



Publication Year	2015
Acceptance in OA	2020-04-10T10:30:53Z
Title	Emergence of a collective crystal in a classical system with long-range interactions
Authors	TURCHI, ALESSIO, Fanelli, D., Leoncini, X.
Publisher's version (DOI)	10.1209/0295-5075/111/30011
Handle	http://hdl.handle.net/20.500.12386/23981
Journal	EUROPHYSICS LETTERS
Volume	111

Emergence of a collective crystal in a classical system with long-range interactions

ALESSIO TURCHI¹, DUCCIO FANELLI² AND XAVIER LEONCINI³

¹ *INAF Osservatorio Astrofisico di Arcetri, Largo Enrico Fermi 5, I-50125 Firenze, Italy*

² *Dipartimento di Fisica e Astronomia, Università di Firenze and INFN and CSDC,s via Sansone, I-50019 Sesto Fiorentino, Firenze, Italy*

³ *Aix Marseille Université, Université de Toulon, CNRS, CPT UMR 7332, 13288 Marseille, France*

PACS 05.20.-y – Classical statistical mechanics
 PACS 05.45.-a – Nonlinear dynamics and chaos

Abstract. - A one-dimensional long-range model of classical rotators with an extended degree of complexity, as compared to paradigmatic long-range systems, is introduced and studied. Working at constant density, in the thermodynamic limit one can prove the equivalence with the Hamiltonian Mean Field model (HMF) and α -HMF: a second order phase transition is indeed observed at the critical energy threshold $\varepsilon_c = 0.75$. Conversely, when the thermodynamic limit is performed at infinite density (while keeping the length of the hosting interval L constant), the critical energy ε_c is modulated as a function of L . At low energy, a self-organized collective crystal phase is reported to emerge. According to numerics, the crystal phase is split into two distinct regimes depending on the value of the energy per particle ε : for small ε , particles are exactly located on the lattice sites. At variance, above a given energy threshold ε^* , such correspondence is broken and particles can travel from one site to the other. However, ε^* does not signal a genuine transition but rather reflects the finite time of observation, as imposed in the simulations. Transport between adjacent crystal sites occurs at an exponentially small rate, and a sufficiently large time, even extraordinarily long, has to be allowed for the mixing to take place at small energies. The equilibrium one particle density function is computed by maximizing the entropy, and returns an analytical insight into the scrutinized phenomena.

The study of long-range interacting systems experienced a renewed interest, in the last decades, due to the increasing computational resources made available on modern computers. Improved numerical simulations allowed to enhance the statistics over previous investigations, enabling for new ideas to be tested and existing theory to be challenged [1, 2]. In a long-range system the two body potential decays with the distance r as $r^{-\alpha}$, the exponent α being smaller than the dimension of the embedding space. Many different systems fall in such a wide category. Among others, it is worth mentioning self-gravitating systems and their applications to astrophysics and cosmology [3, 4], charged plasma dynamics [5, 6], hydrodynamics of two-dimensional vortices [7, 9], spin-wave interactions [10, 11] with their implications for the physics of lasers [12]. As largely testified by recent advances, long-range systems display a rich zoology of peculiar behaviors, ranging from ensemble inequivalence to out-of equilibrium dy-

namics, and constitute an intriguing arena for novel developments, of both applied and fundamental interests.

Several long-range models exist in the literature, that are in principle suitable for elucidating key aspects of both equilibrium and out-of-equilibrium dynamics. Simplified toy models and paradigmatic case studies have been in particular proposed, which allow to progress in the analysis by imposing a substantial reduction in the inherent complexity of the inspected physical problems. Working along these lines, it is instructive to elaborate on the ingredients that drive the spontaneous emergence of coherent self-organized structures, as seen in real experiments. As an example relevant for the topic addressed in this Letter, crystal-like structures have been reported to occur in long-range physical systems. as e.g. in dusty charged plasmas [14, 15], Coulomb-interacting cold atoms [16] or Bose-Einstein condensates [17]. The phenomenology of these latter states is not clearly understood: experiments seem

to suggest the presence of an intermediate phase during the melting of the solid structure, which displays additional ordering at different scales [14]. When long-range interactions are active, the interplay between potential and kinetic energy terms give rise to correlations that do not decay rapidly with the distance. This latter effect may eventually translate into the emergence of non trivial thermodynamic transitions and peculiar symmetries.

