On entropy in eulerian thermodynamics

C. Fronsdal and A. Pathak

Department of Physics and Astronomy, University of California Los Angeles

ABSTRACT To the student of thermodynamics the most difficult subject is entropy. In this paper we examine the actual, practical application of entropy to two simple systems, the homogeneous slab with fixed boundary values of the temperature, and an isolated atmosphere in the presence of the static gravitational field. The first gives valuable insight into the nature of entropy that is subsequently applied to the second system.

It is a basic tenet of thermodynamics that the equilibrium of an extended, homogeneous and isolated system is characterized by a uniform temperature distribution and it is a strongly held belief that this remains true in the presence of gravity. We find that this is consistent with the equations of extended thermodynamics but that entropy enters in an essential way. The principle of equivalence takes on a new aspect.

I. Introduction

To avoid misunderstanding we must take care to define our context, as we have tried to do in the title of this paper; namely, thermodynamics as it is developed in textbooks, including the local extension - including hydrodynamics - that is sometimes referred to as extended thermodynamics, a eulerian field theory in which the dynamical variables are temperature, density and pressure (densities and pressures), that is responsible for the description of equilibrium configurations, as well as dynamical phenomena such as the propagation of pressure waves. Insight derived from kinetic and atomic theory is essential, but it will be used only after translation to thermodynamic field theory.

Because entropy is a difficult concept, we shall narrow the context to the comparatively simple special case of ideal gases. Experiments carried out in the beginning of the 19th century led to the discovery of power laws (Poisson 1835) that relate density, temperature and pressure,

\[ p \propto \rho^\gamma, \quad \rho \propto T^n, \quad \gamma = 1 + \frac{1}{n}, \]

valid under conditions of thermal isolation and equilibrium, in the absence of external forces. This says that \( \rho/T^n \) remains unchanged for an adiabatic process. * A perfect gas

* For example, it remains uniform and constant during sound propagation (Laplace...).
is one for which these laws are exact; for such gases it has been shown that the specific entropy can be expressed, up to an inessential additive constant, as

\[ S = -\mathcal{R} \ln \frac{\rho}{T^n}, \]  

(1.2)

where \( n \), as in Eq.(1), is the adiabatic index of the gas. The value of \( n \) for a specific gas can be derived from kinetic theory, but in our context it is a given datum.

In the case of an equilibrium configuration of a thermally isolated ideal gas, in the absence of external forces, all the fields are uniform, and the entropy is reduced to a real number that effectively parameterizes the adiabats of the system.

Questions that relate to changes in the entropy have to be phrased with care. In this paper we shall discuss entropy within the context of the equations of motion, for stationary configurations.

The equations that are used to describe an isolated, one component thermodynamic system are the equation of continuity, the Navier-Stokes equation and energy conservation. In the simpler case when dissipation and shear forces are neglected they reduce to conservation laws: the equation of continuity (conservation of mass), the Bernoulli equation (conservation of momentum) and energy conservation. To this must be added an equation of state. Here we are going to restrict the attention to equilibrium configurations, for which all the fields are time independent and the velocity field is zero. The full list of equations is then the hydrostatic equation

\[ \text{grad} \, p = 0, \]  

(1.3)

and the equations of state that in the case of an ideal gas is the gas law

\[ p = \mathcal{R} \rho T \]  

(1.4)

and the expression for the internal energy density

\[ u = n\mathcal{R} \rho T. \]  

(1.5)

The polytropic relations can be deduced from the last 2 equations if, in addition, one assumes that the specific entropy is uniform. * In general the set (1.3-5) must be supplemented with some statement about the entropy. To understand the role of entropy it is necessary to go to the foundations of thermodynamics.

* Here we assume that the entropy density \( s = \rho S \), hence attributable to the gas.
The laws of equilibrium thermodynamics of a homogeneous substance, in the absence of external forces, can be summarized as a Gibbsean variational principle,

$$\delta H = 0, \quad H = F(T, V) + ST + PV,$$

where $F$ is the free energy and $V$ is the volume. In the case of an ideal gas

$$F(T, V) = -RT \ln(VT^n).$$

Here $T, V, S$ and $P$ are independent dynamical variables, but only $T$ and $V$ are varied, to give the two principal relations

$$\frac{\delta H}{\delta T}|_{V, S, P} = 0, \quad \text{that is,} \quad \frac{\partial F}{\partial T}|_V + S = 0,$$

and

$$\frac{\delta H}{\delta V}|_{T, S, P} = 0; \quad \text{that is,} \quad \frac{\partial F}{\partial V}|_T + P = 0.$$

Given the structure of the action, there can be no question of variation with respect to $S$ or $P$; these are parameters that distinguish one equilibrium configuration from another, held fixed while $T$ and $V$ are varied.

