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Solving the inverse Ising problem by mean-field methods in a clustered phase space with many states

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Abstract – In this work we explain how to properly use mean-field methods to solve the inverse Ising problem when the phase space is clustered, that is many states are present. The clustering of the phase space can occur for many reasons, e.g. when a system undergoes a phase transition. Mean-field methods for the inverse Ising problem are typically used without taking into account the eventual clustered structure of the input configurations and may lead to very bad inference (for instance in the low temperature phase of the Curie-Weiss model). In the present work we explain how to modify mean-field approaches when the phase space is clustered and we illustrate the effectiveness of the new method on different clustered structures (low temperature phases of Curie-Weiss and Hopfield models).

Introduction. – The Ising inverse problem has been the subject of a large amount of works very recently \cite{1-4}. Although this problem is known since many decades under the name of Boltzmann machine learning (BML), many recent applications and developments in different fields (e.g. biology \cite{6-8}, computer science \cite{9} and physics \cite{10-12}) have renewed the interest in studying such problems. The BML can be investigated under two very different approaches. In the first one, which concerns this work, a set of data is generated according to the Gibbs-Boltzmann measure of a generic Ising model. The input data for the inverse problem are therefore independent and distributed accordingly to the Boltzmann distribution of the system \cite{13}. In a second case, the data are generated according to a stochastic dynamical process which correlates configurations close in time, and this correlations in the input data are exploited in solving the inverse problem \cite{14}. In both cases, the traditional Bayesian approach consists in maximizing the likelihood function of the data. In this work, we focus on the first case which is commonly named “static inverse Ising problem” and is harder than the second case.

In the static case, maximizing the likelihood is a complicated task, because it directly depends on the partition function which is impossible to compute efficiently (in the general case, its complexity grows exponentially with the system size). However, it is still possible to maximize the likelihood by the expectation-maximization method using a Monte-Carlo algorithm and doing a Boltzmann learning procedure \cite{13}. The Monte Carlo is used to evaluate the average value of the observables of the system (here the magnetizations and the correlations) and to update the value of the magnetic fields and the couplings by doing a gradient ascent. Yet, it is known that Monte Carlo estimates do not converge quickly in many cases and may require many steps to obtain accurate mean values. It means that the MC should be run for a long time at each step of the BML procedure making the method quite slow. For this reason, faster methods based on mean-field approximations are commonly used in practical applications.

In a recent work \cite{11} Nguyen and Berg have revisited the problem of finding a good mean-field (MF) approximation for the inverse Ising problem. It was already known that MF methods fail to provide a good couplings reconstruction at low temperatures even for ferromagnetic systems (see Fig. \cite{1} for an example on a ferromagnet and \cite{15} for an example on a MF spin glass). Worst than that, this problem in coupling reconstruction occurs also in cases
where the MF approximation is exact in the thermodynamical limit (e.g. the Curie-Weiss model). This failure in reconstructing couplings in ferromagnetic systems can be understood by looking at the input configurations at low temperatures: below the ferromagnetic transition, indeed, configurations are clustered in two groups of respectively positive and negative magnetization. The naive MF (nMF) approximation is based on the self-consistency equations for the magnetizations, \( m_i = \tanh(\beta \sum_j J_{ij} m_j) \), with \( \beta \) being the inverse temperature, which have 3 solutions for \( \beta > \beta_c \): it is well known that the \( m_i = 0 \) solution is unphysical, while the two solutions with \( m_i \neq 0 \) are thermodynamically stable. However considering all the input configurations together the average magnetizations are zero by symmetry. Therefore, a naive use of MF equations infer the couplings using the unphysical \( m_i = 0 \) fixed point, and lead to a very poor result. Please notice that the same problem arises if one computes correlations in a naive way: using all input data connected correlations would not decay at long distance. Therefore, in order to use properly the nMF equations, it is mandatory to look at the two other solutions characterized by non-zero magnetization. These solutions arise naturally when considering the decomposition of the Gibbs-Boltzmann measure in the configuration space.

