Pattern discovery via entropy minimization

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Abstract

We propose a framework for learning hidden-variable models by optimizing entropies, in which entropy minimization, posterior maximization, and free energy minimization are all equivalent. Solutions for the maximum a posteriori (MAP) estimator yield powerful learning algorithms that combine all the charms of expectation-maximization and deterministic annealing. Contained as special cases are the methods of maximum entropy, maximum likelihood, and a new method, maximum structure. We focus on the maximum structure case, in which entropy minimization maximizes the amount of evidence supporting each parameter while minimizing uncertainty in the sufficient statistics and cross-entropy between the model and the data. In iterative estimation, the MAP estimator gradually extinguishes excess parameters, sculpting a model structure that reflects hidden structures in the data. These models are highly resistant to over-fitting and have the particular virtue of being easy to interpret, often yielding insights into the hidden causes that generate the data.

1 Motivation

In pattern discovery we seek a model that reflects the structure of the data, which we hope in turn reflects the structure of the generating process. The unspoken premise is that the universe is constructed out of relatively small processes; in order to produce an object (dataset) larger than itself, a small process must have some kind of repetition structure, e.g., loops. Therefore a dataset is an “unrolled” record of the process’ internal structure. If we can find a compact model of the data, we feel increasingly confident that 1) we can interpret the model and in doing so learn about the process, and 2) predictions made from the model will be consistent with new samples taken from the process. We now know that #2 is well-founded: Recent theorems in algorithmic complexity assure us that of all possible models (e.g., of all possible Turing machines) the one yielding the smallest two part encoding—model plus data relative to the the model—is the best strategy for hypothesis identification [Vitanyi and Li, 1996] and almost always the best strategy for prediction [Vitányi and Li, 1997].

Sadly, the function which yields the most compact model is not computable. Furthermore, we only know how to search efficiently for accurate models (disregarding compactness) in quite restricted spaces, e.g., for locally optimal parameterizations of stochastic models whose kind and structure we fix in advance. In this paper we broaden that scope by estimating both model structure and parameters in a manner that reveals salient structures in the data. Fortuitously, our framework also yields a fast hill-climbing procedure for finding nearly global optima.

2 Setting

We are interested in learning (point estimating) a single model that is highly predictive and whose structure reflects the structure of the generating process. We must address three important limits: finite data, finite time, and finite precision. In particular, we want to extract as much essential structure as possible from a dataset without modeling any of its accidental structure (e.g., noise and sampling artifacts); we want to do so without any wasted or speculative computation (e.g., models discarded by a selection process); and we want to maximize the information content of all parameters (e.g., bits of evidence supporting each parameter) while being realistic about the information capacity of digitally represented numbers. We begin in a Bayesian setting, but in §6 we shall show that our framework has its clearest interpretation when learning is considered a problem of minimizing entropies.

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We begin with a set of observations \( \mathbf{X} = \{ \mathbf{x}_1, \mathbf{x}_2, \ldots, \mathbf{x}_N \} \) and a hypothesis class of hidden-variable models whose likelihoods are (approximated in practice by) products of densities drawn from the flat-exponential family, e.g., those having minimal sufficient statistics and parameter vectors of equal dimension\(^1\). We use hidden variables because we assume that the data has latent structure and/or is incomplete. The vector \( \mathbf{\theta} = \{ \theta_1, \ldots, \theta_N \} \) parameterizes a model and specifies its structure via a sparse encoding. Starting from a random over-parameterized model \( \mathbf{\theta} \) we seek an optimal embedded model \( \mathbf{\theta}' \) that maximizes the posterior given by Bayes’ rule: \( \mathbf{\theta}' = \arg \max_{\mathbf{\theta}} P(\mathbf{\theta} | \mathbf{X}) \propto P(\mathbf{X} | \mathbf{\theta}) P(\mathbf{\theta}) \), where the likelihood \( P(\mathbf{X} | \mathbf{\theta}) \) measures accuracy in modeling the data and the prior \( P(\mathbf{\theta}) \) measures consistency with our background knowledge.