The equations obtained by minimizing the action are equivalent to those in the standard formulation of thermodynamics in the Helmholtz representation (Legendre transform of energy representation) with the entropy always held fixed. *

In the case of an ideal gas the first of the two equations, Eq. (1.7), gives the formula

$$S = R \ln(VT^n) + nR$$

for the entropy. Contrary to the impression that may be obtained from some texts, this formula is not primarily to be understood as a means of evaluating or defining the entropy for a given thermodynamical configuration, but as an equation of motion or constraint that is to be satisfied by the dynamical variables $T, \rho$, depending on an a priori choice of $S$. The dynamics that results from a particular choice of entropy is referred to as “adiabatic”.

The local extension of the theory is far from being a trivial matter. The dynamical variables are fields over $\mathbb{R}^3$, $F$ and $S$ are interpreted as specific densities (per gram or per particle) and $V = 1/\rho$ is now the specific volume. The total energy is

$$H = \int d^3x \left( \rho \vec{v}^2 / 2 + f(T, \rho) + sT \right) + PV,$$

where

$$f(T, \rho) = \rho F(T, V), \quad \rho = 1/V.$$

* “It is perhaps superfluous at this point to stress again that thermodynamics is logically complete within either the entropy or the energy representations and self-contained within either the entropy or the energy representations and that the introduction of the transformed representations is purely a matter of convenience.” (Callen, 1985, page )
and $s$ is an entropy density. It is this last variable that is the center of interest of this paper. Provisionally we suppose that

$$s = \rho S, \quad (1.10)$$

where the specific entropy $S$ is an independent variable. Both $P$ and $S$ are parameters together their values characterize the adiabatic families of configurations. That is not to say that these parameters can be chosen at random, as we shall see. The dynamical variables are now the fields $T$, $\rho$ and $S$, while $P$ enters only as the pressure at the boundary. *

Variation of $H$ with respect to $T$ and $\rho$, with $S$ and $P$ held fixed, gives local versions of Eq.s (1.7-8). Variation of $T$:

$$\frac{\partial f}{\partial T} + s = 0. \quad (1.11)$$

In the case of an ideal gas

$$f(T, \rho) = R\rho T \ln \frac{\rho}{T^n} \quad (1.12)$$

and (1.11) becomes, under the assumption (1.10),

$$R(\ln \frac{\rho}{T^n} - n) + S = 0; \quad (1.13)$$

The specific entropy $S$ is a field, in principle arbitrary, that is held fixed under the variation of $T$ and $\rho$. An adiabatic motion is one that takes place with no change in the value of this field; that is, a motion for which $S$ is independent of the time. In the case that $S$ is uniform, the last equation is the polytropic condition $\rho \propto T^n$ and the numerical value of $S$ then parameterizes the family of adiabats. **

In the general case, local variation of the density, vanishing at the boundary, with the total mass $\int d^3x \rho$ held fixed gives, for equilibrium configurations ($\vec{v} = 0$ and all fields time independent),

$$\text{grad} \left( \frac{\partial f}{\partial \rho} + T \frac{\partial s}{\partial \rho} \right) = 0, \quad \text{or} \quad \frac{\partial f}{\partial \rho} + T \frac{\partial s}{\partial \rho} = \text{constant}. \quad (1.14)$$

It reduces to the hydrostatic condition $\text{grad} p = 0$ under a certain proviso. The definition of the thermodynamic pressure is

$$p = -\frac{\partial F}{\partial V} = \rho^2 \frac{\partial f}{\partial \rho \rho} = (\rho \frac{\partial}{\partial \rho} - 1)f, \quad (1.15)$$

* When the focus is exclusively on equilibrium configurations it may be said that independent variation of $s$ (with only the total entropy $\int d^3x s$ held fixed) leads to the desired result that $T$ is uniform, but in the general dynamical context this cannot be true; hence variation of the entropy is not to be considered in the present context.

