took into account the conservation of both energy and momentum in the interaction between the electron and the X-ray photon.

\[ \rho = v^2 f \left( \frac{v}{T} \right) \]  \hspace{1cm} \text{(1)}

he was led by this similarity to a farther correspondence with the radiation formula. He discovered, as is known, the formula [Wien’s radiation formula]

\[ \rho = \alpha v^3 e^{-\frac{h\nu}{kT}} \]  \hspace{1cm} \text{(2)}

which is recognized today as the correct limiting formula for large values of \( \frac{v}{T} \). Today we know that no consideration which is based on classical mechanics and electrodynamics can lead to a useful radiation formula; rather that the classical theory leads to the Rayleigh formula.

\[ \rho = \frac{k\alpha}{h} v^2 T \]  \hspace{1cm} \text{(3)}

After Planck, in his ground-breaking investigation, established his radiation formula

\[ \rho = \alpha v^3 \frac{1}{h\nu} \frac{1}{e^{h\nu/kT} - 1} \]  \hspace{1cm} \text{(4)}

\(^{1}\text{Albert Einstein, Physikalische Zeitschrift, 18 (1917), 121-128—trans. Editors.}\)
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on the assumption that there are discrete elements of energy, from which quantum theory developed very rapidly, Wien’s considerations, from which formula (2) evolved, quite naturally were forgotten.

A little while ago I obtained a derivation, related to Wien’s original idea, of the Planck radiation formula which is based on the fundamental assumption of quantum theory and which makes use of the relationship of Maxwell’s curve to the spectral distribution curve. This derivation deserves consideration not only because of its simplicity, but especially because it appears to clarify the processes of emission and absorption of radiation in matter, which is still in such darkness for us. In setting down certain fundamental hypotheses concerning the absorption and emission of radiation by molecules that are closely related to quantum theory, I showed that molecules with a distribution of states in the quantum theoretical sense for temperature equilibrium are in dynamical equilibrium with the Planck radiation; in this way, the Planck formula (4) was obtained in a surprisingly simple and general way. It was obtained from the condition that the quantum theoretic partition of states of the internal energy of the molecules is established only by the emission and absorption of radiation.

If the assumed hypotheses about the interaction of matter and radiation are correct, they will give us more than just the correct statistical partition or distribution of the internal energy of the molecules. During absorption and emission of radiation there is also present a transfer of momentum to the molecules; this means that just the interaction of radiation and molecules leads to a velocity distribution of the latter. This must clearly be the same as the velocity distribution which molecules acquire as the result of their mutual interaction by collisions, that is, it must coincide with the Maxwell distribution. We must require that the mean kinetic energy which a molecule (per degree of freedom) acquires in a Planck radiation field of temperature $T$ be

$$\frac{kT}{2};$$

this must be valid regardless of the nature of the molecules and independent of frequencies which the molecules absorb and emit. In this paper we wish to verify that this far-reaching requirement is, indeed, satisfied quite generally; as a result of this our simple hypotheses about the emission and absorption of radiation acquire new supports.

In order to obtain this result, however, we must enlarge, in a definite way, the previous fundamental hypotheses which were related entirely to the exchange of energy. We are faced with this question: Does the molecule suffer a push, when it absorbs or emits the energy $\epsilon$? As an example we consider, from the classical point of view, the emission of radiation. If a
body emits the energy $\epsilon$, it acquires a backward thrust [impulse] $\frac{\epsilon}{c}$ if all the radiation $\epsilon$ is radiated in the same direction. If, however, the radiation occurs through a spatially symmetric process, for example, spherical waves, there is then no recoil at all. This alternative also plays a role in the quantum theory of radiation. If a molecule, in going from one possible quantum theoretic state to another, absorbs or emits the energy $\epsilon$ in the form of radiation, such an elementary process can be looked upon as partly or fully directed in space, or also as a symmetric (non-directed) one. It turns out that we obtain a theory that is free of contradictions only if we consider the above elementary processes as being fully directed events; herein lies the principal result of the considerations that follow.

