

Machine-learning iterative calculation of entropy for physical systems

Amit Nir (עמית ניר)^{a,b}, Eran Sela (ערן סלע)^a, Roy Beck^{a,b,c,1}, and Yohai Bar-Sinai (יוחאי בר-סיני)^{a,b,d,1}

^aThe School of Physics and Astronomy, Tel Aviv University, Tel Aviv 69978, Israel; ^bThe Center for Physics and Chemistry of Living Systems, Tel Aviv University, Tel Aviv 69978, Israel; ^cThe Center for Nanoscience and Nanotechnology, Tel Aviv University, Tel Aviv 69978, Israel; and ^dGoogle Research, Tel-Aviv 67891, Israel

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Characterizing the entropy of a system is a crucial, and often computationally costly, step in understanding its thermodynamics. It plays a key role in the study of phase transitions, pattern formation, protein folding, and more. Current methods for entropy estimation suffer from a high computational cost, lack of generality, or inaccuracy and inability to treat complex, strongly interacting systems. In this paper, we present a method, termed machinelearning iterative calculation of entropy (MICE), for calculating the entropy by iteratively dividing the system into smaller subsystems and estimating the mutual information between each pair of halves. The estimation is performed with a recently proposed machine-learning algorithm which works with arbitrary network architectures that can be chosen to fit the structure and symmetries of the system at hand. We show that our method can calculate the entropy of various systems, both thermal and athermal, with state-of-the-art accuracy. Specifically, we study various classical spin systems and identify the jamming point of a bidisperse mixture of soft disks. Finally, we suggest that besides its role in estimating the entropy, the mutual information itself can provide an insightful diagnostic tool in the study of physical systems.

entropy estimation | mutual information | machine learning | jamming

E ntropy is a fundamental concept of statistical physics whose computation is crucial for a proper description of many phenomena, including phase transitions (1–3), pattern formation (4), self-assembly (5–7), protein folding (8–10), and many more. In the physical sciences, entropy is typically interpreted as quantifying the amount of disorder of a system or the level of quantum entanglement. Entropy is also a fundamental concept in other fields of thought—statistical learning, economy, inference, and cryptography, among others (11). There it is used to quantify the complexity of statistical distributions. Mathematically, entropy is defined as

$$S = -k_{\rm B} \sum_{i} p_i \log p_i, \qquad [1]$$

where p_i is the probability that the system is in the *i*th microstate, and $k_{\rm B}$ is the Boltzmann constant. For convenience, in what follows we work with units where $k_{\rm B} = 1$.

Analytic calculation of the entropy is achievable only for simple, weakly interacting systems. Experimentally, the entropy can be obtained, for example, by measuring the temperature (T) dependence of the specific heat down to low temperatures (12). Computationally, for all but the simplest systems, a direct calculation of the entropy is computationally infeasible, as it requires computational resources that grow exponentially with system size (13, 14). For example, a classical numerical approach involves integrating the specific heat, which is inferred from energy fluctuations, down to low temperatures (12). This method is computationally costly and can suffer from inaccuracies for systems with numerous ground states at low T. Other methods estimate directly the free energy (15) or embrace additional knowledge on the system, for example from experiment, to reduce the entropic contribution to a manageable computational task (16).

Recently, we and others have shown that using compression algorithms one can compute, to a good approximation, the entropy of fairly complex systems (8, 17, 18). This method is based on Kolmogorov's theorem that states that the optimal compression of data drawn from a distribution is bounded by the distribution's entropy (19, 20). The compression-based methods capitalize on decades of research in computer science, which resulted in fast and efficient compression algorithms, such as the Lempel–Ziv algorithm or variants of it (21) which are widely available. However, these algorithms treat data as a one-dimensional (1D) discrete string, and manipulating higher-dimensional data into a 1D structure results in information loss. For example, it was recently demonstrated that compression-based algorithms misestimate the entropy of systems with long-range correlations and fail to capture delicate transitions in complex systems (17).

