# Irreversible Markov chains, from the TASEP to all-atom Coulomb computations 

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E. P. Bernard, W. Krauth, D. B. Wilson, PRE (2009)

Event-chain algorithms for hard-sphere systems
E. P. Bernard, W. Krauth, PRL (2011)
M. Michel, S. C. Kapfer, W. Krauth, JCP (2014)
S. C. Kapfer, W. Krauth, PRL (2015)
S. C. Kapfer, W. Krauth, PRL (2017)
Z. Lei, W. Krauth (EPL 2018, EPL 2019)
M. F. Faulkner, L. Qin, A. C. Maggs, W. Krauth, JCP (2018)
P. Höllmer, L. Qin, M. F. Faulkner, A. C. Maggs, W. Krauth, arXiv1905XXX JeLLyFysh (Version 1.0) - a Python application for all-atom event-chain Monte Carlo

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# Equation of State Calculations by Fast Computing Machines 

Nicholas Metropolis, Arianna W. Rosenbluth, Marshall N. Rosenbluth, and Augusta H. Teller, Los Alamos Scientific Laboratory, Los Alamos, New Mexico

AND
Edward Teller,* Department of Physics, University of Chicago, Chicago, Illinois (Received March 6, 1953)

A general method, suitable for fast computing machines, for investigating such properties as equations of state for substances consisting of interacting individual molecules is described. The method consists of a modified Monte Carlo integration over configuration space. Results for the two-dimensional rigid-sphere system have been obtained on the Los Alamos MANIAC and are presented here. These resuits are compared to the free volume equation of state and to a four-term virial coefficient expansion.


distinguished by primes. For example, $A_{33}$ is given schematically by the diagram

and mathematically as follows: if we define $f\left(\boldsymbol{r}_{i j}\right)$ by

$$
\begin{array}{lll}
f\left(r_{i j}\right)=1 & \text { if } & r_{i j}<d, \\
f\left(r_{i j}\right)=0 & \text { if } & r_{i j}>d,
\end{array}
$$

then

$$
A_{\mathrm{2}, 3}=\frac{1}{\pi^{2} d^{4}} \int \cdots \int d x_{1} d x_{2} d x_{3} d y_{1} d y_{2} d y_{3}\left(f_{12} f_{23} f_{31}\right)
$$

The schematics for the remaining integrals are indicated in Fig. 6.
The coefficients $A_{3,3}, A_{4,4}$, and $A_{4,5}$ were calculated
were put down at random, subject to $f_{12}=f_{23}=f_{34}$ $=f_{15}=1$. The number of trials for which $f_{45}=1$, divided by the total number of trials, is just $A_{5,5}$.

The data on $A_{4,6}$ is quite reliable. We obtained

## VI. CONCLUSION

The method of Monte Carlo integrations over configuration space seems to be a feasible approach to statistical mechanical problems which are as yet not analytically soluble. At least for a single-phase system a sample of several hundred particles seems sufficient. In the case of two-dimensional rigid spheres, runs made with 56 particles and with 224 particles agreed within statistical error. For a computing time of a few hours with presently available electronic computers, it seems possible to obtain the pressure for a given volume and temperature to an accuracy of a few percent.
In the case of two-dimensional rigid spheres our results are in agreement with the free volume approximation for $A / A_{0}<1$ - 8 -and witb-a-five-term-vicial expansion for 4 th $\geqslant 2.5$. There is no indication of a phase transition.


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## Alder-Wainwright (1962)

## Phase Transition in Elastic Disks*

B. J. Alder and T. E. Wainwright

University of California, Lawrence Radiation Laboratory, Livermore, California
(Received October 30, 1961)
The study of a two-dimensional system consisting of 870 hard-disk urticles in the phase-transition region has shown that the isotherm has a van der Waals-like loop. The nsity change across the transition is about $4 \%$ and the corresponding entropy change is small.

ASTUDY has been made of a two-dimensional sysitem conssisting of 870 hard-disk particles. Simultaneous motions of the particles have been calcuHated by-means-of-an-electronic computer as described previously. ${ }^{1}$ The disks were again placed in a periodically repeated rectangular array. The computer program
interchanges it was not possible to average the two branches.