** The expression for the free energy density can be derived from the two equations of state that characterize an ideal gas, the gas law and the expression for the $u = nRT$ for the internal energy. The expression for the specific entropy density then follows from this condition, that the hamiltonian (1.9) be extremal with respect to variations of the temperature. The fact that the same was derived by kinetic theory is evidence of the compatibility between kinetic theory with thermodynamics, if such were needed.
Using the last 3 equations and (1.11) we find

$$\nabla p = [(1 - \rho \frac{\partial}{\partial \rho}) s] \nabla T - \rho T \nabla \frac{\partial s}{\partial \rho}. \quad (1.16)$$

In the case that \( s = \rho S \), with \( S \) uniform (and independent of \( \rho \)), this leads to the simple statement (1.3),

$$\nabla p = 0.$$

This is the hydrostatic equation, the Bernoulli equation specialized to equilibrium configurations, in the absence of external forces.

In the case of an ideal gas, the pressure defined by (1.15) is \( p = \mathcal{R} \rho T \) and the internal energy density \( f + s T \) reduces, in view of (1.13), to \( n \mathcal{R} \rho T \).

The set of variations that vanish at the boundary is supplemented by variations of density and volume, the total mass held fixed. These variations yield the additional result that the thermodynamical pressure agrees with the external pressure \( P \); this is what justifies the interpretation of the quantity (1.15) as pressure.

The set of equations (1.3-5) is thus in full accord with thermodynamics, in the case that \( s = \rho S \), with \( S \) uniform and independent of \( \rho \). Going beyond the special case of an ideal gas we can always replace the equations of state by an expression for the free energy density. But to justify the hydrostatic condition it is necessary to place the following condition on the entropy:

$$[(1 - \rho \frac{\partial}{\partial \rho}) s] \nabla T - \rho T \nabla \frac{\partial s}{\partial \rho} = 0. \quad (1.17)$$

The traditional approach respects the structure of (extended) thermodynamics only to the extent that this is true. In the polytropic case all the fields are necessarily uniform in stationary configurations. The applications that we shall study in this paper are not of that kind; the entropy plays a more interesting role.

We shall try to advance our understanding of entropy by a careful examination of some simple systems for which the homogeneity of the equilibrium configurations, and the general dynamics, is disturbed by external circumstances, expressed as boundary conditions or as external forces, which shall compel us to allow for a non-uniform entropy and thus to learn how the hydrostatic equation can be justified.

Our first example, in Sections II and III, deal with the stationary conduction of heat through a slab, with the two plates maintained at constant, unequal temperature. The other example, in Section IV, examines the effect of gravity on an isolated column in equilibrium under the assumption that the temperature, at equilibrium, is uniform.

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*** The calculation: We have \( \nabla p = \rho \nabla \frac{\partial f}{\partial \rho} + \frac{\partial f}{\partial \rho} \nabla \rho - \nabla f \), and \( \nabla f = \frac{\partial f}{\partial \rho} \nabla \rho + \frac{\partial f}{\partial T} \nabla T \). Add these two and use (1.11) and (1.14 to get (1.16).
Remark. That the functional (1.9) is a fully fledged hamiltonian is seen as follows. Introduce the Poisson bracket

\[ \{ A, B \} = \int d^3 x \left( \frac{\partial A}{\partial v_i} \frac{\partial B}{\partial \rho} - \frac{\partial B}{\partial v_i} \frac{\partial A}{\partial \rho} \right); \]

then

\[ \dot{\rho} = \{ H, \rho \} = -\partial_i (\rho v_i), \]

\[ \dot{v}_i = \{ H, v_i \} = -\partial_i \frac{\partial h}{\partial \rho} \]

and, of course, \( \dot{H} = \{ H, H \} = 0. \)

II. First example. The slab

We consider the case of an ideal gas that is confined between two parallel (horizontal) plates that are maintained at constant and uniform temperatures \( T_1 < T_2 \), respectively. It is required that the temperature field satisfy the boundary conditions that make it continuous across the boundary; in particular, it can not be uniform. One can envisage a stationary configuration, not in thermal equilibrium, in which the velocity field is zero and the dynamical variables are time independent.

