ENTROPY PRODUCTION IN COLLISIONLESS SYSTEMS. II. ARBITRARY PHASE-SPACE OCCUPATION NUMBERS

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Abstract

We present an analysis of two thermodynamic techniques for determining equilibria of self-gravitating systems. One is the Lynden-Bell (LB) entropy maximization analysis that introduced violent relaxation. Since we do not use the Stirling approximation, which is invalid at small occupation numbers, our systems have finite mass, unlike LB’s isothermal spheres. (Instead of Stirling, we utilize a very accurate smooth approximation for $\ln x$.) The second analysis extends entropy production extremization to self-gravitating systems, also without the use of the Stirling approximation. In addition to the LB statistical family characterized by the exclusion principle in phase space, and designed to treat collisionless systems, we also apply the two approaches to the Maxwell–Boltzmann (MB) families, which have no exclusion principle and hence represent collisional systems. We implicitly assume that all of the phase space is equally accessible. We derive entropy production expressions for both families and give the extremum conditions for entropy production. Surprisingly, our analysis indicates that extremizing entropy production rate results in systems that have maximum entropy, in both LB and MB statistics. In other words, both thermodynamic approaches lead to the same equilibrium structures.

Key words: galaxies: kinematics and dynamics – galaxies: structure

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1. INTRODUCTION

1.1. Motivation

Understanding how collisionless systems attain a specific mechanical equilibrium state is fundamentally important to astrophysics. For example, the cold dark matter structures that exist around galaxies are expected to fall into this class of system. Individual dark matter constituents (whatever they may be) should evolve according to a mean-field gravitational potential, free of the influence of individual encounters.

The range of mechanical equilibria available to a collisionless system is defined by the Jeans equation, which represents the condition that no portion of the system experiences a net force. Unfortunately, the Jeans equation admits an infinity of solutions. Even if the mass distribution is specified, there is an infinite set of acceptable mechanical equilibria, each involving a different velocity distribution. For spherical systems, these velocity distributions differ in their anisotropy profile that quantifies radial versus tangential motion. However, the question remains, how and/or why does any one collisionless system evolve to its particular mechanical equilibrium end state, and what are the properties of such a state?

1.2. Thermodynamic Approaches to the Problem

The statistical mechanics description of thermodynamics provides one path to obtaining the description of the final relaxed state. A fully relaxed system is the most statistically likely state of that system or the one with an entropy maximum. In calculating the most likely state, it is implicitly assumed that all states of the system are equally accessible.

Such an approach was taken by Lynden-Bell (1967) and applied to self-gravitating collisionless systems, with the hope of explaining the observed light distribution of elliptical galaxies. A collisionless system can be thought of as a fluid in the six-dimensional phase space of position and velocity. The “particles” in Lynden-Bell’s (LB’s) analysis are parcels of phase space, i.e., parcels of this fluid, and so the distribution function (DF) representing the phase-space density is defined in terms of energy per unit mass, not energy per particle. The analysis resulted in a DF similar to the Fermi–Dirac (FD) case, but with a different normalization. Lynden-Bell (1967) argued that a non-degenerate limit is appropriate for stellar systems, and thus arrived at a DF similar to that of the Maxwell–Boltzmann (MB) case (an exponential), which resulted in the isothermal sphere representing thermal equilibrium. Since the isothermal sphere has an infinite extent and mass, its emergence from the entropy maximization procedure, which demanded a finite mass system, presented a contradiction. This apparent failure of entropy maximization was puzzling, and it was often argued that such systems do not have states of maximum entropy. Some effort was made to investigate maximizing entropy with additional constraints beyond mass, energy, and angular momentum (Stiavelli & Bertin 1987; White & Narayan 1987). Other routes involving minimum energy states of self-gravitating systems were also developed (e.g., Aly 1994).

Recently, Madsen (1996) (based on earlier work by Simons 1994) has pointed out that the reason for the system’s infinite mass was the use of the Stirling approximation, $\ln n! \approx n \ln n - n$. In contrast to systems usually treated in standard statistical mechanics, self-gravitating systems can apparently have small phase-space occupation numbers $n$, making Stirling a poor approximation. Specifically, these systems have regions of phase space or energy space that are nearly or completely unoccupied, such that $n$ will be small. Spatially, these regions can correspond to the center of the potential as well as its outer edge. Using MB statistics and the exact $\ln n!$, Madsen (1996) has found a DF from entropy maximization that is very similar to King (1966) models. Hjorth & Williams (2010) have shown that the energy-space occupation function $N(E)$ derived using a very accurate smooth approximation to $\ln n!$ closely resembles the results of collisionless N-body simulations (Williams et al. 2010).
1.3. Brief Review of Statistical Representations

From a statistical point of view, entropy is simply a measurement of the number of states accessible to a particular system. This relationship is most commonly expressed quantitatively as

\[ S = k_B \ln \Omega, \]

where \( \Omega \) is the number of accessible states and \( k_B \) is the Boltzmann constant, which serves to give entropy the correct thermodynamic units. As a result, counting procedures are key to determining specific realizations of entropy. Lynden-Bell (1967) discusses how there are four counting types that lead to physically relevant situations. Bose-Einstein statistics follow from counting states for indistinguishable particles that can co-habitate in the same state. When indistinguishable particles are not allowed to share states, FD statistics emerge. Classical MB statistics result from counting states available to distinguishable particles that can share states. Completing the symmetry, systems where distinguishable “particles”—actually, parcels of phase space—cannot co-occupy states obey what has become known as LB statistics (statistics “IV” in LB’s original notation). Each type of statistics will produce different representations of entropy, but we will focus on the two that deal with classical, or distinguishable particles, namely, LB and MB.

We briefly recap the notation used in Lynden-Bell (1967) before proceeding with our discussion. The six-dimensional position–velocity phase space \((x, v)\) is the usual setting for determining the statistics. Imagine phase space to be divided into a very large number of nearly infinitesimal parcels, called micro-cells, each having volume \(\sigma\). Each micro-cell can either be occupied or unoccupied by one of the \(N\) phase-space elements of the system. These elements can be thought of as representing the fine-grained DF, which has a constant density value \(n\). Because collisionless processes imply incompressibility of the fine-grained DF, the phase elements cannot co-habitate. If phase space is also partitioned on a coarser level so that some number \(\nu\) of micro-cells occupy a macro-cell, then we can discuss a coarse-grained DF. The volume of a macro-cell is then \(\nu \sigma\) and the \(i\)th macro-cell contains \(n_i\) phase elements. We assume that while the volume of a macro-cell is much larger than that of a micro-cell, it is still very small compared with the full extent of phase space occupied by the system. The number of ways of organizing the \(n_i\) elements into the \(\nu\) micro-cells without co-habitation is

\[ \frac{\nu!}{(\nu - n_i)!}. \]

If the elements were allowed to multiply occupy micro-cells, as in MB statistics, this number would be given by \(\nu^{n_i}\).

