Math for Machine Learning

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January 25, 2015

Abstract

In this post, I want to discuss the connections between Machine Learning and various other fields (especially Mathematics), citing specific examples where they come up. I have given very high-level explanations below and cut corners at several places as I do not want to get into the depth. My intention here is not to explain any concept precisely, but to merely lay down enough of them on the table to emphasize the role of Mathematics in this fast growing area.

1 Ok, why so much Math?

Machine Learning is an incredibly modern field that borrows heavily from several areas of Mathematics. Having evolved as an inter-disciplinary field which is very applied (driven by data), it has captured concepts, intuition and theory from several places. Being at such an intersection of diverse areas of mathematics and computer science is what makes research in Machine Learning so exciting and challenging!

At this point, I would like to mention Physics as an analogy. I consider Machine Learning to be very similar to Physics as a discipline, primarily because both are applied areas by nature, governed by deep mathematical foundations. It turns out, much of the pre-requisite Math for Machine Learning (Multi-variable calculus, linear algebra) applies to Physics too. Another similarity between the two fields is their philosophy and connection to the real world. In both cases, we try to model the real world phenomena by coming up with hypothesis and backing it up with experiments. While we might not be able to fit most of the happenings in the real world exactly, we try to get as accurate as possible. This in turn leads us to a better understanding of the black-box that generates events in the real world (or data in case of machine learning).

"Imagine being in a battlefield without knowing how to use the weapons you have, (or worse still, not having the weapons at all)!"

That’s exactly how it feels when you set out to do research in machine learning without knowing enough about the fundamental areas underlying it. Without the right intuition, it becomes very hard to build new algorithms or extend existing ones.

Below are the key useful areas:

1.1 Algorithms & Complexity

Knowledge of basic data structures such as arrays/trees/hash tables, programming techniques like dynamic programming, graphs, space and time complexity requirements for a given method, randomized algorithms, sublinear
algorithms.

How do you convince someone that your learning algorithm is more space and/or time efficient (and hence scalable) on big datasets? Exploiting sparsity in the data sets might lead you to better performing algorithms; but how do you qualitatively compare their performance? With data getting more and more massive, going through it entirely is not feasible and therefore sometimes even linear time algorithms maybe too slow. This has led to the field of sub-linear algorithms which work by inspecting only a tiny fraction of the entire data. Property Testing is a closely related topic where the algorithm queries about some property of the input with a time complexity much smaller than the size of the input. These are relatively modern areas in theoretical computer science which have a direct impact on machine learning. Randomized algorithms is another highly useful field that has helped solve several big data problems, for eg large matrix problems have been very successfully dealt with using randomization techniques.

1.2 Linear Algebra

*Rank of a Matrix, Matrix Vector products, Column Spaces and Null Spaces of a matrix, Eigen Values and Vectors, SVD factorization of a matrix, positive-definiteness of a matrix.*

Linear Algebra plays a super heavy role in understanding Optimization methods used for Machine Learning. Let’s take an example to see how. Many problems in machine learning can be expressed as a simple least-squares optimization problem. What is interesting is every least-squares problem can be turned into a Quadratic Program (ie, an optimization problem involving quadratic function of the variables). This is illustrated below:

\[
f(x) = \frac{1}{2} \|Qx - c\|^2
\]

\[
= \frac{1}{2} (Qx - c)^T(Qx - c)
\]

\[
= \frac{1}{2} (x^TQ^TQx - x^TQ^Tc - c^TQx + c^Tc)
\]

Since \(c^Tc\) is a fixed quantity (constant), it is sufficient to solve the Quadratic Programming problem:

\[
f(x) = \frac{1}{2} x^T A x + q^T x
\]

where \(A = Q^TQ\) and \(q = -Q^Tc\). Observe that eqn(2) has a matrix \(A\) in its first term. This matrix \(A\) is the Hessian (generalization of second-order derivatives to higher dimensions) and when it is positive-definite, the quadratic problem takes shape of a ”bowl” in higher dimensions as shown below in Figure 1.

![Figure 1: Three dimensional “bowl” shaped function.](image)

As a consequence, it clearly has only one ”unique” minima (global optimal solution). This concept is termed as ”Convexity” (which is considered a blessing for optimization as such functions are well-understood and have some very attractive properties). Some of these ideas constitute the nuts and bolts of Convex Optimization. Likewise, when the matrix \(A\) is negative-definite, the function takes the shape of an inverted bowl and is not convex (therefore, has no global and unique minimum, but has a global maximum). Finally, when matrix \(A\) is indefinite, then the function takes an interesting form (resembling a saddle) as below in Figure 2. As we can imagine, there

\(^1\)Note that \(Q\) is a matrix, while \(c\) and \(x\) are vectors. This way of writing functions in terms of vectors and matrices is called as vectorizing. This is described in 1.4.
Although the data we obtain in the real world has very large dimensions (hundreds of thousands for example), it can often be reduced to a handful of useful dimensions that we can work with. This is called low-rank approximation (rank of a data matrix determines the true dimensions of your data or how diverse your data actually is). Matrix Factorization methods are based on this and typical recommender systems like the one Netflix uses to predict movie ratings of a user, make use of it. Low-rank approximations are also used in other areas like Computer Vision and Information Retrieval as a tool for extracting correlations in data and removing noise from matrix-structured data.

