Steady State of a Dissipative Flow-Controlled System and the Maximum Entropy Production Principle

Robert K. Niven

School of Engineering and Information Technology, The University of New South Wales at ADFA, Canberra, ACT, 2600, Australia.
Niels Bohr Institute, University of Copenhagen, Copenhagen Ø, Denmark.

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A theory to predict the steady state position of a dissipative, flow-controlled system, as defined by a control volume, is developed based on the Maximum Entropy (MaxEnt) principle of Jaynes, involving minimisation of a generalised free energy-like potential. The analysis provides a theoretical justification of a local, conditional form of the Maximum Entropy Production (MEP) principle, which successfully predicts the observable properties of many such systems. The analysis reveals a very different manifestation of the second law of thermodynamics in steady state flow systems, which provides a driving force for the formation of complex systems, including life.

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1. INTRODUCTION

For over three decades, the Maximum Entropy Production (MEP) principle – more precisely termed the “maximum rate of thermodynamic entropy production” principle – has been found to give successful predictions for the steady state properties of a variety of dynamic, many-degree-of-freedom systems subject to flows of mass, energy, momentum, charge and/or with chemical reactions. Prominent examples include the global general circulation (atmospheric and oceanic) system of the Earth [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13] and possibly other planetary bodies [14, 15]: turbulent convection in a heated fluid (Rayleigh-Bénard convection) [16], mantle convection in the Earth [17, 18, 19] and moons of Jupiter and Saturn [20], global Earth-biosphere water and nutrient cycles [21, 22], vegetation spatial distributions [23], biochemical metabolism [24] and ecosystem operation [25]. The Earth climate models include quite accurate predictions of the mean latitudinal air temperature, meridional heat flux [1], mean vertical air temperature profile, vertical heat flux [1], and historical latitudinal air temperature gradients over decadal and glacial-interglacial time scales [12].

The MEP principle has also been invoked for the explanation of shear turbulence (Couette flow) [17], incompressible and compressible turbulent fluid flow [26], currents in electrical circuits [27, 28, 29, 30], plasma formation [31] and structure [32], crystal growth [33], chemical cycle kinetics [34], diffusion in non-uniform solids [35], photosynthesis mechanism [36], biomolecular motors [37] and economic activity [38]; several detailed reviews are available [9, 11, 39, 40]. The MEP principle has even been invoked as a technical basis for the Gaia hypothesis [10, 21, 41, 42]. The apparent successes of the MEP principle – which can exclude most details of the dynamics – contradict the prevailing paradigm of developing ever-more-complicated dynamic models of complex systems, for example the general circulation models (GCMs) of the Earth climate system. This suggests that in a large class of steady state systems, the dynamics adjust themselves to achieve a state of MEP, rather than the entropy production being a consequence of the dynamics. This implies the action of an as-yet unrecognised physical principle applicable to flow systems – beyond the domain of present-day thermodynamics – which provides the driving force in the system, and so can be used as a “shortcut” in system modelling.

In recent years, several workers have attempted to justify the MEP principle on theoretical grounds. Dewar [43, 44] analysed a time-variant, non-equilibrium system in terms of its available paths in parameter space; the transient and steady state positions are inferred by the maximum entropy (MaxEnt) method of Jaynes [45, 46, 47, 48, 49, 50, 51], using an entropy defined on the set of paths. The analysis has received some criticisms [52, 53], to suggest it might apply only in the near-equilibrium linear (Onsager) regime, in which fluxes are linearly proportional to their driving gradients [54, 55]. Attard [56, 57] also gives an analysis based on an entropy defined on transition probabilities, but cast in the terminology of traditional statistical mechanics (although he denies the MEP principle in favour of an alternative). Beretta [58, 59, 60, 61] examines a steepest entropy ascent principle for transitions between states, based on a quantum thermodynamics formulation. Other arguments for MEP have been given by Županović and co-workers [62], based on the “most probable” increase in entropy during a fluctuation (also criticised [52]), and by Martyushev [63], based on a frame of reference (relative velocity) argument. Of course, this is a well-trodden field, with many historical antecedents to the MEP prin-
principle (see the fascinating review by Martyushev and Seleznev [39]). If the above objections can be overcome, the path-based analyses hold the promise of predicting the behaviour of time-variant systems (transport phenomena). However, they seem unnecessarily complicated for the task of predicting the steady state position, when (as will be shown) a more direct method is available.

The aim of this work is to outline a theory to directly predict the steady state position of a flow-controlled system, based on Jaynes’ MaxEnt method [45, 46, 47, 48, 49, 50, 51]. The analysis provides a theoretical derivation of a local, conditional form of the MEP principle for steady state thermodynamic systems of any type (e.g. heat flow, fluid flow, diffusive flow, electrical flow, chemical and process engineering, biological and human systems). The approach taken here differs from previous analyses [43, 44, 52, 53, 62] in that it considers flux rather than path probabilities, to directly obtain the steady state position. It also employs the local equilibrium assumption commonly used in engineering control volume analysis, but does not appear to be restricted to the linear transport regime; indeed, the linear regime emerges as a first-order approximation to the analysis.

This work is set out as follows. In §2 the principles of Jaynes’ generic theory are outlined, leading in §3 to a discussion of different types of physical systems, which influences the manner in which Jaynes’ method can be applied. Equilibrium systems are first examined, to demonstrate the universality of Jaynes’ method and to highlight the (easily overlooked) equilibrium analogue of the MEP principle. In §4 a theory is developed to predict the steady state position of a flow-controlled thermodynamic system; by comparison to a traditional control volume analysis, this is found to give a local, conditional form of the MEP principle. The implications of the analysis for the formation of complex systems, including life, are discussed in §5.

2. JAYNES’ GENERIC THEORY

We first summarise Jaynes’ generic formulation of statistical mechanics [43, 46, 47, 48, 49, 50, 51], based on the minimum divergence, maximum relative entropy or minimum cross-entropy (MinXEnt) principle, within which the maximum entropy (MaxEnt) principle can be considered a special case. Consider a system of N discrete, distinguishable entities distributed amongst s distinguishable categories or “states” within a system, with constant source (“prior”) probabilities $q_i$ for the filling of each state $i = 1, ..., s$ (the states $i$ may be multivariate, e.g. $\{1, i, j, ..., s\}$). From information-theoretic considerations [64, 65] and/or by combinatorial arguments [66, 67, 68, 69, 70, 71], it can be shown that in the asymptotic limit $N \rightarrow \infty$ (the Stirling approximation [72] or Sanov’s theorem [73]), the “least informative” or “most probable” distribution of the system is obtained by minimising the Kullback-Leibler cross-entropy function $\text{[74, 75]}$ (the negative of the relative entropy function $\text{[47]}$):

$$D = \sum_{i=1}^{s} p_i \ln \frac{p_i}{q_i}$$

subject to the natural and moment constraints on the system, respectively:

$$\sum_{i=1}^{s} p_i = 1,$n(2)$$

$$\sum_{i=1}^{s} p_i f_{ri} = \langle f_r \rangle, \quad r = 1, ..., R,$$n(3)$$

where $p_i$ is the probability of the $i$th category, $f_{ri}$ is the value of property $f_r$ for the $i$th category and $\langle f_r \rangle$ is the mathematical expectation of $f_{ri}$ (generally, each $f_r$ is a “conserved quantity”). Applying Lagrange’s method of undetermined multipliers to (2) gives the stationary or most probable distribution of the system:

$$p^*_i = q_i e^{-\lambda^*_r - \sum_{r=1}^{R} \lambda^*_r f_{ri}} = (Z^*_q)^{-1} q_i e^{-\sum_{r=1}^{R} \lambda^*_r f_{ri}},$$

$$Z^*_q = e^{\lambda^*_0} = \sum_{i=1}^{s} q_i e^{-\sum_{r=1}^{R} \lambda^*_r f_{ri}},$$

where $\lambda^*_r$ is the Lagrangian multiplier for the $r$th constraint, $\lambda^*_0$ is the “Massieu function” [76], $Z^*_q$ is the generalised partition function and $\langle \rangle$ denotes a quantity at the stationary position. Here “stationary” implies a time-invariant distribution [41], with no additional physical interpretation. Subsequent generic analyses pioneered by Jaynes [43, 46, 47, 48, 49, 50, 51, 70] give the relations:

$$D^* = -\lambda^*_0 - \sum_{r=1}^{R} \lambda^*_r \langle f_r \rangle$$

$$\frac{\partial \lambda^*_0}{\partial \lambda^*_r} = -\langle f_r \rangle$$

$$\frac{\partial^2 \lambda^*_0}{\partial \lambda^*_r \partial \lambda^*_m} = \langle f_r f_m \rangle - \langle f_r \rangle \langle f_m \rangle = -\frac{\partial \langle f_r \rangle}{\partial \lambda^*_m} - \frac{\partial \langle f_m \rangle}{\partial \lambda^*_r}$$

$$dD^* = \sum_{r=1}^{R} \lambda^*_r (d\langle f_r \rangle - dW_r) = -\sum_{r=1}^{R} \lambda^*_r \delta Q_r$$

where $dW_r = \langle df_r \rangle = \sum_{i=1}^{s} p^*_i df_{ri}$ and $\delta Q_r = \sum_{i=1}^{s} dp^*_i f_{ri}$ can be identified, respectively, as the increments of “generalised work” on the system and “generalised heat” added to the system during a change in

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1 Use of the continuous form of [41], or the von Neumann entropy based on density matrices, requires different mathematics but gives essentially the same macroscopic results [3-7].
the $r$th constraint $d(f_r)$, in which $\delta x$ denotes a path-dependent differential. (A path is here defined as a specified trajectory in macroscopic coordinates, e.g. constant constraints $\{f_r\}$, constant multipliers $\{\lambda_r\}$ or some mixed form.) To avoid double counting, the $\delta W_r$ terms represent generalised work processes in addition to those introduced by other constraints.