To get the total number of accessible states, the possible ways to distribute the \(N\) phase elements into \(n_i\) chunks must also be included. Lynden-Bell (1967) derives

\[ \Omega_{LB} = \frac{N!}{\prod_i n_i!} \times \prod_i \frac{\nu!}{(\nu - n_i)!}. \]

In the MB case, the only change is that the factorial ratio in the final product term is replaced by \(\nu^{n_i}\).

The LB case disallows two phase-space elements from inhabiting the same phase-space location, so it explicitly takes into account the incompressibility of a collisionless fluid. Since we are primarily interested in dark matter halos, LB is the natural case to consider. For completeness, and for the sake of having a comparison, we also treat the MB case, in the Appendix.

We argue that the lack of an exclusion principle in the MB case is equivalent to allowing collisions between particles. In a collisional system, particles from distant phase-space locations can be scattered into any other phase-space location, thereby increasing the phase-space density at the latter location. In principle, there is no limit to how high the density can get through such scattersings. In practice, the phase-space density probably cannot become very high at most locations, but it can be higher than the original fine-grained DF. We note that it is common to use MB to represent collisional systems. For example, Madsen (1996) argues that it is the correct statistics to use for globular clusters where the relatively small number of stars allows the cluster to relax through two-body interactions. It then makes sense that the energy distribution that the cluster will arrive at will be the same as that in a cloud of gas, which relaxes through collisions between molecules. In the non-degenerate limit, when the micro-cells are very sparsely populated and the density of the coarse-grained DF is very dilute compared with that of the fine-grained function, both LB and MB DFs, and hence density profiles, will look the same.

1.4. This Work

In this paper, we explore two possible approaches to deriving the final equilibrium state of self-gravitating systems, for each of the two types of statistics, LB and MB. In both, we use a very accurate, smooth approximation for \(\ln x\) valid for arbitrary occupation numbers \(x\). However, the price we pay for this improved approximation is the loss of analytic solutions.

The first approach assumes that the final state is the maximum entropy state, an assumption that was first used in the context of self-gravitating systems in 1950’s (Ogorodnikov 1957). The second approach, again in the context of self-gravitating systems, was first taken by Barnes & Williams (2011); it posits that the final state corresponds to the extremum of entropy production rate.

We explore extremizing entropy production because it has not been established beyond a doubt that real or computer-simulated systems do fully relax to maximum entropy states. It is possible that their steady-state configurations do not correspond to maximum entropy. Prior work on thermal non-equilibrium systems, but not in astrophysical contexts, suggests that stationary states—like mechanical equilibrium—occur when entropy production is extremized (e.g., Prigogine 1961; Jaynes 1980; de Groot & Mazur 1984; Grandy 2008). We investigate their applicability to self-gravitating systems in mechanical equilibrium.

The new aspect in the present paper is that we use a very accurate approximation for \(\ln x\), unlike our previous paper that assumed the Stirling approximation. As Madsen (1996) has shown, replacing Stirling with an accurate approximation (1) results in systems with finite total mass and energy, and (2) significantly changes the structure of the systems.

In all, we present four derivations; entropy maximizations for the LB and MB statistics are covered in Section 3.1 and Appendix A.1. Extremizations of entropy production for LB and MB statistics are carried out in Section 3.2 and Appendix A.2, respectively. We develop expressions for the relaxation functions (see Section 2), and use these to better understand the evolution of coarse-grained DF. We compare our results with analogous versions of entropy production derived using the standard Stirling approximation in Barnes & Williams (2011).
The set-up: statistical representations

| (x, v) phase-space | Energy state-space |
|--------------------|-------------------|
| no exclusion       | no exclusion      |
| MB; collisional    | MB; collisionless |
| no exclusion       | with exclusion    |
| LB; collisionless  |                  |
| non-deg.           | degen.            |
| same in non-degen. limit | not relevant |

Attaining final state: maximizing entropy

| (x, v) phase-space | Energy state-space |
|--------------------|-------------------|
| no exclusion       | no exclusion      |
| MB; collisional    | MB; collisionless |
| Stirling           | Stirling          |
| no Stirl.          | no Stirl.         |
| LB67 (M = ∞)       | LB67              |
| M96 BWII           | BWII              |
| discr. cont.       | discr. cont.      |
| in the non-degen. limit | not relevant |

Attaining final state: extremizing entropy production

| (x, v) phase-space | Energy state-space |
|--------------------|-------------------|
| no exclusion       | no exclusion      |
| MB; collisional    | MB; collisionless |
| Stirling           | Stirling          |
| no Stirl.          | no Stirl.         |
| BW1                | BW1               |
| discr. cont.       | discr. cont.      |
| BW1                | BW1               |
| discr. cont.       | discr. cont.      |
| not relevant       |                   |

Figure 1. Schematic summary of the various statistical mechanical approaches to self-gravitating systems. See Section 1.4 for explanation.

Figure 1 puts the present paper (BWII in the figure) in context. It is a schematic summary of the various statistical mechanical approaches to self-gravitating systems. The possible ways to frame the problem appear at the top of the figure; one can formulate the problem in either the regular phase space or the energy space. Below the thick horizontal line we show the two different routes for attaining the final steady state: maximizing entropy and extremizing entropy production. Once these choices are made one has to decide whether small occupation number regime will be important or not, and hence whether to use the Stirling approximation for \( \ln x! \) or not. In the latter case, one must then decide whether to use the discrete (“discr.”) step-like, i.e., exact version of \( \ln x! \), or to approximate it with some smooth function (“cont.”) that remains very accurate down to small \( x \). Note that HW10 and the present paper use different but similar approximations. There is no physical reason to introduce the exclusion principle in the energy state space, hence the corresponding regions are marked as “not relevant.” The bottom entries of some columns in the table contain names of papers where the corresponding options were considered. K66 in parentheses below BWII means that King (1966) results are nearly identical to ours. \( M = \infty \) under LB67 means that Lynden-Bell (1967) final result, the isothermal sphere, had infinite mass.

Much of the background material for this work may be found in Barnes & Williams (2011), and we briefly summarize these previously obtained results in Section 2.