Available long range models often lack the necessary degree of phase space complexity to effectively reproduce the characteristics of such patterns. For instance, in the celebrated Hamiltonian Mean Field (HMF) model the spatial extension of the phase space is bounded in size, each particle's position being mapped onto the circle $\theta \in [0, 2\pi[$. Furthermore, the notion of inter-particles' distance is not explicitly accounted for in the framework of the HMF formulation. Other long-range solvable systems, albeit spatially extended, may lack of translational invariance, like the ϕ^4 model [1, 2], or encode for a peculiar lattice metric, as it is the case for the so called α -HMF model [1, 2]. This latter has a predetermined skeleton that is embedded in the model itself, and which reflects in the observed structures as a natural imprint.

To bridge this gap and eventually describe the appearance of crystal-like long-range structures, we will here introduce and characterize a new model of coupled rotators, closely inspired to the aforementioned α -HMF and HMF models. The model that we are going to introduce will spontaneously freeze into a crystal-like phase, at sufficiently low energies. When the energy gets increased, the self-organized crystal melts, resulting in a disordered gaseous medium. The Letter is organized as follows: we first briefly recall the thermodynamic properties of both the HMF and α -HMF models. We then move to presenting the extended model, focusing on its equilibrium thermodynamical features, and demonstrating the existence of a crystal phase, via combined numerical and analytical means.

The HMF model has imposed itself in recent years as a paradigmatic model for the study of long range interacting systems [18]. Indeed, it displays a rich variety of features, most notably related to its out-of-equilibrium dynamics. Moreover, computing its statistical properties via direct simulations of the microscopic dynamics is both simple and fast. The HMF model is characterized by the following Hamiltonian:

$$H = \sum_{i=1}^N \frac{p_i^2}{2} + \frac{1}{2N} \sum_{i,j=1}^N (1 - \cos(q_i - q_j)), \quad (1)$$

where q_i and p_i stand for the canonically conjugated variables of particle i , and N denotes the total number of particles. The HMF model displays, at equilibrium, a second order phase transition at critical energy $\varepsilon_c = 0.5$, with a characteristic order parameter defined as the global magnetization $\mathbf{M} = \frac{1}{N} \sum_i (\cos(q_i), \sin(q_i))$. The associ-

ated phase space is bounded: the positions of the particles can be mapped on the unitary circle, because of the periodicity of the potential and the mean-field nature of the interaction. In order to explicitly take into account the decay of the interaction potential as due to long-range couplings, an extension of the HMF model referred to as the α -HMF model [19, 20] has been proposed in the literature. In this generalized model, the rotators occupy the sites of a regular lattice and the Hamiltonian writes:

$$H = \sum_{i=1}^N \frac{p_i^2}{2} + \frac{1}{2\tilde{N}} \sum_{i,j=1}^N \frac{1 - \cos(q_i - q_j)}{\|i - j\|^\alpha}, \quad (2)$$

where α is a free parameter, which enables one to adjust the strength of the interaction, and $\|i - j\|$ represents the distance between the lattice sites i and j . The normalization factor reads $\tilde{N} = (2/N)^\alpha + 2 \sum_{i=1}^{N/2-1} (1/i^\alpha)$ and ensures extensiveness. For finite sizes, in order to keep the system statistically invariant by translation along the lattice, it is convenient to impose periodic boundary condition. To accomplish this, it is customary to confine the lattice on a circle of radius $(N - 1)/2\pi$ and then consider in (2) the minimum distance on the circle between any pair of selected sites. For $\alpha < 1$, the α -HMF is thermodynamically equivalent to the HMF model [20–22, 31], albeit it displays a lattice ordering enforced by the model in its microscopic formulation.

