30. THE SPECTRUM OF Si\(^+\) (ONCE IONISED SILICON)

\textit{(Nature, 116, 644, 1925)}

Prof. A. Fowler has recently shown that the spectrum of Si\(^+\) is similar in constitution to that of Al, that is, consists of doublets having \(2p_\alpha, 2p_\beta\) as the highest terms. A number of lines, ascribed by Fowler to Si\(^+\), remain, however, well outside his scheme of classification, and some of these can be arranged in a group of quartets.

**Multiplet I**

<table>
<thead>
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<th>(j)</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>(0)</td>
<td>17154-38 .. 116-0 .. 17038-38 62-19</td>
<td>62-25 (1)</td>
<td></td>
</tr>
<tr>
<td>(2)</td>
<td>17216-57 .. 115-94 .. 17100-63 .. 199-89 .. 16900-74 134-56 ..</td>
<td>134-96 (1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>17255-19 .. 199-49 .. 17035-70 (1)</td>
<td>(3)</td>
<td></td>
</tr>
</tbody>
</table>

Interval ratio

\[\frac{116-00 : 199-89 = 2:9 : 5}{3 : 5} \text{ observed}\]
\[\text{(Landé)}\]

\[\frac{62-20 : 134-76 = 2:3 : 5}{3 : 5} \text{ calculated}\]
\[\text{(Landé)}\]

This is a \(pp^\prime\)-combination.
The intensity rule is only roughly obeyed.

**Multiplet 2**

<table>
<thead>
<tr>
<th>(j)</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1)</td>
<td>18382-63 .. 116-48 .. 18266-15 339-49</td>
<td>339-11 (0)</td>
<td></td>
</tr>
<tr>
<td>(0)</td>
<td>18043-14 .. 116-10 .. 17927-04 .. 199-86 .. 17727-18 (2)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This seems to be an intercombination between doublet \(P\) and quartet \(p\)-terms. But none of the known doublet \(P\)'s has 339 as frequency difference.

The differences occurring in these multiplets occur in other pairs which have not yet been classified, \textit{e.g.}

\[
\begin{align*}
(0) & : 18402-19 — 18286-73 = 115-46
\end{align*}
\]

\[
\begin{align*}
(0) & : 18415-41 — 18280-08 = 135-33
\end{align*}
\]

\[
\begin{align*}
(1) & : 20573-45 — 20373-87 = 199-58
\end{align*}
\]

The group having the successive differences 199 and 116 correspond to \(p\)-terms of a quartet series. Evidently a quartet series is possible for Si\(^+\), but it is only feebly developed under the usual methods of excitation. Without more data on the spectrum of Si\(^+\), it is not possible to find out more information on the point.

Allahabad University.

31. ON THE ABSOLUTE VALUE OF ENTROPY*

Meghnad Saha \& Ramanikanta Sur

\textit{(Phil. Mag., Sr. VII, 1, 279, 1926)}

According to Boltzmann, the entropy of a thermodynamical system is represented by the equation

\[S = k \log W \quad \ldots \quad \ldots \quad \ldots \quad (1)\]

\(S = \text{entropy}, \quad k = \text{Boltzmann's gas constant}, \quad W = \text{probability of the state.}\)

There are different ways of calculating the probability \(W\) for different thermodynamical systems. In previous years, \(W\) was used in a relative sense, in terms of some standard state. By \(W\) was meant the mathematical probability, hence it was always a fraction. Moreover, it remained indeterminate to the extent of an additive constant.

To Planck we owe the conception of the "Thermo-
ON THE ABSOLUTE VALUE OF ENTROPY

This is proportional to the mathematical probability, but not quite equal to it. The mathematical probability is a fraction, while the thermodynamical probability is a whole number. Planck has developed methods for calculating the thermodynamical probability of different systems—e.g., a perfect gas and black body radiation. As is well known, this idea led, in the hands of its author, to the development of the Quantum theory, which is now responsible for progress along all lines in physical science.