2. SUMMARY OF RESULTS FOR LARGE OCCUPATION NUMBERS

Barnes & Williams (2011) have investigated entropy production in self-gravitating systems described by MB and LB and have developed expressions for the entropy production, \( \sigma \) for both the MB,

\[
\sigma_{MB} = -\frac{k_B}{\sigma \eta} \int \Gamma \left[ \ln \left( \frac{f}{\eta} \right) + 1 - \ln N \right] dv, \tag{4}
\]

and LB cases,

\[
\sigma_{LB} = -\frac{k_B}{\sigma \eta} \int \Gamma \left[ \ln \left( \frac{f}{\eta - f} \right) - C \right] dv, \tag{5}
\]

where constant \( C \) in Equation (5) is \( (\ln N-1)+(1/N) \sum_i \nu \ln \nu \). In these expressions, \( f \) is the coarse-grained DF and \( \Gamma \) is the relaxation function and forms the right-hand side of the Boltzmann equation,

\[
\frac{df}{dt} + v \cdot \nabla f + a \cdot \nabla_v f = \Gamma(f). \tag{6}
\]

If one were to assume, for a moment, that in the above equation \( f \) is a fine-grained DF, then the right-hand side would be zero for collisionless systems. In other words \( \Gamma = 0 \), which means that on fine-grained scales there is no change in, or production
of entropy; in collisionless systems, entropy is fixed throughout evolution. Let us be clear, we are not advocating for a specific process as a source for $\Gamma$, like collisions in a gas. The relaxation function simply describes the Lagrangian time rate of change of the DF. Returning to our case where $f$ represents the coarse-grained DF, Equation (6) states that entropy is produced (and as we argue in Barnes & Williams 2011, it happens even in systems that have attained macroscopic steady state) because on microscopic scales the fine-grained DF continues to wind and twist, which, when combined with coarse-graining, gives rise to non-zero entropy change. From a different starting point, Chavanis (1998) develops an expression for the right-hand side of Equation (6) in terms of a “diffusion current” that relates to correlations between fluctuations in the fine-grained DF. While that work details the makeup of this diffusion current, we simply focus on the broad behavior of the relaxation function.

We find extremum entropy production conditions by setting the variation of entropy production, $\delta \sigma$, equal to zero. This operation gives expressions for the relaxation function. For the MB case,

$$\Gamma_{MB}(f) = \frac{(1 - \ln N)\Gamma_{MB}(f = \eta)}{\ln(f/\eta) + 1 - \ln N}. \quad (7)$$

The LB relaxation function is slightly more complex,

$$\Gamma_{LB}(f) = -\frac{C\Gamma_{LB}(f = \eta/2)}{\ln[f/(\eta - f)] - C}. \quad (8)$$

Like Lynden-Bell (1967), the Barnes & Williams (2011) work assumes that the large $n$ Stirling approximation is valid for the systems being investigated. Here, we will be deriving relations analogous to Equations (4), (5), (7), and (8), but using a very accurate approximation, after discussing the results of entropy maximization below.

3. RESULTS FOR ARBITRARY OCCUPATION NUMBERS

The approximation that we utilize in this work is

$$\ln x! = \left(x + \frac{1}{2}\right)\ln(x + 1) - x + \frac{\ln 2\pi}{2} + \lambda_{0,x}. \quad (9)$$

where

$$\lambda_{0,x} = -\frac{x^2 + 2x + \frac{287}{12}}{x^2 + \frac{25}{12}x + \frac{13}{12}}. \quad (10)$$

In the large $x$ limit, this reduces to the usual Stirling approximation $\ln x! = x\ln x - x$. Our approximation is slightly different from the one in Hjorth & Williams (2010). Comparisons between the Stirling approximation, the Hjorth & Williams (2010) approximation, and Equation (9) are shown in Figure 2. The plots illustrate the function $\psi(x + 1) \equiv d\ln x!/dx$ for the three cases. The HW10 and Equation (9) approximations are nearly identical. Most importantly, these two approximations are well behaved at $x = 0$, unlike the Stirling approximation.

3.1. Entropy Maximization

In this section, we follow the overall path taken in Lynden-Bell (1967) to determine the description of entropy for the LB statistics family. Briefly, counting arguments for phase-space macro-cell occupation are combined with the statistical
definition of entropy to give specific representations. Variations in entropy assuming constant mass and energy are then set to zero in order to determine the necessary DFs.

If one demands that multiple phase-space elements cannot simultaneously occupy micro-cells, the multiplicity of states (Equation (3)) combined with the definition of entropy results in,

$$S_{LB} = k_B \left[ \ln N! - \sum_i \ln n_i! + \sum_i \ln v! - \sum_i \ln (v - n_i)! \right],$$

(11)

where again the summations run over the number of macro-cells. We will assume that both $N$ and $v$ are much larger than 1, so that Stirling’s approximation is valid for use in the first and third terms. However, for the $n_i$ and $v - n_i$ terms we will use our improved approximation,

$$\ln n_i! = (n_i + 1/2) \ln (n_i + 1) - n_i + \frac{\ln 2\pi}{2} + \lambda_{0,n_i}.$$  

(12)

Note that this usage does not require either $n_i$ or $v - n_i$ to actually be a small number, rather this keeps the accounting accurate in the event that they do become small. If these values are always large, there will be no difference from the Stirling approximation. The entropy expression now reads,

$$S_{LB} = S_{LB,0} - k_B \sum_i [(n_i + 1/2) \ln (n_i + 1) + (v - n_i + 1/2)$$

$$\times \ln (v - n_i + 1) + \lambda_{0,n_i} + \lambda_{0,(v-n_i)}],$$

(13)

where $S_{LB,0} = k_B [N \ln N - N + M (v \ln v - \ln 2\pi)]$ and $M$ is the total number of macro-cells.

Transforming from the discrete macro-cell occupation number $n_i$ to the continuous coarse-grained distribution function $f$, the entropy becomes

$$S_{LB} = S_{LB,0} - \frac{k_B}{\nu \gamma} \int \left[ \frac{\nu f}{\eta} + \frac{1}{2} \ln \left( \frac{\nu f}{\eta} + 1 \right) 

+ \left( 1 - \frac{\nu f}{\eta} + \frac{1}{2} \right) \ln \left( 1 - \frac{\nu f}{\eta} + 1 \right)

+ \lambda_{0,vf/\eta} + \lambda_{0,v-vf/\eta} \right] dx dv. $$

(14)

Taking the variation of this entropy expression to be zero, with constant mass and energy constraints, leads to the following condition:

$$\ln (F + 1) + \frac{(F + 1/2)}{(F + 1)} - \ln (v - F + 1) - \frac{(v - F + 1/2)}{(v - F + 1)}$$

$$+ \frac{d\lambda_{0,F}}{dF} + \frac{d\lambda_{0,v-F}}{dF} + \mu + \beta \epsilon = 0,$$  

