Generalization: The Hidden Agenda of Learning.
The Past, Present and Future of Neural Networks for Signal

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practical algorithms. One such application is an intelligent user interface agent where a software program is able to adjust its behavior based on observation of users' action as well as interaction with the user through some sort of dialog. For example, in modern speech-recognition software, speaker adaptation is an important component. The current approach is to implement the speaker-adaptation unit as a passive learner that will learn new information about the speaker whenever the speaker points out the mistake in the recognition results. It is possible that the user must perform numerous corrections before the speaker adaptation is fully adapted to the speaker-specific acoustic characteristics. On the other hand, an active speaker-adaptation algorithm would be able to request the speaker to read some specific sentences and quickly adapt to speaker's characteristics. Another example is an intelligent information browser [10] where the browser is able to aid a novice user in locating the most relevant documents by pre-classifying the documents through clustering. Then it would ask user to clarify the ambiguity by presenting the user with sample documents that locate near the present estimate of the decision boundary. An initial prototype of this tool has shown great potential [10].

References

Generalization: The Hidden Agenda of Learning

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Most neural systems are adapted by optimization of a performance index, typically the minimization of a "cost function," based on a finite database (a training set) of \( N \) noisy examples derived from the target system. However, there is always the hidden agenda that the model should perform well, not only on the training set, but on the much larger set of future inputs to the system.

While reading for your finals you solve the previous years' tests, but you know very well that if you then test yourself on last year's test, the result will be biased—too optimistic! Only a test on a fresh data set, a test that was put aside before you started reading, will give you a reliable prediction of the final performance.

Doing well on unseen data may at first seem unattainable, but the ability to generalize in very complex environments is nevertheless one of the most striking properties of neural systems, and indeed one of the reasons that neural networks have shown useful in practical applications.

As an example: in [1], a neural-network system for inspection of handwritten digits was able to classify 99.98% correct after training on a data base of 7291 digits and classify 95% correct on an additional test set of 2007 digits.

When using a super-flexible model family, like neural networks, which in principle can model arbitrarily complex systems, overfit is a major concern, which finds expression in the ubiquitous bias-variance dilemma [2]. The generalization ability of an adaptive system is the quantitative measure of performance on a hypothetical infinite test set. While this quantity cannot be accessed directly, algebraic asymptotic estimates of generalization, valid for large training sets \( (N \rightarrow \infty) \), can be derived [3-8]. Such asymptotic results were earlier derived for supervised learning; however, it was recently shown that generalization ability for unsupervised learning machines (e.g., principal component analysis and clustering schemes) can be analyzed in a similar framework [9].

If sufficient computational capacity is available, empirical resampling schemes can be invoked. The two basic resampling strategies are cross-validation and bootstrap. Cross-validation [10, 11] is based on a random division of the database into disjoint training and validation sets. The procedure can be repeated, leading to more accurate results at the price of increased computation. The so-called leave-one-out cross-validation is based on using only a single example in the test set, and typically resampling \( N \) times. Approximative techniques, by which the computational overhead in leave-one-out is significantly reduced, have been reported [7, 12].

Bootstrap, invented by Efron [13], is based on resampling with replacement. Bootstrap produces pseudo training sets of size \( N \) and, hence, simulates training set fluctuations at the full sample size. It has been applied to control overfit in a number of investigations [14-16].

Optimization of the neural-network architecture may lead to better generalization ability and preferably lower computational burden. Optimizing the network architecture is to optimally trade off bias and variance [2], hence, maximizing generalization ability. This can be done di-
rectly by optimizing the structure of the network by pruning or growing techniques or indirectly by using regularization. Regularization, which goes back to Hadamard, consists of adding a penalty term to the cost function. As an example, consider predicting the sunspot time series shown in Fig. 3. In contrast, Fig. 4 [17] shows that generalization error (test error) is reduced by pruning the network.

