An experimental test of semiclassical radiation theories
by John Milton Wessner

A thesis submitted to the Graduate Faculty in partial fulfillment of the requirements for the degree of
DOCTOR OF PHILOSOPHY in Physics
Montana State University
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Abstract:
An experiment is described which tests the semi-classical radiation theory proposed by E. T. Jaynes
and his collaborators. Their work predicted that the $n = 2$ level of hydrogen will have a lifetime which
varies linearly with the 2s state population for small excited state components. By applying a short
electrical pulse to a 60 keV atomic hydrogen beam with a 2s population of approximately 9%, the
variation of the lifetime could be measured as a function of the 2s occupation after the pulse. A total
change of approximately $+4\%$ is predicted while the observed change was $-.2\%$. 
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in

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MONTANA STATE UNIVERSITY
Bozeman, Montana

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PREFACE

Each of us is attracted to physics by some deep impression made upon him as a child. For me it was the atomic bomb, although the road from that point to now is so circuitous and disconnected that only my most antagonistically humanist friends can still detect the taint. A more tangible starting point is to be found in the reading of a little book by G. P. Thomson called *The Inspiration of Science* and finding that the richness found there could be shared with my students. Unfortunately, the joyous days of string and sealing wax were gone, as rockets went roaring into space from just up the road, and biologists traded in their microscopes for ultracentrifuges.

As a consequence, it is a distinct pleasure to present a piece of research here which seems to test a point with well delineated philosophical roots, using equipment which in some cases was not in existence ten years ago, and yet, in which the data have a rather clean interpretation. One likes to think that had the results
been the opposite that there would now be a run on string and sealing wax.

Because of the amount of historical nonsense I imbibed as a student and my own notion of the significance of this work, I have included in the first chapter a very sketchy history of quantum mechanics up to the enunciation of the principle of complementarity. Things which are common knowledge and true are often omitted, but the outline seemed appropriate in light of my background. The reader may skip to the last six paragraphs if he pleases.

It is customary to thank everyone in sight for their part in the production of a dissertation, and one does well at this point to adhere to custom. Thanks go to Alfred Romer for sending me to Montana and to B. Frank Brown in Melbourne for allowing me to become scholarly at long last. No thanks are due to those who held back; they have their own righteousness for comfort. My hungry children have benefitted from the generous support of the MSU physics department as well as everyone else from whom funds were mulcted.
I am grateful to professor D. K. Anderson for suggesting the original problem and retaining his sanity in the dark days when nothing seemed right, to professor R. T. Robiscoe for some very necessary capital equipment, and to Mr. C. F. Badgley for his advice and direct assistance in building the apparatus. Most of all, may eternal blessings shower upon my wife Sally who, after celebrating her thirtieth birthday at thirty below, three thousand miles from home, has remained steadfast to the revision of the fourth draft.
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An experiment is described which tests the semi-classical radiation theory proposed by E. T. Jaynes and his collaborators. Their work predicted that the $n = 2$ level of hydrogen will have a lifetime which varies linearly with the $2s$ state population for small excited state components. By applying a short electrical pulse to a 60 keV atomic hydrogen beam with a $2s$ population of approximately 9%, the variation of the lifetime could be measured as a function of the $2s$ occupation after the pulse. A total change of approximately +4% is predicted while the observed change was -.2%.
CHAPTER I
WAVES, OSCILLATORS AND PHOTONS

The experiment described in this report is part of a continuing refinement of our understanding of the interaction of matter and electromagnetic radiation. It would take us far beyond the scope of a thesis to give a complete history of this problem, but it is appropriate to outline the essential events. To do so we rely upon several secondary sources\(^1\) but where possible the original references will be cited. We begin by tracing the study of two separate physical phenomena: the photoelectric effect and the blackbody radiation distribution.

Every student of physics knows that James Clerk Maxwell synthesized the concepts of electricity and magnetism.\(^2\) In the process oscillating fields which propagated at the velocity of light were predicted and were subsequently observed by Heinrich Herz.\(^3\) Herz's experiments detected the near field radiation of a spark discharge and, at the time, he observed that the induced spark occurred more readily when the light from the source was allowed to fall on the secondary electrodes.\(^4\) This
phenomenon was studied until by 1905 almost all of the
details of the photoelectric effect were well known to
physicists.

The study of blackbody radiation is obscured some-
what by our lesser familiarity with the thermodynamics
required for its study. In 1859 G. R. Kirchhoff proved that the ratio of the emission of heat to
absorption was independent of the nature of the body
involved. With the invention of the bolometer by
S. P. Langley the study of the wavelength distribution
of cavity radiation could proceed on a firm experimental
basis. In the meantime, theoretical considerations had
verified Stefan's law and established what is now
known as Wien's displacement law.

All this was done without trying to understand the
detailed processes involved. As a first attempt at
dealing with the underlying interaction Wien derived,
on rather weak physical assumptions, a distribution law
for the radiated energy:

$$u_{\nu} = \alpha \nu^3 e^{-\beta \nu/T},$$

(1)

where $u_{\nu}$ is the radiation density in the frequency
interval ν to ν + dν, T is the absolute temperature, and α and β are constants. This was subsequently put on firmer ground by Max Planck.\(^1\) At the same time the Rayleigh distribution law,

\[ u_ν = \frac{8\pi ν^2 kT}{c^3} \]  

was proposed. Equations 1 and 2 each agreed with parts of the data available, but violently disagreed with the rest.

To fully understand the resolution of the problem would require an excessive digression into classical thermodynamics, so we simply state the results. A fundamental relation for any constant volume thermodynamic system is

\[ \frac{\partial S}{\partial U} = T^{-1} \]  

where S is the entropy of the system and U is the internal energy. The physics enters when we write U as a function of T. The different radiation laws involve the equilibria predicted by two forms for the second derivative:

\[ \frac{\partial^2 S}{\partial U^2} \propto U^{-1} \]  

versus

\[ \frac{\partial^2 S}{\partial U^2} \propto U^{-2}. \]

The first leads to the Wien law and the second to
Rayleigh's. The compromise was obvious;

\[ \frac{\partial^2 S}{\partial U^2} = \frac{1}{U(b + U)}, \quad (5) \]

but what justification could be given? Planck was forced to use Boltzmann's statistical interpretation of entropy from which equation 5 followed if the distribution of energy to the radiators was discrete. The quantum was born, albeit reluctantly.\(^{11}\) It should be noted at this point it is matter which is quantized, not the radiation.

In 1905 Einstein succeeded in bringing these two phenomena together by transferring the discreteness of the energy distributions to the radiation.\(^{12}\) The excitement with which this was received is summed up by Planck in his nominating statement for Einstein's admission to the Prussian Academy in 1913:

"That he may sometimes have missed the target in his speculations, as for example, in his hypothesis of light quanta, cannot be held against him, for it is not possible to introduce fundamentally new ideas, even in the most exact sciences, without occasionally taking a risk."

The "photon" would not go away, however. By 1916 Millikan had painstakingly verified all the predictions for the photoelectric effect,\(^{13}\) and in 1923 the
Compton effect\textsuperscript{14} was explained in terms of a "particle-particle" interaction. In the meantime, the understanding of matter had also changed.

The description of atomic processes made incredible progress during this period if one considers that most of the models were purely speculative. As an example of the "old quantum theory", the Landé g-values\textsuperscript{15} were established by assigning to the core of the atom an angular momentum which corresponds to our electron spin. Hence, the persistence in German of calling $j$ the "inner quantum number". Basically, the old quantum theory took the Bohr atom, the correspondence principle, and the established formalism for solving multiply periodic motions to explain the experimental data which had accumulated in atomic physics. It remained for a group of young physicists, born at the same time as the quantum, to provide a quantum mechanics.

For our purposes quantum mechanics has two formulations: matrix mechanics and wave mechanics; the principal parts of the latter overlapped the former. The first modern wave mechanics was developed by
L. de Broglie in a series of papers in 1923 and presented as a doctoral thesis in 1925. The only points that we need be aware of are that the work is done in the framework of special relativity and that one must understand the distinctions between group and phase velocity to treat the theory correctly. With this model de Broglie was able to solve the Bohr hydrogen atom from first principles and suggest to his brother that he look for electron diffraction. (The laboratory was too busy trying to invent television to do so.)

The matrix mechanics begins with W. Heisenberg. Heisenberg was a strong advocate of the discontinuity of all microscopic processes. As a consequence he was willing to reject the notion of a well-defined trajectory, and looked for a way to make position and momentum quantum mechanical variables in the same sense as the energy. The results led to a matrix mechanics which also predicted the experimentally known results for the hydrogen atom and the anharmonic oscillator. The full formalism is found in the papers by Born and Jordan and Born, Jordan and Heisenberg. These papers allow
quite accurate descriptions of atomic phenomena; once again, from first principles.

At this point E. Schroedinger published a sequence of papers\textsuperscript{20-23} which established the wave mechanics as an outgrowth of the Hamilton-Jacobi theory. Starting with a generating function

\[ S = K \ln \Psi \]  

in the Hamilton equation

\[ H(q, \partial S / \partial q) = E \]  

he reduced the quantization of the hydrogen atom to an eigenvalue problem for the time independent Schroedinger equation. With this done he then solved the harmonic oscillator, established the wave mechanical nature of the theory, produced the necessary perturbation theory, arrived at a time dependent equation, and, in an intervening paper,\textsuperscript{24} showed the formal identity of matrix mechanics to wave mechanics. Since it gave an analytic method for obtaining a description of physical systems the wave mechanics soon became the way to solve problems and by 1930 most of the standard results had been obtained.\textsuperscript{17}
In the meantime, the existence of a proper quantum treatment of the harmonic oscillator permitted the photon to become a full physical theory\textsuperscript{25} rather than a heuristic device. Implicit in Dirac's development was an operator formalism which of its nature precludes the study of the instantaneous matter-radiation interaction. While we could trace the history of quantum electrodynamics into the forties,\textsuperscript{26-28} we have reached the point at which this experiment fits.

What of the coupling of matter and radiation? This involves our interpretation of $\psi$ and, more importantly, just what we expect our description of nature to accomplish. Those who wish philosophical discussions of measurement theory are referred elsewhere,\textsuperscript{29-31} but we must treat some of the problem here since the philosophical implications determine how we cut metal.

Schrödinger felt initially that $e^\phi\psi^\dagger$ represented the charge density in configuration space\textsuperscript{23} and he showed briefly how to arrive at the oscillator strengths from this standpoint.\textsuperscript{24} The difficulty with this is apparent in two related ways. Firstly, the temptation is to keep the notion of trajectory by looking at wave packets.
This runs counter to the ascendant philosophical notion that what is not observable must not be included in the theory, although the complete elimination of continuous processes did not occur until the fifth Solvay Conference. More importantly, wave packets do not work in the solution of collision problems since the packet always spreads out—with the exception of the harmonic oscillator functions used by Schroedinger to test the concept.

Born, at the time, was working in the same building as Franck. As a result he firmly believed that the correct interpretation could leave no fuzziness about the nature of an observed electron: it was a particle. Returning to de Broglie's guide wave concept he interpreted $\psi\psi^\dagger$ as giving the probability density of finding the actual particle at the point in configuration space involved, and developed the Born approximation for describing collisions. While Schroedinger pointed out that such an interpretation required an additional hypothesis to his development, the interpretation was too fruitful to be rejected.

