COARSE-GRAINING SCHEMES FOR STOCHASTIC LATTICE SYSTEMS WITH SHORT AND LONG-RANGE INTERACTIONS

MARKOS A. KATSOULAKIS, PETR PLECHÁČ, LUC REY-BELLET, AND DIMITRIOS K. TSAGKAROGIANNIS

Abstract. We develop coarse-graining schemes for stochastic many-particle microscopic models with competing short- and long-range interactions on a $d$-dimensional lattice. We focus on the coarse-graining of equilibrium Gibbs states, and by using cluster expansions we analyze the corresponding renormalization group map. We quantify the approximation properties of the coarse-grained terms arising from different types of interactions and present a hierarchy of correction terms. We derive semi-analytical numerical coarse-graining schemes that are accompanied by a posteriori error estimates for lattice systems with short- and long-range interactions.

1. Introduction

Many-particle microscopic systems with combined short and long-range interactions are ubiquitous in a variety of physical and biochemical systems. They exhibit rich mesoscopic and macroscopic morphologies due to the competition of attractive and repulsive interaction potentials. For example, mesoscale pattern formation via self-assembly arises in heteroepitaxy; other notable examples include polymeric systems, and micromagnetic materials. Simulations of such systems rely on molecular methods such as kinetic Monte Carlo (kMC) or Molecular Dynamics (MD). However, the presence of long-range interactions severely limits the spatio-temporal scales that can be simulated by such direct computational methods.

Received by the editor March 8, 2010 and, in revised form, March 22, 2011 and November 9, 2011.

2010 Mathematics Subject Classification. Primary 65C05, 65C20, 82B20, 82B80, 82-08.

Key words and phrases. Coarse-graining, lattice spin systems, Monte Carlo method, Gibbs measure, cluster expansion, renormalization group map, sub-grid scale modeling, multi-body interactions.

The research of the first author was supported by the National Science Foundation through grants NSF-DMS-0715125, the CDI-Type II award NSF-CMMI-0835673, and the European Commission FP7-REGPOT-2009-1 project “Archimedes Center for Modeling, Analysis and Computation”.

The research of the second author was partially supported by the National Science Foundation under grant NSF-DMS-0813893 and by the Office of Advanced Scientific Computing Research, U.S. Department of Energy under DE-SC0001340; the work was partly done at the Oak Ridge National Laboratory, which is managed by UT-Battelle, LLC under Contract No. DE-AC05-00OR22725.

The research of the third author was partially supported by grant NSF-DMS-06058.

The research of the fourth author has been supported by a Marie Curie Intra European Fellowship within the 7th European Community Framework Program.

©2014 American Mathematical Society
On the other hand, an important class of computational tools used for accelerating microscopic molecular simulations is the method of coarse-graining. By lumping together degrees of freedom into coarse-grained variables interacting with new, effective potentials the complexity of the molecular system is reduced, thus yielding accelerated simulation methods capable of reaching mesoscopic length scales. Such methods have been developed for the study and simulation of crystal growth, surface processes and polymers, e.g., [10,19,21,23,27], while there is an extensive literature on soft matter and complex fluids, e.g., [13,14,30,41]. Existing approaches can give unprecedented speed-up to molecular simulations and can work well in certain parameter regimes, for instance, at high temperatures or low densities of the systems. On the other hand important macroscopic properties may not be captured properly in many parameter regimes, e.g., the melt structures of polymers, [27], or the crystallization of complex fluids, [34]. Motivated in part by such observations we formulated and analyzed, from a numerical analysis and statistical mechanics perspective, coarse-grained variable selection and error quantification of coarse-grained approximations focusing on stochastic lattice systems with long-range interactions, [2,23,25]. We have shown that the ensuing schemes, known as coarse-grained Monte Carlo (CGMC) methods, perform remarkably well even though traditional Monte Carlo methods experience a serious slow-down. In this paper we focus on lattice systems with both short and long-range interactions. Short-range interactions introduce strong correlations between coarse-grained variables, and a radically different approach needs to be employed in order to carry out a systematic and accurate coarse-graining of such systems.

The coarse-graining of microscopic systems is essentially a problem in approximation theory and numerical analysis. However, the presence of stochastic fluctuations on one hand, and the extensive nature of the models (the presence of extensive quantities that scale as \( O(N) \) with the size of system \( N \)) on the other, create a new set of challenges. Before we proceed with the main results of this paper we discuss all these issues in a general setting that applies to both on-lattice and off-lattice systems and present the mathematical and numerical framework of coarse-graining for equilibrium many-body systems.

We denote by \( \sigma \) microscopic states of a many-particle system and by \( \mathcal{S}_N \) the set of all microscopic states (i.e., the configuration space). The energy of a configuration is given by the Hamiltonian \( H_N(\sigma) \) where \( N \) denotes the size of the microscopic system. An example studied in this paper is the \( d \)-dimensional Ising-type model defined on a lattice with \( N = n^d \) lattice points and suitable boundary conditions, e.g., periodic. For both on-lattice or off-lattice particle systems the finite-volume equilibrium states of the system are given by the canonical Gibbs measure at the inverse temperature \( \beta \), describing the most probable configurations

\[
\mu_{N,\beta}(d\sigma) = \frac{1}{Z_N} e^{-\beta H_N(\sigma)} P_N(d\sigma),
\]

where the normalizing factor \( Z_N = \int e^{-\beta H_N} P_N \), the partition function, ensures that (1.1) is a probability measure, and \( P_N(d\sigma) \) denotes the prior distribution on \( \mathcal{S}_N \). The prior distribution is typically a product measure (see for instance [23]) which describes non-interacting particles, or equivalently describes the system at infinite temperature \( \beta = 0 \). At the \( \beta = 0 \) limit the particle interactions included in \( H_N \) are unimportant and thermal fluctuations, i.e., disorder, associated with the product structure of the prior, dominate the system. By contrast at the zero temperature
limit, $\beta \to \infty$, interactions dominate and thermal fluctuations are unimportant; in this case (1.1) concentrates on the minimizers, also known as the “ground states”, of the Hamiltonian $H_N$ over all configurations $\sigma$. Finite temperatures, $0 < \beta < \infty$, describe intermediate states to these two extreme regimes, including possibly phase transitions, i.e., regimes when as parameters, such as the temperature, change, the system exhibits an abrupt transition from a disordered to an ordered state and vice versa, or between different ordered phases.

The objective of (equilibrium) computational statistical mechanics is the simulation of averages over Gibbs states, (1.1) of observable quantities $f(\sigma)$:

$$E_{\mu_{N,\beta}}[f] = \int f(\sigma)\mu_{N,\beta}(d\sigma).$$

Due to the exceedingly high dimension of the integration, even for moderate values of the system size $N$, e.g., $|S_N| = 2^N$ for the standard Ising model, such averaged observables are typically calculated by Markov Chain Monte Carlo (MCMC) methods. Nonetheless, mesoscale morphologies, e.g., traveling/standing waves and patterns, are beyond the reach of conventional Monte Carlo methods. For this reason coarse-graining methods have been developed in order to speed up molecular simulations.

We briefly discuss the mathematical formulation and numerical analysis challenges arising in coarse-graining of an equilibrium system described by (1.1). We denote the configuration space at the coarse level by $\bar{S}_M$ and we denote by $F$ the coarse-graining map $F : S_N \to \bar{S}_M$, $F\sigma = \eta \in \bar{S}_M$. The coarse-grained system size is denoted by $M$, while the microscopic system size is $N = QM$, where we refer to $Q$ as the level of coarse-graining, and $Q = 1$ corresponds to no coarse-graining. At the coarse-grained level one is interested in observables $f(\eta)$ which depend only on the coarse variable $\eta$, and a coarse-grained statistical description of the equilibrium properties of the system should be given by a probability measure $\bar{\mu}_{M,\beta}(d\eta)$ on $\bar{S}_M$ such that the average (the expected value) of such an observable is the same in the coarse-grained as well as fully resolved systems. This motivates the following definition.

**Definition 1.1.** The exact coarse-grained Gibbs measure $\bar{\mu}_{M,\beta}$ is defined by

$$\bar{\mu}_{M,\beta}(A) = \mu_{N,\beta}(F^{-1}(A)),$$

for any (measurable) set $A \subset \bar{S}_M$ or, equivalently,

$$\int f(\eta) \bar{\mu}_{M,\beta}(d\eta) = \int f(F(\sigma)) \mu_{N,\beta}(d\sigma),$$

for all (bounded) $f : \bar{S}_M \to \mathbb{R}$.

Slightly abusing notation we will write $\bar{\mu}_{M,\beta} \equiv \mu_{N,\beta} \circ F^{-1}$ in the sequel. In order to write the measure $\bar{\mu}_{M,\beta}$ in a more convenient form we first compute the exact coarse-graining of the prior distribution $P_N(d\sigma)$ on $S_N$:

$$\bar{P}_M(d\eta) = P_N \circ F^{-1}.$$

The conditional prior probability $P_N(d\sigma \mid \eta)$ of having a microscopic configuration $\sigma$ given a coarse configuration $\eta$ will play a crucial role in the sequel. Recall that for a function $g(\sigma)$ the conditional expectation is given by

$$E[g \mid \eta] = \int g(\sigma) P_N(d\sigma \mid \eta).$$
We now write the coarse-grained Gibbs measure $\bar{\mu}_{M,\beta}$ using a coarse-grained Hamiltonian $\bar{H}_M(\eta)$.

**Definition 1.2.** The exact coarse-grained Hamiltonian $\bar{H}_M(\eta)$ is given by

$$e^{-\beta \bar{H}_M(\eta)} = \mathbb{E}[e^{-\beta H_N} | \eta].$$

This procedure is known as a renormalization group map, [17,20]. Note that the partition functions for $H_N$ and $\bar{H}_M$ coincide since

$$Z_N = \int 1 e^{-\beta H_N} P_N(d\sigma) = \int \int e^{-\beta H_N} P_N(d\sigma | \eta) \bar{P}_M(d\eta) = \int e^{-\beta \bar{H}_M} \bar{P}_M(d\eta) \equiv \bar{Z}_M.$$ 

Hence for any function $f(\eta)$ we have

$$\int f(\eta) \mu_{N,\beta}(d\sigma) = \int f(\eta) \frac{1}{Z_N} e^{-\beta H_N} P_N(d\sigma) = \int \int f(\eta) \frac{1}{Z_N} e^{-\beta H_N} P_N(d\sigma | \eta) \bar{P}_M(d\eta) = \int f(\eta) \frac{1}{Z_M} e^{-\beta \bar{H}_M(\eta)} \bar{P}_M(d\eta),$$

and thus the exactly coarse-grained measure $\bar{\mu}_{M,\beta}(d\eta)$ in (1.3) is given by

$$\bar{\mu}_{M,\beta}(d\eta) = \frac{1}{\bar{Z}_M} e^{-\beta \bar{H}_M(\eta)} \bar{P}_M(d\eta).$$

Although typically $\bar{P}_M(d\eta)$ is easy to calculate, see e.g., (2.4), the exact computation of the coarse-grained Hamiltonian $\bar{H}_M(\eta)$ given by (1.7) is, in general, an impossible task even for moderately small values of $N$.

In this paper we restrict our attention to lattice systems, and our main result is the development of a general strategy to construct explicit numerical approximations of the exact coarse-grained Hamiltonian $\bar{H}_M(\eta)$ in the physically important case of combined and competing short and long-range interactions. Essentially we construct an approximate coarse-grained energy landscape for the original complex microscopic lattice system in Section 2. We show that there is an expansion of $\bar{H}_M(\eta)$ into a convergent series

$$\bar{H}_M(\eta) = \bar{H}_M^{(0)}(\eta) + \bar{H}_M^{(1)}(\eta) + \bar{H}_M^{(2)}(\eta) + \text{error}$$

by constructing a suitable first approximation $\bar{H}_M^{(0)}(\eta)$ and identifying small parameters to control the higher-order terms in the expansion. Truncations including the first few terms in (1.8) correspond to coarse-graining schemes of increasing accuracy. In order to obtain this expansion we rewrite (1.6) as

$$\bar{H}_M(\eta) = \bar{H}_M^{(0)}(\eta) - \frac{1}{\beta} \log \mathbb{E}[e^{-\beta (H_N - B_M^{(0)}(\eta))} | \eta].$$

We need to show that the logarithm can be expanded into a convergent series, uniformly in $N$, eventually yielding an expression of the type (1.8). However, two interrelated difficulties emerge immediately: (a) the stochasticity of the system in the finite temperature case yields the non-linear expression in (1.9) which in turn will need to be expanded into a series; (b) the extensive nature of the microscopic system, i.e., typically the Hamiltonian scales as $H_N = \mathcal{O}(N)$, does not allow the expansion of the logarithm and exponential functions into the Taylor series.
For these reasons, one of the principal mathematical tools we employ is the cluster expansion method; see [38] for an overview and references. As we shall see in the course of this paper cluster expansions will allow us to identify uncorrelated components in the expected value $\mathbb{E}[e^{-\beta(H_N - \bar{H}_M^{(0)}(\eta))} | \eta]$, which in turn will permit us to factorize it, and subsequently expand the logarithm in (1.9) in order to obtain the series (1.8). The coarse-graining of systems with purely long-range interactions was extensively studied using cluster expansions in [2][23][24]. Here we are broadly following and extending this approach. However, the presence of both short and long-range interactions presents new difficulties and requires new methods based on the ideas developed in [4][33]. Short-range interactions induce sub-grid scale correlations between coarse variables, and need to be explicitly included in the initial approximation $\bar{H}_M^{(0)}(\eta)$. To account for these effects we introduce a multi-scale decomposition of the Gibbs state (1.1) into fine and coarse variables, which in turn allows us to describe, in an explicit manner, the communication between scales of the Gibbs state (1.1) into fine and coarse variables, which in turn will permit us to reverse the procedure of coarse-graining in a mathematically systematic manner, i.e., reconstruct spatially localized “atomistic” properties, directly from coarse-grained simulations. We note that this issue arises extensively in the polymer science literature, [31][40].

The paper is organized as follows. In Section 2 we present the microscopic Ising-type models with short and long-range interactions and introduce the coarse-graining maps and the resulting coarse-grained configuration spaces. In Section 3 we discuss our general strategy for the analysis of systems with short and long-range interactions and present our main results. In Section 4 we discuss semi-analytical coarse-graining schemes and their applications to specific examples. Section 5 is devoted to the construction of the cluster expansion and to the proof of convergence of our schemes. In Section 6 we discuss the computational complexity of the proposed coarse-graining schemes.

