The maximum entropy techniques and the statistical description of systems

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Abstract

The maximum entropy technique (MENT) is used to determine the distribution functions of physical values. MENT naturally combines required maximum entropy, the properties of a system and connection conditions in the form of restrictions imposed on the system. It can, therefore, be employed to statistically describe closed and open systems. Examples in which MENT is used to describe equilibrium and non-equilibrium states, as well as steady states that are far from being in thermodynamic equilibrium, are discussed.

Statistical characteristics are widely used in high-energy physics to analyze experimental data \cite{1}. A simple analytical technique is employed by estimating the entropy of nuclear reaction products—a parameter which is sensitive to order-disorder transition, and provides evidence for the formation of ordered systems during a reaction \cite{2}.

Entropy methods for simulation of processes are being increasingly applied to various processes \cite{3}. An example of such an approach is the study of dynamic symmetry and internal chaos in models of interacting bosons \cite{4,5} conducted using the maximum entropy technique (MENT) \cite{6}.

MENT provides an efficient tool for solving inverse problems \cite{7}. Its application to statistical problems is promising. It is desirable, however, to test MEM for models that have already been proven to be reliable for statistical descriptions. In the present study, the applicability of MENT was tested by describing the equilibrium and non-equilibrium state of systems.

The goal of a statistical description of a system is to determine the distribution functions of physical values represented by an integral image in the results of observation. If distribution functions are assessed only on the basis of observation data on a system, then, from a mathematical point of view, an incorrect inverse problem arises in which the requirements of the existence, uniqueness and stability of a solution are not met. The condition of maximum entropy makes it possible to meet this requirement automatically. Only one solution of the problem is selected out of all. This solution has a maximum entropy, is most probable and, therefore, most stable. The resultant solution is obviously positive because the MENT estimate is in exponential form and agrees with one of the main properties of the distribution function — non-negativity. Other properties of the distribution function, e.g. normalization in MENT, can be preset as additional restrictions.

The theory of MENT is based on the second law of thermodynamics which describes the thermodynamic equilibrium of a closed system by maximum entropy. Attempts have been made to generalize this principle by arguing that entropy in internal dissipative
processes is maximum \[8\]. The requirement of maximum entropy, the property of a system and condition of connection in the form of restrictions placed on the system are naturally combined in MENT so that it can be used to statistically describe the equilibrium and non-equilibrium states of open and closed systems.

The mathematical structure of an entropy functional is related to the type of the quantum statistics of a physical data carrier and the extent of filling of quantum degrees of freedom \[7\]. The entropy functional

\[
H = - \sum_{\xi=1}^{N} f(\xi) \ln f(\xi),
\]

where \(f(\xi)\) is the distribution function estimate and \(\xi\) is an independent variable, is used for fermions and bosons with a low probability of filling of energy levels, such as photons of incoherent electromagnetic radiation, hadrons and \(\gamma\)-quanta. When the filling of energy levels is highly probable, a different form of a functional, like the one used in radioastronomy or spectral analysis \[9\], is selected:

\[
H = - \sum_{\xi=1}^{N} \ln f(\xi)
\]

We will use the most common form of an entropy functional (1).

1. **Boltzmann’s distribution**

Let us use MENT to determine the distribution of \(N\)-particles on the \(K\)-levels of energy \(E_i\), which has a maximum entropy, total energy \(E\) and the number of particles \(N\) being retained:

\[
\sum_{i=1}^{K} n_i \cdot E_i = E
\]

\[
\sum_{i=1}^{K} n_i = N
\]

Let us formulate the problem of estimating the probability \(f_i = \frac{n_i}{N}\) of finding a particle on level \(E_i\) as a variation problem on a conditional extremum with the Lagrange factors \(\lambda\) and \(\mu\) in the form:

\[
- \sum_{i=1}^{K} f_i \ln f_i + \lambda \left( NE - \sum_{i=1}^{K} f_i E_i \right) + \mu \left( 1 - \sum_{i=1}^{K} f_i \right) \to \text{max}.
\]

Let us equate the first variation of this relation (4) on \(f_i\) to zero. We will then have a relation:

\[
- \ln f_i - 1 - \mu - \lambda E_i = 0
\]

which gives an expression for probability \(f_i\):

\[
f_i = \frac{1}{Z} \exp(-\lambda E_i).
\]
This distribution function is identical to the Boltzman distribution function if factor $\lambda$ is understood as inverse temperature $\lambda = 1/kT$, and factor $\mu = \ln(Z/e)$.

