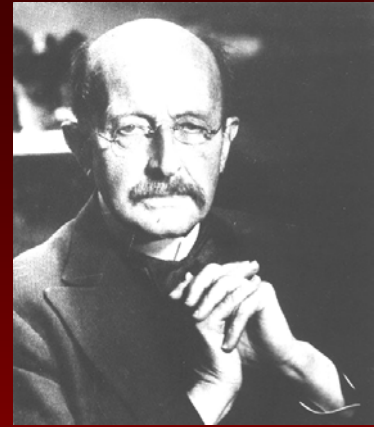


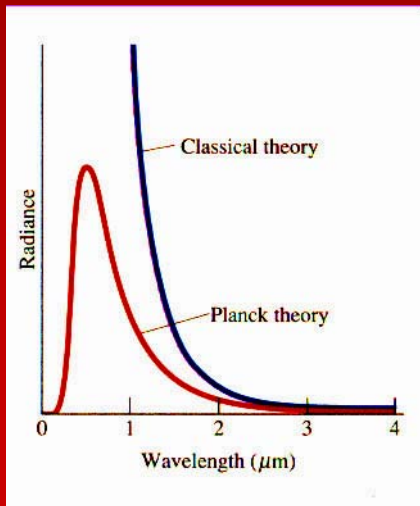
The GENESIS of QUANTUM MECHANICS

The quantum theory had its historical roots in the antithesis between Huyghens's and Newton's theories of light. However it was not yet even suspected that matter might be wavelike- or that light could be corpuscular.

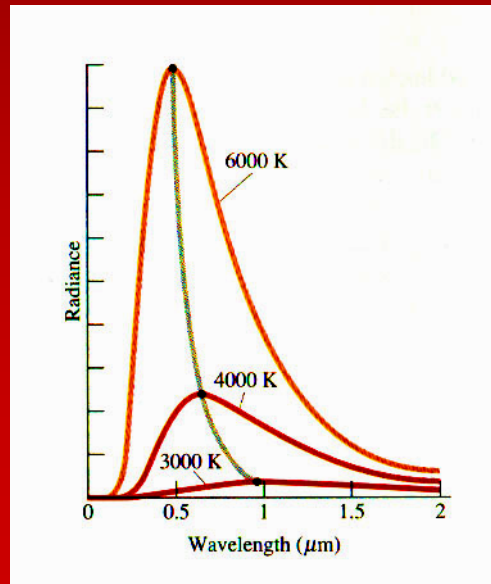
The path to the new physics started with attempts to understand radiant heat and thermal radiation. Maxwell had shown the "equipartition theorem"; in thermal equilibrium, all degrees of freedom had to equally share thermal energy, proportional to temperature **T**. But this led to the Rayleigh-Jeans "ultraviolet catastrophe"; that the infinity of short-wavelength EM modes would contain an infinite thermal energy, at any **T**. This made no physical sense.



Planck (1858-1947)



Thermal energy per unit frequency. Above-quantum vs classical.



Right: quantum for different **T**

In a crucial step, Planck argued that thermal energy should be shared according to a different law, based on the "quantisation" of energy- that it only comes in discrete amounts. This not only gave the correct distribution of thermal energy (quickly verified by Pringsheim), but also gave us a new constant of Nature- **Planck's constant:**

$$h = 6.63 \times 10^{-34} \text{ Joule secs}$$

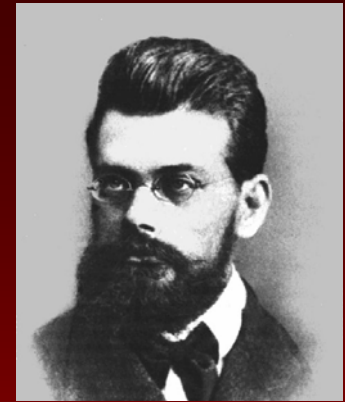
where Joules measure energy.