3 Maximum structure priors

What is our background knowledge? Let us assert that the vector of model parameters \( \mathbf{\theta} \) is itself taken from a random variable \( \mathbf{\Theta} \) which generates all possible processes. We know that on average, these processes are not infinitely unpredictable—otherwise learning would be impossible! Therefore our background knowledge is that, on average, learning will succeed. More formally, we say that the expected entropy of processes from \( \mathbf{\Theta} \) is finite. We write this prior knowledge as \( \xi: \mathbb{E}_\mathbf{\Theta} [H(\mathbf{\Theta})] = h \), where \( H(\mathbf{\theta}) \) is an entropy measure assessed on the model specified by \( \mathbf{\theta} \), \( \mathbb{E}_\mathbf{\Theta} \) is the expectation with regard to \( \mathbf{\Theta} \), and \( h \) is some finite value. The classic maximum-entropy method [Jaynes, 1982, §2] allows us to derive the mathematical form of a distribution from knowledge \( \xi \) about its expectations via Euler-Lagrange equations, yielding

\[
P(\mathbf{\theta} | \xi) \propto \exp[-\lambda H(\mathbf{\theta})] \tag{1}
\]

where the Lagrange multiplier \( \lambda \) depends on \( h \) and is unknown. We shall find meaningful interpretations for several values of \( \lambda \), but we shall concentrate on the assumption that \( \lambda = 1 \). This we shall call the entropic prior

\[
P_\lambda(\mathbf{\theta}) \overset{\text{def}}{=} e^{-H(\mathbf{\theta})} \tag{2}
\]

We assume \( \xi \) henceforth and drop it from notation. We note in passing that by making a similar assertion about the expected perplexity \( e^{H(\mathbf{\theta})} \) and assuming a measure on \( \lambda \), it is also possible to integrate out the Lagrange multiplier and arrive directly at eqn. 2.

We call the reader’s attention to two properties that derive from the definition of entropy: 1) \( P_\lambda(\cdot) \) is a bias for compact models having less ambiguity, more determinism, and therefore more structure. 2) \( P_\lambda(\cdot) \) is invariant to reparameterizations of the model, because the entropy is defined in terms of the model’s joint and/or factored distributions.

We turn now from inferential principles to optimization problems. Models with hidden variables can be very difficult to fit and notoriously prone to over-fitting. Eqn. 2 is a remarkable form which we shall exploit to simultaneously estimate structure and parameters of complex probability models—a mixed combinatorial (structure) and continuous (parameter) optimization problem. We shall develop MAP estimators such that in expectation maximization (EM), the prior drives weakly supported parameters toward extinction, allowing them to be removed from the model without loss of posterior probability. The resulting models can be quite good; however, both hidden-variable and combinatorial optimization problems have notoriously rough energy surfaces and EM is merely a local hill-climber. Before presenting estimators in §5 we shall develop a technique that uses the entropic prior to improve the quality of the local optima found by EM, and which finds a quasi-global optimum in the limiting case.

4 Prior balancing

We now introduce a generalization of the posterior, by rewriting Bayes’ rule with a manipulation of the prior:

\[
\hat{P}(\mathbf{\theta} | \mathbf{X}, T, T_0) \overset{\text{def}}{=} \frac{P(\mathbf{X} | \mathbf{\theta}) P(\mathbf{\theta})^{T_0-T} \delta(T)}{P(T)} \tag{3}
\]

where \( T \) is normally positive and \( \delta(T) \in (0,1] \) is a driving term that monotonically increases as \( T \) approaches zero (e.g., \( \delta(T) \propto e^{-T/2} \)). Varying \( T \) balances the prior against the likelihood, which is useful in iterative parameter estimation because it allows \( \mathbf{\theta} \) to get into the right neighborhood with respect to one constraint before attempting to satisfy the other. Of course, to obtain a proper probability we are obliged to make \( Z = T_0 - T \) converge to a meaningful value (e.g., 1) which can be done by following the gradient \( \Delta T = \frac{\partial}{\partial T} \log \hat{P}(\mathbf{\theta} | \mathbf{X}, T, T_0) \) or by iteratively tracking the MAP estimate of \( T = \arg \max_T \hat{P}(\mathbf{\theta} | \mathbf{X}, T, T_0) \). Using the shorthand \( H = -\log P(\mathbf{\theta}) \), the gradient and MAP estimate for the Gaussian driving function are

\[
\Delta T \propto H - T \quad T = H \tag{4}
\]