(15)

where $F = \nu f/\eta$ is a scaled coarse-grained DF and $\mu$ and $\beta$ are undetermined multipliers associated with mass and energy conservation, respectively. The $\epsilon$ term is the specific energy of a phase element located at position $x$ with velocity $v$, $\epsilon = v^2/2 + \Phi$. The derivative of the $\lambda$ function is

$$\frac{d\lambda_{0,F}}{dF} = -\frac{(F + 1)}{(F + 600/576)}.$$  

(16)

After substituting for these $\lambda$ derivatives and combining terms we have

$$\ln \left[ \frac{F + 1}{v - F + 1} \right] + \frac{\nu f - (v - F + 1)}{(F + 1)(v - F + 1)}$$

$$- \frac{2F^2 - 2Fv - (1 + 600/576)v - 600/288}{F^2 - Fv - (600/576)v + (600/576)^2}$$

$$+ \mu + \beta \epsilon = 0.$$  

(17)

The inelegant ratio terms in this expression are both symmetric about $F = v/2$. The second term on the left-hand side of Equation (17) has values of $-1/2$ when $F = 0$, $0$ when $F = v/2$, and $1/2$ when $F = v$ (the maximum value of $F$ for the LB case). The third term on the left-hand side of Equation (17) is a nearly constant function with a value very close to $-2$ for $0 \leq F \leq v$.

We have not attempted to find an analytic solution for $F$. Graphical solutions of Equation (17) for a series of $\epsilon$ values produce the picture of $f_\eta/\nu$ seen in Figure 3. The overall character of the DF is the same in the non-Stirling and Stirling versions, and both functions are very similar to FD distribution,

$$f_{FD} = \frac{\exp(\mu + \beta \epsilon)}{1 + \exp(-\mu + \beta \epsilon)}.$$  

(18)

This DF of Equation (17) can be transformed into a density distribution using the Poisson equation and the fact that

$$\rho(r) = \int f(r,v)dv = 4\pi \int f(\nu)\sqrt{2|\nu - \Phi(r)|}d\nu.$$  

(19)

Note that this procedure imposes an isotropic velocity distribution for the system.

The density and logarithmic density slope corresponding to the non-Stirling function are presented in Figures 4 and 5, and will be discussed further in Section 4.1.

3.2. Entropy Production Extremization

Just because it is possible to describe thermal equilibrium states for collisionless self-gravitating systems, does it not follow that real systems (either physical or simulated) must achieve them in a Hubble time. It is possible that real systems incompletely relax, leaving them in a thermal non-equilibrium, but long-lived stationary state. Madsen (1996) also points out that in very slowly evolving systems, the maximization of entropy is only temporary. This leads us to infer that the production of entropy may be a useful quantity for discussing quasi-equilibria of collisionless systems. With that in mind, we now proceed to develop the conditions required for a thermal non-equilibrium state to be stationary. Specifically, we find an expression for entropy production in the LB statistical family and then extremize it. (As for the preceding section, an analogous derivation for the MB statistics may be found in Appendix A.2.) As discussed in Section 1.4, existing work on thermal non-equilibrium in non-astrophysical settings suggests that stationary states occur when entropy production is either maximum or minimum (de Groot & Mazur 1984; Grandy 2008, and references therein).

From Equation (14) the entropy density in the LB case can be written as

$$\rho_{SLB} = -\frac{k_B}{\nu \gamma} \int \left[ (F + 1/2) \ln (F + 1) - (v - F + 1/2)$$

$$\times \ln (v - F + 1) + \lambda_{0,F} + \lambda_{0,v-F} - F \frac{S_{LB,0}}{Nk_B} \right] dv,$$  

(20)

where, as before, $\nu = \nu f/\eta$. 

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Figure 3. Lynden-Bell distribution function resulting from the non-Stirling approximation. The Lynden-Bell function derived using the Stirling approximation is also shown.

(A color version of this figure is available in the online journal.)

Figure 4. (a) Logarithmic density and (b) the corresponding $\alpha = -d \log \rho/d \log r$ vs. logarithmic-scaled radius for the Lynden-Bell case. The thick solid black lines correspond to models with $\nu = 100$ while the thick dashed black lines represent models with $\nu = 1000$. The vertical lines mark where the logarithmic slope of the $\nu = 100$ model is isothermal ($\alpha = 2$). The $\alpha$ profiles for NFW, Navarro et al. (2004), and Plummer models are also shown. The profiles for the $\nu = 1000$ model have been horizontally shifted so that the location where $\alpha = 2$ coincides with the other models. The central cusp of the NFW and Navarro et al. (2004) models ($\alpha \to 1$ as $r \to 0$) differs markedly from the core present in the Lynden-Bell case ($\alpha \to 0$ as $r \to 0$). The presence of the core in the Lynden-Bell case is independent of $\nu$.

(A color version of this figure is available in the online journal.)
Equation (21), we get a lengthy expression, where
\[ \gamma = \nu \Gamma / \eta. \]

The acceleration-dependent terms in Equation (22) all disappear under the assumptions that the DF is even in \( \nu \) and disappears when \( \nu = v_{\text{max}} \). The final representation of the time rate of change of the entropy density is
\[ \frac{\partial}{\partial t} (\rho_{\text{SLB}}) = \frac{k_B}{\nu \sigma} \int [(F + 1/2) \ln (F + 1) - (\nu - F + 1/2) \ln (\nu - F + 1) + (\nu - F + 1/2) \ln (\nu - F + 1)] dv \]
\[ + \nu - F + 1/2 - C dF \]

and those in the continuous version of the entropy density evolution equation (Equation 6 in Barnes & Williams 2011),
\[ \frac{\partial}{\partial t} (\rho s) = - \nabla \cdot (\Sigma + \rho s v_0) + \sigma. \]
The entropy flux due to random motions $\Sigma$ is given by the integral in the second term on the right-hand side of Equation (24). The remaining term then makes up the entropy production for the system,

$$
\sigma_{\text{LB}} = -\frac{k_B}{v\sigma} \int \left[ \ln (F + 1) + \frac{F + 1/2}{F + 1} - \ln (\nu - F + 1) + \frac{\nu - F + 1/2}{\nu - F + 1} + d\lambda_{0,F} \frac{dF}{F} + d\lambda_{0,\nu-F} \frac{dF}{F} - C \right] dv.
$$

(26)

As expected, the relaxation function determines the entropy production rate for the system.