The authors of Ref. [11] consider the nMF equations for both states (of positive and negative magnetizations) at the same time. In this way they obtain an over-constrained system of linear equations to be solved. They manage to find a solution by using the pseudo-inverse of a matrix (see [11] for further details). We will see that this approach can be considerably simplified in the case of the Curie-Weiss (CW) model, and then generalized to models with many free-energy minima. In Ref. [11] also the case of the Sherrington-Kirkpatrick (SK) model is considered as a case study with a clustered phase space at low temperatures. We would like to emphasis, however, that the division in metastable states of the SK model is somehow problematic for this approach. The metastable states of the SK model in the glassy phase are highly non-trivial and therefore it is very difficult even to define them properly in a system of limited size. Therefore we claim that the inference algorithm of Ref. [11] as well as the one presented in the present work are not suitable for this kind of models, for which more elaborate techniques (such as the pseudo-likelihood method [16,17]) are required.

In the present work we show that couplings can be well inferred using nMF equations also in the low temperature phase if input configurations are previously clustered and the nMF inference algorithm is applied separately to data in each cluster. We show that our inference procedure based on solving the nMF or TAP equations inside each cluster separately is much simpler than the method proposed in Ref. [11], where self-consistency equations for each cluster need to be solved simultaneously. Therefore the use of complicated numerical algorithms such as the pseudo-inverse is not necessary. In addition, we show that, at variance to what is claimed in Ref. [11], using the present inference procedure one does not estimate wrongly the magnetic fields. It is worth mentioning that, when using one of the MF fixed points with \( m_i \neq 0 \), a spurious magnetic field unavoidably appears due to errors on the inferred couplings. However this magnetic field is very small and decreases when increasing the number of input data.

In order to prove that our method is very efficient we apply it to different kind of models. First we show that in the CW model the results are as good as those from more elaborated methods, like the pseudo-likelihood method. Then we focus on the Hopfield model where the number of different free-energy minima can be controlled and made larger. We show that it is possible to improve the results on the inference process by clustering the set of input configurations and to infer the right number of clusters to be used. We should mention that a previous attempt to infer the couplings in the (sparse) Hopfield model from data collected in a single state was done in [18]. However, in that work, the interaction network was assumed to be known and only couplings intensities were inferred, so a direct comparison with our results is not possible.

**Problem definition and inference algorithms.**

In the static inverse Ising problem one aims at inferring the value of the couplings between the variables and the eventual magnetic fields, given a set of \( M \) equilibrium configurations. More precisely we consider an Ising model with \( N \) spins defined by the Hamiltonian

\[
\mathcal{H}(s) = -\sum_{i<j} J_{ij} s_i s_j - \sum_i h_i s_i ,
\]

where \( i,j = 1, ..., N \). In the static case, the inference process is done by using input data distributed according to the Gibbs-Boltzmann measure

\[
P_{\text{GB}}(s) = \frac{e^{-\beta \mathcal{H}(s)}}{Z} \text{ where } Z = \sum_s e^{-\beta \mathcal{H}(s)} \quad (2)
\]

We remind here that the \( M \) sampled configurations are assumed to be independent.

In the following we will consider two different families of inference methods. For mean-field methods, we shall consider the average magnetizations and correlations of the data

\[
\bar{m}_i = \frac{1}{M} \sum_{a=1}^{M} s_i^a \quad (3)
\]

\[
\bar{c}_{ij} = \frac{1}{M} \sum_{a=1}^{M} s_i^a s_j^a \quad (4)
\]

These observables are the only information needed to infer the parameters of the models when using mean-field methods. We will also consider the pseudo-likelihood methods for which the entire sampled configurations \( \{s_i^a\} \) are
needed. Let us now describe how these methods work and how we will use them in the context of a clustered phase space.

We first consider the naive mean-field approach where the equations can be simply derived by considering the solution of the Curie-Weiss model (where \( J_{ij} = 1/N \)). For this model, the magnetisations and the correlations are given by

\[
m_i = \tanh \left( \beta \sum_{j \neq i} J_{ij} m_j + \beta h_i \right)
\]

(5)

By inverting eq. (6) we can reconstruct directly the couplings \( J_{ij}^\ast \). Then, by using the \( J^\ast \) and eq. (5) we can infer the magnetic fields \( h_i^\ast \)

\[
J_{ij}^\ast = -(e^{-1})_{ij} + \frac{\delta_{ij}}{\beta(1 - m_i^2)}
\]

(7)

\[
h_i^\ast = -\beta^{-1} \left[ \tanh(m^i) - \sum_{j \neq i} J_{ij}^\ast m_j \right]
\]

(8)

We refer to this method as nMF in the rest of the article.