2. MICROSCOPIC LATTICE MODELS AND COARSE-GRAINING

We consider an Ising-type model on the $d$-dimensional square lattice $\Lambda_N := \{x = (x_1, \ldots, x_d) \in \mathbb{Z}^d; 0 \leq x_i \leq n - 1\}$ with $N = n^d$ lattice points. For simplicity we assume periodic boundary conditions throughout this paper, although other boundary conditions can be accommodated. At each lattice site $x$ there is a spin $\sigma(x)$ taking values in $\Sigma = \{+1, -1\}$. A spin configuration $\sigma = \{\sigma(x)\}_{x \in \Lambda_N}$ on the lattice $\Lambda_N$ is an element of the configuration space $\mathcal{S}_N := \Sigma^{\Lambda_N}$. For any subset $X \subset \Lambda_N$ we denote $\sigma_X = \{\sigma(x)\}_{x \in X} \in \Sigma^X$ the restriction of the spin configuration to $X$. Similarly, for a function $f : \mathcal{S}_N \to \mathbb{R}$ we denote $f_X$ the restriction of $f$ to $\Sigma^X$. The energy of a configuration $\sigma$ is given by the Hamiltonian

$$\tag{2.1} H_N(\sigma) = H_N^s(\sigma) + H_N^l(\sigma),$$

which consists of a short-range part $H_N^s$ and a long-range part $H_N^l$. For the short-range part we have

$$H_N^s(\sigma) = \sum_{X \subset \Lambda_N} U_X(\sigma),$$

where the short-range potential $U = \{U_X, X \subset \mathbb{Z}^d\}$, with $U_X : \Sigma^X \to \mathbb{R}$, is translation invariant (i.e., $U_{X+y} = U_X$ for all $X \subset \mathbb{Z}^d$ and all $y \in \mathbb{Z}^d$) and has the finite range $S$ (i.e., $U_X = 0$ whenever $\text{diam}(X) > S$). We define the norm
\[ \|U\| = \sum_{X \supset \{0\} \mid \text{diam}(X) \leq S} \|UX\|_\infty \] where the norm \( \| \cdot \|_\infty \) is the standard sup-norm on the space of continuous functions. A typical case is the nearest-neighbor Ising model

\[ H_N^s(\sigma) = K \sum_{\langle x, y \rangle} \sigma(x)\sigma(y), \]

where by \( \langle x, y \rangle \) we denote summation over the nearest neighbors. For the long-range part we assume the form

\[ H_N^l(\sigma) = -\frac{1}{2} \sum_{x \in \Lambda_N} \sum_{y \neq x} J(x - y)\sigma(x)\sigma(y), \]

where the two-body potential \( J \) has the form

\[ J(x - y) = \frac{1}{L^d} V \left( \frac{1}{L}\|x - y\| \right), \]

for some \( V \in C^1([0, \infty)) \). The factor \( 1/L^d \) in (2.2) is a normalization which ensures that the strength of the potential \( J \) is essentially independent of \( L \), i.e., \( \sum_{x \neq 0} |J(x)| \approx \int |V(r)|dr \). For example, if we choose \( V \) such that \( V(r) = 0 \) for \( r > 1 \), then a spin at the site \( x \) interacts with its neighbors which are at most \( L \) lattice points away from \( x \), and in this case \( L \) is the range of the interaction \( J \). It is convenient to think of \( L \) as a parameter in our model, and more precise assumptions on the interactions will be specified later on.

The finite-volume equilibrium states of the system are given by the canonical Gibbs measure (1.1) and \( P_N(d\sigma) \), the prior distribution on \( \mathcal{S}_N \), is a product measure

\[ P_N(d\sigma) = \prod_{x \in \Lambda_N} P_x(d\sigma(x)). \]

A typical choice is \( P_x(\sigma(x) = +1) = \frac{1}{2} \) and \( P_x(\sigma(x) = -1) = \frac{1}{2} \), i.e., independent Bernoulli random variables at each site \( x \in \Lambda_N \). For the sake of simplicity we consider Ising-type spin systems, but the techniques and ideas in this paper apply also to Potts and Heisenberg models or, more generally, to models where the “spin” variable takes values in a compact space.

2.1. Coarse-graining. In order to coarse-grain our system we divide the lattice \( \Lambda_N \) into coarse cells and define coarse variables by averaging spin values over the coarse cells. We partition the lattice \( \Lambda_N \) into \( M = m^d \) disjoint cubic coarse cells, each containing \( Q = q^d \) microscopic lattice points so that \( N = n^d = (mq)^d = MQ \). We define a coarse lattice \( \Lambda_M = \{k = (k_1, \ldots, k_d) \in \mathbb{Z}^d; 0 \leq k_i \leq m-1\} \) and we set \( \Lambda_N = \bigcup_{k \in \Lambda_M} C_k \) where \( C_k = \{x \in \Lambda_N; k_i q \leq x_i < (k_i + 1)q\} \). Whenever convenient we will identify the coarse cell \( C_k \) in the microscopic lattice \( \Lambda_N \) with the point \( k \) of the coarse lattice \( \Lambda_M \). For any configuration \( \sigma^k \equiv \sigma_{C_k} \) on the coarse cell \( C_k \) we assign a new spin value

\[ \eta(k) = \sum_{x \in C_k} \sigma(x) \]

which takes values in \( \Sigma = \{-Q, -Q + 2, \ldots, Q\} \). We denote the configuration space at the coarse level by \( \tilde{\mathcal{S}}_M = \Sigma^{\Lambda_M} \) and we denote by \( F \) the coarse-graining map

\[ F : \mathcal{S}_N \to \tilde{\mathcal{S}}_M, \quad \sigma = \{\sigma(x)\}_{x \in \Lambda_N} \mapsto \eta = \{\eta(k)\}_{k \in \Lambda_M} \]

which assigns a configuration \( \eta \) on the coarse lattice \( \Lambda_M \) given a configuration \( \sigma \) on the microscopic lattice \( \Lambda_N \).
The exact coarse-grained Gibbs measure is defined in (1.3) for arbitrary Gibbs states having the form (1.7). Since $\eta(k)$ depends only on the spins $\sigma(x)$, with $x \in C_k$, the coarse-grained measure $\bar{P}_M$ is a product measure

$$\bar{P}_M(d\eta) = P_N \circ F^{-1} = \prod_{k \in \bar{\Lambda}_M} P_k(d\eta(k)).$$

For example if $P_x$ is a Bernoulli distribution, then $P_k(\eta(k)) = \left(\frac{1}{2}^{Q(\eta(k)+Q)}\right)^{1/2}$. Similarly, we define the conditional probability measure $P_N(d\sigma|\eta)$ as having a microscopic configuration $\sigma$ on $\Lambda_N$ given a coarse configuration $\eta$ on $\bar{\Lambda}_M$. This measure plays a crucial role in the sequel since it factorizes over the coarse cells

$$P_N(d\sigma|\eta) = \prod_{k \in \bar{\Lambda}_M} P_k(d\sigma^k|\eta(k)),$$

where $P_k(d\sigma^k|\eta(k))$ is the conditional probability of a microscopic configuration $\sigma^k$ on $C_K$ given a coarse configuration $\eta(k)$.

### 3. Approximation strategies for $\bar{H}_M(\eta)$

In this section we present a general strategy for constructing approximations of the exact coarse-grained Hamiltonian $\bar{H}_M(\eta)$ in (1.7). We show how to expand $\bar{H}_M(\eta)$ into a convergent series (1.8) by choosing a suitable first approximation $\bar{H}_M^{(0)}(\eta)$ and identifying small parameters to control the higher-order terms in the expansions. The basic idea is to use the first approximation $\bar{H}_M^{(0)}(\eta)$ in order to rewrite (1.6) as (1.9). We show that the logarithm can be expanded into a convergent series, uniformly in $N$, using suitable cluster expansion techniques. We discuss in detail the case $d = 1$ in order to illustrate general ideas in the case where calculations and formulas are relatively simple. The general $d$-dimensional case is discussed in detail in Section 5.

We recall that the Hamiltonian $H_N(\sigma) = H_N^l(\sigma) + H_N^s(\sigma)$ consists of a short-range part $H_N^s(\sigma)$ with the range $S$ and a long-range part $H_N^l(\sigma)$ whose range is $L$. We choose the coarse-graining level $q$ such that

$$S < q < L.$$

There are two small parameters associated with the range of the interactions,

$$\epsilon_s \propto O\left(\frac{S}{q}\right) \quad \text{and} \quad \epsilon_l \propto O\left(\frac{q}{L}\right).$$

The first approximation is of the form

$$\bar{H}_M^{(0)} = \bar{H}_M^l(0) + \bar{H}_M^s(0),$$

and two distinct separate procedures are used to define the short-range coarse-grained approximation $\bar{H}_M^s(0)$, as well as its long-range counterpart $\bar{H}_M^l(0)$. Due to the non-linear nature of the map induced by (1.9) it is not obvious that (3.1) will be a valid approximation, except possibly at high temperatures, when $\beta \ll 1$. This fact will be established for a wide range of parameters in the error analysis of Theorem 3.3 and in the discussion in Section 4 provided a suitable choice is made for $\bar{H}_M^{(0)}$ and $\bar{H}_M^l(0)$. 

License or copyright restrictions may apply to redistribution; see https://www.ams.org/journal-terms-of-use
3.1. Coarse-graining of the long-range interactions. We briefly recall the coarse-graining strategy of [24] for the long-range interactions. Since the range of the interaction, $L$, is larger than the range of coarse-graining $Q$, a natural first approximation for the long-range part is to average the interaction $J(x-y)$ over coarse cells. Thus we define

$$H_M^{L(0)}(\eta) = E[H_N | \eta],$$

and an easy computation gives

$$H_M^{L(0)}(\eta) = -\frac{1}{2} \sum_{k \in \Lambda_M} \sum_{l \neq k} J(k, l) \eta(k) \eta(l) - \frac{1}{2} \sum_{k \in \Lambda_M} J(k, k) (\eta(k)^2 - Q),$$

where

$$J(k, l) = \frac{1}{Q^2} \sum_{x \in C_k, y \in C_l} J(x-y), \quad J(k, k) = \frac{1}{Q(Q-1)} \sum_{x, y \in C_k, y \neq x} J(x-y).$$

A simple error estimate (see [2, 24] for details in various cases) gives

$$H_N(\sigma) = H_M^{L(0)}(\mathbf{F}(\sigma)) + e_L \quad \text{with} \quad e_L = N \mathcal{O}\left(\frac{q}{L} \|\nabla V\|_{\infty}\right).$$

Using this definition of $H_M^{L(0)}$ we obtain

$$e^{-\beta H_N(\sigma)} P_N(d\sigma | \eta) = e^{-\beta H_M^{L(0)}(\eta)} e^{-\beta \left[H_N^{L(0)}(\sigma) - H_M^{L(0)}(\eta)\right]} P_N(d\sigma | \eta)$$

$$= e^{-\beta H_M^{L(0)}(\eta)} \prod_{j, k \in \Lambda_M} \left(1 + f_{jk}^l\right) P_N(d\sigma | \eta),$$

where

$$f_{jk}^l = e^{\frac{q}{2} \sum_{x \in C_j, y \in C_l, y \neq x} \left(J(x-y) - J(k,l)\right)^2 \sigma(x) \sigma(y) (\delta_{jk} - \delta_{y,k}) - 1}.$$  

Due to the fact that $P_N(d\sigma | \eta)$ has a product structure one can rewrite (3.5) as a cluster expansion, [24] (see also Section 5), as in (1.8). The key element in that cluster expansion is the “smallness” of the quantity

$$|J(x-y) - J(k,l)| \leq 2 \frac{q}{L^{d+1}} \sup_{x' \in C_k, y' \in C_l} |\nabla V(x' - y')|,$$

which yields asymptotics

$$f_{jk}^l = \mathcal{O}\left(q^{2d} \frac{q}{L^{d+1}} \|\nabla V\|_{\infty}\right).$$

The estimate (3.7) follows from regularity assumptions on $V$ and the Taylor expansion.

3.2. Multi-scale decomposition of Gibbs states. This approach provides the common underlying structure of all coarse-graining schemes at equilibrium including lattice and off-lattice models. It is essentially a decomposition of the Gibbs state (1.1) into product measures among different scales selected with suitable properties. We outline it for the case of short-range interactions where we rewrite the Gibbs measure (1.1) as

$$\mu_{N, \beta}(d\sigma) \sim e^{-\beta H_N(\sigma)} P_N(d\sigma) = e^{-\beta H_N(\sigma)} P_N(d\sigma | \eta) \tilde{P}_M(d\eta).$$
We use the notation \( \sim \) meaning up to a normalization constant, i.e., in the equation above we do not spell out the presence of the constant \( Z_N \). We now seek the following decomposition of the short-range interactions:

\[
e^{-\beta H^{l}_N(\sigma)} P_N(d\sigma \mid \eta) = R(\eta) A(\sigma) \nu(d\sigma \mid \eta),
\]

where

(a) \( R(\eta) \) depends only on the coarse variable \( \eta \) and is related to the first coarse-grained approximation \( \bar{H}^{(s,0)}_M(\eta) \) via the formula

\[
R(\eta) = e^{-\beta \bar{H}^{(s,0)}_M(\eta)}, \quad A(\sigma) \nu(d\sigma \mid \eta) = e^{-\beta \left( H^{l}_N(\sigma) - \bar{H}^{(s,0)}_M(\eta) \right)} P_N(d\sigma \mid \eta).
\]

(b) \( A(\sigma) \) has a form amenable to a cluster expansion, i.e.,

\[
A(\sigma) = \prod_{k \in \Lambda_M} (1 + \Phi_k(\sigma)).
\]

The function \( \Phi_k \) is small and moreover \( \Phi_k(\sigma) \) depends on the configuration \( \sigma \) only locally, up to a fixed finite distance from \( C_k \). In the example at hand (for \( d = 1 \)) we have \( \Phi_k(\sigma) = \Phi_k(\sigma^k, \sigma^{k+1}) \).

(c) The measure \( \nu(d\sigma \mid \eta) \) has the general form

\[
\nu(d\sigma \mid \eta) = \prod_{k \in \Lambda_M} \nu_k(d\sigma \mid \eta),
\]

where \( \nu_k(d\sigma \mid \eta) \) depends on \( \sigma \) and \( \eta \) only locally up to a fixed finite distance from \( C_k \). In the example at hand \( \nu_k(d\sigma \mid \eta) \) depends only on the configuration on \( C_{k-1} \cup C_k \cup C_{k+1} \). Even though the measure \( \nu(d\sigma \mid \eta) \) is not a product measure, the fact that this measure has finite spatial correlation makes it adequate for a cluster expansion; see (3.24) and Section 5.

Although here we described the multi-scale decomposition of the Gibbs measure for the case of short-range interactions, the results on the long-range interactions, discussed earlier, can be reformulated in a similar way. In particular, (3.4) and (3.5) can be rewritten as

\[
e^{-\beta H^{l}_N(\sigma)} P_N(d\sigma \mid \eta) = R(\eta) A(\sigma) \nu(d\sigma \mid \eta),
\]

where

\[
R(\eta) = e^{-\beta \bar{H}^{(l,0)}_M(\eta)}, \quad \nu(d\sigma \mid \eta) = P_N(d\sigma \mid \eta), \quad \text{and}
\]

\[
A(\sigma) = e^{-\beta \left( H^{l}_N(\sigma) - \bar{H}^{(l,0)}_M(\eta) \right)} = \prod_{j,k \in \Lambda_M} (1 + f_{jk}^l).
\]

We recall that in analogy to (3.12), the product structure of \( \nu(d\sigma \mid \eta) = P_N(d\sigma \mid \eta) \) allows us to carry out a cluster expansion for the long-range case, and obtain a convergent series such as (1.8), thus yielding an expansion of the exact coarse-grained Hamiltonian \( \bar{H}^{l}_M \). We also note that (3.9), used here as a numerical and multi-scale analysis tool in order to derive suitable approximation schemes for the coarse-grained Hamiltonian, was first introduced in [1][2][3] for the purpose of deriving cluster expansions for lattice systems with short-range interactions away from the well-understood high temperature regime.