The requirement of maximum entropy in a closed system with a constant number of particles and constant energy results in Boltzman distribution.

2. Non-equilibrium distribution

Let us use the MENT algorithm to describe the state of a two-level system (levels 1 and 2) which has magnetic moment $M$ and consists of $N$-particles with magnetic moment $m$. Let us consider that a magnetic field is absent, the system is not in equilibrium and the magnetic moments of the particles located on levels 1 and 2 have opposite directions.

Let $f(1)$ and $f(2)$ are the probabilities of finding a particle on levels 1 and 2, respectively. Equations for the complete probability and polarization $P$ of the system will be regarded as connection conditions:

\[ f(1) + f(2) = 1 \]  
\[ f(1) - f(2) = M/Nm = P. \]  

To estimate $f(1)$ and $f(2)$, we will use a conventional MENT scheme with a functional:

\[-f(1) \ln f(1) - f(2) \ln f(2) + \mu(1 - f(1) - f(2)) + \lambda(P + f(2) - f(1)) \rightarrow \max.\]  

The formulas

\[-\ln f(1) - 1 - \mu - \lambda = 0\]
\[-\ln f(2) - 1 - \mu + \lambda = 0\]

obtained from condition (9) give expressions for probabilities:

\[ f(1) = \exp(-1 - \mu - \lambda) \]
\[ f(2) = \exp(-1 - \mu + \lambda). \]

The substitution of these expressions in conditions (7) and (8) determines a relationship between Lagrange factors:

\[ \exp(-1 - \mu)2\text{ch}(\lambda) = 1 \]
\[ \exp(-1 - \mu)2\text{sh}(\lambda) = P. \]

Dividing the first equation by the second one, we will have expression:

\[ P = \text{th}(\lambda) \]

for the polarization $P$ of the system. Comparing it with an expression for the polarization of the equilibrium state of a two-level system with magnetic moment $M$ in magnetic field $H$ [10]:

\[ P = \text{th}(mH/kT) \]
we conclude that taking $\lambda = mH/kT$, we thus characterize a non-equilibrium state as quasi-equilibrium with magnetic field $H$ which agrees with given polarization $P$ or magnetic moment $M$:

$$H(P) = (kT/2m)\ln[(1 + P)/(1 - P)].$$

(15)

The result obtained implies that if there is a magnetic field which is switched off instantly, the thermodynamically equilibrium state of a two-level system becomes non-equilibrium and cannot change immediately. Therefore, at moments of time that are small in comparison with the relaxation time of the system this state is described as quasi-equilibrium, provided there is a magnetic field which decreases gently in magnitude.

3. Steady states of open systems

The steady states of open systems that are far from equilibrium are observed in many processes. Systems with autocatalytic chemical reactions were among the first to show characteristic types of behaviour of such systems, such as double stability, oscillation and wave generation interpreted by the mechanism of autocatalysis [8]. An example of such a reaction is Belousov-Zhabotinskys reaction with cerium sulphate $Ce_2(\text{SO}_4)_3$, malonic acid $CH_2(\text{COOH})_2$, potassium bromate $K\text{BrO}_3$ and ferroine that are vigorously mixed and supplied to reaction volume from which reaction products are simultaneously removed. Cerium, which catalyzes the reaction, contributes to the production of two molecules from one molecule of an intermediate product of $H\text{BrO}_2$:

$$H\text{BrO}_2 + \text{BrO}_3^- + 3H^+ + 2\text{Ce}^{3+} \rightarrow 2H\text{BrO}_2 + 2\text{Ce}^{4+} + H_2O.$$  

(16)

In this case, the catalyst is not spent, but its valency changes. Ferroine adds a red colour to the solution, if there is an excess of trivalent cerium ions, and a blue colour if there is an excess of quadrivalent cerium ions. If the time of staying of the reagents in the reaction zone is too short to adjust direct and inverse reaction rates, then such phenomena as bistability and chemical hours (periodic alternation of bistable states with an excess of tri- or quadrivalent cerium ions) occur in the system. The colour of the reaction solution varies with a period of several minutes. If the reagents are not mixed, waves are generated and travel.

