testing models, except for the asterisked case, where only 10,000 training models and 2000 testing models were used.
different IoU values. We report the (IoU) value averaged for the entire test set of predictions, with predicted likelihoods thresholded at a value chosen to maximize the average IoU over all the predictions.

Each point of a receiver operating characteristics (ROC) curve (see Figure 7) is based on the number of true positive predictions (vertical axis) and the false positive predictions (horizontal axis) for a particular threshold value. We report the AUC for the predictions, which describes how strong our predictor is. The value ranges from 0.5 to 1.0, where the higher the value the better.

In Table 1, for all data sets (one, two, and four faults in model) AUC exceeds 0.9, which approaches that needed for practical use. Also, IoU surpasses 0.3 for many experiments, which implies that the prediction can be improved in terms of spatial alignment with respect to the ground truth.

The prediction grid size of experiments in Figure 9 and Figure 11 is 32 × 32 × 32, therefore each prediction represents a voxel of 4 × 5 × 5 in the model space. The predictions in Figure 9 follow the expected results, plus some false positives in the bottom-right area of the back slice.

The predictions in Figure 11b follow the expected structure as can be seen in Figure 11a. However, false positives are present in the area where the faults coincide. This is in line with expectations, since a cornucopia of signals and patterns are produced in that area. This case exposes the current limit of our predicting resolution for complex cases. In general, the resolution is limited by the quality of the data.

**Future work**

To evaluate practical usability, we must address how this approach can be scaled to actual production-level seismic data sets. For instance, we have to analyze sensitivity of the predictions to acquisition geometries. Current synthetic input data is based on fixed acquisition geometry. If we use this fixed geometry to train a predictor, one must bin and stack the acquired data so that it matches the acquisition used for training the model. For this reason, we believe that using a fixed dense geometry permits us to accurately bin and stack real data to match the predictor’s input parameters. Denser acquisition means more features, and as a result, more dramatic reduction of the feature space (and/or more complex neural networks) is needed.

Another area where we can improve the prediction quality is by increasing the number of voxels in our downsampled output grid such that we have better resolution in our predictions. This increase in voxel granularity means a more computationally demanding neural network and thus an increase in the cost of tuning and training.

Our workflow is flexible with respect to what can be predicted. The workflow can be repurposed to build one that predicts, for instance, salt bodies instead of fault locations. We change our labeling scheme so that salt bodies are labeled before training. Since salt bodies often create strong signals (at least for the top-of-salt events) we expect that this geologic feature could also be identified using this method. Preliminary results along this line are promising.
Conclusions

We present a novel approach to the challenging multistep seismic model-building problem. It uses a deep learning system to map out a fault network in the subsurface, using raw seismic recordings as input. A distinguishing aspect of the solution is the use of the Wasserstein loss function, which is suited to problems in which the outputs have spatial layout dependency. We demonstrate the system’s performance on synthetic data sets with simple fault networks. The primary challenge going forward will be transitioning to fault networks with more complex 3D geometry. This will lead to more tuned existing networks and/or extensions to our workflows.

The application of machine learning approaches to seismic imaging and interpretation shows great promise in hydrocarbon exploration and can dramatically change how the vast amount of seismic data is used in the future.

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Corresponding author: Mauricio.araya@shell.com

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