The fundamental problem here is psychological, although it may be couched in epistemological terms. The
strict probabilistic interpretation of quantum mechanics requires that we give up knowing or even talking about how one event leads to another, even though we have a definite prescription for predicting the possible sequences of events. Bohr covered this difficulty with the principle of complementarity, but the problem always reverts to our basic notion of what it means "to understand". If one can attach a dynamical significance to $\Psi$ then he understands in a more satisfying sense than if he only calculates ensemble probability averages. Those influenced directly by Bohr and hence by the philosophies of Kierkegaard and James$^{3,5}$ were able to sublimate these difficulties, but each generation of students must face them anew.

A second, more physical problem, arises from the nature of modern quantum electrodynamical theory. It works well,$^{3,6}$ but often appears to sweep difficulties away in a manner reminiscent of the old quantum theory.

The experiment described below looks at an alternative to the canonical interpretation. The photon, whether accepted or not, was part of the world for over twenty years before the quantum theory of matter was
workable. Is it not possible that all photon effects are really in the quantized eye of the beholder? Such explanations account for the photoelectric effect\(^3\)\(^7\) and a number of other supposed QED effects.\(^3\)\(^8\) By the same token, a return to the Schroedinger interpretation of would give time varying sources to produce radiation.

Such a theory--more appropriately, a philosophy--has been advanced by E. T. Jaynes and his collaborators.\(^3\)\(^9\)-\(^4\)\(^1\) We propose a direct test of this theory. In the process we will also be able to look for limitations on the exponential decay law. To fix in the reader's mind this purpose, we digress here to give an overview of the experiment.

The Jaynes' theory predicts a radiative decay law which differs from that of QED. For the excited state densities available to us this would be observed as a linear variation in the lifetime of the \(2p_{1/2}\) state as a function of the presence of the metastable \(2s_{1/2}\) state. To observe this, a beam of 60 keV hydrogen atoms was produced which consisted of about 10% atoms in the metastable state. (We use the usual beam interpretation here.) If a controlled fraction of these was converted to the \(2p_{1/2}\) state
by a short electrical pulse, the subsequent radiation could be monitored to directly measure the changes in the lifetime.

We begin now by looking at the predictions of QED and the Jaynes' theory concerning the behavior of atomic systems.
The statement "light exists as photons" does not alone make a quantum electrodynamics. Between the proposal in 1905 that light must be treated as energy quanta and the full quantization of the radiation field in 1927 no proper formalism could be generated for the simple reason that no well defined quantum mechanical structure existed. As soon as an acceptable theory of the harmonic oscillator was developed the photon was established as part of a working physical theory. Let us now derive the exponential decay law in terms of this theory for the atomic system used in these experiments.

We consider a volume of space $\Omega$ containing an ensemble of atoms and electromagnetic radiation. The atoms obey a Hamiltonian $H_0$ which generates stationary states in the absence of the radiation field:

$$H_0 \phi_k(\mathbf{r}) = \omega_k \phi_k(\mathbf{r}).$$

(1)
(Throughout this paper, the convention $\hbar = 1 = c$ is used, so that energies, frequencies and wave numbers may be used interchangeably.)

The radiation fields are source free and completely described by a transverse vector potential $\hat{A}$. (This is more carefully treated in chapter III.) $\hat{A}$ is to be written in terms of photon states. Consider the possibilities: The "finite" volume $\Omega$ defines a lattice of possible wave vectors $\vec{k}$, each of which may have two orthogonal polarizations $\varepsilon(\vec{k}, \lambda)$. Each of these may be "occupied" by 0, 1, 2... photons. To produce a dynamics, raising and lowering operators, $a^\dagger(\vec{k}, \lambda)$ and $a(\vec{k}, \lambda)$ are defined which have the following properties: Let $| \cdots \text{N} (\vec{k}, \lambda) \cdots \rangle$ correspond to a photon state in which there are N photons of wave vector $\vec{k}$ and polarization $\lambda$ along with other unchanging occupations. Then

$$a^\dagger(\vec{k}, \lambda)| \cdots \text{N} (\vec{k}, \lambda) \cdots \rangle = \sqrt{N+1} | \cdots \text{(N+1)} (\vec{k}, \lambda) \cdots \rangle,$$  \hspace{2cm} (2)

$$a(\vec{k}, \lambda)| \cdots \text{N} (\vec{k}, \lambda) \cdots \rangle = \sqrt{N} | \cdots \text{(N-1)} (\vec{k}, \lambda) \cdots \rangle,$$  \hspace{2cm} (3)

and

$$a^\dagger(\vec{k}, \lambda)a(\vec{k}, \lambda)| \cdots \text{N} (\vec{k}, \lambda) \cdots \rangle = \text{N} | \cdots \text{N} (\vec{k}, \lambda) \cdots \rangle.$$  \hspace{2cm} (4)

This last combination is called the number operator and is used to generate the radiation Hamiltonian.
It then follows that

\[
\hat{\mathbf{A}} = \sum_{\mathbf{k},\lambda} \sqrt{2\pi/\Omega k} (a(\mathbf{k},\lambda) e^{i\mathbf{k}\cdot\mathbf{r}} + a^\dagger(\mathbf{k},\lambda) e^{-i\mathbf{k}\cdot\mathbf{r}}) \hat{\epsilon}(\mathbf{k},\lambda),
\]

where the operator amplitudes contain an implicit time dependence:

\[a(\mathbf{k},\lambda) = a(\mathbf{k},\lambda)|_{t=0} e^{-ikt}.\]

Note that \(\hat{\mathbf{A}}\) is transverse:

\[\nabla\cdot\hat{\mathbf{A}} = 0.\]

The energy of our portion of the world is

\[E = (\frac{\hbar}{2m})(\hat{\mathbf{p}} + e\hat{\mathbf{A}})^2 + E_{\text{atomic}} + E_{\text{rad}},\]

so that to first order in \(\hat{\mathbf{A}}\) the perturbed atomic Hamiltonian is

\[H = H_0 + (e/m)\hat{\mathbf{A}}\cdot\hat{\mathbf{p}} = H_0 + V.\]

Here, \(-e/m\) is the electronic charge to mass ratio.

Hence,

\[V = (e/m)\sum_{\mathbf{k},\lambda} \sqrt{2\pi/\Omega k} (a(\mathbf{k},\lambda) e^{i\mathbf{k}\cdot\mathbf{r}} + a^\dagger(\mathbf{k},\lambda) e^{-i\mathbf{k}\cdot\mathbf{r}}) \hat{p} \cdot \hat{\epsilon}(\mathbf{k},\lambda)\]

is the perturbing potential.

In our work we will be populating three atomic states: \(\phi_1(\mathbf{r})\) will be the ground state; \(\phi_2(\mathbf{r})\) will lie
\(\omega_0 + \omega\) above \(\Phi_1\) but not coupled to it by \(V\); and \(\Phi_3(\mathbf{r})\)
will lie \(\omega\) above \(\Phi_1\) and be coupled to both \(\Phi_2\) and \(\Phi_1\).
Further, \(\omega_0 \ll \omega\). These are illustrated in figure one.
Initially the atom will be in a mixture of \(\Phi_2\) and \(\Phi_3\)
with no photons in the radiation field. The general
state will then be
\[
\Psi(\mathbf{r}, t) = \sum_{\mathbf{k}, \lambda} c_1(\mathbf{k}, \lambda; t)e^{-i\mathbf{k} \cdot \mathbf{r}}|\mathbf{k}, \lambda> + c_2(t)e^{-i(\omega_0 + \omega)t}\Phi_2(\mathbf{r})|\phi>
+ c_3(t)e^{-i\omega t}\Phi_3(\mathbf{r})|\phi>,
\]
where \(|\mathbf{k}, \lambda>\) designates the existence of one photon of
wave vector \(\mathbf{k}\) and polarization \(\lambda\) and no others, and
\(|\phi>\) is the vacuum state. The initial conditions are
\[
c_1(\mathbf{k}, \lambda, 0) = 0 \quad (9a)
\]
\[
|c_2(0)|^2 + |c_3(0)|^2 = 1. \quad (9b)
\]
The Schrödinger equation gives
\[
i\dot{c}_1(\mathbf{k}, \lambda; t) = \sqrt{2\pi/\hbar k}(e/m)\mathbf{E}(\mathbf{k}, \lambda) \cdot <1|e^{-i\mathbf{k} \cdot \mathbf{r}}|3>e^{-i(\omega - k)t}c_3(t), \quad (10)
\]
\[
i\dot{c}_2(t) = 0, \quad (11)
\]
and
\[ \dot{c}_3(t) = \sum_{\mathbf{k}, \lambda} \sqrt{2\pi/\Omega k} (e/m) \hat{e}(\mathbf{k}, \lambda). \]

\[ \langle 3 | e^{i \mathbf{k} \cdot \mathbf{r}_p} | 1 \rangle e^{i(\omega - k)t} c_1(\mathbf{k}, \lambda; t). \] (12)

The coupling between states \( \Phi_2 \) and \( \Phi_3 \) has been ignored since, as we shall see, it is down by a factor of \( 10^{18} \) over the other coupling. To this approximation, we have a linear two state problem; the presence of residual "metastable" atoms does not affect the behavior of the radiating state.

To solve the system of equations we use the fact that 
\[ c_1(\mathbf{k}, \lambda; 0) = 0 \] to obtain

\[ c_1(\mathbf{k}, \lambda; t) = -i\sqrt{2\pi/\Omega k} (e/m) \hat{e}(\mathbf{k}, \lambda) \langle 1 | e^{-i \mathbf{k} \cdot \mathbf{r}_p} | 3 \rangle \times \]

\[ \int_0^t c_3(\tau) e^{-i(\omega - k)\tau} d\tau, \]

which, in turn, is to be substituted into equation 12. The equivalence 
\[ \sum_{\mathbf{k}} \rightarrow (\Omega/8\pi^3) \int d^3k \]
gives

\[ \dot{c}_3(t) = -\int_0^t e^{i \omega(t-\tau)} c_3(\tau) \int_0^\infty |V(k)|^2 e^{-ik(t-\tau)} dk d\tau \] (13)

where
\[ |V(k)|^2 = (e/m)^2 (k/4\pi^2) \sum_\lambda \int d\Omega_k |<1|e^{-ik \cdot \hat{r}_p} \cdot \epsilon(\hat{k},\lambda)|3>|^2 \] (14)

is called the form factor.

Several possibilities present themselves. An exact solution is possible in principle. The expression in equation 13 can be made into a convolution and \( c_3(t) \) obtained by Laplace transforms. The result is exponential decay with several shortlived transients and a small asymptotic \( t^{-2} \) behavior. Another alternative is to do a dipole approximation to \( |V(k)|^2 \) and then proceed "exactly". The traditional approach has been to make the dipole approximation and to assume exponential decay:

\[ c_3(t) \propto e^{-\frac{1}{2} \Gamma t} \] (15)

The rate \( \Gamma \) is adjusted to achieve self consistency in equation 13 for times great compared to \( \omega^{-1} \). This is sufficient for our purposes.