For equal $q_i = s^{-1}$, the foregoing analysis reduces to that of maximising the Shannon entropy $\{h\}$:

$$\mathcal{H} = -\sum_{i=1}^{s} p_i \ln p_i$$

subject to the same constraints $\{2\}-\{3\}$. This gives the same relations $\{1\}-\{9\}$, except that $D^*$ is replaced by $-\mathcal{H}^*$ and the $q_i$ cancel from $\{1\}$.

It is important to note that relations $\{1\}-\{9\}$ only concern the stationary position, i.e. they apply “on the manifold of stationary positions”. Transitions between stationary positions are thus assumed to be quasi-static, i.e. they are idealised as taking place only between stationary positions, without any intermediate non-stationary positions $\mathcal{H}^\dagger$. Comparing $\{9\}$ to the derivative of $\{5\}$ gives the differential:

$$d\phi^* = -d\lambda_0^* = -d\ln Z_0^* = \sum_{r=1}^{R} \lambda_r^* \delta W_r + \sum_{r=1}^{R} d\lambda_r^* \langle f_r \rangle$$

$$= dD^* + \sum_{r=1}^{R} \lambda_r^* d\langle f_r \rangle + \sum_{r=1}^{R} d\lambda_r^* \langle f_r \rangle$$

$\phi^*$ therefore encompasses “all possible changes” in the stationary position, due to changes in the constraints, multipliers or cross-entropy (generalised heat input). For constant multipliers, $d\phi^* |_{\{\lambda_r\}}$ is equal to the weighted change in generalised work on the system; its integrated form $\phi^*$ can therefore be regarded as a generalised, dimensionless free energy-like function or generalised potential $\{43\}$.

We now come to the central tenet of this analysis, the importance of which will become clear in $\{4\}-\{6\}$. If a dynamic system described by $\{1\}-\{11\}$ contains mechanism(s) by which its generalised potential $\phi^*$ can be dissipated (converted to an unrecoverable form), by processes not represented in the constraints, then by successive increments $d\phi^* < 0$ it will converge to a stationary position of no further extractable generalised work $d\phi^* = 0$, equivalent to the minimum position $\phi^* = \phi_{\text{min}}^*$. We here confine the discussion to systems with reproducible dissipative phenomena, whereupon $\phi_{\text{min}}^*$ will be predictable (i.e. $\phi^*$ is a state function). If the incremental changes are restricted in some manner – e.g. the system is confined to a constant-constraint $\{\langle f_r \rangle\}$ or a constant-multiplier $\{\lambda_r\}$ path – then $\phi_{\text{min}}^*$ will be the local (path-dependent) minimum. If there is no such path restriction, $\phi_{\text{min}}^*$ will constitute the global minimum.

It is emphasised that the above analysis $\{1\}-\{11\}$ is generic, and applies to any probabilistic system which can be analysed by the Kullback-Leibler cross-entropy $\{1\}$ or Shannon entropy $\{10\}$. Although its primary application has been to equilibrium thermodynamics, the analysis has far broader power of application $\{45\}$.

For this reason, the symbols used above are generic, and should not be interpreted in terms of particular thermodynamic quantities except when so stated (e.g. the generic entropy $\mathcal{H}$ should not be confused with the thermodynamic entropy $S$).

3. TYPES OF SYSTEMS

3.1. Quantity-Constrained (Equilibrium) Systems

![FIG. 1: (Color online) Different types of physical systems: (a) isolated, (b) diffusive and (c) flow-controlled (control volume).](image)

To understand the application of MinXEnt, we now consider several types of probabilistic systems composed of discrete entities, as illustrated in Figure 1. Historically, Jaynes' method (and its predecessor, traditional statistical mechanics) has been applied to (a) isolated (microcanonical) systems (Figure 1a), held in a state of...
constant mean contents $\langle f_{r} \rangle$ of various physical quantities $f_{r}$ by isolation from the rest of the universe; and (b) various types of open or diffusive (e.g., canonical, grand canonical) systems (Figure 11), open to the diffusion of various quantities $f_{r}$ but with no directed flows, and in contact with infinite *generalised baths* of constant corresponding $\lambda_{r}$ [43]. Clearly, such representations are convenient models to enable the construction of thermodynamic relationships: no system can really be isolated from the rest of the universe, whilst the mechanisms used to maintain the generalised baths are not usually explained. Notwithstanding this criticism, in either case the “least informative” or “most probable” position of the system – the *equilibrium position* – can be calculated by the application of MinXEnt (or, for equal $q_{i}$, by MaxEnt) subject to the natural and moment constraints [43]. Jaynes’ generic formulation ([2]) can then be applied to the analysis of such systems.

As an example, consider an “open” or “heterogeneous” thermodynamic system for which the “entities” are interpreted either as discrete particles (atoms, ions, molecules, etc) or, in the Gibbs-Einstein representation, as duplicates of the entire system [78, 79, 80]. Each entity has the choice of its (quantised) internal energy $U_{i}$, $i = 1, ..., s$; volume element $V_{j}$, $j = 1, ..., t$; and moles of particles $n_{c}$, of each chemical species $c = 1, ..., C$, for which the number of particles $N_{c}$ can range between zero and (effectively) infinity. The system is constrained by its natural constraint $\{i,j,\{N_{c}\}\}$, mean internal energy $\langle U \rangle$, mean volume $\langle V \rangle$ and mean moles of particles $\langle n_{c} \rangle$ of each type $c$. Assuming, from the principle of insufficient reason [51], that each multivariate state $\{i,j,\{N_{c}\}\}$ is equiprobable, we maximise the Shannon entropy:

$$\mathcal{S}_{eq} = - \sum_{i=1}^{s} \sum_{j=1}^{t} \sum_{N_{1}=0}^{\infty} \sum_{N_{2}=0}^{\infty} \cdots \sum_{N_{C}=0}^{\infty} p_{i,j,\{N_{c}\}} \ln p_{i,j,\{N_{c}\}},$$

subject to the constraints, giving the equilibrium position [4]:

$$p_{i,j,\{N_{c}\}} = \frac{\alpha_{i,j,\{N_{c}\}}^{*}}{\Xi^{*}},$$

$$\alpha_{i,j,\{N_{c}\}}^{*} = \exp(-\lambda_{U}U_{i} - \lambda_{V}V_{j} - \sum_{c=1}^{C} \lambda_{c}n_{c}),$$

$$\Xi^{*} = e^{\lambda_{0}} = \sum_{i=1}^{s} \sum_{j=1}^{t} \sum_{N_{1}=0}^{\infty} \sum_{N_{2}=0}^{\infty} \cdots \sum_{N_{C}=0}^{\infty} \alpha_{i,j,\{N_{c}\}}^{*}.$$

From our existing (historical) knowledge of thermodynamics (e.g., by monotonicity arguments [50] or from the zeroth law of thermodynamics [49, 50]), we can identify the Lagrangian multipliers $\lambda_{U} = 1/kT$, $\lambda_{V} = P/kT$ and $\lambda_{c} = -\mu_{c}/kT$ as functions of the Boltzmann constant $k$, absolute temperature $T$, absolute pressure $P$ and molar chemical potential $\mu_{c}$ of each species $c$, whence:

$$\alpha_{i,j,\{N_{c}\}}^{*} = \exp\left(-\frac{U_{i}}{kT} - \frac{PV_{j}}{kT} + \sum_{c=1}^{C} \frac{\mu_{c}n_{c}}{kT}\right),$$

and $\Xi^{*}$ is the grand partition function. Recognising the thermodynamic entropy as $S = k\mathcal{S}_{eq}$, [5] and [6] give:

$$\mathcal{S}_{eq} = k \ln \Xi^{*} + \frac{\langle U \rangle}{T} + \frac{P\langle V \rangle}{T} - \sum_{c=1}^{C} \frac{\mu_{c}\langle n_{c} \rangle}{T}$$

$$d\mathcal{S}_{eq} = \frac{\delta Q_{U}}{T} + \frac{P\delta Q_{V}}{T} - \sum_{c=1}^{C} \frac{\mu_{c}\delta Q_{n_{c}}}{T}$$

$$= \frac{1}{T}(d\langle U \rangle - \delta W_{U}) + \frac{P}{T}(d\langle V \rangle - \delta W_{V})$$

$$- \sum_{c=1}^{C} \frac{\mu_{c}}{T}(d\langle n_{c} \rangle - \delta W_{n_{c}})$$

$$= \frac{d\langle U \rangle}{T} + \frac{Pd\langle V \rangle}{T} - \sum_{c=1}^{C} \frac{\mu_{c}d\langle n_{c} \rangle}{T} - \sum_{c=1}^{C} \frac{\mu_{c}\delta W_{n_{c}}}{T}$$

where $\delta Q_{U}$, $\delta Q_{V}$ and $\delta Q_{n_{c}}$ are the increments in generalised heats, and $\delta W_{U}$, $\delta W_{V}$ and $\delta W_{n_{c}}$ are the increments in generalised work, associated respectively with constraints $\langle U \rangle$, $\langle V \rangle$ and $\langle n_{c} \rangle$. In the first line of (16), the first two terms can be amalgamated into the (actual) heat term $\delta Q/T$ used in thermodynamics; for systems with variable particle numbers, $\delta Q_{n_{c}} = 0$ and (16) reduces to the Clausius equality $d\mathcal{S}_{eq} = \delta Q/T$. Similarly, in the last line of (16), the $\delta W_{V}$ and $P\delta W_{V}$ terms are amalgamated into the combined (actual) work $\sum \delta W$; the latter therefore represents the total (actual) work on the system, in addition to the $Pd\langle V \rangle$ and $\mu_{c}d\langle n_{c} \rangle$ work. For this study, we ignore relativistic or other changes in the “mass levels”, and set $\sum \delta W_{n_{c}} = 0$. It is again emphasised that [13]-[16] apply only to quasistatic transitions, in this case “on the manifold of equilibrium positions”.