\( T \)'s decay rate can be controlled with a variance term on \( \delta \), or by choice of a different driving function. E.g., \( \delta(T) \) may be one-sided (e.g., \( \delta(T) = e^{T^2 - \tau^2} \), \( \Delta T \propto H + 1 - e^{-T^2} \)) or a barrier term (e.g., \( \delta(T) = e^{-T^2/2} \), \( \Delta T \propto H - \sqrt{T} \), \( T = H^2 \)). If \( T_0 \) is not known, the gap from equilibrium \( T \) to the desired \( Z \) can be bridged with a heuristic schedule.

We introduce this as an optimization technique; it is unclear that there is any useful interpretation as an inference.
principle. However, prior balancing does take on a physical meaning with the entropic prior. Taking the negated logarithm of eqn. 3 with the entropic prior $P_e(\theta)$, we obtain

$$-\log \tilde{F}(\theta|X,T,T_0) \geq F = E - (T-T_0)H(\theta) - \log \delta(T)$$

(5)

where $E = -\log P(X|\theta)$ is the error or energy cost of a given parameterization, and $T$ is understood as temperature. The notation $\equiv$ indicates equality assuming additive constants (normalizing terms) that have been omitted. $\tilde{F}$ is an upper bound on the Helmholtz free energy equation of statistical physics, with equality at $\delta(T)=1$, $T_0=0$. Maximizing the modified posterior thus minimizes the free energy, which is analogous to finding an equilibrium configuration in a complex model whose different parts compete to explain the data. In models with factorizable likelihoods and priors, the prior on each independent parameter can be balanced separately, allowing different parts of the model to mature at different rates.

Four interesting cases immediately fall out of $\tilde{F}$: 1) Iteratively re-estimating $\theta$ while $T\to0$ gives deterministic annealing (DA) [Rose et al., 1990; Miller et al., 1996; Hofmann et al., 1998], a pseudo-global optimizer for pock-marked energy surfaces. DA belongs to a family of continuation techniques that convolve (smooth) an energy surface to make it globally convex, then track the optimum as gradual deconvolution reintroduces surface texture\(^2\). The following illustration shows how tracking a local minimum leads to the global minimum of the function in the forefront:

During DA, entropy at high temperatures keeps the system from prematurely committing to nearby local optima and forces it to explore the energy surface’s large-scale

\(^2\)A statistical analogue is robust M-estimation [Huber, 1981] with a shrinking scale parameter [Li, 1996].

structure. In a hidden-variable model, this is equivalent to maximizing almost equally w.r.t. all possible hypotheses within the model (e.g., all possible paths through a hidden Markov model), then concentrating on the most promising hypotheses as the temperature declines. In §5 we’ll introduce MAP estimators that fold DA into EM at no additional computational cost. Note that our modified posterior gives a useful amendment to DA—an automatic annealing schedule that tracks the quality (versely, entropy) of the model via the gradients or MAP estimates w.r.t. $T$.

The remaining cases of interest arise out of different terminal temperatures: 2) At $T-T_0=1$ we obtain a maximum-entropy solution. 3) At $T-T_0=0$ we obtain the maximum-likelihood (ML) solution. 4) At $-Z=T-T_0=-1$ we obtain the maximum structure solution corresponding the the entropic prior in eqn. 2. Conveniently, we have derived a single MAP estimator for all four cases.

5 MAP estimators

We obtain MAP parameter estimators by solving for maxima of the log of the balanced posterior,

$$\hat{\theta} = \arg\max_{\theta} \left[ \log P(X|\theta) - ZH(\theta) \right]$$

(6)

Although this can lead to systems of transcendental equations, we have obtained solutions for most simple distributions and, by subadditivity principles, for all models composed thereof.