The salient point that attracts attention to this simple example is the fact that the boundary conditions are incompatible with a uniform temperature. Since we are definitively dealing with an ideal gas we must use the expression (1.6) for the free energy density

\[ f(T, \rho) = R \rho T \ln \frac{\rho}{T^n}; \] (2.1)

the thermodynamical pressure is thus \( p = R \rho T \) and the formula (1.2) for the specific entropy of the gas applies. In the absence of flow the Bernoulli equation implies that \( p \) must be uniform. Since the temperature cannot be uniform because of the boundary conditions, neither is the density. Then it follows from formula (1.2) that the specific entropy density is not uniform. *

The two fundamental relations of thermodynamics are Eq.s (1.11) and (1.14). For an ideal gas they are

\[ R \rho (\ln k - n) + s = 0, \quad \text{and} \quad RT (\ln k + 1) + \frac{\partial s}{\partial \rho} T = c = \text{constant}, \] (2.2)

where \( k := \rho/T^n \). Multiplying the first by \( T \) and the other by \( \rho \) and subtracting the first from the second we obtain

\[ [(n + 1)R \rho + (\rho \frac{\partial s}{\partial \rho} - s)]T = c \rho. \]

* Here we are still assuming that Eq.(1.10) holds , \( s = \rho S \), but see below.
This shows that, if $s$ is linear in $\rho$ - Eq.(1.10) with $S$ independent of $\rho$ - then the temperature is necessarily uniform, contradicting the boundary conditions. This conclusion is reached irrespective of any assumptions about the specific entropy density; letting it vary from point to point does not help.

This calls for a generalization of the assumption (1.10) and the simplest possibility is to add an extra term to the density

$$s = \rho S + S_{\text{ex}}, \quad (2.3)$$

with both $S$ and $S_{\text{ex}}$ independent of the dynamical variables. Then Eq.s (2.2) become

$$\left(\mathcal{R}(\ln k + 1) + S\right) T = c\mathcal{R}, \quad \text{and} \quad \left[(n + 1)\mathcal{R} - S_{\text{ex}}/\rho\right] T = c\mathcal{R}, \quad (2.4)$$

These two equations give the density and the temperature for any choice of the distributions $S$ and $S_{\text{ex}}$. The boundary conditions restrict the choice; since $T$ is not uniform, $S_{\text{ex}} \neq 0$.

Equation (1.3), $\text{grad} p = 0$, demands that $p$ be uniform. We can reconcile this with thermodynamics by demanding cancelation between the two terms on the right hand side of Eq. (1.16),

$$S_{\text{ex}} \text{grad} T = \rho T \text{grad} S. \quad (2.5)$$

This makes the pressure uniform. Conversely, if $p = \mathcal{R}\rho T$ is uniform, then this equation is a consequence of (2.4). In this case, since $T$ is not uniform neither is $S$.

**Summary.** Every distribution of density and temperature is possible, but when the present boundary conditions apply, then $S_{\text{ex}} \neq 0$ and $S$ is not uniform. There are two interesting special cases:

1. Maintain $\text{grad} p = 0$, accept that the specific entropy is not uniform.
2. Insist that the specific entropy be uniform, then $\text{grad} p = S_{\text{ex}} \text{grad} T \neq 0. \quad (2.6)$

The main conclusion is that the traditional treatment of the slab is consistent with thermodynamics, but that it requires a contribution to the entropy that is not linear in the density. The question remains whether this is allowed by the interpretation of entropy, as understood in statistical mechanics. We think that the answer is affirmative and attempt to justify this as follows.

The total entropy density is $s = \rho S + S_{\text{ex}}$. The term $S_{\text{ex}}$ is the entropy of the “background”, that enters the calculations only through the boundary conditions. The implication of this is that this part of the entropy stands for an interaction of the system under investigation with the background. The dynamics of the background is trivialized in this case by ignoring any effect on it that is due to the contact with our system.

The system (1.3-1.5) is incomplete. In the simplest case it is supplemented by the statement that the specific entropy is uniform and $S_{\text{ex}} = 0$. In general it must be supplemented by some information about the entropy. Once the entropy is specified Eq.s (2.4) are the equations of motion. These 2 equations come from (2.2), where the first is a modified polytropic relation and the second is an integrated form of the hydrostatic condition (1.16).
Both are fundamental equations of thermodynamics, the local extensions of (1.7-8). If the hydrostatic equation is maintained in the original form (grad $p = 0$) then, for any choice of the entropy, Eq.(2.5) must be respected.