There are, however, a few points which are not yet clear. Some of these refer to the fundamental assumptions of the theory, others refer to the mode of application of the fundamental ideas. The exact nature of these obscure points can only be made clear by reference to the actual working, which we now proceed to do.

§ 1. Theory of Perfect Gases

Let us take a large number \( N \) of molecules enclosed within a volume \( V \) in the gas-kinetic sense. To calculate the probability of the system, the volume is divided into a number of cells denoted by \( 1, 2, 3, \ldots, n \), containing \( N_1, N_2, N_3 \ldots N_n \) molecules respectively. The thermodynamical probability is defined as the total number of complexes (i.e., the total number of ways in which this distribution can be effected). It is easy to see that

\[
W_n = \frac{N!}{N_1!N_2!\ldots N_n!} \quad \ldots (2)
\]

The mathematical probability

\[
W_m = \frac{W_n}{\Sigma W_n} = W_n n^{-x}. \quad \ldots (2.1)
\]

The summation \( \Sigma W_n \) is taken over all positive values of \( N_1, N_2, \) etc. consistent with the condition

\[
N_1 + N_2 + \ldots N_n = N.
\]

Let

\[
\frac{N_1}{N} = w_1, \quad \frac{N_2}{N} = w_2, \ldots
\]

Then with the help of Stirling’s formula, it can be easily shown that

\[
\log W_n = -N\sum_{r=1}^n w_r \log w_r \quad \ldots (3)
\]

The actual or equilibrium value of \( \log W_n \) is obtained by making it maximum subject to the prescribed conditions. Let \( \epsilon_r \) = average energy per molecule in the cell \( r \). Then the total energy

\[
E = N_1\epsilon_1 + N_2\epsilon_2 + \ldots N_n\epsilon_n = \Sigma \epsilon w_r = \text{const.} \quad \ldots (4)
\]

and

\[
\Sigma w_r = 1. \quad \ldots (5)
\]

From equations (3), (4), (5), we can deduce that in case of equilibrium

\[
w_r = a e^{-\beta \epsilon_r} \quad \ldots (6)
\]

From the relation \( \frac{\delta S}{\delta E} = \frac{1}{T} \) it can be shown that

\[
\beta = \frac{1}{kT} \quad \ldots (7)
\]

From equations (5) and (6)

\[
a = \frac{1}{\Sigma e^{\epsilon_r/kT}} \quad \ldots (8)
\]

and it can easily be shown that the free energy

\[
F = kN T \log a; \quad (9)
\]

\( a \) however, remains indeterminate, and it cannot be calculated without the introduction of some further hypothesis.

By an application of Liouville’s law, Planck shows that for a system obeying the canonical laws of Hamilton the motion is completely defined if the positional \( (q) \) and the momenta \( (p) \) coordinates corresponding to each individual degree of freedom of each particle are given. In this case he shows that the phase integral

\[
H = \ldots \int \ldots dq dp dq dp dq dp \ldots dq dp \quad (10)
\]

(the integral being taken for all the degrees of freedom for a particle), remains unchanged by subsequent events. \( H \) is known as the extension of the elementary region of the phase space. According to classical theory, \( H \) may have any infinitely small value up to zero, but according to the quantum theory \( H \) has always got a finite, though small, invariant value.

The summation \( \Sigma e^{\epsilon_r/kT} \) can now be effected.

\[
\Sigma e^{\epsilon_r/kT} = \frac{\Sigma e^{\epsilon_r/kT}}{H} dq dp dq dp dq dp, \quad \ldots (11)
\]

For monatomic gases, \( H \) must refer to the representative particle in the cell \( r \), having the coordinates \( x_r, y_r, z_r \), and velocity components \( u_r, v_r, w_r \).