Eqs. [11] and [15]-[16] give the change in the (actual)

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3 As mentioned in [2] the $\delta W_{V}$ refer to work terms in addition to those introduced by the other constraints. Thus in the above equilibrium system, the $Pd\langle V \rangle$ work cannot be introduced within the work term $\delta W_{U}$ associated with $\langle U \rangle$, since it is already included by virtue of the $\langle V \rangle$ constraint. Similarly, $\delta W_{U}$ cannot include the chemical potential work terms $\mu_{c}d\langle n_{c} \rangle$, since these are included by the $\langle n_{c} \rangle$ constraints. This consideration is quite general: the constraints $(f_{r})$ must be linearly independent, but need not be orthogonal; the weighted generalised work terms $\lambda_{m}d(f_{m})$, $m \neq r$. 
free energy function, in energy units:

\[ dJ^* = kT d\phi_{eq}^* = -kT d\ln \Xi^* \]

\[ = -TdS^* + d\langle U \rangle + Pd\langle V \rangle - \sum_{c=1}^{C} \mu_c d\langle n_c \rangle \tag{17} \]

\[ + T \left\{ d\left( \frac{1}{T} \right) \langle U \rangle + d\left( \frac{P}{T} \right) \langle V \rangle - \sum_{c=1}^{C} d\left( \frac{\mu_c}{T} \right) \langle n_c \rangle \right\} \]

in which, from (16), the first four terms on the right correspond to the total work \( \sum \delta W \). Note that – as per Gibbs – we could restrict (17) to these first four terms, by imposing the Gibbs-Duhem relation \( \sum_{c=1}^{R} d\phi_r(f_r) = 0 \). We here wish, however, to consider all possible changes in \( J^* \), allowing both Gibbs-Duhem paths and non-Gibbs-Duhem paths. Integration of (17) from a zero reference state then yields:\n
\[ J^* = kT \phi_{eq}^* = -kT \ln \Xi^* \]

\[ = -TS^* + \langle U \rangle + P \langle V \rangle - \sum_{c=1}^{C} \mu_c \langle n_c \rangle \tag{18} \]

Although less well known, \( J^* \) was reported by Gibbs; in modified form, it forms the basis of the available energy, essergy and exergy concepts. For constant composition, \( J^* \) reduces to the Gibbs free energy \( G^* = -TS^* + \langle U \rangle + P \langle V \rangle \) plus a constant, whilst for constant composition and volume, it gives the Helmholtz free energy \( F^* = -TS^* + \langle U \rangle \) plus a constant.

We first examine (17)-(18) from the original perspective of Gibbs, see also [77, 90, 91]. Following the standard treatment, we consider an open system surrounded by an intensive variable bath (Figure 1), with the double system-bath isolated from the rest of the universe; a single isolated system (Figure 1a) can then be considered as a special case. From (16), (17),

\[ dJ^*|_{T,P,\{\mu_c\}} = \sum \delta W \]

gives the total change in work on the system along a Gibbs-Duhem path, due to processes not represented in the constraints. Thus \( dJ^*|_{T,P,\{\mu_c\}} < 0 \) indicates spontaneous change (work extraction) and \( dJ^*|_{T,P,\{\mu_c\}} > 0 \) a forced change (work input). If the system contains some reproducible dissipative processes, by which work can be expended (e.g. friction or chemical reactions), it will move towards the equilibrium defined by \( dJ^*|_{T,P,\{\mu_c\}} = 0 \), or in other words, to the minimum free energy position \( J^* = J^*_{\min} \). This was implicitly understood by Gibbs, who refers to the equilibrium surface (the “fundamental relation” or Euler surface) as the “surface of dissipated energy”. If no work is extracted in useful form, each increment of lost work is given by [94, 95, 96]:

\[ dJ^*|_{T,P,\{\mu_c\}} = -TdS^*|_{T,P,\{\mu_c\}} - T\delta\sigma|_{T,P,\{\mu_c\}} \leq 0 \tag{19} \]

where \( dS^*|_{T,P,\{\mu_c\}} \) and \( \delta\sigma|_{T,P,\{\mu_c\}} \) respectively give the change in entropy within and outside the system. Eq. (19) represents the net irreversible increase in entropy in the double system-bath (converted to energy units and with change of sign), reflecting the interplay between irreversible change within the system, and the exporting of irreversible change to the bath by the transfer of generalised heats [c.f. 91, 95, 97, 98]. The second term \( \delta\sigma|_{T,P,\{\mu_c\}} \) can be labelled the “entropy produced” by the system, since it materialises outside rather than inside the system. If the incremental losses are unrecoverable (i.e. \( dS^*|_{T,P,\{\mu_c\}} \geq 0 \) and \( \delta\sigma|_{T,P,\{\mu_c\}} \geq 0 \)), the equilibrium position \( J^* = J^*_{\min} \) will correspond to the position of maximum entropy produced, \( \sigma = \sigma_{\max} \) [c.f. 93].

We now go a step farther than Gibbs, to consider systems which can deviate from a Gibbs-Duhem path during dissipation. We again only consider systems with reproducible processes. Here it is preferable to use the potential \( d\psi^* = k d\phi_{eq}^* = dJ^*/T \) (the grand potential form of the negative Massieu function) or negative Planck potential [94, 95, 97, 98, 100]; see Appendix A, rather than \( dJ^* \), to more effectively handle changes in temperature. From (17), each increment \( d\psi^* \leq 0 \) can be divided into two parts: (i) the part \( d\psi^*|_{S^*,\langle U \rangle,V,\{\mu_c\}} < 0 \) which is expended against changes in the intensive variables, causing an irreversible loss of “ability to do generalised work”, manifested as an increase in entropy within or outside the system, and (ii) the part \( d\psi^*|_{T,P,\{\mu_c\}} < 0 \) which – as in (19) – is lost by dissipation within or outside the system. The net loss can thus be written as:

\[ d\psi^* = -dS^*|_{T,P,\{\mu_c\}} - \delta\sigma|_{T,P,\{\mu_c\}} \]

\[ - dS^*|_{S^*,\langle U \rangle,V,\{\mu_c\}} - \delta\sigma|_{S^*,\langle U \rangle,V,\{\mu_c\}} \]

\[ = -dS^* - \delta\sigma \leq 0 \tag{20} \]

The terms \( dS^*|_{S^*,\langle U \rangle,V,\{\mu_c\}} \) and \( \delta\sigma|_{S^*,\langle U \rangle,V,\{\mu_c\}} \) constitute “uncompensated transformations” of Clausius – respectively within and outside the system – since they involve irreversible change(s) in one or more intensive variables. For convenience, the four terms are unified in the second line of (20), into overall entropy changes within and outside the system.

From (20), the system will approach the equilibrium position defined by \( \psi^* = \psi^*_{\min} \), equivalent to the maximum net irreversible increase in entropy within and outside the system. If \( dS^* \geq 0 \) and \( \delta\sigma \geq 0 \), this will again

\[ ^5 \text{It is misleading to call } \sigma \text{ the entropy production. In English, “production” implies an on-going phenomenon, hence a rate process (e.g. a nation’s steel production). This can only apply to a flow system. Confusingly, the symbol } \sigma \text{ is used interchangeably for the amount of thermodynamic entropy produced by a system, its rate of production by a system or the rate per unit volume; these are labelled } \sigma, \bar{\sigma} \text{ and } \dot{\sigma} \text{ respectively here.} \]
correspond to the state at which maximum thermodynamic entropy is produced, \( \sigma = \sigma_{\text{max}} \). If the path of possible transitions is prescribed (e.g. constant \( \{f_r\} \), constant \( \{\lambda_r\} \) or a mixed choice), the system will approach the local minimum \( \psi_{\text{min}} = -\int (dS^* + \delta \sigma) \) along that path; if not, it will converge to the global minimum.

Thus in a dissipative, equilibrium system, the final equilibrium will occur at the position of minimum generalised potential \( \phi_{\text{eq}}^* \). If the incremental increases \( dS^* \geq 0 \) and \( \delta \sigma \geq 0 \) are unrecoverable, this will be equivalent to the position at which maximum thermodynamic entropy is produced. This provides an \textit{equilibrium analogue} of the MEP principle, which is easily overlooked, since it involves a connection between equilibrium states and irreversible changes.

We can also consider the multiplier relations (6), variances (7) and covariances (8), which yield a highly important set of relations for equilibrium systems, as listed in Table I. These include the “Maxwell relations” (7a, 10), which express the connections between various material properties or susceptibilities of the system (e.g. heat capacity, compressibility, coefficient of thermal expansion, activity coefficients, etc). The relations in Table I are valid only at equilibrium; i.e. they apply on the manifold of equilibrium positions. Although discovered long before Jaynes’ generic analysis, such relations demonstrate the power of Jaynes’ method, particularly when applied to new situations.