III. Mechanical model of the slab

Interpretation and clarification of the $\rho S_{ex}$ form for entropy density

Let us summarize the notation: $S$ is the specific entropy of the gas, $s$ the total entropy density, $V$ is the total volume and $N$ is the number of particles. Let $S_T$ be the total measured entropy. $s = S_T/V$. In our considerations of the slab we had to introduce the extra variable $S_{ex}$. This implies that we are postulating an extra degree of freedom (call it $\eta$). Now $s$ will scale correctly only if $\eta$ is also scaled with $U, V$ and $N$.* In the simplest assumption, which is made in this paper, we write the measured entropy as $(S_T - S_{ex}) + S_{ex}$, where $(S_T - S_{ex})$ scales with $(U, V, N)$ while $S_{ex}$ in general scales with $(U, V, N, \eta)$.

In order to clarify the interpretation of the dynamical equations, ** we briefly take a closer look at the form of the energy $H$ that appears in the action and compare it with the expression for $U$ (the internal energy) in the standard Helmholtz formulation. The free energy of the gas is defined as the Legendre transform of its energy, $U_{gas}$, w.r.t. its entropy $S_{gas}$, $S_{gas} = V \rho S$. We have: $F_{gas}(T, \rho) = U_{gas} - TS_{gas}$, so that $U_{gas} = F_{gas} + TS_{gas}$ is the entire contribution of the ideal gas to the total energy $H$. The other term in $H$ is then the contribution of an extra system, always at the same temperature as the gas. $U_{ex} = F_{gas} + TS_{ex}$ is the only part of its energy that is varied during any thermodynamic process. Because the only work delivered by the extradegree of freedom $\eta$ to a reversible work source is $-T dS_{ex}$ and because its temperature is the same as the gas throughout, it may be interpreted as a thermal reservoir as defined in the familiar standard formulations: The 2nd order fluctuations of the extensive variables are zero. *** Alternately, the gas may be a reservoir for this extra system. The bottom line is that the work $(W)$ delivered to a reversible work source is given as: $dW = -dU_{gas} - dU_{ex} = -dU_{gas} + T dS_{gas} = -dF_{gas}$.

An example of such a system, which is considered in the next section, is an ideal gas in contact with a 1-D chain of oscillators placed in between two thermal reservoirs. In this configuration the chain acts as a reservoir and the mechanical work is done by the pressure exerted by the gas.

Thus, we have shown that a class of systems consisting of an ideal gas plus an external system (degree of freedom) that only contributes to $U$ as $TS_{ex}$, allow for a steady state (non divergent) energy flow to be set up between the two thermal reservoirs.****

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* Next sentence is a repetition. Suggest to omit it.
** Omit: that were obtained above upon demanding that the total energy is stationary under variation of $\rho$ and $T$ with both $S$ and $S_{ex}$ held constant
*** Omit the next sentence.
**** I do not understand this.
Example system: Oscillator chain connecting thermal reservoirs

In the previous section we have seen both the necessity to include $S_{ex}$, as well as its interpretation. In this section we analyze a mechanical system with one degree of freedom which can add the term $S_{ex}$ to the ideal gas entropy. A one dimensional chain of harmonic oscillators connecting point particles of equal mass in the presence of an external periodic potential (lattice) is considered. When this chain is placed in contact with an ideal gas, it imposes a constant temperature gradient on the gas.

As shown by Hu, Li and Zhao (20??) using numerical computations, the presence of the external lattice is necessary for Fourier’s heat law to be obeyed. In the absence of the lattice the steady state thermal conductivity depends on the total number of masses (oscillators) $N$ while density is held constant. *This is clearly contrary to the macroscopic phenomena since for example we do not expect the conductivity of a rod to depend on its length. The computations here follow those of Hu et. al and are performed with an external lattice potential.

Consider a chain of point masses connected by oscillators and given by the Hamiltonian:**

$$H = \sum \left( \frac{p_i^2}{2m} + \frac{k}{2}(X_i - X_{i-1} - a)^2 - U(X_i) \right)$$

Here, $U(X_i)$ is an arbitrary periodic potential with a spatial period $b$ and $\omega = \sqrt{\frac{k}{m}}$. We can write the Hamiltonian in dimensionless form with scaled parameters:

$$H = \sum \left( \frac{p_i^2}{2m} + \frac{k}{m}(x_i - x_{i-1} - c)^2 - u(x_i) \right)$$

with $c = a/b$ ($a$ is the equilibrium position).

Also, $T_i = \langle p_i^2 \rangle$, where $\langle \rangle$ denotes the time average. As in Hu et. al the energy current is defined as:

$$J = \frac{\partial x_i}{\partial t} \frac{\partial V}{\partial x_i}(x_i, x_{i+1}),$$

where $V$ is the harmonic oscillator potential.