**FUNDAMENTAL HYPOTHESES OF THE QUANTUM THEORY—CANONICAL DISTRIBUTION OF STATES**

According to the quantum theory, a molecule of a definite kind may, aside from its orientation and its translational motion, be in one of only a discrete set of states $Z_1, Z_2, \ldots, Z_n \ldots$ whose (internal) energies are $\epsilon_1, \epsilon_2, \ldots, \epsilon_n \ldots$. If the molecules of this kind belong to a gas of temperature $T$, then the relative abundance $W_n$ of the state $Z_n$ is given by the statistical mechanical canonical partition function for states

$$W_n = p_n e^{-\frac{\epsilon_n}{kT}}$$  \hspace{1cm} (5)

In this formula $k = \frac{R}{N}$ is the well-known Boltzmann constant, $p_n$ a number that is independent of $T$ and characteristic of the molecule and the state, which we may call the statistical “weight” of the state. Formula (5) can be derived from the Boltzmann principle or purely from thermodynamics. Equation (5) is the expression of the most far-reaching generalization of the Maxwellian distribution of velocities.

The latest important advances in quantum theory deal with the theoretical determination of the quantum theoretical possible states $Z_n$ and their weights $p_n$. For the principal part of the present investigation, it is not necessary to have a more detailed determination of the quantum states.

**HYPOTHESES ABOUT THE ENERGY EXCHANGE THROUGH RADIATION**

Let $Z_n$ and $Z_m$ be two possible quantum theoretical states of a gas molecule whose energies $\epsilon_n$ and $\epsilon_m$ respectively, satisfy the inequality

$$\epsilon_m > \epsilon_n$$
Let the molecule be able to pass from the state $Z_n$ to the state $Z_m$ by absorbing the radiation energy $\epsilon_m - \epsilon_n$; similarly let the transition from state $Z_n$ to the state $Z_m$ be possible through the emission of this amount of energy. Let the radiation emitted or absorbed by the molecule for the given index and combination $(m, n)$ have the characteristic frequency $\nu$.

We now introduce certain hypotheses about the laws which are decisive for these transitions. These hypotheses are obtained by carrying over the known classical relations for a Planck resonator to the unknown quantum theoretical relations.

**Emission**

A Planck resonator that is vibrating radiates energy, according to Hertz, in a known way independently of whether it is stimulated by an external field or not. In accordance with this, let a molecule be able to pass from the state $Z_m$ to the state $Z_n$ with the emission of radiant energy $\epsilon_m - \epsilon_n$ of frequency $\nu$ without being excited by any external cause. Let the probability $dW$ for this to happen in the time $dt$ be

$$dW = A_m^n dt \quad (A)$$

where $A_m^n$ is a characteristic constant for the given index combination.

The assumed statistical law corresponds to that of a radioactive reaction: that elementary process of such a reaction in which only $\gamma$-rays are emitted. We need not assume that this process requires no time; this time need only be negligible compared to the times which the molecule spends in the states $Z_1$, and so on.

**Incident Radiation**

If a Planck resonator is in a radiation field, the energy of the resonator changes because the electromagnetic field of the radiation does work on the resonator; this work can be positive or negative depending on the phases of the resonator and the oscillating field. In accordance with this, we introduce the following quantum theoretical hypothesis. Under the action of the radiation density $\rho$ of the frequency $\nu$ a molecule in state $Z_n$ can go over to state $Z_m$ by absorbing the radiation energy $\epsilon_m - \epsilon_n$ in accordance with the probability law

$$dW = B_n^m \rho dt \quad (B)$$

In the same way, let the transition $Z_m \rightarrow Z_n$ under the action of the radiation also be possible, whereby the radiation energy $\epsilon_m - \epsilon_n$ is emitted according to the probability law

$$dW = B_m^n \rho dt \quad (B')$$
$B_n^m$ and $B_m^n$ are constants. We call both processes “changes of states through incident radiation.”