Equation 13 is now

\[ \frac{1}{2} \Gamma e^{-\frac{1}{2} \Gamma t} = \int_0^t e^{i \omega(t-\tau)} e^{-\frac{1}{2} \Gamma \tau} \int_0^\infty |V(k)|^2 e^{-ik(t-\tau)} dk d\tau \]

\[ = i e^{-\frac{1}{2} \Gamma t} \int_0^\infty \int_0^\infty |V(k)|^2 \left( \frac{1 - e^{i(\omega-k-\frac{1}{2} \Gamma) t}}{\omega - k - \frac{1}{2} \Gamma i} \right) dk, \] (16)
and since \( r \ll \omega \) we have

\[
\frac{1}{2} \Gamma = \int_0^\infty \left\{ \frac{\sin(\omega-k)t}{(\omega-k)} \right\} |V(k)|^2 dk + \\
i \int_0^\infty \left\{ \frac{(1 - \cos(\omega-k)t)/(\omega-k)}{V(k)|^2} \right\} dk.
\]

For \( \omega t \gg 1 \) the oscillatory parts of the integrands have the effect of sharply defining the region about \( \omega \). The real part becomes

\[
\text{Re}\Gamma = \gamma = 2\pi \int_0^\infty \delta(\omega-k) |V(k)|^2 dk = 2\pi |V(\omega)|^2.
\]

Finally, in the dipole approximation

\[
|V(\omega)|^2 = \frac{1}{4\pi^2} \sum_k |d\Omega_k| \langle 1 | [H, \mu] | 3 \rangle \cdot \delta(\vec{k}, \lambda)|^2
\]

\[
= \frac{1}{3\pi} |\mu_{13}|^2
\]

where \( \mu \) is the electronic dipole moment. Hence,

\[
\gamma \approx \frac{4\omega^3}{3} |\mu_{13}|^2.
\]

The atomic system will radiate energy at a rate

\[
R = -\frac{d}{dt} |c_3(t)|^2 \omega = \omega \gamma |c_3(0)|^2 e^{-\gamma t},
\]

an expression which is generally accepted as true and which is tested here. We now turn to a somewhat different analysis of the same problem.
Figure one: Three level atom. This idealized hydrogen structure forms the basis for the research done. A full radiation theory must predict $\omega_0$ since the upper levels are degenerate in the ordinary atomic theory. For the work here the levels should be identified with the hydrogenic $1s_{1/2}$, $2s_{1/2}$ and $2p_{1/2}$ levels, respectively.
CHAPTER III
RADIATION THEORY II
SEMICLASSICAL THEORY

A complete quantum theory of radiation was not uniquely difficult. The consistent treatment of the interaction of classical electromagnetic fields and matter was not fully accomplished until the 1960s.46 The interaction of classical fields and quantized matter has been studied for an equal period and the reader is directed to the review article by Scully and Sargent for a bibliography.38 We concern ourselves here with the developments of E. T. Jaynes and his students,39-41 although we shall use an exposition which follows a slightly different approach.47

The semiclassical radiation theory (SCT) of Jaynes differs from earlier treatments48 in that it studies the dynamics of the interaction in detail. This is done at the expense of the standard interpretation of $\psi^\dagger \psi$ as a probability density over an ensemble of identical atoms. Rather, SCT follows Schroedinger's original notion that

$$\rho = -e\psi^\dagger \psi$$  \hspace{1cm} (1)
is the electronic charge density in configuration space.\textsuperscript{49} For a single electron atom this is a real spatial charge distribution. From this and the usual notion of continuity, the current density is

\[ \mathbf{j} = \text{Re}\{ (-e/m) (\psi^\dagger \mathbf{p} \psi) \}. \]  

(2)

These are point functions, not matrix elements, and act as sources for electromagnetic fields.

This drastically differs from the QED starting point. There the fields derive from photon states which exist independently of the atom. The current density matrix element determines the coupling to the photon states, but cannot be said to be responsible for their existence.

One implication of this difference can be seen quickly. Let us assume we have a dipole transition which would couple the states of a two level atom. In QED if an ensemble of atoms were initially in the excited level, we would see radiation immediately with a rate given by

\[ R = N \omega \gamma e^{-\gamma t}, \]  

(3)

where \( \omega \) is the energy separation of the states and

\[ \gamma = (4/3) \mu^2 \omega^3. \]  

(4)

The dipole strength enters only as the matrix element

\[ \mu^2 = |\langle 1 | e \mathbf{r} | 2 \rangle|^2. \]  

(5)
For an SCT treatment, as we shall see, the radiation rate depends on a physically oscillating dipole and to the same approximation we observe at a large distance,

$$R \propto \left| \int \psi^\dagger \mu \psi d^3r \right|^2.$$  

If $\psi$ is initially in a single parity state this vanishes. Spontaneous decay can occur only if there is some residual ground state present to give a nonvanishing dipole to initiate the process. This clearly precludes a purely exponential decay. The reason such behavior is not observed stems from the impossibility of producing pure excited states.

This point is tricky. In SCT when we speak of an atomic beam which consists of say 12% atoms in some state, $\phi_\alpha$, we do not mean that 12% of the atoms are in the state $\phi_\alpha$ and the remaining 88% in other states. Rather, we mean that each atom has a wave function which on the average is 12% $\phi_\alpha$. This makes the discrepancy from QED difficult to test. Most of the other possible problems are treated by Scully in his Physics Today review. An effect does exist which can be realized experimentally, as we shall now show.
The electric and magnetic field strengths at a point may be written in terms of longitudinal and transverse components:

\[ \vec{E} = \vec{E}_L + \vec{E}_T \]  
\[ \vec{B} = \vec{B}_T \]

where

\[ \nabla \cdot \vec{E}_T = 0, \]  
\[ \nabla \times \vec{E}_L = 0, \]  
\[ \nabla \cdot \vec{B}_T = \nabla \cdot \vec{B} = 0. \]

In terms of these vectors and their sources Maxwell's equations are

\[ \nabla \cdot \vec{E}_L = 4\pi \rho, \]  
\[ \frac{\partial \vec{E}_L}{\partial t} = -4\pi \vec{j}_L, \]  
\[ \nabla \times \vec{E}_T = -\frac{\partial \vec{B}_T}{\partial t}, \]

and

\[ \nabla \times \vec{B}_T = \frac{\partial \vec{E}_T}{\partial t} + 4\pi \vec{j}_T. \]

Since it is the longitudinal part of \( \vec{E} \) which contains all the source dependence, \( \vec{E}_T \) and \( \vec{B}_T \) are sufficient to describe the radiation problem.
This description is best carried out by introducing a vector potential $\mathbf{A}(\mathbf{r},t)$ in the Coulomb gage:

$$\nabla \cdot \mathbf{A} = 0$$  \hspace{1cm} (10a)

$$\mathbf{B}_t = \nabla \times \mathbf{A}$$  \hspace{1cm} (10b)

$$\mathbf{E}_t = -\frac{\partial \mathbf{A}}{\partial t}.$$  \hspace{1cm} (10c)

When this is introduced into equation 9d we have

$$\{\nabla^2 - \frac{\partial^2}{\partial t^2}\} \mathbf{A} = -4\pi \mathbf{j}_t$$  \hspace{1cm} (11)

for which the appropriate solution is

$$\mathbf{A}(\mathbf{r},t) = -\int \mathbf{R}^{-1} \mathbf{j}_t(\mathbf{r}',t-R)d^3\mathbf{r}'$$  \hspace{1cm} (12)

where

$$R = |\mathbf{r} - \mathbf{r}'|.$$  \hspace{1cm} (13)

The retarded solution is deliberately chosen since it is the only meaningful one in this context.

This expression must now be substituted into the Schroedinger equation with the Hamiltonian

$$H = H_0 - \mathbf{A} \cdot \mathbf{j}_t$$  \hspace{1cm} (14)

where $H_0$ is the unperturbed atomic Hamiltonian and $\mathbf{A}$ and $\mathbf{j}_t$ are determined by equations 12 and 2. The result is a nonlinear theory, which is to be used to solve the same
problem as in chapter II with one difference: the two upper levels will be degenerate, \( \omega_0 = 0 \). This system and the couplings are shown in figure one.

The total wave function is written as

\[
\psi(\vec{r}, t) = c_1(t) \phi_1(\vec{r}) + c_2(t)e^{-i\omega t} \phi_2(\vec{r}) + c_3(t)e^{-i\omega t} \phi_3(\vec{r}),
\]

and we are interested in the time evolution of the quantities

\[
N(t) = \sum_i |c_i(t)|^2
\]  
and

\[
E(t) = \sum_i \omega_i |c_i(t)|^2.
\]

where \( N(t) \) is normalization of \( \psi(\vec{r}, t) \) and

\[
H_0 \phi_i(\vec{r}) = \omega_i \phi_i(\vec{r}).
\]

We now show that \( N(t) \) is unity even when the atom radiates and then compare \( dE/dt \) with the same quantity in chapter II.

The Schroedinger equation gives

\[
i\dot{c}_i(t) = - \sum_k c_k(t)e^{-i\omega k t} \int \phi_j^+(\vec{r}) \hat{A} \cdot \hat{j} \phi_k(\vec{r}) d^3r
\]

where

\[
\omega_{kj} = \omega_k - \omega_j.
\]
The current density given in equation 2 can also be written as
\[ j(\mathbf{r}, t) = i\psi^{*}(\mathbf{r}, t)[H, \hat{\mu}_{t}] \psi(\mathbf{r}, t) \] (20)
so that the full expansion of equation 19 is
\[ \dot{c}_{j}(t) = \sum_{k} \sum_{\ell} \sum_{m} c_{k}(t) \omega_{jk} \omega_{\ell m} e^{-i\omega_{jk} t - i\omega_{\ell m} t} M(j, k; \ell, m) \] (21)
where
\[ M(j, k; \ell, m) = \int d^{3}r d^{3}r' c_{\ell}^{*}(t-R) c_{m}^{*}(t-R) R^{-1} e^{-i\omega_{\ell m} t} \{ \phi_{j}^{*}(\mathbf{r}) \mu_{t}^{*} \phi_{k}(\mathbf{r}) \} \cdot \{ \phi_{\ell}^{*}(\mathbf{r}') \mu_{t} \phi_{m}(\mathbf{r}') \}. \] (22)

The first thing to do is eliminate the oscillating terms.

The form of equation 21 is such that we may ignore the degeneracy and obtain
\[ \dot{c}_{j}(t) = i\sum_{k} c_{k}(t) \omega_{jk}^{2} M(j, k; j, k). \] (23)

To evaluate the expressions in equations 16 and 17 we need to find
\[ \frac{d}{dt}|c_{j}(t)|^{2} = c_{j}^{*} c_{j} + c_{j} c_{j}^{*}. \] (24)

We use equation 23 to evaluate this with two further approximations: The variation of $c_{j}(t)$ over the time in
the integration of the retardation is negligible, and we may also use

\[ \frac{\sin \omega_{jk} R}{\omega_{jk} R} \approx 1. \]  \hspace{1cm} (25)

Then we find

\[ \frac{d}{dt} |c_j(t)|^2 = - \frac{4}{3} \omega_{jk}^2 |c_k(t)|^2 |\mu_{jk}|^2 |c_j(t)|^2. \]  \hspace{1cm} (26)

Here, \( \mu_{jk} \) is the total dipole matrix element; the transverse part has been extracted by integrating over all solid angles relative to the direction of propagation of the radiation.

Because \( \omega_{jk} = -\omega_{kj} \), we see that

\[ \frac{dN(t)}{dt} \equiv 0 \]

so that an initially normalized wave function will remain so. For our problem we find

\[ \frac{dE(t)}{dt} = - \frac{4\omega^4}{3} |c_1(t)|^2 |c_3(t)|^2, \]  \hspace{1cm} (27)

which differs from the QED result in that both the initial and final state amplitudes affect the rate.

