The third law of thermodynamics or an absolute definition for Entropy. Part 1 : the origin and applications in thermodynamics.

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Abstract

This article describes the third law of thermodynamics. This law is often poorly known and is often decried, or even considered optional and irrelevant to describe weather and climate phenomena. This, however, is inaccurate and contrary to scientific facts. A rather exhaustive historical study is proposed here in order to better understand, in another article to come, why the third principle can be interesting for the atmosphere sciences.

1 Introduction

Before being able to study in a second part the properties of entropy in the atmosphere, it is necessary to recall in this first part why its calculation poses certain problems in thermodynamics, problems whose solution passes through the invention and the application of the third law of thermodynamics which introduces a kind of absolute in the calculation of entropy. And the idea that certain absolutes may exist had preceded the establishment of the third law.

2 The notion of absolute temperature

Thermodynamics teaches us that the notion of temperature corresponds to the measurement of the energy of the microscopic agitations of atoms or molecules in the solids, liquids or gases which constitute the environment which surrounds us, and therefore in particular in the atmosphere.

Carnot (1824) was able to establish the existence of universal things, supposing that the perpetual motion of thermal machines was impossible. He first highlighted the existence of a maximum efficiency that depends only on the temperatures of the bodies between which these machines operate. He has also shown that the difference of specific heats under constant pressure and volume (the perfect gas constant) is independent of the nature of the bodies studied. He also gathered for the first time the two laws of Mariotte and Gay-Lussac in a single law which he expressed, in a note on page 67, by the equation:

$$P v = c (t + 267).$$

This equation reflects the fact that the product of the pressure \((P)\) by the volume \((v)\) is proportional to the temperature, the coefficient of proportionality being a constant denoted “c”. The temperature is noted here “\((t + 267)\)”, where “\(t\)” is the temperature in degrees centigrade and where “\(267\)” is an approximation of the inverse of the compressibility coefficient previously measured by Gay-Lussac (1802, page 166). We therefore recognize in the equation (1) the perfect gas law, where nowadays the constant \(c\) writes “\(nR\)”, with “\(n\)” representing the number of moles and “\(R\)” the perfect gas constant 8.314 \(\text{J/kg/mol}\).

But how did we arrive at this definition of Carnot, with this number 267, to that of a temperature called “absolute”, where “\(t + 267\)” is now replaced by the absolute temperature “\(T = t + 273.15\)”? Carnot’s work initially had no impact on the scientific community. Things changed with Clapeyron’s (1834) writing of a memoir in which he made Carnot’s works more understandable, by making more
use of mathematical language and using (page 164) the letter “R” instead of “c” in (1). It is then the English and German translations of this article by Clapeyron that will allow Carnot’s ideas to strongly influence the creation of thermodynamic science in England and Germany in the 19th century.

This was the case in England for the work of William Thomson (the future Lord Kelvin), who first knew the work of Carnot via the translation of Clapeyron’s article in 1837, before receiving a copy of the original memoir of 1824, then to make an extensive account of it in 1849 taking advantage of the rigorous new experimental measurements made by Regnault (1847) in France. This prompted Thomson (1848, page 102) to ask the question: “is there a principle upon which to base an absolute thermometric scale”, before answering: yes, by using the ideas of Carnot on the motive power of fire.

This is how Thomson defined the notion of absolute temperature. He first modified the old value of Gay-Lussac 267 in (1) by the value $1/0.00366 = 273.22$ resulting from the works of Regnault (a value he approximated by 273). Then he explained in a note (at the bottom of page 104) that “infinite cold must correspond to a finite number of degrees of air-thermometer below the zero ” (centigrade), since by lowering “$t$” reported in (1) “we should arrive at a point corresponding to the volume of air being reduced to nothing”, a point marked by $-273$ degrees “which cannot be reached at any finite temperature, however low”.

This reasoning can be illustrated by Figure 1 which suggests a prolongation towards low temperatures which seems to point to a temperature around $-273$ degrees where, for a given pressure ($P$), the volume ($v$) would necessarily tend to zero. We now use the name of William Thomson, later Lord Kelvin, to define this scale where the zero of the Celsius scale is $273.15$ degrees on the Kelvin scale. And the last words of Thomson are important, insofar as they explain that this absolute zero must be seen as an inaccessible, asymptotic limit. We will see later the importance of this same vision for the computation of the entropy defined in an “absolute” way (with an inaccessible temperature of 0 K, or asymptotically).