We find that the condition for the entropy production term in Equation (26) to be extremized is

$$
\Gamma_{\text{LB}}(f) = Q \left[ \ln \left( \frac{F + 1}{\nu - F + 1} \right) + \frac{F + 1/2}{F + 1} + \frac{\nu - F + 1/2}{\nu - F + 1} + \frac{d\lambda_{0,F}}{dF} + \frac{\nu - F + 1/2}{\nu - F + 1} - C \right],
$$

(27)

where the integration constant is defined by

$$
Q = \gamma_{\text{LB}}(F = \nu/2) \left[ \frac{\nu + 1}{\nu/2 + 1} - \frac{2(\nu + 2)}{\nu + 600/288} - C \right],
$$

(28)

and the $\lambda$ derivatives are the same as those used in going from Equations (15) to (17). This relaxation function is shown in Figure 6 and is very similar to the corresponding function from Barnes & Williams (2011), their Figure 2.

4. SUMMARY AND DISCUSSION

In an attempt to better understand the evolution of self-gravitating collisionless systems, we have re-investigated a standard statistical mechanics approach to finding equilibria (entropy maximization) and continued to develop a new approach (extremization of entropy production), first applied to self-gravitating systems by Barnes & Williams (2011). Entropy production in non-equilibrium steady-state systems has been previously investigated and found to be useful in several non-astrophysical systems, and so it is interesting to ask if the principle is relevant in gravitational systems.

Both of our approaches use a very accurate approximation for $\ln n!$, and not the standard Stirling formula valid only for large occupation numbers. It has been recently shown that using an approximation that reflects correct behavior for small $n$, and the principle of entropy maximization, leads to density distributions that resemble globular clusters (King profiles; Madsen 1996), and simulated pure dark matter halos (Williams & Hjorth 2010; Williams et al. 2010). Both types of systems have finite mass and energy, in compliance with the entropy maximization constraints.

4.1. Results of Entropy Maximization

Entropy maximization using the LB and MB statistics produces the density profiles and logarithmic profiles slopes, $\alpha = -d \log \rho / d \log r$, depicted in Figures 4 and 7, respectively.

We argue that the LB case accurately represents collisionless systems because co-habitation of the phase-space elements is prohibited. However, the density profiles for any value of $\nu$ are very different from the two cosmological models, Navarro-Frenk-White (NFW; Navarro et al. 1996) and N04 (Navarro et al. 2004), which are fits to the results of cosmological N-body simulations. In contrast to the latter, the LB density profiles have a flat central density slope, while at larger radii $\alpha$ has a steeper rise than that of the Plummer profile (Plummer 1911; Binney & Tremaine 1987), which is an example of a polytropic system. In Figure 5, we compare the profiles to those derived in Appendix A.1, which are nearly identical to King models: LB profiles are poor matches to King models. (The vertical lines in each panel mark the radial position where $\alpha = 2$.) Both the MB and LB density profiles are also different from those found in Hjorth & Williams (2010), as those models can have central density cusps.

We conclude that applying entropy maximization to the LB case coupled with a very accurate approximation for $\ln n!$ produces (isotropic) density profiles that are distinct from any other first-principles function, or fits to the results of N-body simulations. We will return to this point in Section 4.3.

The MB case is intended to represent collisional systems because the co-habitation of the phase-space elements is allowed. The density profiles with $\nu = 100$ and 1000 are given in Figure 7(a). The $r_{\text{max}}$ scaling distance corresponds to the radius where a non-moving particle has an energy for which the DF disappears. The second panel of that Figure 7(b) shows the slope of the logarithmic density profile $\alpha$ along with curves corresponding to three other well known analytical density profiles. These MB solutions are basically identical to King models (see Figure 8). This result comes as no surprise in the light of Madsen (1996), who did not use any approximation for $\ln x!$, and obtained density profiles very similar to King’s. Abandoning the Stirling approximation (and treating the low occupation number limit with the respect it apparently deserves) demonstrates that the King DF is the result of maximizing entropy in MB statistics, under the conditions of fixed mass and energy.

The consequence of using a very good smooth approximation to $\ln x!$, as done here, instead of the exact expression that gives discrete step-like values, as was done in Madsen (1996), is that the former results in spot-on matches to the King profiles, while the latter display modest differences from the King profile shape, especially for low $\Psi(0)/\sigma^2$ values; see Figure 1 of Madsen (1996).

The value of $\nu$ appears to play a role analogous to the King scaling factor $\Psi(0)/\sigma^2$, where $\Psi(0)$ is a relative potential energy at the center, and $\sigma$ is an energy scaling constant (Binney & Tremaine 1987). Increasing $\nu$—or $\Psi(0)/\sigma^2$—increases the concentration of the system. Recall that the higher the concentration, the closer the King model is to an isothermal sphere.

We find that the concentrations of the King models that most closely resemble the MB density profiles increase as the value of $\nu$ increases. Recall that $\nu$ is the number of fine-grained microcells that occupy a coarse-grained macrocell, so as $\nu$ increases the DF becomes “grainer.” At the same time, increasing the concentration of a King model makes it more closely resemble an isothermal sphere. Putting these two observations together implies that increasing the coarseness of the DF (by increasing the number of fine-grained cells contained in a coarse-grained cell) should result in densities that have more isothermal aspects.

Finally, we note that both of the families of profiles obtained here have flat density cores, while cosmologically simulated halos have density cusps.
4.2. Results of Entropy Production Extremization

Some aspect of simulations may delay, or even disallow, the maximizing of entropy necessary to achieve thermal equilibrium. In such a frustrated case, it is possible that a collisionless system finds a stationary state by extremizing its entropy production. As a more concrete example of this situation, imagine a metal bar with one end exposed to a blowtorch and the other end in an ice bath. When the bar reaches a stationary state with a time-independent, but spatially varying, temperature distribution, entropy production in the bar is an extremum (de Groot & Mazur 1984, Chapter 5, Section 3). We have begun to explore the possibility that the mechanical equilibria of simulated collisionless systems can be explained as states of extreme entropy production.

To this end, we have derived expressions for entropy production in collisionless systems using the LB statistics, and in collisional systems, using MB statistics. Further, we have found
Figure 7. (a) Logarithmic density and (b) the corresponding $\alpha = -d \log \rho / d \log r$ vs. logarithmic-scaled radius for the Maxwell–Boltzmann case. The thick solid black lines represent models with $\nu = 100$ while the thick dashed black lines correspond to models with $\nu = 1000$. The vertical lines mark where the logarithmic slope of the $\nu = 100$ model is isothermal ($\alpha = 2$). The $\alpha$ profiles for NFW, Navarro et al. (2004), and Plummer models are also shown. As in Figure 4, the $\nu = 1000$ curve has been shifted horizontally to align the locations where $\alpha = 2$. The central cusp of the NFW and Navarro et al. (2004) models ($\alpha \to 1$ as $r \to 0$) differs markedly from the core present in the Maxwell–Boltzmann case ($\alpha \to 0$ as $r \to 0$). The presence of the core in the Maxwell–Boltzmann case is independent of $\nu$.

(A color version of this figure is available in the online journal.)

