Inference and learning in sparse systems with multiple states

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We discuss how inference can be performed when data are sampled from the non-ergodic phase of systems with multiple attractors. We take as model system the finite connectivity Hopfield model in the memory phase and suggest a cavity method approach to reconstruct the couplings when the data are separately sampled from few attractor states. We also show how the inference results can be converted into a learning protocol for neural networks in which patterns are presented through weak external fields. The protocol is simple and fully local, and is able to store patterns with a finite overlap with the input patterns without ever reaching a spin glass phase where all memories are lost.

I. INTRODUCTION

The problem of inferring interactions couplings in complex systems arises from the huge quantity of empirical data that are being made available in many fields of science and from the difficulty of making systematic measurements on interactions. In biology, for example, empirical data on neurons populations, small molecules, proteins and genetic interactions, have largely outgrown the understanding of the underlying system mechanisms. In all these cases the inverse problem, whose aim is to infer some effective model from the empirical data with just partial a priori knowledge, is of course extremely relevant.

Statistical physics, with its set of well-understood theoretical models, has the crucial role to provide complex, but clear-cut benchmarks, whose direct solution is known and that can therefore be used to develop and test new inference methods.

In a nutshell, the equilibrium approach to inverse problems consists inferring some information about a system defined through an energy function $H(\sigma)$ starting from a set of sampled equilibrium configurations. Suppose $M$ i.i.d sampled configurations $S = \{\sigma^{(i)}\}_{i=1...M}$ generated by a Boltzmann distribution of an unknown energy function $H$, $P(\sigma) = \frac{1}{ZH}e^{-H(\sigma)}$ are given. The posterior distribution of $H$ (also called likelihood) is given by $P(H|S) \propto \exp(-M(\langle H \rangle + \log ZH))P(H)$, where $\langle H \rangle = \frac{1}{M} \sum_{i=1}^{M} H(\sigma^{(i)})$ represents the average of $H$ over the given sample configurations and $P(H)$ the prior knowledge about $H$. The parameter $M$ plays the role of an inverse temperature: when $M$ is very large, $P(H|S)$ peaks on the maximums (with respect to $H$) of the ‘log-likelihood’ $L = -\langle H \rangle - \log ZH$. The problem of identifying the maximum of $L$ can be thought of as an optimization problem, normally very difficult both because the space of $H$ is large and because $\log ZH$ is very difficult to estimate by itself on a candidate solution.

Several methodological advances and stimulating preliminary applications in neuroscience have been put forward in the last few years [1–4]. Still the field presents several major conceptual and methodological open problems related to both the efficiency and the accuracy of the methods. One problem which we consider here is how to perform inference
when data are not coming from a uniform sampling over the equilibrium configurations of a system but rather they are taken from a subset of all the attractive states. This case arises for instance when we consider systems with multiple attractors and we want to reconstruct the interactions couplings from measurements coming from a subset of the attractors.

In what follows, we take as model system the Hopfield model over random graphs in its memory phase (i.e. with multiple equilibrium states) and show how the interaction couplings can be inferred from data taken from a subset of memories (states). This will be done by employing the Bethe equations (normally used in the ergodic phase where they are asymptotically exact) by taking advantage of a certain property of their multiple fixed points in the non-ergodic phase. The method can be used to infer both couplings and external local fields.

We also show how from the inference method one can derive a simple unsupervised learning protocol which is able to learn patterns in presence of week and highly fluctuating input signals, without ever reaching a spin glass like saturation regime in which all the memories are lost. The technique that we will discuss is based on the so called cavity method and leads to a distributive algorithmic implementation generically known as message–passing scheme.

The paper is organized as follows. First, in Section II we define the problem, and make connections with related works. Section III is concerned with the inference problem in non-ergodic regimes, for which a simple algorithmic approached is presented. In Section IV we apply the technique to the finite connectivity Hopfield model in the memory phase. Section V shows how the approach can be turned into an unsupervised learning protocol. Conclusions and perspectives are given in Section VI.

II. THE INVERSE HOPFILED PROBLEM

The Hopfield model is a simple neural network model with pair-wise interactions which behaves as an attractor associative memory [5]. Its phase diagram is known exactly when memories are random patterns and the model is defined over either fully connected or sparse graphs [6, 7]. Reconstructing interactions in the Hopfield model from multi electrode measurements from large collections of neurons in living organisms. The underlying idea is that a statistically consistent model inferred from the data could capture some aspects of the system which are not easy to grasp from the raw data. Pair wise interactions are in fact believed to capture most of the information content of firing activities observed in real systems [1].

In the Hopfield model the couplings $J_{ij}$ are given by the covariance matrix of a set of random patterns which represent the memories to be stored in the system. We will use the model to generate data through sampling and we will aim at inferring the couplings.

The structure of the phase space of the Hopfield model at sufficiently low temperature and for a not too large number of patterns is divided into clusters of configurations which are highly correlated with the patterns. We will proceed by sampling configurations from a subset of clusters and try to infer the interactions. The simple observation that we want to exploit is the fact that fluctuations within single clusters are heavily influenced by the existence of other clusters and thus contain information about the total system.