Starting from these premises, and to eventually explore the possibility of generating a non trivial crystal-like phase, which is not the mere byproduct of the underlying discrete spatial support, we here propose a straightforward generalization of the aforementioned α -HMF model, that we will refer to as to the β -HMF. Particles are no longer forced to sit on specific lattice sites, but can freely explore the hosting support. In practice, we substitute in Hamiltonian (2) the term $\|i - j\|$ by $\|q_i - q_j\|$, a replacement which does not produce divergences in the force field, as long as $\alpha < 1$. Introducing a continuum distance parameter, allows in turn to extend the phase space that particles can explore while, at the same time, removing the artificial constraint of a fixed lattice support. The β -HMF model is in some respects reminiscent of the self-gravitating ring model [30], but, as opposed to this latter, it does not require the introduction of an artificial regularization of the potential at small scales. In this Letter, we will always deal with periodic boundary conditions on a circle of length L . Then, we will consistently define the distance between two particles with position q_i and q_j as $d_{ij} = \min\{|q_i - q_j|, L - |q_i - q_j|\} := \|q_i - q_j\|$ namely, the shortest arc-length on the circle. Based on the above, the Hamiltonian of the β -HMF can be cast in the form [23]:

$$H = \sum_{i=1}^N \frac{p_i^2}{2} + A(N, L) \sum_{i,j=1}^N \frac{1 - \cos(q_i - q_j)}{\|q_i - q_j\|^\alpha} \quad (3)$$

When $0 \leq \alpha < 1$ the system is formally long-range:

this latter parameter can be tuned to control the strength of the long-range interaction. $A(N, L)$ is a normalization constant which guarantees extensiveness, following the Kac prescription [26], and reads $A = \frac{(1-\alpha)}{N} \left(\frac{L}{2}\right)^\alpha$. The above relation can be rigorously derived in the large scale limit and holds in general, as confirmed by numerical simulation. As a consistency argument, we remark that the expected normalization factor are recovered both in the HMF ($\alpha = 0$) and α -HMF ($q_i = 2\pi k + \xi_i$, for $L \rightarrow \infty$, $\xi_i \in [0, 2\pi[$ and $k \in [0, L[$) limits. As a further, preliminary comment, we note that as q_i are assigned to label the particles' positions, in the interaction potential one should write in principle $\cos(z(q_i - q_j))$, where z is a suitable constant that makes the argument of the cosine adimensional. Without loss of generality, we set $z = 1$ in the following. Moreover, in this Letter, we will assume $L = 2\pi l$, $l \in \mathbb{N}$, and introduce the rescaled density $\rho_0 = N/l$ to simplify the notation.

We start our analysis by focusing on the low temperature regime. In this limit we expect to obtain minimal energy states. From the two-body interaction potential, one can imagine that the system will self-organize so to fill distinct available minima. It is however not evident that the process should eventually materialize in a crystal like structure, condensation of all particles in just one minimum being a priori also possible. Indeed, the further apart the particles sit, the wider the potential minimum appears. Hence, for a fixed energy amount, particles are more prone to fluctuate if they populate distant sites, so endowing a global repulsive entropic force. Since we assume particles to populate a bounded circular domain, we expect that the entropic repulsion will oppose condensation, so favoring in turn the spontaneous emergence of the crystal state, at low temperature.

To test the adequacy of this interpretative scenario, we carried out a campaign of N -body simulations. These are performed by using a fifth-order symplectic scheme [28], with a time-step $\delta t = 0.05$. The equation of motion can be derived from Hamiltonian (3). We wish to emphasize that the explicit presence of the inter-particle distance in (3) heavily increases the required computational cost, as compared to the simple HMF case study, or even the α -HMF model [23]). For this reason, our analysis is limited to a maximum of $N = 10^3$ particles. The time needed to converge to equilibrium gets clearly larger, as the population size is increased. As an additional remark, we notice that for $N = 2$ the system is integrable. In this limiting case, one can analytically access the phase space of the system, and use the information to test the numerical implementation. It is indeed extremely useful to benchmark numerical and analytical outcomes to avoid technical pitfalls which might originate when the subtle condition $\|q_i - q_j\| \sim 0$, is dynamically met. Furthermore both total energy and momentum are constants of motion,