Taking

\[
\epsilon_r = \frac{1}{2}(u_r^2 + v_r^2 + w_r^2) + \epsilon_0, \quad \ldots (12)
\]

where \( \epsilon_0 \) denotes the unalterable internal energy of an atom at rest, and

\[
dq dp dq dp dq dp = m^2 du dv dw, \quad \ldots (13)
\]

We obtain

\[
\Sigma e^{\epsilon_r/kT} = V e^{\epsilon_r/kT} (\frac{2\pi m kT}{e})^{\frac{n}{2}} \quad \ldots (13)
\]

With this value of \( \Sigma e^{\epsilon_r/kT} \), neglecting \( \epsilon_0 \),

\[
S = kN \left[ \frac{V}{H} (2\pi m kT)^{\frac{n}{2}} \right] + \frac{3}{2} kN \quad \ldots (14)
\]

\(^1\) The subject matter of this section is merely an abstract of §113-§134 of Planck’s ‘Warme Strahlung,’ 5th edition.
\[
S = kN \log \left( \frac{V T^\frac{3}{2}}{\rho N} \right) + i,
\]  
\[\ldots (14.1)\]
where \(i\) is independent of \(T\) and \(V\), but depends upon \(N\). As far as the calculation of gas laws and specific heat is concerned, the exact value of \(i\) is immaterial; but if we wish to study the chemical behaviour of the gas, the vapour pressure, etc., the knowledge of \(i\) is essential, as was first pointed out by Nernst in connexion with his Heat Theorem.

Now to show how \(i\) depends upon \(N\) we take \(p\) identical vessels, each of volume \(V_i\), containing the same quantity of the same gas side by side. Then suppose the sides be made to collapse suddenly and the \(p\) volumes are made to mix with each other. No change has been made in the system. The entropy can be calculated by adding the entropies of the \(p\) vessels, each of volume \(V_i\). Thus for the entropy of the whole gas occupying the volume \(pV\) we have
\[
S = kNp \log \left( \frac{V T^\frac{3}{2}}{\rho N} \right) + pi(N),
\]  
\[\ldots (14.2)\]
and this must equal
\[
kNp \log \left( \frac{V T^\frac{3}{2}}{\rho N} \right) + i(pN),
\]
which is the entropy of a gaseous mass of \(\frac{pN}{\rho}\) particles occupying the volume \(V\).

Hence
\[pi(N) = kNp \log \rho + i(pN).\]
The above relation is satisfied if
\[i(N) = -kN \log N.\]
Thus we can deduce from the classical theory that
\[
S = kN \log \left( \frac{V}{N} T^\frac{3}{2} \right) \quad \ldots (14.3)
\]
where \(A\) is independent of \(N\), \(T\), and \(V\).

From Planck’s theory, we have
\[
S = kN \log \left( \frac{V}{H} (2\pi mkT)^\frac{3}{2} \right) \quad \ldots (14.4)
\]
Comparing the two expressions, we find that \(H\) must vary as \(N\). Planck puts \(H = Nh^3\). But this is, however, not quite clear from the expression for \(H\), for
\[
H = \left\{ dq_1 dp_1 dq_2 dp_2 dq_3 dp_3 \right\} = h^3,
\]  
\[\ldots (15)\]
if we put, according to the canons of the quantum theory,
\[\left\{ dq dp \right\} = h \text{ (Planck’s constant).}\]

Sackur and Tetrode\footnote{Ehrenfest and Trkal, *Proc. Amst. Akad.* xxiii. (1920).}, who were the first to calculate the value of \(i\), proceeded in a different way. According to Sackur the thermodynamic probability is not equal to our \(W_n\), but it is equal to
\[
\frac{W_n}{N!} \quad \text{or} \quad \frac{1}{N!N_1!N_2!N_3! \ldots N_n!}
\]
\[
= k \log W.
\]
The value of \(H\), according to Sackur, is
\[
\left\{ dq_1 dp_1 dq_2 dp_2 dq_3 dp_3 \right\} = h^3.
\]
Tetrode’s procedure was identical.