Here, we introduce a generic approach which we term machine-learning iterative calculation of entropy (MICE). Our method improves on existing methods in a number of ways: First, it provides state-of-the-art accuracy. Second, it is scalable, in the sense that its computational cost grows logarithmically with system size. Third, it provides estimations of the actual entropy, with physical units, without additive or multiplicative corrections and with no fitting parameters. Fourth, since the underlying computations are performed with artificial neural nets, MICE can be naturally applied to various physical systems by adjusting the network architecture, rather than the digital representation of the system (e.g., flattening high-dimensional systems to one-dimensional byte arrays as in refs. 8, 17, and 18). Finally, it can be applied to both discrete and continuous distributions.

Below we test MICE on several canonical systems: the Ising model on both square and triangular lattices, the XY model

Significance

The calculation of entropy of a physical system is a fundamental step in learning its thermodynamic behavior. However, current methods to compute the entropy are often system specific and computationally costly. Here, we propose a method that is efficient, accurate, and general for computing the entropy of arbitrary physical systems. Our method is based on computing the mutual information between subsystems within the studied system, using a convolutional neural network. This iterative procedure provides accurate entropy evaluation for systems in and out of equilibrium.

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¹To whom correspondence may be addressed. Email: ybarsinai@gmail.com or roy@tauex. tau.ac.il.

with and without an external magnetic field (H), and an athermal system of bidisperse soft disks in two dimensions (2D). We show that our approach provides state-of-the-art accuracy and provides insightful information about the physics as a byproduct.

The Method

Entropy and Mutual Information. In thermodynamics, entropy is considered to be an extensive quantity, i.e., a quantity that scales linearly with system size. This is only approximately true. In fact, the entropy is strictly subextensive. The quantity that measures the subextensiveness is called mutual information.

To be precise, the mutual information (\mathcal{M}) between two random variables A, B is defined by the relation (11)

$$S(A, B) = S(A) + S(B) - \mathcal{M}(A, B),$$
 [2]

where S(A), S(B) are the entropies of A and B, respectively, and S(A, B) is their joint entropy. It is easy to show that $\mathcal{M}(A, B)$ is strictly nonnegative (11). Therefore, if we think of A and B as two halves of a thermodynamical system, this equation tells us that the entropy of the joint system is smaller than the sum of the entropies of its components.

Eq. 2 is the basic relation on which our method relies. It allows calculation of the entropy of a large system by estimating the entropy of each of its halves and the mutual information between them. Since the computational cost of estimating the entropy grows exponentially with the system size, the latter might be a significantly easier problem than the former.

With this in mind, consider a large physical system X_0 , of volume V_0 , which we divide into two equal halves. If we deal with translationally invariant systems, as we assume for the remainder of this work, the two halves are statistically indistinguishable, and we denote both of them by X_1 (Fig. 1A). With this notation, Eq. 2 takes the form

$$S(X_0) = 2S(X_1) - \mathcal{M}(X_1),$$
 [3]

where $\mathcal{M}(X_k)$ is a shorthand notation for the mutual information between two neighboring subsystems X_k . Each of the halves can be further divided into two statistically indistinguishable halves, and this process can be iterated arbitrarily many times. After m iterations, we find that

$$s(X_0) \equiv \frac{S(X_0)}{V} = s_m - \frac{1}{2} \sum_{k=1}^m \frac{\mathcal{M}(X_k)}{V_k},$$
 [4]

where $V_k = 2^{-k} V_0$ is the volume (or area in 2D) of the kth subsystem, and $s_m \equiv S(X_m)/V_m$ is the specific entropy of the *m*th subsystem.

Eq. 4 decomposes the entropy S into contributions from different length scales. At very short scales, the iteration should be carried out only until X_k becomes small enough that its entropy can by directly calculated, either by brute-force enumeration or by using other methods. Since V_k decreases exponentially with k, the number of needed iterations is logarithmic in the system size. In many cases the actual value of the first term in the righthand side of Eq. 4, i.e., the entropy of the smallest subsystem, is an uninteresting additive constant with no physical significance and can be ignored.