We consider a system of $N$ binary neurons $\sigma_i \in \{-1, +1\}$ (or Ising spins) interacting over a random regular graph of degree $K$. The connectivity pattern is defined by the elements $a_{ij} \in \{0, 1\}$ of the adjacency matrix. The (symmetric) interactions between two neighboring neurons are given by the Hebb rule, i.e. $J_{ij} = J_{ji} = \frac{1}{N} \sum_{\mu=1}^{N} \xi_{i}^{\mu} \xi_{j}^{\mu}$, where $\xi_{i}^{\mu}$ are the patterns to be memorized and $P$ is their number. At finite temperature $T = 1/\beta$, we simulate the systems by a Glauber dynamics [8],

$$W(\sigma_i \rightarrow -\sigma_i) = \frac{1 - \sigma_i \tanh \beta h_i}{2},$$

where

$$h_i = \theta_i + \sum_{j \in \partial i} J_{ij} \sigma_j, \quad (2)$$

is the local field experienced by neuron $i$ and $\theta_i$ is an external field. We use $\partial i$ to denote the set of neighbors interacting with $i$. The process satisfies detailed balance and at equilibrium the probability of steady state configurations $\mathcal{g}$ is given by the Gibbs measure

$$\mathcal{P}(\mathcal{g}) = \frac{1}{Z^K} e^{\beta \sum_{i} \theta_i \sigma_i + \beta \sum_{i<j} J_{ij} \sigma_i \sigma_j}, \quad (3)$$
where \( Z[J, \theta] \) is a normalization constant, or partition function. In the memory phase, the system will explore configurations close to a pattern, provided that the initial configuration lies in the basin of attraction of that pattern. In the following we use the wording patterns, basins of attraction or states equivalently. In a given state the average activity (or magnetization) and correlations are denoted by \( m_i^\mu \equiv \langle \sigma_i \rangle_\mu \) and \( c_{ij}^\mu \equiv \langle \sigma_i \sigma_j \rangle_\mu \), where the averages are taken with the Gibbs measure inside that state.

Suppose that starting from some random initial configuration we observe the system for a long time, measuring \( M \) configurations. The standard way to infer interactions couplings \( J \), and external fields \( \theta \) is by maximizing the log-likelihood of \( (J, \theta) \), given the experimental data \([3, 9]\), namely

\[
\frac{1}{\beta M} \mathcal{L}(J, \theta) = \sum_i \theta_i m_i^{exp} + \sum_{i<j} J_{ij}^{\mu} c_{ij}^{exp} + F(J, \theta),
\]

(4)

where \( m_i^{exp} = \frac{1}{M} \sum_{\tau=1}^M \sigma_i^\tau \) and \( c_{ij}^{exp} = \frac{1}{M} \sum_{\tau=1}^M \sigma_i^\tau \sigma_j^\tau \) are the experimental magnetizations and correlations and \( F = -\frac{1}{\beta} \log Z \) is the free energy.

One can exploit the concavity of the log-likelihood and use a gradient ascent algorithm to find the unique parameters maximizing the function. However, this needs an efficient algorithm to compute derivatives of the free energy \( F[J, \theta] \), which in general is a difficult task. A well known technique which can be used for not too big systems is of course the Monte Carlo method (see e.g. \([2]\)). Though under certain limiting assumptions, there exist good approximation techniques which are efficient, namely mean field, small-correlation and large-field expansions \([3, 4, 10–14]\).

In this paper we resort to the mean-field cavity method, or Belief Propagation (BP), to compute the log-likelihood (see e.g. \([15–18]\)). This technique is closely related to the Thouless-Anderson-Palmer (TAP) approximation in spin glass literature \([19, 20]\). The approximation is exact on tree graphs and asymptotically correct as long as the graph is locally tree-like or the correlations between variables are sufficiently weak. In spin glass jargon, the approximation works well in the so called replica symmetric phase.

In the BP approach, the marginals of variables and their join probability distribution (which is assumed to take a factorized form where only pair correlations are kept) are estimated by solving a set of self-consistency functional equations, by exchanging messages along the edge of the interaction graph. A message (typically called “cavity belief”) \( \pi_{i \rightarrow j}(\sigma_i) \) is the probability that neuron \( i \) takes state \( \sigma_i \) ignoring the interaction with its neighbor \( j \). We call this probability distribution a cavity message. The BP equations read (see Ref. \([18]\) for comprehensive review):

\[
\pi_{i \rightarrow j}(\sigma_i) \propto e^{\beta \theta_i} \prod_{k \in \partial i \setminus j} \left( \sum_{\sigma_k} e^{\beta J_{ij} \sigma_i \sigma_j \pi_{k \rightarrow i}(\sigma_k)} \right).
\]

(5)

The equations are solved by starting from random initial values for the cavity messages and updating them in some random sequential order till a fixed point is reached. Upon convergence the cavity messages are used to obtain the local marginals or “beliefs”:

\[
\pi(\sigma_i, \sigma_j) \propto e^{\beta J_{ij} \sigma_i \sigma_j \pi_{i \rightarrow j}(\sigma_i) \pi_{j \rightarrow i}(\sigma_j)}.
\]

(6)

These marginals are enough to compute the magnetizations \( m_i \) and correlations \( c_{ij} \) and thus can be used for maximizing the log-likelihood by updating the parameters as

\[
\theta_i = \theta_i + \eta (m_i^{exp} - m_i^\mu), \]

\[
J_{ij} = J_{ij} + \eta (c_{ij}^{exp} - c_{ij}^\mu),
\]

(7)

with \( \eta \ll 1 \) and positive. Repeating this procedure for sufficient times leads to an estimate of the unknown parameters. Assuming that the external fields are absent, the inference error can be written as:

\[
\Delta_j = \sqrt{\frac{\sum_{i,j \in \partial i} (J_{ij}^{true} - J_{ij})^2}{KN}}.
\]

(8)

A more accurate estimate of the correlations can be obtained by exploiting the Fluctuation-Response theorem \( c_{ij} = \partial m_i / \partial \theta_j \). This method, called Susceptibility Propagation \([12, 21]\), uses derivatives of cavity messages (cavity susceptibilities). Its time complexity grows as \( KN^2 \), to be compared with the \( KN \) complexity of BP equations.