The solution of the coupled equations in equation 26 is a straightforward exercise. We first introduce a density matrix notation

\[ \sigma_{k\ell}(t) = c_k(t)c_{\ell}^*(t), \]  \hspace{1cm} (28)
and then write

\[ \dot{\sigma}_{11}(t) = \gamma \sigma_{11}(t) \sigma_{33}(t), \quad (29a) \]

\[ \dot{\sigma}_{22}(t) = 0, \quad (29b) \]

\[ \dot{\sigma}_{33}(t) = - \gamma \sigma_{33}(t) \sigma_{11}(t), \quad (29c) \]

where

\[ \gamma = (4w^3/3)\mu_1^3 \]

is the same characteristic rate as in QED. These coupled equations are subject to the conditions

\[ \sigma_{11}(t) + \sigma_{22}(t) + \sigma_{33}(t) = 1 \quad (30a) \]

and

\[ \sigma_{22}(0) = \sigma_2. \quad (30b) \]

We can solve equation 29b and then use equation 30a to eliminate \( \sigma_{11}(t) \) from equation 29c:

\[ \dot{\sigma}_{33}(t) = - \gamma \sigma_{33}(t) \eta \sigma_{33}(t), \quad (31) \]

where

\[ \eta = 1 - \sigma_2. \quad (32) \]

This integrates to

\[ \eta \gamma (t - t_0) = \ln(\eta - \sigma_{33}(t))/\sigma_{33}(t) \quad (33) \]

where

\[ \eta \gamma t_0 = \ln(\sigma_{33}(0)/\sigma_{11}(0)). \quad (34) \]
Hence,

\[ \sigma_{33}(t) = \frac{\eta}{1 + e^{\eta \gamma (t-t_0)}}. \]  

\(35\)

The presence of the residual metastable component and the initial excited state population both affect the time evolution of the system. The time \(t_0\) in a typical experiment where

\[ \sigma_{33}(0)/\sigma_{11}(0) \approx 0.1 \]

and

\[ \eta \approx 1 \]

is

\[ t_0 \approx -2.3/\gamma \]

so that \(\sigma_{33}(t)\) is described by its asymptotic behavior

\[ \sigma_{33}(t) \sim \eta \{\sigma_{33}(0)/\sigma_{11}(0)\} e^{-\eta \gamma t}. \]  

\(36\)

We expect to see exponential decay, but both the rate and the apparent initial population are affected by a quantity which was totally irrelevant in the QED derivation. It is on the basis of this prediction that we can test SCT, for we can produce a beam of atomic hydrogen with a metastable component of about ten percent. By dumping different fractions of these into the \(2p_{3/2}\) state and looking at the
subsequent decay rate we shall be able to look for changes in the rate as a function of $\sigma_2$ after the dump. The next two chapters describe the apparatus necessary; and the results appear in chapter VI.
CHAPTER IV
APPARATUS

So much for the nature of the problem; now to the equipment designed to test it. This apparatus was required to produce a well-collimated beam of hydrogen atoms with a known velocity and metastable fraction. It had to convert a variable fraction of the metastables into the $2p_{\frac{1}{2}}$ state in a time short compared to the $2p_{\frac{1}{2}}$ lifetime and without significantly populating any other states; and it was equipped to monitor the subsequent radiation in a statistically reliable way. The conversion problem will be treated in the following chapter; the rest of the apparatus is the subject here.

A prerequisite for a fast hydrogen beam is a fast proton beam. This was produced by a Cockcroft-Walton accelerator and analyzing system described elsewhere.\textsuperscript{52} Figures two and three show the rest of the system.

At the entrance to the charge exchange cell X, the 60 keV protons entered a region containing $H_2$ gas at a pressure of about 3.0 $\mu$ where approximately 20% picked up an electron and became neutral hydrogen in some quantum state. From there the mixed beam entered a region D containing two parallel plates spaced 1.0 cm apart and
Figure two: Schematic view of the apparatus. The beam is collimated and mass analyzed when it enters the exchange cell (X). Differential pumping on either side of the exchange cell was accomplished by a 4" main diffusion pump operating on a manifold and assisted on the downstream side by a 2" pump. The "As" are the apertures.
FIG 2

To MKS Gage

A₁

A₂

D

A₃

H⁺

H₂

2" Pump

4" Pump

C
Figure three: Experimental chamber interior. For clarity the light collector (C) is shown as a cone; the actual geometry appears in figure nine. The beam monitor (B) detail is shown in figure twelve. The plates (Q) following the electrodes were added after the decay curves were taken.
39.5 cm long. These were charged just enough to sweep all the remaining protons from the beam. The pressure there was maintained at $5 \times 10^{-6}$ T. Next the beam passed through an aperture into the experimental chamber C. The instrumentation there permitted the measurement of the radiation per atom and the indirect measurements of the 2p and 2s components of the beam. It consisted of a prequench electrode P, conversion electrodes E, a movable Lyman alpha detector C, and the beam current detector B. After the basic experiment was finished two long parallel plates Q were installed after the conversion electrodes for calibration purposes. These processes will now be discussed in detail beginning with the exchange cell.

The beam of protons entered the exchange region through a .041" movable aperture which further collimated the beam. The other end of the exchange cell had a similar aperture .036" in diameter and was located 10 cm downstream. The exchange cell gas, in this case H$_2$, was admitted through a Granville Phillips Variable Leak and monitored by an MKS Baratron pressure meter referenced to the throat of the beam tube diffusion pump. The exchange cell pressure could be conveniently maintained between 0.5 and 10 \(\mu\). The systematic production of
metastable atoms required that beam atoms suffer only one collision. The graph in figure four illustrates the relative metastable production as a function of exchange cell pressure. The initial rise shows the gradual predominance of H₂ over the residual gases; the plateau is the single collision region; and collisional quenching becomes a factor thereafter. The differential pumping was not adequate to pursue the details of this process.

The largest uncertainty in this experiment has to do with the percentage of the neutral beam in the metastable 2s₁/₂ state. Figure five shows this beam richness versus energy as inferred from the absolute cross section measurements of Bayfield. Bayfield worked at exchange cell thicknesses on the order of 0.2 μ-cm, but the plateau shown in figure four justifies the 30 μ-cm value used to obtain a large count rate. He lists an uncertainty other than calibration of 20%. The potential calibration error is given as 55%. The least value for the richness consistent with other work is about 2% at 30 keV. Since the destruction cross section seems to be decreasing and the production is definitely increasing, it is appropriate to infer 2% as a lower limit but to quote results in terms of the 11.15% shown in figure five.
Figure four: Exchange cell characteristics. The data taken over the first micron correspond to a mixture of $\text{H}_2$ and background gas. Beyond that the system is flushed. The nonlinearity at 3.0 $\mu$ is approximately 10% as measured by the $L_\alpha$ counts per second.
Figure five: Metastable fraction. The absolute cross section measurements given in reference fifty-four were used to infer the beam richness as a function of energy. The "richness" is defined as $\sigma_{2S}/\sigma_{\text{neutral}}$. 
FIG 5
with a 20% uncertainty. As it happens the final results of the experiment are conclusive to well below a 1% level.

Upon leaving the charge exchange cell the mixed beam entered a long sweep region. Here an electric field of approximately 30 V/cm swept out all the residual protons. This was necessary since it was discovered that the presence of any protons destroyed the integrity of the counting statistics. This electric field is the smallest consistent with the requirement of pure Poisson statistics. The graph of the counting rate observed versus the potential across the plates shown in figure six is consistent with the low field limit developed in appendix A. From this graph it is seen that 78% of the metastables produced actually enter the experimental region through the last aperture.

This last aperture had a .020" hole which collimated the beam completely within the .041" hole in the electrode assembly. Each of these apertures could be grounded through an ammeter to help locate the beam. The three inches of beam tube prior to the first aperture was clear plastic so that at pressures around $10^{-4}$T the actual shape of the beam there could be observed.
Figure six: Sweep destruction. By plotting the photon count rate as a function of the sweep voltage squared, the fraction of the metastable atoms destroyed in the sweep region was determined.
FIG 6
A schematic view of the experimental chamber is shown in figure three. All the parts were mounted on an aluminum plate which could be leveled and rotated for optimum alignment. The chamber itself was a 10\(\frac{1}{2}\)" ID by 6" brass pot which was mounted directly over the gate to a 4" diffusion pump system. The description of the parts follows in the order of their appearance to the beam.

The prequench electrode permitted the effects of a neutral ground state beam to be observed. It was a single flat piece of copper placed 0.83 cm below the top of a piece of square wave guide. (See figure seven.) When this was charged to about 1500 volts negative the two-inch length was sufficient to eliminate any metastables which might be observed. This prequench was used to differentiate between the background signal due to the beam presence alone and that caused by inadvertent quenching of metastables. It could also be used in the calibration process for the electrodes and the beam richness measurements shown in figures four and six. The background signal measured this way was very stable and was also position independent over about ten lifetimes along the beam.
Figure seven: Prequench electrode. The ground plane formed by standard .720" square waveguide was 0.83 cm above the copper electrode. The structure was 5 cm long, and the electrode was 1.82 cm wide.
The electrode structure was located 2.5 inches downstream from the last aperture. The details of the actual electrode assembly are deferred to chapter V but some description is necessary for clarity. The assembly was constructed on a piece of .032" copper which was then mounted on a brass support. This support was \( \frac{3}{4} \)" thick and isolated from ground by a piece of lucite. As a consequence the fraction of the beam which hit the structure could be determined to within a factor of two or three which is due to secondary electron emission characteristics. The beam at the third aperture was about two millimeters broad and as that aperture was moved across this profile a pronounced minimum was observed in the electrode current. With a secondary electron current at the beam detector of 200 nA the electrode "current" was held between 0.1 and 0.2 nA. The beam filled between 80 and 90\% of the aperture.

The potential difference across the electrodes applied a short electrical pulse to the atoms. This converted, or mixed depending on the interpretation, some of the metastables into the \( 2p_{3/2} \) state. This fraction subsequently decayed to the ground state by
emitting Lyman alpha ($L_\alpha$) radiation in the field free downstream region. The light was detected by a Bendix channeltron electron multiplier photon detector (CEM). (The use of a "photon" detector to test a classical radiation theory was made possible by the fact that the photoelectric effect can be explained in terms of classical light waves and quantized matter states.) This device is an excellent uv light detector and is even better as a charged particle detector. As a consequence, it had to be completely enclosed in a metal case except for an entrance slit covered by a LiF window. The response for a typical combination of window and detector is shown in figure eight. The only spectral line which will show a position and beam intensity dependence and be detected is $L_\alpha$.

The light collection geometry is shown in figure nine, and the effect of the finite slit width is discussed in appendix B. An incident uv light pulse gives an electrical output pulse on the order of 50 mV with negative polarity. This was fed into a standard NIM scintillation preamplifier which gave it the appropriate shape for use in an NIM pulse amplifier. This then fed a scaler/timer.
Figure eight: Detector response. The transmission properties of LiF\textsuperscript{57} and the sensitivity of the CEM detector\textsuperscript{58} can be combined to give the optical window of the L\textsubscript{\textalpha} detector.
Figure nine: Light collection geometry. The CEM aperture was nominally 1.0 mm. The slit was 0.32 cm high and constructed of .002" shim stock pieces. The LiF window was between the slit and the detector.
These steps are outlined in figure ten. The electrical circuit for the channeltron is shown in figure eleven.

Light from any point along the beam could be monitored by moving the detector which was mounted on an aluminum cart that traveled on linear bushings. The cart was pushed upstream by a rod which entered the vacuum through an O-ring grease seal and returned by the pull of a weight which hung into the throat of the chamber diffusion pump. The position of the cart was set with a micrometer head at the end of the rod. With the use of spacers a total throw of five inches could be used if desired.