### 3.2. Flow-Controlled (Steady-State) Systems

We now examine a third kind of probabilistic system, which much more closely matches our experience of real systems and of life on Earth: the \textit{flow-controlled system} (Figure 1), constrained by flows of various physical quantities \( f_r \) through the system. Flow-controlled systems are familiar to fluid mechanists and engineers, as defined by the \textit{control volume}, a geometric region through which fluid(s) can flow (the Eulerian description), bounded by its \textit{control surface}. In the language of statistical mechanics, flow-controlled systems constitute a separate “ensemble”, very different to those examined in equilibrium thermodynamics. For the simplified flow system shown in Figure 1, the mean rate of change of each conserved quantity \( f_r \) with respect to time \( t \) within the system is:

\[
\frac{d\langle f_r \rangle}{dt} = \langle f_r \rangle_{\text{in}} - \langle f_r \rangle_{\text{out}} + \langle \dot{f}_r \rangle_{\text{prod}}
\]  

(21)

where \( \langle f_r \rangle_{\text{in}} \), \( \langle f_r \rangle_{\text{out}} \) and \( \langle \dot{f}_r \rangle_{\text{prod}} \) represent, respectively, the mean flow rates of \( f_r \) into and out of the control volume, and rate of production within the control volume. At steady state, \( d\langle f_r \rangle/dt = 0 \), and so for a system with no production terms, (21) reduces to:

\[
\langle f_r \rangle_{\text{in}} = \langle f_r \rangle_{\text{out}} = \langle f_r \rangle
\]  

(22)

The mean flow rates \( \langle f_r \rangle \) can then be interpreted as constraints on the distribution of instantaneous flow rates \( f_{ri} \) through the system. By information-theoretic reasoning [64, 65] and/or the combinatorial basis of entropy [66, 67, 68, 69, 70, 71], the MinXEnt principle can then be used to calculate the stationary or “most probable” distribution of flow rates within the system – its \textit{steady state position} – subject to the constraints on the system. Moment constraints are just moment constraints, regardless of their physical manifestation. In consequence, the entirety of Jaynes’ generic approach (22) can be applied to the analysis of steady state, as well as equilibrium, systems. Of course, flow-controlled systems may be subject to variable or cyclic flow rates, causing deviations from the steady state position; such effects are not considered further here.

Notwithstanding the broad applicability of Jaynes’ method to both equilibrium and steady state systems, it is important to understand their differences. In equilibrium systems, the physical “entities” represent individual particles (e.g. molecules, ions, oscillators, etc), or entire systems of such particles, which can choose different values \( f_{ri} \) of various physical parameters \( f_r \). In contrast, in the Eulerian description of a steady state system, the “entities” represent individual fluid elements (in the

| Table I: Multiplier relations (6), variances (7) and covariances (8) for the example equilibrium thermodynamic system (3.3) at equilibrium (note \( \text{var}(a) = \langle a^2 \rangle - \langle a \rangle^2 \), \( \text{cov}(a,b) = \langle ab \rangle - \langle a \rangle \langle b \rangle \), and \( \mu_b \) implies \( b \neq c \)). |
|------------------|------------------|-------------------|
| Multiplier relations |
| \( \langle U \rangle = kT^2 \frac{\partial \ln \Xi^*}{\partial T} \) \( T,P,\{\mu_c\} \) |
| \( \langle V \rangle = -kT \frac{\partial \ln \Xi^*}{\partial P} \) \( T,\{\mu_c\} \) |
| \( \langle n_c \rangle = kT \frac{\partial \ln \Xi^*}{\partial \mu_c} \) \( T,P,\{\mu_b\} \) |

| Variances |
| \( \text{var}(U) = kT^2 \frac{\partial^2 \langle U \rangle}{\partial T^2} \) \( P,\{\mu_c\} \) |
| \( \text{var}(V) = -kT \frac{\partial^2 \langle V \rangle}{\partial P^2} \) \( T,\{\mu_c\} \) |
| \( \text{var}(n_c) = kT \frac{\partial^2 \langle n_c \rangle}{\partial \mu_c^2} \) \( T,P,\{\mu_b\} \) |

| Covariances (Maxwell relations) |
| \( \text{cov}(U,V) = -\frac{\partial \langle U \rangle}{\partial P} \frac{\partial \langle V \rangle}{\partial T} \) \( T,\{\mu_c\} \) |
| \( \text{cov}(U,n_c) = \frac{\partial \langle U \rangle}{\partial \mu_c} \) \( T,P,\{\mu_b\} \) |
| \( \text{cov}(V,n_c) = \frac{\partial \langle V \rangle}{\partial \mu_c} \) \( T,\{\mu_c\} \) |
| \( \text{cov}(n_c,n_b) = \frac{\partial \langle n_c \rangle}{\partial \mu_b} \) \( T,P,\{\mu_{c,b}\} \) |

The above expressions are applicable to situations where the flow rates \( f_{ri} \) are conserved, i.e. the system is \textit{flow-controlled}.
The MEP principle has been criticised for appearing to lack a unique stationary position. For a given constraint set, the stationary position of a system is usually considered unique within its parameter space. However, this depends on the type of constraints; e.g. power-law constraints are known to give multiple stationary positions. Uniqueness of the stationary position, in general, is therefore not claimed here, but is correct in the case of linear moment constraints.

A unique stationary position $D^*$ or $S^*$ in parameter space does not imply uniqueness of the dynamic structure(s) which can produce it; indeed, many such structures may be possible. The occurrence of any given structure will depend on the interplay between the system history (hysteresis phenomena) and the dynamics (the extent of fluctuations, or “jitter”, in the system), both of which lie outside the domain of Jaynes’ method per se. This feature is also well known in equilibrium systems: e.g. the equilibrium position of a supersaturated solution does not provide any information about the particle size distribution or geometric form of the precipitant, which instead will depend on localised (history-dependent) nucleation, crystallisation and diffusion phenomena.

The MEP principle has been criticised for appearing to be a means of selection between a few isolated optima, rather than a true variational principle. However, many equilibrium systems also exhibit a discontinuous approach to equilibrium for dynamic reasons, especially those with a phase change; e.g. the rapid solidification of a saturated solution of sodium acetate trihydrate – which can dissolve in its own water of crystallisation – when tapped. The equilibrium between nitroglycerine and oxygen also illustrates this point. In flow-controlled systems, we might expect any locally optimal steady states to be similarly narrowly defined, since they may require special coordination of flows in different domains. Thus although derived by variational means, there is nothing wrong with the minimisation of $\phi^*$ as a selection principle.

The rationale for the analysis of a steady state system by Jaynes’ method is therefore identical – with many of the same caveats – to that for equilibrium systems.

4. ANALYSIS OF FLOW-CONTROLLED SYSTEMS

4.1. Control Volume Analysis

Calculation of the steady state position of a flow-controlled system can now proceed, firstly involving a traditional engineering thermodynamic analysis. Consider the control volume shown in Figure 2 divided into infinitesimal volume elements, each of which is assumed to satisfy the “local equilibrium assumption”. This allows the definition of local specific (per unit mass) quantities including specific internal energy $u$, specific volume $\rho^{-1}$, specific moles $n_c$ of each species $c$ and specific entropy $s$ of each element, where $\rho$ is the fluid density, as well as intensive variables such as temperature $T$, pressure $P$ and molal chemical potentials $\mu_c$ of each species $c$ on its boundary. The elements are thus sufficiently small for local equilibrium to hold, but obviously repre-
sent averages over regions large enough to be considered a thermodynamic continuum. In general, for an element subject to the total entropy flux \( \dot{J}_{s,tot} \), the volumetric rate of entropy production within the element (SI units: J K\(^{-1}\) m\(^{-3}\) s\(^{-1}\)) and overall rate of production (J K\(^{-1}\) s\(^{-1}\)) are \( [92] [105] [106] [107] \):

\[
\dot{\sigma} = \frac{\partial \rho s}{\partial t} + \nabla \cdot \dot{J}_{s,tot} \geq 0
\]

\[
\dot{\sigma} = \iiint_{CV} \hat{\sigma} dV \geq 0
\]

where \( \nabla = [\partial/\partial x, \partial/\partial y, \partial/\partial z]^\top \) is the Cartesian gradient operator; \( \mathbf{a} \cdot \mathbf{b} \) is the vector scalar product (dot product); and \( \mathbf{a}^\top \) is the transpose of \( \mathbf{a} \). For an element with mean local heat flux \( \dot{J}_Q \), diffusive mass fluxes \( j_c \) of each species \( c \) (relative to the mass-average velocity \( \mathbf{v} \) through the element), viscous stress tensor \( \tau \) and molar rate per unit volume \( \dot{\xi}_d \) of each chemical reaction \( d = 1, ..., D \), from the conservation of energy, mass, momentum and chemical species, the total entropy flux and rate of entropy production per volume reduce to [92] [105] [106] [107]:

\[
\dot{J}_{s,tot} = \rho s \mathbf{v} + \dot{J}_Q \left( \frac{1}{T} \right) - \sum_{c=1}^{C} \left( j_c \left( \frac{\mu_c}{M_c T} \right) - \sum_{c=1}^{C} \left( \nabla \left( \frac{\mu_c}{M_c T} \right) - \mathbf{g}_c \right) \right)
\]

\[
\dot{\sigma} = \dot{J}_Q \cdot \nabla \left( \frac{1}{T} \right) - \sum_{c=1}^{C} \left( \nabla \left( \frac{\mu_c}{M_c T} \right) - \mathbf{g}_c \right)
\]

\[
-\tau : \nabla \left( \frac{\mathbf{v}}{T} \right)^\top - \sum_{d=1}^{D} \dot{\xi}_d \frac{A_d}{T} \geq 0
\]

where \( \mathbf{A} : \mathbf{B} \) is the tensor scalar product; \( \mathbf{a} \mathbf{b}^\top \) is the vector dyadic product (often simply written \( \mathbf{ab} \)); \( M_c \) is the molecular mass of species \( c \); \( \mathbf{g}_c \) is the specific body force on each species \( c \); and \( A_d = \sum_{c=1}^{C} \nu_{cd} \mu_c \) is the molar chemical affinity of reaction \( d \), wherein \( \nu_{cd} \) is the stoichiometric coefficient of species \( c \) in the \( d \)th reaction \( (\nu_{cd} > 0 \text{ for a product and } A_d < 0 \text{ for a reactant}) \). Alternative formulations of \( [26] \) are available for different situations [102] [106] [107]: the present formulation is sufficiently broad for the present discussion.