For the multivariate Gaussian distribution with mean $\mu$ and covariance matrix $K$ the entropy is \(\frac{1}{2} \log(2\pi)^{N/2} |K|\). The entropic prior is thus $P_e(\mu,K) \propto |K|^{-3/2}$, which is uniform in $\mu$ and inversely proportional to the volume of $K$. To find the entropic MAP estimator we set the gradient of the log-posterior to zero. Without loss of generality we assume a zero mean to simplify the derivation:

$$0 = \frac{d}{dK} \log \left( \prod_{n} \mathcal{N}(x_n;K) \right) P_e(K)^2$$

$$= \frac{1}{dK} \left[ \sum_{n} \left[ -\frac{1}{2} x_n^T K^{-1} x_n - \frac{1}{2} \log(2\pi)^{N/2} |K| \right] ight]$$

$$-Z \frac{d}{dK} H(\mathcal{N}(K))$$

$$= \frac{1}{dK} \left[ \sum_{n} \left[ -x_n^T K^{-1} x_n - \log |K| - Z \log |K| - c \right] ight]$$

$$= \frac{1}{2} \sum_{n} \left[ 2K^{-1} x_n x_n^T K^{-1} - K^{-1} x_n x_n^T K^{-1} \circ I \right]$$

\(^3\)We thank the anonymous reviewers for pointing out work by [Ukeda and Nagano, 1995], who developed this special case directly via maximum entropy considerations, albeit without estimators or annealing schedule.
\[-\frac{1}{N} \sum_{n=1}^{N+Z} 2K^{-1} - K^{-1} \circ I \]
\[= \sum_{n=1}^{N} K^{-1} x_n x_n^T K^{-1} - \frac{N}{N+Z} K^{-1} \]
\[-\frac{1}{N} \sum_{n=1}^{N} (K^{-1} x_n x_n^T K^{-1} - \frac{N}{N+Z} K^{-1}) \circ I \quad (11)\]

Left- and right-multiplying by \( K \) reveals the entropic MAP estimator for covariances, which is essentially an \( N+Z \) normalization of the scatter of \( N \) samples:
\[K = \frac{\sum_{n=1}^{N} x_n x_n^T}{N+Z} \quad (12)\]

Note that the maximum structure (\( Z=1 \)) MAP estimator is the best mean squared-error estimate, while the maximum-entropy (\( Z=\infty \)) MAP estimator is the best unbiased estimate. Similarly normalized estimators give the scale parameters of related distributions, e.g., exponential, Laplace, and gamma.

A multinomial distribution has entropy \( H(\theta) = - \sum_i \theta_i \log \theta_i \) and entropic prior \( P_0(\theta) \propto \theta^\alpha \prod_i \theta_i^{\beta_i} \). For the MAP estimator given a vector \( \omega \) of evidence for each alternative, we set the derivative of the log-posterior to zero, using a Lagrange multiplier to ensure \( \sum_i \theta_i = 1 \),
\[0 = \frac{\partial}{\partial \theta_i} \left( \log \prod_i \theta_i^{\omega_i+Z\theta_i} + \lambda \sum_i \theta_i \right) \]
\[= \frac{\omega_i}{\theta_i} + Z \log \theta_i + Z + \lambda \quad (14)\]

We solve this a system of simultaneous transcendental equations for \( \theta_i \) using the Lambert \( W \) function [Corless et al., 1996], an inverse mapping satisfying \( W(y)e^{W(y)} = y \) and therefore \( \log W(y) + W(y) = \log y \).