ATOMS & ELECTRONS

Dalton, in 1802, showed how to understand chemical combinations, if some elementary chemicals came in atomic units. Despite the evidence for atoms, and elements (classified by Mendeleev in the periodic table), their existence was still doubted in 1900. It was finally established by theoretical work of Maxwell, Boltzmann & Einstein- they showed how one could quantitatively understand gas dynamics, as well as the random “Brownian” motion of small particles in liquids, in terms of atomic motion and collisions. Atomic sizes were found to be $\sim 1 \text{ \AA} = 10^{-10} \text{ m}$. Electrons were discovered by JJ Thompson in 1895 in gas discharges- their mass is $9.11 \times 10^{-31} \text{ kg}$.



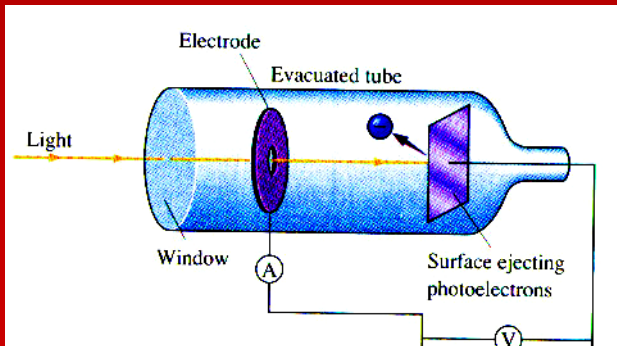
Dalton (1766-1844)



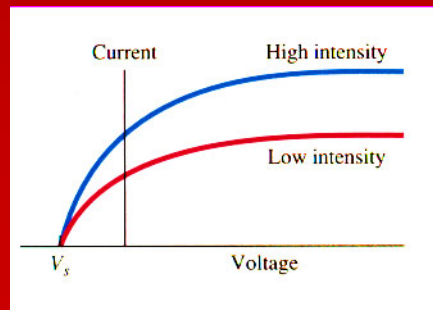
Boltzmann (1844-1906)

Einstein went much farther. He explained the “photoelectric effect” (the emission of electrons by a metal when light is shone on it), using Planck’s quantum hypothesis. He argued that light came in

packets (“photons”) of energy $E = h\nu$ where ν is the frequency of the light. This explained why the electronic energy depended only on the light frequency, & not its intensity.



Apparatus for observation of the photoelectric effect.

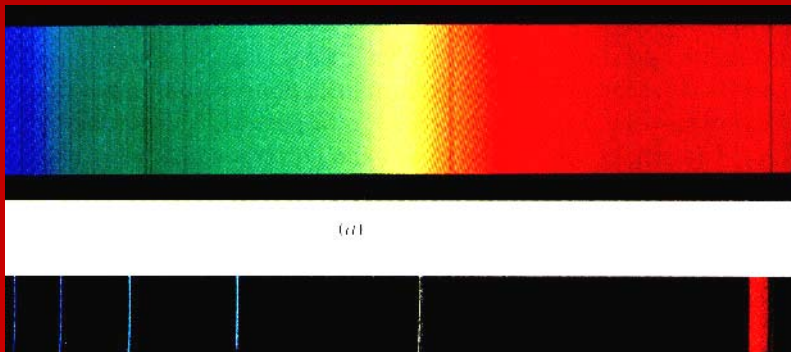


Photoelectric effect- the threshold V is invariant.