Another interesting Dimensionality Reduction algorithm is PCA (Principal Components Analysis). A simple way to understand PCA is to visualize a bunch of red and blue data points dispersed in a 3-D space (assume the red points represent spam emails and blue represent legit emails). These points can be projected onto a 2-D surface in different ways (for eg: onto the XY plane, onto the YZ plane or onto the XZ plane). If we examine each of these projections, we will realize that the distribution of points in each of them are slightly different. Some of these projections separate the red and blue points better than the rest. The job of PCA is to identity such projections which yield maximal separation (or capture maximum variance in the data) as they are the most useful to us. And how can we do this? Well, that’s where eigen vectors and values come to our rescue. The principal eigen vector of the covariance matrix (computed using the data points we have) corresponds to the axis that provides maximum separation / variance in our data. PCA is used widely in Computer Vision and Image Compression, to mention a few.

Algorithms in machine learning involve dozens of vector-vector multiplications (dot-products) and matrix-vector, matrix-matrix multiplications. All of these operations can be extremely costly and a bottleneck when trying to scale to big data. However, if we can cleverly manipulate or take advantage of special matrices which contains lots of zeros (“sparsity”), we can reduce such computations significantly.

1.3 Probability Theory & Statistics

- **Probability Theory:** Counting and Combinatorial methods, Bayes’ Theorem, Random Variables, Expectation, Variance, Conditional and Joint Distributions, Moment Generating Functions, Exponential Family of Distributions
- **Statistics:** Maximum Likelihood Estimation, MAP, Prior and Posterior, Sampling methods, Gibbs

As you would expect, this is the single-most important field which also conveys the essence of machine learning, namely - estimating the parameters of the data-generating process. Several machine learning methods have probabilistic interpretations and its common to hear the words frequentist and bayesian ways of doing things. One way to look at the difference between them is that the frequentist methods are concerned with estimating the parameters of their model that have the highest chance of generating the "current data"; this is called the Maximum Likelihood Estimation (MLE) and written as:

$$\arg\max_{\theta} \log \mathcal{L}(\theta) = \arg\max_{\theta} \log P(\text{Data}|\text{Parameters})$$ (3)

MLE has the tendency to overfit (generalize poorly on unseen data) and hence the Bayesian approach proposes incorporating historical evidence (based on "past data") into the current model. This is called the prior. The task
of estimation then boils down to using Bayes’ rule as below:

\[ P(\text{Parameters}|\text{Data}) = \frac{P(\text{Data}|\text{Parameters})P(\text{Parameters})}{P(\text{Data})} \]  

(4)

This can be equivalently written as:

\[ \text{Posterior} = \frac{\text{Likelihood} \times \text{Prior}}{\text{Normalizing Constant}} \]  

(5)

Posterior gives us a probability distribution over the parameters and this is used in various ways to make predictions on the new data. As is evident by now, random variables play a huge part in estimation and we often deal with independence assumptions between them, work with their expected values and variances. It is also important to know the functional forms of some key probability distributions, for instance the most popularly used Gaussian Distribution (or normal distribution) \( \sim N(\mu, \sigma^2) \), which (in its univariate form) can be expressed as:

\[ \frac{1}{\sigma \sqrt{2\pi}} \exp \left\{ -\frac{(x - \mu)^2}{2\sigma^2} \right\} \]  

(6)

Methods of Sampling play an important role in optimization algorithms. Often the gradient (generalization of the derivative - this has been described in a bit more detail further down below) needs to be calculated over the entire data set and this is very expensive to compute in every iteration. To avoid this, algorithms like Stochastic Gradient Descent (SGD) randomly sample a data point and update its gradient alone, this makes the algorithm independent of the number of data points which means it will scale well. But, now we only selected specific data points; so our method has lot of variance. How can we reduce this and make sure randomly selecting one point in each iteration will in the long run mimic the same behavior that we would have got if we had selected all the data points and computed the exact gradient? This is where techniques of designing unbiased sampling mechanisms come up.