The neutral beam current was measured by the secondary electrons ejected from an aluminum target. The ejection efficiency for neutral atoms in the kilovolt range runs from 1.10 to 1.25 times the easily measured value for protons.\textsuperscript{59} For a fixed energy experiment this variation is unimportant. The Faraday cage system is shown in figure twelve. It could be set to read the current to the target and suppress the electron emission or to enhance the secondary electron current. In the former setting a neutral beam appeared to be a slightly negative current. This has been observed before,\textsuperscript{60} and
Figure ten: Electronic flow chart. The signals to be processed were a varying dc current and random negative 50 mV pulses. The recorder output of the ammeter was converted to a proportional frequency which was counted to integrate the beam charge on scaler B. Scaler A would either count a preset number of pulses while displaying the elapsed time or would count for a preset length of time.
Figure eleven: CEM circuit. The current pulse put out by the CEM is converted to a voltage by the 1 MΩ load resistor. The capacitor blocks the 2600 V potential applied to the CEM and provides a low impedance path for the pulse. The voltage divider made possible the use of two detectors during the preliminary studies.
FIG 12

2.9 kV in

2.2 M

1 M

.003 μf

4 M

To CEM

Signal out
Figure twelve: Beam detector. The neutral beam was detected by a modified Faraday cage. When the potential applied to the cup was reversed, the secondary electrons were repelled back to the target giving a true measure of the charged beam component. Only the two relevant switch positions are shown. The collection characteristics are given in figure thirteen.
Figure thirteen: Collector characteristics. The current registered by the meter is shown for a fixed proton current and a variable cup potential. Note the break in the scale.
FIG 13

With increasing cup potential (V), the meter current (μA) decreases (Suppress) until a point is reached where it begins to increase sharply (Enhance).
may be due to neutrals which bounce off the target and eject electrons from the cup. A spot observed on the inside of the cup supports this idea. The second configuration was used for the neutral beam.

In either case, the meter shown in figure twelve was a Kiethley electrometer, and the shielding was sufficient to permit sub nanoampere readings. The proton currents were about 200 nA as were the secondary electron currents for the neutral beam. The electrometer has a recorder output of zero to one volt. This was fed into a VIDAR voltage to frequency converter which puts out an asymmetrical wave form at 10 kHz/V. When this signal was counted by a scaler it effectively integrated the beam current. As a result, the light measurements could be normalized to the number of beam atoms past the detector. All rates shown are in counts/nA-sec of secondary electrons.

The data taking proceeded as follows: The beam was adjusted until the current to the electrode structure was a minimum. The electrode voltage was set and the CEM located at the furthest point along the beam desired. A total radiation count of 10 000 was made and the time
and total secondary electron charge was recorded. The scalers could then be set to either count charge or tenths of seconds to one significant digit, i.e. $N \times 10^M$. Whichever gave a better approximation to the actual elapsed time was used to determine the background. The micrometer head was then moved upstream 0.100" and the process repeated. A total sweep of 0.500" corresponded to 2.3 lifetimes and gave the decay rate to 0.5%.

After the lifetime measurements had been made it was necessary to modify the chamber to calibrate the electrode conversion efficiency. This was done by installing two plates immediately after the electrode structure. When a voltage was applied across these, radiation was observed corresponding to the Stark mixing of the 2s and 2p states. This was very roughly approximated by the large field treatment in appendix A. The number of quenched atoms inferred from this was compared to the arbitrary working scale assigned to the electrode quench curve. (See figure fifteen.) This comparison will be treated in detail when it is made.
The discussion in chapter III established that SCT could be tested if a beam of radiating atoms with a variable metastable component could be produced. It is a simple matter to produce a hydrogen beam containing some metastable atoms, but the discussion of the charge exchange cell indicates some of the difficulties involved in knowing the actual fraction obtained. These problems pale before those involved in producing a radiating beam of known composition.

The problem is threefold. The mixing electric field must be confined in space to a length short compared to a decay length for the p state; for a beam energy of 58 keV, this is 0.53 cm. On the other hand, the beam aperture must be on the order of 0.1 cm in order to have reasonable counting rates. Secondly, the field must be sufficiently homogeneous to give a "textbook" appearance: not only must the field variations be smooth, but they must be small enough so that the beam average bears some relationship to the expected behavior in a completely
uniform field. Finally, the observed $L_\alpha$ count as a function of applied voltage must be related to a fractional quench of the incident metastables. Figure fourteen shows the electrode structure used and figure fifteen shows the observed count rate as a function of voltage. Empirically, the conditions have been met, but the calibration of a scale for the rise and understanding of the observed oscillations remain.

The problems of immediate concern are to establish that such mixing meets the requirements of SCT and a calibration. The problem of variations in the mixing fraction from atom to atom is treated in appendix C. That the field was confined to the region limited by that ground planes was verified by looking at the signal roll off as the detector moved across the electrode structure. The observed width matched that predicted by geometry, as is seen in figure sixteen.

The calibration was accomplished, such as it was, by comparing the signal at the first maximum of the quench curve to that obtained by looking at radiation in the presence of a constant electric field. The steps involved are as follows: The number of $2p$ atoms created in the
Figure fourteen: Electrodes. Neither view is drawn to scale. (a) The upstream ground plane ($G_1$) was .032" copper and supports the rest. The insulation (I) was .00075" mylar, while the electrode parts (E) were .010" brass shim stock. The downstream ground plane ($G_2$) was .002" shim stock. The electrode separation was .045". (b) The .041" apertures through the ground planes were aligned on a wire drill and centered in the nominal 0.10" slit in the insulation. The entire structure was held fixed by two 2-56 screws which pulled the ground planes together. The total structure is 0.5" wide.
Figure fifteen: Quantum oscillations. When a potential difference of $2E$ was applied to the electrodes the photon count gave a measure of the number of metastable atoms driven into the $2p_{\frac{1}{2}}$ state. As the field increases the net number of radiating atoms produced passes through several maxima and minima.
Net Rate (Cts/nA-sec)

Zero Field Rate

Applied Voltage (V)

FIG 15
Figure sixteen: Signal roll off. The points show the fraction of the expected signal observed as the detector opening was obscured by the electrode structure. The smooth curve is the predicted behavior from appendix B if no radiation is emitted for $x < 0$. 
FIG 16

Fraction Transmitted vs. Window Position (in)
electrode region is essentially proportional to the count-rate immediately after leaving that field divided by the decay rate. However, a number of corrections must be made. Some of the 2p atoms will decay in the electrode region and not be included in the field free counts. If we assume that the metastables are quenched at a constant rate during the transit, then this correction is 0.9. Other reasonable estimates do not differ significantly. If \( p \) is the fraction of the metastable atoms observed to be quenched, and \( p_Q \) is the actual fraction, then

\[
p = 0.9p_Q \tag{1}
\]

and

\[
\sigma_2 \approx \sigma_D \{1 - (p/0.9)\}. \tag{2}
\]

Here \( \sigma_D \) is the fraction of the beam entering the electrode region in the metastable state, i.e. 78% of that at the end of the charge exchange cell.

The problem is to calibrate \( p \). The analysis in appendix A suggests that for small constant fields the radiation rate determined some distance into a uniform transverse electric field could be used to determine \( \sigma_D \), but in practice the extrapolation extended too far back to obtain consistent results. Going to the other extreme, large fields produce mixing which gives a single
decay rate right up to the origin if the description can be pushed that far. However, for large fields significant amounts of the $2p_{3/2}$ level are mixed in and the analysis is marginal at best. None the less, it was observed that the initializing count rate divided by the fitted decay constant was independent of the voltage applied to the plates, even though the measured rate continued increasing beyond the $\gamma$ value predicted by the theory. The low field values also extrapolated to this general value. From this it is concluded that the total obtained for large fields may be used to normalize $p$, and further, that the appropriate polarization is $-0.31$ as has been measured for low fields.\textsuperscript{61}

One more correction must be made before the observed quench percentage can be related to $\sigma_2$. Tucked away along the axis in figure fifteen is a horizontal line which represents the quench signal level due to the physical presence of the electrodes in the beam path. This light was emitted by atoms which had suffered collisions and involved a state preparation independent of that used to test SCT. If $f$ is the fraction of $\sigma_D$ which is struck, then $\sigma_D(1 - f)$ is the effective incident...
beam and only radiation from those atoms was counted. The fraction of these converted is \( (p - f) \) so that
\[
\sigma_2 = \sigma_D(1 - f)(.9 + f - p)/.9. \tag{3}
\]

The maximum peak on the curve in figure fifteen corresponds to \( p = .64 \pm .06 \). Further, \( f = .006 \), so that collisional effects were negligible. As will be shown in the discussion of the results, the observed decay rate in SCT should be written as
\[
\gamma = \gamma_0 + \beta p \tag{4}
\]
where
\[
\gamma_0 = A(1 - \sigma_D) \tag{5}
\]
and
\[
\beta = A\sigma_D/.9; \tag{6}
\]
A is the Einstein coefficient for spontaneous emission. For \( \sigma_D = .78 \times .1115 = .087 \) and \( A = 6.25 \times 10^8 \text{ sec}^{-1} \),
\[
\beta = (6.03 \pm 1.46) \times 10^7 \text{ sec}^{-1},
\]
where the 10% error associated with the normalization of \( p \) has been added to the 20% uncertainty inherent in the determination of \( \sigma_D \). The least value of \( \beta \) imaginable is \( 1.6 \times 10^7 \), which is obtained by taking \( \sigma_D = .025 \) and using the lowest conversion fraction obtained.
Appendix C establishes that enough is known about the composition of the radiating beam to test the theory. However, it would be interesting at least to be able to predict something similar to the oscillations shown in figure fifteen. To do this a 20:1 mock up of the electrodes was tested in an electrolytic tank. From the considerations elaborated in appendix D, the distribution of potential is described by

\[ V(x, z) = E_0 m_0 \sinh(z/l) e^{-l_2u(x/m_0)^2} \quad (7) \]

where

\[ u(z) = e^{2|z/h|}, \quad (8) \]

\( x \) is measured along the beam axis, and \( z \) increases from the center to the positive electrode. The values of the parameters for the mock up are given in the appendix.

If the fields appropriate to this are used in the Schrödinger equation and the radiation emitted is averaged over the beam cross section the result will be a theoretical prediction for the wiggles in figure fifteen. This calculation is described in appendix E, and the result is shown in figure seventeen. The data points are taken from figure fifteen and the horizontal scale
is modified so that the peaks coincide. No hyperfine structure effects are included.