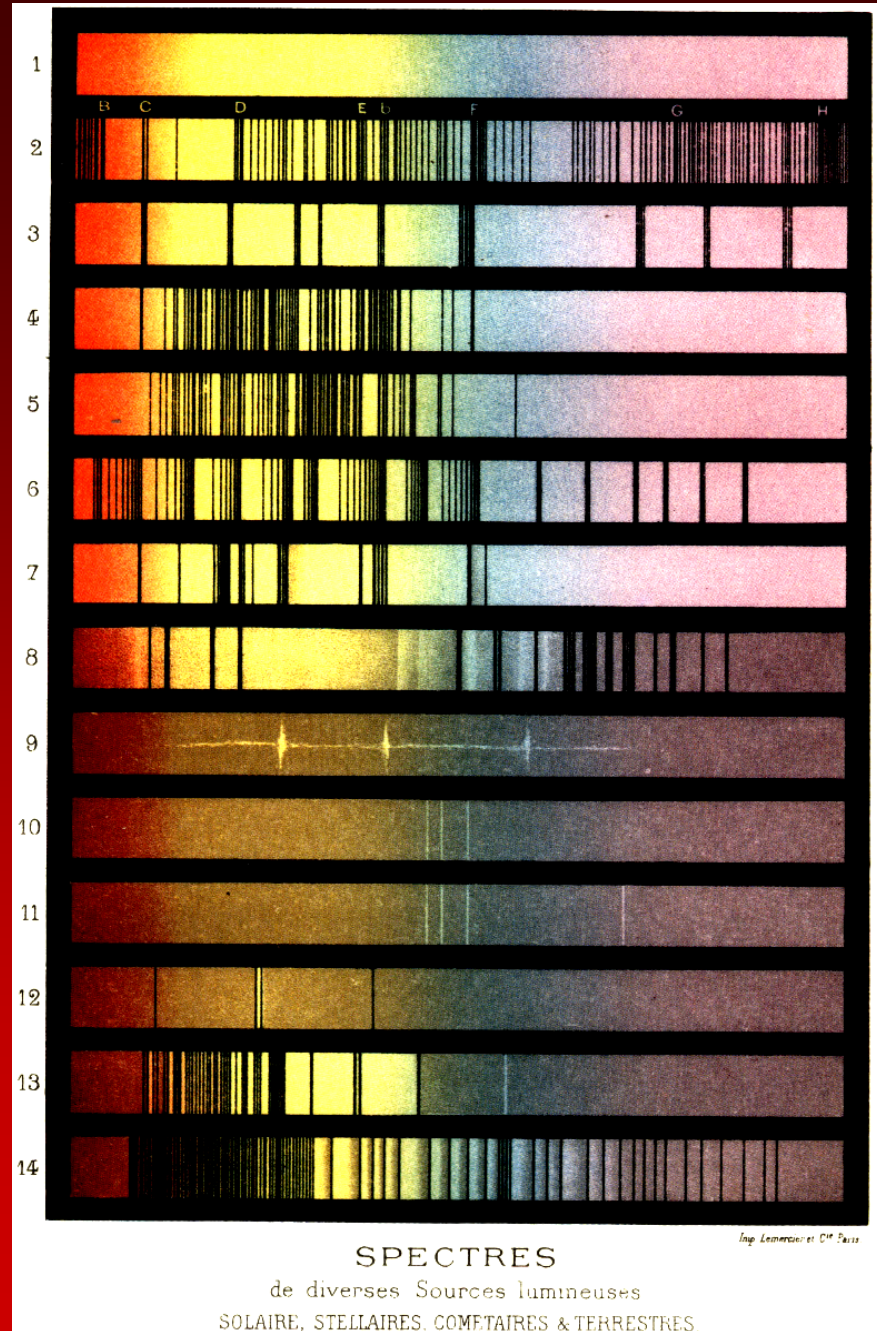
ATOMIC SPECTRA

It was confidently claimed by the French positivist philosopher A. Comte that nobody could ever know what the stars were made of. However the discovery that the “Fraunhofer lines” in the spectra of stars including the sun were the same as those in the spectra of elements on earth allowed a chemical analysis of the stars (reinforced when unknown lines in the solar spectrum were identified by Lockyer in 1868 to be from a newly-discovered element- he called it **He**).

The question was- what was causing these lines?



Above: emission and absorption spectra of **H**
 Right: spectra of stars (from Flammarion, 1880)



RADIOACTIVITY

A development which seemed to be completely unrelated began with the discovery of radioactivity by H. Becquerel in 1896, and the demonstration by P. and M. Curie, in a series of painstaking experiments, that radioactivity was the transmutation of certain elements to others with the concomitant emission of high-energy invisible rays.