Attention to this point has recently been called by Ehrenfest and Trkal\footnote{Ehrenfest and Trkal, *Proc. Amst. Akad.* xxiii. (1920).}, who have introduced a new method of dealing with the thermodynamical problems, \textit{e.g.} dissociation equilibrium and vapour pressure, in all these difficulties are avoided. We shall return to this method shortly. Planck has justified his assumption, viz., \(H = N h^3\), by taking into account the permutability of the molecules, but the reasoning is rather difficult to follow.

\[\S 2. \text{Ehrenfest’s Method.}\]

Ehrenfest has introduced a very general method for dealing with the thermodynamical behaviour of different and complex systems, like mixtures of gases reacting with each other. We shall confine our attention to the case of perfect gases.

Ehrenfest replaces the entropy function \(S\) by another function \(\{\gamma\}\) which is allied but not equal to \(W\). \(\{\gamma\}\) is supposed to represent the total phase space described by the system and is defined in the following way:—Let the system consist of \(N\) particles. Then each particle describes a subphase space \(\{\mu\}\), where
\[
\{\mu\} = \ldots \{ dq_1 dp_1 dq_2 dp_2 \ldots dq_r dp_r \} \quad \ldots (16)
\]
The total \(\{\gamma\}\)-space described by the system is then given by
\[
\{\gamma\} = P \frac{N!}{N} \{\mu\}, \quad \ldots (17)
\]
the product extending over all the particles. \(P\) represents the permutability of the particles.

Now, in our case, \textit{i.e.}, for monatomic gases,
\[
\{\gamma\} = \left\{ \ldots \{ dx_1 dy_1 dy_2 dz_1 dm_1 dm_2 \} \ldots \right\}, \quad \ldots (17.1)
\]
\[
u = p_{1r}, \quad m_r = p_{2r}, \quad m_w = p_{3r}.
\]
Now
\[
\Sigma p_{1r}^2 + p_{2r}^2 + p_{3r}^2 = 2mE, \quad \ldots (17.2)
\]
where \(E = \text{total kinetic energy and } m = \text{mass of a particle.}\)

Hence
\[
\{\gamma\} = \frac{N! (2\pi mE)^\frac{3N}{2}}{\Gamma (\frac{3N}{2})} \quad \ldots (18)
\]
Putting now
\[
E = \frac{3N}{2} kT,
\]
\[
\log \{\gamma\} = N \log \left( \frac{V}{2\pi m k T} \right)^\frac{3N}{2} \quad \ldots (19)
\]
According to Sackur-Tetrode:
\[
S = kN \log \left( \frac{V}{Nh^3} \left( \frac{2\pi mkT}{h} \right)^\frac{3N}{2} \right)
\]
\[
= k \log W.
\]
Comparing the two expressions,
\[ W = \frac{[\gamma]}{k^{3N}N!} \] \hspace{1cm} (20)

§ 3

Ehrenfest and Trkal have not traced any connexion between \([\gamma]\) and \(W\). But from equations (18), (19), and (20), it is possible not only to connect \(W\) and \([\gamma]\) for any general system, but also to lay down a general theorem for the calculation of the probability of any system. This we now proceed to do.

According to Planck, the probability \(W\) is a whole number. But whenever in physics we wish to determine the absolute value of any quantity, we must as well lay down a "unit" for it. Now let us see what is meant by "Unit Probability."

The idea of "Unit Probability" is intimately connected with the zero of entropy; for \(S = k \log W\), and when \(W = 1\), \(S = 0\). According to the second law of thermodynamics, this takes place when a condensed system is reduced to absolute zero; for a reversible engine acting between this system and another system at a finite temperature \(T\) will be able to convert all the heat energy abstracted into mechanical work.