The error estimates for the approximating coarse-grained measure \( \bar{\mu}_{\beta,M} \), which is defined by the expansion of the coarse-grained Hamiltonian truncated at the \( k+1 \)st term, is quantified in terms of the specific relative entropy \( \frac{1}{N} R(\bar{\mu}_{\beta,M}^{(0)} \mid \mu_{N,\beta} \circ F^{-1}) \). The measure \( \mu_{N,\beta} \circ F^{-1} \) denotes the exactly coarse-grained measure \( \mu_{N,\beta} \) induced
by the coarse-graining map $F$. The relative entropy of the measure $\mu$ with respect to the measure $\nu$ is

$$R(\mu \mid \nu) \equiv \int \log \left( \frac{d\mu}{d\nu} \right) d\mu.$$ 

3.3. Coarse-graining schemes in one spatial dimension. We sketch how to obtain a decomposition such as \ref{eq:cluster expansion} for $d=1$ and construct a suitable $R(\eta)$. We first focus on the short-range interactions with radius $S$. Using that $S < q$, we write the corresponding Hamiltonian as

$$H^S_N(\sigma) = \sum_{k \in \Lambda_M} H^S_k(\sigma) + \sum_{k \in \Lambda_M} W_{k,k+1}(\sigma),$$ \hspace{1cm} (3.15)

where

$$H^S_k(\sigma) = \sum_{X \subset C_k} U_X(\sigma), \quad W_{k,k+1}(\sigma) = \sum_{X \cap C_k \neq \emptyset, X \cap C_{k+1} \neq \emptyset} U_X(\sigma),$$

i.e., $H^S_k$ is the energy for the cell $C_k$ which does not interact with other cells, i.e., under the free boundary conditions, and $W_{k,k+1}$ is the interaction energy between the cells $C_k$ and $C_{k+1}$. Thus, we can split the one-dimensional lattice into non-communicating components, for instance, even- and odd-indexed cells, and write

$$e^{-\beta H^S_N} P_N(d\sigma \mid \eta) = \prod_{k: \text{odd}} \left[ e^{-\beta(W_{k-1,k}+W_{k,k+1})} e^{-\beta H^S_k} P_k(d\sigma^k \mid \eta(k)) \right]$$

$$\times \prod_{k: \text{even}} e^{-\beta H^S_k} P_k(d\sigma^k \mid \eta(k)).$$ \hspace{1cm} (3.16)

In \ref{eq:partition function}, we will normalize the factors for $k$ odd by dividing each factor with the suitably defined corresponding partition functions for the regions $C_k$ and $C_{k-1} \cup C_k \cup C_{k+1}$.

**Definition 3.1.** We define the partition function with boundary conditions $\sigma^{k-1}$ and $\sigma^{k+1}$, i.e.,

$$Z_k(\eta(k); \sigma^{k-1}, \sigma^{k+1}) = \int e^{-\beta(W_{k-1,k}+W_{k,k+1})} e^{-\beta H^S_k} P_k(d\sigma^k \mid \eta(k)).$$ \hspace{1cm} (3.17)

In order to decouple even and odd cells we define the partition function with free boundary conditions on $C_{k-1}$ and boundary condition $\sigma^{k+1}$ on $C_{k+1}$, i.e.,

$$Z_k(\eta(k); 0, \sigma^{k+1}) = \int e^{-\beta W_{k,k+1}} e^{-\beta H^S_k} P_k(d\sigma^k \mid \eta(k)), \hspace{1cm} (3.18)$$

and similarly $Z_k(\eta(k); \sigma^{k-1}, 0)$, as the partition function with free boundary conditions on $C_{k+1}$ and boundary condition $\sigma^{k-1}$ on $C_{k-1}$. We also denote by $Z_k(\eta(k); 0, 0)$ the partition function for $C_k$ with free boundary conditions. We define the three-cell partition function with free boundary conditions

$$Z_{k-1,k,k+1}(\eta(k-1), \eta(k), \eta(k+1); 0, 0)$$

$$= \int e^{-\beta(H^S_{k-1}+W_{k,k-1}+H^S_k+W_{k,k+1}+H^S_{k+1})}$$

$$\times P_{k-1}(d\sigma^{k-1} | \eta(k-1)) P_k(d\sigma^k | \eta(k)) P_{k+1}(d\sigma^{k+1} | \eta(k+1)). \hspace{1cm} (3.19)$$

The key to the decomposition and eventually to the cluster expansion is the introduction of a “small term” analogous to \ref{eq:cluster expansion}. 

Definition 3.2.
\begin{equation}
(3.20) \quad f^{k-1}_{k+1}(\eta(k);\sigma^{k-1},\sigma^{k+1}) = \frac{Z_k(\eta(k);\sigma^{k-1},\sigma^{k+1})Z_k(\eta(k);0,0)}{Z_k(\eta(k);0,\sigma^{k+1})Z_k(\eta(k);\sigma^{k-1},0)} - 1.
\end{equation}

An important element in the cluster expansion in Section \[5\] is the estimation of the terms \( f^{k-1}_{k+1} \). However, a straightforward estimate yields
\begin{equation}
(3.21) \quad f^{k-1}_{k+1}(\eta(k);\sigma^{k-1},\sigma^{k+1}) \simeq \beta |U|,
\end{equation}
while for a sharper estimate relying on the spatial correlation length of the short-range interactions, we refer to (4.9) and (4.10) below.

We can now rewrite
\begin{equation}
Z_k(\eta(k);\sigma^{k-1},\sigma^{k+1}) = (f^{k-1}_{k+1}(\eta(k);\sigma^{k-1},\sigma^{k+1}) + 1) \times \frac{Z_k(\eta(k);0,\sigma^{k+1})Z_k(\eta(k);\sigma^{k-1},0)}{Z_k(\eta(k);0,0)}.
\end{equation}

In (3.16) we now divide and multiply each factor with \( k \) odd by \( Z_k(\sigma^{k-1},\sigma^{k+1}) \) and use the formula (3.22). Furthermore, we multiply and divide each factor with even \( k \) by \( Z_{k-1,k,k+1}(0,0) \) and obtain
\begin{equation}
e^{-\beta H_N} P_{\sigma}(d\sigma | \eta)
= \prod_{k: \text{odd}} Z_k(0,0)^{-1} \prod_{k: \text{even}} Z_{k-1,k,k+1}(0,0) \prod_{k: \text{odd}} (f^{*}_{k-1,k+1} + 1) \equiv R(\eta)
\end{equation}
\begin{equation}
\times \prod_{k: \text{odd}} e^{-\beta (H_{k-1,k+1}^{*} + W_{k-1,k+1})} Z_k(\sigma^{k-1},\sigma^{k+1}) P_{k}(d\sigma^{k} | \eta(k)) \prod_{k: \text{even}} e^{-\beta H_{k}^{*} Z_{k+1}(\sigma^{k},0) Z_{k-1}(0,\sigma^{k})} P_{k}(\sigma^{k} | \eta(k)) \equiv \nu(d\sigma | \eta)
\end{equation}
where we have used that
\begin{equation}
\prod_{k: \text{odd}} Z_k(0,\sigma^{k+1})Z_k(\sigma^{k-1},0) = \prod_{k: \text{even}} Z_{k+1}(\sigma^{k},0) Z_{k-1}(0,\sigma^{k}).
\end{equation}

It is easy to verify that \( \nu(d\sigma | \eta) \) defined in (3.24) is a normalized measure and has the form required in condition (c) of the multi-scale decomposition of the Gibbs measure. The factor \( R(\eta) \) defined in (3.23) gives the first-order corrections induced by the correlations between adjacent cells. Putting together the analysis for short and long-range interactions we obtain the main result formulated as a theorem.

Theorem 3.3. Let
\begin{equation}
\tilde{H}^{(0)}_M(\eta) = \tilde{H}_M^{(0)}(\eta) + \tilde{H}^{s,(0)}_M(\eta),
\end{equation}
where \( \tilde{H}_M^{(0)}(\eta) \) is given in (3.2) and (3.3) and
\begin{equation}
\tilde{H}^{s,(0)}_M(\eta) = - \sum_{k: \text{odd}} U^{s,(0)}_k(\eta(k)) + \sum_{k: \text{even}} U^{s,(0)}_{k-1,k,k+1}(\eta(k-1),\eta(k),\eta(k+1)),
\end{equation}
with the one-body interactions
\begin{equation}
U^{s,(0)}_k(\eta(k)) = -\frac{1}{\beta} \log Z_k(\eta(k);0,0),
\end{equation}
and the three-body interactions
\[ U^{s,(0)}_{k-1,k,k+1}(\eta(k-1), \eta(k), \eta(k+1)) \]
\[ = -\frac{1}{\beta} \log Z^{(0)}_{k-1,k,k+1}(\eta(k-1), \eta(k), \eta(k+1); 0, 0), \]
where \( Z_k \) and \( Z^{(0)}_{k-1,k,k+1} \) are given by (3.18) and (3.14) respectively. Then:

1. We have the error bound
\[ |\bar{H}_M - \bar{H}^{(0)}_M| = NO\left(\frac{\beta S\|U\|}{q} + \frac{q\beta\|\nabla V\|_{\infty}}{L}\right), \]
for a short-range potential with the range \( S \ll q \ll L \). The loss of information when coarse-graining at the level \( q \) is quantified by the specific relative entropy error
\[ \frac{1}{N} R(\bar{\mu}_{M,\beta}^{(0)}|\mu_{N,\beta} \circ F^{-1}) = O\left(\frac{\beta S\|U\|}{q} + \frac{q\beta\|\nabla V\|_{\infty}}{L}\right). \]

2. There exist \( \delta_1 > 0 \) and \( \delta_2 > 0 \) such that if
\[ \sup_{k} \sum_{\sigma} \sup_{i \neq k} |f_{jk}(\bar{\sigma}, \sigma_k)| \leq \delta_1, \] and \( \sup_{k} \sup_{\sigma^{k-1}, \sigma^{k+1}} |f^{s,(0)}_{k-1,k+1}(\eta(k); \sigma^{k-1}, \sigma^{k+1})| \leq \delta_2, \]
where \( f_{k-1,k+1} \) and \( f_{jk} \) are given by (3.20) and (3.6) respectively, then \( \bar{H}_M - \bar{H}^{(0)}_M \) is expanded in an absolutely convergent series in the parameters \( \beta\|U\|S \) and \( \beta \frac{q^2}{L}\|\nabla V\|_{\infty} \):
\[ \bar{H}_M(\eta) = \bar{H}^{(0)}_M(\eta) + \bar{H}^{(1)}_M(\eta) + \ldots + \bar{H}^{(p)}_M(\eta) + M O((\max(\beta\|U\|S, \beta \frac{q^2}{L}\|\nabla V\|_{\infty}))^{p+1}), \]
where the exact form of the higher-order terms \( \bar{H}^{(p)}_M(\eta) \) can be directly computed by writing explicitly the terms in the power series (3.24).

**Remark 3.1.** Note that once the absolute convergence of (3.31) is established, one can further estimate the terms of the expansion, in which case it is expected to obtain an absolutely convergent series in the parameters \( \beta\|U\|S \) and \( \beta \frac{q^2}{L}\|\nabla V\|_{\infty} \). This has been shown in [24] for the long-range case only, but a similar result is expected to hold in the present case. In such a case the corresponding error in (3.31) would read \( NO((\max(\beta\|U\|S, \beta \frac{q^2}{L}\|\nabla V\|_{\infty}))^{p+1}) \).

To motivate the reader we next give an outline of the proof containing the basic ideas, and for the full proof we refer to the general d-dimensional case in Theorem 3.7.

**Outline of the proof.** We define \( \bar{H}^{(0)}_M \), from (3.22) and \( \bar{H}^{s,(0)}_M \) as in (3.28) and (3.24). Then with \( \bar{H}_M^{(0)} := \bar{H}^{(0)}_M + \bar{H}^{s,(0)}_M \), from (1.6) we obtain
\[ \bar{H}_M(\eta) = \bar{H}^{(0)}_M(\eta) - \frac{1}{\beta} \log \int A(\sigma)\nu(d\sigma|\eta), \]
where \( \nu \) is determined in (3.24) and \( A \) is given by
\[ A(\sigma) \equiv \prod_{i<j} (1 + f_{ij}) \prod_{i: \text{odd}} (1 + f_{i-1,i+1}) = \sum_{\mathcal{L} \in \mathcal{A}_G} \prod_{i \in \mathcal{L}} f_{i-1,i+1} \sum_{G \in \mathcal{G}_M} \prod_{\{i,j\} \in E(G)} f_{ij}. \]
where $G_M$ is the set of all graphs on $M$ vertices, where $M$ is the total number of coarse cells.

Substituting $A(\sigma)$ from (3.33) in the second term of the right hand side of (3.32) we need to integrate with the non-product measure $\nu(d\sigma|\eta)$. Hence, every term $f_{i-1,i+1}^s$ for $i$ odd has to be integrated with the measures on the coarse cells $\bigcup_{j=i-2}^{i+2} C_j$. Such a set we call a link $l_i$. Similarly, every $f_{ij}^l$ gives rise to a similar link, where if $i$ is even then it needs to be integrated over $\bigcup_{j=i-2}^{i+2} C_j$ as before, while if it is odd the integration is over $\bigcup_{j=i-1}^{i+1} C_j$. Given $R_0 \subset \tilde{\Lambda}_M$ we define the link $\ell \equiv \bigcup_{i \in R_0} l_i$. We define a polymer $R$ to be a collection of these two types of links, namely $R := (l_1, \ldots, l_p; \ell)$ (from which we can read the short $f_{i-1,i+1}^s$ and the long $f_{ij}^l$ range contributions). Two polymers $R_1$ and $R_2$ are called compatible if $\text{supp}(R_1) \cap \text{supp}(R_2) = \emptyset$. With these definitions we have:

\begin{equation}
\int A(\sigma)\nu(d\sigma|\eta) = \sum_{\{R_1, \ldots, R_n\}_{\text{comp}}} \prod_{i=1}^n w(R_i)
\end{equation}

where

\begin{equation}
w(R) := \int \prod_{i \in I} f_{i-1,i+1}^s \sum_{G \in C_{R_0}} \prod_{(i,j) \in E(G)} f_{ij}^l \nu(d\sigma|\eta),
\end{equation}

where $C_{R_0}$ is the set all connected graphs on the vertices of $R_0 \subset \tilde{\Lambda}_M$. The form on the right hand side of (3.34) is amenable to the cluster expansion theorem which we will use in order to write the logarithm as an absolutely convergent series; see Theorem 5.3. Thus, while the error estimate in (3.29) comes directly from the uniform estimates (3.8) and (3.21), the absolute convergence of the series (3.31) is the result of Theorem 5.4 with the small parameters to be determined by the convergence condition, given in Lemma 5.6 and being directly related to the uniform estimates (3.8) and (3.21). The complete proof in all space dimensions is carried out in Section 5.