A second approximation commonly used is to consider the pseudo-likelihood method (PLM). PLM is based on the maximization of the marginals probability of one spin \( s_i \) given that the rest of the spins are fixed: \( p(s_i|s_{\bar{i}}) \). The PLM consists in maximizing the sum of all the log-pseudo-likelihood

\[
P_L = \frac{1}{NM} \sum_{i,a} \log(p(s_i^a|s_{\bar{i}}^a))
\]

(9)

In this method, we need to have access to all the configurations \( \{s_i^a\} \). The advantage of this method is that it deals also with high order correlations and thus provides much better performances on finite dimensional systems, but it also can handle directly clustered phase space. Moreover it has a polynomial complexity at variance to using the true likelihood of the data.

Clustering methods and inference with clustered phase space. Here we describe the clustering algorithms that we use to divide configurations in clusters before applying the nMF method. These clustering algorithms group configurations together based on their distances: configurations are put in the same group if they are “close” enough and “far” from the other clusters, where the concepts of “close” and “far” usually need to be determined in a self-consistent way. We use the Hamming distance defined by \( d_{ab} = 1/(4N) \sum_i (s_i^a - s_i^b)^2 \). In the present work we use two different clustering methods. First we consider the soft \( K \)-means clustering. This method clusterizes the space of configurations by assigning each configuration to the closest of the \( k \) centers “softly” (a configuration is assign to a center with a given probability). Then the position of the \( k \) centers is updated accordingly to the position of the configurations inside each cluster. The procedure is repeated until convergence. This method is very fast, the complexity scale as \( O(M) \), but the results can depend strongly on the initial conditions (i.e. on how the \( k \) centers are chosen at the beginning).

A second method is based on density clustering. The density clustering algorithm we consider first defines the density around each point. In our case the density is the number of configurations within a given range. Then, each data point is associated to its closest neighbour with higher density. This process naturally separates the phase space into a number of clusters which depend on the range used for defining the neighborhoods. Therefore by using this algorithm we do not need to specify the number of clusters. Thus this second clustering algorithm has the advantage of finding by itself the number of clusters. It suffers however of a larger complexity, scaling as \( O(M^2) \).

After clustering the configurations we have to used them properly to infer the parameters of the model. We define the observables of the \( k \)-th cluster by

\[
\bar{m}_{i}^{(k)} = \frac{1}{M_k} \sum_{a \in C_k} s_i^a
\]

(10)

\[
\bar{c}_{ij}^{(k)} = \frac{1}{M_k} \sum_{a \in C_k} s_i^a s_j^a
\]

(11)

where \( C_k \) is the set of indices of configurations belonging to the \( k \)-th cluster and \( M_k = |C_k| \). We now apply the nMF equations separately for each cluster and obtain a different estimate of the parameters for each cluster \( J_{ij}^{(k)} \). Finally, to obtain the best estimate for the couplings we take the weighted average of all the different estimates

\[
J_{ij} = \frac{1}{M} \sum_{k} M_k J_{ij}^{(k)}
\]

(12)

To estimate the magnetic field, we first compute them within each cluster: \( h_i^{(k)} \) is obtained from eq. (8) with the estimates \( J_{ij}^{(k)} \). The final estimate for the magnetic fields is again given by the weighted average over the clusters

\[
h_i^{\ast} = \frac{1}{M} \sum_{k} M_k h_i^{(k)}
\]

(13)

Results on the Curie-Weiss model. – The Curie-Weiss (CW) model is a fully connected ferromagnet with \( J_{ij} = 1/N, \forall i \neq j \). The model has a paramagnetic phase (\( m_i = 0 \)) at high temperature \( \beta < \beta_c = 1 \) and a ferromagnetic phase (\( m_i \neq 0 \)) above \( \beta_c \). In the ferromagnetic phase, two states of positive and negative magnetizations coexist. In the limit of very large system sizes \( (N \to \infty) \) magnetizations and correlations can be computed analytically by eqs. (50), which are exact up to \( O(1/N) \) corrections. It means that, by using eqs. (78) one should obtain the best possible estimate of the parameters \( J_{ij} \)
and $h_i$, but in the ferromagnetic phase, only the solution
with non-zero magnetization of the eq. (5) should be con-
sidered (as discussed in the Introduction). We evaluate
how the following three inference algorithm perform
in the estimate of couplings in the CW model: (i) the
nMF method used naively, without clustering the con-
figurations; (ii) the nMF method on configurations clustered
using two clusters; (iii) the PLM on the original con-
figurations.