The fit is not earthshaking, but it is worth showing and discussing briefly. It is tempting to suggest that inclusion of hfs would improve the fit, but some thought will convince the reader that multiplying the number of component oscillations by four is not going to narrow the structure. What is more likely the case is that the inhomogeneities as reflected in the beam profile average are not so great as the model predicts. Unfortunately, the cost of producing this curve prohibited the variation of any of the parameters.
Figure seventeen: Predicted quantum oscillations. The points are data values taken from figure fifteen and shifted to match the first maximum. The curve is a beam average of the transitions predicted by integrating the Schroedinger equation for a series of applied fields. The vertical scale is unaltered. The beam was assumed to fill 85% of the aperture.
FIG 17

Applied Voltage

Relative Intensity
QED and SCT make specific predictions about the decay of the $2p_{1\over2}$ state of hydrogen if we are careful in how we interpret the mathematics. In QED we consider an ensemble of atoms whose wave functions are written as

$$\Psi(\mathbf{r},t) = \sum c_i(t) \phi_i(\mathbf{r}).$$  \hspace{1cm} (1)$$

The coefficients are interpreted to mean that if a measurement is made at time $t$, $|c_i(t)|^2$ atoms will be found in the state $\phi_i(\mathbf{r})$. For the hydrogen atoms in this experiment only three states were involved—neglecting hfs:

$$\Psi(\mathbf{r},t) = c_1(t) \phi_{1s}(\mathbf{r}) + c_2(t)e^{-i(\omega_o+\omega)t}\phi_{2s}(\mathbf{r})$$

$$+ c_3(t)e^{-i\omega t}\phi_{2p}(\mathbf{r})$$  \hspace{1cm} (2)$$

where only the $j = \frac{1}{2}$ substates were occupied. QED predicted that an ensemble of $N_o$ atoms would radiate energy at a rate

$$R(t) = N_o \gamma \omega |c_3(0)|^2 e^{-\gamma t}$$  \hspace{1cm} (3)$$

where $\gamma$ is given by the Einstein coefficient connecting the two states.
SCT interprets the ensemble differently. The magnitude of the coefficients $|c_i(t)|^2$ may be measured from the ensemble, but the individual atoms really exist in the mixed state of equation 1. This is no longer a mathematical figment which serves only to predict the results of measurements. For the same set of hydrogen atoms as before we expect a radiation rate

$$R(t) = N_0 (\eta \gamma) \omega |c_3(0)/c_1(0)|^2 e^{-\eta \gamma t}$$

(4)

where

$$\eta = 1 - |c_2(0)|^2.$$  

(5)

Since $|c_2(0)|^2$ can be made to vary over several percent such changes should be observable in the measurement of the rate constant in equation 4.

For the atomic beam described in chapter IV $|c_2(0)|^2$ may vary in principle from zero to 0.087, and with the electrodes described in chapter V was varied from 0.031 to 0.073. However, this is not the best way to look at the experiment. Instead, let us assume some fixed value for the metastable fraction and then look at the percentage $p$ of these quenched.
The observed radiation rate constant will be given by
\[ \Gamma = \Gamma_0 + (\beta/v)p \]  \hspace{1cm} (6)
where the value of \( \beta \) depends on the initial richness of the beam and the value of \( A \) as discussed in chapters IV and V. The velocity \( v \) converts the times to beam positions, and \( \Gamma = \gamma/v \).

For QED we expect \( \beta = 0 \). The radiation was measured along the beam at positions given in inches so the natural experimental units for \( \Gamma \) are in \(^{-1}\). From the previous discussion and a measured beam energy of 58.1 keV we expect \( \beta/v = 0.46 \) in \(^{-1}\) if the prediction of SCT is true.

The procedure seems simple enough: Select a sequence of electrode voltages which give a distribution of \( p \) values and then fit the measured rates to equation 6. The problem of measuring \( \Gamma \) to an appropriate precision is discussed in appendix F. Figure eighteen shows the radiation rate as a function of \( x \) over a much greater interval than we used, while table I shows the measured values of \( \Gamma \) as a function of \( p \). The uncertainty assigned to a given measurement of \( \Gamma \) is taken from all the residuals in the forty-two curve fits. The value \( \sigma_\Gamma = .025 \) in \(^{-1}\) corresponds to the expected error from counting sources.
Figure eighteen: Exponential decay. The strictly exponential decay of the \( L_\alpha \) radiation was observed over more than five lifetimes, and during the experiment measurements could be made another lifetime closer to the electrodes. The error associated with the higher count rates is one third that shown on the lower values.
FIG 18
The uncertainties in the values of \( p \) are inferred from the initial counts in the \( \Gamma \) determinations with the counting uncertainties removed. They reflect to some extent a drift in the electrode efficiency since some degradation definitely occurs. As we shall see this uncertainty is immaterial.

The bottom row of table I shows results of a preliminary fit of the data to equation 6. The values of \( p \) were treated as exact to obtain preliminary values of \( \beta/v \) and \( \Gamma_0 \) for use in a more careful regression. However, the resulting values of \( \beta/v \) are so small that such corrections are unnecessary. The uncertainty in \( \Gamma \) completely determines the weight to be assigned each point.

Finally, the weighted mean value of \( \beta/v \) is

\[ \beta/v = -0.021 \pm 0.021 \text{ in}^{-1}, \]

compared to

\[ (\beta/v)_{\text{SCT}} = 0.46 \pm 0.11, \]

and

\[ (\beta/v)_{\text{QED}} = 0. \]

This rules out SCT if the systematic errors are fully
understood. These will be discussed in the following chapter. All experimental uncertainties are quoted as one standard deviation.
Table I: Rate values. The $2p_{1/2}$ decay rate in reciprocal inches is given for each set of quench percentages. The uncertainty in $\Gamma$ is 0.025 and this determines the uncertainty quoted for $\beta$. The uncertainties shown for the $p$ values come from the count rates observed in measuring $\Gamma$. 
**TABLE I**

<table>
<thead>
<tr>
<th>p</th>
<th>Observed Decay Rates T/v (in.⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.640±0.008</td>
<td>4.791 4.746 4.812 4.774 4.754 4.765 4.777 4.743</td>
</tr>
<tr>
<td>0.535±0.009</td>
<td>4.768 4.738 4.748 4.715 4.802 4.797 4.764 4.766</td>
</tr>
<tr>
<td>0.424±0.015</td>
<td>4.833 4.753 4.777 4.791 -- -- -- --</td>
</tr>
<tr>
<td>0.350±0.007</td>
<td>-- -- -- -- 4.752 4.784 4.777 4.741</td>
</tr>
<tr>
<td>0.340±0.014</td>
<td>4.809 4.783 -- 4.752 -- -- -- --</td>
</tr>
<tr>
<td>0.244±0.012</td>
<td>4.755 -- 4.759 4.754 4.790 4.735 4.819 4.744</td>
</tr>
<tr>
<td>0.147±0.008</td>
<td>4.829 4.753 4.774 4.729 4.776 4.767 4.832 4.819</td>
</tr>
<tr>
<td>β/v (in.⁻¹)</td>
<td>-0.040 -0.039 +0.049 +0.040 -0.018 +0.048 -0.125 -0.088</td>
</tr>
<tr>
<td></td>
<td>±0.058 ±0.064 ±0.049 ±0.062 ±0.059 ±0.059 ±0.059 ±0.059</td>
</tr>
</tbody>
</table>
As in any experiment the interpretation of the results in the preceding chapter depends upon an understanding of their limitations. These may be caused by instrumental problems or be inherent in the processes involved. An example of the latter is to be found in the uncertainty quoted on the measurement of \( \Gamma \). If we had been "successful" in measuring ten thousand counts to a precision of better than one hundred counts we would have been forced to search for a systematic error source in the counting process. Similarly, the electrode voltage could be reproduced only to within about a third of a volt, and this as well as other factors appear in the uncertainty assigned to \( p \).

These problems can be broken down into three categories. The first involves problems in calibrating the apparatus which were discussed in detail in the descriptions of the electrodes and the charge exchange process. They result in what is best described as
a 25% uncertainty in the expected results if SCT is correct. The second category involves purely random errors. Included in this are parameter changes which occur in times comparable to a single count. Examples would include the counting distribution just mentioned as well as fluctuations in the beam velocity and the exchange cell pressure. The final category is long term drifts and effects dependent upon the electrode voltage. These become intermingled in that the lower voltage counts take longer.

Let us begin with the second category. The value of $\Gamma$ is primarily determined by the velocity of the beam which in turn depends upon a combination of the accelerator voltage, the analyzing magnet current, and the quadrupole lens fields. The magnet power supply is regulated to 0.01%, so we may consider only voltage uncertainties and the angular acceptance of the aperture system. If the beam were to sweep over the entire width of the last aperture the velocity would change by 0.2%. Such detuning was not observed, but it serves to give an idea of the spread of velocities from which a signal was obtained. The fine tuning adjustments on the high
voltage suggest that the central velocity fluctuations were less than 0.1%. The quickest measure of these is provided by $\sigma_T$. The variation over a sequence of runs was negligible while the day to day changes were mainly responsible for the larger variance associated with the average of all the $T$ measurements: 0.029 compared to 0.025 out of 4.773.

The other possible short term fluctuation would be in the richness of the beam. This would be caused by variations in the exchange cell pressure. The observed fluctuations in $P_X$ were always less than 0.1 μ and were adjusted away as soon as they were spotted. The Baratron would show a zero drift over a data taking sequence of less than 0.2 μ and both of these seem to lie well within the limits of the plateau in the variation of beam richness with $P_X$ shown in figure four. The random error in the measuring device is rated at 0.05%.

Random errors were indeed pushed down to where the necessary precision could be obtained in a finite time. It is the effects in the third category which determine the significance of the result.
Systematic errors come in three varieties: time dependent, electrode dependent, and independent effects associated with the lifetime measurements. Let us take care of the last first. The decay rate observed is affected by ambient fields; the earth's magnetic field comes to mind as the principal source, but the wiring in the chamber could also be a source of stray fields. The magnetic field strength in the experimental chamber with the magnet on was ~0.5 Gauss which corresponds to a motional electric field

$$E_{\text{mot}} = 1.5 \text{ V/cm} = 0.003E_0,$$

where $E_0 = 477.3 \text{ V/cm}$ is the characteristic mixing field for the two upper levels. For small perturbations the decay rate is decreased by an amount proportional to $(E/E_0)^2$ so that the rate measured is true to the limits shown.

Another possible measurement problem is presented by the finite segment of the beam observed. It is shown in appendix B that an exponential functional form is unaffected by a finite window. The efficiency does change very slightly with $\Gamma$ but the rate measurement is independent of the absolute magnitudes involved.
The last item in this set is cascading. The detector response shown in figure eight suggests that only \( L_\alpha \) is to be observed. The linearity of the graph in figure eighteen indicates that the problem does not exist over the five lifetimes shown.

Time dependent uncertainties are a bit more difficult to eliminate. Items which could affect the measurement of \( \beta \) are drifts in the light detection efficiency, in the secondary electron emission characteristics of the neutral beam detector, the beam velocity, the richness of the beam, and the background rate. Two things were done to minimize the effects of these: they were eliminated so far as possible, and the order of data taking was randomized with respect to \( p \); just in case. Some discussion is in order, however.

For a given electrode setting variations in the counting rates are shown in table I of chapter VI as a variation in the value of \( p \) listed there. There was indeed some drift over the course of approximately forty hours of data taking. The effect was a decrease in the count rate and could be caused by an electrode degradation, a counting efficiency decrease---either
the window or the detector, or an increase in the number of secondary electrons per neutral atom. This last can be ruled out since the secondary electron emission was observed to decrease with contamination. The probable cause is primarily electrode contamination with a small amount of additional loss in the LiF window. The effect was about five percent over the entire sequence of measurements and was only apparent in the final analysis.

A stickier problem was changes in the background counting rate. Radiation induced by the electrodes when they were grounded had to be subtracted out since it was not produced in a way consistent with SCT. This would increase as insulation in the electrodes or in the drift region became charged. The latter had the effect of defocusing the beam and was greatly decreased by cleaning the sweep plates and installing a metal shield over the exposed insulation on the third aperture. The electrode problem was never completely solved. When the mylar was moved far enough from the beam hole to be untouchable, the metal parts would short. Ultimately, the drift was reduced to less than ten percent of the net background count and the background count itself was
pushed down to a maximum of ten percent of the lowest count rate. This effect was not so much systematic as random; when the background became excessive the measured decay rates became untrustworthy.