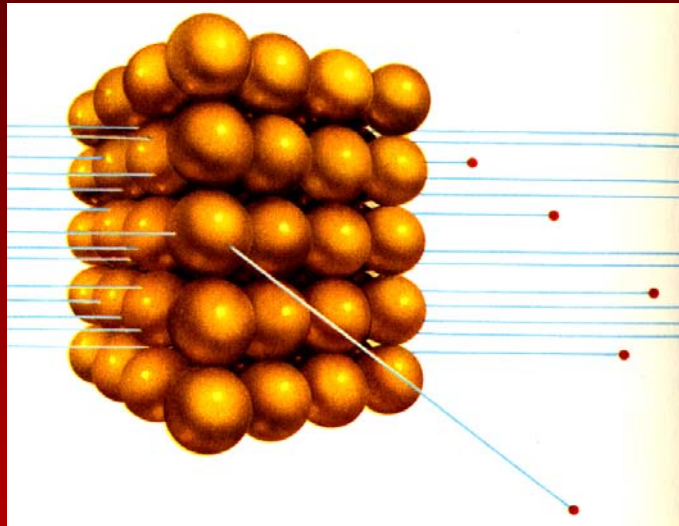
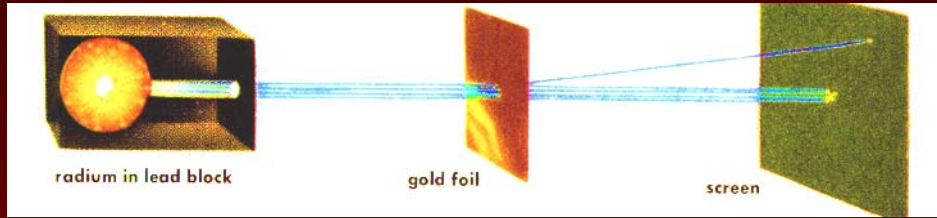
Above: the shed in the Ecole Normale where they worked



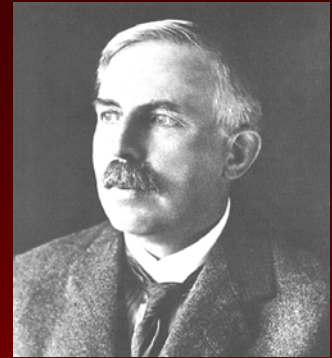
Pierre (1859-1906) and Marie Curie (1867-1934)

The isolation of new radioactive elements, and the mystery of the underlying physics, caused a sensation- particularly as the conservative French establishment refused to properly acknowledge the role of Marie Curie. By the time she was awarded her 2nd Nobel prize (in the middle of a scandalous love affair with Langevin), she was famous around the world- and had to hide in England for 6 months from the French press.

Discovery of the ATOMIC NUCLEUS

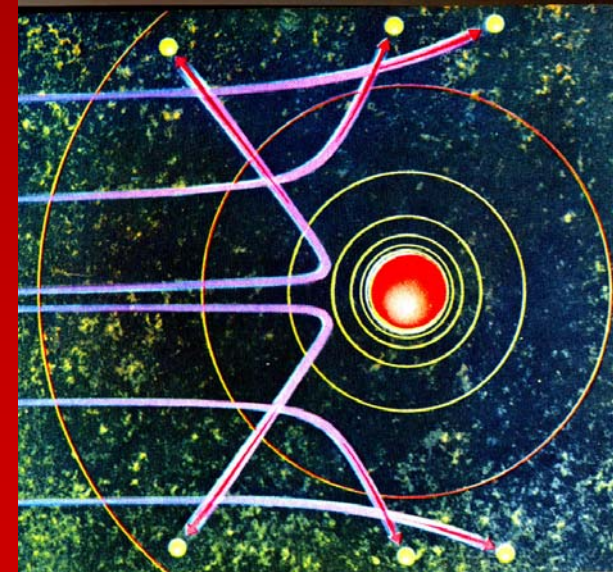


The mystery of the radioactive elements was solved in brilliant work by Rutherford, mostly done with Soddy in McGill in 1904-07, between periods in Cambridge and Manchester.