In summary, the crux of our method is replacing the problem of evaluating the entropy by that of calculating the mutual information between subsystems of varying sizes (Fig. 1A). It is left to understand how to actually calculate the mutual information, which is the topic of the next section.

Estimating the Mutual Information. Recently, Belghazi et al. (25) proposed a method to calculate the mutual information between high-dimensional random variables with neural networks. Their



Fig. 1. (A) Schematic illustration of MICE. By dividing the system into smaller subsystems and calculating the mutual information between them we reconstruct the entropy of the whole system. The entropy of the smallest subsystem is calculated directly by enumeration. Dashed red lines mark the length of interface (ℓ_i) between two subsystems in the *i*th iteration. (*B*-D) The difference between MICE estimations of s and known benchmarks. Note that the units are chosen such that $k_{\rm B} = 1$. We present three estimation methods: MICE, naive extrapolation from a system of 16 spins (main text), and a compression-based method (8). MICE shows superior performance in all cases. B–D show results for (B) the ferromagnetic Ising model on a square lattice, (C) the antiferromagnetic Ising model on a triangular lattice, and (D) the XY model on a square lattice. In B and C we benchmark against known analytical results for infinite systems (22, 23), respectively. In D, we benchmark against the calculation of ref. 24.

idea is simple and elegant: Following a theorem by Donsker and Varadhan (26), the mutual information between two variables, A and B, can be expressed as a solution to a maximization problem:

$$\mathcal{M} = \sup_{\theta \in \Theta} \left[\left\langle \mathcal{F}_{\theta}(A, B) \right\rangle_{P_{A,B}} - \log \left\langle e^{\mathcal{F}_{\theta}(A, B)} \right\rangle_{P_{A \times B}} \right].$$
 [5]

Here, $\mathcal{F}_{\Theta}: A \times B \to \mathbb{R}$ is a family of functions parameterized by a vector of parameters θ , $P_{A,B}$ is the joint distribution of A and B, and $P_{A \times B}$ is product of their marginal distributions. In our case, since A and B are subsystems of a bigger system, $\langle \cdot \rangle_{P_{A,B}}$ means averaging over samples of A and B taken from the same sample of the bigger system, while $\langle \cdot \rangle_{P_{A \times B}}$ means averaging over samples of A and B taken independently. Heuristically, the reason that this representation works is that the mutual information measures how much the joint distribution differs from the product of marginal distributions. In fact, $\mathcal{M}(A, B)$ equals the Kubleck-Leibler divergence between these two distributions (11).

While there is much to be said about Eq. 5, for the purpose of this work it suffices to note that it reduces the problem of PHYSICS

calculating \mathcal{M} to an optimization problem, which naturally suggests the prospect of using artificial neural networks (ANNs) to parameterize the function \mathcal{F}_{θ} . This is the core idea of Belghazi et al. (25), which we adopt. In machine-learning language, Eq. 5 is taken to be the loss function of the network.

For the complete implementation details see *SI Appendix*, section 1. In broader strokes, the process is as follows: First, using standard methods, a sizable dataset of samples of the system is produced. Then, for each size of subsystem pair we generate two datasets: one in which the two subsystems are taken from the same larger sample (the "joint" dataset) and another in which each subsystem is sampled independently (the "product" dataset). Then, each of the datasets is fed to an ANN, the two averages in Eq. 5 are calculated, and the weights of the ANN are updated to maximize the loss. This process is repeated until the loss stops improving and \mathcal{M} saturates. We found the exponential moving average useful to reduce noise when estimating \mathcal{M} over the final training epochs. Finally, \mathcal{M} is calculated from the trained ANN by averaging Eq. 5 over a separate dataset, different from the one used to train the network.

Results

To demonstrate the performance and versatility of MICE we chose four systems representing different classes of collective behavior: 1) the 2D ferromagnetic Ising model on a square lattice with coupling constant J = 1, a canonical example of a system with a second-order phase transition; 2) the antiferromagnetic Ising model on a triangular lattice (J = -1), a canonical example of a frustrated system with degenerate ground states (27); 3) the continuous XY model on a square lattice, which has a continuous symmetry and features a topological phase transition (27); and 4) finally, an athermal system of a bidisperse mixture of elastic particles which undergoes a jamming transition when its density is increased above a certain threshold (28). For all these systems our method achieves state-of-the-art performance. In addition, in some cases it provides physical insights about the structure and scales of the emergent behavior, as discussed below.

