

Information from fine structure lines: How to excite C⁺, O and N⁺?

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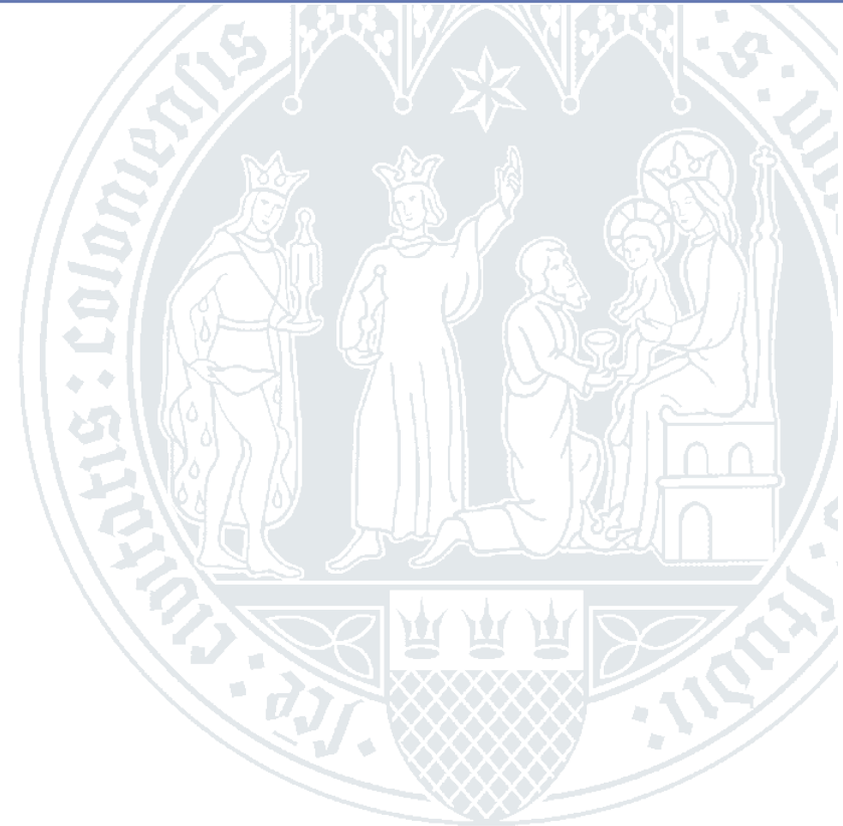
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Repetition of basic atomic physics:

The H-atom

- Simple problem: $p + e^-$

- Pure Coulomb problem:

$$\hat{H} = \frac{\vec{p}^2}{2m_r} - \frac{(Z)e^2}{4\pi\epsilon_0|\vec{r}|}$$

- With reduced mass

$$m_r = \frac{m_e m_p}{m_e + m_p}$$

Z = charge for more general case of multiple protons

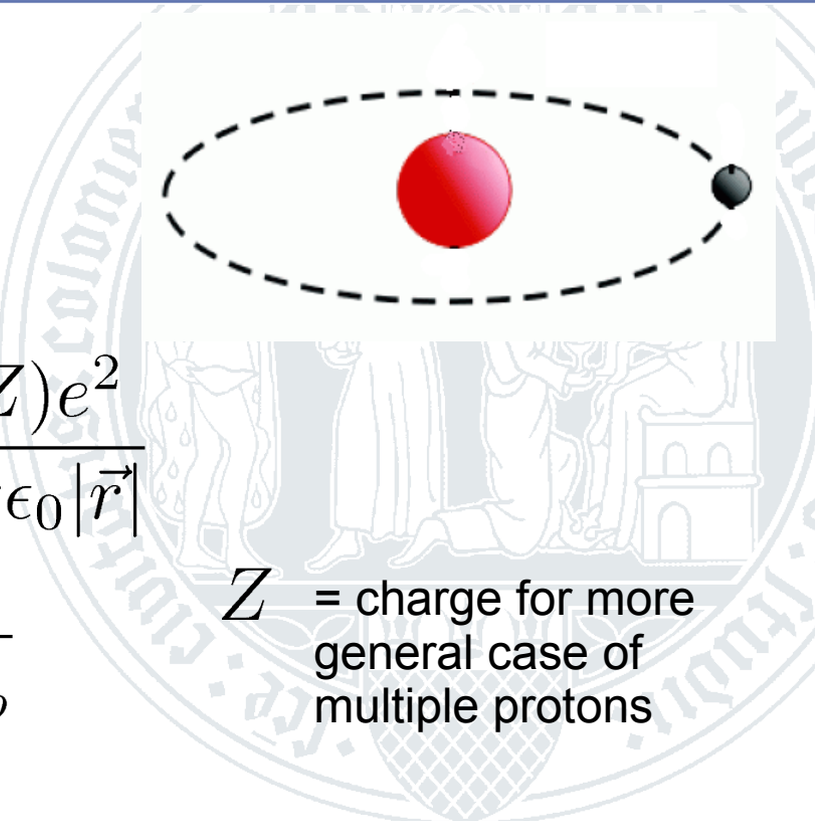
- Solution has eigenvalues

$$E = -R \frac{(Z^2)}{n^2}$$

n = main quantum number

- With Rydberg constant

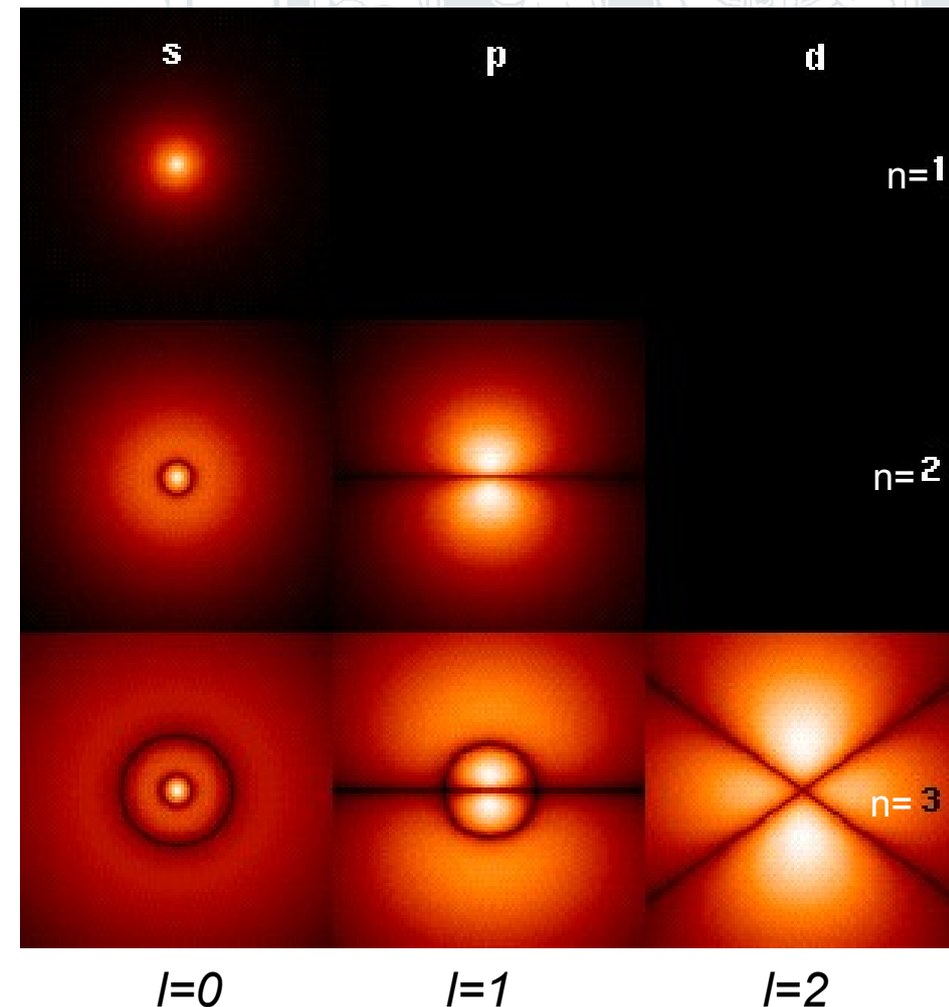
$$R = \frac{1}{2} \left(\frac{e^2}{4\pi\epsilon_0} \right)^2 \frac{m_r}{\hbar^2} \approx 13.605\text{eV}$$



Repetition of basic atomic physics:

The H-atom

- Orbits: Solution of the full spatial problem
 - Provides in total 3 quantum numbers:
 n, l, m_l
 - l = orbital quantum number $l=0\dots n-1$
 - m_l = projection of l on z-axis: $m_l=-l\dots l$
 - Energy levels n are degenerate with respect to l and m_l
 - Corresponding eigenvector: $|n, l, m_l\rangle$



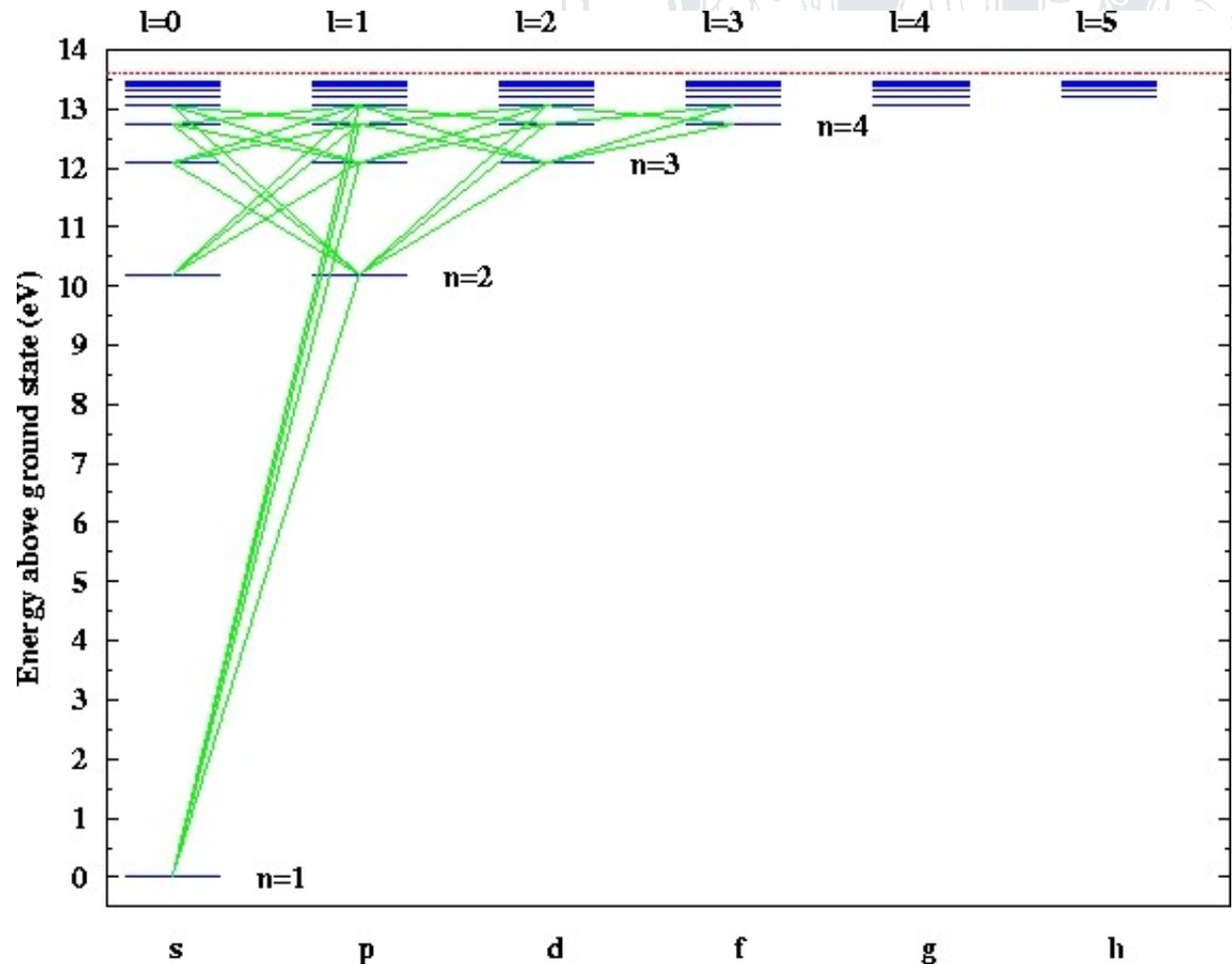
Basics of spectroscopy: The hydrogen atom

Repetition of basic atomic physics:

The H-atom

- Possible transitions
 - from level n to n'

$$E_n - E_{n'} = -(Z^2)R \left(\frac{1}{n^2} - \frac{1}{n'^2} \right)$$



Basics of spectroscopy: The hydrogen atom

Repetition of basic atomic physics:

The H-atom

- Possible transitions
 - from level n to n'
- Gives series for different n'
 - $n'=1$ = Lyman
 - $n'=2$ – Balmer
 - $n'=3$ – Paschen
 - $n'=4$ – Brackett
 - $n'=5$ – Pfund

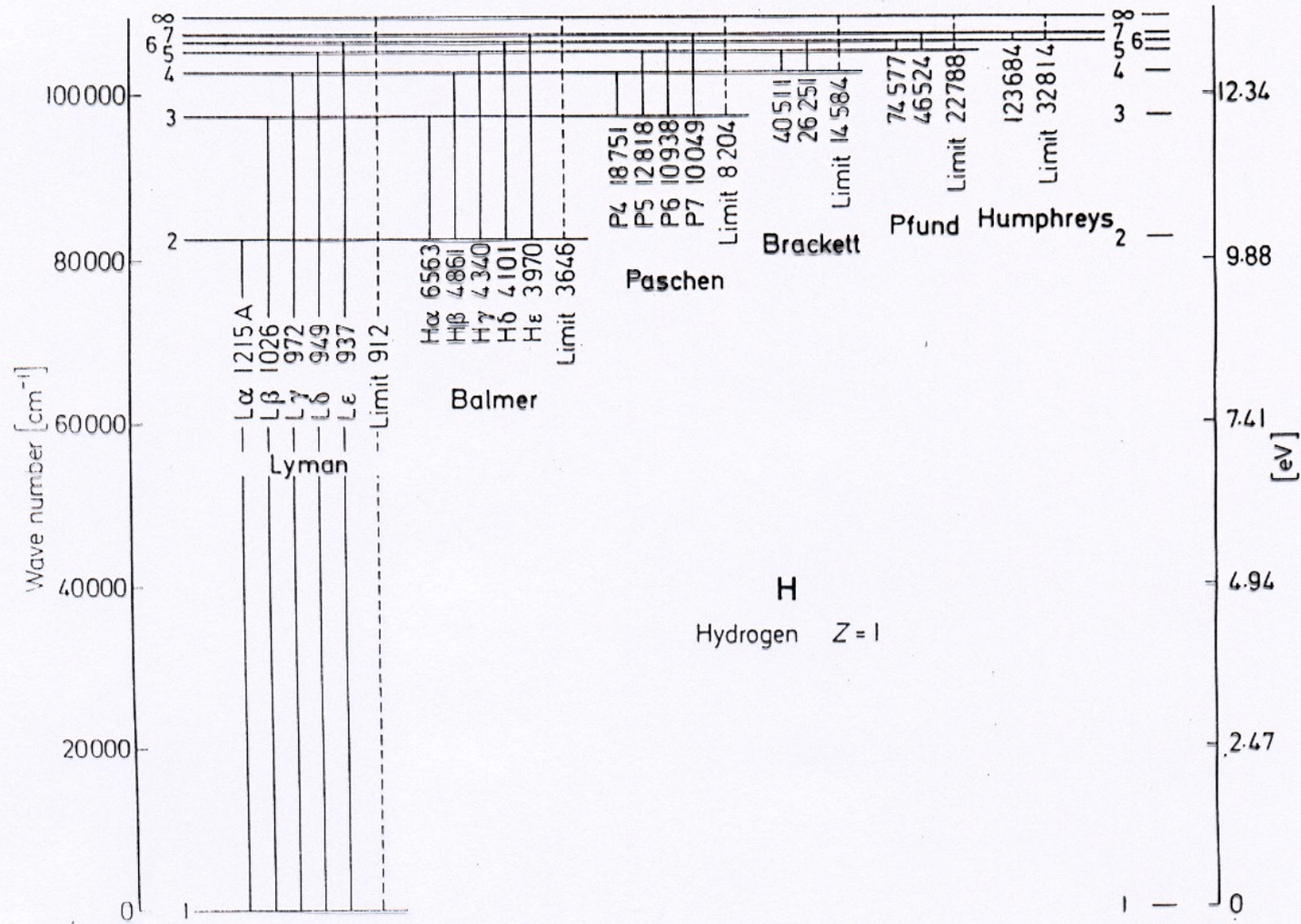


Fig. 13. Grotrian diagrams for H, He I, He II, C II, C III, C IV, O I, O II, O III, N II, Mg I, Mg II, Ca I, Ca II, Fe I, and Fe II. (After MOORE and MERRILL, 1968)

