Using Machine Learning to Optimize Storage Systems

Dr. Kiran Gunnam
Outline

1. Overview
2. Building Flash Models using Logistic Regression.
3. Storage Object classification
4. Storage Allocation recommendation engines for elastic storage systems.
5. Building an end-to-end model
Overview
What is Machine Learning?

- **AI**: capability of a machine to imitate intelligent human behavior. (Artificial general intelligence (AGI)-not yet there, Artificial narrow intelligence (ANI)- specific tasks such as playing GO, playing chess, self-driving cars)
- **Machine learning**: subset of AI. Algorithms to analyze data to make and improve the decision making process.
- **Formal Definition**: A computer program is said to learn from experience E with respect to some class of tasks T and performance measure P if its performance at tasks in T, as measured by P, improves with experience E.
- **Deep Learning**: Subset of Machine Learning. Analyze data using multiple layers of abstraction. Examples are Convolutional Neural Networks, Recurrent Neural Networks etc.
Machine learning techniques

- Supervised Learning involved constructing trained models with known input data and output data.
- Unsupervised Learning finds patterns within data about which nothing is known prior.
- Reinforcement learning is a reward based learning where the algorithm learns to make specific decisions based on broader guideline.
Supervised vs Unsupervised

SUPERVISED
- Evidence Based Prediction
- Takes a known set of input data and responses to predict responses to new input data.
- Classification - discrete responses
- Regression - continuous responses

UNSUPERVISED
- Draw inferences from data sets consisting of input data without strictly labelled responses.
- Clustering - analyze data to find patterns and groupings
### Algorithms

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<th>Unsupervised</th>
<th>Reinforcement</th>
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<td>Policy-based</td>
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<td>(applied for games such as Chess</td>
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<td>and GO where rules of game are</td>
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<td>known and model is perfect)</td>
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</tbody>
</table>

8/7/2017 Kiran Gunnam, Flash Memory Summit 2017
Flash Model and Parameter Optimization
Flash Parameter Optimization

- Device Manufacturers tune the flash parameters to achieve reasonable specification for broad
- However it is possible to optimize the flash parameters based on the operating state of the flash memory.
- Advantage of flash parameter optimization: Improve the flash endurance.
- However the challenge is that many registers has dependencies. It is critical to determine a set of register settings that satisfy all the criterion.
Logistic Regression to build Flash Models

- Why use logistic regression?
- Estimation by maximum likelihood
- Interpreting coefficients
- Hypothesis testing
- Evaluating the performance of the model
The main approaches are:

• **Forward selection**, which involves starting with no variables in the model, testing the addition of each variable using a chosen model fit criterion, adding the variable (if any) whose inclusion gives the most statistically significant improvement of the fit, and repeating this process until none improves the model to a statistically significant extent.

• **Backward elimination**, which involves starting with all candidate variables, testing the deletion of each variable using a chosen model fit criterion, deleting the variable (if any) whose loss gives the most statistically insignificant deterioration of the model fit, and repeating this process until no further variables can be deleted without a statistically significant loss of fit.

• **Bidirectional elimination**, a combination of the above, testing at each step for variables to be included or excluded.
Model Building

Start: Constant term only, no variable terms yet

Stage I: test
Linear terms

Stage II: test
Cross-product terms

Stage III: test
Univariate 2nd, 3rd, 4th order terms

Sufficient?

Yes

Delete insignificant variable terms

Model selected; done.

Linear terms added

Cross-product terms added

Higher order terms added

No

Necessary?

Yes

No
Why use logistic regression for Flash Models?

- There are many important research topics for which the dependent variable is "limited."
- There are several independent variables and combinations of independent variables (flash device registers) affect the dependent variable (flash page error)
- Binary logistic regression is a type of regression analysis where the dependent variable is a dummy variable: coded 0 (error) or 1 (no error)
The Linear Probability Model

In the OLS (ordinary least squares (OLS) or linear least squares regression):

\[ Y = \gamma + \varphi X + e \; ; \text{where } Y = (0, 1) \]

- The error terms are heteroskedastic
- \( e \) is not normally distributed because \( Y \) takes on only two values
- The predicted probabilities can be greater than 1 or less than 0
More:

- The logistic distribution constrains the estimated probabilities to lie between 0 and 1.
- The estimated probability is:

  \[ p = \frac{1}{1 + \exp(-\alpha - \beta X)} \]

- if you let \( \alpha + \beta X = 0 \), then \( p = .50 \)
- as \( \alpha + \beta X \) gets really big, \( p \) approaches 1
- as \( \alpha + \beta X \) gets really small, \( p \) approaches 0
Comparing LP and Logit Models

LP Model

Logit Model
Maximum Likelihood Estimation (MLE)

- MLE is a statistical method for estimating the coefficients of a model.
- The likelihood function (L) measures the probability of observing the particular set of dependent variable values \((p_1, p_2, ..., p_n)\) that occur in the sample:
  \[ L = \text{Prob} (p_1 \times p_2 \times \cdots \times p_n) \]
- The higher the L, the higher the probability of observing the ps in the sample.
MLE involves finding the coefficients \((\alpha, \beta)\) that makes the log of the likelihood function \((LL < 0)\) as large as possible.

Or, finds the coefficients that make \(-2\) times the log of the likelihood function \((-2LL)\) as small as possible.

The maximum likelihood estimates solve the following condition:

\[
\{Y - p(Y=1)\}X_i = 0
\]

summed over all observations, \(i = 1,\ldots,n\)
An interpretation of the logit coefficient which is usually more intuitive is the "odds ratio"

Since:

\[ \frac{p}{1-p} = \exp(\alpha + \beta x) \]

\( \exp(\beta) \) is the effect of the independent variable on the "odds ratio"
Storage Object Classification
## Storage Object Classes

<table>
<thead>
<tr>
<th>File class</th>
<th>Example policy</th>
</tr>
</thead>
<tbody>
<tr>
<td>File size is zero</td>
<td>Allocate with directory</td>
</tr>
<tr>
<td>$0 &lt; \text{size} \leq 16\text{KB}$</td>
<td>Use RAID1 for availability</td>
</tr>
<tr>
<td>File lifespan $\leq 1\text{sec}$</td>
<td>Store in NVRAM</td>
</tr>
<tr>
<td>File is write-only</td>
<td>Store in an LFS partition</td>
</tr>
<tr>
<td>File is read-only</td>
<td>Aggressively replicate</td>
</tr>
</tbody>
</table>

**Table 1. Classes we want to predict.**
Architecture overview of self-learning

Source: http://repository.cmu.edu/pdl/69/
# Features

<table>
<thead>
<tr>
<th>Request Features</th>
<th>Workload Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Types of current and x previous requests</td>
<td>Number of reads, number of writes, and read/write ratio</td>
</tr>
<tr>
<td>Individual request size</td>
<td>Average request size</td>
</tr>
<tr>
<td>Sequential or random</td>
<td>Sequential/random ratio</td>
</tr>
<tr>
<td>Arrival times of current and x previous requests</td>
<td>Average request arrival rate</td>
</tr>
<tr>
<td>Number of processes (issuing requests)</td>
<td>Average number of processes</td>
</tr>
<tr>
<td>Think time for each request</td>
<td>Average think time</td>
</tr>
<tr>
<td>Inter-request block number distances between current request and x previous requests</td>
<td></td>
</tr>
<tr>
<td>Logical block number of each request</td>
<td></td>
</tr>
</tbody>
</table>
Example Data and Tree

### ID3 Algorithm

1. Select attribute A as node to split on (based on relative ranking).
2. Split samples according to values they take on attribute A.
3. If leaf nodes are "pure", done.
4. Else, if attributes remaining, goto 1.

---

**Figure 1. Example data and tree.**
**k Nearest Neighbor**

- Three things are required
  - Feature Space (Training Data)
  - Distance metric - to compute distance between records.
    - Euclidean Distance
      \[ d(x_i,x_j) = \sqrt{\sum_{k}(x_{i,k} - x_{j,k})^2} \]
    - Manhattan Distance
      \[ d(x_i,x_j) = \sum_{k} |x_{i,k} - x_{j,k}| \]
  - The value of k – the number of nearest neighbors to retrieve from which to get majority class.