Does this mean that this concept of extremely low temperature and impossible to reach is easy to accept? It is quite the opposite, because one often opposes to this concept the strange and difficult side to admit that the laws which govern the changes of the gases at the usual atmospheric temperatures (say between $-100$ C and $+50$ C) can depend on something happening at $-273.15$ C!
And yet, the application of the formula \[ (1) \] with the more accurate value 273 (instead of 267) would lead to very different results if we took another absolute origin of the temperatures, as for example the value 200. Thus, let us imagine that at a constant volume \( (v) \) we increase the temperature of a gas from 5 C to 50 C. According to the formula \[ (1) \] an initial pressure of 1000 hPa would increase up to 1000 times 323/278, or 1162 hPa, if one uses the Kelvin scale, or up to the slightly different value of 1165 hPa with the old value 267 of Gay-Lussac, but up to a much higher value of 12200 hPa if we used the value 200 in \[ (1) \]. The experiment is easy to carry out, and it will give reason to the use of the absolute scale of the temperatures with the 273.15 value, which thus influences in the life of every day: the physics of the usual world “feels” what happens at zero Kelvin, around \(-273\) C.

The same “remote” influence is involved in the calculation of the efficiency of the thermal machines, which is equal to \( 1 - T_C/T_H \) and which depends on the ratio of the absolute temperatures of the systems operating between a cold source at \( T_C \) and a hot source at \( T_H \), with both \( T_C \) and \( T_H \) expressed with the Kelvin scale and not another. And the same is true for statistical physics and for the distribution of quantized states \( \exp \left[ -E/(k T) \right] \), which depends on the inverse of the temperature \( 1/T \), with the obligation to take for \( T \) the absolute definition of the temperature with an origin at \(-273.15 \) Celsius, excluding all other definitions.

3 The definition of entropy

It is again through the writings of Clapeyron, but published in German in 1843 this time, and in the knowledge of the works of Thomson, that Rudolf Clausius (1850) was able to know and use Carnot’s ideas to establish a “mechanical theory of the heat”. Clausius then introduced in 1865 (page 390) in German and in 1867 (page 357) in English, like energy, a new function which he called “entropy” and which he noted “\( S \)”. This function is defined by Clausius by the differential equation

\[
dS = \frac{\delta Q}{T},
\]

where “\(dS\)” is an elementary evolution of entropy, \(\delta Q\) is an elementary exchange of heat that occurs during a reversible transformation, and \(T\) is the Kelvin absolute temperature introduced in the previous paragraph (excluding all other possible definitions).

Then, to know the entropy \( S \) for the ambient temperature \( T \), it is necessary to integrate this equation \[ (2) \] between a certain temperature \( T_0 \) and \( T \). To do this, we must add the “infinitely small” \(dS\) between \(T_0\) and \(T\), so we need to know “\(\delta Q\)” for each temperature between \(T_0\) and \(T\). Moreover, the entropy \(S_0\) at \(T_0\) must be known. The knowledge of \(S(T)\) is therefore only relative, since it is subordinate to that of \(S_0(T_0)\) which is called “integration constant”. It appears that, at constant pressure \(p_0 = 1000\) hPa, the quantity \(\delta Q = C_p\,dT\) is proportional to the elementary evolution of the temperature \((dT)\) and to the heat capacity \(C_p(T)\), which depends a priori on the temperature. To calculate the entropy at ambient temperatures \((T)\) we must therefore compute the following sum:

\[
S(T) = S_0(0 \, \text{K}) + \left[ S_1 - S_0 \right] + \frac{L_1}{T_1} + \left[ S_2 - S_1 \right] + \frac{L_2}{T_2} + \left[ S_3 - S_2 \right] + \frac{L_3}{T_3}.
\]

The quantities \(L_1\), \(L_2\) and \(L_3\) are the latent heat of change of phase (solid/solid), solid/liquid and liquid/vapour) which are divided by the respective temperatures \(T_1\), \(T_2\) and \(T_3\). The differences \([S_1 - S_0]\), \([S_2 - S_1]\) and \([S_3 - S_2]\) represent the mathematical integrals of \(C_p(T)/T\) between the temperature limits \(T_0\) and \(T_1\), then \(T_1\) and \(T_2\), then \(T_2\) and \(T_3\).