The electron spin

- Adds spin system $|s, m_s\rangle$ with $s=1/2$
- Provides additional coupling term between orbital momentum and spin

$$\hat{H}_{fs} = \frac{e}{2m_e^2 c^2} \left(-\frac{1}{r} \frac{\partial}{\partial r} \frac{(Z)e^2}{4\pi\epsilon_0 |\vec{r}|} \right) \hat{\vec{s}} \cdot \hat{\vec{l}}$$

- Solution has eigenvalues

$$E_{FS} = \alpha^2 R \frac{(Z^4)}{n^3} \frac{j(j+1) - l(l+1) - s(s+1)}{l(l+1)(2l+1)}$$

- j = total angular momentum from $\vec{j} = \vec{l} + \vec{s} \rightarrow j=l-s \dots l+s$

- $\alpha = \frac{e^2}{4\pi\epsilon_0 \hbar c} \approx \frac{1}{137}$ Sommerfeld fine structure constant

The electron spin

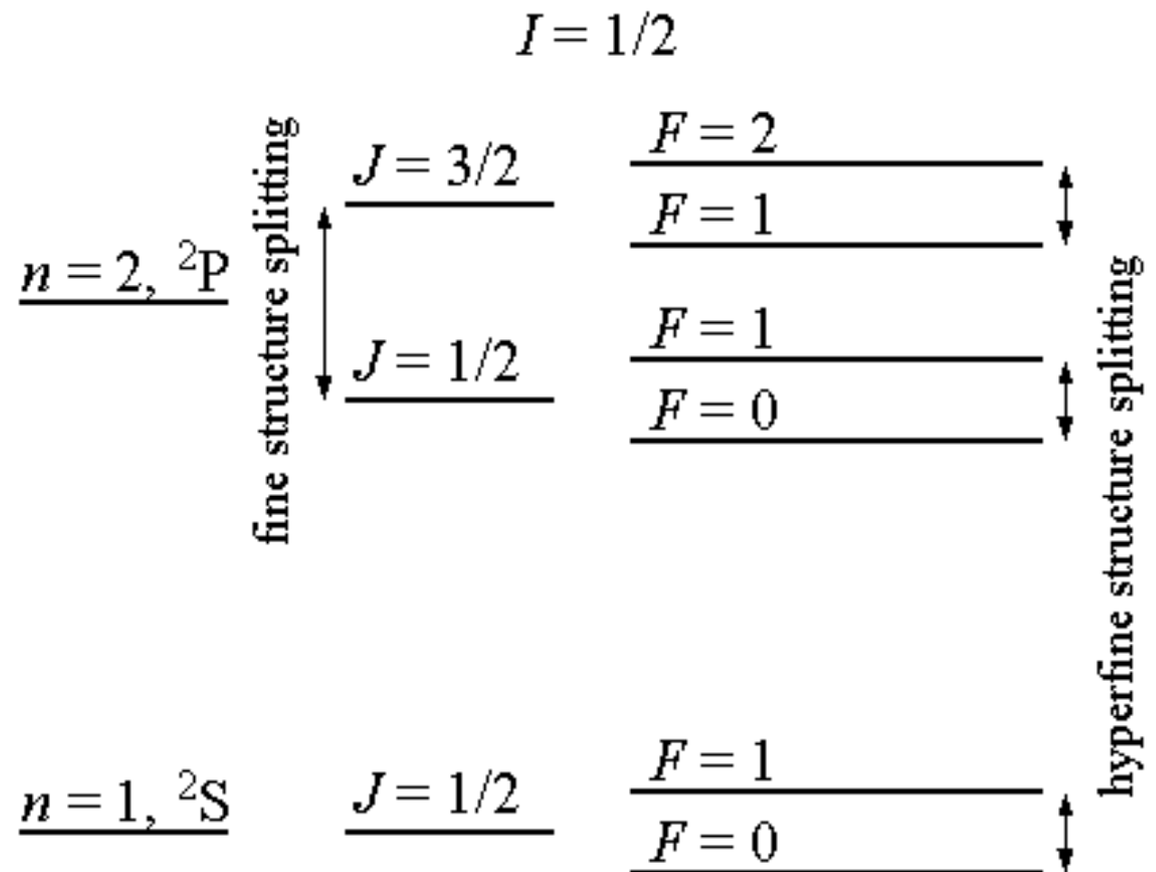
- Fine structure energies are scaled by α^2 relative to main level energies
- H-atom
 - No split for $n=1, l=0$ because $j=1/2$ independent of m_s

- First split for $n=2, l=1$
 $\rightarrow j=1/2, 3/2$

$$\Delta E_{fs} = 4.53 \cdot 10^{-5} \text{eV}$$

$$= 10.9 \text{GHz}$$

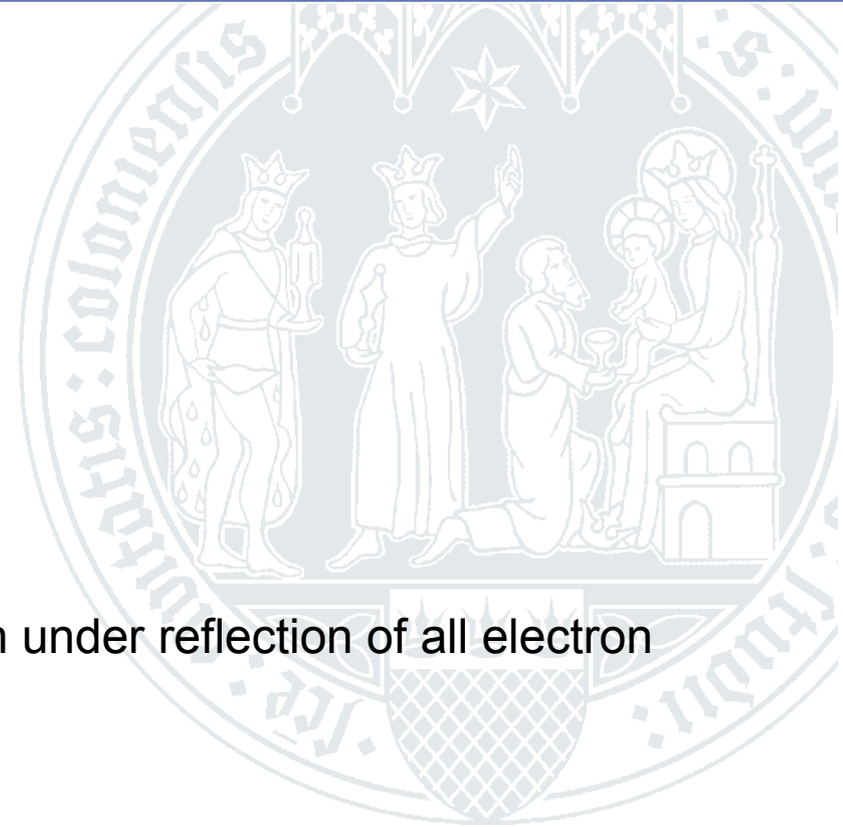
- Requires $n=2$ excitation:
 $10.2 \text{eV} = 120000 \text{K}$
- Not directly observable
 due to other splits



Spectroscopy notation

$$2s+1 l_j^{(p)}$$

- l encrypted in letters
 - $0 = s, 1 = p, 2 = d, 3 = f, 4 = g, 5 = h, \dots$
- p = parity
 - Blank for parity = even, o for parity = odd
 - Characterizes whether wave function changes sign under reflection of all electron positions through the origin.
- **Examples:**
 - $^2s_{1/2}$ = ground state
 - $^2p_{1/2}, ^2p_{3/2}$ = first excited state (Balmer level)



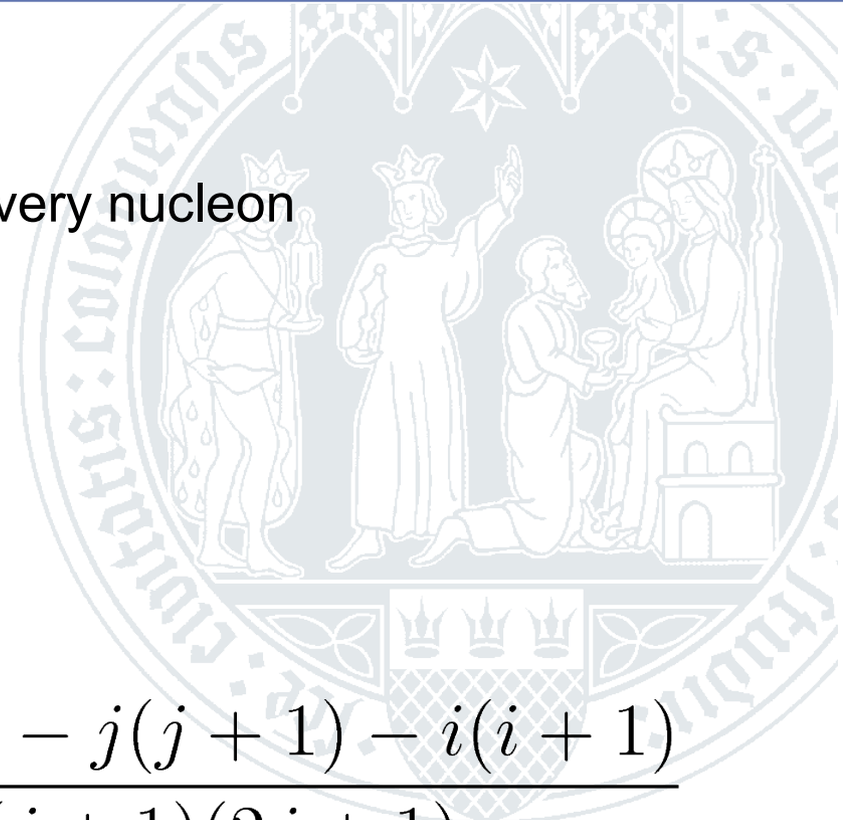
The nuclear spin

- Adds nuclear spin system $|i, m_i\rangle$ with $i=1/2$ for every nucleon
- Treatment equivalent to fine-structure splitting
- New quantum number $\vec{f} = \vec{j} + \vec{i}$
 - f = total angular momentum from $\rightarrow f=j-i\dots j+i$

- Eigenvalues of hyperfine coupling

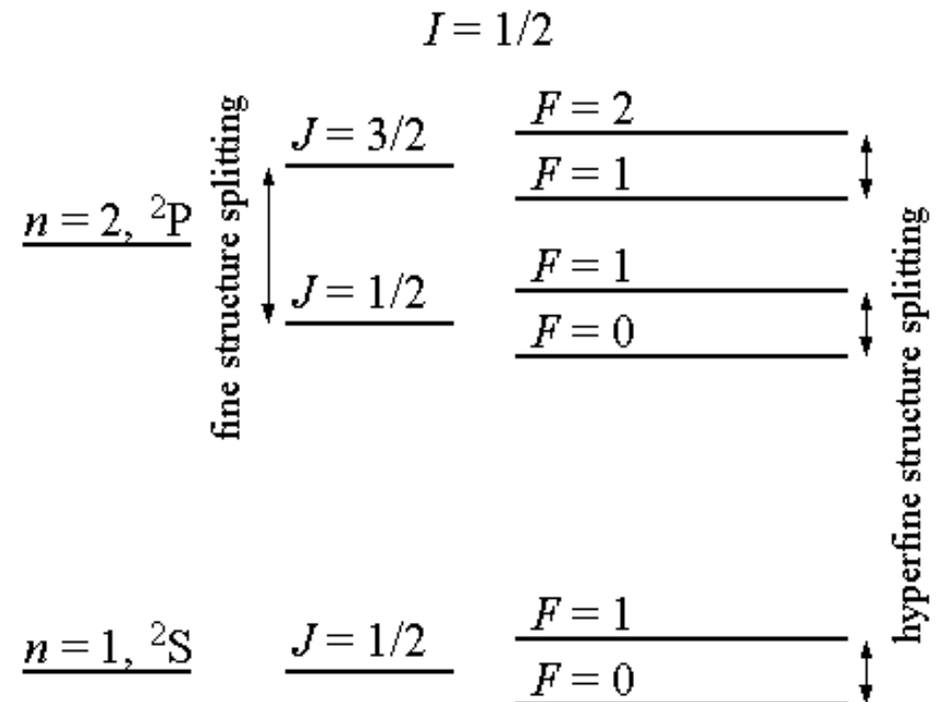
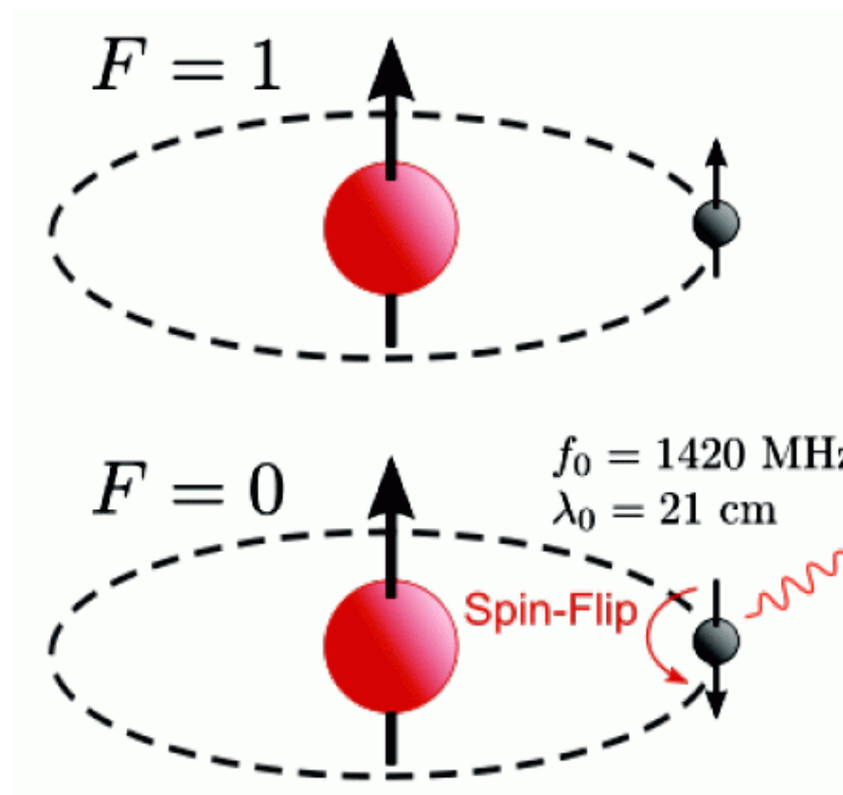
$$E_{HF} = g_N \frac{m_e}{m_p} \alpha^2 R \frac{(Z^4)}{n^3} \frac{f(f+1) - j(j+1) - i(i+1)}{j(j+1)(2j+1)}$$