Spin Models. All three spin models were simulated for a system of 64×64 spins with periodic boundary conditions. The distribution was sampled using standard, well-established methods: The Ising models were simulated using Metropolis Monte Carlo simulations as in ref. 8 and the XY model was simulated using the Wolff algorithm as in ref. 29 (*SI Appendix*, section 2).

Lattice systems can naturally be represented as 2D arrays (the triangular lattice can be represented on a square lattice with diagonal interactions) (27). This allows the usage of one of the most successful ANN architectures to parameterize \mathcal{F} of Eq. 5: feedforward convolutional nets (30, 31). We use one to three convolutional layers, each of 8 to 16 filters of size 3×3 , followed by two fully connected layers, using RELU (rectified linear unit) activation, implemented in PyTorch (32). Complete details about the hyperparameters for each model are given in *SI Appendix*, section 1. We calculate \mathcal{M} between subsystems of sizes ranging from a pair of spins to system size. The entropy of a single spin was trivially calculated using brute-force enumeration.

The deviations of our entropy estimations from known results (22-24) are shown in Fig. 1 *B–D*. In all three cases we see impressive quantitative agreement, to a fraction of k_B , with no fitting parameters. We also benchmark our results against the recently proposed compression-based algorithm (8). Relying on highly optimized code and treating the system as effectively 1D, the compression-based algorithm is obviously much faster, about one to two orders of magnitude in terms of runtime. However, while it captures the trend, it offers substantially inferior accuracy in some cases. For example, the low-temperature behavior of the

antiferromagnetic Ising model (Fig. 1C) is governed by a thermodynamic number of ground states with long-range correlations. There, the error of MICE is smaller by an order of magnitude than that of the compression algorithm method.

It is insightful to compare the performance against another very efficient, albeit naive, estimation of *s*—calculating *s* for a small collection of spins by direct enumeration and neglecting the mutual information (i.e., the last term in Eq. 4). In other words, this is assuming that *S* is extensive. This estimation, which we refer to as "naive extrapolation," provides only slightly worse accuracy than the compression method, as seen in Fig. 1. In all cases, MICE provides the most accurate calculation with a maximal error of 0.06 $k_{\rm B}$ per spin for all of the systems and across all temperatures. In *SI Appendix*, section 3 we also use MICE to estimate the heat capacity, showing it outperforms the standard method based on energy fluctuations, since the latter is hard to sample at low temperatures or near a phase transition.

As presented above, our method requires training an ANN for every temperature. This is computationally costly. For example, a single training run for calculating \mathcal{M} between two 64×32 systems of the ferromagnetic Ising model takes several minutes on a standard personal computer. If we were to generate all points in Fig. 1 in this method, the computation time would reach 1 to 2 d. However, drastic improvements in the calculation time can be obtained by leveraging the similarity of the systems between different temperatures. This is done by using the weights (Θ in Eq. 5) that were obtained by training for a given temperature as the initial conditions of the training process of a different temperature or size. This technique is ubiquitous in the field of machine learning, where it is called "transfer learning" (33). In our case it reduces the training time by one to two orders of magnitude. For additional information see *SI Appendix*, section 1F.

Mutual Information as a Probe. The main purpose of MICE is providing an accurate estimation of S. In addition, the byproduct of the calculation, namely the mutual information between systems at different sizes, which is essentially a decomposition of the entropy to contributions from different length scales, can be an interesting observable in its own right. Here we briefly discuss how it captures insightful aspects of the thermodynamics and can be used to assess the accuracy of the MICE against known limiting behaviors. In passing we note that the mutual information between different scales was shown to be informative in analysis of disordered systems (34, 35).