1 There are 3 solid phases \((\alpha, \beta, \gamma)\) and 2 solid/solid changes for O2; 2 solid phases \((\alpha, \beta)\) and 1 solid/solid change for N2. See the diagrams for \(c_p(T)\) in the Fig.1 of Marquet and Geleyn (2015), available in arXiv: [https://arxiv.org/pdf/1510.03239.pdf](https://arxiv.org/pdf/1510.03239.pdf)
This method of calculating the entropy of all bodies is called “calorimetric” because we have to integrate the values of $C_p(T)/T$, where the specific heats $C_p(T)$ must be measured with “calorimeter” devices. The difficulties in calculating $\Delta S$ are therefore important. We must first know the values of the latent heats $L_1$, $L_2$ and $L_3$, also those of $C_p(T)$ for each of the temperatures between 0 K and $T$, including for the temperatures (very) close to 0 K. Finally, we need to know the value of the “integration constants” $S_0$ at 0 K, which must a priori depend on each given chemical species.

4 The importance of integration constants

The problem of these “integration constants” was clearly posed for the first time by Le Chatelier (1888) in the context of the determination of chemical equilibria. His goal was to use the characteristic functions introduced by Massieu (1869, 1876), then modified and popularized by Gibbs (1876-78) after his passage in Paris. It was a question of calculating the variation of the free enthalpy ($\Delta G$). It is one of the characteristic functions defined by Gibbs, whose variation induced by any chemical reaction at constant temperature is written in the form:

$$\Delta G = \Delta (H - T S) = \Delta H - T \Delta S.$$  \hspace{1cm} (4)

$$\Delta G$$ is to be computed for chemical reactions such as

$$N_2 + 2 O_2 \iff 2 NO_2,$$  \hspace{1cm} (5)

the symbol “$\Delta$” in (4) refers to the difference of the quantities $H$ and $S$ evaluated for the right-hand side of (5), i.e. for NO$_2$, minus those evaluated for the member on the left, i.e. for N$_2$ and O$_2$, with weighting factors depending on the stoichiometric coefficients (here: $-1$ and $-2$ for N$_2$ and O$_2$; $+2$ for NO$_2$).

Gibbs’ contribution is to show that the sign of $\Delta G$ determines the equilibrium and meaning of chemical reactions such that (5): reaction in equilibrium if “$G$” is minimal and if $\Delta G = 0$, move to the right if $\Delta G < 0$ (product NO$_2$ in majority), move to the left if $\Delta G > 0$ (reactants N$_2$ and O$_2$ in majority).

However, since the entropy variation $\Delta S$ in (4) is multiplied by the temperature $T$, the presence of the integration constants $S_0(0 \text{ K})$ in (3), which depends both on the reactants and the products, induce an impact which then depends on this temperature $T$. And because of this, the sign of $\Delta G$ in (4) is modified by all the changes of the values of the integration constants $S_0$ which are, a priori, specific to each of the components of (5). For this chemical reaction (5), the remaining sum in factor of $T$ writes: “$2 S_0(\text{NO}_2) - S_0(\text{N}_2) - 2 S_0(\text{O}_2)$”. For this reason, the nature of the equilibrium of the chemical reactions depends on the values of $S_0$ for each of the products and reactants. And that is why the thermochemistry tables do not treat $\Delta H$ and $\Delta S$ equivalently. The enthalpies of reaction $\Delta H$ are directly given for all usual bodies, the integration constants for $H$ having no chemical impact. Differently, these tables give the absolute values of the entropies $S$, which must be summed for all the products to the right of (5), then for all the reactants to the left of (5), before making the difference of these two sums and multiplying by $T$, to form the quantity “$-T \Delta S$”.