- Choosing the value of k:
  - If k is too small, sensitive to noise points
  - If k is too large, neighborhood may include points from other classes
  - Choose an odd value for k, to eliminate ties

\[
\begin{align*}
  &\text{k=4 the unknown purple symbol belongs to red square class} \\
  &\text{k=11 the unknown symbol belongs to yellow diamond class} \\
  &\text{K = 17 it belongs to the yellow diamond class still.}
\end{align*}
\]
Identity (absolute) error

\[ d_j(x_{ij}, x'_{ij}) = I(x_{ij} \neq x'_{ij}) \]

Squared distance

\[ d_j(x_{ij}, x'_{ij}) = (x_{ij} - x'_{ij})^2 \]

\( L_q \) norms

\[ L_{qi'q'} = \left( \sum_j |x_{ij} - x'_{ij}|^q \right)^{1/q} \]

Canberra distance

\[ d_{ii'} = \sum_j \frac{|x_{ij} - x'_{ij}|}{|x_{ij} + x'_{ij}|} \]

Data point:

\[ x_i = [x_{i1} \ldots x_{ip}]^T \]

Correlation

\[ \rho(x_i, x_{i'}) = \frac{\sum_j (x_{ij} - \bar{x}_i)(x'_{ij} - \bar{x}_{i'})}{\sqrt{\sum_j (x_{ij} - \bar{x}_i)^2 \sum_j (x'_{ij} - \bar{x}_{i'})^2}} \]
k Nearest Neighbor

- The straightforward algorithm has a cost of $O(n \log(k))$, not good if the dataset is large.
- We can use indexing with k-d trees as follows
- For eg. We have the following training data
  - \{(1,9),(2,3),(4,1),(3,7), (5,4), (6,8), (7,2), (8,8), (7,9), (9,6)\}
- Build a K-D tree
  - Pick a dimension, find median, split data, repeat.
- Find nearest neighbors for new point (7,4)
  - Find region contacting (7,4)
  - Compare to all point in region
Approximation of discrete functions by a decision tree. In the nodes of trees are attributes and in the leaves are values of discrete function.

Deriving a tree

Until each leaf node is populated by as homogeneous a sample set as possible: Select a leaf node with an inhomogeneous sample set. Replace that leaf node by a test node that divides the inhomogeneous sample set into minimally inhomogeneous subsets, according to an entropy calculation.
Using Boosted Decision Trees

- Boosting means that each tree is dependent on prior trees, and learns by fitting the residual of the trees that preceded it.
- Flexible: can deal with both continuous and categorical variables
- How to control bias/variance trade-off
  - Size of trees
  - Number of trees
- Boosting trees often works best with a small number of well-designed features
- Boosting “stubs” can give a fast classifier
- Supervised learning method and needs labeled dataset
Allocation recommendation engines in Elastic Storage Systems
Block diagram of a storage allocation system
More on storage allocation

Partitioned into Node vs Cluster shared
Application Optimized (semantics consistent with data types)
Application Direct Accessed (use space IO)
Intelligent Data placement for scalability
An unsupervised clustering algorithm

“$K$” stands for number of clusters, it is typically a user input to the algorithm; some criteria can be used to automatically estimate $K$

It is an approximation to an NP-hard combinatorial optimization problem

$K$-means algorithm is iterative in nature

It converges, however only a local minimum is obtained

Works only for numerical data

Easy to implement
$x_1,\ldots, x_N$ are data points or vectors of observations

Each observation (vector $x_i$) will be assigned to one and only one cluster

$C(i)$ denotes cluster number for the $i^{th}$ observation

Dissimilarity measure: Euclidean distance metric

$K$-means minimizes within-cluster point scatter:

$$W(C) = \frac{1}{2} \sum_{k=1}^{K} \sum_{C(i)=k} \sum_{C(j)=k} \|x_i - x_j\|^2 = \sum_{k=1}^{K} N_k \sum_{C(i)=k} \|x_i - m_k\|^2$$

where

$m_k$ is the mean vector of the $k^{th}$ cluster

$N_k$ is the number of observations in $k^{th}$ cluster
**K-means Algorithm**

For a given cluster assignment $C$ of the data points, compute the cluster means $m_k$:

$$m_k = \frac{\sum_{i:C(i)=k} x_i}{N_k}, \ k = 1, \ldots, K.$$

For a current set of cluster means, assign each observation as:

$$C(i) = \arg \min_{1 \leq k \leq K} \|x_i - m_k\|^2, \ i = 1, \ldots, N$$

Iterate above two steps until convergence
K-means clustering example

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Kiran Gunnam, Flash Memory Summit 2017
How to build an end to end application