Rutherford (1871-1937)

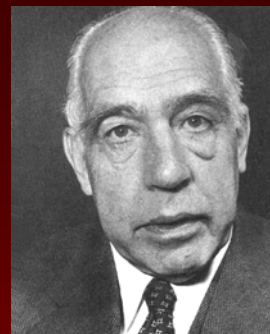
He tried to probe the distribution of charge inside the atom by firing fast radioactive particles at a very thin sheet of Gold leaf (only about 100,000 atoms thick). To his astonishment almost all the particles went straight through- except for a few that were very strongly scattered (see left).



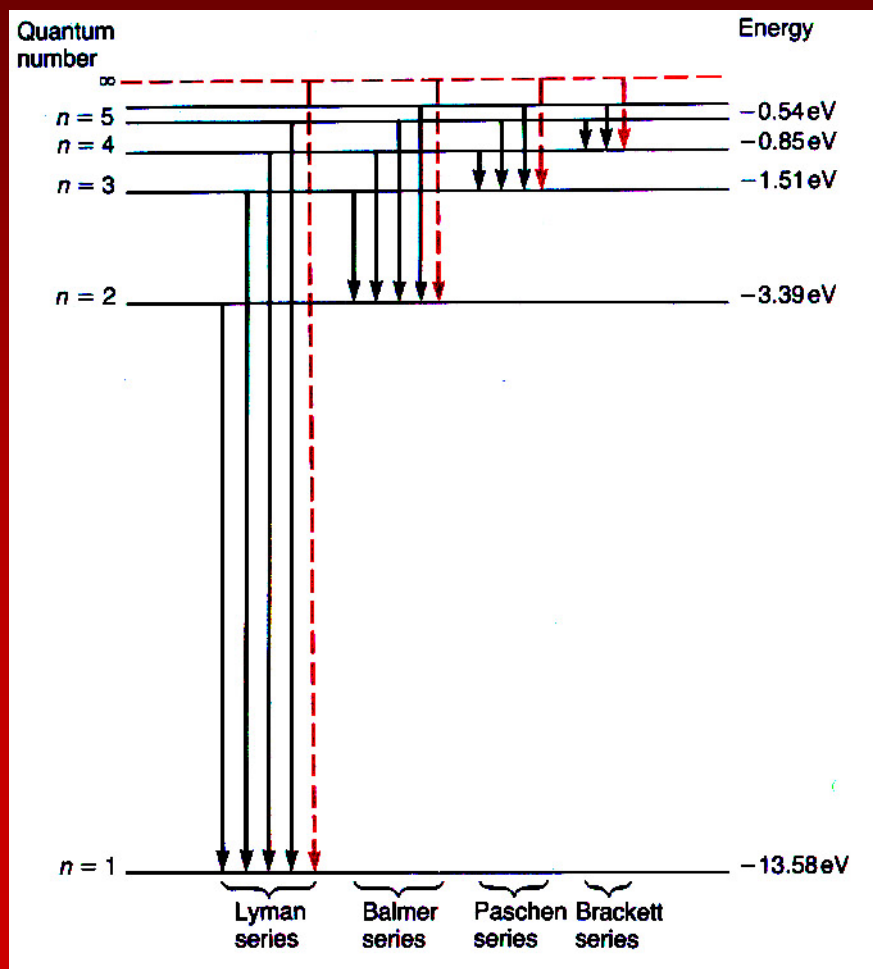
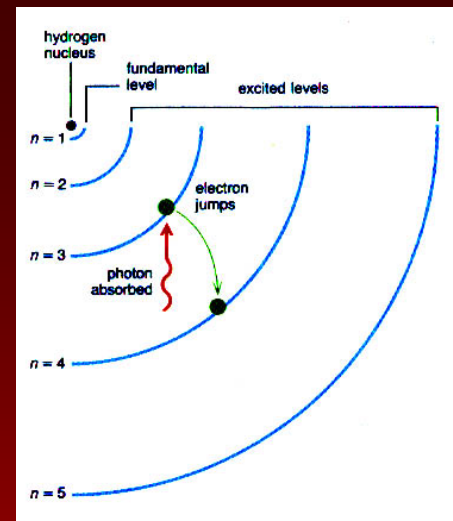
Unlike most experimentalists then (and now), Rutherford did not labour under the delusion that experimentalists should leave the thinking to the theorists. He promptly worked out a model theory which explained the results quantitatively- the positively charged particles were being repelled by a positively charged atomic nucleus, as shown at right. The nucleus was fantastically small- roughly 100,000 times smaller than the atom, or 10^{-15} m across.