Clausius (1865 pages 392-397, 1867 pages 359-363) and Gibbs (1876-78, pages 151-152) mentioned the existence of integration constants for energy, entropy and potentials (characteristic functions), but without noting a possible impact on physical or chemical processes. It is Le Chatelier who first explained in 1888 in the chapter XI entitled “Integration constant” (page 184) that, for entropy: “The determination of (…) this constant of integration (…) would bring complete knowledge of the laws of equilibrium. It would make it possible to determine, a priori, independently of any new experimental
data, the complete equilibrium conditions corresponding to a given chemical reaction”. Indeed, if we could define $\Delta S$ up to an arbitrary constant depending on each species, we could, after multiplication by $T$, modify the sign of $\Delta G$, and thus the sense of equilibrium of (5) at will, and thus remove all oxygen molecules from the atmosphere in favour of nitrogen dioxide!

5 Values of the integration constants

The first answer to the question clearly posed by Chatelier was provided by the Nernst’s “Heat Theorem” (“Wärmesatz” 1906 in German, 1907 in English, “Théorème de la chaleur” 1910 in French).

![Figure 2: Graphical translation of the Nernst’s theorem as expressed in his articles of 1906, 1907, 1910, but with modern notations. The curves for the change in free enthalpy $\Delta G$ and in enthalpy $\Delta H$ are plotted in ordinate as a function of the absolute temperature on the abscissa.](image)

The essence of Nernst’s theorem is illustrated in Figure 2. Experimental facts show that it is the situation described in (b) that is observed. It appears first that $\Delta G$ and $\Delta H$ tend toward a common value for temperatures tending towards absolute zero, and therefore the product “$T \Delta S$” (the gap between the curves of $\Delta G$ and $\Delta H$) tends to zero both in (a) and (b). The more important and new feature is that the two functions $\Delta G$ and $\Delta H$ converge towards each other like in (b) with the same horizontal tangent. This means that the case (a) where $\Delta S$ tends toward a constant $A$ (inducing a linear variation $AT$ for $T \Delta S$) is not observed. These facts observed by Nernst led him to state a theorem which indicates that the heat capacity $C_p$ must vary with temperature so that $\Delta S$ must tend to zero “fast enough” for $T$ approaching the absolute zero (namely faster than a linear law in $T$).

Einstein (1907) was able to reinforce this prediction by establishing a theoretical formula for the heat capacity $C_p$ of the solids which possessed this behavior, via a decay of the type “$x^2 \exp(-x)$” when $1/x$ (proportional to $T$) tends to zero (and $x$ tends to infinity). However, this formulation did not agree with the experimental data because the decay was too fast with temperature. This formulation has been improved by Nernst and Lindemann (1911) by making the half-sum with the first harmonic: “$0.5 \left[ x^2 \exp(-x) + (x/2)^2 \exp(-x/2) \right]$”. However, this formulation was not yet fully consistent with the observations, and its introduction seemed a little bit ad-hoc.

It is precisely to better understand these disagreements on the calculations of the heat capacities
$C_p$ at low temperatures close to absolute zero that the first Solvay congress was organized in 1911 (see the report by Paul Langevin and Maurice de Broglie, 1912). It was Nernst who suggested this international meeting to Ernest Solvay, who gave him carte blanche to invite the greatest scientists of the time. During this congress, the new formulation of Nernst and Lindemann was criticized by Lorentz and Einstein, the latter (page 302) also criticizing Nernst’s theorem by saying: “we can not deduce (it) from the fact that specific heats ($C_p$) disappear near absolute zero”; even so, in his eyes: “its legitimacy becomes more likely”. And Einstein hypothesized that “Nernst’s theorem amounts to stating the hypothesis, moreover quite plausible, that a change (...sufficiently close to absolute zero ...) is always possible in a purely statistic way, from the point of view of wave mechanics”.

And indeed Debye (1912) found, on the basis of reasoning using quanta and statistical physics, the correct theoretical formulation for the heat capacities $C_p$ of solids, showing (page 800) that: “for sufficiently low temperatures the specific heat becomes proportional to the third power of the absolute temperature”. In doing so, we can calculate via (3) the entropy of the non-metallic solids, which depends on the cube of the temperature and leading to the internal energy being in fourth power of the temperature, in agreement with the Planck’s law of radiation.