First, we look at \mathcal{M} between subsystems at various sizes for the ferromagnetic Ising model on a square lattice, plotted in Fig. 2. \mathcal{M} manifestly shows the phase transition (36, 37). Indeed, $d\mathcal{M}/dT$ peaks exactly at the theoretical infinite-system critical temperature $T_c = 2.269J$ (Fig. 2B).*

In addition, the accuracy of our calculation can be corroborated against known limits at both high and low temperatures. For $T \ll T_c$, all spins essentially point in the same direction. To be precise, in the low-T limit the ground-state entropy of the whole system, or any subsystem, is exactly log(2). This implies that the mutual information between any two subsystems is also log(2) which we indeed observe for all subsystem sizes (Fig. 24).

For $T \gg T_c$, the mutual information between two subsystems can be obtained by a rigorous high-T expansion. The calculation is straightforward but lengthy, and for the sake of brevity its details are given in *SI Appendix*, section 4A. However, the result is short and intuitive: The leading-order behavior at high T is

^{*}In second-order phase transitions the entropy is continuous but its temperature derivative (which is proportional to the heat capacity) (1) diverges. Since S is a sum over $\mathcal{M}(X_i)$ (Eq. 4), we expect $d\mathcal{M}/dT$ to diverge, rather than \mathcal{M} .



Fig. 2. Analyzing \mathcal{M} for the 2D Ising model on a square lattice for different subsystem sizes. (A) \mathcal{M} complies with two known limits: At low temperatures $\mathcal{M} = \log(2)$. At high temperatures \mathcal{M} approaches the theoretical value of Eq. **6**, as shown in *Inset* (dashed line). (B) The derivative of the mutual information peaking at the theoretical value $T_c \approx 2.269J$ (23). (C) \mathcal{M} normalized by the interface length for varying subsystem sizes (i.e., number of spins). For visual clarity, all curves are normalized to start at unity at zero area. (D) \mathcal{M} per area as function of area for the ferromagnetic Ising model on a square lattice (squares) and the antiferromagnetic triangular lattice model (triangles) at various temperatures. \mathcal{M} decays faster for the ferromagnetic model, as the correlation lengths are much shorter.

$$\mathcal{M} = \frac{1}{2} \frac{\ell}{T^2}, \quad \text{for Ising model}$$

$$\mathcal{M} = \frac{1}{4} \frac{\ell}{T^2}, \quad \text{for XY model},$$
 [6]

where ℓ is the interface size between the subsystems, i.e., the number of spins in one system that directly interact with spins in the other. As seen in Fig. 2 *A*, *Inset*, our method shows excellent agreement with this prediction, again with no fitting parameters. In passing we note that Eq. 6 is akin to the famous area law in quantum entanglement (38).

That is, when $T > T_c$, the mutual information per interface length is independent of the system size, as expected. However, for $T < T_c$ the entropy is not extensive, and \mathcal{M}/ℓ decays quickly with the size of the subsystem (Fig. 2C). This means that the summands in Eq. 4, which are \mathcal{M} normalized by the 2D volume (i.e., area), decay quickly for large subsystems. This is visualized in Fig. 2D. Fig. 2D also shows that in the antiferromagnetic model the summands decay more slowly, which is expected since it features long-range correlations.

Next, in Fig. 3 we examine the entropy and the mutual information in the XY model. At high temperatures \mathcal{M} decays as described in Eq. 6. Below the critical temperature, the famous Kosterlitz–Thouless transition temperature $T_{KT} = 0.8J$, \mathcal{M} approaches a *T*-independent plateau for $H \neq 0$ and diverges logarithmically when H = 0. This divergence is due to the continuous degeneracy of the XY model, which is lifted in the presence of an external field. In the transition between these limits, \mathcal{M} features a pronounced peak, which becomes smaller and shifts to higher temperatures with increasing H (Fig. 3*C*).