References
On-Line Step-Size Selection for Training of Adaptive Systems

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An adaptive system is by its very nature time-varying. The rate at which such a system changes its internal parameters determines its capabilities to adjust to and to obtain useful information from an unknown physical environment. One of the simplest and most-popular techniques for adjusting an adaptive system’s parameters is the gradient-descent method, in which the parameters are changed according to the derivatives of a particular cost function with respect to the current parameter values. Both the least-mean-square (LMS) algorithm for adaptive filters [1] and the back-propagation algorithm for multilayer perceptrons [2] are gradient-descent methods. Much is known about the behavioral characteristics of gradient adaptation, and the algorithms are usually numerically robust. In gradient descent, the step sizes control the magnitudes of the changes in the parameters in the negative direction of the gradient.

The Kalman filter forms the basis for another class of parameter-estimation techniques that employ second-order information about the cost function being minimized. Recursive least-squares (RLS) techniques are widely used in linear estimation tasks and [3] explore the connections between RLS and Kalman techniques. The extended Kalman filter is a linearized version of the Kalman filter for nonlinear state-space estimation tasks [4], and it has been successfully applied to multilayer perceptron training [5]. Many simplified and approximate versions of this algorithm have been studied. Note that in RLS and linearized least-squares methods with forgetting factor \( \lambda \), the parameter \( (1 - \lambda) \) plays the role of the step size, whereas in Kalman techniques the step size is automatically determined from its underlying Bayesian problem formulation.

**Goal of Step-Size Selection**

The performance of any adaptive system that is attempting to drive its adjustable parameters to an optimum fixed set of parameters is governed by two quantities: (i) its convergence rate and (ii) the misadjustment in steady state. The convergence rate refers to the transient behavior of the parameters as they approach their optimum values. Misadjustment refers to the additional error in the output of the system caused by the random fluctuations of the parameters in steady-state. Generally speaking, the convergence rate of a system increases for step sizes that are somewhat less than one-half of the maximum value of the step size that provides stable adaptive behavior. In contrast, the misadjustment generally decreases as the step size is decreased.

The goal of any time-varying step size procedure is to increase the step size to a large but stable operating value when the parameters are some distance from their optimum settings and to systematically decrease the step size to reduce the misadjustment when the parameters are in the vicinity of their optimum settings. Since stability is often not explicitly ensured within a time-varying step-size method, it is generally necessary to limit the range of step sizes to guarantee stable operation of the system.

When the desired parameter settings vary with time, the system must continually readjust its parameters to follow these variations. The error induced in the model parameters by any time-variation of the unknown optimal parameters is called the lag error. In some cases where the velocity of the model parameters is constant, an optimum step-size value exists that minimizes the contributions of the misadjustment and the lag error after all initial transients in the parameters have died out.

We can classify a step-size selection method as adaptive or nonadaptive depending on its form. Nonadaptive methods calculate a time-varying step size according to a priori knowledge about the signals being processed, the cost function being optimized, and/or the parameter structure of the system. These techniques include asymptotically optimal methods as derived via the theory of stochastic-approximation [6] methods based on a statistical analysis of the particular system [7, 8] and heuristic approximations to these methods, commonly known as “search-and-converge” [9] or “gear-shifting.” By contrast, adaptive methods are based on on-line measurements of the state of the adaptive system, usually as characterized by the outputs or by the parameter updates of the system.

Nonadaptive step-size methods usually require more information about the adaptive system and the problem context than do adaptive step-size methods. However, nonadaptive step-size methods usually outperform adaptive step-size methods because of this increased knowledge.

**On-Line Adaptive Step-Size Selection**

Gradient adaptation has proven to be quite useful for parameter estimation. It can also be used to optimize the step-size parameters in an on-line fashion. This idea has appeared and reappeared in the scientific literature. One of the earliest descriptions of the methodology appears in [10], and it was later reintroduced to both the neural-network [11] and signal-processing [12, 13] communities, where it has become known as the “delta-bar-delta rule” and “gradient step-size method,” respectively. An alternative version of