Remark 3.2. The error estimate (3.29) suggests qualitatively an estimate on the regimes of validity of the method, and on the “optimal” level, $q = q_{\text{opt}}$, when we restrict ourselves to the regime $S < q < L$, where $S$ and $L$ are the respective interaction ranges for short and long-range potentials. The corresponding error is then

\begin{equation}
q_{\text{opt}} \simeq \sqrt{SL \frac{||U||}{||\nabla V||_\infty}}, \quad \frac{1}{N} R(\mu_{M,\beta}^{(0)} | \mu_{N,\beta} \circ F^{-1}) = O \left( \beta \sqrt{\frac{S}{L} ||U|| ||\nabla V||_\infty} \right).
\end{equation}

Remark 3.3. The application of Theorem 3.8 requires that we check the validity of (3.30), which is certainly guaranteed by the conditions (3.8) and (3.21) in suitable regimes. More interestingly, for specific examples (3.30) can be verified directly, through a sharper estimate than (3.21), thus improving (3.29). This estimate relies on the spatial correlation length of the short-range interactions; we refer to (4.9) and (4.10) below. Furthermore, in (4.9) we even obtain an upper bound that depends only on the coarse observables. This allows us to check the conditions (3.30) (dictated by the cluster expansions) computationally in the process of a Monte Carlo simulation involving only the coarse variables $\eta$. We note here that in [32, 33], the short-range condition in (3.30) is taken as an assumption. Overall, in
one dimension, this condition holds up to very low temperatures while in dimension \( d \geq 2 \) this condition can be satisfied in the high-temperature regime; see for example the analysis in [4] where similar conditions are used for the nearest-neighbor Ising model in the dimension \( d = 2 \) all the way down to the critical temperature.

### 3.4. A posteriori error estimates.

In [24] we introduced the use of cluster expansions as a tool for constructing a posteriori error estimates for coarse-graining problems, based on the rather simple observation that higher-order terms in [3.31] can be viewed as errors that depend only on the coarse variables \( \eta \). Following the same approach an a posteriori estimate immediately follows from (3.31).

**Corollary 3.1.** We have

\[
R(\bar{\mu}_{M,\beta}^{(0)} | \mu_{N,\beta} \circ \mathbf{F}^{-1}) = \beta \mathbb{E}_{\bar{\mu}_{M,\beta}^{(0)}}[S(\eta)] + \log \left( \mathbb{E}_{\bar{\mu}_{M,\beta}^{(0)}}[e^{-\beta S(\eta)}] \right) + O(\delta^2),
\]

where the residuum operator is \( S(\eta) = \bar{H}_M^{(1)}(\eta) \).

In [22] we already employed this type of estimate for stochastic lattice systems with long-range interactions, in order to construct adaptive coarse-graining schemes. These tools operated as an “on-the-fly” coarsening/refinement method that recovers accurately phase diagrams. The estimates allowed us to change adaptively the coarse-graining level within the coarse-graining hierarchy once sufficiently large or small errors were detected, thus speeding up the calculations of phase diagrams. Earlier work that uses only an upper bound and not the asymptotically sharp cluster expansion-based estimate can be found in [7,8].

### 3.5. Microscopic reconstruction.

The reverse procedure of coarse-graining, i.e., reproducing “atomistic” properties, directly from coarse-grained simulation methods is an issue that arises extensively in the polymer science literature, [31,40]. The principal idea is that computationally inexpensive coarse-graining algorithms will reproduce large scale structures and subsequently microscopic information will be added through *microscopic reconstruction*, for example the calculation of diffusion of penetrants through polymer melts, reconstructed from coarse-grained simulation, [31].

In this direction, the CGMC methodology discussed in this section can provide a framework to mathematically formulate microscopic reconstruction and study related numerical and computational issues. Indeed, the conditional measure \( A(\sigma) \nu(d\sigma | \eta) \) in the multi-scale decompositions (3.9) and (3.13) can also be viewed as a microscopic reconstruction of the Gibbs state (1.1) once the coarse variables \( \eta \) are specified. The product structure in (3.11) and (3.12) allows for easy generation of the fine scale details by first reconstructing over a family of domains given only the coarse-grained data and gradually moving to the next family of domains given now both the coarse-grained data and the previously reconstructed microscopic values.

In view of this abstract procedure based on multi-scale decompositions such as (3.9), we readily see that the particular product structure of the explicit formulas (3.23) and (3.24) for the case of the dimension \( d = 1 \) yields a hierarchy of reconstruction schemes. A first-order approximation can be based on the approximation \( A(\sigma) \approx 1 \) (cf. (3.21), (3.23)):

(a) first, \( R(\eta) \) defined in (3.23) provides the coarse-graining scheme, which will produce coarse variable data \( \eta(k) \) for all \( k \);
(b) next, we reconstruct the microscopic configuration $\sigma^{\text{even}}$ consisting of the $\sigma^k$’s in all boxes (coarse cells) with $k$ even using the measure $\nu_k(d\sigma|\eta) := e^{-\beta H^s_k} Z_k(\sigma_k(0) = 0) P_k(d\sigma^k | \eta(k))$, conditioned on the coarse configuration $\eta(k)$ from (a) above;

(c) finally, we reconstruct the microscopic configuration in the remaining boxes with $k$ odd using $\nu_k(d\sigma|\eta) := e^{-\beta \left( u^k + w_{k-1, k} + w_{k+1, k} \right)} Z_k(\sigma^k_{k-1, k+1}) P_k(d\sigma^k | \eta(k))$, given the coarse variable $\eta(k)$ from step (a), and the microscopic configurations $\sigma^{\text{even}}$ from step (b).

We note that this procedure is local in the sense that the reconstruction can be carried out in only the “subdomain of interest” of the entire microscopic lattice $\Lambda_N$; this is clearly computationally advantageous because microscopic kMC solvers are used only in the specific part of the computational domain, while inexpensive CGMC solvers are used in the entire coarse lattice $\bar{\Lambda}_M$. Further discussion on the numerical analysis issues related to microscopic reconstruction for lattice systems with long-range interactions can be found in [23,25,26,39].

4. Semi-analytical coarse-graining schemes and examples

Next we discuss the numerical implementation of the effective coarse-grained Hamiltonians derived in Theorem 3.3. We begin with a general implementation scheme and we subsequently investigate further reductions of the computational complexity for particular examples in one and two space dimensions.

4.1. Semi-analytical splitting schemes and inverse Monte Carlo methods.

One of the main points of our method is encapsulated in (3.25): the computationally expensive long-range part for conventional Monte Carlo methods can be computed by calculating the explicit formula given in (3.3) in the spirit of our previous work [24]. Then we can turn our attention to the short-range interactions where Monte Carlo methods, at least for reasonably sized domains, are inexpensive. More specifically, for the evaluation of the short-range contribution in (3.25) we introduce the normalized measure

$$\hat{P}_k(d\sigma^k | \eta(k)) = \frac{1}{Z_k(\eta(k); 0, 0)} e^{-\beta H^s_k} P_k(d\sigma^k | \eta(k)),$$

where the sum is computed with free boundary conditions on $C_k$ and $Z_k(\eta(k); 0, 0)$ is accordingly defined as in (3.18). Thus (3.25) can be rewritten as

$$\hat{H}^{s, (0)}_M = \sum_{k \in \bar{\Lambda}} \hat{U}^{s, (0)}(\eta(k)) + \sum_{k: \text{even}} \hat{V}^{s, (0)}_{k-1, k+1}(\eta(k-1), \eta(k), \eta(k+1)),$$

where, based on (3.26) and (4.1), we defined the three-body coarse interaction potential

$$\hat{V}^{s, (0)}_{k-1, k, k+1}(\eta(k-1), \eta(k), \eta(k+1)) = \frac{1}{\beta} \log \int e^{-\beta (W_{k-1, k}(\sigma) + W_{k+1, k}(\sigma))} \times \hat{P}_{k-1}(d\sigma^{k-1} | \eta(k-1)) \hat{P}_k(d\sigma^k | \eta(k)) \hat{P}_{k+1}(d\sigma^{k+1} | \eta(k+1)).$$

The main difficulty in the calculation of (4.3) is that for the three-body integral one needs to perform the integration for all possible combinations of the multi-canonical constraint. On the other hand all simulations involve only short-range interactions and need to be carried out only on three coarse cells, rather than
the entire lattice. Practically, the calculation of (4.3) can be implemented using the so-called inverse Monte Carlo method, [27]. We sample the measure $\hat{P}_k$ using Metropolis spin flips and subsequently we create a histogram for all possible values of $\eta(k) = \sum_{x \in C_k} \sigma(x)$. Then we compute the above integral by using the samples which correspond to the prescribed values $\eta(k-1), \eta(k)$ and $\eta(k+1)$.

A complementary approach in order to further increase the computational efficiency of the schemes presented in Theorem 3.3 is to rearrange the splitting based on the size of the error in (3.29). Indeed, these estimates suggest a natural way to decompose the overall interaction potential into: (a) a short-range piece $J_s$ including possible singularities originally in $J$, e.g., the non-smooth part in the Lennard-Jones potential, and (b) a locally integrable (or smooth) long-range decaying component, $J_l$. Thus, if $K(x,y)$ is the short-range potential in (2.1) we can rewrite the overall potential as

\begin{equation}
K(x,y) + J(x,y) = J_s(x,y) + J_l(x,y).
\end{equation}

In this way the accuracy can be enhanced by implementing the analytical coarse-graining (3.3) for the smooth long-range piece $J_l(x,y)$, and the semi-analytical scheme (3.26) for the “effective” short-range piece $J_s(x,y)$.

4.2. Multi-body interactions. Existing coarse-graining methods, e.g., [16], employ an inverse Monte Carlo computation involving both short and long-range interactions, and due to computational limitations have to disregard multi-body terms such as the ones derived in this paper, e.g. Theorem 3.3. The splitting approach developed here allows us to calculate analytically the approximate effective Hamiltonian for the costly long-range interactions, (3.3) in (3.25) or (4.4), and in parallel carry out the inverse Monte Carlo step for (1.2). The necessity to include multi-body terms in the effective Hamiltonian was first discussed in [2] together with their role in the proper coarse-graining of singular short-range interactions. For nearest-neighbor lattice models, coarse-graining schemes involving only the single-body interactions such as (3.27) were proposed in [9], while two-body interactions were considered in [11]. As we will see next, such two-body coarse-grained correlations are contained in the three-body terms in (3.28). Furthermore, we can precisely quantify the regimes where multi-body terms (3.28) are necessary in the context of a specific example.

4.3. A typical example: improved schemes and a posteriori estimation. We examine the derived coarse-graining schemes in the context of a specific, but typical example. We consider the Hamiltonian

\begin{equation}
H_N(\sigma) = H_s^N(\sigma) + H_l^N(\sigma) := K \sum_{\langle x, y \rangle} \sigma(x)\sigma(y) - \frac{1}{2} \sum_{\langle x, y \rangle} J(x - y)\sigma(x)\sigma(y),
\end{equation}

where by $\langle x, y \rangle$ we denote summation over the nearest neighbors, i.e., $|x - y| = 1$, and by $\langle x, y \rangle$ the long-range summation as in (2.22). Although we follow the splitting strategy discussed in the previous paragraph we present a simplified numerical algorithm by carrying out further analytical calculations. Not surprisingly, such calculations allow not only for easier sampling in the semi-analytical calculations of the inverse Monte Carlo, but give additional insight on the nature of multi-body, coarse-grained interactions.
For the short-range contributions, given a coarse cell $C_k$ with $q$ lattice points, we denote by $x_1, \ldots, x_q$ the lattice sites in $C_k$. With this notation, following (4.3) the short-range three-body interaction is given by

$$
\tilde{V}_{k-1,k,k+1}^{s,(0)}(\eta(k-1), \eta(k), \eta(k+1)) = \frac{1}{\beta} \log \int e^{-\beta K(\sigma^{k-1}(x_q)\sigma^k(x_1) + \sigma^k(x_2)\sigma^{k+1}(x_1))}
$$

(4.6) \quad \times \hat{P}_{k-1}(d\sigma^{k-1} | \eta(k-1))\hat{P}_k(d\sigma^k | \eta(k))\hat{P}_{k+1}(d\sigma^{k+1} | \eta(k+1)) .

The main difficulty in computing the second term is the conditioning on the coarse-grained values $\eta(k-1), \eta(k), \eta(k+1)$ over three coarse cells. At first glance this requires that we run multi-constrained Monte Carlo dynamics for every given value of the $\eta$’s, i.e., for $q^3$ variables. However, as we show in the sequel, when dealing with a particular example, e.g., the nearest neighbor interactions, the computationally expensive three-body term reduces to a product of one-body terms. We first rewrite

$$
e^{-\beta K(\sigma^{k-1}(x_q)\sigma^k(x_1))} = a - b\sigma^{k-1}(x_q)\sigma^k(x_1),
$$

where we set

$$a = \cosh(\beta K), \quad b = \sinh(\beta K), \quad \lambda = \tanh(\beta K).$$

Moreover, we introduce the one- and two-point correlation functions

$$\Phi^x_k(\eta_k) := \int \sigma(x)\hat{P}_k(d\sigma^k | \eta(k)) \quad \text{and} \quad \Phi^x:y_k(\eta_k) := \int \sigma(x)\sigma(y)\hat{P}_k(d\sigma^k | \eta(k)).$$

By symmetry we have that $\Phi^{x_1}_k = \Phi^{x_2}_k$ and similarly, consider $\Phi^{x_1,x_q}_k$ for $x = x_1$ and $y = x_q$. Furthermore, these functions depend on $k$ only via the coarse variable $\eta_k$, hence we now define

$$\Phi^1(\eta_k) := \int \sigma(x_1)\hat{P}_k(d\sigma^k | \eta(k)) \quad \text{and} \quad \Phi^2(\eta_k) := \int \sigma(x_1)\sigma(x_q)\hat{P}_k(d\sigma^k | \eta(k)).$$

It is a straightforward computation to show that

$$
\tilde{V}_{k-1,k,k+1}^{s,(0)}(\eta(k-1), \eta(k), \eta(k+1)) = -\frac{2}{\beta} \log a
$$

$$
-\frac{1}{\beta} \log \left( 1 - \lambda \Phi^1(\eta(k-1))\Phi^1(\eta(k)) - \lambda \Phi^1(\eta(k))\Phi^1(\eta(k+1))
$$

(4.8) \quad + \lambda^2 \Phi^1(\eta(k-1))\Phi^2(\eta(k))\Phi^1(\eta(k+1)) \right).

Although these are three-body interactions, the additional analytical calculations reduce their computation to the nearest-neighbor Monte Carlo sub-grid sampling of (4.7). Moreover, from (3.20) we have

$$f^{s}_{k-1,k+1}(\eta(k); \sigma^{k-1}, \sigma^{k+1}) = \frac{\lambda^2 \sigma^{k-1}(x_q)\sigma^{k+1}(x_1)[\Phi^2(\eta(k)) - (\Phi^1(\eta(k)))^2]}{(1 - \lambda \sigma^{k-1}(x_q)\Phi^1(\eta(k)))(1 - \lambda \sigma^{k+1}(x_1)\Phi^1(\eta(k)))},$$

thus the following estimate holds for some $C > 0$:

$$
\sup_{\sigma^{k-1}, \sigma^{k+1}} |f^{s}_{k-1,k+1}| \leq C\lambda^2 |\Phi^2(\eta(k)) - (\Phi^1(\eta(k)))^2| \equiv \Theta(\eta_k; \lambda),
$$

(4.9)

where the right hand side $\Theta$ is an a posteriori functional in the sense that it can be computed from the coarse-grained data.
4.3.1. Analytical formulas for CG interactions and complexity reduction. We can further reduce the complexity of the Monte Carlo sub-grid sampling of the quantities $\Phi_1(\eta_k)$ and $\Phi_2(\eta_k)$ in (4.8) and the a posteriori error indicator (4.9), through their calculation or approximation by analytical formulas, when they are available. This approach will be especially relevant in higher dimensions as we also discuss in Section 6. We can derive such analytical approximations in two different ways.