In this paper we will work exclusively with the BP estimate which is simple and accurate enough for our studies. Actually, if one is interested only on correlations along the edges of a sparse graph, the BP estimation would be as good as the one obtained by susceptibility propagation.
III. INFERENCE IN THE NON-ERGODIC PHASE

In an ergodic phase a system visits all configuration space. Sampling for a long time is well represented by the measure in (3).

In a non-ergodic phase, as happens for the Hopfield model in the memory phase, the steady state of a system is determined by the initial conditions. Starting from a configuration close to pattern $\alpha$, the system spends most of its time (depending on the size of system) in that state. We indicate the probability measure which describes such a situation by $P_\alpha(\mathcal{Z})$, that is the Gibbs measure restricted to state $\alpha$. If configurations are sampled from one state, then the expression for the log-likelihood in (4) should be corrected by replacing $F$ with $F_\alpha$, the free energy of state $\alpha$. Still the log-likelihood is a concave function of its arguments and so there is a unique solution to this problem.

It is well known that the Bethe approximation is asymptotically exact in the ergodic phase ([18]). In this case, the Gibbs weight can be approximately expressed in terms of one- and two-point marginals $P_{ij}(\sigma, \sigma)$, $P_i(\sigma)$ as follows:

$$P(\mathcal{Z}) \simeq \prod P_i(\sigma) \prod_{i<j} P_{ij}(\sigma_i, \sigma_j).$$

(9)

The above equation is exact only asymptotically (on a replica-symmetric system); it can be used for inference in at least two ways: The simplest one is by replacing $P_{ij}(\sigma, \sigma)$ and $P_i(\sigma)$ in the above expression by their experimental estimation (given as input of the inference process), equating (9) to (3) and solving for $J$ and $\theta$. This is known as the “independent pairs” approximation. A second one, often more precise but computationally more involved, is to search for a set of $J, \theta$ and a corresponding $J, \theta$-fixed point of BP equations, such that the Bethe estimation $\pi_{ij}(\sigma, \sigma)$, $\pi_i(\sigma)$ of the two- and one-point marginals matches the experimental input as accurately as possible.

In a non-ergodic phase, it is known however that BP equations typically do not converge or have multiple fixed points. This is normally attributed to the fact that the BP hypothesis of decorrelation of cavity marginals fails to be true. When a BP fixed point is attained, it is believed to approximate marginals inside a single state (and not the full Gibbs probability), as the decorrelation hypothesis are satisfied once statistics are restricted to this state [18].

The fact that BP solutions correspond to restriction to subsets of the original measure may suggest that there is little hope in exploiting (9) on such systems. Fortunately, this is not the case. For every finite system, and every BP fixed point $\alpha$ the following holds,

$$P(\mathcal{Z}) = \frac{Z^\alpha_{\text{Bethe}}}{Z_{\text{Bethe}}} \prod \pi^\alpha_i(\sigma_i) \prod_{i<j} \pi^\alpha_{ij}(\sigma_i, \sigma_j) \pi^\alpha_j(\sigma_j).$$

(10)

A proof of a more general statement will be given in appendix A. As in the ergodic case, (10) can be exploited in at least two ways: one is by replacing $\pi^\alpha_{ij}(\sigma, \sigma)$ and $\pi^\alpha_i(\sigma)$ by their experimental estimation inside a state and solving for $J, \theta$ the identity between (10) and (3), exactly like in the independent pairs approximation, as if one just forgets that the samples come from a single ergodic component. A second one is by inducing BP equations to converge on fixed points corresponding to appropriate ergodic components. In this paper we will take the latter option.

Please notice that our method, as an algorithm, is more flexible with respect to the first one; indeed, there is no reason to have a BP fixed point for any set of experimental data, especially when the number of samples or size of system is not too large. It means that matching exactly the data with those of a BP fixed point is not always possible. Therefore, a better strategy would be to find a good BP solution which is close enough to the experimental data.