In engineering modelling, the local mass, momentum and energy conservation equations and local entropy production [26] are usually non-analytic, and so are calculated on a two- or three-dimensional grid of finite rather than infinitesimal domains (e.g., by the finite element method) such that the local solutions are self-consistent and match the boundary conditions and source-sink terms of the overall control volume [89] [105] [106] [107]. This approach assumes that the calculated solution is unique. However, for many flow systems, the entropy production [26] is indeterminate, since one or more of the fluxes \( (\dot{J}_Q, j_c, \tau, \dot{\xi}_d) \) and/or gradients \( (\nabla (1/T), \nabla (\mu c / M c T), g_c / T, \nabla (\mathbf{v} / T)^\top) \) across the overall control volume are unknown, there being no principle within equilibrium thermodynamics by which they can be determined. In such cases, \( [26] \) has been solved by assuming the near-equilibrium linear (Onsager) regime, for which closure of the equations can be attained by the use of the equilibrium Gibbs-Duhem relation, Curie postulate, tensor symmetry and specification of the (linear) mass, momentum and energy diffusion coefficients [102] [106] [107]. Outside the linear regime, such as in turbulent fluid flow, these assumptions can produce serious computational difficulties (e.g. the need for extremely small elements) and are commonly handled by broad simplifications or empirical rules. Indeed, for many flow systems, the entropy production is completely disregarded. Eqs. [23]–[26] therefore apply to a flow-controlled thermodynamic system at steady state, but in many systems, may not uniquely define the steady state.

4.2. Jaynes Analysis

We now examine a flow-controlled system from Jaynes’ perspective [22]. In general, for an infinitesimal fluid element of a control volume, Jaynes’ relations for the steady state position are exactly as given in \( [4]–[11] \), but this time with mean flux constraints \( \mathbf{j}_i \) instead of quantity constraints \( \langle f_i \rangle \) (for convenience the \( \langle \cdot \rangle \) notation is dropped), and increments in generalised heat fluxes \( \delta q_r \) instead of in the generalised heats \( \delta Q_r \). The Lagrangian multipliers, say \( \zeta_r \), will of course be different to those of equilibrium thermodynamics \( \lambda_r \), conjugate to the flux constraints considered. We should also denote the cross-entropy or entropy by some special symbol, e.g. \( D_{st} \) or \( S_{st} \), to emphasise that it applies to a steady state system, and is quite different to that for an equilibrium system, \( D_{eq} \) or \( S_{eq} \). Our generic analysis of the steady state position of a flow-controlled system is now complete, requiring only the substitution of relevant parameters for the system under consideration.

Returning to the example of \( [11] \) we consider that each element can experience instantaneous values of the heat flux \( \dot{J}_Q \), mass fluxes \( \dot{J}_m \), stress tensor \( \tau \) and reaction rates \( \dot{\xi}_d \), where the indices \( I, J, L_d \) and \( N_c \) can (in principle) take any value. The system is constrained by mean values of the heat flux \( \dot{J}_Q \), mass fluxes \( \dot{J}_m \), stress tensor \( \tau \) and reaction rates \( \dot{\xi}_d \) through or within the element, and by the natural constraint \( [22] \), where expectations are taken with respect to the joint probability \( \pi_{I,J,L_d,N_c} \) of instantaneous fluxes. As for the equilibrium analysis \( [33, 11] \) we assume equiprobable

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6 We adhere strictly to engineering convention: a flux is measured in SI units of quantities s\(^{-1}\), whilst a flux is expressed in quantities m\(^{-2}\) s\(^{-1}\).

7 The viscous stress tensor \( \tau \) is equal to the net or molecular stress tensor \( \Pi \) less the uniform pressure component \( P \), whence \( \tau = \Pi - P \delta \), where \( \delta \) is the Kronecker delta tensor [103] [106] [107]. The stress notation of Bird et al. [107] is adopted here, in which \( P > 0 \) and \( \tau_{ij} > 0 \) denote compression.
subject to its constraints. From (13) and (9):

$$\pi^*_I = \frac{\beta^*_I}{Z^*},$$

$$\beta^*_I = \exp \left( -\zeta Q \cdot j_Q, I - \sum_{c=0}^{C} \zeta c \cdot j_N c - \zeta \tau : \tau - \sum_{d=1}^{D} \zeta_d \xi_d \right)$$

$$Z^* = e^{\zeta^*} = \sum_{I=1}^{\infty} \sum_{\xi=1}^{\infty} \sum_{d=1}^{\infty} \beta^*_I$$

$$\Delta st = \ln Z^* + \zeta Q \cdot j_Q + \sum_{c=1}^{C} \zeta c \cdot j_c + \zeta \tau : \tau + \sum_{d=1}^{D} \zeta_d \xi_d$$

$$d\Delta st^*_I = \zeta Q \cdot \delta q_Q + \sum_{c=1}^{C} \zeta c \cdot \delta q_c + \zeta \tau : \delta q_{\tau} + \sum_{d=1}^{D} \zeta_d \delta q_d$$

where the multipliers $\zeta Q, \zeta c, \zeta \tau$ and $\zeta d$ are associated with the heat, species, stress tensor and reaction constraints ($\zeta Q$ and $\zeta c$ are vectors and $\zeta \tau$ a second order tensor); $\delta q_Q, \delta q_c, \delta q_{\tau}$ and $\delta q_d$ are the corresponding changes in generalised heat fluxes; $\zeta^*$ is the Massieu function and $Z^* = e^{\zeta^*}$ the partition function. We see that $\Delta st^*$ is a dimensionless flux entropy. Eqs. (11) and (29)-(30) then give the change in a generalised potential function for steady state systems:

$$d\phi^*_st = -d\zeta^*_0 = -d\ln Z^*$$

$$= d\Delta st^*_I + \zeta Q \cdot \delta j_Q + \sum_{c=1}^{C} \zeta c \cdot \delta j_c + \zeta \tau : \delta \tau + \sum_{d=1}^{D} \zeta_d \delta \xi_d$$

$$+ d\zeta Q \cdot j_Q + \sum_{c=1}^{C} d\zeta c \cdot j_c + d\zeta \tau : \tau + \sum_{d=1}^{D} d\zeta_d \xi_d$$

Again – with Gibbs – we could exclude a net change in the multipliers $\zeta$, giving $d\phi^*_st|_{\zeta^*} = \sum_{r=1}^{R} \zeta c \delta w_r$, where $\delta w_r$ are changes in generalised work fluxes, in addition to those incorporated by other constraints. However, as with equilibrium systems (8.1), we wish to consider all possible variations of $\phi^*_st$. Integration of (31), from the zero flux, zero multiplier (equilibrium) position $\phi^*_st0 = -\Delta st^*$ yields:

$$\phi^*_st = -\Delta st^*_I + \zeta Q \cdot j_Q + \sum_{c=1}^{C} \zeta c \cdot j_c + \zeta \tau : \tau + \sum_{d=1}^{D} \zeta_d \xi_d$$

This is the steady state analogue of the equilibrium generalised potential $\phi^*_eq$. Relations (28)-(32) apply only to quasistatic transitions “on the manifold of steady state positions”. However, in this case – since this is a flow system – they always involve irreversible processes.

4.3. Synthesis

We can now combine the control volume (8.1) and Jaynesian (8.2) analyses of the example problem (Figure 2). Since the fluxes in (26) and (32) are linearly independent (not necessarily orthogonal), it is possible to equate similar terms, giving the following identities:

$$\phi^*_st = -\Delta st^*_I - \frac{\theta \nu}{k} \hat{\sigma}$$

$$\zeta Q = \theta \nu \nabla \left( \frac{\mu c}{M \cdot T} - \frac{q}{T} \right)$$

$$\zeta \tau = \theta \nu \nabla \left( \frac{v}{T} \right)$$

$$\zeta = \frac{\theta \nu A_d}{k} \frac{T}{T}$$

where $\theta$ and $\nu$ respectively are characteristic time and volume scales for the system considered, which emerge from the fact that $\phi^*_st, \Delta st^*_I$ and each product in (32) must be dimensionless.