Two-dimensional systems were then studied, since the number of particles required to form clusters of particles of one phase of any given diameter is less than in three dimensions. Thus, an 870 hard-disk system is

## $2 D$ melting transition



$$
\eta=0.48
$$


$\eta=0.72$

- Generic 2D systems cannot crystallize (Peierls, Landau 1930s) but they can turn solid (Alder \& Wainwright, 1962).
- Nature of transition disputed for decades.


## Kosterlitz-Thouless (1973)

## Ordering, metastability and phase transitions in two-dimensional systems

## J M Kosterlity and D J Thouless

Departmentofiviathematical'Physics, Unitiversity of Birmingham, Birmingham B15 2TT, UK

Received 13 Novembe

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1972
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## 1. Introduction

Peierls (1935) has argued that thermal motion of long-wavelength phonons will destroy the long-range order of a two-dimensional solid in the sense that the mean square deviation of an atom from-its equilibrium position increases logarithmically with the size of the system, and the Bragg peaks of the diffraction pattern formed by the system are broad instead of sharp. The absence of long-range order of this simple form has been shown by Mermin (1968) using rigorous inequalities. Similar arguments can be used to show that there is no spontaneous magnetization in a two-dimensional magnet with spins with more than one degree of freedom (Mermin and Wagner 1966) and that the expectation value of the superfluid order parameter in a two-dimensional Bose fluid is_zero (Hohenberg 1967).
On the other hand there is inconclusive evidence from the numerical work on a two-dimensional system of hard discs by Alder and Wainwright (1962) of a phase transition between a gaseous and solid state. Stanley and Kaplan (1966) found that hightemperature series expansions for two-dimensional spin models indicated a phase

Possible phases in two dimensions


| Phase | positional order | orientational order |
| :--- | :--- | :--- |
| solid | algebraic | long-range |
| hexatic | short-range | algebraic |
| liquid | short-range | short-range |

## Correlation time in larger simulations

disk $k$

same disk

$t=25600000000$

- $\tau$ exists, but it is large $(\tau \gg 25600000000)$.


## Detailed balance - global balance


global balance

detailed balance

maximal global balance

- flow into $a=$ Boltzmann weight $\pi(a)$ (global balance condition):

$$
\underbrace{\sum_{k} \pi(k) p(k \rightarrow a)}_{\text {flow into a } \sum_{k} \mathcal{F}(k \rightarrow a)}=\pi(a)
$$

- flow $\mathcal{F}(a \rightarrow b) \equiv$ flow $\mathcal{F}(b \rightarrow a)$ (detailed balance condition):

$$
\underbrace{\pi(b) p(b \rightarrow a)}_{\text {from } b \text { to } \operatorname{F} \mathcal{F}(b \rightarrow a)}=\underbrace{\pi(a) p(a \rightarrow b)}_{\mathcal{F}(a \rightarrow b) \text { flow from } a \text { to } b}
$$

## Detailed balance - global balance


global balance

detailed balance

maximal global balance

- flow into $a=$ Boltzmann weight $\pi(a)$ (global balance condition):

$$
\underbrace{\sum_{k} \pi^{(t-1)}(k) p(k \rightarrow a)}_{\text {flow into a } \sum_{k} \mathcal{F}(k \rightarrow a)}=\pi^{(t)}(a)
$$

- flow $\mathcal{F}(a \rightarrow b) \equiv$ flow $\mathcal{F}(b \rightarrow a)$ (detailed balance condition):

$$
\underbrace{\pi(b) p(b \rightarrow a)}=\underbrace{\pi(a) p(a \rightarrow b)}
$$

flow from $b$ to a $\mathcal{F}(b \rightarrow a) \quad \mathcal{F}(a \rightarrow b)$ flow from $a$ to $b$

## Detailed balance - global balance


global balance

detailed balance

maximal global balance

- flow into $a=$ Boltzmann weight $\pi(a)$ (global balance condition):

$$
\underbrace{\sum_{k} \pi(k) p(k \rightarrow a)}_{\text {flow into a } \sum_{k} \mathcal{F}(k \rightarrow a)}=\pi(a)
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- flow $\mathcal{F}(a \rightarrow b) \equiv$ flow $\mathcal{F}(b \rightarrow a)$ (detailed balance condition):