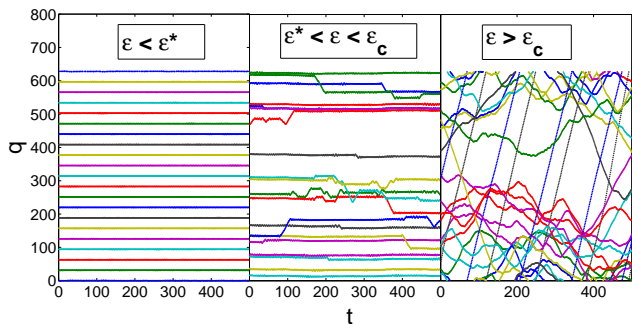


Fig. 1: A few tracer particles positions in the q -space as function of time. The analysis of the recorded trajectories clearly show that three dynamical regimes exist for the β -HMF model (3), depending on the energy per particle ε . At $\varepsilon = \varepsilon^*$, the perfect crystal like structure is lost, and the particles can wander to neighboring sites. Above ε_c the self-organized crystal melts, resulting in a disordered gaseous medium. Here $N = 500$, $L = N$ and $\alpha = 0.5$.

due to the translational invariance, and were numerically monitored over time. All simulations presented in this Letter assume an initial Gaussian distribution of momenta, with total momentum $P = 0$. For any given value of the energy, the system is observed to converge to the same equilibrium state, independently of the distribution positions q , assigned at $t = 0$.

To characterize the thermodynamic fate of the system one can in principle rely on the magnetization $M = |M|$. When particles are uniformly spread over the hosting domain, we have $M = 0$. Conversely, $M = 1$ is equally associated to bunching on a single cluster, the condensed phase to which we alluded above, or to a distribution peaked on sites which are 2π -periodic, namely the sought crystal state. To remove this degeneracy and discriminate between a crystal ordering phase and a condensate solution with all particles close to each other in space, one needs to compute the distribution of relative particles' positions.

We first start by qualitatively looking at some particle trajectories, q versus time, t , as obtained through direct N -body simulations. Those are displayed in Fig. 1. Three rather different equilibrium regimes can be clearly detected. At very low energy densities, the system self-organizes into a crystal in which particles are bounded in the vicinity of what appear to be lattice sites. When increasing the energy per particle above a threshold amount that we termed ε^* , the system enters a partially disordered phase: particles are nevertheless localized over finite windows of time, and the global magnetization is still positive. At higher energies, for $\varepsilon > \varepsilon_c$, the system loses any traces of regularity and the crystal melts into a gaseous medium.

To cast this observation on solid grounds, we calculated analytically the single particle distribution as follows a

maximization of the thermodynamic entropy, subject to the constraints of the dynamics. As compared to the HMF and α -HMF models, we have one more degree of freedom in choosing the thermodynamic limit. The system now depends on both the rescaled length l and the average particle density ρ_0 . When performing the limit $N \rightarrow \infty$ we can consider $\rho_0 \rightarrow \infty$ at fixed l , or, conversely, the limit $l \rightarrow \infty$ at constant ρ_0 . Indeed, the dependence on both ρ_0 or L can be made explicit in the scaling constant A . The Hamiltonian in the infinite density limit ($N \rightarrow \infty$ with l constant) reads:

$$\mathcal{H}(q, p) = \frac{p^2}{2} + \frac{1 - \alpha}{2\pi L} \left(\frac{L}{2}\right)^\alpha \int_L \frac{1 - \cos(q - q')}{\|q - q'\|^\alpha} dq'. \quad (4)$$

In the dual limit, when the density is kept constant and $l \rightarrow \infty$, it is not so straightforward to analytically recover the analogue Hamiltonian. The problem lies in that the potential is modulated by the cosine with a periodicity of 2π , but it is not periodic. So we are not allowed to perform trivial simplifications without any knowledge on the form of the distribution function. Still we can solve the system for finite length, and observe the scaling of the associated thermodynamic quantities for $l \rightarrow \infty$. Let us denote with $f(q, p, t)$ the single particle distribution function: $f(q, p, t)dqdp$ represents the fraction of particles in the interval $[q, q + dq][p, p + dp]$ at time t . The system conserves the energy density $\varepsilon[f] = \int_\Gamma f(q, p, t)\mathcal{H}(q, p)dqdp$ where $\mathcal{H}(q, p)$ follows equation (4). Also the total momentum $P[f] = \int_\Gamma f(q, p, t)pdqdp$ is conserved together with normalization condition $\mathcal{G}[f] = \int_\Gamma f(q, p, t)dqdp = 1$ where Γ represents the phase space volume. Assume then a strictly Boltzmannian entropy $S[f] = -\int_\Gamma f(q, p, t)\log(f(q, p, t))dqdp$. To obtain the equilibrium function $f(q, p)$ which maximizes the entropy $S[f]$, under the constraints of the dynamics imposed, we have to solve the maximization problem $\max_f \{S[f] \mid \int_\Gamma f(q, p, t)dqdp = 1, \varepsilon[f] = E, P[f] = P\}$. By introducing the Lagrange multipliers β , γ and μ , and requiring stationarity of the entropy functional, one readily obtains:

$$f(q, p) = D e^{-\beta(\frac{p^2}{2} + V(q))}, \quad (5)$$

where

$$V(q) = \int \int f(q', p) \frac{1 - \alpha}{2\pi L} \left(\frac{L}{2}\right)^\alpha \frac{1 - \cos(q - q')}{\|q - q'\|^\alpha} dpdq'. \quad (6)$$

and $D = e^{-1-\mu}$. In the above calculation we made use of $\gamma = 0$, as $P[f] = 0$, for our class of initial conditions. Expression (5) is the standard Maxwellian distribution, where β is the inverse temperature and μ stands for the chemical potential. As for the HMF model, one obtains an implicit system, since $V(q)$ depends in turn on $f(q, p)$. To solve the problem we use an iterative numerical scheme introduced in [30], which generalizes the standard Newton technique by ensuring that entropy steadily increases towards the maximum value at each iteration step. From

equation (5) we define the equilibrium density function $\rho(q)$ as:

$$\rho(q) = \int f(q, p)dp = D \sqrt{\frac{2\pi}{\beta}} e^{-\beta V(q)}. \quad (7)$$

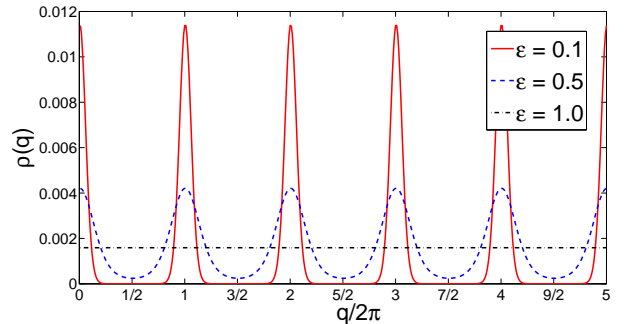


Fig. 2: Analytically predicted equilibrium distribution densities $\rho(q)$ for different dynamical regimes and $\alpha = 0.5$. Here we plot a small portion of the q axis with 5 periods for a distribution function computed for $l = 100$ periods. The crystal lattice appears as expected.

A comparison between the theoretically predicted $\rho(q)$ and the result of numerical simulations can be easily drawn for $\varepsilon < \varepsilon_c$, when particles occupy the self-emerging lattice, and for $\varepsilon > \varepsilon^*$, when particles are homogeneously spread. In both cases, we observe a good matching between theory and simulations, even with a low number of particles and low densities (data not shown). In the intermediate phase, averaging effects due to particles relocations makes the direct comparison more cumbersome, since one has to cope with the numerical artifacts associated to the lack of local translational invariance, at finite times. To overcome these limitations, in figure 3 we confront the analytical predictions for the distribution of the inter-particle distances $\mathcal{C}(d) = \int \rho(q + d)\rho(q)dq$ to the simulated data. $\mathcal{C}(d)$ is by definition translational invariant and the agreement between theory and simulations is almost perfect, for all the explored energy regimes.

As an additional test, we proceed to compute the curve for the equilibrium magnetization $M(\varepsilon)$, for different values of the scaling exponent α and for $l = 1000$. A second order phase transition between ferromagnetic $M > 0$ and paramagnetic $M = 0$ phases is found to occur at a critical energy density ε_c , see Figure 4. Remarkably, for $\alpha < 0.6$ (for the chosen value of l) the data collapse on the HMF transition curve and $\varepsilon_c = 0.75$. For larger values of α the curves shifts towards the left and, consequently, ε_c gets reduced. The energy threshold ε^* (as determined numerically) which putatively separates regular and disordered (soft) crystals, does not leave a trace in the magnetization curves. We shall return on this point later on. For both large and small values of the scaling exponent α the agreement between theory and simulations is convincing.