UNDERSTANDING CHALLENGES AND SELECTING RIGHT MACHINE LEARNING ALGORITHM, DATA PREPROCESSING, EVALUATING MODEL
CHALLENGES IN MACHINE LEARNING

- Heterogeneous Data
  - Dataset could have images, text and sensor signals as composite data.

- Preprocessing of data
  - Numeric datasets could have data lying in different data ranges and would require to be normalized before they are used.

- Picking the best suitable model
  - Based on the kind of application we are dealing with we need to pick a suitable algorithm. This decision may not be very intuitive.
Selecting an algorithm

- Selecting a machine learning algorithm deals with having insight about the kind of data with which we are dealing.
- If we know the nature of the data and need to make a future prediction then we should select supervised machine learning algorithms.
- If we do not know the nature of the data and would like to discover the patterns within it, we should select an unsupervised machine learning algorithm.
The perfect classification algorithm

- Objective function: encodes the right loss for the problem
- Parameterization: makes assumptions that fit the problem
- Regularization: right level of regularization for amount of training data
- Training algorithm: can find parameters that maximize objective on training set
- Inference algorithm: can solve for objective function in evaluation
Selecting Model

Try simple classifiers first

Better to have smart features and simple classifiers than simple features and smart classifiers

Use powerful classifiers if more training data is available (bias-variance tradeoff)
Bias-Variance Trade-off

- Models with too few parameters are inaccurate because of a large bias (not enough flexibility).
- Models with too many parameters are inaccurate because of a large variance (too much sensitivity to the sample).

Source: D. Hoiem
Bias-Variance Trade-off

\[ E(\text{MSE}) = \text{noise}^2 + \text{bias}^2 + \text{variance} \]

- Unavoidable error
- Error due to incorrect assumptions
- Error due to variance of training samples

See the following for explanations of bias-variance (also Bishop’s “Neural Networks” book):

- [http://www.inf.ed.ac.uk/teaching/courses/mlsc/Notes/Lecture4/BiasVariance.pdf](http://www.inf.ed.ac.uk/teaching/courses/mlsc/Notes/Lecture4/BiasVariance.pdf)
Bias-Variance Trade-off

Few training examples

Many training examples
Effect of Training Size

![Graph showing the effect of training size on error]

- Fixed prediction model
- Error vs. Number of Training Examples
- Testing vs. Training
- Generalization Error
Regularization

- When the model fits the training data but does not have a good predicting performance and generalization power, we have an overfitting problem.
- Regularization is a technique used to avoid this overfitting problem.
- The idea behind regularization is that models that overfit the data are complex models that have for example too many parameters.
Data Preprocessing
Data Pre-processing

- Data Cleaning
  - Correct bad data, filter some incorrect data out of the data set and reduce the unnecessary detail of data.

- Data Transformation/normalization/Feature scaling
  - Feature scaling is a method used to standardize the

\[
x' = \frac{x - \bar{x}}{\sigma}
\]

Where \(x\) is the original feature vector, \(\bar{x}\) is the mean of that feature vector, and \(\sigma\) is its standard deviation.

\[
x' = \frac{x - \min(x)}{\max(x) - \min(x)}
\]
If we are in a data-rich situation, the best approach for both *model selection* and *model assessment* is to randomly divide the dataset into three parts: training set, validation set, and test set.

The *training set* is used to fit the models. The *validation set* is used to estimate prediction error for model selection. The *test set* is used for assessment of the prediction error of the final chosen model.

A typical split might by 50% for training, and 25% each for validation and testing.
Resampling methods – a simple list

- The validation set approach,
- The bootstrap method for assessing statistical accuracy.

- Please see http://tinyurl.com/ybojkt9f
- And http://tinyurl.com/y8oxwq9s
- For more details.
Thanks!