Ignoring the information that our samples come from a single ergodic component would result in a large inference error due to the maximization of the wrong likelihood. As an example, let us take a tree graph with Ising spins interacting through random couplings $-1 \leq J_{ij} \leq +1$, in zero external fields $\theta_i = 0$. Let us assume the system follows the following measure:

$$P_d(\mathcal{Z}) \propto I(\mathcal{Z} \in \Omega_d(\xi)) e^{\beta \sum_{i<j} J_{ij} \sigma_i \sigma_j},$$

(11)

where $I(\mathcal{Z} \in \Omega_d(\xi))$ is an indicator function which selects configurations in the subspace $\Omega_d(\xi)$, a sphere of radius $d$ around pattern $\xi$. One can fix a fraction $q$ of the boundary spins to the values in pattern $\xi$. For $q = 0$ and $q = 1$ the system mimics ergodic and non-ergodic phases, respectively. By the BP approximation we can compute the magnetizations $m^{(d)}_i$ and the correlations $c^{(d)}_{ij}$, see Appendix B for more details. Taking these as experimental data, we may perform the inference by assuming that our data represent the whole configuration space. Figure 1 displays the inference error that one obtains in this way. Notice that in principle the error would be zero if we knew $\Omega_d(\xi)$. In contrast to the paramagnetic phase, in the non-ergodic case the error remains nonzero even if sampling is performed for very long times.
The Hopfield model can be found in three different phases. For large temperatures the system is in the paramagnetic phase where, in the absence of external fields, magnetizations $m_i$ and overlaps $O^\mu = \frac{1}{N} \sum_i \xi_i^\mu m_i$ are zero. If the number of patterns is smaller than the critical value $P_c$, for small temperatures the system enters the memory phase where the overlap between the patterns and the configurations belonging to states selected by the initial conditions can be nonzero. For $P > P_c$, the Hopfield model at low temperature enters in a spin glass phase, where the overlaps are typically zero. In fully connected graphs $P_c \simeq 0.14N$ whereas in finite connectivity graphs it is about the average degree (or $K$ in random regular case) [22].

Take the Hopfield model with zero external fields and in the memory phase. We measure samples from a Glauber dynamics which starts from a configuration close to a pattern $\nu$. The system will stay for a long time in the state $\nu$ and is thus well described by the restricted Gibbs measure $\mathcal{P}_\nu(\sigma)$. In a configuration $\sigma$, the local field seen by neuron $i$ is $h_i = \sum_{j \in \partial i} J_{ij}\sigma_j + \frac{1}{N} \sum_{\mu \neq \nu} \xi_i^\mu + \frac{1}{N} \sum_{j \in \partial i} \xi_{ij}^\nu \sigma_j$. If $\nu$ corresponds to the retrieved pattern, the first term (signal) would have the dominant contribution to $h_i$. The last term (noise) is a contribution of the other patterns to the local field. To exploit this information, we look for a set of couplings that result to a Gibbs state equivalent to the observed state of the system. One way to do this is by introducing an auxiliary external field pointing to the local field. To exploit this information, we look for a set of couplings that result to a Gibbs state equivalent to the observed state of the system. One way to do this is by introducing an auxiliary external field pointing to the experimental magnetizations, i.e. $\theta_i = \lambda m_i^{\text{exp}}$, for a positive $\lambda$; we may set the couplings $J_{ij}$ at the beginning to zero and compute our estimate of the correlations $c_{ij}$ by the BP algorithm. This can be used to update the couplings by a small amount in the direction that maximizes the likelihood, as in (7) (we do not update the external fields which for simplicity are assumed to be zero). This updating is repeated iteratively, decreasing $\lambda$ by a small amount in each step. The procedure ends when $\lambda$ reaches the value zero.

The auxiliary field is introduced only to induce convergence of the equations towards a fixed point giving statistics inside a particular state. Figure 2 compares the inference error obtained with the above procedure for several values of temperature in the memory phase. For the parameters in the figure, the inferred couplings from one basin were enough to recover the other two patterns. In the figure we also see how the error decreases by taking larger number of samples from the system.

In general we may have samples from different states of a system. Let us assume that in the Hopfield model we stored $P$ patterns by the Hebb rule but the samples are from $Q$ basins. The estimated correlations $c_{ij}^{\mu}$ in any state $\mu \in \{1, \ldots, Q\}$ should be as close as possible to the experimental values $c_{ij}^{\text{exp},\mu}$.

A natural generalization of the previous algorithm is the following: As before we introduce external fields $\theta_i^\mu = \lambda m_i^{\text{exp},\mu}$ for each state $\mu$. At fixed positive $\lambda$ we compute the estimated BP correlations for different states. Each of these estimations can be used to update the couplings as in the single state case. Specifically, this amounts to make a single additive update to the couplings by the average vector $\eta \Delta \sigma$ given by $\Delta \sigma_{ij} = (\bar{c}_{ij}^{\text{exp},\mu} - \bar{c}_{ij}^\mu)$, where $\bar{c}_{ij}^{\text{exp},\mu} = \frac{1}{Q} \sum_{\mu=1}^Q c_{ij}^{\text{exp},\mu}$ and $\bar{c}_{ij}^\mu = \frac{1}{Q} \sum_{\mu=1}^Q c_{ij}^\mu$. Indeed, the addends of $\Delta \sigma$ will be typically linearly independent, so $\Delta \sigma = 0$ will imply $c_{ij}^{\text{exp},\mu} = c_{ij}^\mu$ for $\mu = 1, \ldots, Q$. 

FIG. 1: Inference error in ergodic ($q = 0$) and non-ergodic ($q = 1$) cases, if the data come from subspace $\Omega_d(\xi)$ and we do not use this information.

IV. INFERENCE IN THE HOPFIELD MODEL
We then decrease $\lambda$ and do the BP computation and update steps. Again we have to repeat these steps until the external field goes to zero. Figures 3 and 4 show how the inference error changes with sampling from different states. Notice that if we had an algorithm that returns exact correlations, an infinite number of samples from one state would be enough to infer the right interactions in the thermodynamic limit. However, given that we are limited by the number of samples, the learning process is more efficient if this number is taken from different states instead of just one.