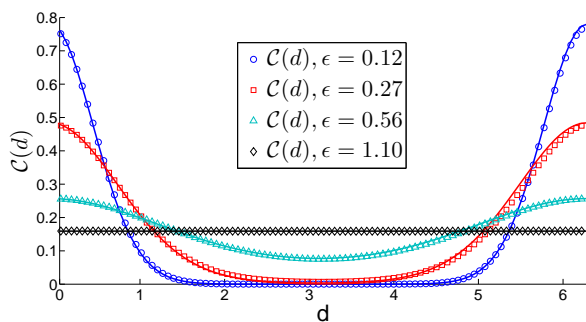


Fig. 3: The theoretical distribution $\mathcal{C}(d)$ of inter-particle distances for infinite density (solid lines) is compared to direct simulations for $\rho_0 = 1$ (symbols). Here $l = 500$ and $\alpha = 0.5$. Different curves correspond to distinct energy densities ε . The horizontal axis is rescaled in a $\text{mod}(2\pi)$ representation and the curves are normalized so to have unitary area. $\mathcal{C}(d)$ is translationally invariant: an almost perfect matching is observed.

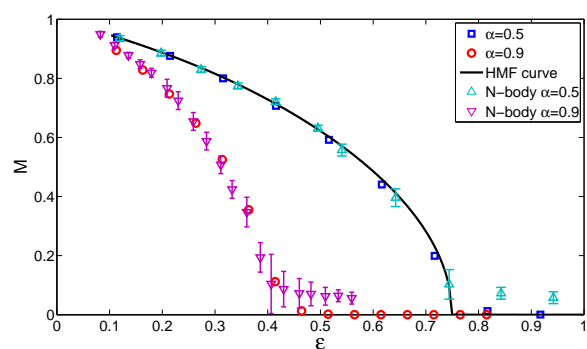


Fig. 4: Comparison between the analytic magnetization curves and the data recorded via N -body simulations, for a random flat initial condition over the space positions. Data refer to $l = 1000$. An average over a time window $\Delta t = 5000$ is performed. As expected data are noisy close to the critical transition energy ε_m . The black thick curve represents the HMF equilibrium curve.

A dedicated campaign of investigations suggests that, for all values of α , the transition energy ε_c converges to the asymptotic value 0.75, in the limit $l \rightarrow \infty$. In this limit, the transition curves of the β -HMF and HMF (and clearly α -HMF) superpose to each other. In the following we will provide an independent argument to show that the β -HMF reduces to the α -HMF, when $l \rightarrow \infty$.

Let us start by simplifying the expression of the distance term which appears in Hamiltonian (3). Define $q_j = 2\pi h_j + \delta_j$, with $h_j \in \{0, \dots, n\}$ playing the role of a lattice index and $0 < \delta_j < 2\pi$. Given the equilibrium distribution $\rho(\delta, h)$ computed above, in the limit $L \rightarrow \infty$, we ignore the contribution to the potential coming from pair of particles i and j which are close and have $h_i = h_j$. Under these working assumptions, for $h_i \neq h_j$ the distance term in equation (3) can be simplified as:

$$\begin{aligned} & \frac{1}{\left\| \frac{2\pi h_i}{L} - \frac{2\pi h_j}{L} + (\delta_i - \delta_j)/L \right\|^\alpha} \approx \\ & \approx \frac{1}{\left\| \frac{2\pi h_i}{L} - \frac{2\pi h_j}{L} \right\|^\alpha} - \alpha \frac{(\delta_i - \delta_j)}{L \left\| \frac{2\pi h_i}{L} - \frac{2\pi h_j}{L} \right\|^{\alpha+1}}. \end{aligned} \quad (8)$$