Independently, Nernst (1912) responded to Einstein’s criticisms by justifying his 1906 theorem differently, assuming that “(...) der absolute Nullpunkt (...) nicht zu erreichen”, namely that “the absolute zero cannot be reached”, which led to the “principle of unattainability of the absolute zero” clearly expressed by Simon (1927). We recognize the same idea expressed by Thomson (Kelvin) in 1848 on the point marked by $-273$ degrees: “which cannot be reached at any finite temperature, however low”. This property is analogous to the new Nernst’s general principle expressed in 1912, and it corresponds to an experimental truth that has never been denied until today.

![Figure 3](image)

**Figure 3:** The entropy curves ($S$) are plotted as ordinates as a function of the absolute temperature ($T$) as abscissa. The aim is to provide a graphical illustration of the principle of “unattainability of absolute zero” as expressed by Nernst in 1912 and Simon (1927). This principle means that $S_{01} = S_{02}$ in (a). Then, Planck (1917) postulated that this common value is a universal value $S_0$ like in (b) for the more stable solid state of all species at $T = 0$ K. The next contribution of Planck (1917) was to postulate a zero entropy $S_0 = 0$ for $T = 0$ K for all species, without loss of generality (the yellow arrow).

This principle can easily be understood from the two diagrams in Figure 3. We find a situation in (a) where we can reach the absolute zero in three steps: an isentrope up to the point 1 followed by an isotherm up to the point 2, then a last isentrope up to the point 3. This situation corresponds to
two formulations for the entropy of a system having an internal parameter with two distinct states $X_1$ and $X_2$ (for example magnetization), and with different entropies at zero Kelvin ($S_{01}$ different from $S_{02}$). But only the situation described in (b) is in agreement with the experimental facts, because if the entropy $S_0$ is the same at zero Kelvin for all the parameters, then it obviously takes an infinity of steps to get closer and closer to the zero Kelvin. We deduce that the entropy of a body cannot have several distinct values at zero Kelvin, and that $S_{01} = S_{02} = S_0$ like in (b). More recently, Massanes and Oppenheim (2017) consider that they have proved, in general terms, this “principle of unattainability of absolute zero”.

6 The third law of thermodynamics

Max Planck (1870) passed his thesis on the second principle and the applications of entropy, less than 5 years after the paper of Clausius. Moreover, Planck (1900, 1901) applied Boltzmann’s (1877) little-known ideas very soon, to express the formula of the black body thanks to a calculation of the entropy of the radiation written in the form:

$$S = k_B \log(W) + S_0 (0K).$$

In this formula (6) derived by Planck, the entropy $S$ depends on the logarithm of the number $W$ of configurations of the system, a formula that Boltzmann had suggested differently (see the English translation in Sharp and Matschinsky, 2015) and in two stages. Boltzmann computed on the one hand the “permutability measure” $\Omega = \log(W) + C_1$ close to his equation (34) page 403, with an interest mentioned by Boltzmann to set $C_1 = 0$, and he computed on the other hand the change in entropy $S - S_2 = K \Omega + C_2$ close to his equations (62) and (65) on pages 427-428, where Boltzmann mentioned a proportionality coefficient ($K = 2/3$ with certain units $R = 1$ and for a monoatomic gas) corresponding to $“k_B”$ in (6). This formula was obtained by Boltzmann “with a suitable choice of constant” $C_2 = 0$, leading to (6) for $K = R/N = k_B$. The entropy of Boltzmann was thus $S = K \ln(W) + S_2 + C_2 + K C_1$, and the formula of Planck (1917) corresponds to $S_0 = S_2 + C_2 + K C_1$. Planck then evaluated in his two articles of 1900 and 1901 (about the black-body radiation law) both the numerical value of the constant “$h$” (which bears his name) and of the constant “$k_B$” of (6), giving it the name of “Boltzmann constant” in honour of the magnificent (but often obscure) work of Boltzmann.

In formula of Planck (6) appears the constant $S_0 (0K)$ which has disappeared from the famous Planck formula inscribed on Boltzmann’s tomb: “$S = k \log W$”. This constant $S_0$ corresponds to absolute zero temperature where, by hypothesis, all the statistical microstates are degenerated into a unique configuration of maximum probability $W = 1$, leading to $\log(W) = 0$ by definition of the logarithm, and therefore to $S = 0$ if the constant $S_0$ is set to zero.