From (8.3) and the foregoing analysis (2 and 8.1), if the flow-controlled system contains some process(es) which dissipate its generalised potential $\phi^*_st$, it will move by increments:

$$d\phi^*_st = -d\Delta st^*_I - \frac{\theta \nu}{k} \hat{\sigma} \leq 0$$

until it converges to a final steady state position given by $d\phi^*_st = 0$, hence at minimum $\phi^*_st$. Eq. (33) is analogous to (20) for equilibrium systems, accounting for the loss of generalised potential within and outside the system. If $d\Delta st^*_I \geq 0$ and $\hat{\sigma} \geq 0$, the steady state position $\phi^*_st = \phi^*_{st,min}$ will correspond to the position of maximum entropy production $\hat{\sigma} = \hat{\sigma}_{max}$. The analysis therefore provides a theoretical justification for a local form of the MEP principle, conditional on the assumption that incremental increases in $\Delta st^*$ and $\hat{\sigma}$ are irreversible. As with equilibrium systems (8.1), the dissipation may be confined to a constant-multiplier $\{\zeta^*_c\}$, constant-constraint $\{j_c\}$ or mixed constraint-path, leading to a local (path-dependent) minimum $\phi^*_{st,min}$, or may be free to choose its own path, whereupon it will attain a global minimum.

Processes for which $\hat{\sigma} > 0$ (to which the MEP principle applies) are here termed exoentropogenic, since they

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8 An alternative view, not considered further here, is to interpret $(\theta \nu/k) (g/J) \cdot \hat{j}$ as a generalised work flux, which must be subtracted from a redefined generalised heat flux $\delta q_j = (\theta \nu/k) \nabla(\mu c/M \cdot T) \cdot \hat{j}$.

9 Note $\theta \nu$ has SI units of m$^3$ s, equivalent to a four-dimensional space-time element, or action divided by pressure.

10 From ancient Greek: ezo-, outer or external (obverse endo-, within or internal); tropos, transformation (used by Clausius [83]); and -genic, generating or producing.
result in the production of entropy and its export to the rest of the universe. Exoentropogenic processes are one class of processes leading to the formation of steady-state flow systems, in the same way that exothermic reactions – in which the heat $\delta Q < 0$ – are one class of processes which generate equilibrium systems. It is left as an open question here whether exoentropogenic processes (and hence the MEP principle) are universal in application, or are merely an important class of processes applicable to flow systems at steady state.

### 4.4. Implications

The analysis has several important implications. Firstly, under conditions in which the MEP principle applies, from (20) and (53):

- If the fluxes or rates $\vec{j}_Q, \vec{j}_c, \tau$ and $\xi_d$ in (20) are indeed constant, the system will achieve MEP by maximising the magnitudes of the gradients or forces $|\nabla (T^{-1})|$, $|\nabla (\mu_c/T)|$, $|\nabla (\nu/T)^{\top}|$, and $|A_d/T|$ across or within each element, each weighted by its conjugate flux term. Since the gradients can be interpreted as measures of disequilibrium of the system, we see that a flow-controlled system is driven to a steady state position of maximum disequilibrium.

- If, on the other hand, the gradients or forces $\nabla (T^{-1})$, $\nabla (\mu_c/T)$, $\nabla (\nu/T)^{\top}$ and $A_d/T$ in (20) are fixed, the system will be driven to maximise the fluxes or rates $\vec{j}_Q, \vec{j}_c, \tau$ and $\xi_d$, again in the form of a weighted sum. In other words, the most probable response of a system to gradients or forces in one or more intensive variables $\lambda_r$ is the occurrence of flow of the corresponding physical quantities $f_r$, at their maximum mean flow rates.

This is also consistent with maximum disequilibrium of the system.

The distinction between fixed flux $\vec{j}_r$ (Neumann) or intensive variable gradient $\nabla \lambda_r$ (Dirichlet) boundary conditions is well known in the solution of differential equations. In the MaxEnt analysis of a steady state system, the choice between such boundary conditions plays exactly the same role as the choice between fixed content ($f_r$) or intensive variable constraints in an equilibrium system, or in other words, between corresponding micro-canonical and canonical representations. Although the two representations within each pair are different, the mathematical apparatus used to examine either set of constraints is the same [c.f. 45]. Flow-controlled systems can also be subject to composite flux-gradient (Robin) boundary conditions, or a mixed set of conditions, giving rise to steady states which have no equilibrium analogue in quantity-constrained systems.

To achieve maximum $\hat{\delta}$ in (20), the conjugate vectorial fluxes and gradients should be collinear. A consequence of the MEP principle is that the free fluxes or gradients will try to orient themselves to attain collinearity, to the maximum extent allowable by competition between different flows and/or by anisotropy within the system (e.g. due to the velocity gradient). Similarly, the system will endeavour to align its principal stresses with the principal directions of the velocity gradient. This may provide an explanation for the many simplifications used in control volume analysis, such as tensor symmetry.

Whilst (20) is in differential form, in systems with simple boundary conditions it may be possible to apply its finite difference approximation to much larger domains. This appears to be the case for the Earth climate system, which has been analysed (on a whole-Earth scale) using simple two-box or ten-box models [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13]. The analysis also indicates MEP to be a local principle, which need not apply to the universe as a whole. Only the entropy production of a system – not the entire universe – is maximised, since it is only within a system that there exist mechanisms by which the MEP state can be attained. This is a characteristic feature of previous applications of the MEP principle, which has incited much debate [1, 2, 3, 4, 9, 11, 50].

Tidying up the analysis, (33)-(37) can be substituted into (28)-(32). Steady state analogues of the four laws of equilibrium thermodynamics can also be obtained from Jaynes’ approach [49, 50], as set out in Appendix B. We can also consider the set of multiplier relations (3), variances (7) and covariances (5) for steady state systems, examined further in Appendix C. The analysis therefore provides a set of testable laws and transport relations for the behaviour of steady state thermodynamic systems. Furthermore, as shown in Appendix C the Onger linear regime emerges as a first-order approximation to the analysis, in the vicinity of equilibrium. The analysis therefore provides a general mathematical formalism for the analysis of flow systems – including non-equilibrium thermodynamic systems – both near and far from equilibrium.

This is as far as our comparative analysis of steady state thermodynamics can take us, using the parameters of equilibrium thermodynamics, and it is important that it not be taken too far. For example, in many systems the local equilibrium assumption – which forces the MEP principle to adopt a local formulation – may be too restrictive; such considerations lead into the domain of more complicated approaches, such as extended irreversible thermodynamics [109, 110]. For more comprehensive treatments it may be necessary to abandon the connection with equilibrium parameters, and conduct the steady state analysis using the “raw” Lagrangian multipliers $\zeta_r$ – or some functions thereof – obtained directly by MinXEnt. The raw multipliers (the “ideal” gradients) could then be correlated to the actual gradients by gradient coefficients, analogous to the activity and fugacity coefficients of equilibrium thermodynamics. In some systems, it may even be necessary to adopt process- and direction-specific time and volume scales $\theta_{rj}$ and $\nu_{rj}$ (possibly also varying with position), leading to weighted scales $\theta_{sys}$ and $\nu_{sys}$ for the system as a whole. The anal-
ysis extends naturally to further developments, for example the use of a “local steady state assumption” in the analysis of transient phenomena. This leads into a higher-order theory of acceleration phenomena, in which a maximum is sought in the sum of products of accelerating transport terms $\partial j_r/\partial t$ and gradients of the gradients $\nabla \zeta^*_r$ (a conditional, maximum $\hat{\sigma}$ principle) \cite{111}, an idea best examined elsewhere.

5. DISCUSSION

The foregoing analysis would be incomplete without a brief account of its implications. From \cite{111} MEP emerges as a local, conditional principle, in which each local control volume behaves as an actor or agent which seeks to minimise its generalised potential $\phi^*_st$, and hence (conditionally) to maximise its local entropy production. This provides a driving force for the formation and reinforcement of “emergent” self-organisation of the system as a whole, since by such “cooperative federalism”, each local “selfish” control volume can achieve much higher entropy production than it could of its own volition. We therefore see that the MEP principle drives the formation of structure and function. Of course, this does not in any way preclude the development of competition between control volumes – or even between associations of control volumes or entire systems – for a greater share of generalised potential; such competitive forces are certainly well known to us. These twin effects, a predominant, higher level cooperation and a lower level (but occasionally overwhelming) competition, are the hallmark of the “dynamic steady state” of a dissipative, flow-controlled system.

The analysis therefore confirms a number of aspects of the philosophy of Prigogine and co-workers on nonequilibrium, dissipative systems \cite{92, 93, 112, 113}, even though his “minimum entropy production principle” (valid only in the linear regime) is quite different \cite{111, 113}. It also provides an explanation for the “constructal law” of Bejan \cite{114}: “For a finite size flow system to persist in time (to survive) its configuration must evolve ... [to] provide easier and easier access to the currents that flow through it.” Furthermore, it confirms the essence of the (non-mathematical) gradient theory of Schneider and Sagan \cite{115}, who argue that “Nature abhors a gradient”; i.e. the occurrence of flow is a natural response to a physical gradient.

Finally, the analysis goes to the heart of the “riddle of life” posed by many scientists \cite[e.g.][]{116, 117}, concerning the perceived contradiction between the second law of thermodynamics and the existence of life. To recap: it is one thing to suggest that life can form, in that its ability to increase the thermodynamic entropy of the universe exceeds the reduction of entropy associated with its structure. But if life were merely an accident – a fluctuation – why should it not just die out? Why should it be so resilient to extreme events, as evidenced by its regrowth after the many mass-extinction episodes in the history of the Earth? As shown, the existence and evolution of life is driven by a deeper, purely probabilistic form of the second law \cite{71}:

“A system tends towards its most probable form.”