First, exact analytical formulas for the correlation function $\Theta(\eta_k; \lambda)$, as well as $\Phi_1(\eta_k)$ and $\Phi_2(\eta_k)$, can be easily obtained with direct calculation replicating the unconstrained case, e.g. [17] Chapter 3.5; or alternatively via the analytically available correlation function of the nearest neighbor Ising model in the presence of an external field, and the equivalence of ensembles manifested in the coverage/external field phase diagram, [3,17,38]. Both approaches yield

\[
\Theta(\eta_k; \lambda) = |\Phi_2(\eta(k)) - [\Phi_1(\eta(k))]^2| \simeq \exp \left( - \frac{|x_1 - x_q|}{\xi_{\text{n.n.}}} \right) = \exp \left( - \frac{q}{\xi_{\text{n.n.}}} \right),
\]

where $\xi_{\text{n.n.}}$ is the correlation length on the nearest-neighbor Ising model for an external field corresponding to coverage $\eta(k)$ through the corresponding phase diagram. In general, relations such as (4.10) hold away from phase transition regimes, and can also be used in coarse-graining schemes for higher-dimensional lattice systems; see in addition Section 6.

On the other hand, a high temperature expansion of the quantities $\Phi_1(\eta_k)$ and $\Phi_2(\eta_k)$ yields much simpler approximations, which however are more restricted in their applicability:

\[
\Phi_1(\eta) = E[\sigma(x) | \eta] + O(\lambda) = \frac{\eta q}{q} + O(\lambda),
\]

\[
\Phi_2(\eta) = E[\sigma(x)\sigma(y) | \eta] + O(\lambda) = \frac{\eta^2 - q}{q(q-1)} + O(\lambda).
\]

Then,

\[
\Theta(\eta; \lambda) \simeq \lambda^2 |\Phi_2 - (\Phi_1)^2| = \lambda^2 \frac{q^2 - \eta^2}{q^2(q-1)} + O(\lambda^3).
\]

Thus the condition (3.30) of Theorem 3.3 and the derived coarse-grained approximations can be conditionally checked during simulation by

\[
\sup_{\sigma_{k-1},\sigma_{k+1}} |f_{k-1,k+1}^s| \leq C \frac{\lambda^2}{q-1} \left(1 - \frac{\eta^2}{q^2} \right) + O(\lambda^3).
\]

We note that (4.14) suggests a quantitative understanding of the dependence of the coarse-graining error for the nearest-neighbor Ising model. The error increases, (a) when the parameter $\lambda^2$ increases, i.e., at lower temperatures/stronger short-range interactions, (b) when the level of coarse-graining $q$ decreases, and (c) at regimes where the local coverage $\eta$ is not uniformly homogeneous, i.e., away from the regime $\eta \approx \pm q$. Such a situation occurs, for example, around an interface in the phase transition regime. This is the case even in one dimension if long-range interactions are present in the system. Similarly, (4.9) and (4.10) along with (8.8) provide an even more detailed understanding than (8.36) of the parameters affecting a suitable coarse-graining level $q$, for systems with both short- and long-range interactions.
5. Coarse-graining schemes in higher dimensions and proofs

In this section we prove Theorem 3.3 and its generalization in higher dimensions. We formulate the proofs in full generality, assuming a $d$-dimensional lattice. As in the one-dimensional case considered in Section 3.3, our approach here will also consist of two main steps: (a) a multi-scale decomposition of the Gibbs states in all dimensions, and (b) a rigorous cluster expansion to obtain a convergent series such as (3.31).

Here coordinates of lattice points are understood as multi-indices in $\mathbb{Z}^d$. We start by constructing the a priori coarse-grained measure induced by the short-range interaction. We perform a block decimation procedure following the strategy in [33] and partition $\Lambda_M$ into $2^d$-many sublattices of spacing 2. Let $e_\alpha, \alpha = 2, 3, \ldots, 2^d$ be

$$
\begin{array}{cccccccc}
4 & 3 & 4 & 3 & 4 & 3 \\
1 & 2 & 1 & 2 & 1 & 2 \\
3 & 4 & 3 & 4 & 3 \\
2 & 1 & 2 & 1 & 2 \\
1 & 2 & 1 & 2 & 1 & 2
\end{array}
$$

Figure 1. The sublattices $\Lambda_\alpha^\mu$ covering the coarse lattice $\Lambda_M$. The vectors $e_\alpha$ defining translations of the first sublattice $\Lambda_1^\mu$ are depicted for $d = 2, 3$. The cells on the two-dimensional lattice are numbered with values of $\alpha = 1, \ldots, 4$ according to which sublattice $\Lambda_\alpha^\mu$ they belong.

We write the coarse lattice as union of sublattices

$$
\Lambda_M = \bigcup_{\alpha=1}^{2^d} \Lambda_\alpha^\mu,
$$

where $\Lambda_1^\mu = 2\Lambda_M$, $\Lambda_2^\mu = \Lambda_1^\mu + e_2$ and $\Lambda_\alpha^{\alpha+1} = \Lambda_\alpha^\mu + e_{\alpha+1}$, for $\alpha = 1, \ldots, 2^d - 1$.

Given a coarse cell $C_k$ we define the set of neighboring cells by

$$
\partial C_k := \bigcup_{\{l : \|l - k\| = 1\}} C_l,
$$

where $\|l - k\| := \max_{i=1, \ldots, d} |l_i - k_i|$. We also let $D_k := C_k \cup \partial C_k$. Given a sublattice $\Lambda_\alpha^\mu$, we denote by $\sigma^\alpha$ the microscopic configuration in all the cells $C_k \in \Lambda_\alpha^\mu$ and by $\sigma^{\alpha >}$ the configuration in $\Lambda_\beta^\mu$ for all $\beta > \alpha$. We also define a function $p : \Lambda_M \to \{1, \ldots, 2^d\}$ such that for $k \in \Lambda_\alpha^\mu$, we have $p(k) = \alpha$ if $C_k \in \Lambda_\alpha^\mu$.

We split the short-range part of (2.1),

$$
H_N^s(\sigma) = \sum_{\alpha} \sum_{k \in \Lambda_\alpha^\mu} H_k^s(\sigma^\alpha) + \sum_{\alpha} \sum_{k \in \Lambda_\alpha^\mu} W_k(\sigma^{\alpha}; \sigma^{\alpha >}),
$$

License or copyright restrictions may apply to redistribution; see https://www.ams.org/journal-terms-of-use
where, for \( k \in \tilde{\Lambda}^1_M \), the terms \( H_k(\sigma^\alpha) \) are the self-energy on the boxes \( C_k \) given by

\[
H_k^*(\sigma^\alpha) = \sum_{X \subset C_k} U_X(\sigma^\alpha).
\]

Moreover, the energy due to the interaction of \( C_k \) with the neighboring cells is given by

\[
W_k(\sigma^\alpha; \sigma^{>\alpha}) = \sum_{X \subset D_k} U_X(\sigma^\alpha \vee \sigma^{>\alpha}),
\]

where \( \sigma^\alpha \vee \sigma^{>\alpha} \) is the concatenation on \( \tilde{\Lambda}^1_M \) and \( \tilde{\Lambda}^{>\alpha}_M \).

### 5.1. Multi-scale decomposition in two-dimensions.

To better illustrate the construction of the decomposition \((3.9)\) we first present the two-dimensional case. Next, we construct the reference conditional measure \( \nu(d\sigma|\eta) \) in \((3.9)\), under the constraint of a fixed averaged value \( \eta = \{\eta(k)\}_{k \in \tilde{\Lambda}_M} \) on the coarse cells.

**Step 1.** The starting point is a product measure on \( C_k \) for \( k \in \tilde{\Lambda}^1_M \). We let \( A_1(k) \equiv C_k \) and after appropriate normalization we obtain

\[
e^{-\beta H_k^*(\sigma)} \prod_{k \in \tilde{\Lambda}_M} P_k(d\sigma) = \prod_{\alpha \geq 2 \ k \in \tilde{\Lambda}^\alpha_M} \left( e^{-\beta H_k^*(\sigma^\alpha)} e^{-\beta W_k(\sigma^\alpha; \sigma^{>\alpha})} P_k(d\sigma^\alpha) \right) \times \prod_{k \in \tilde{\Lambda}^1_M} Z(A_1(k); \sigma^{>1}; \eta(k)) \nu^1_{>1}(d\sigma^1),
\]

where

\[
\nu_{>1}^1(d\sigma^1) := \prod_{k \in \tilde{\Lambda}^1_M} \left[ \frac{1}{Z(A_1(k); \sigma^{>1}; \eta)} e^{-\beta W_k(\sigma^1; \sigma^{>1})} e^{-\beta H_k(\sigma^1)} P_k(d\sigma^1) \right]
\]

is the new prior measure on \( \tilde{\Lambda}^1_M \) with boundary conditions \( \sigma^{>1} \) and the canonical constraint \( \eta(k) \), \( k \in \tilde{\Lambda}^1_M \). The partition function

\[
Z(A_1(k); \sigma^{>1}; \eta(k)) = \int e^{-\beta H_k^*(\sigma)} e^{-\beta W_k(\sigma^1; \sigma^{>1})} P_k(d\sigma^1)
\]

depending on the boundary conditions \( \sigma^{>1} \) on the set \( \partial A_1(k) \) couples the configurations in \( C_l \) with \( l \in \partial A_1(k) \). In particular, it couples the configurations \( \sigma^2 \) and gives rise to a new interaction between them for which it will be shown that it is small due to Condition 5.1.

**Step 2.** Moving along the vector \( e_2 \) we seek the measure \( \nu_{>2}^2 \) on \( \{+1, -1\} \bigcup_{k \in \tilde{\Lambda}^2_M} C_k \). Given the partition function \( Z(A_1(k); \sigma^{>1}; \eta(k)) \) we denote by \( S_{k,e_2}^+ Z \) the partition function on the same domain \( A_1(k) \) as \( Z \), but with new boundary conditions which are the same as \( Z \) in the \( +e_2 \) direction, free in the \( -e_2 \) direction and unchanged in all the other directions. Similarly, we denote by \( S_{k,e_2}^- Z \) the partition function with free boundary conditions in the direction \( +e_2 \) and by \( S_{k,e_2}^0 Z \) with free boundary conditions in both \( \pm e_2 \) directions. With these definitions we have the identity

\[
Z(A_1(k); \sigma^{>1}; \eta(k)) = \frac{(S_{k,e_2}^+ Z)(S_{k,e_2}^- Z)}{(S_{k,e_2}^0 Z)} (1 + \Phi_k^1),
\]
where we have introduced the function $\Phi^1_k$ which contains the interaction between the variables $\sigma^{>1}$, and it is given by

$$\Phi^1_k := \frac{Z(A_1(k); \sigma^{>1}; \eta(k))(S^0_{k,e_2}Z)}{(S^+_{k,e_2}Z)(S^-_{k,e_2}Z)} - 1.$$  

In this way we split the partition function $Z$ into a part where the interaction between the cells $C_{k-e_2}$ and $C_{k+e_2}$ is decoupled and an error part which is to be small.

The next step is to index the new partition functions $(S^+_{k,e_2}Z)$ and $(S^-_{k,e_2}Z)$ (which are functions of $\sigma^2$ indexed by $k \in \bar{\Lambda}_M^1$) with respect to $k \in \bar{\Lambda}_M^2$. We have

$$\prod_{k \in \bar{\Lambda}_M^2} (S^+_{k,e_2}Z)(S^-_{k,e_2}Z) = \prod_{k \in \bar{\Lambda}_M^2} (S^+_{k-e_2,e_2}Z)(S^-_{k+e_2,e_2}Z).$$

Then if we neglect for a moment the error term $(1 + \Phi^1_k)$, in order to define $\nu^2_{>2}$ we have to deal with the following terms:

$$\prod_{k \in \bar{\Lambda}_M^2} (S^0_{k,e_2}Z)^{-1} \prod_{k \in \bar{\Lambda}_M^2} \left[ e^{-\beta H_k^2(\sigma^2)} e^{-\beta W_k(\sigma^2; \sigma^{>2})} (S^+_{k-e_2,e_2}Z)(S^-_{k+e_2,e_2}Z) P_k(d\sigma^2) \right].$$

The terms in the second product contain all possible interactions in the set

$$A_2(k) = C_{k-e_2} \cup C_k \cup C_{k+e_2}$$

for $k \in \bar{\Lambda}_M^2$ with the corresponding partition function being given by

$$Z(A_2(k); \sigma^{>2}; \eta(k)) = \int e^{-\beta H_k^2(\sigma^2)} e^{-\beta W_k(\sigma^2; \sigma^{>2})} (S^+_{k-e_2,e_2}Z)(S^-_{k+e_2,e_2}Z) P_k(d\sigma^2).$$

By normalizing with this function we obtain the measure

$$\nu^2_{>2}(d\sigma^2) = \prod_{k \in \bar{\Lambda}_M^2} \left[ \frac{1}{Z(A_2(k); \sigma^{>2}; \eta(k))} \right] e^{-\beta H_k^2(\sigma^2)} e^{-\beta W_k(\sigma^2; \sigma^{>2})} (S^+_{k-e_2,e_2}Z)(S^-_{k+e_2,e_2}Z) P_k(d\sigma^2).$$

Note that the factor $(S^0_{k,e_2}Z)^{-1}$ depends on $\eta$ as well as on $\sigma^{>2}$ and hence we will need to further split it when we define a measure on the variables on which it depends. Summarizing the first two steps we have obtained that the left hand side of (5.2) is equal to

$$\left[ \prod_{k \in \bar{\Lambda}_M^1} Z(A_2(k); \sigma^{>2}; \eta(k)) \prod_{k \in \bar{\Lambda}_M^1} (S^0_{k,e_2}Z)^{-1} \prod_{k \in \bar{\Lambda}_M^1} (1 + \Phi^1_k) \right] \nu^2_{>2}(d\sigma^2) \nu^1_{>1}(d\sigma^1).$$

If we are interested in the case $d = 1$, this would be the final expression. However, for higher dimensions we need to repeat the above steps. We give one more step in order to obtain more intuition on the relevant terms and then we give the final expression in agreement with the result in [33]. The proof of the general formula is done with a recurrence argument on the number of steps, and for the details we refer to [33].
Step 3. To proceed in the next step along direction $e_3$ we split $Z(A_2(k); \sigma^2; \eta(k))$ (which couples the configurations in $C_k$ with $p(k) = 3$) in the same fashion as before. We have

$$Z(A_2(k); \sigma^2; \eta(k)) = \frac{(S_{k,e_3}^+ Z)(S_{k,e_3}^- Z)}{(S_{k,e_3}^0 Z)} (\Phi_k^3 + 1),$$

where

$$\Phi_k^3 := \frac{Z(A_2(k); \sigma^2; \eta(k))(S_{k,e_3}^0 Z)}{(S_{k,e_3}^+ Z)(S_{k,e_3}^- Z)} - 1.$$

We further change the indices in such a way that they are expressed with respect to $k \in \Lambda^3_M$ and then we glue the partition functions on $C_k$, $A_2(k-e_3)$ and $A_2(k+e_3)$. We define

$$A_3(k) := C_k \cup A_2(k-e_3) \cup A_2(k+e_3)$$

and

$$Z(A_3(k); \sigma^3; \eta(k)) := \int e^{-\beta \mathcal{H}_k^s} e^{-\beta m_k(\sigma^3; \eta)} (S_{k-e_3,e_3}^+ Z)(S_{k+e_3,e_3}^- Z) P_k(d\sigma^3).$$