V. FROM INFERENCE TO UNSUPERVISED LEARNING

Hebbian learning is a stylized way of representing learning processes. Among the many oversimplifications that it involves there is the fact that patterns are assumed to be presented to the networks through very strong biasing signals. On the contrary it is of biological interest to consider the opposite limit where only weak signals are allowed and retrieval takes place with sizeable amount of errors. In the spin language we are thus interested in the case in which the system is only slightly biased toward the patterns during the learning phase. In what follows we show that
FIG. 4: Effect of sampling from $Q$ states: inference error and number of states which are stable and highly correlated with the patterns. Here $N = 1000$, $\beta = 2$ and $\eta = 0.02$.

one can “invert” the inference process discussed in the previous sections and define a local learning rule which copes efficiently with this problem.

As first step we consider a learning protocol in which the patterns are presented sequentially and in random order to the system by applying an external field in direction of the pattern, that is a field $\theta^\mu = \lambda \xi^\mu$ with $\lambda > 0$. We may assume that initially all couplings are zero.

Depending on the strength of the field, the system will be forced to explore configurations at different overlaps with the presented pattern $\mu$. A small $\lambda$ corresponds to a weak or noisy learning whereas for large $\lambda$ the system has to remain very close to the pattern. For any $\lambda$ we can compute the correlations $c_{ij}^{\lambda,\mu}$ by either sampling from the Glauber dynamics or by directly running BP. At the same time we can compute correlations $c_{ij}^\mu$ by the BP algorithm in the absence of external fields and try to find couplings which match the correlations in the two cases, namely we can update the couplings by a quantity $\eta (c_{ij}^{\lambda,\mu} - c_{ij}^\mu)$. The process is repeated for all couplings and for $t_L$ iterations with the same pattern $\mu$. Next we switch to some other randomly selected pattern $\nu$ and the whole process is repeated for $T$ learning steps. Notice that here $\lambda$ is fixed from the beginning.

The above learning protocol displays a range of interesting phenomena. Firstly one notices that for $c_{ij}^{\lambda,\mu} \approx \xi^\mu \xi^\mu_j$ (i.e. for very large external fields) and $c_{ij}^\mu \approx 0$ (i.e. for very high temperature or isolated neurons) the above learning results to the Hebb couplings of the Hopfield model. In Figure 5 we compare the histogram of learned couplings for small and large $\lambda$ with the Hebbian ones.

The number of patterns $P_s$ which are highly correlated with stable configurations depends on the strength of external fields. We consider that pattern $\mu$ is “learned” if there is a Gibbs state with nonzero overlap $O^\mu$ that is definitely larger than the other ones $\{O^\nu | \nu \neq \mu\}$. Figures 6 and 7 show how these quantities evolve during the learning process and by increasing the magnitude of external filed.

For small $P$ nearly all patterns are learned, whereas, for larger $P$ some patterns are missing. A large number of patterns can thus be learned at the price of smaller overlaps and weaker states. That is, the average overlap in successfully learned patterns decreases continuously by increasing $P$, approaching the paramagnetic limit. In Figure 8 we compare this behavior with that of Hebb couplings.

As the figure shows, there is a main qualitative difference between Hebbian learning of the Hopfield model and the protocol discussed here. In the former case when the number of stored patterns exceeds some critical value the systems enters in a spin glass phase where all memories are lost and the BP algorithm does not converge anymore. On the contrary, in our case many patterns can be stored without ever entering the spin glass phase (for a wide range of choices of $\lambda$). The BP algorithm always converges, possibly to a wrong fixed point if the corresponding pattern is not stored.
A. Population dynamics analysis of the learning protocol

Population dynamics is usually used to obtain the asymptotic and average behavior of quantities that obey a set of deterministic or stochastic equations [23]. For instance, to obtain the phase diagram of the Hopfield model with population dynamics one introduces a population of $N_P$ messages representing the BP cavity messages in a reference state, e.g. pattern $\xi^\nu = +1$ [22]. Then one updates the messages in the population according to the BP equations: at each time step a randomly selected cavity message is replaced with a new one computed by $K - 1$ randomly selected ones appearing on the r.h.s. of the BP equations. In each update, one generates the random couplings $J_{ij} = \frac{1}{K} + \frac{1}{K} \sum_{\mu \neq \nu} \xi_i^\mu \xi_j^\nu$ by sampling the other $P - 1$ random patterns. After a sufficiently large number of updates, one can compute the average overlap with the condensed pattern $\nu$ to check if the system is in a memory phase. The stability of condensed state would depend on the stability of the above dynamics with respect to small noises in the cavity messages. If $N_P$ is large enough, one obtains the phase diagram of Hopfield model in the thermodynamic limit averaged over the ensemble of random regular graphs and patterns. We used the above population dynamics to obtain the phase diagram of the Hopfield model on random regular graphs, see Figure 9.

In order to study the new learning protocol we need a more sophisticated population dynamics. The reason is that
FIG. 7: Average number of successfully learned patterns for different value of $\lambda$. Here $N = 1000$, $\beta = 2$, $\lambda = 0.2$ and $\eta = 0.003$. The average is taken over 10 realizations of the patterns.