Setting \( y = e^{\theta} \) and working backwards towards eqn. 14,
\[0 = -W(e^{\theta}) - \log W(e^{\theta}) + x \quad (15)\]
\[= \frac{-1}{W(e^{\theta})} - \log W(e^{\theta}) + x + \log q - \log q \quad (16)\]
\[= \frac{-q}{W(e^{\theta})} + \log q/W(e^{\theta}) + x - \log q \quad (17)\]

Setting \( x = 1 + \lambda/Z + 4\log q \) and \( q = -\omega_i/Z \), eqn. 17 simplifies to \( Z \times \lambda \times \theta_i \):
\[0 = \frac{\omega_i}{W(-\omega_i e^{1+\lambda/Z} Z)} + \log W(-\omega_i e^{1+\lambda/Z} Z) + 1 + \lambda/Z \quad (18)\]

which implies that
\[\theta_i = \frac{-\omega_i/Z}{W(-\omega_i e^{1+\lambda/Z} Z)} \quad (19)\]

where eqn. 14 and eqn. 19 form a set of fix-point equations for \( \lambda \) that typically converge in 2-5 iterations. This derivation generalizes that given in [Brand, 1997a]. Details on computing \( W \) are given in [Brand, 1997b].

Differentiating eqn. 14 again we find that the posterior is concave in any parameter provided \( \omega_i > Z \theta_i \), which is always true except for near-degenerate cases when \( Z > \max_i \omega_i \) (e.g., there is almost no evidence).

Note that we have assumed that the likelihood and the prior are factorizable, which is fortunately the case for many popular hidden-variable models.

### 6 Minimization of entropies

For flat exponential distributions, the principle of maximum likelihood is equivalent to minimizing cross-entropy (directed Kullback-Leibler distance) between the data’s sufficient statistics and the distribution’s estimated parameters. Our framework identifies three entropies associated with modeling, and the MAP estimator minimizes their sum. This is most clearly seen by considering the negated log-posterior of a multinomial parameter in the maximum structure (\( Z=1 \)) case:
\[-\log P(\omega | \theta) e^{-H(\theta)} = -\log \prod_i \theta_i^{\omega_i+\theta_i} \quad (20)\]
\[= -\sum_i (\omega_i + \theta_i) \log \theta_i \quad (21)\]
\[= -\sum_i (\omega_i \log \theta_i + \theta_i \log \theta_i - \omega_i \log \omega_i + \omega_i \log \omega_i) \]
\[= -\sum_i \omega_i \log \theta_i + \sum_i \omega_i \log \frac{\omega_i}{\theta_i} - \sum_i \theta_i \log \theta_i \quad (23)\]
\[= H(\omega) + D(\omega || \theta) + H(\theta) \quad (24)\]

Each of these entropies has a useful interpretation: \( H(\omega) \) measures uncertainty in the expected sufficient statistics, and is linearly related to their coding length. The cross-entropy \( D(\omega || \theta) \) measures error in the model’s fit to the data, and is linearly related to the coding costs of aspects of the data not captured by the model. (Ideally, this will dwindle to purely random noise.) Together, these terms give the expected coding length of the data relative to the model. \( H(\theta) \) measures uncertainty in the model’s component distributions, as well as its coding length. In short, MAP estimation reduces all entropies and relative entropies between the model and the data’s sufficient statistics. In practice, it does so by extinguishing parts of the model that are ill-matched to the structure of the data, and therefore weakly supported. More evidence concentrates on the surviving parameters (equivalently

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4In these cases we may either use slightly more elaborate methods to find the global optimum or simply accept a local optimum and ignore its typically negligible effect on the joint distribution.
$H(\omega)$ is minimized, and their information content is thereby maximized.

### 6.1 Variations

One may introduce other entropies into eqn. 24, perhaps to symmetrize the cross entropy (equiv. KL divergence)

$$H(\omega) + D(\omega||\theta) + D(\theta||\omega) + H(\theta).$$  \hspace{1cm} (25)

One might want to conserve information contained in previous estimates

$$H(\omega) + D(\omega||\theta) + D(\theta||\omega_{\text{est}}) + H(\theta),$$  \hspace{1cm} (26)

or stay close to (or far from) a reference model $\theta^*$

$$H(\omega) + D(\omega||\theta) + D(\theta||\theta^*) + H(\theta).$$  \hspace{1cm} (27)

Happily, small variations on the MAP formula given above can accommodate any combination of these constraints.