This rich behavior of \mathcal{M} can be understood in simple terms. The high-temperature behavior is accurately described by Eq. 6, which is a further corroboration of our method (Fig. 3B). The low-temperature behavior can be understood, much like in the case of the Ising model, in terms of collective behavior. For $H \neq 0$ and $T < T_{KT}$ all spins are mostly aligned with the field, even if it is relatively small, because of the broken symmetry. In this case, spins fluctuate mildly around their ground state and a harmonic approximation can be made. Within the harmonic approximation the mutual information, \mathcal{M}_h (the subscript *h* stands for harmonic), can be obtained analytically in terms of block determinants of the Hamiltonian, a derivation which is given in detail in *SI Appendix*, section 4B. The results of this calculation are presented in Fig. 3C and show good quantitative agreement.

Finally, we remark that the generic behavior of \mathcal{M} —a T-independent plateau at low T followed by a peak and a powerlaw decay at large T—is also present in very small systems. In fact, even a system of two spins behaves in a qualitatively similar way, although the transition temperatures between the regimes are quite different due to the collective behavior of the spins (Fig. 3D and SI Appendix, section 5).

A Continuous, Out-of-Equilibrium System

One of the main advantages of MICE is that it is very versatile in terms of the systems it can operate on. As long as a well-defined distribution exists and samples can be drawn from it, and as long as the system can be digitally represented in a manner compatible with ANNs, MICE should be, at least potentially, applicable. In particular, the scheme presented above can be applied to out-of-equilibrium systems, whose entropy calculation is a challenge both technically and conceptually (8, 15, 17, 18, 39, 40). Clearly, the result of MICE will be an estimate of the entropy defined in Eq. 1, which is the information-theoretic definition of entropy. Relating the result to other thermodynamic properties would depend on the details of the system, which is always the case in calculating thermodynamic properties of out-of-equilibrium systems.

Jammed solids are a prominent class of out-of-equilibrium systems whose entropy plays a crucial role in their dynamics (41). In these systems the entropy, which stems from steric interactions, is geometric in nature and measures the number of ways the



Fig. 3. Analysis of the XY model under the external field (*H*) using MICE. (*A*) Entropy as a function of temperature for various external fields. *Inset* shows ds/dT, and T_{KT} is marked with a dashed line. (*B*) Mutual information between two systems of size 32×16 spins, for varying fields. The arrow marks the peak in $\mathcal{M}(H=0)$ at T_{max} . The blue line is the high-temperature limit, Eq. **6**. (C) Two features of the curves in *B* are replotted: the low-*T* plateau value (evaluated at T = 0.1J), compared to the analytically calculated values at $T \rightarrow 0$ in the harmonic approximation, \mathcal{M}_h (black line). T_{max} is plotted in orange circles. (*D*) Exact numerical calculation of \mathcal{M} between two isolated spins for varying *H*, showing qualitatively similar behavior to that in *B* (although note that the temperature axis is logarithmic, unlike in *B*).

system's constituents can be ordered in space without overlap. When this depends sensitively on the density, jamming occurs. The jamming transition is also important as it is thought that understanding it would guide us in understanding one of the most important open problems in condensed-matter physics—the glass transition, which is also intimately related to entropic effects (41–43).

As a representative example, we study here a bidisperse mixture of soft disks. This system exhibits a jamming transition at high densities (44). Several works have attempted to identify the jamming transition of this system, using dynamic properties such as the jamming length scale or the effective viscosity (45) and using static properties such as pair correlations or fraction of jammed particles (44, 45). Recently, Zu et al. (17) tried to measure the entropic signature of the jamming transition and have shown that compression-based methods have failed to do so. The authors of ref. 17 have generously shared their dataset with us, to test our method on, which we do below.

The system is an equimolar bidisperse system of disks with one-sided harmonic interactions (Fig. 4*A*). The simulation is performed in a finite box with periodic boundary conditions. The area density of the particles, ϕ , is a control parameter which is changed by changing the number of particles, *N*. Further details about the simulation are given in *SI Appendix*, section 6. The system is expected to undergo a jamming transition at $\phi_J \approx 0.841$ (28, 45).