Figure 8. Comparisons between Maxwell–Boltzmann (solid) and King (dashed) density profiles for different $\nu$ values: (a) $\nu = 100$, (b) $\nu = 1000$, (c) $\nu = 5000$. The King profile scale factor $\Psi(0)/\sigma^2$ is given in each panel. The dashed lines represent King density profiles while the solid lines show the Maxwell–Boltzmann distributions. The comparison between these two profile types reveals more similarity than when the Madsen (1996) and King models are compared.

(A color version of this figure is available in the online journal.)
the form of the relaxation function required to guarantee an entropy production extreme. Recall that the relaxation function is the right-hand side of the Boltzmann equation, Equation (6), that describes the coarse-grained DF evolution.

As in our previous work which was valid only for large phase-space occupation numbers, the MB relaxation function (pictured in Figure 9) is positive for all values of \( F \) (as long as \( N > 1 \)) and becomes more constant as the number of particles \( N \) increases. It is clear that the basic form of the relaxation function is not dependent on the specifics of the DF. A visual comparison between Figure 6 and Figure 1 in Barnes & Williams (2011) reinforces this point.

The general form of the LB relaxation function shown in Figure 6 has similarities and differences with its large occupation number version derived in Barnes & Williams (2011). The similarities are that both the functions increase with \( f/\eta \) upto just short of \( f/\eta \approx 1 \), then spike toward large positive values of \( \Gamma_{\text{LB}} \), go through a singularity at \( f/\eta \approx 1 \), then become negative, and eventually asymptote to \( \Gamma_{\text{LB}} = 0 \) as \( f/\eta \to 1 \) (see the bottom panel of Figure 6). The difference is that the large occupation number \( \Gamma_{\text{LB}} \) is independent of \( \nu \), while its arbitrary occupation number analogue is not. In the latter case the values of \( C \) and \( \nu \) that produce the singularity are linked through the relationship,

\[
\log \frac{\nu_{\text{min}}}{100} = 0.4348(C - 4.1518),
\]

where \( \nu_{\text{min}} \) is the minimum value of \( \nu \) that will cause a singularity for a given \( C \). This expression is valid as long as \( \nu_{\text{min}} \gtrsim 10 \). The importance of the singularity in both cases is that it signals that the relaxation function will reach zero when the coarse-grained distribution function \( f \) reaches the fine-grained value \( \eta \), bringing the evolution of the DF to a halt.

To sum up, extremizing entropy production in the LB case leads to a relaxation function \( \Gamma_{\text{LB}} \) versus \( f/\eta \) shape that drives \( \Gamma_{\text{LB}} \), and hence the entropy production rate, \( \sigma_{\text{LB}} \), to zero, which in turn means that the endpoint of evolution has a maximum (more correctly, an extremum) entropy. In other words, the entropy production extremization procedure that we followed in Section 3.2 tell us that the final state of a self-gravitating collisionless system is a state of maximum entropy. Since in Section 3.1 we derived such a state, using entropy maximization, it must be the same state.

The important point in both the LB and the MB case is that extremizing entropy production leads to relaxation functions that drive a coarse-grained DF to behave like a fine-grained DF.

To test the hypothesis that for self-gravitating systems the state of entropy production extreme coincides with the state of maximum entropy, we are undertaking a comparison between these analytical descriptions of entropy behavior in collisionless systems and results of \( N \)-body and semi-analytical simulations (e.g., the extended secondary infall model, ESIM; Williams et al. 2004; Austin et al. 2005). This further work will help settle exactly what role entropy production plays in determining collisionless equilibria.
4.3. Outstanding Questions

Aside from the future work described above, to quantify entropy production rate in simulated numerical systems, there are also a few questions that remain unanswered.

4.3.1. Maximizing Entropy in Phase Space versus Energy Space

Hjorth & Williams (2010) and the present paper maximized entropy, but in different state spaces, energy and (x, v) phase space, respectively. The results of the two studies, for example in terms of the density profiles, are very different from each other. It is not surprising that they are different, but it is not immediately obvious which one of the two approaches would produce a better description of the results of collisionless N-body simulations. After looking at all the density profiles it is seen that Hjorth & Williams (2010) profiles are similar to those of simulated systems (Williams & Hjorth 2010; Williams et al. 2010), while those from the LB case in the present work (Section 3.1, Figures 4 and 5) are not.

We note that our LB entropy maximization results are what Lynden-Bell (1967) would have obtained had he not used the Stirling approximation. The procedure does not rely on any approximations and is well motivated from first principles, assuming that all states are equally accessible and efficient mixing in phase space can be achieved. This beg the question, do such systems exist? More work with simulations is needed to address this question.

4.3.2. The Necessity of the Low Occupation Number Regime

Our MB entropy maximization results of Appendix A.1 (and the very similar results of Madsen 1996) describe some globular clusters well (Elson et al. 1987; Meylan & Heggie 1997; L. L. R. Williams et al. 2012, in preparation), and the results of Hjorth & Williams (2010) describe N-body and ESIM halos. These facts lead us to the following chain of reason. First, globular clusters, along with N-body and ESIM halos, are self-gravitating systems. Second, the entropy maximization procedures that produce successful models of these systems require correct treatment of low phase-space occupation numbers. Therefore, it appears that the low occupation number regime is an important distinction between self-gravitating and non-self-gravitating systems.

Why is this the case? We speculate that self-gravitating systems are special not only because they are finite in extent due to the long-range nature of the force, but also because of correlations between spatial regions and energies of particles. Specifically, the center of the potential contains phase-space elements with the most bound energies, while the outskirts are populated by elements with the least bound or even zero energies. These requirements are peculiar to self-gravitating systems and apparently necessitate low occupation numbers because the numbers of particles in these regions must approach zero.

However, the above hypothesis does not explain why the discrete version of ln x! must be smoothed to represent real systems. This smoothing, though apparently necessary, is still not well justified.

4.3.3. Should Maximum Entropy Imply Constant Temperature?

The steady state of self-gravitating systems can apparently be calculated as the maximum entropy state. Though these systems are in mechanical equilibrium, one could argue that they are not in conventional thermal equilibrium because their kinetic temperature is manifestly different across the system. Can we reconcile maximum entropy defining thermodynamic equilibrium with the resulting non-constant temperature? (Note that a realistic self-gravitating system cannot have a constant kinetic temperature, because there is no finite-mass solution to the Jeans equation with constant σ.) This contradiction is part of the reason why it made sense to conclude, based on Lynden-Bell (1967) work, that there is no maximum entropy state for self-gravitating systems.