The goal of the numerical computations is to compute the spatial variation of the entropy $S$. Note that $T$ is the only thermodynamic variable defined above in terms of the dynamical quantities. The density does not enter these considerations (this point is discussed further later***). So to find variation of $S$ we use the equation of state relating $S$ and $T$ (in the canonical formalism). For a single harmonic oscillator in contact with a thermal reservoir at temperature $T$, we can derive starting with canonical partition function:

$$\frac{S}{k} = 1 + \ln(kT) - \ln \frac{k}{\omega}$$

Here $k$ is the Boltzmann constant. This equation is obtained by using the quantum formula for the energy oscillator energy levels and then keeping terms to first order in $\hbar/\omega kT$ since our treatment is classical. The important point about the above equation is the logarithmic dependence of $S/k$ on $kT$.

* Confusing; what $N$ is this?

** Best not to use the letters $U, u$

*** Is it?
**Calculations**

Fixed boundary conditions are used for the chain. Thermostats at $T_+$ and $T_-$ are connected to the penultimate masses $(1, n-1)$ on either side. The action of the thermostats is given by the equations: $\ddot{x}_1 = f_1 - f_2 - \gamma_1 \dot{x}_1; \ddot{x}_n = f_{n-1} - f_n - \gamma_n \dot{x}_n$, with $f_i = -\partial V(x_i, x_{i-1})/\partial x_i$, $V$ is harmonic potential. $\gamma_1 = (x_1)^2 T_+ - 1; \gamma_n = (x_n)^2 T_- - 1$.

This system is solved using 8th order Runge-Kutta algorithm in MATLAB, to find $x_i(t)$. Then the temperature is found using $T_i = \langle (p_i)^2 \rangle$, where the average is taken over 10000 time units. The results for the spatial profile of $kT$ are plotted below:

![Temperature profile](image)

**Figure 1:** The vertical axis is $kT$ in scaled units on the x axis, $i$ is the mass (oscillator) number and $n$ is the total number of masses. The red line (slope = -80) was fitted to the computation results.

It is now straightforward to calculate the spatial profile of $S/k$ using the above result for temperature: (Only the $\ln(kT)$ term from the equation above for $Sk$ is plotted. The constants can be absorbed into the definition of $S/k$).

![Entropy profile](image)

**Figure 2:** Entropy Profile. Each point represents the total entropy within bin located at corresponding $i$. 
IV. Second example, the vertical column

After the initial success of hydro-thermodynamics in accounting for (or at least elegantly summarizing) the early laboratory experiments, in which the role of the ambient gravitational field could safely be ignored, it is natural to apply the theory to the conditions in the earthly atmosphere. Because this is a complicated system, in which both radiation and convection play important roles, we shall examine the somewhat academic problem of an isolated, vertical column at equilibrium. *

The role of gravity is believed to be understood and defined, if not uniquely, by the equivalence principle of General Relativity: In a non relativistic context it is enough to include a term that represents the gravitational interaction in the hamiltonian, **

\[ H = \int d^3x \left( \phi \rho + \rho \vec{v}^2 + f(T, \rho) + sT \right), \]

where \( \phi = gz \), \( z \) the vertical coordinate and \( g \) constant.

The variational equations associated with this model are precisely those of the earliest model atmospheres, with \( s = \rho S \) and the specific entropy \( S \) taken to be constant.

Variation with respect to \( T \) and \( \rho \) gives Eq.s (2.2) as before, modified only by the gravitational term, when \( \vec{v} = 0 \),

\[ \mathcal{R} \rho (\ln k - n) + s = 0, \quad \text{and} \quad \phi + \mathcal{R} T (\ln k + 1) + \frac{\partial s}{\partial \rho} T = c = \text{constant}. \quad (4.1) \]

Multiplying the first equation by \( T \) and the other by \( \rho \) and subtracting the first from the second we obtain

\[ \rho \phi + [(n + 1)\mathcal{R} \rho + (\rho \frac{\partial s}{\partial \rho} - s)]T = c \rho. \quad (4.2) \]

The situation encountered with the conducting slab is reversed. If \( s \) is linear in \( \rho \); that is, if \( s = \rho S \), whether or not \( S \) is uniform, then this implies a constant, non zero lapse rate, \( \text{grad} T = \text{constant} \neq 0 \). This constant temperature lapse is consistent with observation in real atmospheres, but it represents a failure in the attempt to describe an isolated column of gas, for it is contradicts the axiom of uniformity of the temperature - if this axiom is extended so as to apply in the presence of a gravitational field.