- Hyperfine splitting typically lower than fine-structure splitting by factor $\frac{m_e}{m_p}$
- g_N – Gaunt factor of the core
- Spectroscopic notation: explicit writing of f



Hyperfine structure

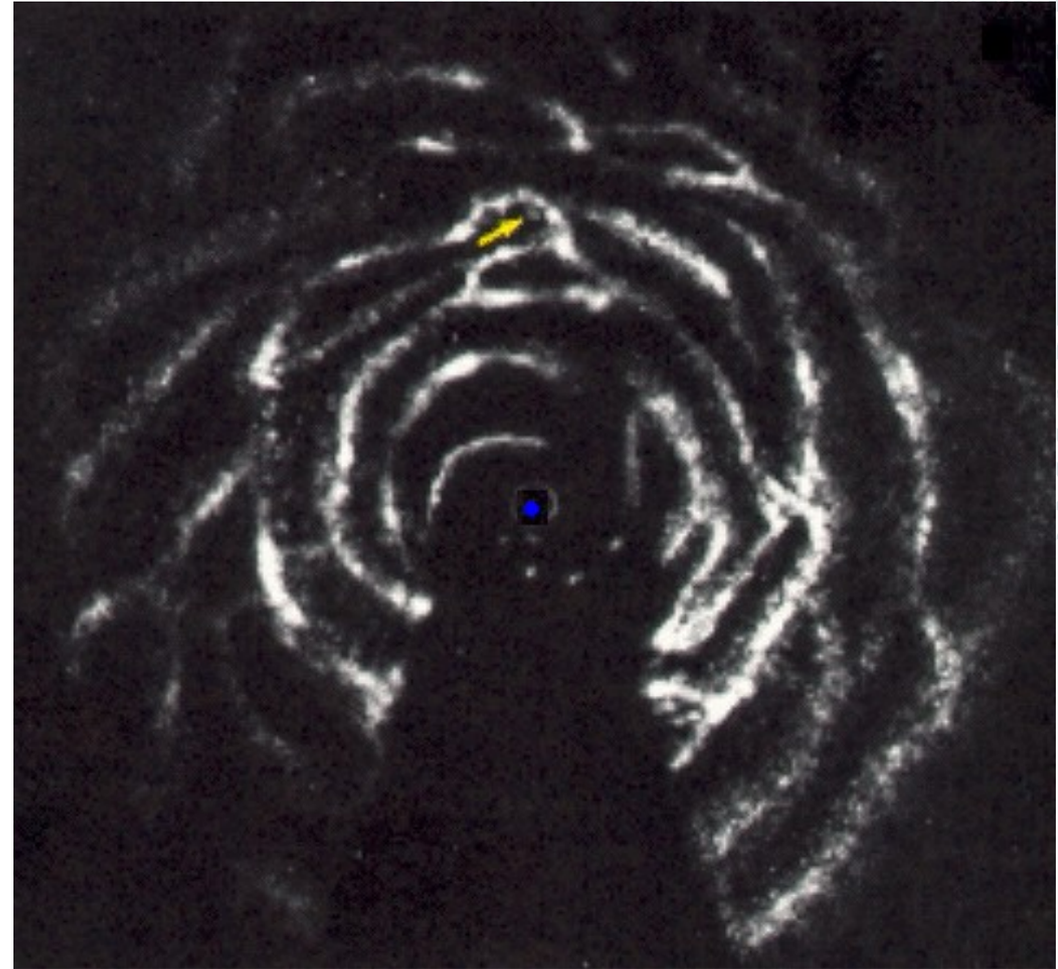
- H-atom
 - for $n=1$ (ground state, $l=0$): $^2s_{1/2}$: $f=1-0$
 - Can be observed for “cold hydrogen”



- famous 21cm line = 1.42GHz
- allows for mapping of the Milky Way
- magnetic dipole transition \rightarrow forbidden
 - $A = 2.9 \cdot 10^{-15} \text{ s}^{-1}$
 - $t = 10^7 \text{ a}$

Hyperfine structure

- H-atom
 - for $n=1$ (ground state, $l=0$): $^2s_{1/2}$: $f=1-0$
 - Can be observed for “cold hydrogen”
- allows for mapping of neutral interstellar gas



1951 Lyman Spitzer: Mapping of the Milky Way

Multiple electrons

- Arrangement in subshells with increasing energy level
 - = increasing main and orbital quantum numbers n, l
- Example: $C^+ = 1s^2 2s^2 2p^1$
 - 3 subshells occupied
 - Closed shells: Sum of orbital momenta and spins always = 0
 - No contribution to radiative interaction
 - Can be ignored
 - 3rd subshell is open:
 - can take 6 electrons: $m_l = -1, 0, 1, m_s = -\frac{1}{2}, \frac{1}{2}$
 - dominates radiative interaction as
 - Only electrons in open shell need to be considered



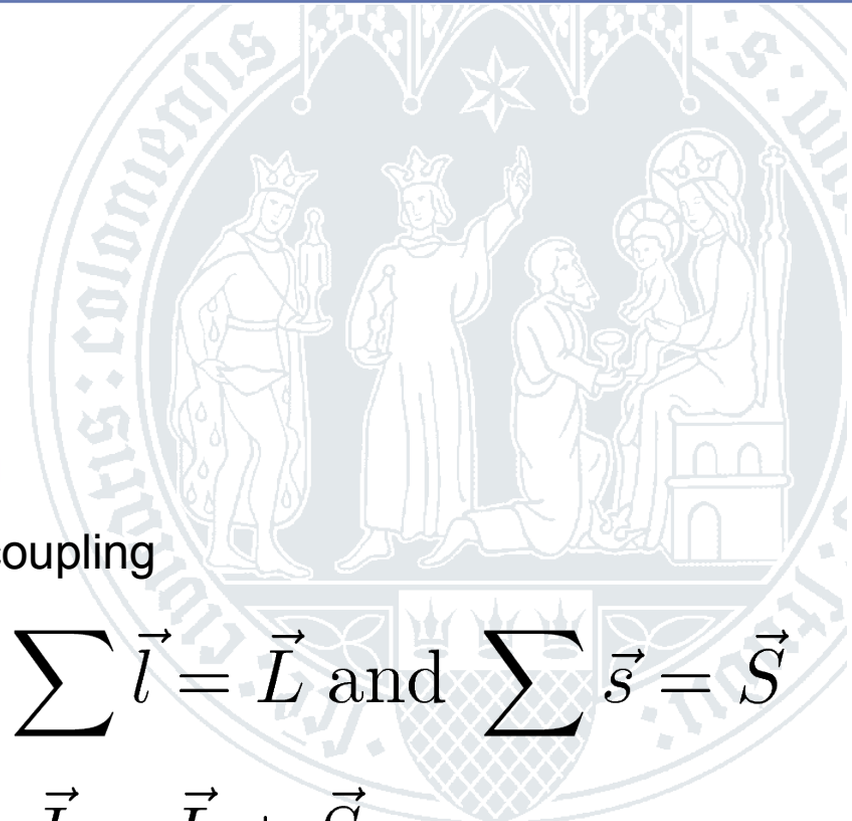
Multiple electrons

- Coupling between different electrons
 - Provides additional term to Hamiltonian
 - No analytic solution
 - Approximation through **Russell-Sounders coupling**
 - Inter-electron coupling stronger than spin-orbit coupling

- Individual orbital momentums and spins add up $\sum \vec{l} = \vec{L}$ and $\sum \vec{s} = \vec{S}$

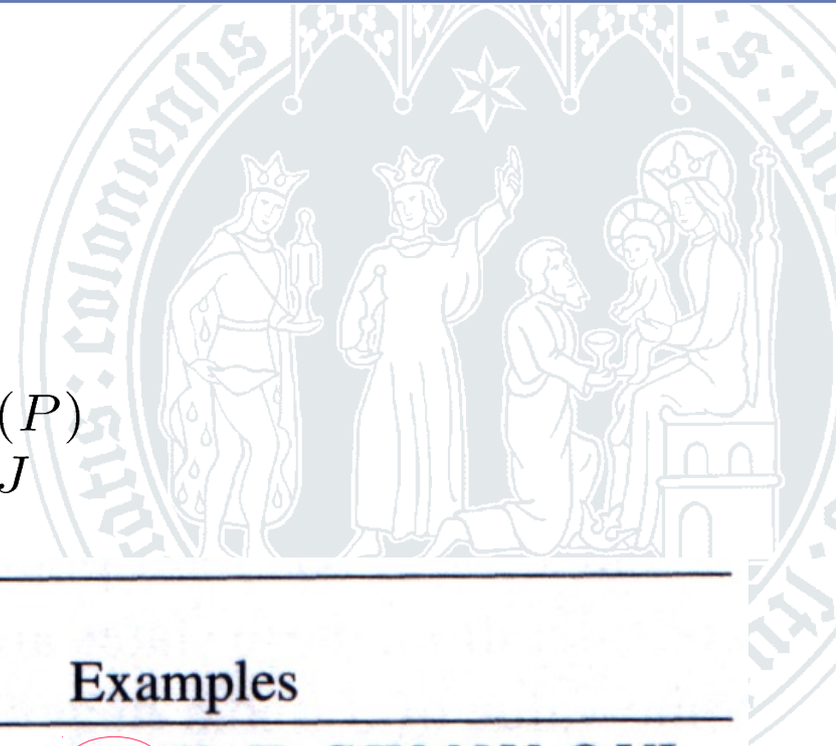
$$\vec{J} = \vec{L} + \vec{S}$$

- **Remember:** approximation only $\rightarrow L, S, J$ are no “good” quantum numbers
- Different approximation for heavy elements
- Spectroscopic notation:
 - capital letters for sum over multiple electrons in open shell
 - explicit writing of F for sum of total angular momentum including nuclear spin



Astrophysically relevant subterms

- Remember H-atom: $2s+1 l_j^{(p)}$
- Equivalently for multi-electron atoms $2S+1 L_J^{(P)}$



Ground configuration	Terms (in order of increasing energy)	Examples
$\dots ns^1$	$^2S_{1/2}$	H I , He II, C IV, N V, O VI
$\dots ns^2$	1S_0	He I, C III, N IV, O V
$\dots np^1$	$^2P_{1/2,3/2}^{\circ}$	C II , N III , O IV
$\dots np^2$	$^3P_{0,1,2}, ^1D_2, ^1S_0$	C I , N II , O III , Ne V, S III
$\dots np^3$	$^4S_{3/2}^{\circ}, ^2D_{3/2,5/2}^{\circ}, ^2P_{1/2,3/2}^{\circ}$	N I, O II, Ne IV, S II, Ar IV
$\dots np^4$	$^3P_{2,1,0}, ^1D_2, ^1S_0$	O I , Ne III, Mg V, Ar III
$\dots np^5$	$^2P_{3/2,1/2}^{\circ}$	Ne II, Na III, Mg IV, Ar IV
$\dots np^6$	1S_0	Ne I, Na II, Mg III, Ar III

Russel-Sounders coupling

- Solution

- “Hund rules”:
 - 1) higher S → lower energies
 - 2) higher L → lower energies
 - 3) lower J and shell \leq half-full → lower energies
 shell $>$ half-full → higher energies

- Example: $C^+ = 1s^2 2s^2 2p^1$

- 2p electron has $S = 1/2, L = 1$

- gives $J = 1/2$ ($m_J = -1/2, 1/2$)

- or $J = 3/2$ ($m_J = -3/2, -1/2, 1/2, 3/2$) = **2-level system**

- Only 3rd Hund rule applies:

$$\Delta E = 7.86 \text{meV}$$

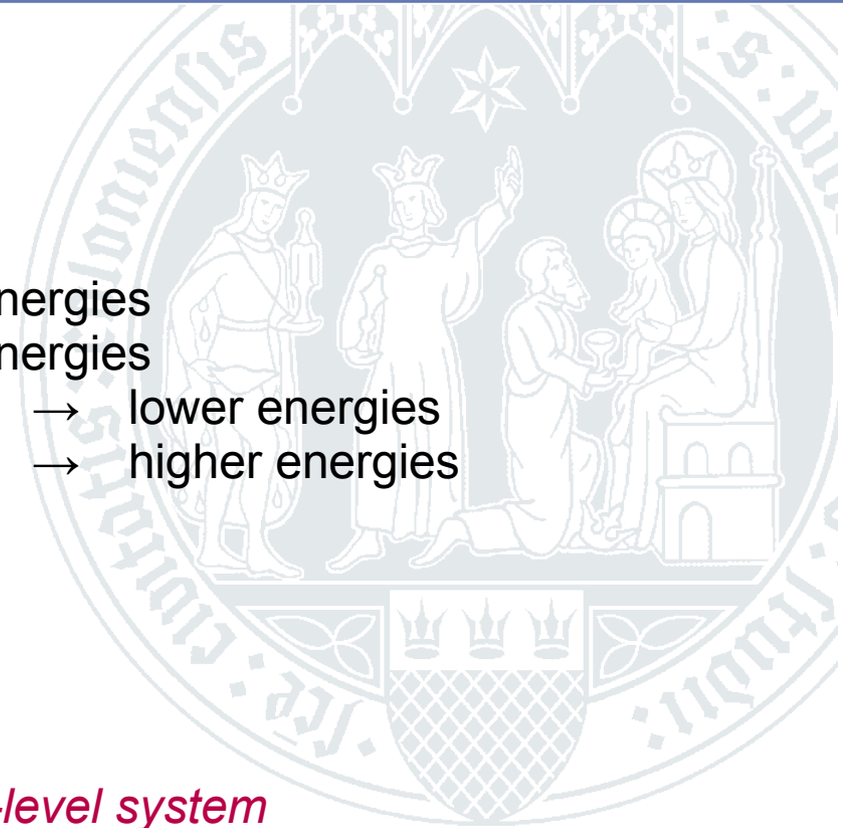
$$= 1900.536 \text{GHz}$$

$$= 157.7 \mu\text{m}$$

$${}^2P_{3/2} \quad g = 4$$

$${}^2P_{1/2} \quad g = 2$$

- Equivalent for N^{++} : $\Delta E = 57 \mu\text{m}$



Hyperfine structure

- Superimposed if nuclear spin does not sum up to 0

Table 17. A table of hyperfine transitions at radio frequencies¹

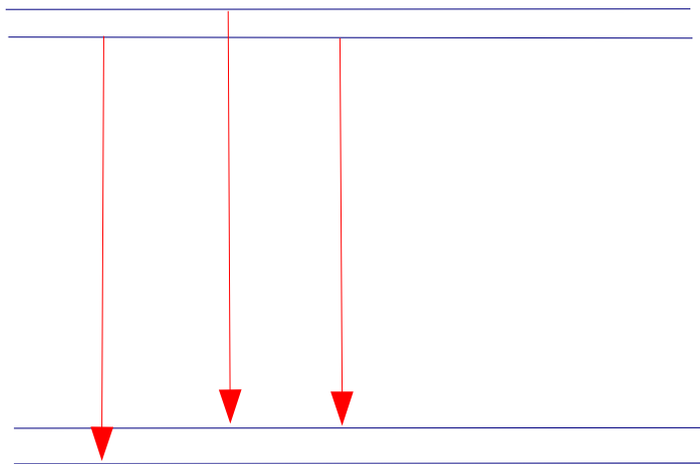
Atom or molecule	Spin	Transition	Frequency (Hz)	A_{mn} (sec ⁻¹)
H I neutral hydrogen	$\frac{1}{2}$	$^2S_{1/2}, F=0-1$	$1.420405751.786 \times 10^9 \pm 0.01$	2.85×10^{-15}
D deuterium	1	$^2S_{1/2}, F=\frac{1}{2}-\frac{3}{2}$	$3.27384349 \times 10^8 \pm 5$	4.65×10^{-17}
He II singly ionized helium	$\frac{1}{2}$	$^2S_{1/2}, F=1-0$	$8.66566 \times 10^9 \pm 1.8 \times 10^5$	6.50×10^{-13}
N VII ionized nitrogen	1	$^2S_{1/2}, F=\frac{1}{2}-\frac{3}{2}$	5.306×10^7	1.49×10^{-19}
N I neutral nitrogen	1	$^4S_{3/2}, F=\frac{3}{2}-\frac{5}{2}$	2.612×10^7	1.78×10^{-20}
		$F=\frac{1}{2}-\frac{3}{2}$	1.567×10^7	3.84×10^{-21}
H ₂ ⁺ ionized molecular hydrogen	1	$F_2, F \frac{3}{2}, \frac{5}{2} - \frac{1}{2}, \frac{3}{2}$	$1.40430 \times 10^9 \pm 10^7$	2.75×10^{-15}
		$F_2, F \frac{3}{2}, \frac{3}{2} - \frac{1}{2}, \frac{3}{2}$	$1.41224 \times 10^9 \pm 10^7$	2.80×10^{-15}
Na I neutral sodium	$\frac{3}{2}$	$^2S_{3/2}, F=1-2$	1.77161×10^9	5.56×10^{-15}

¹ From TOWNES (1957), FIELD, SOMERVILLE, and DRESSLER (1966), and KERR (1968).