We propose that for gravitationally bound collisionless systems, one must carefully separate kinetic temperature T_k from thermodynamic temperature T_r. To be clear, the kinetic temperature we are referring to is related to the peculiar velocity in a system, T_k = β(v_p)^2. The thermodynamic temperature is linked to the energy scale β that serves as a Lagrange multiplier through β = 1/(k_B T_r). By definition, T_r must be a constant, and there is no contradiction as in Lynden-Bell (1967)—systems in the maximum entropy state are characterized by a constant thermodynamic temperature, even as they have a varying kinetic temperature.

We can also address a related question involving the temperature and energy change that appear in the thermodynamic definition of entropy, dS = δQ/T. This temperature must be a constant, so we would associate this with the thermodynamic temperature T_r. δQ (normally called heat) refers to all non-work exchanges of energy, and must be replaced by δε because in collisionless self-gravitating systems what is being transferred is total energy, potential and kinetic, not just kinetic. Entropy changes can occur due to changes in mass distribution as well as kinetic energy transport.

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APPENDIX

MAXWELL–BOLTZMANN STATISTICS

A.1. Entropy Maximization

Using the results of Section 1.3, the multiplicity function for a system obeying MB statistics combined with the definition of entropy (Equation (1)) produces

\[ S_{MB} = k_B \left[ \ln N! - \sum_i \ln n_i! + N \ln v \right]. \]  \hspace{1cm} (A1)

where the summation involving the macro-cell occupation n_i runs over the number of macro-cells. We will assume that N >> 1, so that ln N! = N ln N - N. However, we will not use the Stirling approximation for the second term on the right-hand side of Equation (A1). With our approximation, Equation (12), we can now rewrite the entropy as

\[ S_{MB} = S_{MB,0} - k_B \sum_i \left( (n_i + 1/2) \ln (n_i + 1) + \lambda_{0,n_i} \right), \]  \hspace{1cm} (A2)

where S_{MB,0} = Nk_B[ln(Nv) - M ln 2\pi/2N] is a constant and M is the total number of macro-cells.
We now replace the discrete macro-cell occupation number \( n_i \) with the coarse-grained distribution function \( f \) to produce

\[
S_{\text{MB}} = S_{\text{MB},0} - \frac{k_B}{v \sigma \tau} \int \int \left[ \frac{(v f/\eta) + 1/2}{v} \ln \left( \frac{v f/\eta + 1}{v} \right) + \frac{\lambda_{0,F}}{v} \frac{\partial \lambda_{0,F}}{\partial f} + \mu \right] d x \ d v.
\]

(A3)

To maximize this entropy function, we set \( \delta S_{\text{MB}} = 0 \), subject to constant mass and energy constraints. The expression that results is

\[
\ln \left( \frac{v f/\eta + 1}{v} \right) + \frac{(v f/\eta + 1/2)}{(v f/\eta + 1)} + \ln \left( \frac{v f/\eta + 1}{v} \right) + \frac{\lambda_{0,F}}{v} \frac{\partial \lambda_{0,F}}{\partial f} + \mu + \beta \epsilon = 0,
\]

(A4)

where \( \mu \) and \( \beta \) are undetermined multipliers associated with mass and energy conservation, respectively. The \( \epsilon \) term is the specific energy of a phase element located at position \( x \) with velocity \( v \), \( \epsilon = v^2/2 + \Phi \). Note that if the standard Stirling approximation had been employed, the second and third terms would be absent and the logarithmic term would simply read \( \ln (v f/\eta) \). In this case, the usual, physically inconsistent, MB distribution function would result.

As before, we make a change of variables, \( F \equiv v f/\eta \), and the derivative of the \( \lambda \) function is given by Equation (16). Equation (A4) can then be re-cast as

\[
\ln (F + 1) - \frac{(264F + 276)}{576(F + 1)(F + 600/576)} + \mu + \beta \epsilon = 0.
\]

(A5)

Again, we have not searched for an analytical solution, but graphical solutions for \( F \) for a range of \( \epsilon \) values can be combined to produce a plot of \( F(\epsilon) \). Specifically, Figure 10 illustrates the behavior of the normalized coarse-grained distribution function \( f/\eta \) for a particular value of \( v \). Values of the Lagrange multipliers \( \mu \) and \( \beta \) were adjusted so that \( \epsilon \) is always positive. Because MB statistics have no exclusion principle, \( f/\eta \) can have values above 1.

A.2. Entropy Production Extremization

We begin by writing entropy in terms of entropy density,

\[
S = \int \rho s \ dx,
\]

(A6)

where \( \rho \) is mass density, \( s \) is the specific entropy, and the integral is taken over the spatial extent of the system. From the entropy form given in Equation (A3), the entropy density can now be written as

\[
\rho s_{\text{MB}} = -\frac{k_B}{v \sigma \tau} \int \left[ (F + 1/2) \ln (F + 1) + \lambda_{0,F} F \right] \left[ \frac{S_{\text{MB},0}}{N k_B} \right] d v.
\]

(A7)

where, as earlier, \( F = v f/\eta \).

Taking a partial time derivative of Equation (A7) results in

\[
\frac{\partial}{\partial t} (\rho s_{\text{MB}}) = -\frac{k_B}{v \sigma \tau} \int \left[ \ln (F + 1) + \frac{(F + 1/2)}{(F + 1)} + \frac{\partial \lambda_{0,F}}{\partial F} - B \right] d v,
\]

(A8)

where \( B = S_{\text{MB},0}/N k_B \) is a constant.

Substituting \( \partial F/\partial t \) from the Boltzmann equation into Equation (A8) results in a lengthy expression, similar to Equation (22) in the LB case. We will deal with this expression
term by term. For reference, the expression after substitution is
\[
\begin{align*}
\frac{\partial}{\partial t} (\rho s_{\text{MB}}) &= -k_B\frac{v}{\sigma T} \int (-v \cdot \nabla F) \ln (F + 1) + (F + 1/2) \frac{(F + 1)}{(F + 1)} \\
&\quad + \frac{\partial \lambda_{0,F}}{\partial F} - B \int v \cdot \nabla v F \\int (F + 1) \\
&\quad + \frac{\partial \lambda_{0,F}}{\partial F} - B \int v \cdot \nabla v F \\int (F + 1) \\
&\quad + \frac{\partial \lambda_{0,F}}{\partial F} - B \int v \cdot \nabla v F \\int (F + 1) \\
&\quad + \frac{\partial \lambda_{0,F}}{\partial F} - B \int v \cdot \nabla v F \\int (F + 1) \\
&\quad + \frac{\partial \lambda_{0,F}}{\partial F} - B \int v \cdot \nabla v F \\int (F + 1) \\
&\quad + \frac{\partial \lambda_{0,F}}{\partial F} - B \int v \cdot \nabla v F \\int (F + 1) \\
&\quad + \frac{\partial \lambda_{0,F}}{\partial F} - B \int v \cdot \nabla v F \\int (F + 1) \\
&\quad + \frac{\partial \lambda_{0,F}}{\partial F} - B \int v \cdot \nabla v F \\int (F + 1) \\
\end{align*}
\]
where \( \gamma = (v\Gamma)/\eta \).