To exemplify the steady states of open systems, let us discuss a simple model of autocatalysis [8] represented by two conjugate chemical reactions:

$$A + 2X \overset{k_1}{\underset{k_2}{\rightleftharpoons}} 3X, \quad X \overset{k_3}{\underset{k_4}{\rightleftharpoons}} B.$$  

(17)

The concentration of intermediate product $X$ is accepted as the only state variable. Substances $A$ and $B$ are either continuously supplied to the system or are removed from it, thereby providing the constancy of their concentrations $a$ and $b$. Substance $A$ is a catalyst. It increases the amount of intermediate product $X$ which, at the same time, is transformed to substance $B$.

At equilibrium, the rates of direct and inverse reactions are equal:

$$k_1ax^2 = k_2x^3$$

$$k_3x = k_4b.$$  

(18)
This relation determines in the only manner the concentration of intermediate product $x$ and the reagent concentration ratio $a/b$:

$$
x = \frac{k_4b}{k_3} = \frac{k_1a}{k_2}
$$

(19)

$$
\frac{b}{a} = \frac{k_1k_3}{k_2k_4}
$$

When a steady state is far from equilibrium, it is characterized by the balancing of the total effects of direct and inverse reactions in the form of a cubic equation:

$$
-k_2x^3 + k_1ax^2 - k_3x + k_4b = 0.
$$

(20)

Using the conventional MENT scheme to describe such a steady state with connection condition (20)

$$
-x \ln x + \lambda(-k_2x^3 + k_1ax^2 - k_3x + k_4) \to \text{max},
$$

(21)

we come to the equation:

$$
\ln x = \lambda(-3k_2x^2 + 2k_1ax - k_3)
$$

(22)

which can easily be solved graphically.

It is much more difficult now to trace a relationship between Lagrange factor $\lambda$ and concentration $x$ because of a lack of an analytical expression. However, the characteristics of the process described can be determined from relation (22). This equation can have either one or two solutions. In the latter case, bistability exists in an open chemical system and the phenomenon of chemical hours can occur. In this case, the values of concentrations $x$ corresponding to these states are found to be quantitatively dependent and obviously positive and the states are most stable in accordance with MENT.

A relationship between Lagrange factor $\lambda$, concentration $x$ and other parameters of the problem can easily be shown for the case when equation (22) has one solution. The graphs of the logarithm in the left part and those of a parabola in the right part of the equation touch each other and, consequently, have a common tangent. Let us transform the equality of the derivatives of these functions

$$
\frac{1}{x} = \lambda(-6k_2x + 2k_1a)
$$

(23)

into a quadratic equation:

$$
6k_2\lambda x^2 - 2k_1a\lambda x + 1 = 0.
$$

(24)

Because the solution is unique, the discriminant of this quadratic equation must be equal to zero:

$$
4k_1^2a^2\lambda^2 - 24k_2\lambda = 0.
$$

(25)

Lagrange factor $\lambda$ can thus be expressed by the relation:

$$
\lambda = \frac{6k_2}{k_1^2a^2}.
$$

(26)
and concentration \( x \), which passes to a bistable mode, by the relation:

\[
x = \frac{k_1a}{6k_2}.
\]

(27)

The value. It has been shown experimentally that transition to a bistable mode occurs before thermodynamic equilibrium sets in.

MENT is a convenient technique for simulation of systems because it allows to use not only empirical but also a priori information. One can estimate the sensitivity of the distribution function to one or another condition. To model complex statistical systems on a computer, special MENT algorithms [11, 12] are used.

Conclusions

1. MENT is an up-to-date and promising technique, which can efficiently be used to solve statistical problems. The MENT estimate of the distribution function agrees with the density properties of probability. The non-negativity property is satisfied automatically because the MENT solution has an exponential form, and normalization can be considered one of additional conditions observed when applying the technique.

2. Generalizing the second law of thermodynamics for open systems, one can use MENT for their statistical description.

3. When describing equilibrium and non-equilibrium states on the basis of conventional statistical models, MENT gives known results and new results when used to describe the steady states of open systems with autocatalytic reactions.

4. A simple scheme, observation data and a priori information used as restrictions make MENT a convenient tool for describing and modelling various processes, including those in high-energy physics.

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