Hyperfine structure

- Example: $^{13}\text{C}^+$

- $I = \frac{1}{2}$ combines with $J = \frac{1}{2}$ to $F=0$ ($m_F=0$) or $F=1$ ($m_F=-1, 0, 1$)
- with $J = \frac{3}{2}$ to $F=1$ ($m_F=-1, 0, 1$) or $F=2$ ($m_F=-2, -1, 0, 1, 2$)
- C^+ levels split up



$${}^2P_{3/2} \quad F = 2 \quad g = 5$$

$${}^2P_{3/2} \quad F = 1 \quad g = 3$$

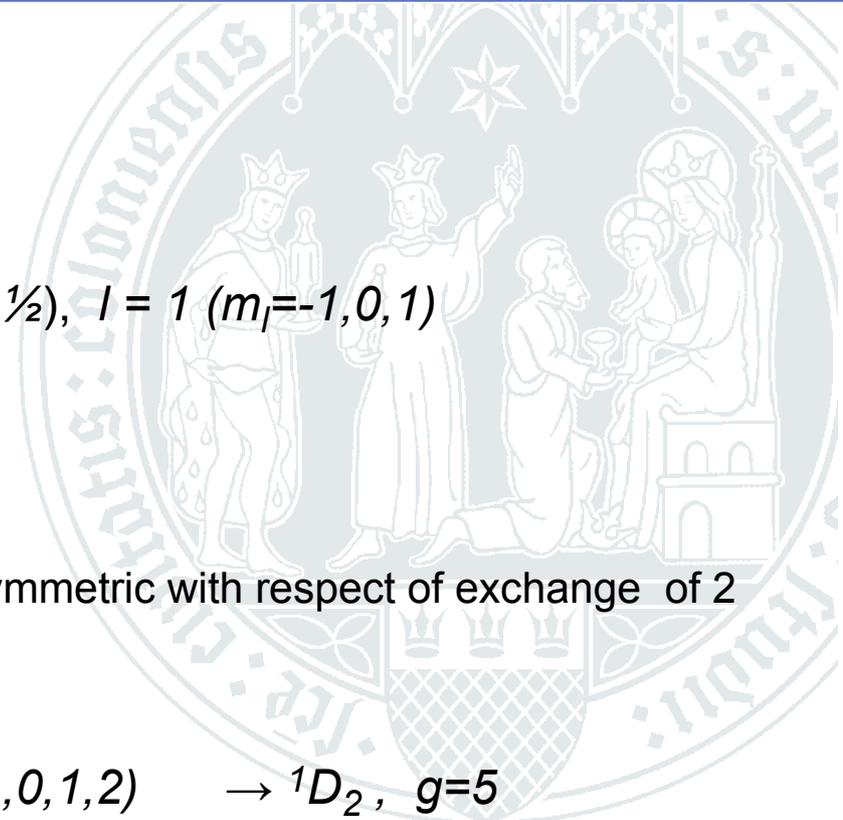
$${}^2P_{1/2} \quad F = 1 \quad g = 3$$

$${}^2P_{1/2} \quad F = 0 \quad g = 1$$

- 3 possible transitions, $F = 2 \rightarrow 0$ is forbidden

Combination of states

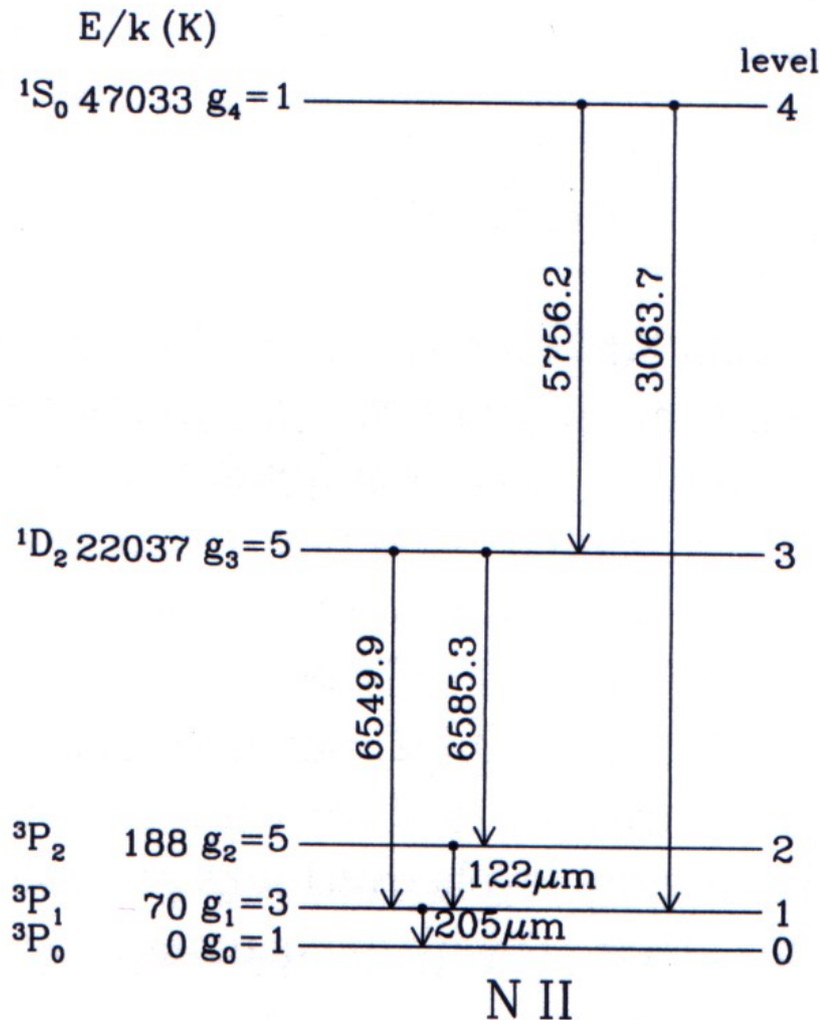
- Example: $N^+ = 1s^2 2s^2 2p^2$
 - Combination of two electrons with $s = \frac{1}{2}$ ($m_s = -\frac{1}{2}, \frac{1}{2}$), $l = 1$ ($m_l = -1, 0, 1$)
→ gives $6 \times 6 = 36$ possible combinations
 - Pauli exclusion principle:
 - 2 electrons never in same state
 - Electrons indistinguishable, i.e. wave functions antisymmetric with respect of exchange of 2 electrons
 - 15 allowed combinations remaining
 - $l_1 \uparrow\uparrow l_2, s_1 \uparrow\downarrow s_2 : L=2, S=0 \quad \rightarrow J=2 (m_J=-2,-1,0,1,2) \quad \rightarrow {}^1D_2, g=5$
 - $l_1 \uparrow\downarrow l_2, s_1 \uparrow\downarrow s_2 : L=0, S=0 \quad \rightarrow J=0 (m_J=0) \quad \rightarrow {}^1S_0, g=1$
 - $l_1 \perp l_2, s_1 \uparrow\uparrow s_2 : L=1, S=1$
 - $\rightarrow J=0 (m_J=0) \quad \rightarrow {}^3P_0, g=1$
 - $\rightarrow J=1 (m_J=-1,0,1) \quad \rightarrow {}^3P_1, g=3$
 - $\rightarrow J=2 (m_J=-2,-1,0,1,2) \quad \rightarrow {}^3P_2, g=5$



Basics of spectroscopy: Combination of states



- “Hund rules”:
 - 1) higher S → lower energies
 - 2) higher L → lower energies
 - 3) lower J and shell \leq half-full → lower energies
 shell $>$ half-full → higher energies



The $S=0$ states need very high excitation energy
 → usually not relevant for ISM physics

→ **treat N^+ as 3-level system**

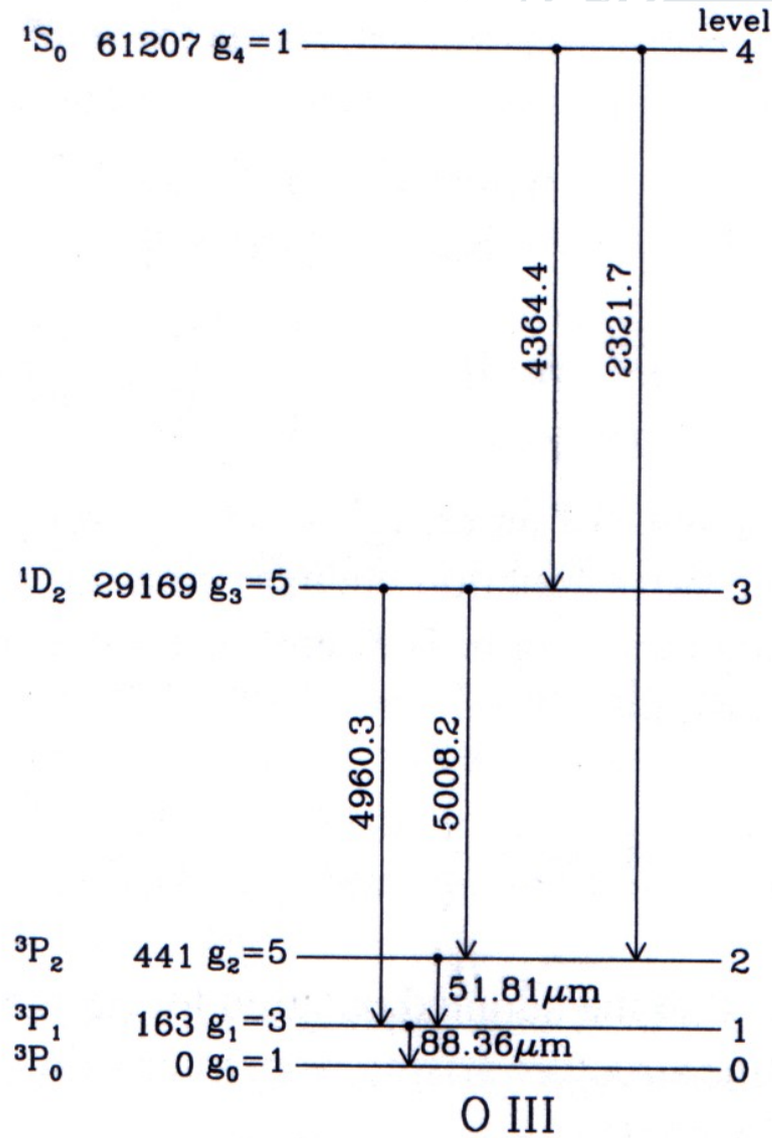
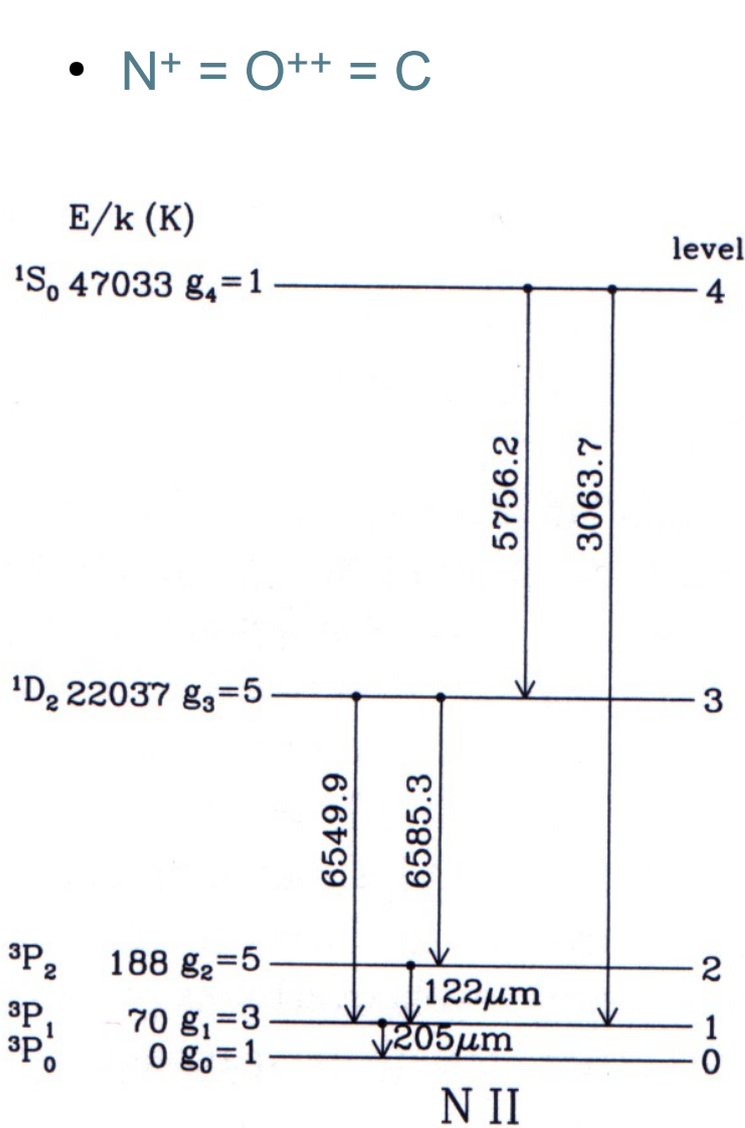
Only 2 radiative transitions. $J = 2 \rightarrow 0$ forbidden