For simplicity of exposition we have identified $H(\theta)$ with the entropy of the model’s component distributions, leading to an extremely efficient form of description length minimization. One may also choose other entropy measures for $H(\theta)$. For example, one may prefer to minimize the joint entropy

$$H(X,\theta) = H(X) + H(\theta|X) = c + \sum_j p_j \sum_k \theta_{jk} \log \theta_{jk}$$  \hspace{1cm} (28)

where $H(X) = c$ is fixed and $p_j$ are the probabilities of hidden variable values given the data. These are estimated from the data during the E-step of EM, and in some cases can be computed from the parameters (e.g., in Markov models, $p_j$ is the stationary probability of state $j$). This case leads to another small variation on the MAP estimator, etc., but the result is only approximate because the entropy is minimized w.r.t. $p_j$ calculated from the previous rather than the current parameter estimates. Nonetheless, we find in practice that this case is very well behaved, especially in deterministic annealing, where $p_j$ changes quite gradually.

### 7 Trimming

One problem with entropy minimization on digital computers is that numerical error often intrudes before parameters are fully extinguished. Eqn. 14 obliges us to add numbers to their logarithms, which becomes troublesome when parameter values are not near 1. Fortunately, the prior allows us to identify excess parameters that can be trimmed from a model without loss of posterior probability, such that

$$\hat{p}_j(\theta_d|X,Z) \geq \hat{p}_j(\theta|X,Z)$$  \hspace{1cm} (29)

Expanding via Bayes’ rule, taking logarithms, and rearranging, we obtain

$$Z[H(\theta) - H(\theta|\theta_i)] \geq \log P(X|\theta) - \log P(X|\theta_i)$$  \hspace{1cm} (30)

Operationally, a parameter is trimmed by setting it to a target value which makes the model simpler or easier to interpret. Typically the target will be the minimal entropy extinction value, e.g., setting a multinomial parameter $\theta_i \leftarrow 0$, but it may simply be a usefully interpreted value, e.g., setting a variance parameter $\kappa_{ij} \leftarrow 1$ (equivalently $\log \kappa_{ij} \leftarrow 0$). When zeroing, eqn. 30 can be approximated via differentials, yielding

$$Z \frac{\partial H(\theta)}{\partial \theta_i} \theta_i > \theta_i \frac{\partial \log P(X|\theta)}{\partial \theta_i}$$  \hspace{1cm} (31)

One may observe from this that a parameter can be trimmed when varying it increases the entropy faster than the log-likelihood. The sides of eqns. 30 and 31 can be mixed and matched to obtain mathematically convenient forms. For the multinomial trim test, we set l.h.s. eqn. 30 ($-Z\theta_i \log \theta_i$) against r.h.s. eqn. 31 to obtain:

$$\theta_i < \exp \left[ \frac{1}{Z} \frac{\partial \log P(X|\theta)}{\partial \theta_i} \right]$$  \hspace{1cm} (32)

The gradient of the log-likelihood is normally computed for re-estimation, so these trimming tests are very cheap.

Solving eqn. 30 via W gives the variance trimming criteria:

$$k_{ij} \geq -\frac{k_{ii}}{W_{ij}}(-\frac{k_{ij} \exp k_{ij}}{\log \kappa_{ij}}) \text{ iff } \kappa_{ij} > 1$$  \hspace{1cm} (33)

$$k_{ij} \leq -\frac{k_{ii}}{W_{ij}}(-\frac{k_{ij} \exp k_{ij}}{\log \kappa_{ij}}) \text{ iff } \kappa_{ij} < 1$$  \hspace{1cm} (34)

There is also a trimming test for covariances ($k_{ij} \leftarrow 0$), but it is beyond the scope of this paper.

Trimming is a local heuristic that increases but does not maximize the posterior. It makes sense in iterative estimation of models containing hidden variables because it can radically sparsify the model, which in turn reduces ambiguity in the data’s expected sufficient statistics. In short, it accelerates learning. In our experiments we have used trimming mainly to speed parameter extinction that would otherwise happen gradually via MAP estimation. As noted above, this becomes particularly desirable when parameters are near extinction and floating-point calculations introduce numerical round-off and underflow errors. In addition, trimming near convergence can “bump” the model out of the local probability maximum and into a parameter subspace of simpler geometry, thus enabling further training.