There are a few differences between this system and the spin models discussed above. First, it is not a lattice system with discrete states. Rather, here the state space is continuous, parameterized by the positions of the particles. This requires a careful treatment of the discretization scheme. The choice of discretization scheme, and specifically the spatial resolution of discretization, affects the results in a nontrivial manner. Finally, in the analysis of the spin models we employed MICE on subsystems of all sizes, between one spin and the whole system. However, the soft disk systems are so large that doing so will be both impractical and unnecessary (adequate resolution requires $\sim 3 \times 10^6$ pixels, as discussed below). Before describing the results, we briefly discuss how these challenges are resolved, since they are common to many physical systems of interest, both in and out of equilibrium.

Continuous Systems (Differential Entropy). Since the system is continuous, the summation in Eq. 1 should be replaced by integration:

$$\tilde{S} = -\int p(x)\log p(x)dx.$$
[7]

This definition is known as differential entropy. Note that $\log p(x)$ is ill defined since it depends on the choice of units of x in a nonmultiplicative manner.

This nonmultiplicative component, which depends logarithmically on the length unit, is fundamentally related to the fact that the digital representation of the system is discrete and thus the differential entropy of Eq. 7 differs from the discrete entropy of Eq. 1 by a factor that diverges logarithmically with the resolution of the discretization. For a detailed derivation see *SI Appendix*, section 7.

Moreover, we also show there that, quite conveniently, the representation of S in terms of Eq. 4 offers a well-defined way to remove this divergence. While \tilde{S} of a continuous system depends logarithmically on the resolution, \mathcal{M} becomes independent of it



Fig. 4. (*A*) Snapshots from the bidisperse mixture simulation below and above the jamming transition density (ϕ_J). (*B*) A blowup of the marked region in *A*. We discretized the system (colored circles) as Boolean 2D images (black and white pixels). *Top* and *Bottom* show a spatial resolution of $\mathcal{R} = 5$ and $\mathcal{R} = 9.5$, respectively. The pixels are the input to MICE. (*C*) The effect of discretizing with various resolutions (\mathcal{R}) and various densities: \mathcal{M} between two subsystems of size $2\sigma \times 1\sigma$ (*Left*) and $4\sigma \times 2\sigma$ (*Right*). At high resolutions, \mathcal{M} becomes independent of \mathcal{R} . Green and red arrows indicate the resolutions represented in *B*, *Top* and *Bottom*, respectively. Different markers correspond to different densities; see key in *E*. (D) \mathcal{M}/ℓ as a function of the area of the subsystem (\mathcal{A}) at various densities; see key in *E*. For large enough ℓ , \mathcal{M} becomes linear in ℓ . (*E*) \mathcal{M}/A as a function A at various densities. \mathcal{M} becomes negligible for large subsystems. The dashed colored lines represent the extrapolation of Eq. 9, based on the subsystem at the size represented by the black dashed line. (*F*) The density dependence of the excess entropy. *Inset* shows the results of MICE (blue) and the linear trend of \tilde{s}/N at low densities (dashed orange line). For visual clarity, the linear trend in ϕ is subtracted in the main panel. The dashed black line represents the theoretical jamming transition point.

in the limit of very fine resolution. In fact, the necessary resolution is such that no physically relevant information is lost by the discretization, i.e., when all continuous configurations that map to the same discrete representation are equiprobable.

Therefore, when we estimate S according to Eq. 4, we can avoid the logarithmic divergence simply by omitting the first term in the right-hand side. That is, in what follows we do not present \tilde{s} but rather

$$\Delta \tilde{s} \equiv \tilde{s} - \frac{S(X_m)}{V_m} = -\sum_{k=1}^m \frac{\mathcal{M}(X_k)}{2V_k}.$$
[8]

As a side note, we remark that the omitted term, $S(X_m)/V_m$, is simply the entropy density of the smallest subsystem. It corresponds to the entropy of an "ideal gas" composed of copies of the smallest subsystem. Subtracting the entropy of an ideal gas is common in entropy calculations of thermodynamic systems (17, 39). The result of the subtraction is commonly referred to as "excess entropy."