In Fig. 1 we report the error achieved by different meth-
ods in the temperature range $\beta \in [0.1, 2]$ with $M = 10^4$
and $M = 10^5$ in inferring the couplings using the following
definition

$$
\varepsilon^2 = \frac{2}{N(N-1)} \sum_{i<j} (J_{ij} - \hat{J}_{ij})^2
$$

(14)

For $\beta < \beta_c$, the paramagnetic fixed point is correct
and therefore the reconstruction achieve by nMF is the
best possible. However, for $\beta > \beta_c$ the nMF error (red
curves) suddenly raises, because the $m_i = 0$ fixed point
is no longer the physical one. On the contrary, using the
nMF method on the data clustered with exactly 2 clusters
(green curves), provides a small error in the ferromagnetic
phase, but fails badly in the paramagnetic phase. The
inference methods that provide the best estimate in the
whole temperature range are the PLM (blue curves) and
the nMF with data clustered via density clustering (purple
curve), that automatically split the input data in one or
two clusters, depending on symmetries in the input data.
It is worth stressing that these two methods have essen-
tially the same error at any temperature: that is even the
nMF approximation provides the best possible estimates
if applied to properly clustered data.

In Fig. 1 we show results obtained with $M = 10^4$ and
$M = 10^5$ in order to make evident whether the uncertain-
ties in the couplings estimates are due to the noise in the
input data or to an intrinsic limitation of the inference al-
gorithm. For example deep in the ferromagnetic phase the
nMF method has an error decreasing only slightly when
$M$ increases, because the error is mainly due to a limita-
tion of the method. On the contrary, PLM and nMF with
properly clustered data provide a result whose uncertainty
is mainly due to noise in the input data: indeed the error
decreases as $1/\sqrt{M}$.

To confirm the correctness of the inference algorithm
based on data clustering and nMF equations, we also
looked a the inferred value of the magnetic field by using
eqs. (8) and (13). We see clearly in the insets of Fig. 1
that, in the low temperature phase, the clustering+nMF
method does not predict any anomalously large magnetic
field, thanks to the fact that, clustering the input data,
we are actually using the magnetized solutions of eq. (5).
In our numerical experiments, we have found too large in-
ferred magnetic fields only if either system size was too
small or the input data were too noisy: in the former case
the problem resides in the fact eq. (5) is crudely approxi-
mate, while in the latter case it is a consequence of large
errors in couplings reconstruction.

Results on the Hopfield model. — We now extend
our analysis to a more complicated case by considering
the Hopfield model. The Hopfield model has been intro-
duced long time ago [22] to model neural networks: it is
a fully-connected Ising model, whose couplings can be
chosen such that the model free-energy has $2P$ different
minima (that act has attractors for the pattern recovery
dynamics). In some sense, the Hopfield model can be seen
as a generalization of the Curie-Weiss model, which is in-
deed equivalent to the $P = 1$ case. We are interested in
studying the inverse Ising problem in the Hopfield model,
because configurations sampled at low temperature in the
Hopfield model are typically clustered around the $2P$ free-
energy minima: consequently naive MF methods face even
more severe limitations than in the low temperature phase
of the CW model, and we want to study how much MF
methods for the inverse Ising problem can be improved by
clustering input configurations.

The Hamiltonian of the Hopfield model reads

$$
\mathcal{H}(\sigma) = -\frac{1}{N} \sum_{ij} \frac{1}{P} \sum_{\alpha=1}^P \xi^\alpha \eta^\alpha s_i s_j
$$

(15)

where the $P$ patterns $\xi^\alpha$ identify the directions of the free-
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Fig. 2: Main panels: errors in inferring couplings in Hopfield models with \( P = 2 \) uncorrelated (top), correlated (center) and anti-correlated (bottom) patterns. The comparison is between MF methods with clustered data (either \( K \)-means or density clustering) and PLM. In the inset of the top panel, we show that the likelihood of the clustering algorithm suggests to take one cluster below \( \beta_c \approx 1.1 \) and 4 clusters above \( \beta_c \). In the inset of the bottom panel we show the magnetic field inferred by nMF+clustering, which is very small in both phases.