Not only do parameter extinction and trimming protect against over-fitting, but by sculpting the model to fit the data, they often reveal a simple machine that explains the data (e.g., a finite-state machine from an HMM: a circuit from a NN). We shall give many examples in $\exists \theta$.

### 8 Related ideas

Model selection criteria such as MML [Wallace and Boulton, 1968; Wallace and Freeman, 1987], AIC
Consequences; the figure below contrasts the discontinuous on discretization of continuous values has some severe with probability close to 1. Unfortunately, MDL based it is MDL in the strong sense of [Vitányi, 1978] it minimizes stochastic complexity and, remarkably, that derived a binomial parameter estimator and showed that to approximate estimators and occasionally yielding a point estimator for discrete sample spaces. E.g., [Vovk, 1995] natural interpretations at a bit-level of granularity, leading to approximate estimators and occasionally yielding a point estimator for discrete sample spaces. E.g., [Vovk, 1995] derived a binomial parameter estimator and showed that it minimizes stochastic complexity and, remarkably, that it is MDL in the strong sense of [Vitányi and Li, 1997], with probability close to 1. Unfortunately, MDL based on discretization of continuous values has some severe consequences; the figure below contrasts the discontinuous Vovk estimator with equivalent entropic MAP estimator.

Computer scientists have tried to get more leverage out of model selection criteria by proposing search heuristics that reduce the squandered computation that dominates generate-and-test “structure-learning” algorithms. [Ikeda, 1993] and [Stolcke and Omohundro, 1994] presented search algorithms for HMM structure that generate rival models by splitting or merging states that are likely to reduce description length. Whereas run-times were reported in days, entropic estimation will induce HMM topologies from similar datasets in a matter of minutes. [Friedman, 1998] advertised a “Structural EM” algorithm for Bayesian networks, but the M-step is approximate and there is a penalized generate-and-test search inside the EM loop. We would propose entropy minimization algorithm for Bayesian networks in which parameter extinction makes subgraphs deterministic and thus trimmable. Unfortunately the E-step for dense graphs can be computationally prohibitive, which makes starting with over-complete structure problematic.

Most exponential density forms have conjugate priors that can be used to obtain estimators with some description-length reducing properties. For example, for multinomial parameters one might use Dirichlet priors with exponents $\alpha_i < 1$. Of course, one must choose values for these parameters a priori; this is equivalent to inventing extra observations before any actual samples have been taken, a potentially dangerous business.

9 Examples

We have used entropic estimation to obtain low entropy models in a variety of model classes. These have been tested on benchmark datasets as well as real-time data obtained directly from computer vision and speech analysis systems. The resulting models have consistently been smaller, more discriminative, better generalizing, and more predictive than conventionally (e.g., ML) estimated models, except when both are under-fit. Typically the resulting model is interpretable, often providing insight into the causal structure of the process that generated the data. Here we give a variety of examples

9.1 Mixture models

A mixture model was fitted to an ring of Cartesian samples taken from a uniform distribution over the polar coordinates $r \in [1, 2]; \theta \in [0, 2\pi]$. The figures show the model estimated conventionally, entropically with trimming, and entropically with trimming and deterministic annealing. Initial conditions were identical for all three cases. Ellipses indicate iso-density contours for the Gaussian components.

In the first case, over-fitting has resulted in a model of the accidental properties of the data (e.g., the clumpiness of the sample). In the second case, trimming has removed excess components and the resulting model looks much more like the essential structure of the data, but a large under-sampled region near the top still affects the model. In the third case, DA circumvents this local optimum and finds a model which generalizes even better.
9.2 Radial basis function networks (RBFNs)

Vowel classification: We obtained the British English vowel recognition dataset from the CMU Neural-Bench Archive. Each example of a vowel is characterized by LPC coefficients from two time-frames. This dataset has been treated several times using perceptrons, neural networks, Kanerva models, radial basis function networks, and non-parametric methods (nearest neighbor). The last yielded the best classification of the test set (56%), while RBFNs peaked at 53% with 528 Gaussian basis functions [Robinson, 1989]. We combined entropically estimated mixture models of each vowel’s training data into an RBFN and obtained 58.7% correct classification on the test set. Entropic estimation found mixtures of 1-3 components per vowel, resulting in an RBFN of only 22 basis functions.