The question presents itself now as to the momentum that is transferred to the molecule in these changes of state. We begin with the events associated with incident radiation. If a directed bundle of rays does work on a Planck resonator, then an equivalent amount of energy is removed from the bundle. This transfer of energy results, according to the law of momentum, to a momentum transfer from the beam to the resonator. The latter therefore experiences a force in the direction of the ray of the radiation beam. If the energy transferred is negative, the force acting on the resonator is opposite in direction. In the case of the quantum hypothesis, this clearly means the following. If, as the result of incident radiation, the process $Z_n \rightarrow Z_m$ occurs, then an amount of momentum

$$\frac{\epsilon_m - \epsilon_n}{c}$$

is transferred to the molecule in the direction of propagation of the bundle of radiation. If we have the process $Z_m \rightarrow Z_n$ for the case of incident radiation, the magnitude of the transferred momentum is the same, but it is in the opposite direction. If a molecule is simultaneously exposed to many bundles of radiation, we assume that the total energy $\epsilon_m - \epsilon_n$ is taken from or added to just one of these bundles, so that even in this case the momentum

$$\frac{\epsilon_m - \epsilon_n}{c}$$

is transferred to the molecule.

In the case of emission of energy by radiation by a Planck resonator, there is no net transfer of momentum to the resonator because, according to classical theory, the emission occurs as a spherical wave. However, we have already noted that we can arrive at a contradiction-free quantum theory only if we assume that the process of emission is a directed one. Every elementary process of emission ($Z_m \rightarrow Z_m$) will then result in a transfer to the molecule of an amount of momentum

$$\frac{\epsilon_m - \epsilon_n}{c}.$$  

If the molecule is isotropic, we must take every direction of emission as equally probable. If the molecule is not isotropic, we arrive at the same result if the orientation changes in a random way in the course of time. We must, in any case, make such an assumption also for the statistical laws
(B) and (B') for incident radiation since otherwise the constants $B_n^m$ and $B_m^n$ would have to depend on direction, which we can avoid by assuming isotropy or pseudo-isotropy (through setting up temporal mean values).

**DERIVATION OF THE PLANCK RADIATION LAW**

We now enquire about those effective radiation densities $\rho$ which must prevail in order that the energy exchange between molecules and radiation as a result of the statistical laws (A), (B) and (B') shall not disturb the distribution of molecular states present as a consequence of equation (5). For this, it is necessary and sufficient that on the average, per unit time, as many elementary processes of type (B) take place as processes (A) and (B') together. This condition gives as a result of (5), (A), (B), (B'), for the elementary processes corresponding to the index combination ($m$, $n$) the equation

$$p_n e^{-\frac{\epsilon_n}{kT}} B_n^m \rho = p_m e^{-\frac{\epsilon_m}{kT}} (B_m^n \rho + A_m^n)$$

If, further, $\rho$ is to become infinite as $T$ does, the constants $B_n^m$ and $B_m^n$ must satisfy the relation

$$p_n B_n^m = p_m B_m^n$$  \hspace{1cm} (6)

We then obtain as the condition for dynamical equilibrium the equation

$$\rho = \frac{A_m^n / B_m^n}{e^{\frac{\epsilon_m - \epsilon_n}{kT}} - 1}$$  \hspace{1cm} (7)

This is the dependence of the radiation density on the temperature that is given by the Planck law. From the Wien displacement law (1) it then follows immediately that

$$\frac{A_m^n}{B_m^n} = a \nu^3$$  \hspace{1cm} (8)

and

$$\epsilon_m - \epsilon_n = h\nu$$  \hspace{1cm} (9)

where $a$ and $h$ are universal constants. To obtain the numerical values of $a$ and $h$ we must have an exact theory of electrodynamic and mechanical
processes; we content ourselves for the moment with the Rayleigh law in the limit of high temperatures, where the classical theory is valid in the limit.

Equation (9) is, as we know, the second principal rule in Bohr's theory of spectra, about which we may assert, following upon Sommerfeld's and Epstein's completion of the theory, that it belongs to the most fully verified domain of our science. It also contains implicitly the photochemical equivalent law, as I have already shown.