The BOHR ATOM

The “old quantum theory” of Bohr gave a partial understanding of the Hydrogen spectrum. Bohr supposed a set of “allowed” orbits of electrons around a **H** nucleus, labeled by a “quantum number” **n**.



N. Bohr (1885-1962)



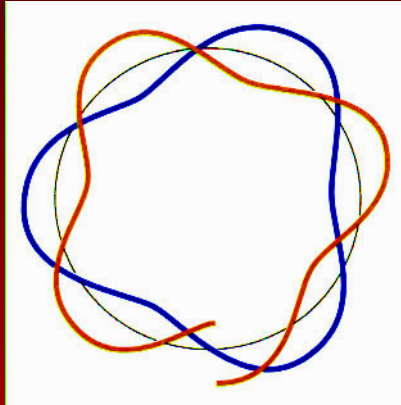
An atom could emit or absorb radiation in this theory only in discrete amounts- in “quantum jumps” between allowed orbits (shown at left for **H**). The energy of the radiation is

$$E_{nm} = h\nu_{nm}/2\pi = (\epsilon_n - \epsilon_m)$$

where ν_{nm} is the frequency of the radiation, and ϵ_n & ϵ_m are the allowed energies of the electrons. This gave an explanation of spectral lines- although there was no explanation of why the electron orbits had to be “quantized”.

MATTER WAVES

To take the crucial next step it helped to not know too much. Prince Louis de Broglie, studying for his PhD in Paris, advanced the idea that matter was associated with waves, and the momentum \mathbf{p} of the waves was given by

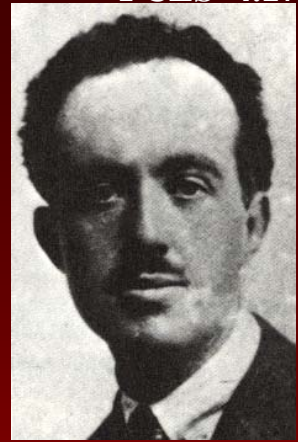


Atomic wave-functions
Only the blue closes on itself.

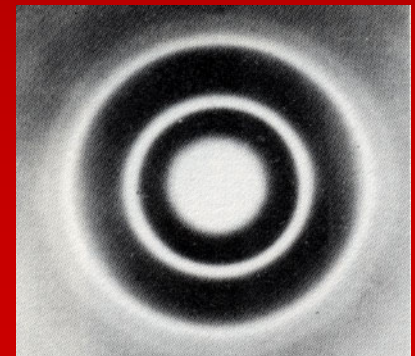
$$\mathbf{p} = \mathbf{h}/\lambda$$

where λ is the wavelength of the matter wave. Because Planck's constant is so small, the wave effects are undetectable except for very light systems. Suppose, eg., we have a mass **1 kg**, moving at speed **1 m/s.**, so it has unit momentum (remember $\mathbf{p} = \mathbf{mv}$ in classical physics). Then, since $\lambda = \mathbf{h}/\mathbf{p}$, we find the very small wavelength of **6.63×10^{-34} m**. This is **$> 10^{19}$** times smaller than the atomic nucleus! However an electron moving at the same **1 m/s** will have a wavelength $\lambda = 6.63 \times 10^{-34} / 9.1 \times 10^{-31}$ m, just under a millimetre. Thus we should see wave interference and diffraction effects with electrons. These were found a few years after de Broglie's prediction, by Davisson & Germer in the USA, and Thompson in Cambridge (see right).