### 1.4 Multi-variable Calculus (Matrix-Vector Calculus)

*Vector-valued functions, Partial-Derivatives, Gradient, Directional Gradient, Hessian, Jacobian, Laplacian, Lagrange Multipliers*

We know from our high-school calculus that *derivatives* represent the "rate of change" of the function at a given point. And, in order to minimize or maximize a function you set its derivative equal to zero. Such a way of obtaining the solution is called the "closed-form" solution and this was an easy thing to do because back then, our functions then involved just handful of dimensions (often just a single variable). However, in machine learning its very common to deal with functions that operate on variables having hundreds of dimensions. What does derivative or slope even mean in such situations? That’s where partial-derivatives come into the picture. You can look at them as derivatives of the function in each dimension of the variable. Combine these partial-derivatives into a vector and that gives us what is called the "Gradient". Similarly, taking the second-order derivative of Gradient gives us a matrix termed as the "Hessian". Also, in reality it is often not possible to obtain a "closed-form" solution (by setting the gradient to zero) because the dimensions of the gradient could be extremely high and storage/time required for this might be expensive. In such cases, knowledge of gradients and hessians help us define things like directions of descent and rate of descent which tell us how should we travel in our function space in order to get to the bottom most point (which represents the optimal solution). Thus at this point, we work using more abstract objects - "Vectors" and "Matrices" and should be reasonably comfortable with applying differential and integral calculus on them. This is also called "Vectorizing the equations", which Prof. Andrew Ng discusses in detail in one of his ML lectures (Refer Coursera). Below is an example of a simple objective function (linear regression with weights \( x \), \( N \) data points and \( D \) dimensions) both in the vectorized and expanded notation.

**Expanded notation:**

\[ J(w) = \frac{1}{2} \sum_{i=1}^{N} \left( \sum_{j=1}^{D} a_{ij}x_j - b_i \right)^2 \]  

(7)

**Vectorized notation:**

\[ J(w) = \frac{1}{2} \|Ax - b\|^2 \]  

(8)

As you can see the vectorized notation is not only compact and gets rid of the annoying summations but also is more expressive and makes us think more abstractly in terms of higher level objects like vectors (x and b) and
matrices (A). Such representations also help us leverage all the rich properties that vectors and matrices provide us (that speed up our computation and make things easily parallelizable, to mention a few). What this also means is that, we need to now get comfortable with doing arithmetic in this new space. For instance, while moving terms around, we cannot divide by a matrix and instead have to compute the inverse (At this point, I should also mention that computing inverses of big matrices are expensive too and there are workarounds and techniques to deal with this). The method of Lagrange multipliers is a standard way in calculus to maximize or minimize functions when there are constraints involved. This comes up a lot in Optimization.

1.5 Real Analysis

*Properties of Sets and Sequences, Convergence of Sequence, Cauchy Sequences, Concepts from Topology like Open and Closed Sets, Metric Spaces, Limits and Continuity of functions, Differentiability*

While this is the most rigorous of all mathematical fields I have mentioned, its importance is highly understated. For a person in Machine Learning, (quoting a friend of mine in his exact words) Real Analysis is a long-term investment. It gives you sufficient practice with the art of writing mathematics proofs in a rigorous manner and making precise statements without gaps and holes in your arguments. This is an extremely useful skill to acquire that helps you when writing and reading research papers. Although intuition and pictures are often good ways to describe and understand ideas; compressing them in a clear mathematical way is sometimes valuable as it avoids ambiguity. Analysis also helps one navigate more comfortably through some fundamental concepts in numerical optimization such as - sequences, what does it mean for a sequence to converge to a limit point, how do you define the rate of convergence, what does it mean for a sequence or set to have a sup and inf? While fields of Linear Algebra and Multi-variate calculus help you design optimization methods for machine learning, tools from Real Analysis will help you verify their correctness theoretically and provide guarantees on convergence, etc.

1.6 Information Theory

*Entropy, Mutual Information, Information Gain, KL Divergence*

This branch of applied mathematics deals with studying how to quantify information. Entropy for example quantifies the uncertainty involved in predicting the value of a random variable and intersects with other fields like Probability Theory. KL-Divergence is a widely used metric to measure how different two probability distributions are. A simple example is to consider Decision Trees which is a popular classification method in machine learning. The way decision tree works is very similar to the famous Twenty Questions Game, where the player of the game is expected to guess a person’s name by asking twenty yes-no answer questions. If you have played this game before, you will realize that the first few questions one would ask would be the ones such as - "Is the person male or female?", "Is the person dead or alive?", "Is the person real or animated character?", etc as opposed more specific ones like "Did the person act in movie XYZ?", "Is the person tall?", etc. This is because the former set of questions divide your mental search space to the maximum, or in information theory terms - they provide the maximum information gain (or highest reduction in entropy). The computation that happens so rapidly in a split-second in our brain is picking all possible paths in the decision tree (in our analogy of the game, this is equivalent to considering all possible questions that can be asked), computing the reduction in entropy assuming we had gone down that path and then picking the one that gave maximum reduction in entropy. The mathematical tools that are used to analyze all of this have roots in information theory.

For the sake of completeness, let me mention that there are few other areas such as Differential Geometry and Measure Theory which make cameo appearances once in a while, but the show is predominantly run by the ones above!

**Note:** If you find any mistakes or have suggestions on ways I could improve this document please send me an email at: params@ucsc.edu. Thanks for reading!