Discretization. Since convolutional ANNs show state-of-the-art capabilities in extracting information from images, we discretize phase space by mapping a state of the system to a 2D image, whose pixels are black if they contain a center of a particle (Fig. 4B).^{\dagger} The spatial resolution of the image is a hyperparameter of our method. We measure the resolution with the dimensionless number $\mathcal{R} = \sigma/p$, where p is the spatial extent of a pixel and σ is the diameter of the smaller disk. Based on the discussion above, we expect the estimation of \mathcal{M} to converge to a constant value when \mathcal{R} is increased. This is indeed the case, as demonstrated in Fig. 4C. In what follows, we use $\mathcal{R} = 10$, for which \mathcal{M} is converged. We note that in terms of resources, the computational cost of discretizing the system is negligible compared to simulating the system or training the ANN. In addition, as shown below, the ANN does not have to be applied on the whole system, so a fine discretization does not lead to a memory bottleneck, at least not in 2D.

Extrapolating the Mutual Information. The resolution required for convergence necessitates $\sim 10^6$ pixels to discretize the whole system. Feeding such a large image to an ANN might be possible, but requires unreasonable computational resources for the task at hand. Luckily, this is not necessary.

As discussed above, for large enough subsystems, that is, scales much larger than the longest correlation length of the system, we expect \mathcal{M} to grow linearly with the interface length (Fig. 2C). In precise terms, we expect

$$\mathcal{M}(X_k) = \frac{\ell_k}{\ell_n} \mathcal{M}(X_n).$$
 [9]

If we assume this is obeyed for all systems larger than X_k , this relation can be used to replace the summands in Eq. 4, and the summation can be done analytically without calculations on subsystems larger than X_k . Fig. 4D shows that this happens for subsystems of length $\sim 4\sigma$. In Fig. 4E we show that Eq. 9, based on the values of \mathcal{M} for this size, quantitatively reproduces the

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values of the summands of Eq. 4 for sizes larger than 4σ , i.e., a 2D volume of $A = 16\sigma^2$.

Results. We are now in position to calculate the entropy of the whole system for various densities. Assuming that Eq. 9 is satisfied for n > m, Eq. 4 can be analytically summed, yielding (*SI Appendix*, section 8)

$$s = s(x_m) - 2\frac{\mathcal{M}(X_m)}{V_m}.$$
[10]

Fig. 4 *F*, *Inset* shows $\Delta \tilde{s}/N$ as a function of ϕ . It is seen that at low densities $\Delta \tilde{s}$ depends roughly linearly on the density (dashed orange line). To emphasize the phase transition, in the main panel of Fig. 4*F* we plot the same data with this linear trend subtracted. The change in the behavior of $\Delta \tilde{s}$ around the expected jamming point is evident. Importantly, we remind the reader that compression-based entropy estimations were less successful in showing this transition (section 3.5 of ref. 17). A more detailed comparison with the results of ref. 17 is given in *SI Appendix*, section 9.

Discussion and Conclusion

Machine-learning algorithms in general, and neural networks in particular, offer an effective tool to identify patterns in high-dimensional data with complex correlation structure. We have shown that these capabilities can be leveraged to tackle another important challenge—computing the entropy of physical systems.

The crux of the method is mapping the problem of entropy calculation to an iterative process of mutual information estimation. By doing so we were able to estimate the entropy of canonical statistical physics problems, both discrete and continuous, both in and out of equilibrium, outperforming compression-based entropy estimation methods. Finally, we demonstrated that MICE naturally allows us to decompose the entropy into contributions from different scales, providing an insightful diagnostic for the thermodynamics of physical systems.

We surmise that MICE could be a promising tool for the study of many important systems, such as the configurational entropy of amorphous solids (46), the entropy crisis of glassy systems (42), entropy of active matter (40), and more. The main limit of the proposed method would depend on the minimal system size for which Eq. 9 applies, which determines the largest input for which an ANN should be trained. This is the dominant factor in the computational cost of our method. In addition, we believe that with adequate modifications MICE could be used on quantum systems, for which the mutual information is fundamentally related to entanglement of quantum states (47). A relevant direction could be the extraction of entropy from quantum Monte Carlo simulations. These directions will be explored in future works.

Data Availability. All study data are included in this article and SI Appendix.

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