Indeed we have been made to understand, or at least to accept, that the equilibrium configuration of an isolated homogeneous substance is isothermal, and with especial emphasis, that this remains true notwithstanding the influence of an ambient gravitational field.

We are thus in an uncomfortable position that we may try to remedy by the same strategy that was used to account for the boundary conditions associated with slab geometry. That is, we must give up the (admittedly unmotivated) idea that the column is isentropic and instead postulate that it is isothermal, at equilibrium.

* It would seem natural to study this simplest case first, as a preliminary to developing a theory of atmospheres that are subject to additional complications.

** See for example Eckert (1987) page 260. Not the best reference.
We are led to try the strategy that led to a solution of the slab problem, setting

\[ s = \rho S + S_{\text{ex}}. \]

The equations of motion (4.1–2) are now

\[ \mathcal{R} \rho (\ln k - n) + \rho S + S_{\text{ex}} = 0 \] (4.3)

and

\[ \phi + \mathcal{R} T (\ln k + 1) + ST = c = \text{constant}, \] (4.4)

combining to give

\[ gz + (n + 1) \mathcal{R} T = S_{\text{ex}} T / \rho + c, \] (4.5)

In this case the pressure is not expected to be uniform. The thermodynamic pressure is again equal to \( \mathcal{R} \rho T \), but the hydrostatic condition (1.5) is modified by the action of gravity

\[ \text{grad} p + \rho \text{grad} \phi = 0. \] (1.5’)

As in the case of the slab, the thermodynamical relations (4.3) and (4.4) imply a modification, exactly as in Eq. (1.16),

\[ \text{grad} p + \rho \text{grad} \phi = [(1 - \rho \frac{\partial}{\partial \rho})s] \text{grad} T - \rho T \frac{\partial s}{\partial \rho}. \] (4.6)

That \( T \) must be uniform is an almost universally accepted axiom. The first term on the right hand side is therefore zero. The hydrostatic condition of the standard theory then requires that the second term vanish as well, hence \( S = \partial s / \partial \rho \) is uniform. Then Eq. (4.4) gives the standard result: the density, at equilibrium, takes the form

\[ \rho = \rho_0 = e^{-1 - \mathcal{R} T_0 T} e^{(c - gz) / \mathcal{R} T_0}. \] (4.7)

The other equation of motion, Eq. (4.5), tell us that \( S_{\text{ex}} \) can not be taken to vanish. At equilibrium, on the assumption that \( \text{grad} T = 0 \),

\[ S_{\text{ex}} = \frac{g(z - z_0)}{T_0} \rho_0. \] (4.8)

We can fix the specific entropy \( S \) (a constant) and the temperature (another constant) and calculate the external entropy (variable) and the density (also variable). But that is not how we understand Gibbs’ variational principle, or the two principal relations of thermodynamics. Instead we have been taught to fix the entropy to define the adiabatic system and calculate the extremal of the energy function with respect to variations of \( T \) and \( \rho \). Equations (4.3) and (4.4) are equations of motion to be satisfied by \( \rho \) and \( T \), for an entropy fixed in advanced; Eq. (4.8) is a constraint on the allowed choice of \( S_{\text{ex}} \) arising from the additional requirement that the temperature at equilibrium be uniform.
This makes it difficult to satisfy the axiom of uniform temperature. It may be possible to accept the idea that, given time, the entropy will adjust until a uniform temperature is achieved. But this change of entropy is required to take place, for the system under consideration, only in the presence of gravity. We thus end up with a modified set of instructions for incorporating the effect of gravity into the theory:

1. Add the term $\rho \phi$ to the Hamiltonian and
2. Add the term $S_{\text{ex}}$, precisely as in (4.8), to the entropy.

The influence of gravity is a little more complicated than expected.

It is fortunate that in actual atmospheres, there has been no need to introduce this external contribution to the entropy. The prevailing view is that simplicity is restored ($S_{\text{ex}}$ is taken to vanish and the atmosphere becomes approximately isentropic) through the agency of radiation and convection. The ideal situation, in which gravity acts alone on the isolated gas is seldom considered theoretically, nor is it seen as being worthy of experimental investigation.