The process to evaluate these integrals is very similar to what has been discussed in the LB case. Using the fact that \( v \cdot \nabla F = \nabla \cdot Fv \), the first integral on the right-hand side of Equation (A9) can be re-written as
\[
-\nabla \cdot \int [(F + 1/2) \ln (F + 1) + \lambda_{0,F} - FB] v dv. \quad (A10)
\]
Note that this expression is reminiscent of the form of the entropy density in Equation (A7).

We next turn our attention to the terms involving acceleration in Equation (A9). The fact that \( a \) is velocity independent implies that \( a \cdot \nabla v = \nabla v \cdot a \), a fact that will be used often in dealing with these terms. Let us start with the first part of the second term on the right-hand side of Equation (A9),
\[
\int (-a \cdot \nabla v, F) \ln (F + 1) \, dv = -\int (\nabla v \cdot a F) \ln (F + 1) \, dv. \quad (A11)
\]
Integrating by parts produces two terms, one with the form,
\[
\nabla v \cdot [a F \ln (F + 1)] \, dv. \quad (A12)
\]
which equals zero after using the divergence theorem and the fact that any physical DF must vanish for large velocities. The other term resulting from the integration by parts is
\[
a \cdot \int (\nabla v F) F \frac{F}{F + 1} \, dv = a \cdot \int \nabla v [(F + 1) - \ln (F + 1)] \, dv. \quad (A13)
\]
A single component of this integral will appear as
\[
\int_{v_{1,\text{min}}}^{v_{1,\text{max}}} \frac{\partial}{\partial v_1} [(F + 1) - \ln (F + 1)] \, dv_1 \\
= (F + 1) - \ln (F + 1)|_{v_{1,\text{min}}^{v_{1,\text{max}}}}. \quad (A14)
\]
where \( v_{1,\text{min}} = -v_{1,\text{max}} \) representing the maximum speed possible for the system. Assuming that \( F \) is even in \( v_{1,\text{max}} \) and that \( F(v_{1,\text{max}}) = 0 \) so that the DF disappears at the maximum speed, this integration results in zero. Similar manipulations can be applied to the rest of the acceleration-dependent parts of the second term on the right-hand side of Equation (A9), resulting in the entire second term being equal to zero. Equation (A9) now has the form,
\[
\frac{\partial}{\partial t} (\rho s_{\text{MB}}) = -k_B\frac{v}{\sigma T} \int [-\nabla \cdot \int [(F + 1/2) \ln (F + 1) \\
+ \lambda_{0,F} - FB] v dv + \int \gamma (F) \left[ \ln (F + 1) \\
+ \frac{F + 1/2}{F + 1} + \frac{\partial \lambda_{0,F}}{\partial F} - B \right] dv]. \quad (A15)
\]
Assuming the velocity field to be composed of mean and peculiar components \( v = v_0 + v_p \), we can re-cast Equation (A15) as
\[
\frac{\partial}{\partial t} (\rho s_{\text{MB}}) = -\nabla \cdot (\rho s_{\text{MB}} v_0 - k_B\frac{v}{\sigma T} \int [-\nabla \cdot \int [(F + 1/2) \ln (F + 1) \\
+ \lambda_{0,F} - FB] v dv + \int \gamma (F) \left[ \ln (F + 1) \\
+ \frac{F + 1/2}{F + 1} + \frac{\partial \lambda_{0,F}}{\partial F} - B \right] dv]. \quad (A16)
\]
We now equate the terms in Equation (A16) to those in the continuous version of the entropy density evolution equation, Equation (25),
\[
\frac{\partial}{\partial t} (\rho s) = -\nabla \cdot (\Sigma + \rho s v_0 + \sigma). \quad (A17)
\]
The entropy flux \( \Sigma \) is given by the integral in the second term on the right-hand side of Equation (A16) and represents randomly fluxed entropy. The remaining term is the entropy production for the system,
\[
\sigma_{\text{MB}} = -k_B\frac{v}{\sigma T} \int \gamma (F) \left[ \ln (F + 1) + \frac{F + 1/2}{F + 1} \\
+ \frac{\partial \lambda_{0,F}}{\partial F} - B \right] dv. \quad (A18)
\]
This equation explicitly demonstrates how the non-collisionless nature of the coarse-grained DF leads to changes in entropy.

As mentioned in Section 1, thermodynamic non-equilibrium systems can have steady states described by extrema of entropy production. We then set \( \delta \rho_{\text{MB}} = 0 \). Taking the variation of Equation (A18), we find
\[
\delta \rho_{\text{MB}} = -k_B\frac{v}{\sigma T} \int \delta F \left[ \frac{d\gamma}{dF} \left[ \ln (F + 1) + \frac{F + 1/2}{F + 1} \\
+ \frac{\partial \lambda_{0,F}}{\partial F} - B \right] \\
+ \frac{2}{F + 1} - \frac{F + 1/2}{(F + 1)^2} + \frac{d^2 \lambda_{0,F}}{dF^2} \right] dv. \quad (A19)
\]
Since \( \delta F \) is arbitrary, the variation disappears only when the term in curly braces is zero. The condition for an extremum in entropy production is
\[
\frac{d\ln \gamma}{dF} \left[ \ln (F + 1) + \frac{F + 1/2}{F + 1} + \frac{\partial \lambda_{0,F}}{\partial F} - B \right] \\
+ \left[ \frac{2}{F + 1} - \frac{F + 1/2}{(F + 1)^2} + \frac{d^2 \lambda_{0,F}}{dF^2} \right] = 0. \quad (A20)
\]
The solution for the MB relaxation function is
\[
\gamma_{\text{MB}}(F) = P \left[ \ln (F + 1) + \frac{F + 1/2}{F + 1} + \frac{\partial \lambda_{0,F}}{\partial F} - \frac{S_{\text{MB},0}}{Nk_B} \right], \quad (A21)
\]
where the constant can be expressed as
\[
P = \gamma_{\text{MB}}(F = v) \left[ \ln (v + 1) + \frac{v + 1/2}{v + 1} \\
- \frac{v + 1}{v + 600/576} - \frac{S_{\text{MB},0}}{Nk_B} \right]. \quad (A22)
\]
The $\gamma_{MB}(F = \nu)$ term is the relaxation function value when the coarse-grained DF is equal to the constant fine-grained DF value ($f = \eta$).

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