**METHOD FOR CALCULATING THE MOTION OF MOLECULES IN RADIATION FIELDS**

We now turn our attention to the investigation of the motion imparted to our molecules by the radiation field. We make use in this of a method that is known to us from the theory of Brownian motion and which I have often used in investigating motions in a region containing radiation. To simplify the calculation, we shall carry it through for the case in which the motion occurs only along the X-direction of the coordinate system. We further content ourselves with calculating the mean value of the kinetic energy of the translational motion, and thus dispense with proof that these velocities $v$ are distributed according to the Maxwell law. Let the mass $M$ of the molecule be large enough so that higher powers of $\frac{v}{c}$ can be neglected relative to lower ones; we can then apply the usual mechanics to the molecule. Moreover, without any loss in generality, we may carry out the calculation as though the states with indices $m$ and $n$ were the only ones the molecule can be in.

The momentum $Mv$ of a molecule undergoes two kinds of changes in the short time $\tau$. Even though the radiation is the same in all directions, the molecule, because of its motion, will experience a resistance to its motion that stems from the radiation. Let this opposing force be $Rv$, where $R$ is a constant to be determined later. This force would ultimately bring the molecule to rest if the randomness of the action of the radiation field were not such as to transfer to the molecule a momentum $\Delta$ of alternating sign and varying magnitude; this random effect will, in opposition to the previous one, sustain a certain amount of motion of the molecule. At the end of the given short time $\tau$ the momentum of the molecule will equal

$$Mv - Rv\tau + \Delta$$

Since the velocity distribution is to remain constant in time, the mean of the absolute value of the above quantity must equal that of the quantity
$M \nu$; thus, the mean values of the squares of both quantities averaged over a long time or over a large number of molecules must be equal:

$$(M \nu - R \nu \Delta + \Delta)^2 = (M \nu)^2$$

Since we have taken into account the influence of $\nu$ on the momentum of the molecule separately, we must discard the mean value $\nu \Delta$. On developing the left-hand side of the equation we thus obtain

$$\Delta^2 = 2RM\nu^2$$

(10)

The mean value $\nu^2$ which the radiation of temperature $T$ by its interaction imparts to the molecule must just equal the mean value $\nu^2$ which the gas molecule acquires at temperature $T$ according to the gas law and the kinetic theory of gases. For otherwise the presence of our molecules would disturb the thermal equilibrium between thermal radiation and an arbitrary gas of the same temperature. We must therefore have

$$\frac{M\nu^2}{2} = \frac{kT}{2}$$

(11)

Equation (10) thus goes over into

$$\frac{\Delta^2}{\tau} = 2RkT$$

(12)

The investigation is now to be carried through as follows. For a given radiation density ($\rho(\nu)$) we shall be able to compute $\Delta^2$ and $R$ by means of our hypotheses about the interaction between radiation and molecules. If we put this result into (12), this equation will have to be identically satisfied when $\rho$ is expressed as a function of $\nu$ and $T$ by means of Planck's equation (4).

**Computing $R$**

Let a molecule of the given kind be in uniform motion with speed $\nu$ along the $X$-axis of the coordinate system $K$. We inquire about the momentum transferred on the average from the radiation to the molecule per unit time. To calculate this we must consider the radiation from a coordinate system $K'$ that is at rest with respect to the given molecule. For we have formulated our hypotheses about emission and absorption only for molecules at rest. The transformation to the system $K'$ has often been performed in the literature. Nevertheless, I shall repeat the simple considerations here for the sake of clarity.
Relative to $K$ the radiation is isotropic, that is, the quantity of radiation in a solid angle $d\kappa$ in the direction of the radiation in a frequency range $dv$ is

$$\rho dv \frac{d\kappa}{4\pi}$$

(13)

where $\rho$ depends only on the frequency $v$ but not on the direction of the radiation. This special beam corresponds to a special beam in the system $K'$ which is also characterized by a frequency range $dv'$ and a solid angle $d\kappa'$. The volume density of this special beam is

$$\rho'(v',\phi') dv' \frac{d\kappa'}{4\pi}$$

(13')

This defines $\rho'$. It depends on the direction of the radiation which, in the familiar manner, is defined by the angle $\phi'$ it makes with the $X'$ axis and which its projection on the $Y'$, $Z'$ plane makes with the $Y'$ axis. These angles correspond to the angles $\phi$ and $\psi$ which in an analogous manner determine the direction of $d\kappa$ in $K$.