This “MaxProb” principle \cite{66, 67, 68, 69, 70, 71}, applied to an isolated or open, dissipative equilibrium system, drives the system towards the equilibrium position $\delta^*eq$ or $S^*$ of minimum generalised potential $\phi^*_eq$ (e.g. minimum free energy $F^*$, $G^*$ or $J^*$), since this is more probable than other forms of the system. In contrast, in a flow system it provides a driving force for the formation of complex, dissipative structures, including life, to attain the local steady state position $\delta^*_st$ of minimum generalised potential $\phi^*_st$, hence (conditionally) of maximum $\hat{\sigma}$, since this position – not the equilibrium position – is more probable than other realisations of the system. The above probabilistic statement of the second law therefore contains within it both a “force of death” and a “force of life”, associated respectively with constraints on the mean contents $\langle f_r \rangle$ or fluxes $\langle j_r \rangle$ (or their corresponding multipliers). It also implies a surprising inevitability to the evolution of life in a flow-controlled system, whenever the conditions are suitable, and indeed, of other complex systems such as fluid turbulence, biological and ecological structures, transport and communications networks, economic systems and human (or sentient) civilisation.

6. CONCLUSIONS

In this study, a clear distinction is made between (i) quantity-controlled systems, constrained by a set of mean physical quantities $\langle f_r \rangle$ and/or their corresponding multipliers $\lambda_r$, which converge towards an equilibrium position, and (ii) flow-controlled systems, as defined by a control volume (the Eulerian description), constrained by a set of mean fluxes $\langle j_r \rangle$ and/or their corresponding multipliers $\zeta_r$, which converge towards a steady state position. A theory to determine the steady state position of a flow-controlled thermodynamic system is derived using the generic MaxEnt principle of Jaynes \cite{45, 46, 47, 48, 49, 50, 51, 65}. The analysis is shown to yield a local, conditional form of the MEP principle. It also yields steady state analogues of the four laws of equilibrium thermodynamics, and sets of multiplier, variance and covariance (Maxwell-like) relations applicable to flow-controlled systems at steady state. The derivation is limited to reproducible flow-controlled systems, but does not appear to be restricted to the near-equilibrium linear regime; indeed, the latter can be recovered as a special case \cite[see Appendix C]{3}. The analysis reveals a very different manifestation of the second law of thermodynamics in steady state systems, which provides a driving force for the formation of complex, dissipative systems, including life.
Further work is required on the scope of the present analysis, and the relationship between concepts developed here and those of others; e.g. between the steady state generic entropy $S^*_{st}$, the path-based entropies of Dewar [43, 44] and Attard [56, 57], and the quantum formulation of Beretta [58, 59, 60, 61]. Greater attention should also be paid to the tremendous power of the MinXEnt / MaxEnt method pioneered by Jaynes [45, 46, 47, 48, 49, 50, 51], and its combinatorial (probabilistic) basis [62, 63, 64, 65, 66].

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APPENDIX A: TERMINOLOGY

From the foregoing discussion (especially §2-41), it is evident that there are difficulties in terminology concerning the negative Massieu function $\phi^* = -\lambda_0^* = -\ln Z_q^*$. Although described herein as a free energy-like concept or “generalised potential”, from §3 it has more in keeping with an entropy-related quantity. In equilibrium systems, $\phi^*_q$ is usually multiplied by $kT$ to give the free energy; but as shown in §3 it is more appropriate to multiply it by $k$, to avoid complications with changes in the intensive variables. The free energy concept of Gibbs [83], designed purely for changes in the extensive variables, is thus less versatile than the slightly older concept of Massieu [76], adopted by many well-known thermodynamicists e.g. [77, 78, 79] for this reason. Concepts related to the free energy, such as exergy and exergy fluxes [80, 81, 82, 83, 84, 85], suffer from the same disadvantage (a further disadvantage is their need for an unchanging reference system). In steady state systems, we could multiply $\phi^*_q$ by $kT/\theta V$ to give the “free power”, or by $k/\nabla(T^{-1})\theta V$ to give a flux analogue, but neither choice makes much sense, and would involve the same awkwardness in the handling of intensive variables as does the free energy. If $\phi^*_q$ must have units, it is preferable to multiply it by $k/\theta V$ to give it the same units as $\dot{\phi}$; being of opposite sign, we could call this (as would Gibbs [82]) the “available capacity for entropy production” or (as would Brillouin [118]) the “free negentropy production”. Provided that the dissipative processes are macroscopically reproducible, the fact that this quantity has entropy units (a non-conserved quantity) instead of energy units (a conserved quantity) will not affect its application.

APPENDIX B: STEADY STATE “LAWS”

Steady state analogues of the “laws” of equilibrium thermodynamics are readily obtainable from Jaynes’ generic approach [49, 50]. Although unsurprising, they are included here for completeness. Firstly, two control volumes which are joined (side-by-side) such that their fluxes are combined (i.e. $\langle j_m \rangle_{tot} = \langle j_m \rangle_1 + \langle j_m \rangle_2$ for $\langle j_m \rangle \in \{ \{ j_r \} \}$) will attain a steady state position of identical multipliers $\{ \zeta_m \}$, and so will share common $m$th gradients; this is the zeroth law. The first law is simply a definition of generalised heat and work for the flux of a conserved quantity, which at steady state is given by $d\langle j_r \rangle = \delta q_r + \delta w_r$; thus $\delta q_r$ gives a (spontaneous) change in the distribution of instantaneous fluxes, whilst $\delta w_r$ represents a (recoverable) change in the instantaneous fluxes themselves. The second law can be stated in many ways (see §5); in a state function sense, it is given by the generalised Clausius equality (30). The third law – arguably more of a definition of a convenient reference state – can be stated as “the steady state entropy $S^*_{st}$ approaches zero at the position of zero gradients”, i.e. when it relaxes to an equilibrium position.

APPENDIX C: JAYNES’ RELATIONS

In linear transport theory, it is common to employ the so-called “Curie postulate”, in which the scalar, vector or tensor fluxes $j_r$, $\mathbf{j}$, or $\mathbf{J}$, are assumed to depend on forces or gradients of the same type $\Delta \lambda_r$, $\nabla \lambda$, or $\nabla \lambda^i$, but not on those of other types [105, 106, 107]. This postulate cannot be assumed to apply in general. In consequence, Jaynes’ relations (6-8) and higher derivatives for scalar and each component of vector and tensor multipliers in a steady state system will be of generalised form:

\[
\frac{\partial \zeta_0^*}{\partial \zeta_{ij}^*} = - (j_{ij}^*), \quad (C1)
\]

\[
\frac{\partial^2 \zeta_0^*}{\partial \zeta_{ij}^* \partial \zeta_{kl}^*} = (j_{ij}^*)^2 - (j_{ij}^*)^2 = - \frac{\partial (j_{ij}^*)^2}{\partial \zeta_{ij}^*}, \quad (C2)
\]

11 Tribus [42] would distinguish them as “laws of thermostatics” and “laws of thermodynamics”. 

12
state positions. Note that these relations do not impose
derivative is taken at constant other Lagrangian multi-
and all directions redundant for scalars). Each partial

\[ \langle \partial \phi \rangle_{\zeta, n, p, \theta} = - \langle \partial (j_{rj}) \rangle_{\zeta, n, p, 0} - \langle \partial (j_{mn}) \rangle_{\zeta, 0, 0} \]

(C3)

\[ \frac{\partial^2 \phi}{\partial \zeta \partial \zeta} = - \langle \partial (j_{rj}) \rangle_{\zeta, 0, \tau} = - \langle \partial (j_{mn}) \rangle_{\zeta, 0, 0} - \langle \partial (j_{n, p}) \rangle_{\zeta, 0, 0} \]

(C4)

where the expectation notation \( \langle \cdot \rangle \) is reinstated;
\( t, j, k, \ell, \varphi, \theta \in \{ x, y, z \} \) (with \( j, \ell, \theta \) redundant for vectors
and all directions redundant for scalars). Each partial
derivative is taken at constant other Lagrangian multi-
pliers (see Appendix B), yielding (77):

\[ \langle j_{rj} \rangle = \sum_{m, n, p} L_{rj, m, n, p} \zeta_{m, n, p} \]

(C5)

with the equilibrium-limit derivatives:

\[ L_{rj, m, n, p} = \langle \partial (j_{rj}) \rangle_{\zeta = 0} \]

By the use of Jaynes’ method, we therefore recover the
near-equilibrium linear regime (C6) with, from (C4), the
Onsager reciprocal relations \( L_{rj, m, n, p} = L_{0, m, n, p} \)
(54, 55). For simple boundary conditions, these relations may apply
to the overall system as well as at local scales. How-
ever, the analysis goes well beyond Onsager’s, indicating that the reciprocal relations (C3) and higher derivatives
(C4) and onwards) also apply well away from equilib-
rium, at least when expressed in terms of the raw multi-
pliers (see §4.4) at local scales, although they will not then enter into the expansions (C5) - (C6). Of course,
the expansion (C5) (hence (C6)) will become more and
more inaccurate with increasing distance from equilib-
rium, even if successively higher order terms are included.
From (C4) and onwards, each set of higher order deriv-
atives will also obey a symmetry relation, both near and
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TABLE II: Multiplier relations (C1), variances (C2) and covariances (C3) for the steady state thermodynamic system of Figure 2, at steady state (here in each partial derivative can be judged by context.