The quantization of momentum also gave a simple explanation of Bohr's quantization rules for atomic orbits (see above); only orbits which could fit a wave in, and thus having definite wavelengths, were allowed .



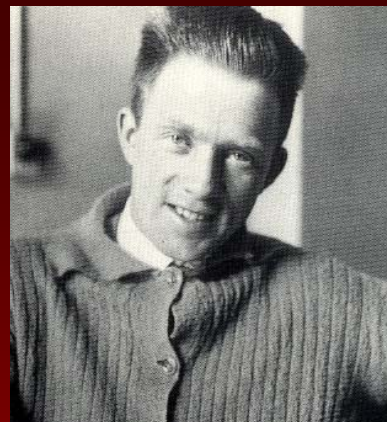
L. De Broglie
(1892-1987)



Electron diffraction by a small hole (Thompson)

The DISCOVERY of QUANTUM MECHANICS

The great theoretical denouement came in 1925. I do not give the details of the theory here, but just sketch the historical development. This began with the “matrix mechanics” of Heisenberg, who found a mathematical way of relating results of measurements on momentum & position- which gave the ‘Uncertainty principle’. His scheme was based on mathematics then unfamiliar to most. Soon after Schrodinger also



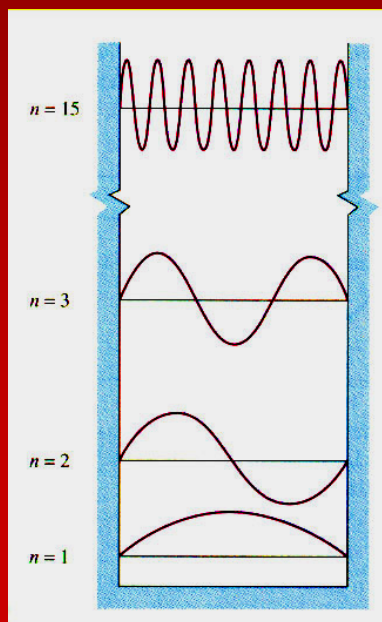
W Heisenberg (1901-76)



E Schrodinger (1887-1961)

gave what seemed a quite different theory, based on a “wave equation” for a matter wave. He was led to this via a deep analysis of the mathematics underlying particle and wave motion, extending old work of WR Hamilton. Although also difficult, this was more familiar territory for most physicists. Then, in what was a real surprise, Schrodinger showed the 2 theories were equivalent.

Schrodinger’s equation showed how to find the “wave function” $\psi(\mathbf{r},t)$ as a function of position \mathbf{r} and time t for any given potential energy. An example is shown (left); a particle is confined to a box, and we see the allowed states, whose energy increases as their wavelength gets shorter.