To begin with, it is clear that the same transformation law between (13) and (13') must hold as between the amplitudes $A^2$ and $A'^2$ of a plane wave moving in the corresponding direction. Hence, to our desired approximation we have

$$\frac{\rho'(v',\phi') dv' d\kappa'}{\rho(v) dv d\kappa} = 1 - 2 \frac{v}{c} \cos\phi$$

(14)

or

$$\rho'(v',\phi') = \rho(v) \frac{dv}{dv'} \frac{d\kappa}{d\kappa'} \left( 1 - 2 \frac{v}{c} \cos\phi \right)$$

(14')

The relativity theory further gives the formulae, valid to the desired approximation,

$$v' = v \left( 1 - \frac{v}{c} \cos\phi \right)$$

(15)

$$\cos\phi' = \cos\phi - \frac{v}{c} + \frac{v}{c} \cos^2\phi$$

(16)

$$\psi' = \psi$$

(17)

From (15) it follows, to the same approximation, that

$$v = v' \left( 1 + \frac{v}{c} \cos\phi' \right).$$
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Hence, again to the desired approximation

$$\rho(v) = \rho \left( v' + \frac{v}{c} v' \cos \phi' \right)$$

or

$$\rho(v) = \rho(v') + \frac{\partial \rho(v')}{\partial v} \left( \frac{v}{c} v' \cos \phi' \right)$$  \hspace{1cm} (18)

Further, according to (15), (16), and (17)

$$\frac{dv}{dv'} = \left( 1 + \frac{v}{c} \cos \phi' \right)$$

$$\frac{dk}{dk'} = \frac{\sin \phi' \, d\phi' \, d\psi'}{\sin \phi \, d\phi \, d\psi} = \frac{d(\cos \phi)}{d(\cos \phi')} = 1 - 2 \frac{v}{c} \cos \phi'$$

As a result of these two equations and equation (18), equation (14') goes over into

$$\rho'(v', \phi') = \left[ (\rho)_v + \frac{v}{c} v' \cos \phi' \left( \frac{\partial \rho}{\partial v} \right)_{v'} \right] \left( 1 - 3 \frac{v}{c} \cos \phi' \right)$$  \hspace{1cm} (19)

With the aid of (19) and our hypotheses about the radiation from and radiation onto molecules, we can easily calculate the average momentum transferred to the molecule per unit time. Before we can do this, however, we must say something to justify our procedure. It may be objected that equations (14), (15), (16) are based on Maxwell's theory of the electromagnetic field that is not consistent with the quantum theory. This objection deals, however, more with the form than with the substance of the problem. For, no matter how the theory of electromagnetic processes may be formulated, in any case the Doppler principle and the law of aberration still remain, and hence also the equations (15) and (16). Moreover, the validity of the energy relationship (14) certainly extends beyond that of the wave theory; this transformation law is also valid, for example, according to relativity theory, for the energy density of a mass of infinitesimally small rest density that is moving with the [quasi-] speed of light. We may therefore assert the validity of equation (19) for any theory of radiation.