In the standard Hopfield model, the \( \xi \)'s are drawn from the bimodal distribution, that is \( \xi^\alpha = \pm 1 \) with probability 1/2 independently. In our study we also consider the case where the pattern \( \xi \) are correlated by setting 10% of their components equal (\( \xi^\alpha = \xi^\beta \) \( \forall \alpha, \beta \)), and anti-correlated (only when \( P = 2 \)) by setting 10% of their components in an opposite way (\( \xi^\alpha = -\xi^\beta \)). This model presents a paramagnetic phase at high temperature, and an ordered phase at low temperature defined by the states around the patterns \( \{ \xi \} \) if the number of patterns is not too high [23]. The ordered phase is characterized by a Gibbs-Boltzmann measure clustered around one of the \( 2P \) available states (for a given \( P \) there will be \( 2P \) stable states in the low temperature region due to the spin flip symmetry).

We show now our results on inferring the Hopfield couplings by using MF methods on clustered data. In Fig. 2 we consider systems with \( N = 100 \) spins, \( P = 2 \) (therefore 4 states) in all the three possible cases (standard, correlated and anti-correlated patterns). We observed that MF methods with the right number of clusters perform similarly to the PLM, which is at present the best possible algorithm to solve the inverse Ising problem. The right number of clusters can be obtained either by density clustering or by maximizing the likelihood of the clustering obtained by \( K \)-means (see top panel inset in Fig. 2).

As in the CW model, also for the Hopfield model the magnetic fields inferred by MF methods on clustered data are very small, and independent on the eventual long range order present in the model (see inset in lower panel of Fig. 2).

In Fig. 3 we show the results on inferring couplings of Hopfield models with \( P = 3 \) patterns (and thus 6 free-energy minima). Again MF methods applied on input data clustered with the right number of clusters perform very similarly to PLM, and much better than standard MF methods applied directly to all input data. It is worth noticing that the best result by the clustering+nMF algorithm as been obtained by running the clustering procedure several times with different initial conditions (data labeled ‘many IC’ Fig. 3) and then picking the clustering having the largest likelihood. This is expected since a clustering algorithm as \( K \)-means is not very stable for large \( K \) and its outcome strongly depends on the initial condition.

Let us finally discuss the time complexity of the three algorithms we have used: PLM, \( K \)-means+nMF and dens.clus.+nMF. Regarding the system size dependence, all three algorithms have a time complexity \( O(N^3) \), either because of the inversion of a \( N \times N \) matrix in MF methods, either because of the computation of the gradient of the pseudo-likelihood (PL), which is \( O(N) \), in a space of \( O(N^2) \) variables. Their dependence on the number \( M \) of input configurations is different: PLM is linear in \( M \), but the search for the maximum of the PL, requires to compute PL and its derivatives many times; \( K \)-means is linear in \( M \), but often a search for the optimal clustering requires to run it with many different initial conditions; density clustering is \( O(M^2) \), so, although it provides a
robust result, it is impracticable when the number of samples is very high (however we are aware that the authors of Ref. [21] are developing a faster version of the density clustering algorithm). In practice, we observe it is better to use PLM when the number $M$ of input configurations is small and nMF with $K$-means clustering when $M$ becomes large.

Conclusions. – In this work we have presented a very simple way to make mean field approximations to the inverse Ising problem effective also in the low temperature phase, where symmetries get usually broken and, correspondingly, input data get clustered. The idea is to cluster the input data and to apply mean-field methods to each data cluster. We have tested this clustering+nMF algorithm on the Curie-Weiss and Hopfield models, comparing results with the most sophisticated and state-of-the-art pseudo-likelihood method.

Results are very promising and redeem mean-field approximations to inverse problems, even in those cases where the structure of the input data is such that a straightforward application of mean-field methods would be ineffective.

The natural follow-up to this work is application of clustering+nMF methods to inverse problems based on real data. It is worth remembering that often in solving inverse problems based on real and noisy data, the robustness of simple MF methods is more valuable than the putative higher accuracy of more sophisticated methods: see e.g. the case of inferring protein contacts [4]. From this point of view, enlarging the range of applicability of MF methods by data clustering is certainly very useful and maybe better than developing higher order approximations (that strongly depends on the model used to describe the data).

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