The background effect was partly electrode voltage dependent, and hence, a more crucial problem. Two other such difficulties were conceived and must be accounted for. The first and simpler of these involved the ejection of electrons from the electrode structure by beam atoms. These would have large cross sections for the excitation of the background gas and would also have an electrode voltage dependent trajectory. This problem did not exist in practice. The current to ground from the electrode structure was less than 0.1% of the beam current and the observed background for a prequenched beam was independent of position along the beam and the electrode voltage.

The final effect was the most important from the standpoint of testing SCT. This was the leaking of the electric field beyond the region defined by the electrodes. Since any perturbation mixing the $2p_{\frac{1}{2}}$ and $2s_{\frac{1}{2}}$ states increases the apparent lifetime of the $2p_{\frac{1}{2}}$ state,
the results could favor QED when in fact SCT was true. For simplicity we consider the constant field necessary to change the decay rate by one percent. An analysis similar to that in appendix A but with more general initial conditions gives as the effective rate constant for small perturbations

\[ \gamma' = \gamma \{ 1 - \left( \frac{E}{E_0} \right)^2 \}. \]

The desired one percent variation requires a field strength of about 50 V/cm from shielded objects at a maximum of 75 V and a minimum distance of about 0.5 cm away.

It seems reasonable to expect a small negative value for \( \beta \) if QED is true, but we cannot attribute the failure of SCT to any of the effects discussed here.
CONCLUSIONS

As was pointed out in the introduction, two conclusions may be drawn from the experiments described herein. As a test of the semiclassical radiation theory proposed by E. T. Jaynes the conclusion is rather obvious. The theory predicts a specific dependence of the excited state lifetime on the metastable component and that effect is not observed. Since the theory also fails to predict the correct intensity ratios in Raman scattering and the relative polarizations in cascade processes it must be rejected as formulated. However, we shall return to this in a moment.

Another interpretation of this work, although not the one which initiated it, is that we have investigated the dependence of the exponential decay law on the method of preparation of the radiating state. Within the narrow range of preparations used we observed no first order effects.

A modification of the semiclassical theory has been proposed by R. K. Nesbet which has the effect of saving the phenomenon in the case of Raman scattering and also, apparently, in this instance. It is not appropriate to
to go into detail here but by second quantizing the coefficients of the stationary states in the expansion of the wave function he succeeds in eliminating the final state dependence of the radiation rate—see equation 27 of chapter III. Since this does not invalidate the essential feature of Jaynes' program, namely to provide an alternative to the QED treatment of the radiation field, we must reserve judgment on the feasibility of semiclassical theories in general until all the implications of this modification have been investigated.
APPENDIX A
METASTABLE QUENCH IN A UNIFORM FIELD

We consider here the time evolution of a mixed state

\[ \Psi(\mathbf{r}, t) = a(t) \phi_s(\mathbf{r}) + b(t) e^{i \omega t} \phi_p(\mathbf{r}) \]  

which we will couple heuristically to a ground state through the decay of the \( p \)-component. (These are \( j = \frac{1}{2} \) states.) The mixing will be caused by an electric field \( E = 2E \).

From the Schroedinger equation we have

\[ i \dot{a}(t) = E e^{i \omega t} s |z| p ^{b(t)} \]  

and

\[ i \dot{b}(t) = E e^{-i \omega t} p |z| s ^{a(t)} \]  

We then modify equation 3 to give the correct behavior in the absence of the perturbation.

\[ i \dot{b}(t) = E e^{-i \omega t} p |z| s ^{a(t)} - \frac{1}{2} i \gamma b(t) \]  

The most convenient initial conditions are

\[ a(0) = 1, \ b(0) = 0. \]  

The time evolution of the mixed state is given by

\[ N(t) = |a(t)|^2 + |b(t)|^2, \]  

so we begin the process of solving equations 2 and 3'.
The two first order equations combine to give
\[ \dot{a} - i(\omega + \frac{1}{2}i\gamma)\dot{a} + M^2a = 0 \] (6)
where
\[ M = eE|s|z|p> = -\sqrt{3}e\alpha_o E \]
and \( \alpha_o \) is the Bohr radius. This we will solve in the approximation \( \gamma \ll \omega \). The exact solution is
\[ a(t) = a_+ e^{r_+ t} + a_- e^{r_- t} \]
with
\[ r_\pm = \frac{1}{2}i(\omega + \frac{1}{2}i\gamma) \pm \frac{1}{2}i((\omega + \frac{1}{2}i\gamma)^2 + 4M^2)^{\frac{1}{2}} \]
\[ \approx \frac{1}{2}i\omega(1 \pm Q) - \frac{1}{2}\beta_\pm \gamma. \]
The parameters \( Q \) and \( \beta_\pm \) are defined by
\[ Q = \{1 + (4M^2/\omega^2)\}^{\frac{1}{2}}, \] (7a)
and
\[ \beta_\pm = (Q \pm 1)/2Q. \] (7b)
From the initial conditions we obtain
\[ a_\pm = \beta_\pm \]
so that
\[ a(t) = e^{\frac{1}{2}i\omega t}\{\beta_- e^{\frac{1}{2}iQ\omega t} e^{-\frac{1}{2}\beta_+ \gamma t} + \beta_+ e^{\frac{1}{2}iQ\omega t} e^{-\frac{1}{2}\beta_- \gamma t}\}, \] (8)
and
\[ |a(t)|^2 = \beta_-^2 e^{-\beta_+ \gamma t} + \beta_+^2 e^{-\beta_- \gamma t} + 2\beta_+ \beta_- e^{-\frac{1}{2}\gamma t}\cos Q\omega t. \] (9)
If equation 8 is differentiated and substituted into equation 2 we have

\[ b(t) = -\left(\frac{\omega}{2M}\right)e^{\frac{-i}{2}\omega t}\{\beta_-(Q+1)e^{\frac{i}{2}\omega t}e^{-\frac{i}{2}q_+\gamma t} - \beta_+(Q-1)e^{\frac{i}{2}\omega t}e^{-\frac{i}{2}q_-\gamma t}\}, \]

whence,

\[ |b(t)|^2 = \beta_+\beta_-\{e^{-\beta_+\gamma t} + e^{-\beta_-\gamma t} - 2e^{-\frac{1}{2}\gamma t}\cos Q\omega t\}. \]  

The excited state amplitude is

\[ N(t) = \beta_-e^{-\beta_+\gamma t} + \beta_+e^{-\beta_-\gamma t}, \]

where we have used the fact that

\[ \beta_+ + \beta_- = 1. \]

We need the behavior of this in two limits:

\[ E \ll E_0 = \omega/\sqrt{3}a_0 \]  

and

\[ E \gg E_0, \]

although the condition in equation 13 is restricted to

\[ E/E_0 < 8 \]

for the two state analysis to be applicable to the first excited state of hydrogen.\(^6\)

For the first case \( Q = 1, \beta_+ = 1 \) and \( \beta_- \approx (E/E_0)^2 \)
so that

\[ N(t) \approx e^{-\left(E/E_0\right)^2\gamma t}. \]
This behavior was observed in sweeping protons from the beam and was verified earlier by Sellin.\textsuperscript{66}

In the other extreme we have $Q \gg 1$, and hence, $\beta_- \sim \frac{1}{2} \sim \beta_+$, so that

$$N(t) = e^{-\frac{1}{2}\gamma t}.$$  

The observed radiation rate will be half that for an unperturbed $2p$ state.
APPENDIX B
WINDOW FUNCTION

Consider the counts observed at a position $x$ along the beam. The CEM observes a finite segment of beam more or less centered at $x$ and detects light emitted from an infinitesimal portion at $x + \xi$ with efficiency $f(x, \xi)d\xi$. The count rate is then given by

$$r(x) = n\gamma\int_{-\infty}^{\infty} e^{-\gamma(x+\xi)/v} f(x, \xi)d\xi$$

(1)

where $\gamma$ is the decay constant of the atoms, $v$ is the beam velocity and $n$ is the excited atom density at $x = 0$. If the beam is properly oriented, the moments of the efficiency function are independent of $x$ and

$$r(x) = C(\gamma/v)e^{-\gamma x/v}$$

(2)

where

$$C(\gamma/v) = n\int_{-\infty}^{\infty} e^{-\xi(\gamma/v)} f(x, \xi)d\xi.$$  

(3)

The window depends only on the rate, in this case $\gamma/v$.

The total window width could be measured by obscuring the beam with the electrode structure. The effective width of .07" agrees with the geometry and the 0.024" measured
slit size. If we use a triangle centered at $x$ for $f(x,\xi)$ we have

$$f(x,\xi) = \begin{cases} 
\frac{(a + \xi)}{a^2} & -a < \xi < 0 \\
\frac{(a - \xi)}{a^2} & 0 < \xi < a \\
0 & |\xi| > a
\end{cases}$$

(4)

where $a = .035''$ in this experiment. The normalization is chosen so that a fixed rate would be seen unaltered. The resulting window is

$$C(\gamma/v) = 2(\cosh(\gamma a/v) - 1)/(\gamma a/v)^2.$$  

(5)

For $\gamma a/v \ll 1$ we have

$$C(\gamma/v) \approx 1 + (1/12)(\gamma a/v)^2,$$  

(6)

and in this instance a 100% variation in $\gamma/v$ will cause a 1% change in $C(\gamma/v)$. 


APPENDIX C
QUENCH DISTRIBUTIONS

The SCT predicts that the 2p-state decay rate for an individual atom is given by
\[ \gamma = A(1 - \sigma) \]  
where \( A \) is the Einstein coefficient and \( \sigma \) is the residual 2s occupation. We do not look at individual atoms, but rather measure the average amount of 2p conversion for the entire beam.

Let \( s \) be the residual 2s occupation for a differential cross section of the beam. We then observe for an electrode potential \( E \)
\[ \sigma(E) = \int_0^1 f(E,s)sds, \]  
where the distribution \( f \) of \( s \) values is clearly included in \([0,1]\), and further
\[ \int_0^1 f(E,s)ds = 1 \]  
If the atomic rate is given by
\[ \gamma = A(1 - s), \]  
do we observe a beam average given by equation 1?
The observed count rate is given by

\[ r(x) = r_0(E) \int_0^1 e^{-A(1-s)x/v} f(E,s) ds \]
\[ = r_0(E) e^{-Ax/v} \int_0^1 e^{Asx/v} f(E,s) ds. \] (5)

Over the range of positions used and metastable occupations available we have

\[ Asx/v \ll 1 \]

whence

\[ \int_0^1 e^{Asx/v} f(E,s) ds = \]

\[ \int_0^1 (1 + Asx/v + \frac{1}{2}(Asx/v)^2 + \ldots) f(E,s) ds \]
\[ \approx 1 + (A\sigma(E)x/v + \frac{1}{2}(A\sigma(E)x/v)^2 + \frac{1}{2}(Ax/v)^2 (\frac{s^2}{\sigma} - \sigma^2)) \]
\[ \approx e^{A\sigma(E)x/v} \]

with an error determined by the width of the distribution. If this is 10% then the error is 1%. In practice this seems a likely upper limit--see appendix E.

We see then that the observations made in terms of the beam average reflect the variations in atomic behavior to better than 1%.
APPENDIX D
ELECTRODE FIELD MEASUREMENTS

The electrode structure shown in figure fourteen does not permit an analytic determination of the voltage distribution. As a result electrolytic tank measurements were used to determine whether the fields were sufficiently confined to meet the criteria of SCT.