The radiation belonging to the solid angle $d\kappa'$ would, according to (B), give rise to

$$B_{n, m} \rho'(v', \phi') \frac{d\kappa'}{4\pi}$$

elementary processes per second of radiation events of the type $Z_n \rightarrow Z_m$ if the molecule after each such process immediately returned to state
\[ Z_n. \text{ Actually, however, the time of lingering in state } Z_n, \text{ according to (5), is} \]

\[ \frac{1}{S} p_n e^{-\frac{\epsilon_n}{kT}} \]

where we have used the abbreviation

\[ S = p_n e^{-\frac{\epsilon_n}{kT}} + p_m e^{-\frac{\epsilon_m}{kT}} \]

(20)

The number of these processes per second is therefore actually

\[ \frac{1}{S} p_n e^{-\frac{\epsilon_n}{kT}} B_n^m \rho'(v', \phi') \frac{d\kappa'}{4\pi}. \]

In each of these elementary processes the momentum

\[ \frac{\epsilon_m - \epsilon_n}{c} \cos \phi' \]

is transferred to the molecule in the direction of the \( X' \)-axis. In an analogous manner we find, based on (B') that the corresponding number of elementary processes of radiation events of type \( Z_m \rightarrow Z_n \) per second is

\[ \frac{1}{S} p_m e^{-\frac{\epsilon_m}{kT}} B_n^m \rho'(v', \phi') \frac{d\kappa'}{4\pi} \]

and in each such elementary process the momentum

\[ -\frac{\epsilon_m - \epsilon_n}{c} \cos \phi' \]

is transferred to the molecule. The total momentum transferred to the molecule per unit time by incident radiation is, keeping in mind (6) and (9),

\[ \frac{h v'}{c S} p_n B_n^m \left( e^{-\frac{\epsilon_n}{kT}} - e^{-\frac{\epsilon_m}{kT}} \right) \int \rho'(v', \phi') \cos \phi' \frac{d\kappa'}{4\pi} \]

where the integration is to be taken over all solid angles. Carry this out, and we obtain with the aid of (19) the value

\[ -\frac{h v}{c^2 S} \left( \rho - (\frac{1}{2}) v \frac{\partial \rho}{\partial v} \right) p_n B_n^m \left( e^{-\frac{\epsilon_n}{kT}} - e^{-\frac{\epsilon_m}{kT}} \right) v. \]
Here we have represented the effective frequency again with \( \nu \) and not with \( \nu' \). This expression gives, however, the total momentum transferred on the average to a molecule moving with speed \( \nu \). For it is clear that those elementary processes of emission of radiation not induced by the action of the radiation field have no preferred direction as seen from system \( K' \) and hence, on the average, cannot transfer any momentum to the molecule. We thus obtain as the final result of our considerations

\[
R = \frac{\hbar \nu}{c^2 S} \left( \rho - \frac{1}{3} \nu \frac{\partial \rho}{\partial \nu} \right) p_x B_n e^{-\frac{e_n}{kT}} \left( 1 - e^{-\frac{kT}{\hbar \nu}} \right) \tag{21}
\]

**CALCULATING \( \Delta^2 \)**

It is much easier to calculate the random effect of the elementary processes on the mechanical behavior of the molecule. For we can calculate this for a molecule at rest for which the approximation which we have been using applies.

Let some event cause the momentum \( \lambda \) to be transferred to a molecule in the X direction. This momentum is to be of varying magnitude and direction from moment to moment. However, let \( \lambda \) obey a statistical law such that its average value vanishes. Then let \( \lambda_1, \lambda_2 \ldots \) be the momenta which are transferred to the molecule in the X-direction by various operating causes that are independent of each other so that the total momentum that is transferred is

\[
\Delta = \Sigma \lambda
\]

We then have (if for the individual \( \lambda \), their mean values \( \lambda \) vanish)

\[
\Delta^2 = \Sigma \lambda^2 \tag{22}
\]

If the mean values \( \lambda^2 \) of the individual momenta are all equal to each other \( (= \lambda^2) \) and if \( l \) is the total number of processes giving rise to momenta, we have the relation

\[
\Delta^2 = l \lambda^2 \tag{22a}
\]