Consider now a particular equilibrium state. Choose a value for $S$ and $T$ and determine the values of the distributions $\rho$ and $S_{\text{ex}}$. Having thus chosen the entropy we can approach dynamical problems, and especially the propagation of sound on the adiabatic hypothesis. Let us make the common assumption that conduction makes no appreciable contribution, then there is no need to invoke the heat equation.

Since the propagation of sound implies motion we now include the kinetic energy density $\rho \vec{v}^2$ in the Hamiltonian density. Of the two basic equations (4.3) and (4.4) only the second (the hydrostatic equation) is affected; including the kinetic energy and taking the gradient we obtain the Bernoulli equation

$$\frac{D\vec{v}}{DT} + \text{grad } p + \rho \text{ grad } \phi = S_{\text{ex}} \text{ grad } T - \rho T \text{ grad } \frac{\partial s}{\partial \rho}.$$  \hspace{1cm} (5.1)

We have assumed that the right hand side vanishes at equilibrium, and we used the additional requirement of uniform equilibrium temperature to place restrictions on the entropy. But once the entropy has been chosen in accord with this prescription all the equations of motion must continue to hold, and the second term on the right hand side of (5.1) remains zero, but the first term remains.

To evaluate the speed of sound we can use any complete set of equations of motion and the simplest set consists of the continuity equation and the two equations (4.3) and (4.4), the latter modified by the kinetic energy,

$$\dot{\rho} + \text{div}(\rho \vec{v}) = 0,$$

$$\mathcal{R} \rho (\ln k - n) + \rho S + S_{\text{ex}} = 0 \hspace{1cm} (4.3)$$

$$\dot{\vec{v}} + \text{grad} \left( \phi + \vec{v}^2/2 + \mathcal{R} T (\ln k + 1) + ST \right) = \text{constant}, \hspace{1cm} (4.4')$$
The propagation of sound is seen, since first proposed by Laplace, as an adiabatic process, and as a first order perturbation of the state of equilibrium. Without the presence of gravity it leads to the result that the speed of propagation is \( c = \sqrt{\gamma RT} \). In the case of the isolated column the equations that govern the perturbation are, to first order in \( \delta \rho \) and \( \delta T \) are

\[
\delta \dot{\rho} + \text{div}(\rho \vec{v}) = 0,
\]

\[
\frac{\delta k}{k} = S_{\text{ex}} \frac{\delta \rho}{\rho^2}, \quad \frac{\delta T}{T} = \frac{1}{n} \left( 1 - \frac{S_{\text{ex}}}{R \rho} \right) \frac{\delta \rho}{\rho},
\]

\[
\dot{\vec{v}} + \text{grad} \delta \left( RT (\ln k + 1) + ST \right) = 0.
\]

With the neglect of terms of order \( g^2 \) we obtain

\[
\ddot{\rho} = -\text{div}\left( \dot{\rho} \vec{v} \right) = RT \text{div} \left( \rho \text{grad} \left[ \left( \gamma - \frac{2g(z - z_o)}{nT} \right) \frac{\delta \rho}{\rho} \right] \right).
\]

Besides the appearance of damping, we note the dependence of the speed of propagation on the altitude.

The propagation of sound in a polytropic atmosphere (taken as a good representative of real atmospheres) is discussed by many authors, see e.g. Stanyukovich (...). It appears that there has been no study of the propagation of sound in the ideal atmosphere (uniform temperature in the presence of the gravitational field). This may be because the mere statement that \( T \) is uniform in the equilibrium configuration is not a dynamical statement that applies to adiabatic changes; it is more of the nature of a boundary condition. In this paper we have found the underlying dynamical statement, and that puts us in the position of being able to analyze questions of dynamics.

VI. Discussion

The extra contribution \( S_{\text{ex}} \) was invented, first, to understand the slab. We expect that it is uncontroversial, although the physical justification may be put into question. Using only the basic relations of thermodynamics we showed that the assumption \( s = \rho S \) is inconsistent with the boundary conditions. The analysis raise an interesting question - panel (2.6): is the pressure uniform across the slab? This should be easy an easy question to subject to observation.

The case of the isolated atmosphere in a gravitational field is different, since a physical interpretation of the extra piece in the entropy is lacking. Indeed, the physical model that was invoked in the case of the slab is much less convincing in the case of the atmosphere. We have attempted to incorporate the assumption of a uniform temperature into adiabatic thermodynamics. If the attempt is found to be unconvincing then the issue remains open. We believe that it deserves to be debated and that, perhaps, after all, the question is worthy of experimental investigation.
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