The corresponding measure is

$$\nu_{2,3}^3(d\sigma^3) = \prod_{k \in \Lambda^3_M} \left[ \frac{1}{Z(A_3(k); \sigma^3; \eta(k))} \times e^{-\beta \mathcal{H}_k^s(\sigma^3)} e^{-\beta m_k(\sigma^3, \sigma^3)} (S_{k-e_3,e_3}^+ Z)(S_{k+e_3,e_3}^- Z) P_k(d\sigma^3) \right],$$

and the left hand side of (5.2) is now equal to

$$\prod_{k \in \Lambda^3_M} \left[ e^{-\beta \mathcal{H}_k^s(\sigma^4)} e^{-\beta m_k(\sigma^4, \sigma^4)} P_k(d\sigma^4) \right] \prod_{k \in \Lambda^4_M} Z(A_3(k); \sigma^3; \eta(k)) \prod_{k \in \Lambda^4_M} (S_{k,e_3}^0 Z)^{-1} \times \prod_{k \in \Lambda^2_M} (S_{k,e_2}^0 Z)^{-1} \prod_{k \in \Lambda^1_M} (1 + \Phi_k^3) \prod_{k \in \Lambda_M^1} (1 + \Phi_k^1) \nu_{2,3}^3(d\sigma^3) \nu_{2,2}^2(d\sigma^2) \nu_{1,1}^1(d\sigma^1).$$

As in the previous steps we can perform the usual actions on the partition function $Z(A_3(k); \sigma^3; \eta(k))$ which will give rise to a new element $A_4(k)$ with $k \in \Lambda^3_M$ and new error terms $\Phi_k^4$ with $k \notin \Lambda^4_M$. Furthermore, we have a similar splitting for the factor $(S_{k,e_2}^0 Z)^{-1}$ which also depends on $\sigma^4$, since the zero boundary condition involves only the direction $e_2$. For dimensions higher than two, related calculations will involve all the terms of similar origin as we move to new sublattices $\Lambda^\alpha_M$, with $\alpha > 4$. Finally, the reference conditional measure $\nu(d\sigma|\eta)$ in (3.9), in the two-dimensional case is

$$\nu(d\sigma|\eta) := \nu_{2,3}^3 \nu_{2,2}^2 \nu_{1,1}^1.$$

Concluding, in the case of the two-dimensional nearest neighbor Hamiltonian, we obtain that the leading term in the approximation of the coarse-grained Hamiltonian $\bar{H}_M^s$ consists of terms that refer to four different types of multi-cell interactions,

$$\bar{H}_M^{s,(0)} = \sum_{k \in \Lambda^1_M} \log Z(A_4(k)) - \sum_{k \in \Lambda^2_M} \log Z(A_4(k))$$

$$+ \sum_{k \in \Lambda^3_M} \log Z(A_4(k)) - \sum_{k \in \Lambda^4_M} \log Z(A_4(k)),$$

in the coarse-grained Hamiltonian $H^s_M$.
where $A_4(k)$ is a collection of coarse cells centered in $k \in \Lambda^\alpha_M$ and it is different depending on the sublattice to which the reference cell $k$ belongs. For $\alpha = 1, 2, 3, 4$ we have

$$A_4(k) = \begin{cases} 
\bigcup_{i,j \in \{-1,0,+1\}} C_{k+ie_2+je_3}, & k \in \Lambda^4_M, \\
\bigcup_{j \in \{-1,0,+1\}} C_{k+je_3}, & k \in \Lambda^3_M, \\
C_k, & k \in \Lambda^2_M, \\
\bigcup_{i \in \{-1,0,+1\}} C_{k+ie_2}, & k \in \Lambda^1_M.
\end{cases}$$

Figure 2 depicts the index sets $A_4(k)$ for the reference cell $k$ belonging to $\Lambda^\alpha_M$ for $\alpha = 1, \ldots, 4$.

5.2. Multi-scale decomposition in higher dimensions. We focus on the relevant quantities which are the reference measure $\nu_{\geq \alpha}^\alpha(d\sigma^\alpha)$, the error term $\Phi^\alpha_k$, with $k \in \Lambda_M$, and the sets $A_{\alpha}(k)$ and $B_{\alpha}(k)$, with the latter being the relevant boundary of $A_{\alpha}$. The index $\alpha$ indicates the sublattice we are considering.

Definition 5.1. The sets $A_{\alpha}(k)$ and $B_{\alpha}(k)$ for $k \in \Lambda^\alpha_M$ are

$$A_{\alpha}(k) = \bigcup_{l: \|l-k\|=1, \rho(l) \leq \alpha} C_l, \quad B_{\alpha}(k) = \bigcup_{l: \|l-k\|=1, \rho(l) > \alpha} C_l.$$

Definition 5.2. Given $\alpha = 1, \ldots, 2^d$ we define the normalized Bernoulli measure on $\Lambda^\alpha_M$,

$$(5.9) \quad \nu_{\geq \alpha}^\alpha(d\sigma^\alpha) = \prod_{k \in \Lambda^\alpha_M} \nu_{\geq \alpha}^B_{\alpha}(k)(d\sigma^\alpha),$$

where

$$(5.10) \quad \nu_{\geq \alpha}^B_{\alpha}(k)(d\sigma^\alpha) = \frac{e^{-\beta H^\alpha_k(\sigma^\alpha)} e^{-\beta W^\alpha_k(\sigma^\alpha; \sigma^{>\alpha})}}{Z(A_{\alpha}(k); \sigma^{>\alpha}; \eta(k))} Z(A_{\alpha}(k)/\{k\}; \sigma^{>\alpha}; \eta(k)) \prod_{l \in B_{\alpha}(k)} P_l(d\sigma^\alpha).$$
As we have seen in Step 3 we have two kinds of error terms $\Phi^\alpha_k$, in particular, those with $k \in \Lambda^\alpha_M$ and others with $k \notin \Lambda^\alpha_M$. In order to describe the latter we need to introduce additional notation.

For $\alpha = 1, \ldots, 2^d$ we denote by $\Gamma_\alpha$ the family of parallel hyperplanes of dimension $d - 1$ orthogonal to $e_{\alpha+1}$ passing through all the points $k \in \Lambda^\alpha_M$. Note that for any $\alpha$, we have that $\Lambda^\alpha_M = \Gamma_\alpha \cup (\Gamma_\alpha + e_{\alpha+1})$. In the next definition we introduce a new parameter $\epsilon_\alpha(k) \in \{\pm 1\}$ depending on whether we should perform gluing or unfolding as discussed before. This is determined as follows: for fixed $\alpha \in \Lambda^\alpha_M$ let $d(\alpha, \beta)$ be the distance between the sublattices $\Lambda^\alpha_M$ and $\Lambda^\beta_M$ in the metric $\|\alpha - \beta\|_\infty = \sum_{i=1}^d |\alpha_i - \beta_i|$. Moreover, we can find orthogonal vectors $\{v_j\}_{j=1,\ldots,d(\alpha,\beta)}$ and a family of signs $\{\epsilon_j\}_{j=1,\ldots,d(\alpha,\beta)}$ such that

$$\Lambda^\alpha_M = \Lambda^\beta_M + \gamma(\alpha, \beta)$$

with $\gamma(\alpha, \beta) = \sum_{j=1}^{d(\alpha,\beta)} \epsilon_j v_j$.

Note also that $|\gamma(\alpha, \beta)| = d(\alpha, \beta)$. Then the exponents $\epsilon_\alpha(k)$ with $p(k) = \beta$ are given by

$$\epsilon_\alpha(k) := (-1)^{|\gamma(\alpha, \beta)|}.$$

Furthermore, we denote by $Y(k, \gamma(\alpha, \beta))$ the affine hyperplane of codimension $|\gamma(\alpha, \beta)|$ orthogonal to the connecting vectors $\{v_j\}_{j=1,\ldots,|\gamma(\alpha, \beta)|}$ and passing through the point $k$.

$$Y(k, \gamma(\alpha, \beta)) = \bigcap_{j=1}^{N} Y(k, v_j),$$

where $Y(k, v)$ is the hyperplane of dimension $d - 1$ passing through $k$ and being perpendicular to the vector $v$. From the set of coarse-lattice points belonging to $Y(k, v)$ we define the corresponding set by

$$\mathcal{Y}(k, \gamma(\alpha, \beta)) := \bigcup_{l \in Y(k, \gamma(\alpha, \beta))} C_l.$$

Then, letting $l \in \Lambda^\alpha_M$, for $l$ such that $C_l \subset \partial C_k$ and with $l \in \Lambda_M^\beta$, for some $\beta$, we define

\begin{align*}
A_\alpha(l) &= \begin{cases}
\emptyset & \text{if } p(l) > 2d(\alpha), \\
A_\alpha(k) \cap \mathcal{Y}(l, \gamma(\alpha, \beta)) & \text{otherwise},
\end{cases} \\
B_\alpha(l) &= B_\alpha(k) \cap \mathcal{Y}(l, \gamma(\alpha, \beta)).
\end{align*}

With the above definitions we can determine the error terms in the general expansion.

**Definition 5.3.** For any $k \in \Lambda^\alpha_M$ and for $k \in \Gamma_\alpha$ the error terms are given by

$$\Phi^\alpha_k = -1 + \frac{Z(A_\alpha(k); \sigma^{\alpha}; \eta(k))Z(A_{\alpha+1}(k); \sigma^{\alpha+1}; \eta(k))}{(S^+_{k,e_{\alpha+1}}Z)(S^-_{k,e_{\alpha+1}}Z)}.$$

Moreover, if $k \in \Gamma_\alpha + e_{\alpha+1}$ and $k \notin \Lambda^\alpha_M$ we have

$$\Phi^\alpha_k = -1 + \frac{Z(A_\alpha(k); \sigma^{\alpha}; \eta(k))Z(A_{\alpha+1}(k); \sigma^{\alpha+1}; \eta(k))}{(S^+_{k-e_{\alpha+1},e_{\alpha+1}}Z)(S^-_{k+e_{\alpha+1},e_{\alpha+1}}Z)} - \epsilon_\alpha(k).$$

Furthermore, if $k \in \Lambda^\alpha_{M+1}$ we replace $Z(A_{\alpha+1}(k); \sigma^{\alpha+1})$ by $Z(A_{\alpha+1}(k)/|k|; \sigma^{\alpha+1})$. 
From Proposition 2.5.1 in [33] we have that the general $d$-dimensional formulation of the a priori measure induced by the short-range interactions is

$$e^{-\beta H_N^s(\sigma)} \prod_{k \in \Lambda_M} P_k(d\sigma) = R^s(\eta) A(\sigma) \nu(d\sigma|\eta) ,$$

where we have the following factors:

(i): a product of partition functions (depending only on the coarse-grained variable $\eta$) over finite sets of coarse cells with supports $A_2^{d}(k)$, with $k \in \Lambda_M^\alpha$ and $\alpha = 1, \ldots, 2^d$

$$R^s(\eta) := \prod_{\alpha=1}^{2^d} \prod_{k \in \Lambda_M^\alpha} \left[Z(A_2^{d}(k); \eta(k))^{2^d} \right],$$

(ii): error terms in the form of a gas of polymers (with the only interaction to be a hard-core exclusion)

$$A(\sigma) := \prod_{\alpha=1}^{2^d} \prod_{j \leq 2^d(\alpha)} \prod_{k \in \Lambda_M^j} (1 + \Phi_k^\alpha),$$

(iii): a reference measure induced by only the short-range interactions once we neglect the reference system and the error terms

$$\nu(d\sigma|\eta) := \nu^{2^d} \ldots \nu_2^{\mu} \nu_1^{\mu} .$$

With this expansion for the short-range interactions, going back to the general strategy presented in Section 3, if we also consider the long-range contribution from (3.2), we obtain

$$e^{-\beta H_M(\eta)} = \int e^{-\beta H_N^s} \prod_{k} P_k(d\sigma) = e^{-\beta H^{l,(0)}(\eta)} R^s(\eta) \int e^{-\beta (H_N^s - H^{l,(0)})} A(\sigma) \nu(d\sigma|\eta) ,$$

which implies that

$$\bar{H}_M(\eta) = H^{l,(0)}(\eta) - \log R^s(\eta) - \frac{1}{\beta} \log \mathbb{E}_\nu[e^{-\beta (H_N^s - H^{l,(0)})} A(\sigma)|\eta] .$$

5.3. Cluster expansion and effective interactions. The goal of this section is to expand the term $\mathbb{E}_\nu[e^{-\beta (H_N^s(\sigma) - \bar{H}^{l,(0)}(\eta))} A(\sigma)|\eta]$ in (5.11) into a convergent series using a cluster expansion. By the construction given previously the terms in $A(\sigma)$ are already in the form of a polymer gas with hard-core interactions only. For the long-range part we first write the difference $H^l_N(\sigma) - \bar{H}^{l,(0)}(\eta)$ as

$$H^l_N(\sigma) - \bar{H}^{l,(0)}(\eta) = \sum_{k \leq l} \Delta_{kl} J(\sigma) ,$$

where

$$\Delta_{kl} J(\sigma) := -\frac{1}{2} \sum_{x \in C_k \atop y \in C_l, y \neq x} (J(x,y) - \bar{J}(k,l))\sigma(x)\sigma(y)(2 - \delta_{kl}) .$$
We also define \( f_{kl}(\sigma) := e^{-\beta \Delta_{kl} J(\sigma)} - 1 \) and we obtain
\begin{equation}
\mathbb{E}_\nu[e^{-\beta (H_N(\sigma) - H^{1,(0)}(\eta))} A(\sigma|\eta)] = \int \prod_{k \leq l} (1 + f_{kl}) \prod_{\alpha = 1}^{2^d} \prod_{j \leq 2^d(\alpha)} (1 + \Phi_k^\alpha) \nu(d\sigma|\eta).
\end{equation}

We define the polymer model which contains combined interactions originating from both the short and long-range potential. By expanding the products in (5.16) we obtain terms of the type
\begin{equation}
\prod_{j=1}^p \Phi_{k_j}^{\alpha_j} \prod_{i=1}^q f_{l_i,m_i} \quad \text{where } k_j, l_i, m_i \in \Lambda_M \text{ and } \alpha_j \in \{1, \ldots, 2^d\}
\end{equation}
for some \( p \) and \( q \). The factors \( \Phi_{k_j}^{\alpha_j} \) are functions of the variables which are on the boundary of the corresponding sets \( A_{\alpha_j}(k_j) \). This boundary is described by the set
\begin{equation}
C_0^\alpha(k) = \begin{cases} B_\alpha(k) & \text{if } k \in \Gamma_\alpha, \\ B_{\alpha+1}(k) & \text{if } k \in \Gamma_\alpha + e_{\alpha+1}. \end{cases}
\end{equation}

Furthermore, since the measure \( \nu(d\sigma|\eta) \) is not a product measure but instead a composition of measures each one parametrized by variables which are integrated by the next measure, we need to create a “safety” corridor around the sets \( C_0^\alpha \) depending on the level of \( \alpha \). This is given in the next definition. For a given integer \( \beta \) with \( 1 < \beta < 2^d - \alpha \) we define
\begin{equation}
C_\beta^\alpha(k) = \bigcup_{\epsilon_1, \ldots, \epsilon_\beta \in \{\pm 1\}^\beta \mid C_1 \subset \partial(C_{k+\epsilon_1 e_{\alpha+1} + \ldots + \epsilon_\beta e_{\alpha+\beta}). p(l) > \alpha + \beta} C_l.
\end{equation}