The second term in expression (8) vanishes, when performing the sum in the limit $L \rightarrow \infty$. One loses therefore the dependence on δ . In other words, the numerator and denominator of the potential in equation (3) become independent and the potential scales as $\sum_{j=1}^N (1 - \cos(\delta_i - \delta_j)) / (\|h_i - h_j\|^\alpha)$. Since the equilibrium distribution function is homogeneous in the index h and since each interval (of length 2π) in which we have subdivided the system, contains the same number of particles, we can reorder the sum in the potential by avoiding to individually count the particles which share the same index h_i . We then perform the replacement $h_i \rightarrow i$. Apart for a multiplying constant depending on ρ_0 , we recover hence the form of the α -HMF potential (2). Since the steps car-

ried out above are independent of α , it can be speculated that the α -HMF equilibrium solution, equivalent to that associated to the original HMF model, is also found within the β -HMF setting, for any value $0 \leq \alpha < 1$, provided l is sufficiently large.

We now turn back to Fig. 3. For low energies the distribution appears to display a classical crystalline structure. Particles are confined on lattice sites, as enforced by the fact that $\rho(q)$ displays a periodic set of minima with $\rho \sim 0$. On the other hand, for higher energies the minimum ρ_{min} of $\rho(q)$ is positive: even though the crystal seems to preserve a well defined structure, this latter reflects an average collective behaviour. Indeed there is no one-to-one correspondence between particle positions and lattice sites, and fluctuations instigate an effective mixing between adjacent lattice sites. In order to check whether a transition between the two aforementioned crystalline phases exist, one must determine if $\rho_{min} = 0$, below a given finite energy density. For this purpose, we build on the observed analogy between the β -HMF and α -HMF models, and thus the expected equivalence with the HMF, in the limit for $l \rightarrow \infty$. The distribution of densities obtained for the β -HMF model superposes exactly (data not shown) to those obtained for the the HMF, which are known almost explicitly. The interest of this conclusion are manyfolds. On the one side, this observation implies that the universality class of the HMF extends to embrace non trivial models, as the β -HMF is. Then, $\rho_{min} \sim \exp(-\beta M)$ for $q = \pi$, meaning that ρ_{min} formally vanishes only at zero temperature. Hence, at zero energy density (zero temperature), we have a perfect crystal lattice, particles being *de facto* trapped on specific lattice sites. For a non zero, but still small, energy density, we end up with a collective crystal structure with particles that can in principle move from one site to another, while preserving a macroscopic crystal-like self-organization. The lattice emerges as a collective property

of the distribution, and the trapping is effective for finite, though potentially large, times. The critical energy ε^* separating perfect and soft crystal regimes reflects therefore the finite time of observation rather than representing a genuine transition between two distinct equilibrium states.

Summing up, we have here introduced and studied a novel long-range model, to shed further light on those spatial self-organized structures that can emerge when long-range coupling are at play. A continuous inter-particles distance term adds a degree of complexity with respect to the simplified HMF and α -HMF models, while reproducing similar thermodynamics behaviours. In the extended phase space of the β -HMF it is possible to observe a more rich phenomenology of equilibrium features. Specifically, we unravelled three different dynamical states, encompassing a regular crystal phase, a disordered soft crystal phase and a gaseous phase at high energies, which could not be seen in the reduced phase space of previous long-range models. The crystal phases manifest as a generalized self-organized state, in which the local lattice emerge from the collective behaviour of the particles. A solid to gas transition manifests at a critical energy which is sensitive to the scaling size L , and converges to 0.75 in the limit $L \rightarrow \infty$. The distinction between perfect and soft crystal is instead meaningful, only if the time of observation is finite. We were able to analytically obtain a form for the equilibrium distribution function and apply it to describe the observed system behavior. Assuming the validity of the latter solution, the system can be formally reduced to the HMF in the limit $l \rightarrow \infty$, thus reproducing the same equilibrium transition between magnetized and homogeneous states. When operating at constant l , but infinite density, the β -HMF returns a different equilibrium transition curve as compared to the HMF model. Further studies should aim at evaluating the interest of the disordered crystal phase predicted within the realm β -HMF model, in light of its potential applications to real long-range systems of experimental relevance. For instance the fact there is no direct correspondence between emerging crystal sites and effective particle position could be reminiscent of supersolid properties. Another open question has to do with the possible presence of quasi-stationary out-of-equilibrium states which were non investigated in this paper.

* * *

X. L. thanks M. Adda-Bedia and T. L. Van den Berg for useful discussions during the setting up of the β -HMF model. X. L. is partially supported by the FET project Multiplex 317532.