Planck was in fact a specialist in the computation of entropy in all areas of physics, and he was able to deliver a synthesis of his vision of thermodynamics in his 1917 treatise. Planck added, in the third German edition, and even more in the fifth edition, a chapter on the third law where he wrote in the preface: “The Nernst theorem in its extended form has in the interval received abundant confirmations and can now be regarded as well established.”

The current “extended form” of the third principle is the one introduced by Planck (1917, page 274) which admits, in two stages, that the absolute value of the thermodynamic entropy of any homogeneous solid at 0 Kelvin is: (i) “a universal constant”; (ii) “that one can set to zero without loss of generality”. This second action (ii) corresponds to the yellow arrow in the Figure (6)(b). Note that this choice corresponds to the cancellation of the additive constant $S_0$ in the Planck-Boltzmann statistical formula (6). An important caution is that this definition only concerns solids, and is therefore not applicable to perfect gases.

By a lack of luck, the water which constitutes an important component of the atmosphere possesses
The dry air entropy is set in "law of thermodynamics. The water vapour entropy is set in "

Table 1: Standard molar entropies (in cal/K/mol, under one atmosphere and at 298.15 K) for: Nitrogen (N₂), Oxygen (O₂), Water vapour (H₂O), Carbon dioxide (CO₂) and Argon (Ar). The results are given for ST12 (Sackur, Tetrode, 1912); LG17 (Lewis and Gibson 1917); LR23 (Lewis and Randall, 1923); K32 (Kelley, 1932); G34 (Gordon, 1934); GR96 (Gokcen and Reddy, 1996) and C98 (Chase, 1998)

|       | ST12/S | LG17/C | LR23/C | K32/C | G34/S | GR96/C | GR96/S | C98/S |
|-------|--------|--------|--------|-------|-------|--------|--------|-------|
| N₂    | 45.59  | 45.6   | 45.8±0.3 | 45.94±0.2 | 45.78 | 45.796±0.005 |
| O₂    | 48.23  | 48     | 49.1±0.1 | 49.12±0.1 | 49.02 | 49.031±0.008 |
| H₂O   | 51.1   | 50.0±2 | 45.1   | 44.31 | 45.12 | 45.132±0.010 |
| CO₂   |        |        |        | 51.13±0.1 | 51.09 | 51.098±0.029 |
| Ar    | 37.0   | 36.43  | 36.4   | 36.96±0.2 | 37.00 | 37.000±0.001 |

the singular property of having a residual entropy still present at 0 K. The work of Pauling (1935)
completed by Nagle (1966) was used to estimate the value at \( S_0 = 0.82 \text{ cal/K/mole} \), by calculating
the impacts due to the transfer of charges via the hydrogen bonds still at work at 0 K within the molecule
H₂O. So, just for water, you need to include that difference \( S_0 \) to have a reference that is “the most
stable” at zero Kelvin.

As a conclusion, the third principle corresponds to setting \( S = 0 \) for the “more stable state” of all
solid states at \( T = 0 \text{ K} \). This is equivalent to set \( S_0 (0 \text{ K}) = 0 \) in the formula \( W \) of Planck, leading to the Boltzmann’s tomb formula: “\( S = k \log W \)

It is important to note that the caloricimetric values of entropy (calculated by integrating “\( C_p(T)/T \)”) are consistent with calculations made from quantum mechanics and statistical physics (computations of \( W \)). Indeed, the work of Tetrode and Sackur conducted from 1912 (see Grimus, 2013) resulted in a purely theoretical calculation of the entropy for monoatomic gases using the Boltzmann-Plank equation (6) with \( S_0 = 0 \) and by a meticulous computation of the number of complexions \( W \) for the quantified energies related to the translation movements of the atoms. The same theoretical work was then carried out for the more complex molecules for polyatomic gases such as steam water (Gordon 1934), taking into account rotational and possible vibrational motions.