Then for given \( \alpha \in \{1, \ldots, 2^d\} \) we call a “bond” of type \( C^\alpha \) the set
\begin{equation}
C^\alpha(k) = \bigcup_{\beta=0}^{2^d-\alpha} C_\beta^\alpha(k).
\end{equation}

With this definition, any factor \( \Phi_{k_j}^{\alpha_j} \) has a region of dependence which is given by the bond \( C^\alpha(k) \). Similarly, for the factors \( f_{l_i,m_i} \) originating from the long-range interactions the initial domain of dependence is \( C_{l_i} \cup C_{m_i} \). However, due to the non-product structure of the measure we need to introduce a safety corridor in the same way. Given \( k \in \Lambda_M \) for \( \beta \) an integer with \( 1 < \beta < 2^d - p(k) \) we define
\begin{equation}
C_\beta(k) = \bigcup_{\epsilon_1, \ldots, \epsilon_\beta \in \{\pm 1\}^\beta \mid C_1 \subset \partial(C_{k+\epsilon_1 e_{\alpha+1} + \ldots + \epsilon_\beta e_{\alpha+\beta}). p(l) > p(k) + \beta} C_l.
\end{equation}

Then for a given \( f_{kl} \) we define
\begin{equation}
C(k,l) = \bigcup_{\beta=1}^{2^d-p(k)} C_\beta(k) \bigcup_{\beta=1}^{2^d-p(l)} C_\beta(l).
\end{equation}

With a slight abuse of notation we define for \( R_0 = \{k_1, \ldots, k_{|R_0|}\} \)
\begin{equation}
C(R_0) = \bigcup_{i=1}^{|R_0|} \bigcup_{\beta=1}^{2^d-p(k_i)} C_\beta(k_i).
\end{equation}
A bond \( l \) will be either a \( C^\alpha_k \) bond for some \( \alpha, k \), called a bond of type 1, or a \( C(R_0) \) bond, where \( R_0 \) is any subset of \( \hat{\Lambda}_M \), and we call it a bond of type 2 and we denote it by \( \ell \). We say that two bonds \( l_1 \) and \( l_2 \) are connected if \( l_1 \cap l_2 \neq \emptyset \). We call a polymer \( R \) a set of bonds \( (l_1, \ldots, l_p, \ell) \) where \( l_1, \ldots, l_p \) are bonds of type 1 and \( \ell \) is a bond of type 2, i.e., there is a unique \( R_0 \) such that \( \ell \equiv C(R_0) \). A polymer is called connected if for any two elements there exists a chain of connected bonds in \( R \) joining them. The support \( \text{supp}(R) \) of \( R \) is \( \text{supp}(R) = \bigcup_{p=1}^p l_i \cup \ell \) and the cardinality \( |R| := p + |\ell| \). Let \( \mathcal{R} \) be the set of all such polymers. Two polymers \( R_1, R_2 \) are said to be compatible if \( \text{supp}(R_1) \cap \text{supp}(R_2) = \emptyset \) and we write \( R_1 \sim R_2 \).

Given a polymer \( R = (l_1, \ldots, l_p; \ell) \) we define the activity of \( R \) to be the function \( w : \mathcal{R} \to \mathbb{C} \) given by

\[
(5.23) \quad w(R) = \int \nu(d\sigma|\eta) \left( \prod_{j=1}^p \Phi^\alpha_{k_j} \sum_{g \in \mathcal{C}_{R_0}} \prod_{\{k,l\} \in E(g)} f_{kl} \right),
\]

where \( \mathcal{C}_{R_0} \) is the collection of connected graphs on the vertices of \( R_0 \subset \hat{\Lambda}_M \) and \( E(g) \) is the set of edges of the graph \( g \). Note also that from each \( l_j \) we read \( \alpha_j, k_j \) and from \( \ell \) we read \( \alpha, k \). Let

\[
\mathcal{D}_\mathcal{R} = \bigcup_{n=0}^{|\mathcal{R}|} \{(R_1, \ldots, R_n) \subset \mathcal{R} : \forall i \neq j, R_i \sim R_j \};
\]

then the partition function \( Z \) can be written as

\[
Z = \sum_{G \in \mathcal{D}_\mathcal{R}} \prod_{R \in G} w(R),
\]

which is the abstract form of a polymer model. Thus we can apply the general theorem of the cluster expansion once we check the convergence condition. The condition is stated as a theorem in [5].

**Theorem 5.4** ([5]). Let \( a : \mathcal{R} \to \mathbb{R}_+ \). Consider the subset of \( \mathbb{C}^\mathcal{R} \),

\[
P^a_\mathcal{R} := \{ w(R), R \in \mathcal{R} : \forall R \in \mathcal{R} : |w(R)| e^{a(R)} < 1 \text{ and } \sum_{R' \sim R} (-\log(1 - |w(R')| e^{a(R')})) \leq a(R) \}.
\]

Then on \( P^a_\mathcal{R} \), \( \log Z \) is well defined and analytic and

\[
(5.24) \quad \log Z = \sum_{I \in \mathcal{I}(\mathcal{R})} c_I \prod_{R \in \text{supp}(I)} w(R)^{I_R},
\]

where \( I = (I_R)_{R \in \mathcal{R}} \), \( \mathcal{I}(\mathcal{R}) \) is the collection of all multi-indexes \( I \), i.e., integer valued functions on \( \mathcal{R} \), and

\[
c_I = \frac{1}{I_{R_1}! \ldots I_{R_{|\mathcal{R}|}}!} \left. \frac{\partial^{I_{R_1} + \ldots + I_{R_{|\mathcal{R}|}}}}{\partial I_{R_1} w(R_1) \ldots \partial I_{R_{|\mathcal{R}|}} w(R_{|\mathcal{R}|})} \log Z \right|_{\{w(R_i) = 0\}_i}.
\]

For the proof we refer to [5]. Thus we need to check the condition of convergence. The following estimate for the long-range potential was proved in [24].
Lemma 5.5. Assume that $J$ satisfies (2.2). Then there exists a constant $C_1 \simeq \frac{2d+1}{L} \|\nabla V\|_{\infty}$ such that
\begin{equation}
\sup_{k \in \Lambda_M \setminus \{\pm k\}} \sum_{l \neq k} |\Delta_{kl} J(\sigma)| \leq C_1 ,
\end{equation}
for every $\sigma$.

For the short-range interaction we follow the analysis of [33] and we consider the following condition.

Condition 5.1. Let $e$ be a vector in one of the directions of the lattice $\Lambda_M$ and $Z_U(\Lambda; \sigma_-, \sigma_+, \tau; \eta_V)$ be the partition function for the interaction $U$ in the space domain $\Lambda$. We consider boundary conditions $\sigma_{\pm}$ in the directions $\pm e$ and $\tau$ in all other directions. Moreover, we impose multi-canonical constraints $\eta(k)$ for $k \in V \subset \Lambda_M$ with $\Lambda = \bigcup_{k \in V} C_k$. For a given $q > r_0$, with $|C_k| = q^d$, the following inequality holds:
\begin{equation}
\sup_{\sigma_{\pm}, \tau} \sup_{\Lambda} \sup_{\eta_V} |Z_U(\Lambda; \sigma_-, \sigma_+, \tau; \eta_V) Z_U(\Lambda; 0, 0, \tau; \eta_V) - 1| \leq C_2 ,
\end{equation}
where given the numbers $r = 2^{2d}[3(2^{d+1} + 1)]^d$ and $\delta_2$ determined in Lemma 5.6 the upper bound $C_2$ satisfies
\[ r C_2 e < \delta_2 . \]

Notice that we work with the same condition as Condition $C_L$ defined in [33], where in our notation $L$ is $q$, yet similar analysis applies in order to prove convergence of the cluster expansion under the milder condition of Condition $C'_L$, again as in [33]. We skip the analysis of such issues since it goes beyond the goal of the present work, however we note that in the one-dimensional case, (4.9) and (4.10) allow us to estimate $C_2$. Furthermore, these conditions are related to the ones presented in [12] in order to ensure that a given system belongs to the class of completely analytical interactions. For further details we refer the reader to [32] and [4] and to the references therein.

We next prove the convergence condition.

Lemma 5.6. There are $\delta_1$ and $\delta_2$ small enough such that if $C_1 < \delta_1$ and $C_2 < \delta_2$, then the set $\mathcal{P}_R$ is non-empty.

Proof. We take $a(R) = c|R|$, where $c$ is a constant to be chosen later. Note that $-\log(1 - x) \leq 2x$, so it suffices to show that
\begin{equation}
\sum_{R' \sim R} 2 |w(R')| e^{a(R')} \leq a(R) .
\end{equation}
Suppose that the generic polymer $R'$ is given by $R' = (l_1, \ldots, l_p; \ell')$, for some $p \geq 0$, where $l_j \equiv C^{a_j}(k_j')$ for $j = 1, \ldots, p$ and $\ell' \equiv C(R'_0)$ with $|R'_0| = n$ for some $n \geq 0$. For $|w(R')|$ we have
\[ |w(R')| \leq \int \nu(da|\eta) \prod_{j=1}^{p} |\Phi_{k_j'}^{\alpha_j}| \cdot \sum_{g \in \mathcal{G}_R} \prod_{\{k,l\} \in E(g)} f_{kl} | . \]
By the graph-tree inequality we have that for all $\sigma$, $\eta$ and with $|R'_0| = n$
\[ \sum_{g \in \mathcal{G}_R} \prod_{\{k,l\} \in E(g)} f_{kl} \leq e^{nC_1} \sum_{\tau_0 \in \mathcal{T}_R} \prod_{\{k,l\} \in \tau_0} |\beta \Delta_{kl} J(\sigma)| , \]
where from Lemma 5.5 we have that \( C_1 \simeq 2^{d+1} \| \nabla V \|_{\infty} \). We also let \( \sup_{\sigma} |\Delta_{kl} J(\sigma)| \leq \Delta_{kl} \) with \( \Delta_{kl} \equiv q^{-2d+1} \frac{1}{L^r} \| \nabla V \|_{\infty} 1_{(k,l) : |l-k| \leq \frac{L}{q}} \). Moreover, from Condition 5.1 we have

\[
\int \nu(d\sigma|\eta) \prod_{j=1}^{p} |\Phi_{k_j}^{\alpha_j}| \leq (C_2)^p.
\]

Then for the activity \( w(R') \) we obtain

\[
|w(R')| \leq e^{nC_1} \sum_{\tau^0 \in T_n} \prod_{(k,l) \in \tau^0} (\beta \Delta_{kl}) \cdot (C_2)^p.
\]

Thus to satisfy the sufficient condition for the convergence of the cluster expansion we first bound the sum \( \sum_{R' \sim R} \) by

\[
\sup_{k_0 \in \text{supp}(R)} |\text{supp}(R)| \sum_{R': \text{supp}(R') \supset \{k_0\}} \sum_{p \geq 1} \sum_{l_1,\ldots,l_p} |w(R')|e^{\alpha_{l_p}} + \sum_{n \geq 2} \frac{1}{(n-1)!} \sum_{k_1=k_0, k_2,\ldots,k_n} |w(R')|e^{\alpha_n}
\]

\[
+ \sum_{n \geq 2} \frac{1}{(n-1)!} \sum_{k_1=k_0, k_2,\ldots,k_n} \sum_{p_1,\ldots,p_n} \sum_{l_1,\ldots,l_{p_1}} \sum_{l_2,\ldots,l_{p_2}} \cdots \sum_{l_n,\ldots,l_{p_n}} |w(R')|e^{(p+n)}.
\]

Next, we use (5.27) where for every tree \( \tau^0 \) we have that

\[
\sup_{k \in R'} \sum_{k_1, k_2,\ldots,k_n} \prod_{i,j \in \tau^0} (\beta \Delta_{k_i,k_j}) \leq (\beta C_1)^{n-1}.
\]

By Cayley’s formula \( \sum_{\tau^0 \in T_n} 1 = n^{n-2} \) and the fact that the cardinality of the sum \( \sum_{\bigcup_{i=1}^{n-1} l_i \supset \{k_0\}} \) can be bounded by \( r^n \), where \( r = 2^{d+1} [3(2d+1)+1]d \) is an upper bound for the maximum number of \( C^\alpha(k) \) bonds that can pass through a point, as showed in [33], we obtain the geometric series

\[
\sum_{p \geq 1} \sum_{l_1,\ldots,l_p} |w(R')|e^{\alpha_p} \leq \sum_{p \geq 1} (rC_2e)^p = \frac{rC_2e}{1 - rC_2e}
\]

when there are only links of type 1 (short range),

\[
\sum_{n \geq 2} \frac{1}{(n-1)!} \sum_{k_1=k_0, k_2,\ldots,k_n} |w(R')|e^{\alpha_n} \leq e^{\beta C_1 + \epsilon} \sum_{n \geq 2} (\beta C_1 e^{\beta C_1 + \epsilon})^{n-1}
\]
for only links of type 2 (long range) and
\[
\sum_{n \geq 2} \frac{1}{(n-1)!} \sum_{k_1=k_0} \sum_{k_2, \ldots, k_n} \sum_{p_1 \geq 1} \sum_{\ldots} \sum \left| w(R') \right| e^{c(p+n)}
\leq \left( rC_2 e \right)^n \sum_{n \geq 2} \frac{1}{(n-1)!} \sum_{k_1=k_0} \sum_{k_2, \ldots, k_n} \left| w(R') \right| \left( \frac{rC_2 e}{1 - rC_2 e} \right)^n
\]
for the mixed case. Choosing $c = 1$, there are $\delta_1$ and $\delta_2$ sufficiently small such that if $C_1 < \delta_1$ and $rC_2 e < \delta_2$, then
\[
\frac{rC_2 e}{1 - rC_2 e} + e^{\beta C_1 + e} \frac{\beta C_1 e^{C_1 + 1}}{1 - \beta C_1 e^{C_1 + 1}} + e^{\beta C_1 + e} \frac{rC_2 e}{1 - rC_2 e} - e^{\beta C_1 + e} \frac{rC_2 e}{1 - rC_2 e} < \frac{1}{C};
\]
hence (5.26) holds. \hfill \square

Summarizing, one can formulate the $d$-dimensional version of Theorem 3.3.

**Theorem 5.7.** Let
\[
\tilde{H}_M^{(0)}(\eta) = \tilde{H}_M^{(t)}(\eta) + \tilde{H}_M^{(s)}(\eta),
\]
where $\tilde{H}_M^{(0)}(\eta)$ and $\tilde{H}_M^{(s)}(\eta) := -\log R^s(\eta)$ are given in (3.3) and (5.13) respectively (for $d = 2$ the latter is also given explicitly in (5.8)). Then, we have the error bound
\[
\left| \tilde{H}_M - \tilde{H}_M^{(0)} \right| = N\mathcal{O} \left( \frac{C_2 q^d}{q} + \frac{q^3 |\nabla V|_\infty}{L} \right).
\]

The loss of information when coarse-graining at the level $q$ is quantified by a specific relative entropy error similar to (3.29). Moreover, there exist $\delta_1(\beta, d) > 0$ and $\delta_2(\beta, d) > 0$ such that if $C_1 < \delta_1$ and if Condition 5.1 holds with $C_2 < \delta_2$, then $\tilde{H}_M - \tilde{H}_M^{(0)}$ is expanded in an absolutely convergent series in the parameters $C_1$ and $C_2$.

\[
\tilde{H}_M(\eta) = \tilde{H}_M^{(0)}(\eta) + \tilde{H}_M^{(1)}(\eta) + \ldots + \tilde{H}_M^{(p)}(\eta) + M\mathcal{O}((\max \{C_1, C_2\})^{p+1}),
\]
where the terms $\tilde{H}_M^{(p)}$ are determined by evaluating the terms of the series (5.24).