Home value prediction: We obtained the publicly available Boston home value database from StatLib and set out to predict the median home value from 13 other potentially relevant attributes. The features are numeric or boolean-valued and have different (arbitrary) dynamic ranges; we normalized each dimension to unit variance before mixture modeling. In each trial, all models were identically initialized and trained on half of the examples, randomly selected, then used to predict home values for the remaining test examples. We use the performance of the ML model as a baseline in each trial; our measure of performance is how much an entropically estimated model reduced the mean squared-error of predictions on the test set. The following table shows mean improvement in RBFN prediction accuracy over maximum-likelihood models, as a function of # of RBFs at initialization.

<table>
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<th>entropic MAP</th>
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<th>+annealing</th>
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</tbody>
</table>

All values are at a $p<10^{-5}$ level of statistical significance except for those marked with a ‘↑’ ($p<0.01$) or a ‘*’ ($p<0.05$). The improvement is quite significant given a large initialization (recall that these scores are in terms of unit-variance prices). This can be attributed to entropic estimation’s resistance to over-fitting. The advantage disappears or becomes statistically insignificant when the initializations are too small, but notice that even in these under-fitted models, annealing is finding superior local optima.

9.3 Hidden Markov models (HMMs)

Handwriting analysis and classification: We obtained handwriting samples by 10 writers from the UNIPEN archive. The diagrams below show entropic and maximum-likelihood models of the pen-strokes for the digit “5,” estimated from pen-position data taken at 5msec intervals from 10 different individuals via an electro-magnetic resonance sensing tablet. Initial conditions were identical.

Ellipses indicate show iso-density contours for each state; ‘x’ and arcs indicate state dwell and transition probabilities, respectively, by their thicknesses. Entropic estimation induces an interpretable automaton that captures essential structure and timing of the pen-strokes, as well as variations in their ordering between writers. 50 of the original 80 dynamical parameters were trimmed. Estimation without the entropic prior results in a wholly opaque model, in which none of the original dynamical parameters were trimmed. Over 10 trials with different parts of the database held out, entropic models yielded 96% correct classification; ML models yielded 93%.

Video analysis: In this example a gigabyte of video randomly taken from an office setting was filtered to extract motion vectors and then modeled entropically with a hidden Markov model [Brand, 1997a]. The resulting HMM state machine is compact enough to read:

9
8
7
6
5
4
3
2
1

Roughly 2/3 of the transitions were trimmed. We have taken the liberty of labeling states by using forward-backward analysis to find their support in new video. The states mapped nicely to typical work activities of the office occupant. An HMM conventionally estimated from the same initial conditions is fully connected and thus too bushy to profitably illustrate or interpret.

Prediction in Bach chorales: We obtained a dataset of melodic lines from 100 of J.S. Bach’s 371 surviving
chorales from the UCI repository [Merz and Murphy, 1998], and transposed all into the key of C. We compared
entropically and conventionally estimated HMMs in prediction and classification tasks, training from identical
random initial conditions and trying a variety of different
initial state-counts. We trained with 90 chorales and testing
with the remaining 10. In ten trials, all chorales were
rotated into the test set. The results neatly chart the
sparsification, classification, and prediction superiority of
entropically estimated HMMs.

9.4 Recurrent neural networks (RNNs)

In this rather speculative example, an entropic prior was
designed for the weights of a recurrent neural network by
assuming that $\Theta$ generates neural networks with weights
from a Gaussian distribution with functionally related mean
and variance. This yields a modified back-propagation
rule and a trim test. The figures show a semi-recurrent
neural network (a DAG with self-connections for memory;
activation propagates one link per cycle) trained to compute
XOR using conventional back-prop with weight decay
(left) and entropic back-prop with trimming (right). Initial
conditions were identical.

Note that the entropically estimated topology is an
amalgam of the two minimal feed-forward XOR circuits.