FIG. 8: Average number of successfully learned patterns in the learning algorithm and Hebb rule. The inset shows the average overlap. Here $N = 1000$, $\beta = 2$, $\lambda = 0.2$ and $\eta = 0.003$. The average is taken over 10 realizations of the patterns.

in contrast to Hebb couplings, we do not know in advance the learned couplings. In Appendix C we explain in more details the population dynamics that we use to analyze the learning process studied in this paper. The algorithm is based on $P$ populations of BP messages and one population of couplings. These populations represent the probability distributions of BP messages in different states and couplings over the interaction graph. For a fixed set of patterns $\{\epsilon^\mu|\mu = 1, \ldots, P\}$ we update the populations according to the BP equations and the learning rule, to reach a steady state. Figure 10 displays the histogram of couplings obtained in this way. In the figure we compare two cases of bounded and unbounded couplings. In the first case the couplings have a magnitude less than or equal to 1 whereas in the second case they are free to take larger values. We observe a clear difference between the two cases; when $\lambda$ is small, the couplings are nearly clipped in the bounded case whereas the unbounded couplings go beyond $\pm 1$. However, in both cases there is some structure in the range of small couplings. Increasing the magnitude of $\lambda$ we get more and more structured couplings. For very large fields they are similar to the Hebb couplings. For small fields the histogram of the couplings is very different from the Hebb one, though the sign of the learned and the Hebbian couplings is the same.

There are a few comments to mention here; in the population dynamics we do not have a fixed graph structure and to distinguish $P$ patterns from each other we have to fix them at the beginning of the algorithm. Moreover, we have...
to modify the BP equations to ensure that populations are representing the given patterns, see Appendix C. And finally the outcome would be an average over the ensemble of random regular graphs, for a fixed set of patterns.

Having the stationary population of couplings, one can check the stability of each state by checking the stability of the BP equations at the corresponding fixed point. The maximum capacity that we obtain in this way for the learned couplings is the same as the Hebb one whereas on single instances we could store much larger number of patterns. The reason why we do not observe this phenomenon in the population dynamics resides in the way that we are checking stability; the fixed patterns should be stable in the ensemble of random regular graphs. In other words, checking for stability in the population dynamics is stronger than checking it in a specific graph.

The main result of our analysis consists in showing that the distribution of the couplings arising from the BP learning protocol is definitely different from the Hebbian one.

VI. DISCUSSION AND PERSPECTIVES

We studied the finite connectivity inverse Hopfield problem at low temperature, where the data are sampled from a non-ergodic regime. We showed that the information contained in the fluctuations within single pure states can be used to infer the correct interactions.

We also used these findings to design a simple learning protocol which is able to store patterns learned under noisy conditions. Surprisingly enough it was possible to show that by demanding a small though finite overlap with the patterns it is possible to store a large number of patterns without ever reaching a spin glass phase. The learning process avoids the spin glass phase by decreasing the overlaps, as the number of patterns increases. A separate analysis which is similar to the one presented in ref. [25] (and not reported here) shows that the equations can be heavily simplified without losing their main learning capabilities.

In this paper we focused on a simple model of neural networks with symmetric couplings. It would be interesting to study more realistic models like the integrate and fire model of neurons with general asymmetric couplings. Moreover, instead of random unbiased patterns one may consider sparse patterns which are more relevant in the realm of neural networks.

The arguments presented in this paper can also be relevant to problem of inferring a dynamical model for a system by observing its dynamics. In this case, a system is defined solely by its evolution equations and one cannot rely on the Boltzmann equilibrium distribution. Still it is possible to try to infer the model by writing the likelihood for the model parameters given the data and given the underlying dynamical stochastic process. A mean-field approach has been recently described in [24]. We actually checked this approach in our problem and observed qualitatively the same behavior as the static approach. In fact, which method is best heavily depends on the type of data which are available.
FIG. 10: The histogram of learned couplings in the population dynamics with the BP approximation. Here \( N_p = 1000, \beta = 1, \) and \( \eta = 0.01. \)

VII. ACKNOWLEDGEMENTS

APPENDIX A: PROOF OF EXACTNESS OF THE BETHE EXPRESSION FOR ARBITRARY BP FIXED POINTS

A \( \beta \to \infty \) limit version of this result (except the determination of the value of the constant \( Z_{\text{Gibbs}} / Z_{\text{Bethe}} \)) appeared in [26]. This result is valid for general (non-zero) interactions. For a family of “potentials” \( \Psi_a : z_a \mapsto \Psi_a(z_a) > 0, \) where we denote by \( z_a \) the subvector of \( z \) given by \( \{ x_i : i \in \partial a \} \). We will use the shorthand \( i \in a \) or \( a \in i \) to mean \( i \in \partial a \).