According to our hypothesis, in each process of incident radiation and outflowing radiation, the momentum

\[
\lambda = \frac{h \nu}{c} \cos \phi
\]
is transferred to the molecule. Here $\phi$ is the angle between the X-axis and some randomly chosen direction. Hence, we obtain

$$\overline{\lambda^2} = \frac{1}{3} \left( \frac{h\nu}{c} \right)^2.$$

Since we assume that all the elementary processes that are present are to be considered as events that are independent of each other, we may apply (22a); $l$ is then the number of all elementary processes that occur in the time $\tau$. This is twice as large as the number of radiation-incident processes $Z_n \rightarrow Z_m$ in the time $\tau$. We thus have

$$l = \frac{2}{6} p_n B_n^m e^{-\frac{\xi_n}{kT}} \rho \tau$$  \hspace{1cm} (24)

From (23), (24) and (22) we thus obtain

$$\frac{\overline{A^2}}{\tau} = \frac{2}{3S} \left( \frac{h\nu}{c} \right)^2 p_n B_n^m e^{-\frac{\xi_n}{kT}} \rho$$  \hspace{1cm} (25)

**RESULTS**

In order now to show that the momenta transferred from the radiation to the molecule according to our basic hypotheses never disturb the thermodynamic equilibrium, we need only introduce the values for $\frac{\overline{A^2}}{\tau}$ and $R$ calculated in (25) and (21) respectively after the quantity

$$\left( \rho - \left( \frac{1}{6} \right) \nu \frac{\partial \rho}{\partial \nu} \right) \left( 1 - e^{-\frac{h\nu}{kT}} \right)$$

in (21) is replaced by

$$\frac{\rho hv}{3RT}$$

from (4). We then see that our fundamental equation (12) is satisfied identically.

The above consideration lends very strong support to the hypotheses introduced earlier for the interaction between matter and radiation by means of absorption and emission, and through incident and outgoing radiation. I was led to these hypotheses in trying to postulate in the simplest possible way a quantum behavior of molecules that is analogous to
the Planck resonators of classical theory. We obtained, without effort, from the general quantum assumption for matter, the second Bohr rule (equation (9)) as well as Planck's radiation formula.

Most important, however, appears to me the result about the momentum transferred to the molecule by incoming and outgoing radiation. If one of our hypotheses were altered, the result would be a violation of equation (12); it appears hardly possible, except by way of our hypotheses, to be in agreement with this relationship which is demanded by thermodynamics. We may therefore consider the following as pretty well proven.

If a beam of radiation has the effect that a molecule on which it is incident absorbs or emits an amount of energy $h\nu$ in the form of radiation by means of an elementary process, then the momentum $h\nu/c$ is always transferred to the molecule, and, to be sure, in the case of absorption, in the direction of the moving beam and in the case of emission in the opposite direction. If the molecule is subject to the simultaneous action of beams moving in various directions, then only one of these takes part in any single elementary process of incident radiation; this beam alone then determines the direction of the momentum transferred to the molecule.

If, through an emission process, the molecule suffers a radiant loss of energy of magnitude $h\nu$ without the action of an outside agency, then this process, too, is a directed one. Emission in spherical waves does not occur. According to the present state of the theory, the molecule suffers a recoil of magnitude $h\nu/c$ in a particular direction only because of the chance emission in that direction.

This property of elementary processes as expressed by equation (12) makes a quantum theory of radiation almost unavoidable. The weakness of the theory lies, on the one hand, in its not bringing us closer to a union with the wave theory, and, on the other hand, that it leaves the time and direction of the elementary processes to chance; in spite of this, I have full confidence in the trustworthiness of this approach.

Only one more general remark. Almost all theories of thermal radiation rest on the considerations of the interaction between radiation and molecules. But, in general, one is satisfied with dealing only with the energy exchange, without taking into account the momentum exchange. One feels justified in this because the momentum transferred by radiation is so small that it always drops out as compared to that arising from other dynamical processes. But for the theoretical considerations, this small effect is on an equal footing with the energy transferred by radiation because energy and momentum are very intimately related to each other; a theory may therefore be considered correct only if it can be shown that the momentum transferred accordingly from the radiation to the matter leads to the kind of motion that is demanded by thermodynamics.