Wave-functions in a box, & energy levels

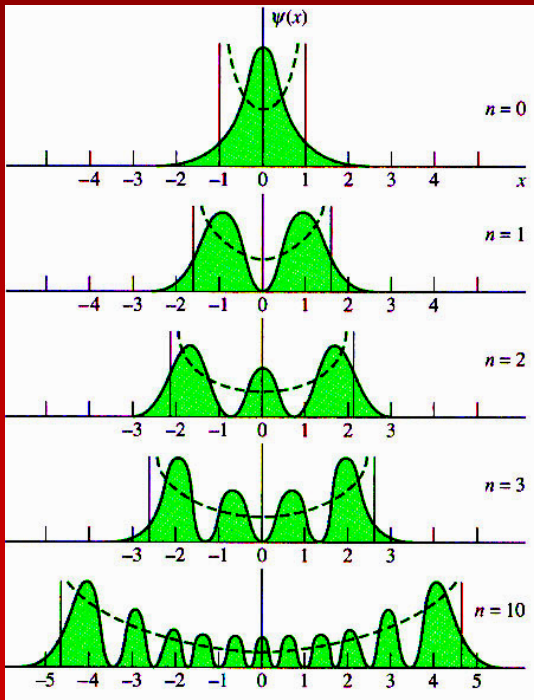
The PROBABILITY INTERPRETATION



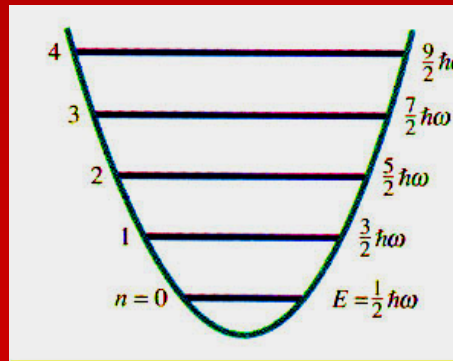
M Born (1882-1970)

The question remained, after the quantum theory was complete- what was the connection between the quantum amplitudes and the physical world? The answer given by M Born was very startling, and changed physics irrevocably. He argued that the

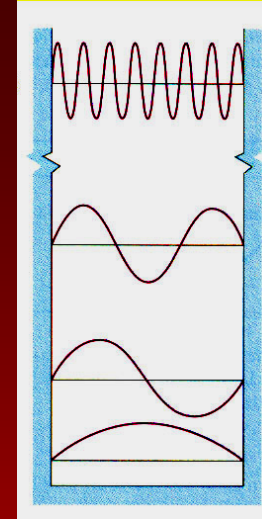
wave amplitudes could be used only to describe the **PROBABILITY** that some phenomenon would occur. Formally, the probability that we see a system at, say, a position **r**, is given by



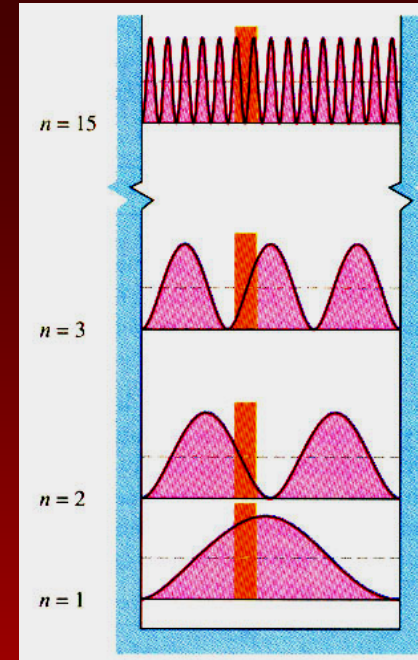
Oscillator probability density



Oscillator energy levels



Energy levels & wave-fns in hard-walled box potential



Probability densities for same potential. The orange band is a position resolution.

$$P(r) = |\psi(r)|^2$$

which means that we take the **MAGNITUDE** of the wave-function, & square it, to get the probability.

QUANTUM MECHANICS of the ATOM

Once quantum mechanics was formulated as a theory, it was almost trivial to find the electronic structure of the **H** atom. The potential is known- it is just the Coulomb interaction we saw back on page 3.11, viz, $V = -kq/r$, where here the charge $q = e$, the electronic charge. So one just solves the Schrodinger equation. For more complicated atoms with lots of electrons one uses powerful computers. The results for the **H** atom are shown below.

The intuitive picture that emerges is that we have a whole set of different wave-functions, with different energies $\epsilon_j = h\nu_j$, oscillating at frequencies ν_j . Only a certain set of energies are allowed- but the system can jump from one “energy level” to another provided the energy difference is taken or given by a photon. This can happen by the absorption of a photon by an electron- increasing its energy- or by emission of a photon, in which case the electron jumps down to a lower energy level (cf. page 4.16).

Thus quantum mechanics opened the way for an understanding of chemistry- details to follow later.



Probability densities for different **H** electronic wave-functions