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$$

## Reversible Metropolis algorithm, 1d (detailed balance)



- Local Metropolis: $x_{i} \rightarrow x_{i} \pm \epsilon$ (reject if overlap, $\epsilon>0$ )
- Detailed balance:

$$
\pi_{a} p(a \rightarrow b)=\pi_{b} p(b \rightarrow a)
$$

## Reversible Metropolis algorithm, 1d (detailed balance)



- Local Metropolis: $x_{i} \rightarrow x_{i} \pm \epsilon$ (reject if overlap, $\epsilon>0$ )
- Detailed balance:

$$
\pi_{a} p(a \rightarrow b)=\pi_{b} p(b \rightarrow a)
$$

## Reversible Metropolis algorithm, 1d (global balance)



- Reversible Metropolis: $x_{i} \rightarrow x_{i} \pm \epsilon$ (reject if overlap, $\epsilon>0$ )
- Global balance:

$$
\mathcal{F}_{a}^{\mathrm{rev}}=\frac{1}{2 N} \sum_{i} \underbrace{\left(\mathcal{A}_{i}^{+}+\mathcal{R}_{i}^{+}+\mathcal{A}_{i}^{-}+\mathcal{R}_{i}^{-}\right)}_{=2 \text { for any } \epsilon}=1
$$

- NB: $\mathcal{A}_{i}^{+}(\epsilon)+\mathcal{R}_{i}^{-}(\epsilon)=1$ also $\mathcal{A}_{i}^{-}(\epsilon)+\mathcal{R}_{i}^{+}(\epsilon)=1$.


## Sequential Metropolis algorithm, 1d (global balance)

a


- Sequential Metropolis: Update 0 , then 1 , then $2, \ldots$
- Global balance:

$$
\mathcal{F}_{a}^{\text {seq }}=\frac{1}{2}\left(\mathcal{A}_{i}^{+}+\mathcal{R}_{i}^{+}+\mathcal{A}_{i}^{-}+\mathcal{R}_{i}^{-}\right)=1
$$

Our method in this respect is similar to the cell method except that our cells contain several hundred particles instead of one. One would think that such a sample would be quite adequate for describing any onephase system. We do find, however, that in two-phase systems the surface between the phases makes quite a nerturbation. Also. statistical fluctuations mav be
configurations with a probability $\exp (-E / k T)$ and weight them evenly.

This we do as follows: We place the $N$ particles in any configuration, for example in a regular lattice. Then we move each of the particles in succession according to the fortowing prescription:

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- Forward Metropolis: $x_{i} \rightarrow x_{i}+\epsilon$ (reject if overlap, $\left.\epsilon>0\right)$
- 

$$
\mathcal{F}_{a}^{\text {forw }}=\frac{1}{N} \sum_{i} \underbrace{\left(\mathcal{A}_{i}^{+}+\mathcal{R}_{i-1}^{+}\right)}_{=1 \text { for any } \epsilon}=1,
$$

- NB: Forward sequential Metropolis is wrong (but there is a variant that is OK )


## Lifted Forward Metropolis algorithm, 1d (global balance)

- Move $i$ forward until it is rejected by $i+1$.
- Then move $i+1$ forward until it is rejected, etc.

- $\mathcal{F}_{(a, i)}^{\text {lift }}=\mathcal{A}_{i}^{+}+\mathcal{R}_{i-1}^{+}=1$.
- NB: 1 time step: 1 particle move OR 1 lifting move.
- Infinitesimal $\epsilon \rightarrow 0$ version: Event-chain algorithm.

| Algorithm | mixing | discrete analogue |
| :--- | :--- | :--- |
| Rev. Metropolis | $N^{3} \log N$ | Symmetric SEP |
| Forward Metropolis, Lifted | $N^{5 / 2}$ | TASEP |
| Event-chain, Lifted (restarts, a) | $N^{2} \log N$ | lifted TASEP |
| Event-chain, Lifted (restarts, var) | $N^{2}$ | lifted TASEP (var.) |

- For Symmetric SEP mixing of Lacoin (2014).
- For TASEP mixing cf Baik \& Liu (2016).
- All others cf Kapfer \& Krauth (2017).
- Rigorous Proofs for event-chain cf Lei \& Krauth (2018).