We can suppose this to take place in another way. Suppose we have a gas at a finite temperature. The particles are moving in a chaotic way and at a distance from each other. Suppose all of them suddenly begin to move with identical velocity in the same direction, with the proviso, however, that the total kinetic energy remains the same. Then if this system, on coming into contact with another body and suffering inelastic collision, transfers the whole of its energy to that body, we can say that the heat motion has been completely converted into mechanical work. Thus the idea of absolute zero, and zero of entropy, presupposes a system of particles absolutely devoid of all motion. But this is not sufficient. If the particles which are devoid of all motion remain at a distance from each other, then, owing to mutual attraction and there being no motion, they will begin to move towards each other, and kinetic energy will again be developed and can again be converted into mechanical work. For an attracting system, motion, and with it the energy available from the system, will entirely stop when the particles are packed together in the closest manner possible.

We assume that such a system possesses unit probability. Let the value of \(\gamma\)-space for such a system be denoted by \([\gamma]\).

We may remark here that these considerations apply only if we regard the atoms as the final constituents of matter. This, however, is not the case, and therefore, even when the state pictured above has been reached, the electrons of one atom will react mechanically on the other atoms and electrons; and motion will never be entirely absent. Thus we cannot conceive of an absolute zero of temperature unless we picture to ourselves a state in which the protons and electrons have combined in some unknown way and annihilated all matter. We can therefore talk of absolute zero in a world where there is no matter. But according to the generalized theory of relativity, there can be neither space time, nor any physical quantity in a world which is entirely devoid of matter. Hence we come to a conclusion which has been previously reached by Nernst, viz., in the phenomenal world it is impossible to reach the absolute zero of temperature.

We shall now proceed to calculate the thermodynamical probability of a system in the terms of \([\gamma]\), as unity. The probability of the state of a system is proportional to the phase space \([\gamma]\) described by the system. Therefore the thermodynamical probability at a finite temperature

\[ W = \frac{[\gamma]}{[\gamma]_0} \] \hspace{1cm} (21)

We have now to find out the value of \([\gamma]_0\).

Let us suppose that in the system there are \(N\) particles, each particle having \(f\) degrees of freedom. We shall follow Ehrenfest's considerations in calculating \([\gamma]_0\). According to the definition, for a single particle

\[ \{\mu\} = \ldots \{dq_1dp_1, dq_2dp_2\ldots dq_fdq_f\} \] \hspace{1cm} (16)

\[ = \hbar^f. \]

Let us suppose that each particle is a Planck-resonator and confine our attention to one degree of freedom only. According to the quantum theory its phase-point \((q,p)\) must lie at \(p = 0, q = 0\), or on one of the ellipses \(h, 2h, \ldots\), the area between two consecutive ellipses being given by

\[ \oint dq dp = \hbar^f. \]

Ehrenfest attaches a "weight" \(\hbar\) to each one of these ellipses, and in particular also to the point \((p = 0, q = 0)\). Now, in our case, \(i.e.,\) for a system at absolute zero, none of the degrees of freedom of any particle is excited; or the phase of every particle for each degree of freedom is at \((p = 0, q = 0)\).

Hence, for a single particle,

\[ \{\mu\} = \ldots \{dq_1dp_1, dq_2dp_2\ldots dq_fdq_f\} \]

\[ = \hbar^f. \]

For the whole system

\[ \pi^N_{\{\mu\}} = \hbar^{fN} \] \hspace{1cm} (22)

Again the \(N\) particles can be arranged amongst themselves in \(N!\) ways. Combining this with (22), we obtain

\[ [\gamma]_0 = \hbar^{fN}N! \] \hspace{1cm} (23)

To sum up, the thermodynamic probability in Planck's sense is given by

\[ W = \frac{|\gamma|}{h^{\text{sym}} N} \]  

(20·1)

This will cover all cases.

On the basis of this law we shall now calculate the entropy of a system consisting of diatomic molecules.

§ 4. Entropy of a Gas consisting of Diatomic Molecules.

In this case we shall ignore the rotation of molecules about the axis of symmetry and also all internal motions of atoms in the molecule. Therefore, each molecule has only five degrees of freedom, three of these being translational and two rotational.