**Proof.** It is a direct consequence of Theorem 5.1 and Lemma 5.6. \hfill \square

**Remark 5.1.** As we also discussed in Remark 3.3 in a one-dimensional setting, estimates such as the ones in Theorem 5.7 relying on the constant $C_2$ of Condition 5.1 are sharper than the estimates in (3.29) which are derived from a counting argument such as (3.21).

**Remark 5.2.** For more general boundary conditions than periodic, e.g., zero, we need to confine the system in a box $\Lambda$ and fix a configuration outside. With minor modifications in the proof (essentially splitting into cells well inside the bulk and cells that can be affected by the boundary conditions), Theorems 3.3 and 5.7 still hold true, but with an extra error coming from the cells interacting with the boundary of the box which will be even smaller by a factor $|\partial \Lambda|/|\Lambda|$. 

License or copyright restrictions may apply to redistribution; see https://www.ams.org/journal-terms-of-use
6. Computational complexity of coarse-graining schemes

One of the outcomes of the numerical analysis of coarse-graining schemes via the cluster expansion analysis (3.31) is the reduction of the complexity in the calculation of each of the terms in the series into a combination of low-dimensional local potentials, e.g. (3.3) and (3.26). Therefore, a key challenge in the implementation of these coarse-graining methods is to develop strategies that efficiently address the computational complexity of pre-computing local multi-body terms, for instance (3.28) or (5.8). We already showed in Section 4.3 in the case of one dimension that the computational complexity of local terms such as (3.28) can be greatly reduced by exact analytical calculations and/or additional approximations, yielding semi-analytical splitting schemes which are effectively multi-body combinations of easily computable one-body terms such as (4.7), or in Section 4.3.1 through analytical expressions such as (4.10), as well as (4.11) and (4.12) in the high-temperature approximation of each of the local terms.

In two dimensions the same strategy is also applicable as we briefly outline next. Following the approach in (4.2) we rewrite (5.8) by replacing the partition functions \( Z(A_4(k)) \) by \( \tilde{Z}(A_4(k)) \), which denotes a partition function with respect to the normalized measure (4.1). As in the case of dimension one in Section 4 our goal is to simplify as much as possible the terms \( \tilde{Z}(A_4(k)) \), for \( k \in \bar{\Lambda}^\alpha_M \), \( \alpha = 1, 3, 4 \), and give an algorithm to compute them efficiently. Note that in the two-dimensional case the many-body terms seem intractable due to the multi-canonical constraints, e.g. for the case \( \alpha = 4 \), see Figure 2, we have to compute \( \tilde{Z}(A_4(k)) \) for all possible combinations of 9 values of \( \eta \).

However, for the nearest-neighbor Ising model we can perform an analytical calculation and/or approximation and simplify the terms with significant reduction of the computational complexity: similarly to Section 4.3 we denote the lattice sites within a coarse cell \( C_k \) by \( x_{i,j} \), \( i, j = 1, \ldots, q \), with \( x_{1,1} \) being the upper-left one. We present in full detail only the computation of \( \tilde{Z}(A_4(k)) \), with \( k \in \bar{\Lambda}^1_M \). We also denote by \( k^l \) and \( k^r \) the left and right coarse cell to \( C_k \). We have:

\[
\tilde{Z}(A_4(k)) = \int \prod_{i=1}^{q} \left[ (1 - \lambda \sigma^{k^l}(x_{i,q}) \sigma^{k^r}(x_{i,1}))(1 - \lambda \sigma^{k^l}(x_{i,q}) \sigma^{k^r}(x_{i,1})) \right] \hat{P}_{k^l} \hat{P}_{k^r} = 1 - \lambda \sum_{i=1}^{q} \left( \Phi^{i,q}_{k^l} \Phi^{i,1}_{k^r} + \Phi^{i,q}_{k^r} \Phi^{i,1}_{k^l} \right)
\]

(6.1)

\[+ \lambda^2 \sum_{i<j} \left( \Phi^{i,q}_{k^l} \Phi^{(i,1),(j,q)}_{k^r} + \Phi^{i,q}_{k^r} \Phi^{(i,1),(j,q)}_{k^l} + \Phi^{(i,q),(j,1)}_{k^r} \Phi^{i,1}_{k^l} \right) + O(\lambda^3),\]

where, e.g. \( \Phi^{i,q}_{k^l} := \int \sigma^{k^l}(x_{i,q}) \hat{P}_{k^l} \) is a one-point correlation function and \( \Phi^{(i,1),(j,q)}_{k^r} \) := \( \int \sigma^{k^r}(x_{i,1}) \sigma^{k^r}(x_{j,q}) \hat{P}_{k^r} \) the two-point correlation.

The computation of \( \tilde{Z}(A_4(k)) \) for \( k \in \bar{\Lambda}^3_M \) is identical with the \( k \in \bar{\Lambda}^1_M \) case discussed above, while the computation of \( \tilde{Z}(A_4(k)) \) for \( k \in \bar{\Lambda}^4_M \) is somewhat more involved since one has to take into account all the possible interactions within the 9-cell \( A_4(k) \), with \( k \in \bar{\Lambda}^4_M \).

These calculations allow us to reduce the 9-body term in (5.8) into combinations of one-body terms; see also (4.5) for the one-dimensional analogue, which in turn can
be easily calculated: for every value of \( \eta \) we create a library with the values of the above functions; for each one we need to run a constrained Monte Carlo algorithm but it should converge quite fast since it is restricted only on one coarse cell. Note also that due to the symmetries several of the correlation functions appearing above are the same and need to be computed once. Imposing periodic boundary conditions on the coarse cells (instead of free) and committing an additional error can further simplify all such calculations.

Further approximations for the local multi-body terms such as high temperature expansions, (4.11) and (4.12), as well as analytical formulas for the correlations (4.10) (in higher dimensions such formulas hold away from phase transitions), can give rise to further simplifications in the multi-body terms. Finally, we can approximate \( \log \hat{Z}(A_4(k)) \) in yet a different way by carrying out an additional cluster expansion of the interactions within the set \( A_4(k) \).

In view of the preceding discussion regarding pre-computing multi-body terms, it seems necessary to re-examine whether we should coarse-grain at all! We showed in this paper that coarse-graining with controlled-errors typically requires multi-body terms derived via cluster expansions. On the other hand, in order to avoid the computations with the multi-body terms we may consider an approximate microscopic scheme with similar error control features. More specifically, we may consider an approximating Hamiltonian where only the long-range potential is compressed by a local averaging over each coarse cell, i.e., we define the new Hamiltonian,

\[
(6.2) \quad H_N(\sigma) = H_N^s(\sigma) + H_M^{L,(0)}(F(\sigma)),
\]

where the coarse-graining map \( F \) is defined in (1.3) and the coarse-grained long-range Hamiltonian \( H_M^{L,(0)} \) in (3.3). It is straightforward to show that the corresponding microscopic measure has the same error estimates as the one in Theorem 3.3, where in addition the error due to the short range interactions is absent since the corresponding part is computed exactly. This approach would be similar in spirit to various interaction compression techniques such as the multi-pole method.

We briefly discuss how this microscopic algorithm with compressed long-range interactions compares, in terms of computational complexity, with the fully coarse-grained simulation. First, the multi-body CG interactions involve only the short-range microscopic interactions and can be either pre-computed off-line as in the current literature in polymeric and bio-molecular systems or one can employ the semi-analytical controlled-error approximations, discussed in the paper. In either case they will not affect the computational complexity or the CPU count of a coarse-grained algorithm. Similarly, in (6.2) the cost of simulating microscopic short-range interactions is overwhelmed by the long-range ones. Thus we can focus on a complexity and CPU comparison by ignoring the nearest-neighbor interactions. Indeed, such CPU comparisons between microscopic models with piecewise constant potentials, as the one in (6.2), and coarse-grained models were already carried out in earlier publications; see for example Figure 9 in [21]. In these comparisons we see a very substantial speed-up in the coarse-grained model, which is at least quadratic in the coarsening factor \( q \); see also the operation count in Table 1.

In general, when comparing coarse-graining schemes to the direct numerical simulation of the microscopic variables \( \sigma \) such as (6.2), we readily see that they reduce the computational complexity of the simulation of lattice systems in two ways. First, long-range interactions are compressed through the effective interactions in
Table 1. Computational complexity of evaluating the Hamiltonian on the $d$-dimensional lattice for the interaction range $L$ and the coarse-graining level $q$ (24).

<table>
<thead>
<tr>
<th>Hamiltonian</th>
<th>Count</th>
<th>Speed-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microscopic $q = 1$: $H_N^{(\sigma)}$</td>
<td>$O(NL^d)$</td>
<td>1</td>
</tr>
<tr>
<td>Coarse-graining scheme $H_M^{(0)}$:</td>
<td>$O(ML^d/q^d)$</td>
<td>$O(q^{2d})$</td>
</tr>
</tbody>
</table>

Table 2. Computational complexity on the $d$-dimensional lattice for the interaction range $L$ and the coarse-graining level $q$: number of classes of lattice sites with equal rates for $\{-1, 1\}$-spins.

<table>
<thead>
<tr>
<th>Microscopic CG with $q$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$O(2^{L^d})$</td>
</tr>
</tbody>
</table>

(3.3): see Table 1 for the corresponding operation count for the evaluation of a Hamiltonian. Second, Kinetic Monte Carlo simulations of lattice systems with such complex interactions at a microscopic level have a high number of rejections when one considers null-event algorithms, 29, while rejection-free methods such as the Bortz, Kalos, Lebowitz (BKL) algorithm, 29, have an exponentially large number of classes of sites with equal rates, rendering this sophisticated algorithm impractical. On the other hand, due to the compression of the configuration space, coarse-graining yields a vastly reduced number of such classes; see Table 2 for an exact comparison with the microscopic case, which in turn can allow for an efficient implementation of the BKL method.

7. Conclusions

In this paper, we developed coarse-graining schemes for stochastic many-particle microscopic models with competing short- and long-range interactions on a $d$-dimensional lattice. We focused on the coarse-graining of equilibrium Gibbs states, and using cluster expansions we analyzed the corresponding renormalization group map. We quantified the approximation properties of the coarse-grained terms arising from different types of interactions and presented a hierarchy of correction terms. Finally, we derived semi-analytical numerical coarse-graining schemes along with a posteriori error estimates for lattice systems with short and long-range interactions.

An important outcome of the cluster expansion analysis for the approximation of coarse-grained Hamiltonian by the series (1.8) is the reduction of the complexity in the calculation of each of the terms in the series into a combination of low-dimensional local potentials, e.g. (3.26). In turn, we show in Section 4 and Section 6 that exact analytical calculations and/or additional approximations can greatly reduce the computational complexity of such local terms, yielding semi-analytical splitting schemes for the coarse-graining of lattice systems with short and long-range interactions. Furthermore, these schemes can be split, within a controllable approximation error, into a long and a short-range calculation; see (3.25). The
long-range part, which is computationally expensive for conventional Monte Carlo methods, can be cheaply simulated using the analytical formula given in (3.2) in the spirit of our previous work [24]. In this case computational savings are due to the reduction in the degrees of freedom by \( Q = N/M \) and compressing the range of interactions. For the short-range interactions we use the semi-analytical formulas (4.2) which involve pre-computing coarse-grained interactions with Monte Carlo simulation. However, the simulation is done for a single subdomain of adjacent coarse cells, while for Ising-type models, see Section 4 and Section 6, one can reduce the pre-computation on a single coarse cell. The error estimates in Theorem 3.3 and Theorem 5.7 also suggest an improved decomposition to short and long-range interactions. Indeed, they imply splitting and rearranging the overall combined short and long-range potential into a new short-range component that includes possible singularities originally in the long-range interaction, e.g., the non-smooth part in a Lennard-Jones potential, and a locally integrable (or smooth) long-range decaying component. Presumably similar strategies could be applied for off-lattice systems such as the coarse-graining of polymers.

In contrast to the splitting approach developed here that allows us to analytically calculate the long-range effective Hamiltonian (3.3) in (3.25) and in parallel carry out the semi-analytical step for (4.2), existing methods, e.g., [16,27], employ semi-analytical computations involving both short, as well as costly long-range interactions. Thus, multi-body terms, which are believed to be important at lower temperatures, [16], have to be disregarded. A notable result of our error analysis is the quantification of the role of multi-body terms in coarse-graining schemes, and the relative ease to implement them using the aforementioned splitting schemes. Theorem 3.3 addresses this issue, while in Section 4 we further quantify the regimes where multi-body terms are necessary in the context of a specific example. In [2] the necessity to include multi-body terms in the effective coarse-grained Hamiltonian was first discussed in a numerical analysis context for systems with singular (at the origin) long-range interactions.

Cluster expansions such as (1.8) can also be used for constructing a posteriori error estimates for coarse-graining problems, based on the rather elementary observation that higher-order terms in (3.31) can be viewed as errors that depend only on the coarse variables \( \eta \). In [22] we already employed this type of estimate for stochastic lattice systems with long-range interactions in order to construct adaptive coarse-graining schemes. These tools operated as an “on-the-fly” coarsening/refinement method that accurately recovers phase diagrams. The estimates allowed us to adaptively change the coarse-graining level within the coarse-graining hierarchy once suitably large or small errors were detected, and thus to speed up the calculations of phase diagrams. Adaptive simulations for molecular systems have also been recently proposed in [36], although they are not based on an a posteriori error analysis perspective. Finally, the cluster expansions necessary for the rigorous derivation and error estimates of the schemes developed here rely on the smallness of a suitable parameter introduced in Theorem 3.3; see (3.30). In Section 4 we construct an a posteriori bound for this quantity that can allow us to track the validity of the cluster expansion for a given resolution in the course of a simulation. This approach is, at an abstract level, similar to conditional a posteriori estimates proposed earlier in the numerical analysis of geometric partial differential equations, [15,28].
Further challenges for systems with short and long-range interactions not discussed here include: error estimates for observables/quantities of interest, the development of coarse-grained dynamics from microscopics, phase transitions and estimation of physical parameters, such as critical temperatures. Work related to these directions for systems with long-range interactions have been carried out in [25], [6] and [5].

ACKNOWLEDGMENTS

The authors acknowledge valuable comments and suggestions by the anonymous referees that led to a substantially improved manuscript.

REFERENCES


DEPARTMENT OF MATHEMATICS, UNIVERSITY OF MASSACHUSETTS, AMHERST, MASSACHUSETTS 01003 AND DEPARTMENT OF APPLIED MATHEMATICS, UNIVERSITY OF CRETE AND FOUNDATION OF RESEARCH — AND — TECHNOLOGY-HELLAS, GREECE

E-mail address: markos@math.umass.edu

DEPARTMENT OF MATHEMATICAL SCIENCES, UNIVERSITY OF DELAWARE, NEWARK, DELAWARE 19716

E-mail address: plechac@math.udel.edu

DEPARTMENT OF MATHEMATICS, UNIVERSITY OF MASSACHUSETTS, AMHERST, MASSACHUSETTS 01003

E-mail address: lr7q@math.umass.edu

HAUSDORFF CENTER FOR MATHEMATICS, UNIVERSITY OF BONN, D-53115 BONN, GERMANY

E-mail address: dtsagkar@gmail.com