Proposition. Given a factorized probability function

\[
\mathcal{P}(z) = \frac{1}{Z_{\text{Gibbs}}} \prod_a \Psi_a(z_a)
\]

and a BP fixed point \( \{ b_{ia} \}_{i \in a, a \in A} \) and plaquette marginals \( b_a(x_a) = z^{-1}_a \Psi_a(z_a) \prod_{i \in a} b_{ia}(x_i) \) and single marginals \( b_i(x_i) = \sum_{\{ x_j : j \in a \setminus i \}} b_a(x_a) \) for every \( a \in i \), then

\[
\mathcal{P}(z) = \frac{Z_{\text{Bethe}}}{Z_{\text{Gibbs}}} \prod_a \frac{b_a(z_a)}{\prod_{i \in a} b_i(x_i)} \prod_i b_i(x_i)
\]
Proof. Using the fact that \( b_i(x_i) = z_i^{-1} \prod_{a \in i} b_{a}(x_i) \) and \( b_{ia}(x_i) = z_{ia}^{-1} \prod_{a \in i \setminus a} b_{a}(x_i) \), we obtain \( b_{ia}(x_i) = b_i(x_i) z_{ia}^{-1} x_i^{-1} \), then using the definitions:

\[
\prod_a b_a(x_a) \prod_i b_i(x_i) = \prod_a \Psi_a(x_a) z_a^{-1} \prod_{i \in a} b_{ia}(x_i) \prod_i b_i(x_i) \\
= \prod_a \Psi_a(x_a) z_a^{-1} b_{ia}(x_i) \prod_{i \in a} b_{ia}(x_i) \\
= \prod_a \Psi_a(x_a) z_a^{-1} b_{ia}(x_i) \prod_{i \in a} b_{ia}(x_i) \\
= Z_{\text{Gibbs}} P(\bar{\Psi}) \prod_a \frac{z_a^{-1}}{z_a} \prod_{i \in a} z_{ia}^{-1} x_i^{-1} \\
= Z_{\text{Gibbs}} P(\bar{\Psi}) Z_{\text{Bethe}}
\]

This proves that a fixed point can be interpreted as a form of reparametrization of the original potentials. In fact, a sort of converse also holds:

**Proposition:** If \( P(\bar{\Psi}) \propto \prod_a \Psi_a(x_a) \) satisfies a Bethe-type expression

\[
P(\bar{\Psi}) \propto \prod_a b_a(x_a) \prod_i b_i(x_i) 
\]

with \( \sum_{\{x_i \in i \setminus a\}} b_a(x_a) = b_i(x_i) \) for every \( i \in a \). Then there exists a BP fixed point \( \{ b_{a}(x_a) \}_{i \in a, a \in A} \) such that

\[
\Psi_a(x_a) \propto \prod_{i \in a} b_{ia}(x_i) 
\]

**Proof:** Choose any configuration \( \bar{\Psi} \). We will use the following notation: \( \mathcal{I}_a = \{ i \} \cap \partial a, \) and \( \mathcal{I}_x = \{ i \} \setminus \mathcal{I}_a \). Define \( b_{ia}(x_i) \propto \Psi_a^{-1} (x_i, x_{\mathcal{I}_a}) b_i(x_i, x_{\mathcal{I}_a}) \), normalized appropriately. Afterwards, we can define \( b_{ia}(x_i) \propto b_i(x_i) b_{ia}(x_i) \).

By definition of \( P \) we have \( P(x_i | \bar{\Psi}) = \prod_{a \in i} \Psi_a(x_i, x_{\mathcal{I}_a}) \). Similarly, but using (A1), and noting by \( n_i = |\partial i| \),

we have also \( P(x_i | \bar{\Psi}) \propto b_i(x_i) b_{ia}(x_i) \prod_{a \in i} b_{ia}(x_i) \). Thus \( b_i(x_i) b_{ia}(x_i) \propto b_i(x_i) b_{ia}(x_i) \prod_{a \in i} b_{ia}(x_i) \).

By definition of \( P, P(x_i | \bar{\Psi}) \propto \Psi_i(x_i) \prod_{a \in i} c_i(x_i) \) where \( c_i(x_i) = \prod_{a \in i \setminus a} \Psi_a (x_i, x_{\mathcal{I}_a}) \). Moreover using (A1), we can conclude that also \( P(x_i | \bar{\Psi}) \propto b_i(x_i) \prod_{a \in i} c_i(x_i) \) where \( c_i(x_i) = \prod_{a \in i \setminus a} b_{ia}(x_i) \). This implies that \( b_i(x_i) \propto \Psi_i(x_i) \prod_{a \in i} c_i(x_i) \). But we also have that \( \frac{c_i(x_i)}{d_i(x_i)} \propto b_i(x_i) b_{ia}(x_i) \propto \Psi_i(x_i) \prod_{a \in i} b_{ia}(x_i) \) as desired. Now \( b_i(x_i) = \sum_{\mathcal{I}_a} b_{ia}(x_i) \) by hypothesis, so \( b_{ia}(x_i) \propto b_i(x_i) b_{ia}(x_i) \prod_{a \in i \setminus a} \Psi_a(x_a) \prod_{j \in a \setminus i} b_{aj}(x_j) \) and this proves that the second BP equation is also satisfied.

**APPENDIX B: COMPUTING THERMODYNAMIC QUANTITIES IN A RESTRICTED SPACE**

Consider the Ising model on a tree graph of size \( N \) with couplings \( J \) and external fields \( h \). Suppose that we are given a reference point \( \xi \) in the configuration space \( \{-1,+1\}^N \) and the following measure

\[
P_d(\bar{\Psi}) \propto \prod_{i < j} d_{i \rightarrow j} (1 - \delta_{\sigma_i \xi, \xi_j}),
\]

where \( \Omega_d(\xi) \) is a sphere of radius \( d \) centered at \( \xi \). By distance of two configurations we mean the Hamming distance, i.e. number of spins which are different in the two configurations. The aim is to compute thermodynamic quantities like average magnetizations and correlations in an efficient way. We do this by means of the Bethe approximation and so BP algorithm.