REFERENCES

- [1] A. Campa, T. Dauxois, S. Ruffo, Physics Reports **480**, 57-159 (2009).
- [2] A. Campa, T. Dauxois, D. Fanelli, S. Ruffo, Physics of Long Range Interacting Systems, Oxford University Press (2015).
- [3] D. Lynden-Bell, R. Wood, Mon. Not. R. Astron. Soc. **138** 495 (1968).
- [4] P. H. Chavanis, I. Ispolatov, Phys. Rev. E **66**, 036109 (2002).
- [5] M. K. H. Kiessling and T. Neukirch, Proc. Nat. Acad. Sci. USA **100**, 1510 (2003).
- [6] P. H. Chavanis, Eur. Phys. J. B **52**, 61 (2006).
- [7] R. A. Smith and T. M. O'Neil, Phys. Fluid. B **2**, 2961 (1990).
- [8] E. Caglioti, P. L. Lions, C. Marchioro and M. Pulvirenti, Commun. Math. Phys. **174**, 229 (1990).
- [9] R. S. Ellis, K. Haven, and B. Turkington, Nonlinearity **15**, 239 (2002).
- [10] J. Barré, D. Mukamel, S. Ruffo, Phys. Rev. Lett. **87**, 030601 (2001).
- [11] D. Mukamel, S. Ruffo and N. Schreiber, Phys. Rev. Lett. **95**, 240604 (2005).
- [12] J. Barré T. Dauxois, G. de Ninno, D. Fanelli, S. Ruffo, Phys. Rev E **69**, 045501(R) (2004).
- [13] A. Pluchino, V. Latora, A. Rapisarda, Phys. Rev. E **69**, 056113 (2004).
- [14] H. M. Thomas, G. E. Morfill, Nature **379**, 806 - 809 (1996).
- [15] M. H. Thoma, M. Kretschmer, H. Rothermel, H. M. Thomas, G. E. Morfill, Am. J. Phys. **73**, 5 (2005).
- [16] A. Bermudez, J. Almeida, F. Schmidt-Kaler, A. Retzker, M. B. Plenio, arXiv:1108.1024v1 (2011).
- [17] Kazimierz Lakomy, Rejish Nath, Luis Santos, arXiv:1107.3132v1 (2011).
- [18] M. Antoni and S. Ruffo, Phys. Rev. E **52**, 2361 (1995).
- [19] C. Anteneodo, C. Tsallis, Phys. Rev. Lett. **80**, 5313 (1998).
- [20] F. Tamarit, C. Anteneodo, Phys. Rev. Lett. **84**, 208 (2000).
- [21] T. L. Van den Berg, D. Fanelli, X. Leoncini, EPL **89**, 50010 (2010).
- [22] R. Bachelard, T. Dauxois, G. De Ninno, S. Ruffo, F. Staniscia, Phys. Rev. E **83**, 061132 (2011).
- [23] T. L. Van den Berg, *Systèmes avec interaction à longue portée*, Master thesis, director X. Leoncini (2009).
- [24] W. Braun and K. Hepp, Commun. math. Phys. **56**, 125 (1977).
- [25] A. Antoniazzi, F. Califano, D. Fanelli, S. Ruffo, Phys. Rev. Lett. **98**, 150602 (2007).
- [26] M. Kac, G. E. Uhlenbeck, P. C. Hemmer, J. Math. Phys. **4**, 216 (1963).
- [27] X. Leoncini, T. L. Van den Berg, D. Fanelli, EPL **86** 20002 (2009).
- [28] R. I. McLachlan, P. Atela, The accuracy of symplectic integrators, Nonlinearity **5**, 541-562 (1992).
- [29] A. Antoniazzi, D. Fanelli, J. Barré, P. H. Chavanis, T. Dauxois, S. Ruffo, Phys. Rev. E **75**, 011112 (2007).
- [30] T. Tatekawa, F. Bouchet, T. Dauxois, S. Ruffo, Phys. Rev. E **71**, 056111 (2005).
- [31] A. Turchi, D. Fanelli, X. Leoncini, Comm. Nonlin. Sci. Num. Sim. **16**, 4718-4724 (2011).