NB: All algorithms converge towards equilibrium.

## Faster algorithm: Event-chain algorithm again



- Bernard, Krauth, Wilson (2009).
- Infinitesimal moves: No multiple overlaps, consensus.
- Michel, Kapfer, Krauth (2014) (smooth potentials).


## Event-chain algorithm, general d (global balance)



## Checking global balance (hard spheres, general d)



## Checking global balance (hard spheres, general d)



## Checking global balance (hard spheres, general d)



## Hard-disk configuration



- $1024^{2}$ hard disks
- Bernard, Krauth (PRL 2011)


## equation of state




- Many confirmations (PRE Milestone (2013)).


## Possible phases (again)

| Phase | positional order | orientational order |
| :--- | :--- | :--- |
| solid | algebraic | long-range |
| hexatic | short-range | algebraic |
| liquid | short-range | short-range |

## Spatial correlations at $\eta=0.718$ and 0.720



- Two-dimensional pair correlations, sample-averaged.
- At $\eta=0.718$; hexatic: First-order liquid-hexatic transition.
- At $\eta \sim 0.720$ : KT-type hexatic-solid transition.
- Bernard \& Krauth (PRL 2011).
- Many confirmations.
- Soft disks: $V \propto(\sigma / r)^{n}$.


- Kapfer \& Krauth (PRL 2015).
- Two melting scenarios depending on softness $n$ of potential


## Factorized Metropolis algorithm



- Metropolis algorithm

$$
p^{\mathrm{Met}}(a \rightarrow b)=\min \left[1, \prod_{i<j} \exp \left(-\beta \Delta V_{i, j}\right)\right]
$$

- Factorized Metropolis algorithm (Michel, Kapfer, Krauth 2014)

$$
\begin{aligned}
& p^{\text {Fact. }}(a \rightarrow b)=\prod_{i<j} \min \left[1, \exp \left(-\beta \Delta V_{i, j}\right)\right] . \\
& X^{\text {Fact. }}(a \rightarrow b)=X_{1,2} \wedge X_{1,3} \wedge \cdots \wedge X_{N-1, N}
\end{aligned}
$$

## All-Atom Coulomb problem 1/5



- 3D water model: bond, bending, Lennard-Jones, Coulomb (SPC/Fw)


## All-Atom Coulomb problem 2/5



- 3D water model: bond, bending, Lennard-Jones, Coulomb (SPC/Fw)
- Factors and types.


## All-Atom Coulomb problem 3/5



- Factor $M=\left(I_{M}, T_{M}\right):\left|I_{M}\right|=6$, two molecules. $T_{M}=$ 'Coulomb'.


## All-Atom Coulomb problem 4/5



- Water model: bond, bending, Lennard-Jones, Coulomb (SPC/Fw)


## All-Atom Coulomb problem 5/5



- Complexity $\mathcal{O}(1)$ per 'lifting' move.


## ECMC for all-atom water simulations

- Oxygen-Oxygen distance for 32 water molecules.


See: Faulkner, Qin, Maggs, Krauth (2018).

## ECMC - JeLLyFysh



- cf Höllmer, Qin, Faulkner, Maggs \& Krauth:
- arXiv1905XXX
- 'JeLLyFysh' Open-source Python application for irreversible Markov chains


## Conclusions

- Hard disks and two-dimensional melting
- Detailed balance - global balance
- Sampling $\exp (-\beta U)$ without knowing $U$


## Conclusions

- Hard disks and two-dimensional melting
- Equilibrium - steady state
- Sampling $\exp (-\beta U)$ without knowing $U$


## Conclusions

- Hard disks and two-dimensional melting
- Equilibrium - steady state (indistinguishable from equilibrium)
- Sampling $\exp (-\beta U)$ without knowing $U$