Multiplier relations
\[-\langle \dot{Q}_1 \rangle = - \frac{\partial \dot{Q}_1}{\partial (K \frac{\partial (1/T)}{\partial T})}\]
\[-\langle \dot{Q}_2 \rangle = - \frac{\partial \dot{Q}_2}{\partial (K \frac{\partial (1/T)}{\partial T})}\]
\[-\langle \dot{Q}_3 \rangle = - \frac{\partial \dot{Q}_3}{\partial (K \frac{\partial (1/T)}{\partial T})}\]
\[-\langle \dot{Q}_4 \rangle = - \frac{\partial \dot{Q}_4}{\partial (K \frac{\partial (1/T)}{\partial T})}\]
\[-\langle \dot{Q}_5 \rangle = - \frac{\partial \dot{Q}_5}{\partial (K \frac{\partial (1/T)}{\partial T})}\]

Variances
\[\text{var}(\dot{Q}_1) = \frac{\partial \dot{Q}_1}{\partial (K \frac{\partial (1/T)}{\partial T})}\]
\[\text{var}(\dot{Q}_2) = \frac{\partial \dot{Q}_2}{\partial (K \frac{\partial (1/T)}{\partial T})}\]
\[\text{var}(\dot{Q}_3) = \frac{\partial \dot{Q}_3}{\partial (K \frac{\partial (1/T)}{\partial T})}\]
\[\text{var}(\dot{Q}_4) = \frac{\partial \dot{Q}_4}{\partial (K \frac{\partial (1/T)}{\partial T})}\]
\[\text{var}(\dot{Q}_5) = \frac{\partial \dot{Q}_5}{\partial (K \frac{\partial (1/T)}{\partial T})}\]

Covariances (Maxwell-like relations)
\[\text{cov}(\dot{Q}_1, \dot{Q}_2) = \frac{\partial \dot{Q}_1}{\partial (K \frac{\partial (1/T)}{\partial T})} \frac{\partial \dot{Q}_2}{\partial (K \frac{\partial (1/T)}{\partial T})}\]
\[\text{cov}(\dot{Q}_1, \dot{Q}_3) = \frac{\partial \dot{Q}_1}{\partial (K \frac{\partial (1/T)}{\partial T})} \frac{\partial \dot{Q}_3}{\partial (K \frac{\partial (1/T)}{\partial T})}\]
\[\text{cov}(\dot{Q}_1, \dot{Q}_4) = \frac{\partial \dot{Q}_1}{\partial (K \frac{\partial (1/T)}{\partial T})} \frac{\partial \dot{Q}_4}{\partial (K \frac{\partial (1/T)}{\partial T})}\]
\[\text{cov}(\dot{Q}_1, \dot{Q}_5) = \frac{\partial \dot{Q}_1}{\partial (K \frac{\partial (1/T)}{\partial T})} \frac{\partial \dot{Q}_5}{\partial (K \frac{\partial (1/T)}{\partial T})}\]

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TABLE III: Nomenclature used in this study (ES = equilibrium system; CV = control volume).

| Symbol | Meaning |
|--------|---------|
| $A_d$ | molar chemical affinity of reaction $d$ ($A_d < 0$ for spontaneous forwards reaction) |
| $C$ | total number of chemical species |
| $D$ | Kullback-Leibler cross-entropy function (generic) |
| $f_{r,i}$ | value of property $f_r$ in the $i$th category (generic) |
| $\langle f_r \rangle$ | mathematical expectation of property $f_r$ (generic) |
| $F_r$ | flow rate of property $f_r$ (CV) |
| $\mathcal{F}_{r,i}$ | instantaneous flow rate of $f_r$ (CV) |
| $g_c$ | specific body force on chemical species $c$ (CV) |
| $S$ | entropy function (generic) |
| $J_{N}e$ | instantaneous mass flux of chemical species $e$ through element (relative to $v$) (CV) |
| $\mathcal{J}_{j,\text{tot}}$ | total thermodynamic entropy flux through element (CV) |
| $\mathcal{J}_{s}$ | mean mass flux of chemical species $s$ through element (relative to $v$) (CV) |
| $\mathcal{J}_{Q}$ | mean local heat flux through element (CV) |
| $\mathcal{J}_{Q,r}$ | instantaneous value of heat flux through element (CV) |
| $J$ | free energy for system with $\langle U \rangle$, $\langle V \rangle$ and $\langle n_c \rangle$ constraints (ES) |
| $k$ | Boltzmann constant |
| $m_c$ | specific (per unit fluid mass) moles of particles of chemical species $c$ (CV) |
| $M_c$ | molecular mass of chemical species $c$ |
| $\langle n_c \rangle$ | mean moles of particles of chemical species $c$ (ES) |
| $n_{N,e}$ | moles of particles of chemical species $c$ (ES) |
| $N$ | total number of particles or entities |
| $p_i$ | probability of the $i$th category (generic) |
| $p_{i,j,(N_c)}$ | joint probability of each category in ES |
| $P$ | absolute pressure ($P > 0$ for compression) |
| $q_i$ | source (prior) probability of $i$th category (generic) |
| $\delta q_r$ | increment in “generalised heat” flux associated with $r$th constraint (CV) |
| $\delta Q$ | increment in actual (thermodynamic) heat (ES) |
| $\delta Q_r$ | increment in “generalised heat” associated with $r$th constraint (generic, ES) |
| $R$ | total number of moment constraints |
| $s$ | number of categories (generic, ES); specific thermodynamic entropy (CV) |
| $S$ | thermodynamic entropy (ES) |
| $T$ | absolute temperature |
| $t$ | time |
| $u$ | specific (per unit fluid mass) internal energy (CV) |
| $\langle U \rangle$ | mean internal energy (ES) |
| $U_i$ | internal energy level (ES) |
| $v$ | mass-average velocity through element (CV) |
| $V$ | volume |
| $\langle V \rangle$ | mean volume (ES) |
| $V_j$ | volume element “level” (ES) |
| $\mathcal{V}$ | characteristic volume length scale (CV) |
| $\delta w_r$ | increment in “generalised work” flux associated with $r$th constraint (CV) |
| $\delta W$ | increment in actual (thermodynamic) work (ES) |
| $\delta W_r$ | increment in “generalised work” associated with $r$th constraint (generic, ES) |
| $Z^*_i$ | partition function (generic) |
| $Z$ | partition function (CV) |
TABLE III: | Symbol Meaning |
|----------------|
| **Greek symbols** |
| $\alpha$, $\beta$, $\gamma$, $\delta$ | defined in [13] (ES) |
| $\epsilon$ | Kronecker delta tensor |
| $\zeta_0$ | Massieu function = Lagrangian multiplier for the natural constraint (CV) |
| $\zeta_r$ | Lagrangian multiplier for the $r$th constraint (CV) |
| $\theta$ | characteristic time length scale (CV) |
| $\lambda_0$ | Massieu function = Lagrangian multiplier for the natural constraint (generic, ES) |
| $\lambda_r$ | Lagrangian multiplier for the $r$th constraint (generic, ES) |
| $\mu_c$ | molar chemical potential of chemical species $c$ |
| $\nu_{c,d}$ | stoichiometric coefficient of chemical species $c$ in the $d$th reaction ($\nu_{c,d} > 0$ for a product) |
| $\xi_d$ | extent of chemical reaction $d$ |
| $\hat{\xi}_d$ | mean molar rate per unit volume of chemical reaction $d$ (CV) |
| $\hat{\xi}_{c,d}$ | instantaneous molar rate per unit volume of chemical reaction $d$ (CV) |
| $\Xi$ | grand partition function (ES) |
| $\Pi$ | joint probability of each category in CV |
| $\rho$ | fluid density |
| $\sigma$ | entropy produced and exported from system (ES) |
| $\hat{\sigma}$ | rate of production of thermodynamic entropy in control volume (CV) |
| $\tau$ | mean viscous stress tensor in element (CV) |
| $\tau_{ij}$ | instantaneous value of viscous stress tensor in element (CV) |
| $\psi$ | negative Planck potential for system with $\langle U \rangle$, $\langle V \rangle$ and $(n_c)$ constraints (ES) |
| $\phi$ | potential function (generic) |

**Superscripts**

| Symbol Meaning |
|----------------|
| $*$ | at a stationary position |

**Subscripts**

| Symbol Meaning |
|----------------|
| $c$ | chemical species index (ES); pertaining to the $j_c$ constraint (CV) |
| $CV$ | control volume (CV) |
| $d$ | chemical reaction index (ES); pertaining to the $\hat{\xi}_d$ constraint (CV) |
| $eq$ | an equilibrium (quantity controlled) system (ES) |
| $i$ | index of categories (generic); index of energy levels (ES) |
| $I, J$ | Cartesian coordinate indices |
| $I$ | index of instantaneous heat flux (CV) |
| $k, \ell$ | Cartesian coordinate indices |
| $L_d$ | index of instantaneous rate per volume of chemical reaction $d$ (CV) |

**Mathematical symbolism**

| Symbol Meaning |
|----------------|
| $\langle x \rangle$ | mathematical expectation of $x$ |
| $\delta x$ | total differential of $x$ |
| $\nabla$ | Cartesian gradient (vector) operator |
| $\nabla \cdot$ | Cartesian divergence (scalar) operator |
| $\var{a}$ | variance of $a$ |
| $\text{cov}(a, b)$ | covariance of $a$ and $b$ |
| $a^\top$ | transpose of $a$ |
| $a \cdot b$ | vector dyadic product |
| $A : B$ | tensor scalar product |
| $x|_{a, b,...}$ | quantity $x$ taken at fixed values of $a, b,...$ |
| $\sum_{a} x$ | sum of $x$ from $a$ to $b$ |