Thus

\[ |\gamma|_0 = h^{\text{sym}} N \]

\[ = h^{\text{sym}} N, \]  

(24)

where \( N \) denotes the total number of molecules considered.

Now, for a single molecule,

\[ |\mu| = \int dx dy dz dt dp_{1x} dp_{2x} dp_{3x} dp_{4x} dp_{5x} \]

\[ = V^x (4\pi)^x \int dp_{1x} dp_{2x} dp_{5x} \]

(25)

Therefore \( \gamma \)-space for the total system is given by

\[ |\gamma| = \prod_{x} |\mu| \]

\[ = V^x (4\pi)^x \int dp_{1x} dp_{2x} dp_{5x} \]  

(26)

the integrals being taken for all molecules, \( dp_{1x}, dp_{2x}, \) etc., denote the momenta of the different molecules.

The total kinetic energy of the molecules being given, the integration is to be taken over all possible values of the momenta which are consistent with it. If \( E \) be the kinetic energy given, we have

\[ \sum \frac{p_{1x}^2}{2m} + \frac{p_{2x}^2}{2m} + \frac{p_{3x}^2}{2a} + \frac{p_{4x}^2}{2b} = E, \]  

(27)

where \( m \) is the molecular mass and \( a \) and \( b \) represent the moments of inertia. The molecules being all similar, these quantities \( (m, a, \) and \( b) \) have the same values for all the molecules. The total number of terms in the left-hand side of equation (27) is \( 5N \).

Hence

\[ |\gamma| = V^x (4\pi)^x \int dp_{1x} dp_{2x} dp_{5x} \]

\[ = V^x (4\pi)^x \sqrt{2\pi E^{5/2}} \left( \frac{5N}{2} \right) \]

\[ = \frac{V^x (4\pi)^x \sqrt{2\pi E^{5/2}} \left( \frac{5N}{2} \right)}{h^{\text{sym}} N} \] 

(28)

since \( N \) is very large, we may put \( 5N \) for \( 5N - 1 \).

Hence

\[ W = \frac{|\gamma|}{|\gamma|_0} = \frac{|\gamma|}{h^{\text{sym}} N !} \]

\[ = \frac{V^x (4\pi)^x \sqrt{2\pi E^{5/2}} \left( \frac{5N}{2} \right)}{h^{\text{sym}} N !} \] 

(29.1)

Using Stirling's formula and putting \( E = \frac{1}{2} kT \), we obtain

\[ W = \left( \frac{e^{\frac{1}{2}} 4\pi V \left( (2\pi kT)^{5/2} \right) m^{3/2} a^{1/2} b^{1/2}}{N h^5} \right)^{\frac{5N}{2}} \] 

(29.2)

\[ S = k \log W \]

\[ = N k \log \left( \frac{e^{\frac{1}{2}} 4\pi V \cdot m^{3/2} a^{1/2} b^{1/2} \cdot (2\pi kT)^{5/2}}{N h^5} \right) \] 

(30)

It may be emphasized here that the formula \( E = \frac{1}{2} kT \) for monatomic gases and \( E = \frac{1}{3} kT \) for diatomic gases have not been assumed here, but followed directly from the theory. The calculations have not been reproduced here.

32. ON ENTROPY OF RADIATION II*

MEGHINAD SAHA & RAMANI KANTA SUR

(Phil. Mag., Sr. VII, t, 890, 1926)

In a previous paper, it has been shown that the thermodynamical probability of a system is given by the law,

\[ W = |\gamma| / |\gamma|_0 \]

(1)

Where \( |\gamma| = \) phase-space described by the system at temperature \( T_0 \), \( |\gamma|_0 = \) phase-space described at absolute zero. It was shown that the absolute value of entropy of perfect gases could be deduced from this theorem.

The same formula can be applied for deducing the entropy of radiation. Since the time when Bartoli deduced the existence of radiation pressure from thermodynamical reasoning, it has been customary to look upon radiation as