This equivalence can be verified by comparing the entropy measurements made with the experimental
and theoretical values with reference to the table in Marquet’s Appendix A (2017), an extended version
of which is given here in Table 1. The experimental method (calorimetric, /C) is obtained by calculating
\[ \log W \text{ with } S_0 = 0 \text{ and a careful calculation of } W \text{ for quantized energies for translational motion, rotation or vibration, depending on the nature of the atoms or molecules.} \]

We see in Table 1 that the agreement between experimental (calorimetric /C) and theoretical (quantum and statistical physics /S) values improves with time (from 1912 to 1998). This was largely due to measurements made by Giauque (1949) of values of \( C_p(T) \) for most known species and for values very close to absolute zero. This work was sanctioned by a Nobel Prize (see Tiselius, 1949). Giauque insists on the agreement of his measurements with the third principle, which together, as imagined by Le Chatelier in 1888, make it possible to predict the stability regimes of all chemical reactions.

It may be mentioned that the third principle is only partially used in some official definitions of thermochemical tables used in atmospheric and oceanic studies, such as those of IAPWS (2010) and Feistel et al. (2010), with however an explicit mention of the possibility to use the third law expressed in Feistel and Wagner (2006) for the water Ice-Ih, and also by Lemmon et al. (2000) who used a third-value for the reference entropy of the dry air formed of a mixture of N₂, O₂ and Ar.² It is therefore

² It is easy to modify the TEOS-10 software in order to compute the absolute entropy of moist air based on the third law of thermodynamics. The water vapour entropy is set in “init_iapws95” in “Flu_1.F90”, where the present value \( n_{i}^{0} = 8.32044648374969 \) can be replaced by \( -15.94 \), with a difference given by the ratio \( -s_{i}^{0}/R \approx -3517/461.52 \approx -7.62 \). The dry air entropy is set in “init_iapws10” in “Air_1.F90”, where the present value \( n_{i}^{0} = 9.7450251743948 \) can be
easy to modify two lines in the software TEOS-10 of the IAPWS to agree with the third law.

Some of the values of the entropies listed in Table 1 have been recalculated in Marquet (2015), the aim being to validate the experimental values used to calculate the thermal enthalpy with respect to absolute zero. The agreement was correct for the entropies of N\(_2\) \((46.0\pm0.2)\) and H\(_2\)O \((45.2\pm0.1)\), with a larger value for O\(_2\) \((49.7\pm0.4)\). This larger value is obtained by taking into account the second order transition forming a kind of Dirac peak for the \(C_p\) of O\(_2\) in solid phase at 23.85 K. I was able to observe on this occasion that the biggest error in the value of the entropy comes from the terms depending on the latent heats of phase change, and much less of the integrals of \(C_p(T)\) from 0 K to 273.15 K.

Finally, let us recall the Schrödinger’s synthesis (1944, pages 15-17, “Discussion of the Nernst theorem”) read during his seminars in Dublin where he taught courses in statistical physics: i) the important thing for this integration constant (i.e. \(S_0\)) is not the zero value set by Planck; ii) the important thing is that it is a quantity that is independent of all the physical parameters (temperature, pressure, magnetism, electric charge, ...) and also of the nature of the bodies in general; iii) this constant is, in the end, independent of all the bodies and it is a quantity that we can set to zero without loss of generality, as Planck did, just to avoid any confusion and temptation to choose another possibly variable and inappropriate value.

7 Conclusions

Mistrust of the validity of the third principle of thermodynamics continues. This is the reason why it seemed important to recall why the greatest scientists, such as Nernst, Einstein and Planck as early as 1911, till Schrödinger in 1944, considered this scientific principle as verified by all the experimental facts and in agreement with the best theoretical developments based on statistical physics and quantum mechanics.

This article will form the basis for a second part (Marquet, 2019) which will present the different ways of calculating the moist-air entropy of the atmosphere consisting of a mixture with variable proportions of dry air, water vapour, cloud liquid water, cloud ice and liquid or solid precipitations.

It will be seen in the second part that the same questions and doubts about the validity of the third law, with a possible influence of a hypothesis made at the absolute zero of temperatures, are at work in meteorology and in the climate sciences. However, the entropy defined with the third law corresponds to new, singular and very interesting properties, and should be taken into account to better understand certain properties of the moist-air atmosphere consisting of a gas with a variable quantity of water vapour.

8 References

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