For purposes of description we establish a coordinate system at the center of the field region: \( \hat{x} \) shall point in the beam direction, \( \hat{z} \) from the negative to the positive electrode, and \( \hat{y} \) along the electrodes in a righthanded way. The variation of \( V(x,y,z) \) in each direction was investigated systematically. Briefly, \( E_x \) and \( E_z \) are comparable while \( E_y \) may be ignored. Let us now consider \( V(x,z) \).

Initially it was thought that the \( x \) dependence would satisfy some sort of multipole law

\[
V(x,y) \propto (1 + a^2 x^2)^{-\nu/2}
\]

but a sequence of fits for larger and larger \( \nu \) improved too slowly, and the numbers involved soon exceeded the capacity of the calculator being used. A graph of some
representative multipole fits showed that, for a given peak value and width, the wings fell off much faster than the fitting function. However, since the fit clearly improved for increasing \( v \) the limit of

\[
f(v, x) = (1 + a^2 x^2)^{-v/2}
\]

for \( v \to \infty \) was determined subject to the restriction that the width be held fixed. The limit function turns out to be a Gaussian. Figure nineteen shows the \( x \) dependence of \( V(x, z) \) for three values of \( \xi = z/r; r \) is the aperture radius. It can be seen that the parameters of the Gaussian both depend upon \( \xi \). The limit function behaves somewhat better than the wings. For \( \xi < .5 \) which includes 75% of the beam cross section the disagreement becomes insignificant.

From the values obtained in the fits to the form

\[
V(x, z) = A(z) \exp \left( -\frac{1}{2} \left( \frac{x}{m(z)} \right)^2 \right)
\]

the two remaining functions could be obtained. Figures twenty and twenty-one show that

\[
A(z) = A_0 \sinh \left( \frac{z}{\lambda} \right)
\]  \( \text{(3a)} \)

and

\[
m(z) = m_0 e^{-|z/h|}.
\]  \( \text{(3b)} \)
Figure nineteen: Gaussian fits. The potential induced by the electrode configuration shown in figure fourteen was approximated by a Gaussian function along the beam axis. This fit is shown for three off axis positions.
\[ \xi = 1 \]
\[ (x \times \frac{1}{2}) \]

\[ \xi = 0.7 \]

\[ \xi = 0.4 \]
Figure twenty: Field inhomogeneity I. The amplitude of the Gaussian potential function had a hyperbolic sine dependence on the distance off axis. No points could be established for \( \xi < .4 \), but an independent direct measurement of \( V \) versus \( \xi \) gave the same relationship.
FIG 20
Figure twenty-one: Field inhomogeneity II. The width of the Gaussian decreased exponentially with the distance off axis. This is partly due to the fact that for larger amplitudes the wings become less important. The values on the logarithmic vertical axis are in millimeters on the mock up.
Hence, the working description of the electrode field is

$$V(x,z) = E_0 m_0 \sinh(z/\lambda) e^{-\frac{1}{2}u(x/m_0)^2}$$

where

$$u = e^{2|z/h|}.$$  \hspace{1cm} (4)

The parameters involved are related as follows:

$$m_0/\lambda = 2.34, \quad m_0/h = 0.494, \quad r/\lambda = 3.615, \quad r/h = 0.730,$$

and $$m_0 = 0.337 \text{ mm}$$. This last value is for the structure seen by the atoms.
APPENDIX E

BEAM AVERAGING

The fields derived in appendix D must be used to obtain a solution to the Schroedinger equation for the atomic system as it passes through the aperture. If we write the general atomic state as

\[ \psi(r, t) = c_1(t)\langle \hat{r} | 2, 0, \frac{1}{2}, \frac{1}{2} \rangle + c_2(t)\langle \hat{r} | 2, 0, \frac{1}{2}, -\frac{1}{2} \rangle + c_3(t)\langle \hat{r} | 2, 1, \frac{1}{2}, \frac{1}{2} \rangle + c_4(t)\langle \hat{r} | 2, 1, \frac{1}{2}, -\frac{1}{2} \rangle, \]

where the notation is \(|n, \ell, j, m_\ell\rangle\), the coefficients must satisfy the matrix equation

\[
\begin{bmatrix}
\dot{c}_1 \\
\dot{c}_2 \\
\dot{c}_3 \\
\dot{c}_4
\end{bmatrix} = \begin{bmatrix}
0 & 0 & -X & -Z \\
0 & 0 & X & Z \\
-Z & X & -\omega & 0 \\
-X & Z & 0 & -\omega
\end{bmatrix}
\begin{bmatrix}
c_1 \\
c_2 \\
c_3 \\
c_4
\end{bmatrix}
\]

where

\[ X = \sqrt{3} ea E_x, \]

\[ Z = \sqrt{3} ea E_z, \]

\[ \omega = \text{the Lamb shift, and } a \text{ is the Bohr radius.} \]

Calling the coefficient matrix \(M\) and setting
\[ C = A + iB \]  
(3)
gives two coupled first order equations:
\[ \dot{A} = MB \]  
(4a)
and
\[ \dot{B} = -MA. \]  
(4b)
To the approximations that no decays take place during the pulse and that hfs is an intrusion, the radiation observed will be given by
\[ R \propto |c_3(\infty)|^2 + |c_4(\infty)|^2. \]  
(5)
The solution needs one further parametrization.
Let the maximum width of the fields be \( m_0 \) and write
\[ m_0 = vT \]  
(6)
where \( v \) is the particle velocity. In terms of \( \theta = t/T \) the fields are
\[ E_x = E_0 \theta \sinh(\xi/\lambda)e^{-\frac{1}{2}u\theta^2} \]  
(7)
and
\[ E_z = E_0 \left( \frac{m_0}{\hbar} \right)^2 \theta \sinh(\xi/\lambda) - \frac{m_0}{\lambda} \cosh(\xi/\lambda)e^{-\frac{1}{2}u\theta^2} \]  
(8)
where \( \lambda = \lambda/r \), and the other terms are defined in appendix D. This system of equations was solved numerically by a modified Euler method\(^6\) and the solutions
for each value of \( E_0 \) were averaged over the cross section by varying \( \xi \). Note the relativeness of infinity. The infinity appearing in equation 5 is the zero of the radiative process—in practice infinity was \( \Theta = 6 \).

The quench curve generated appears in figure seventeen. Figure twenty-two shows the component curves for three values of \( \xi \) while figure twenty-three shows the variation of the quench over the beam for fixed \( E_0 \).
Figure twenty-two: Radiation intensity components I. The quantum oscillations depend not only on the applied potential but on the position of the atom relative to the axis. This graph shows the oscillations for three different positions off axis. The failure of the swings to reach zero is apparently real and not an integration error.
FIG 22

Relative Intensity

$\xi = .6$

$\xi = .4$

$\xi = .2$

$E_0$
Figure twenty-three: Radiation intensity components II. For a given applied field the distribution of quench fractions is seen to be quite broad.
APPENDIX F

STATISTICAL OPTIMIZATION

We are faced with the problem of measuring one percent changes in the lifetime of the excited hydrogen atom. Clearly we want to do this as efficiently as possible, so some consideration must be made of the method of measuring $\gamma$.

The raw data are a conversion percentage, a position, photon counts with the electrodes charged and grounded, and the corresponding total number of secondary electrons collected. This last quantity has an uncertainty related to the meter response and drift which is negligible compared to the other uncertainties. This quantity is called the beam charge. A series of photon counts were made in which the beam charge was held fixed and the position reset each time. The data collected had a sample variance predicted by the Poisson distribution. Therefore the only statistical uncertainties are in the conversion and in the photon count. The latter is relevant to our consideration now.

We are to fit the relation

$$\ln r = -\Gamma x + I,$$

(1)
where
\[ \Gamma = \gamma/v, \]  
\[ r = N/e_2, \]  
(2a)  
(2b)

\( N \) is the photon count, \( e_2 \) is the beam charge, and \( x \) is the position of the detector. We ignore the background for the moment since it was less than 10% and as low as 0.5% of the net rate.

In order to ascribe a specific uncertainty to the value of \( \Gamma \) obtained we must assign weights to the values of \( r \). We minimize

\[ S = \sum_{i=1}^{n} (\ln(r_i + \Gamma x_i - I))^2 w_i \]  
(3)

where the subscript ordinates the data and

\[ w = \sigma_{\ln r}^{-2}. \]  
(4)

We have then

\[ \sigma_{\ln r}^2 = (\partial \ln r / \partial N)^2 \sigma_N^2 + (\partial \ln r / \partial e_2)^2 \sigma_{e_2}^2 = N^{-1}. \]  
(5)

Hence, the terms in equation 3 will have a fixed weight if we take our data with a fixed photon count.

This turns out to be more convenient mathematically than holding the time fixed. The value \( \Gamma \) obtained by
minimizing $S$ is

$$T = \frac{\langle x \ln r \rangle - \langle x \rangle \langle \ln r \rangle}{\langle x^2 \rangle - \langle x \rangle^2}$$

(6)

where

$$\langle q \rangle = n^{-1} \sum_{i=1}^{n} q_i .$$

By the same technique as in equation 5

$$\sigma_T^2 = \{nN(\langle x^2 \rangle - \langle x \rangle^2)\}^{-1}. \quad (7)$$

Having fixed $\sigma_T$, how long will it take to obtain the data necessary to determine $\Gamma$? Let $R$ be the number of photons counted per second at $x = 0$. We want to count $N$ photons at intervals $\Delta x$. For $n$ measurements we will require

$$T = \frac{N^{n-1}}{R} \sum_{k=0}^{\infty} e^{k\Gamma \Delta x}$$

or

$$T = \frac{(N/R)(1 + e^{\Gamma \Delta x})}{(1 + e^{\Gamma \Delta x})} . \quad (8)$$

In these terms equation 7 becomes

$$\sigma_T^2 = \frac{12}{(\Delta x)^2} n(n^2 - 1)N . \quad (9)$$

The problem is reduced to minimizing $T$ subject to the constraint provided by equation 9.

This was done numerically for the values of $\Gamma$ and $\Delta x$ appropriate to the experiment. The result was
that a series of three widely spaced long counts would be best. However, a more psychologically tenable six counts per \( \Gamma \) determination was adopted at the expense of about half an hour over the series of runs. These were made at intervals of 0.100" which corresponds to 0.76 nsec and for 10 000 counts. The precision on a given measurement of \( \Gamma \) was 0.5\%.

The uncertainty ascribed to the final data was the average \( s_{\ln r} \) as determined by

\[
s^2_{\ln r} = \frac{1}{n-1} \sum_{i=1}^{n} (\ln r_i + \gamma x_i - 1)^2.
\]

This turned out to match the predicted value when the background is included.
FOOTNOTES


10. This is part of a review in M. Planck, *Ann. Physik*, 1, 69 (1900).


17. W. Heisenberg, The Physical Principles of the Quantum Theory, trans. by C. Eckart and F. C. Hoyt (Dover, New York). These 1929 lectures at the University of Chicago provide an excellent insight into how quickly the quantum mechanics fell into place.


27. J. Schwinger, Phys. Rev. 73, 416 (1948).

29. M. Jammer, Ref. 1, Chap. 6, et seq.
35. M. Jammer, Ref. 1, p. 166ff.
46. The complete development is found in F. Rohrlich, *Classical Charged Particles* (Addison-Wesley, Reading, Mass., 1965). The appropriate primary references—usually to Rohrlich—are given there.


49. Ref. 23, p. 135.

50. Ref. 40, p. 1257.

51. Ref. 46, p. 68, 69.


56. Ibid., p. 100, 101.

57. A. Baer, private communication.


60. Ref. 54, p. 118.


66. Ibid., p. 1250.

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radiation theories