First we express the global constraint \( I(\bar{\Psi} \in \Omega_d(\xi)) \) as a set of local constraints by introducing messages \( d_{i \rightarrow j} \) that each node sends for its neighbors. For a given configuration \( \bar{\Psi} \),

\[
d_{i \rightarrow j} = \sum_{k \in \partial i \setminus j} d_{k \rightarrow i} + (1 - \delta_{\sigma_i \xi, \xi_j}),
\]
denotes the distance of \( \mathbf{z} \) from \( \xi \) in the cavity graph \( \mathcal{G}_{i\rightarrow j} \) which includes \( i \) and all nodes connected to \( j \) through \( i \). With these new variables we can write BP equations as

\[
\pi_{i\rightarrow j}(\sigma_i, d_{i\rightarrow j}; \sigma_j, d_{j\rightarrow i}) \propto e^{\beta J_{ij} \sigma_i \sigma_j} \sum_{\{\sigma_k, d_{k\rightarrow i}\}_{k\in \partial \setminus \{j\}}} I_i \prod_{k \in \partial \setminus \{j\}} \pi_{k\rightarrow i}(\sigma_k, d_{k\rightarrow i}; \sigma_i, d_{i\rightarrow k}),
\]

where \( I_i \) is an indicator function to check the constrains on \( d_{i\rightarrow k} \) and \( \sum_{k \in \partial i} d_{k\rightarrow i} + (1 - \delta_{\sigma_i, \xi_i}) \leq d \). Starting from random initial values for the BP messages we update them according to the above equation. After convergence the local marginals read

\[
\pi(\sigma_i, \sigma_j) \propto e^{-\beta J_{ij} \sigma_i \sigma_j} \sum_{d_{i\rightarrow j},d_{j\rightarrow i}} I_{ij} \pi_{i\rightarrow j}(\sigma_i, d_{i\rightarrow j}; \sigma_j, d_{j\rightarrow i}) \pi_{j\rightarrow i}(\sigma_j, d_{j\rightarrow i}; \sigma_i, d_{i\rightarrow j}),
\]

where in \( I_{ij} \) we check if \( d_{i\rightarrow j} + d_{j\rightarrow i} \leq d \). These marginals will be used to compute the average magnetizations and correlations. Notice that when the graph is not a tree we need to pass the messages \( d \) along the edges of a spanning tree (or chain) which is selected and fixed at the beginning of the algorithm.

APPENDIX C: POPULATION DYNAMICS

Consider \( P \) patterns \( \xi^\mu \in \{-1, +1\} \), where \( \mu = 1, \ldots, P \) and \( a \) goes from 1 to \( N_p \), which is equivalent to the size of system. The patterns, learning rate \( \eta \) and parameter \( \lambda \) are fixed at the beginning of the algorithm. To each pattern we assign a population of messages \( \pi^\mu_{a, i}(\sigma) \) where \( l = 1, \ldots, K \) (is the node degree). These are to represent the normalized BP messages that we use in the learning algorithm. Besides this we have also a population of couplings \( J_{ab} \).

The population dynamics has two update steps: updating the \( P \) populations of messages and updating the population of couplings.

To update the messages in population \( \mu \) we do the following:

i) select randomly \( (a_0, l_0) \) and \( \{(a_1, l_1), \ldots, (a_{K-1}, l_{K-1})\} \),
ii) use messages \( \{\pi^\mu_{a_1, l_1}^\mu, \ldots, \pi^\mu_{a_{K-1}, l_{K-1}}^\mu\} \) and couplings \( \{J_{a_0,a_1}, \ldots, J_{a_0,a_{K-1}}\} \) to compute a new BP message \( \pi^\mu_{a_0, l_0} \),
iii) replace message \( \pi^\mu_{a_0, l_0}(\xi^\mu) \) with \( \max(\pi^\mu_{a_0, l_0}(\xi^\mu), \pi^\mu_{a_0, l_0}(-\xi^\mu)) \).

Notice to the maximum we are taking in the last step. This is to ensure that BP messages in population \( \mu \) are related to pattern \( \xi^\mu \). We do these updates for \( t_{BP} \) iterations, where in each iteration all members of a population are updated in a random sequential way.

To update the couplings we go through the \( P \) populations and do the following:

i) select randomly \( (a, l_a) \) and \( (b, l_b) \),
ii) use messages \( \pi^\mu_{a, l_a}^\mu, \pi^\mu_{b, l_b}^\mu \) and coupling \( J_{ab} \) to compute correlation \( c^\mu_{ab} \), i.e. in presence of external fields \( \lambda^\mu_{a} \) and \( \lambda^\mu_{b} \),
iii) use messages \( \pi^\mu_{a, l_a}^\mu, \pi^\mu_{b, l_b}^\mu \) and coupling \( J_{ab} \) to compute correlation \( c^\mu_{ab} \), i.e. in absence of the external fields,
iv) update the coupling as \( J_{ab} = J_{ab} + \eta (c^\mu_{ab} - c_{ab}) \)

The learning updates are done for \( t_L \) iterations. All together the population dynamics will have \( T \) learning steps each one consist of \( P t_{BP} + P t_L \) update iterations. In practice we set \( t_{BP} \simeq 10 \) and \( t_L \simeq 1 \).