$$122 \mu\text{m} = 2457 \text{ GHz}$$

$$205 \mu\text{m} = 1458 \text{ GHz}$$

Other atoms with same configuration: ${}^3P_{0,1,2}$

- $N^+ = O^{++} = C$

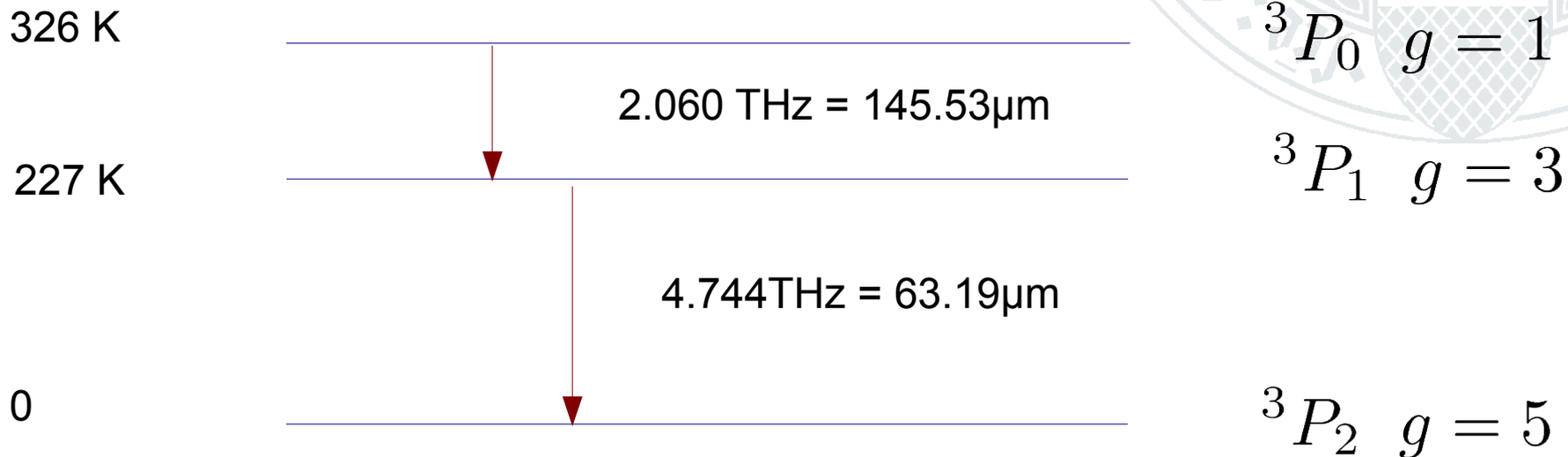


17600		
62		
24		370 μm
0		609 μm
		CI

Basics of spectroscopy: Combination of states

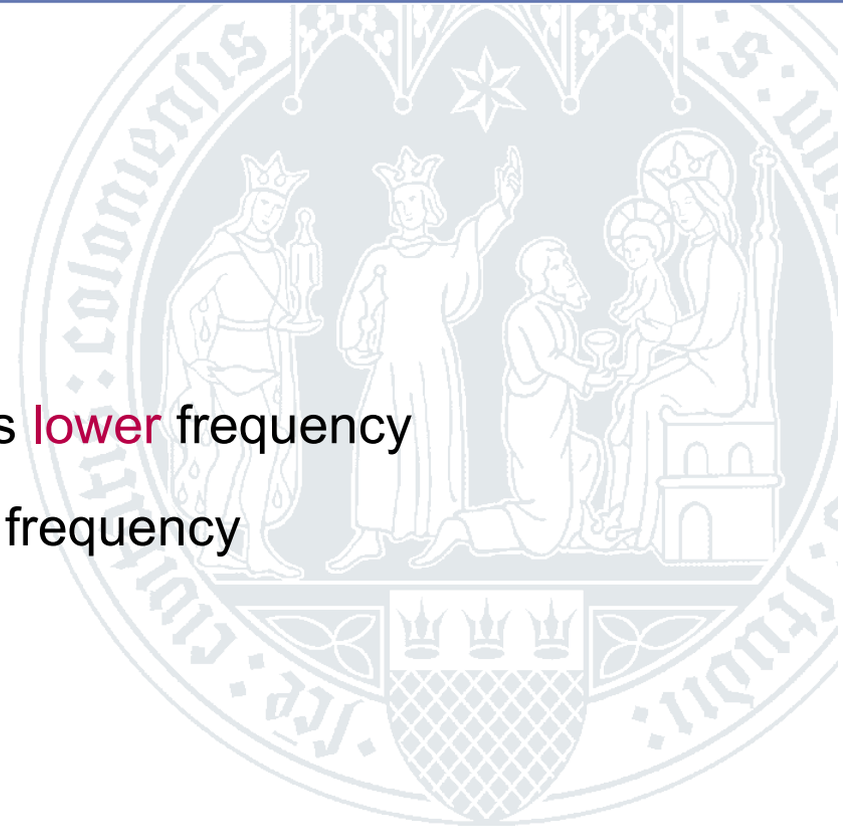


- Similar addition of configurations for all 4 electrons
- Now third Hund rule with “shell > half-full” applies → reverse order: ${}^3P_{2,1,0}$
- Only $S=1$ states excited at moderate temperatures (below 20000K)
 - Effectively **3-level system** as well
 - Radiative transitions only $J = 2 \rightarrow 1$ and $J = 1 \rightarrow 0$



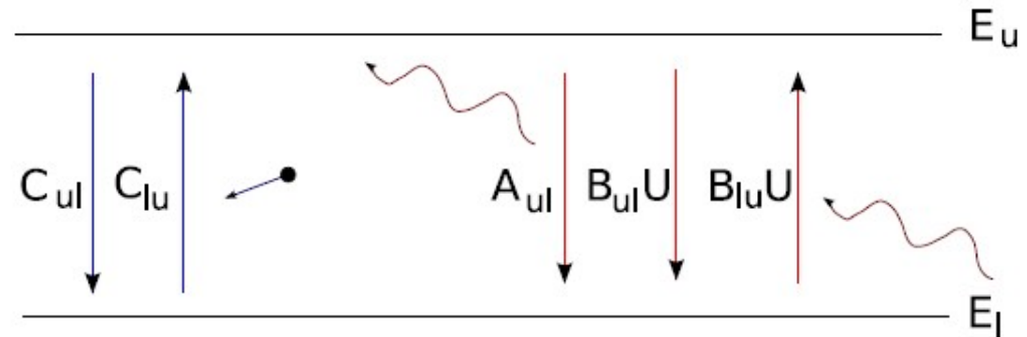
Few cases to distinguish:

- **2-level systems:** HI, C⁺, N⁺⁺
- **3-level systems:** C, N⁺, O⁺⁺, O
 - $^2P_{0,1,2}$ (C, N⁺, O⁺⁺): ground-state transition has **lower** frequency
 - $^2P_{2,1,0}$ (O): ground-state transition has **higher** frequency
- **4-level systems:** $^{13}\text{C}^+$
- **$2p^3$ systems:** N, O⁺ → “Darkness”
 - no fine-structure transitions excited at temperatures $\ll 20000\text{K}$



- Consider transitions between 2 levels:

- Spontaneous emission: A_{ul}
- Stimulated emission: $B_{ul}U$
- Absorption: $B_{lu}U$
- Collisional transitions: C_{ul}, C_{lu}



$$U = \iint \frac{I_\nu \phi(\nu)}{c} d\nu d\Omega = \frac{4\pi}{c} \langle I_{\text{line}} \rangle$$

- Rate coefficients are mutually dependent:

- Number conservation:

$$C_{lu} = C_{ul} \frac{g_u}{g_l} \exp\left(-\frac{h\nu}{kT_{\text{kin}}}\right)$$

$$B_{lu} = B_{ul} \frac{g_u}{g_l}$$

- Quantum mechanics:

$$B_{ul} = A_{ul} \frac{c^3}{8\pi h\nu^3}$$

General case: transitions between multiple levels

- To determine the excitation of the system the matrix of balance equations had to be solved

$$n_i \left(\sum_{j < i} A_{ij} + \sum_{j \neq i} B_{ij} u_{ij} + \sum_{j \neq i} C_{ij} \right) = \sum_{j > i} A_{ji} n_j + \sum_{j \neq i} B_{ji} u_{ij} n_j + \sum_{j \neq i} C_{ij} n_j$$

- n_i - level population of level i
- A_{ij}, B_{ij} - spontaneous, induced radiative rate coefficients
 u_{ij} - radiative energy density at transition frequency
- $C_{ij} = \sum_{i_{coll}}^{n_{coll}} n_{i_{coll}} \gamma_{ij, i_{coll}}$ - collisional rate coefficients for different collision partners i_{coll} :
 $H_2, H, H^+, He, \text{electrons}, \dots$

Description of level populations by excitation temperature T_{ex}

$$\frac{n_j}{n_i} = \frac{g_j}{g_i} \exp\left(-\frac{h\nu_{ij}}{kT_{ex,ij}}\right)$$

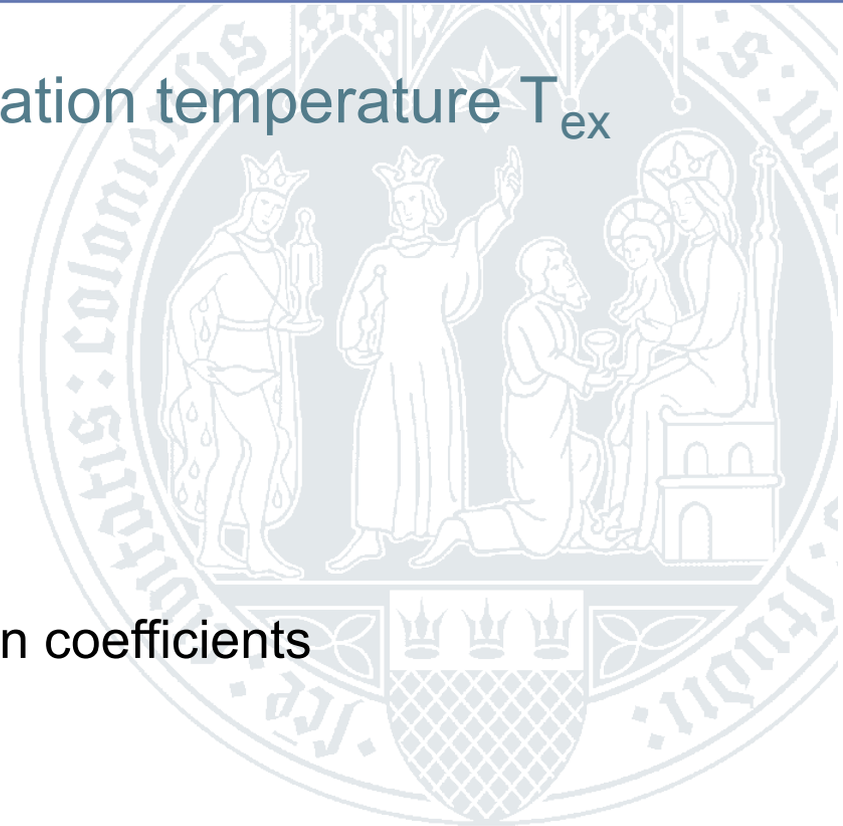
- Usually different for every pair of levels i, j
- Can be obtained exploiting properties of Einstein coefficients

- Radiation: $B_{ij}g_i = B_{ji}g_j$

- Collisions: $C_{ij} = C_{ji} \frac{g_j}{g_i} \exp\left(-\frac{h\nu_{ij}}{kT_{kin}}\right)$

- for $j > i$

→ 3 limiting cases



Limiting cases

- Collision-dominated ($n_{i_{coll}}$ large): $C_{ij} \gg A_{ij}, B_{ij}u_{ij}$ for all i,j

→ Balance equation:
$$\frac{n_i}{g_i} \sum_{j \neq i} C_{ij} = \sum_{j \neq i} \frac{n_j}{g_j} C_{ij} \exp\left(\frac{h\nu_{ij}}{kT_{kin}}\right)$$

– Solution:

- $T_{ex,ij} = T_{kin}$ for all i,j

= LTE (local thermodynamical equilibrium)

Limiting cases

- Radiation dominated (u_{ij} large): $B_{ij}u_{ij} \gg A_{ij}, C_{ij}$ for all i, j

→ Balance equation:
$$\frac{n_i}{g_i} \sum_{j \neq i} B_{ij} u_{ij} = \sum_{j \neq i} \frac{n_j}{g_j} B_{ij} u_{ij}$$

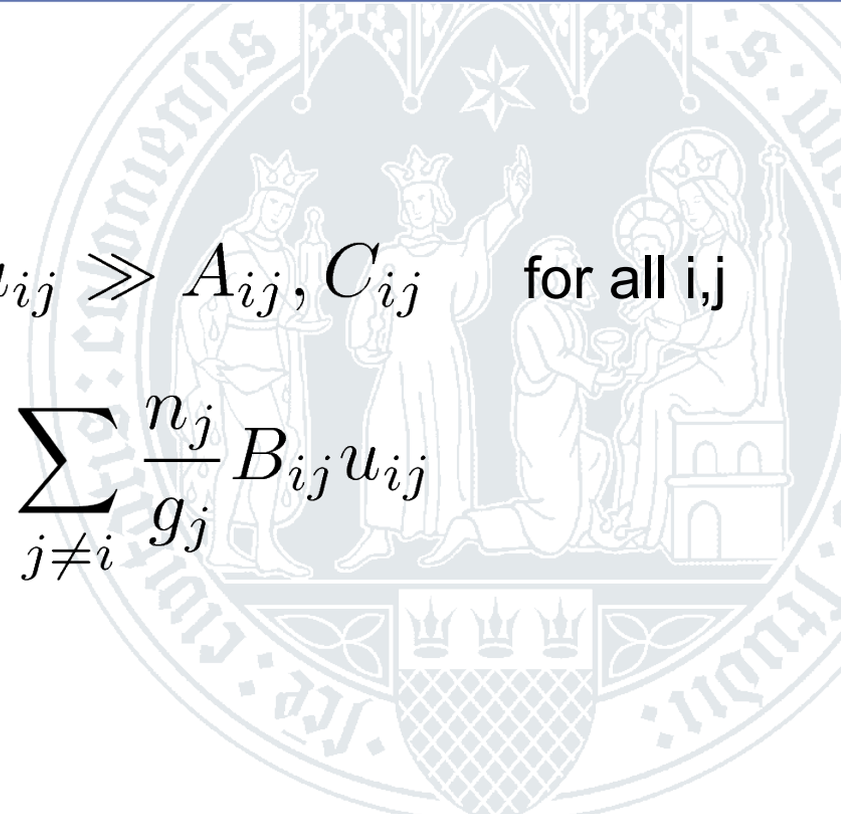
– Solution:

- $n_i/g_i = \text{const.}$
- $T_{ex,ij} = \infty$

- Vacuum ($n_{i_{coll}}, u_{ij}$ small): $A_{ij} \gg B_{ij}u_{ij}, C_{ij}$ for all i, j

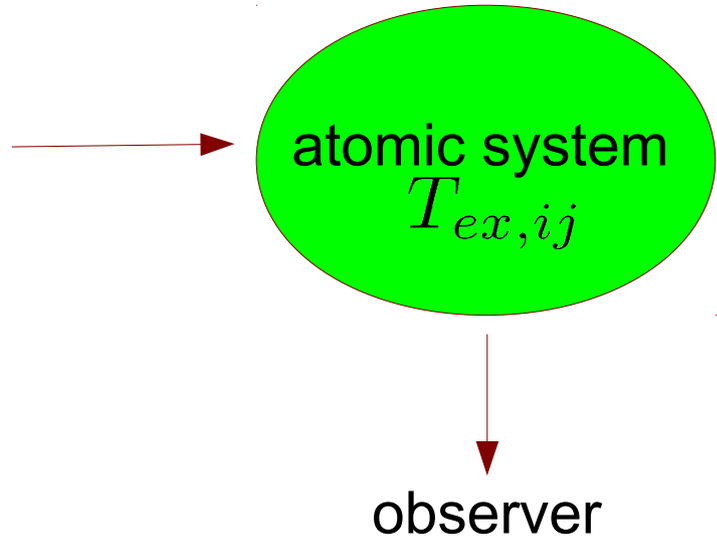
– Solution:

- $n_i \ll 1 \quad \forall i > 0$
- $T_{ex,ij} = T_{bg}$

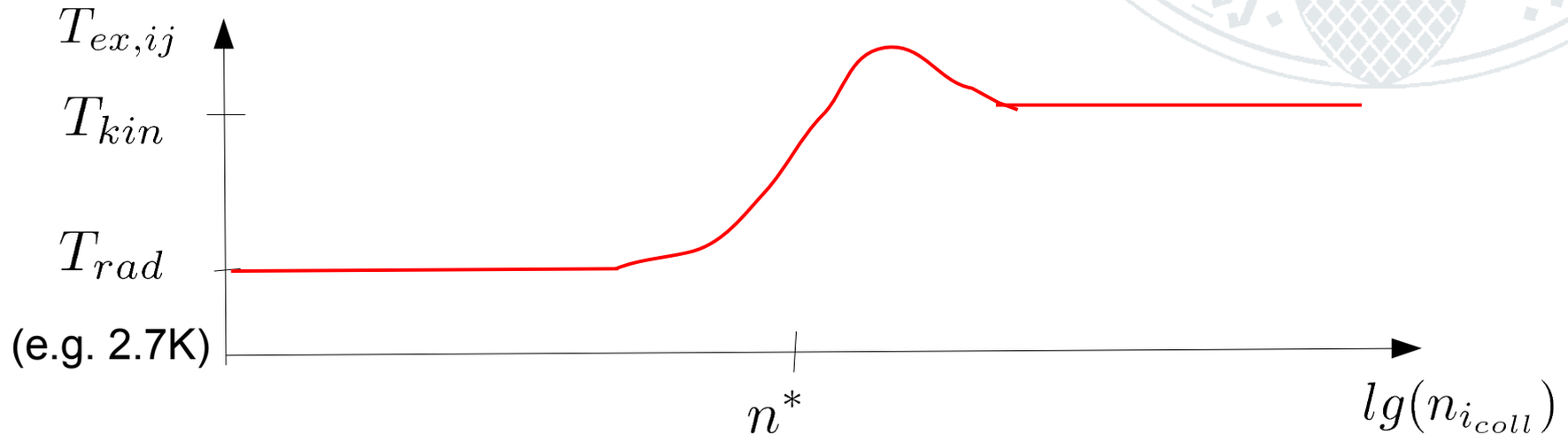


General case

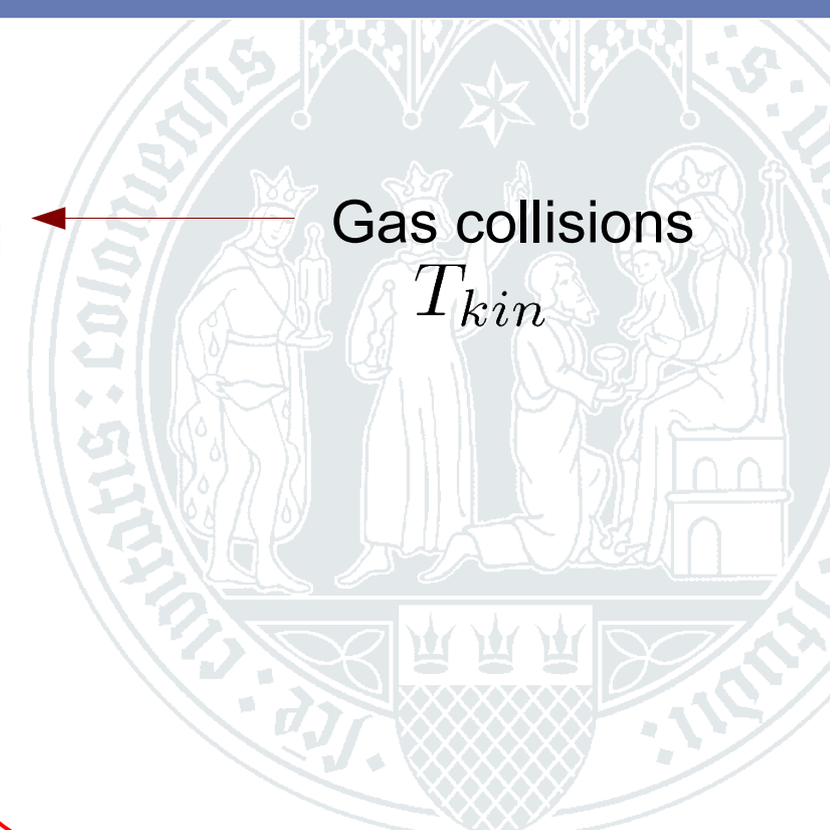
- Radiation field
 T_{rad}



- Observable excitation temperature:



- Transition roughly around critical density: $n_{i_{coll}}^* = A_{ij} / \gamma_{ij, i_{coll}}$



Gas-phase conditions

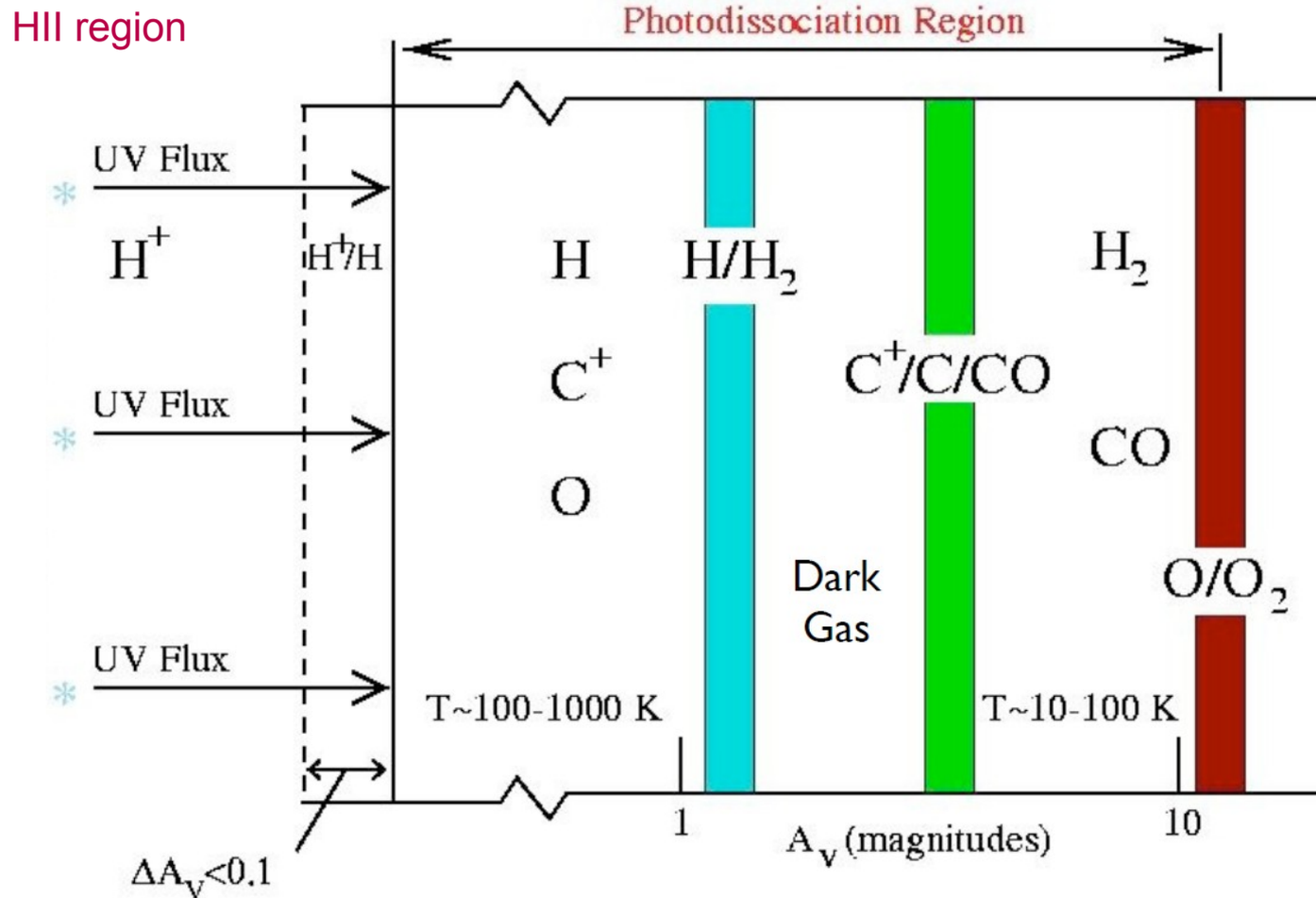
Element	number abundance	1 st ionization	2 nd ionization potential
- H	100	13.6 eV	
- He	9	24.6 eV	54.4 eV
- O	0.026	13.6 eV	35.1 eV
- C	0.012	11.3 eV	24.4 eV
- N	0.008	14.5 eV	29.6 eV

Wakelam & Herbst (2008)

- O is neutral whenever H is neutral
- Only C can be (singly) ionized in regions where H is neutral
- OIII traces very energetic UV fields

Spatial distribution

HII/OIII HII/NII/CII H/O/CII H₂/O/CII H₂/O/C



- Determines the main collision partners

Layering:

HII/OIII HII/NII/CII H/O/CII $\text{H}_2/\text{O/CII}$ $\text{H}_2/\text{O/C}$

determines the main collision partners:

- **OIII**

- Electrons: density given by number of H atoms

- **NII**

- Electrons: density given by number of H atoms

- **CII**

- Electrons: density given by H atoms or C atoms

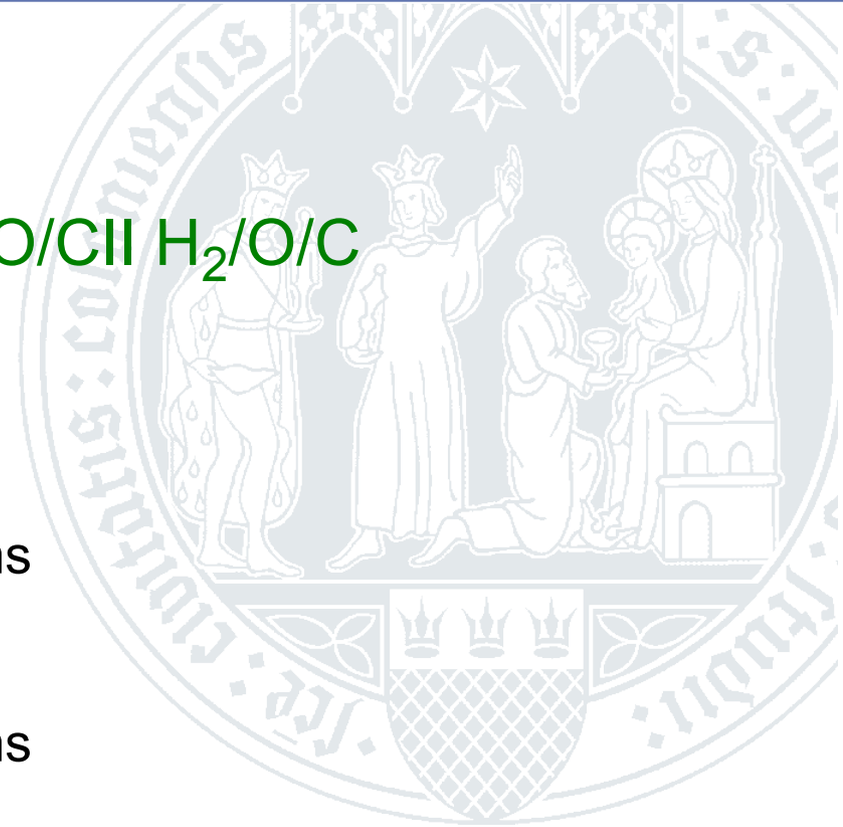
- H-atoms

- H_2 molecules

- **CI**

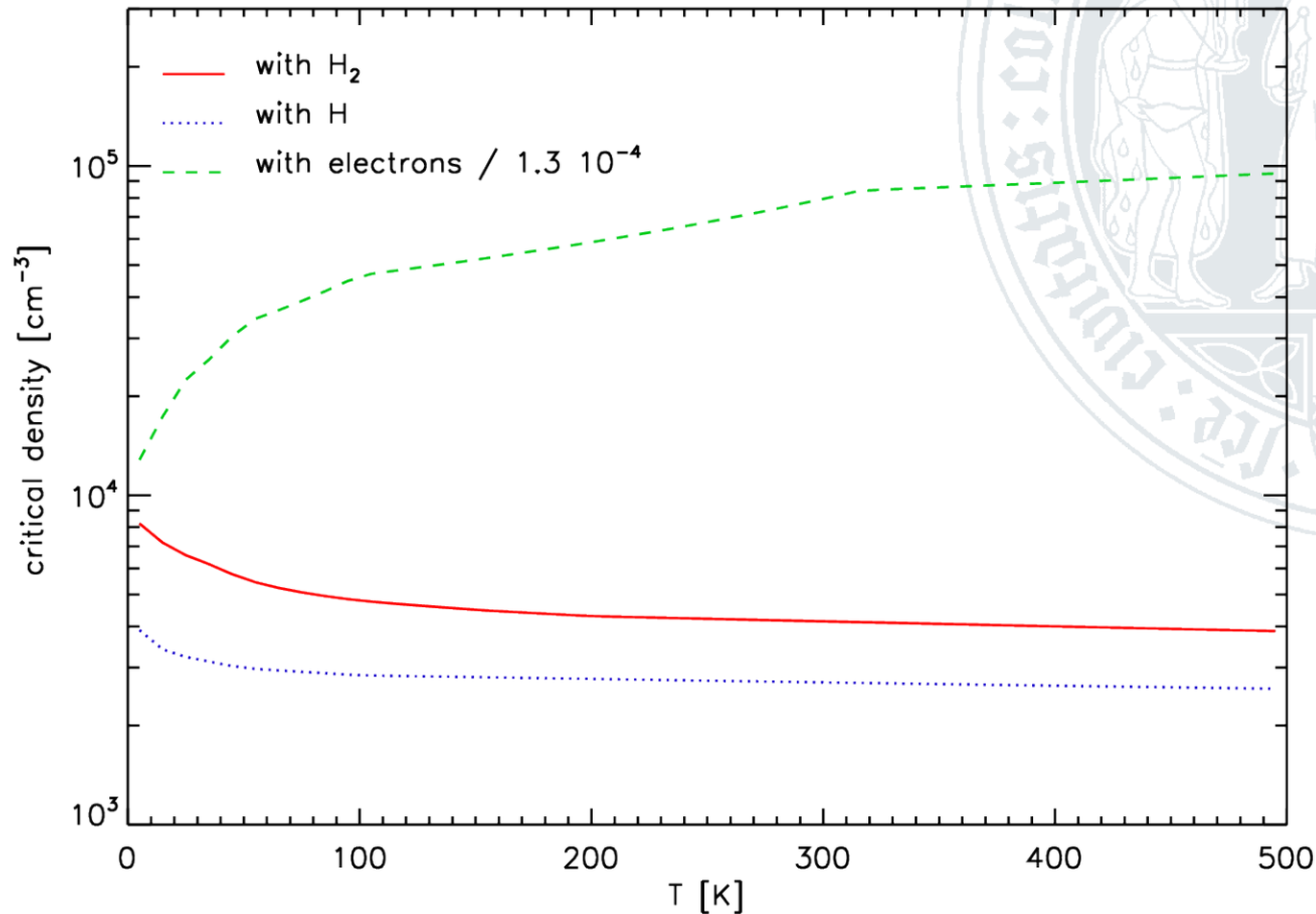
- H_2 molecules

(all have additional minor contribution from He)



[CII] excitation:

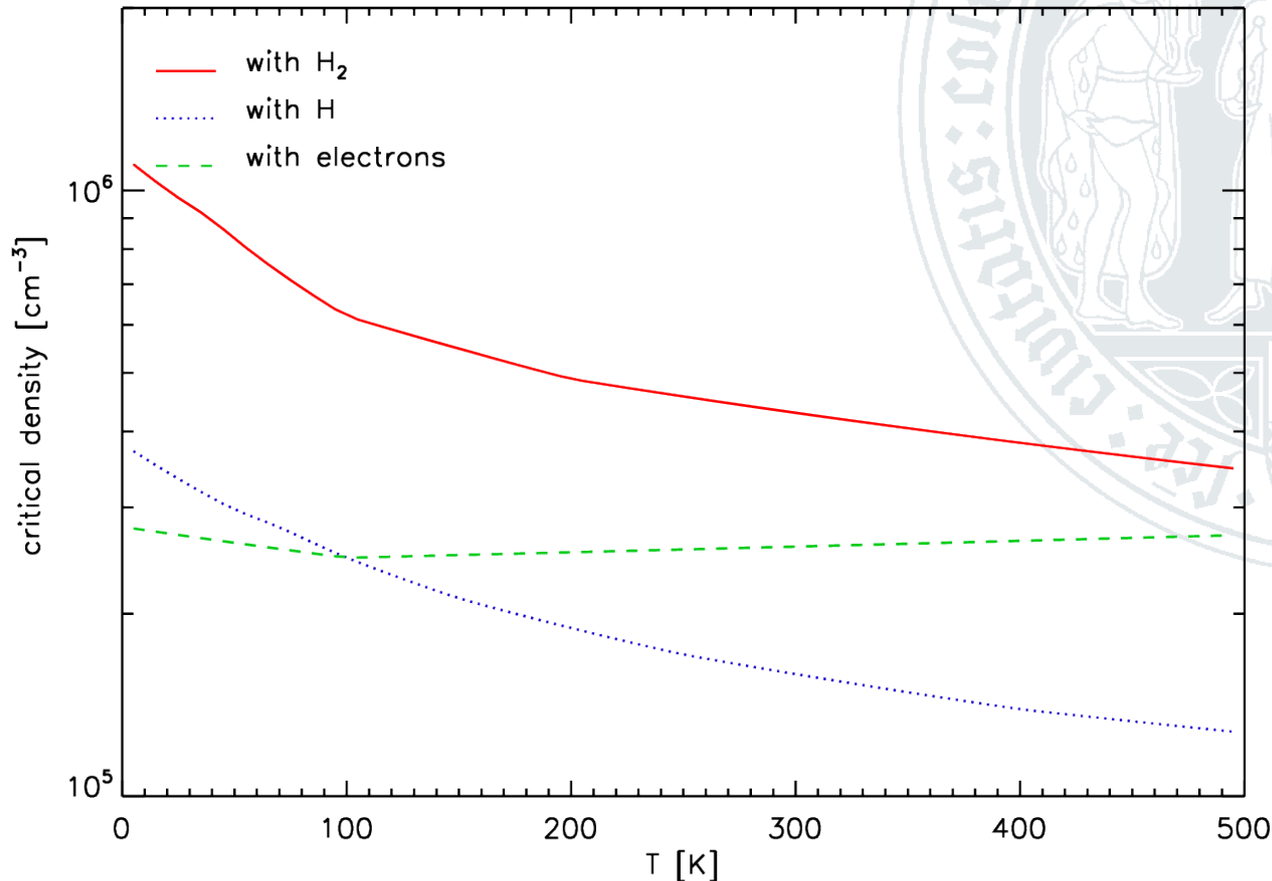
- Critical densities for different relevant collision partners



- Electrons only play a role if more than the carbon is ionized ($X[\text{C}/\text{H}] = 1.3 \cdot 10^{-4}$), i.e. in HII regions

[OI] excitation:

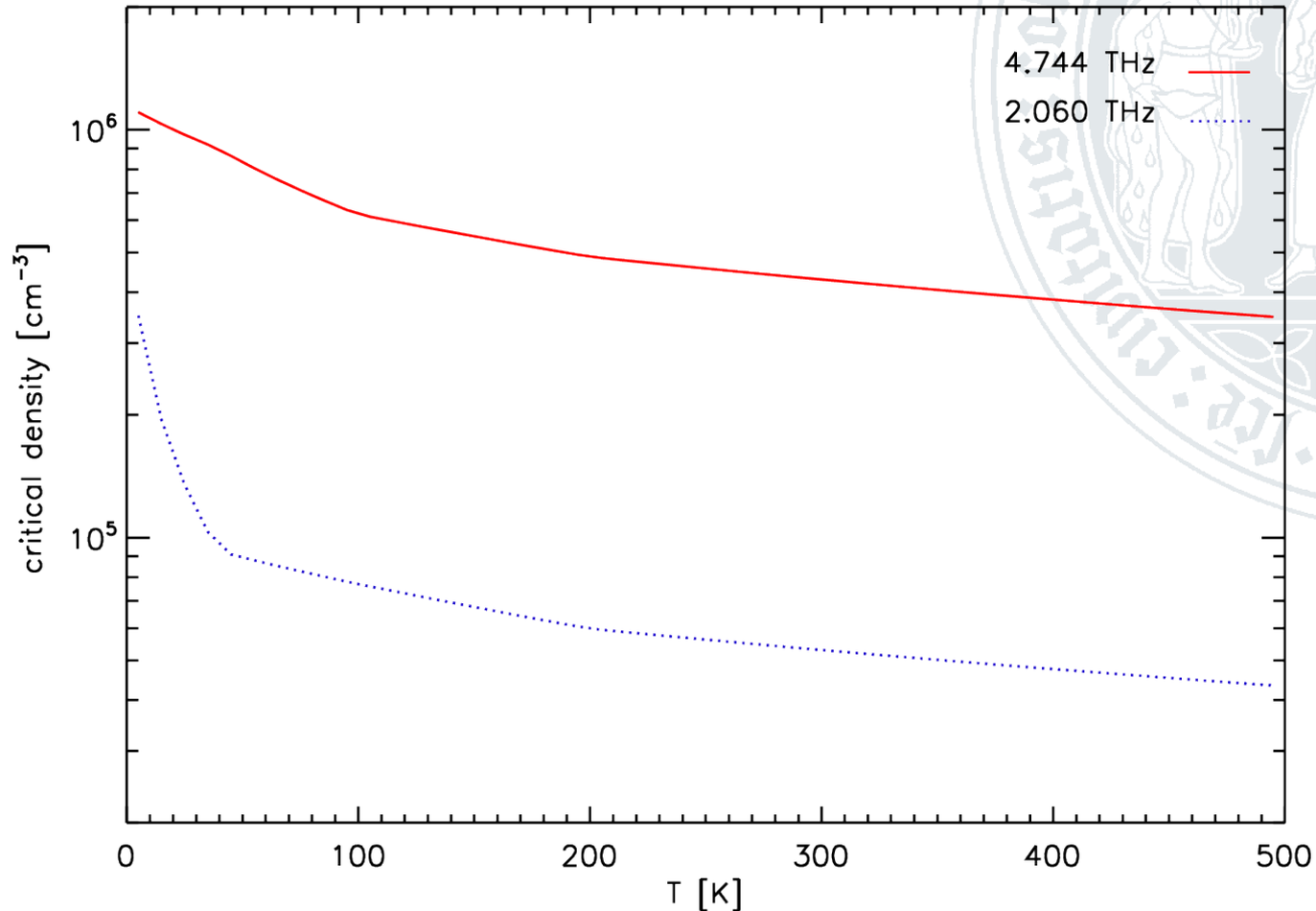
- 63 μm ground-state transition



- H-excitation is most efficient, but regions with high fraction of atomic H always have low densities (or volumes) so that there O⁺ is never in LTE
- High densities associated with H₂ – factor 3 less efficient

[OI] excitation:

- Full 3-level system - critical densities for two transitions:



- Critical densities are temperature-dependent!
- Non-trivial behavior for n-level systems (n > 2)!

n_i from balance equations

- Emission coefficient: $\epsilon_\nu(\vec{n}) = h\nu_{ij}n_iA_{ij}\frac{\phi_\nu(\vec{n})}{4\pi}$
(spontaneous emission)
 - Absorption coefficient: $\kappa_\nu(\vec{n}) = h\nu_{ij}(n_jB_{ji} - n_iB_{ij})\frac{\phi_\nu(\vec{n})}{c}$
(absorption and induced emission)
 - With line profile $\phi_\nu(\vec{n}) = \int \delta\left(\nu - \nu_{ij}\left[1 + \frac{\vec{v}\vec{n}}{c}\right]\right) p(\vec{v}) d^3\vec{v}$
$$\approx \frac{1}{\sqrt{2\pi}\sigma} \frac{c}{\nu_{ij}} \exp\left(-\frac{c^2}{2\sigma^2} \left[\frac{\nu - \nu_{ij}}{\nu_{ij}}\right]^2\right)$$
- for Maxwellian velocity distribution
- normalized so that omission of $\phi_\nu(\vec{n})$ gives line-integrated quantities

Alternative expression

- $\epsilon_\nu(\vec{n}) = \kappa_\nu(\vec{n}) \times B_{\nu_{ij}}(T_{ex,ij})$
 - $B_{\nu_{ij}}(T_{ex,ij})$ - Planck function (blackbody emissivity)

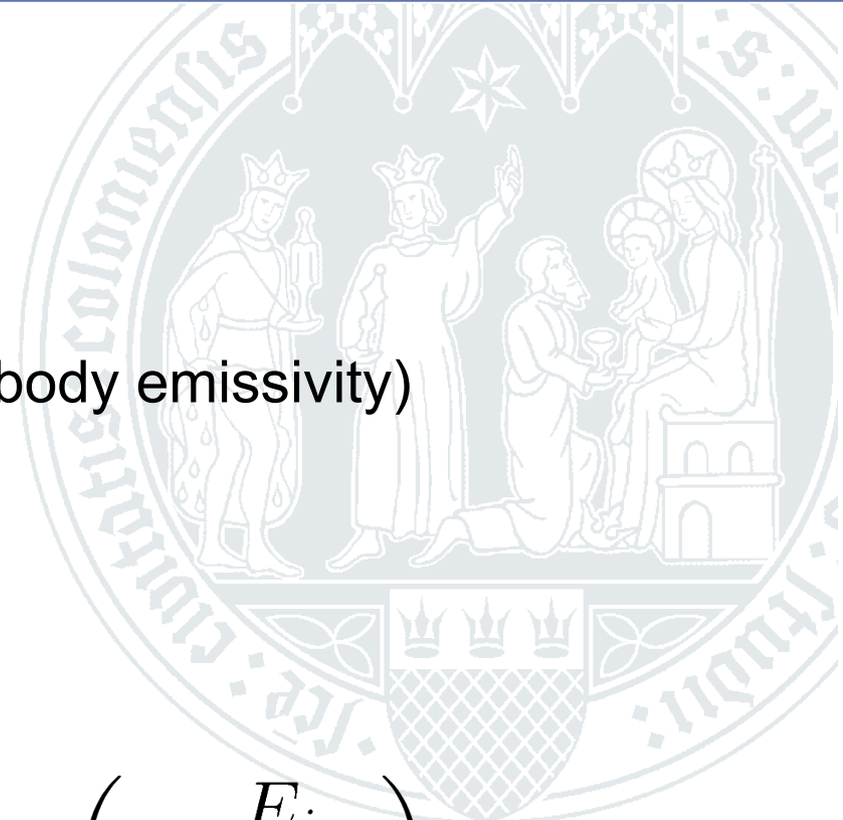
Practical computation:

- LTE

- Boltzmann distribution of level populations n_i :

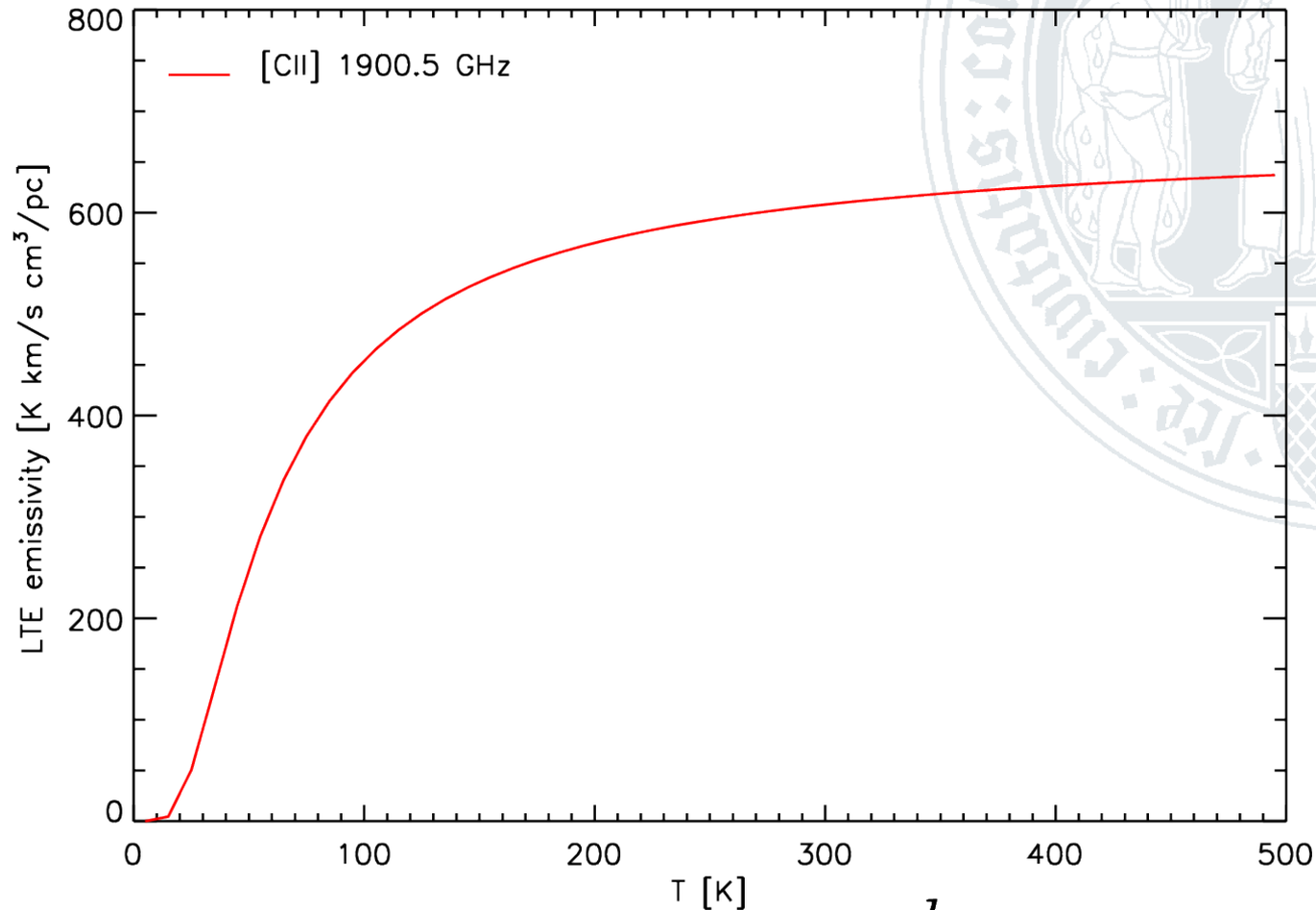
$$n_i \propto g_i \times \exp\left(-\frac{E_i}{kT_{kin}}\right)$$

- $T_{ex,ij} = T_{kin}$ i.e. $B_{\nu_{ij}}(T_{ex,ij}) = B_{\nu_{ij}}(T_{kin})$



LTE emissivity:

- Most simple: [CII] as two-level system



- Constant emissivity at temperatures above $\frac{h\nu_{ij}}{k} = 91.2\text{K}$

[CII] 1.9THz:

- Analytic description for emissivity

$$\int \epsilon dv = \frac{hc^3 A}{8\pi k\nu^2} \times N_{C^+} \frac{g_u \exp(-\Delta E/kT_{\text{ex}})}{g_l + g_u \exp(-\Delta E/kT_{\text{ex}})}$$

$$\approx 1011 \frac{\text{K kms}^{-1}}{\text{cm}^{-3} \text{ pc}} \times N_{C^+} \frac{2 \exp(-91.2\text{K}/T_{\text{ex}})}{1 + 2 \exp(-91.2\text{K}/T_{\text{ex}})}$$

and absorption/optical depth

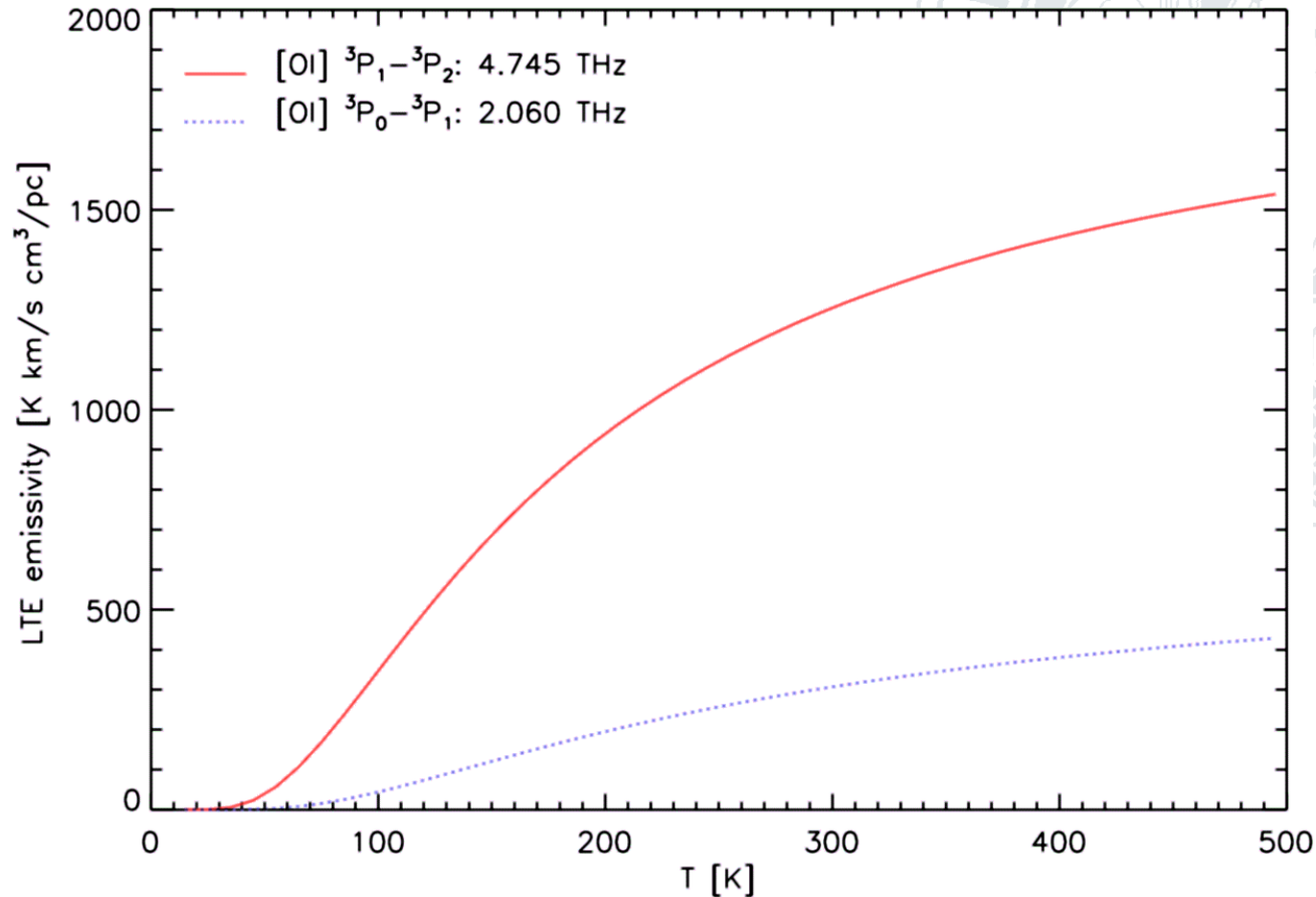
$$\int \tau dv = 7.15 \times 10^{-18} \frac{\text{kms}^{-1}}{\text{cm}^{-2}} \times N_{C^+} \frac{1 - \exp(-91.2 \text{ K}/T_{\text{ex}})}{1 + 2 \exp(-91.2 \text{ K}/T_{\text{ex}})}$$

$$\approx 7.15 \times 10^{-18} \frac{\text{kms}^{-1}}{\text{cm}^{-2}} \times N_{C^+} \frac{32.9 \text{ K}}{T_{\text{ex}}}$$

→ With $X(\text{C}^+/\text{H}) = 1.3 \cdot 10^{-4}$, [CII] turns optically thick for $N_{\text{H}} \sim 2 \cdot 10^{21} \text{ cm}^{-2}$, i.e. $A_{\text{V}} \sim 1$

LTE emissivity:

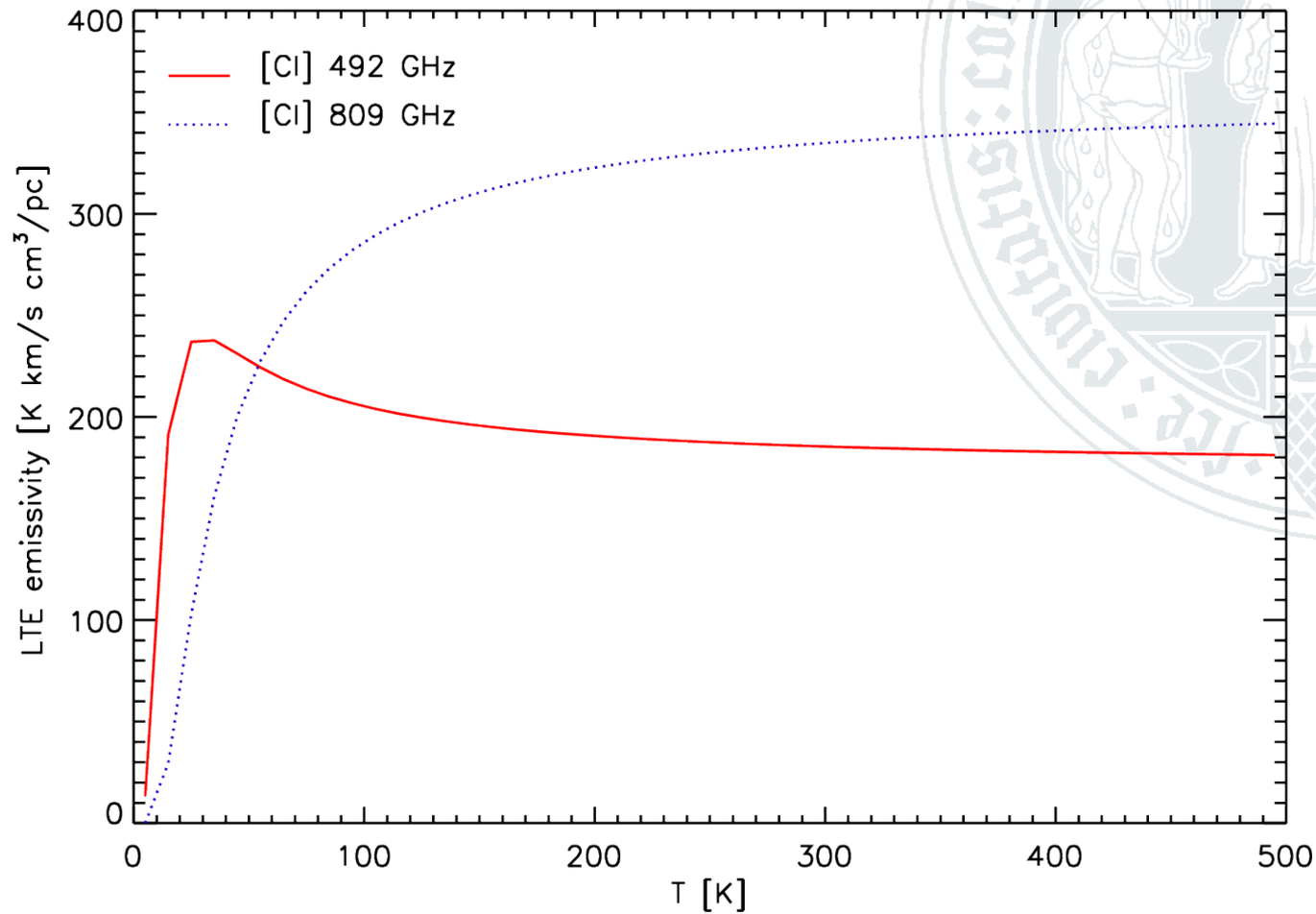
- [OI] as three-level system



- Transition to constant emissivity shifted by factor 2 due to higher level energies

LTE emissivity:

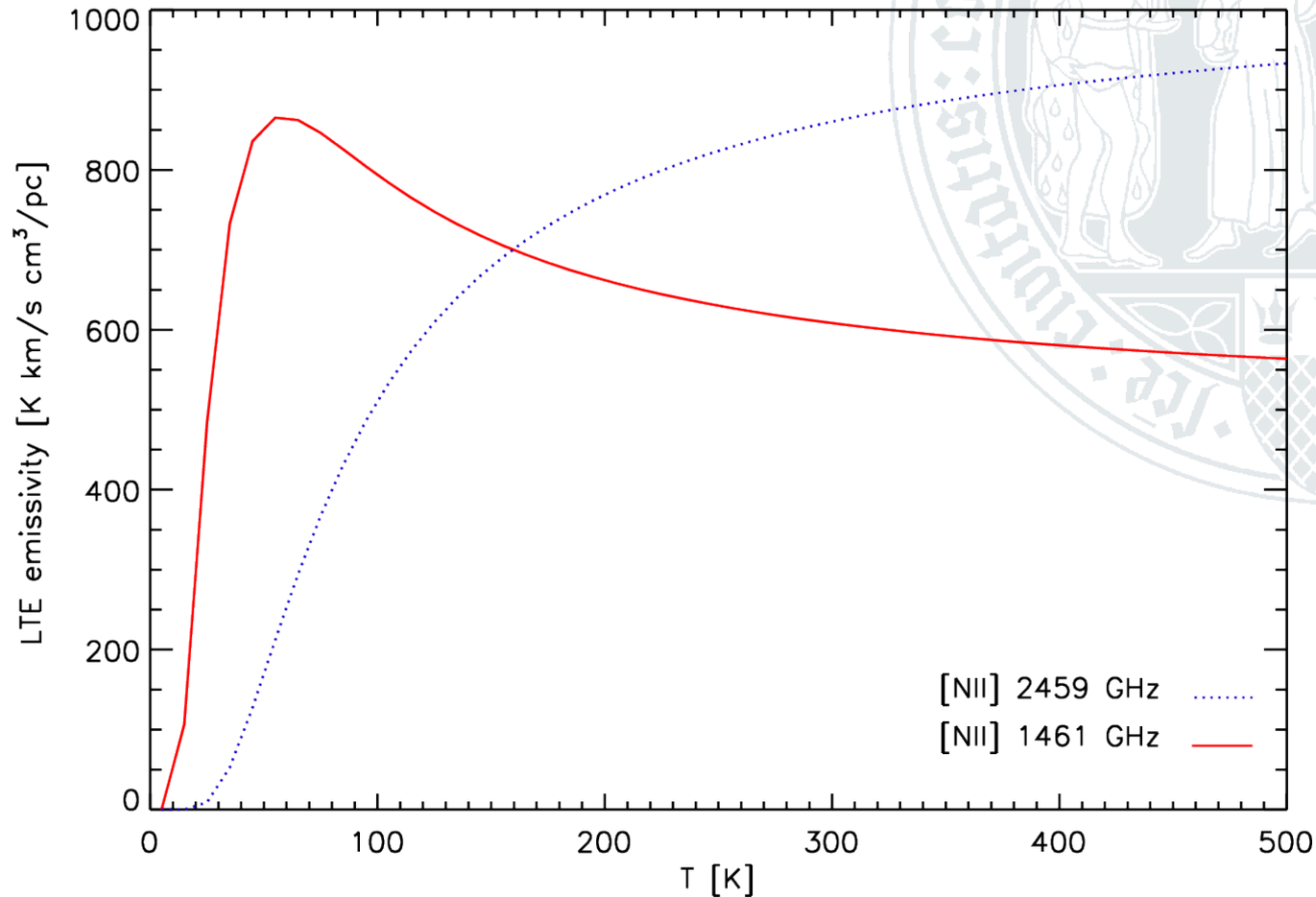
- [CI] as three-level system



- “Overshooting” in ground-state line for all $^3P_{0,1,2}$ systems

LTE emissivity:

- [NII] as equivalent system



- [OIII] fully equivalent (not shown here)

Next-step approximation:

- **Negligible radiative excitation:** $B_{ij}u_{ij} \ll A_{ij}, C_{ij}$

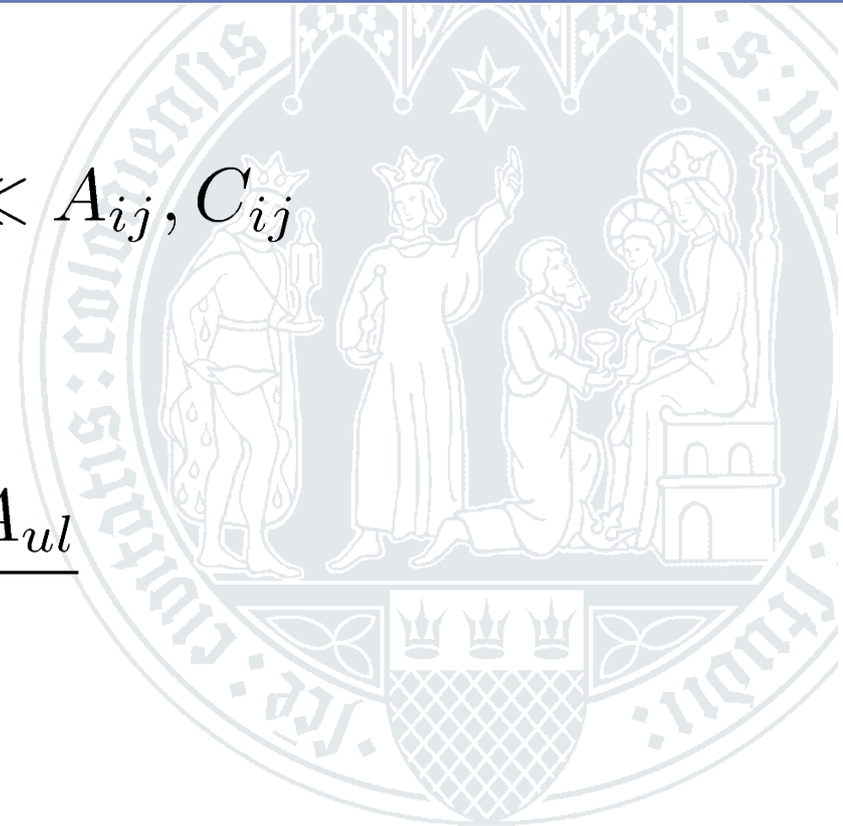
- Implies optically thin geometry

- Analytic solutions for 2-level system:

$$n_u = \frac{g_u n_l}{g_l} \exp\left(-\frac{h\nu_{ul}}{kT_{kin}}\right) \frac{C_{ul} + A_{ul}}{C_{ul}}$$

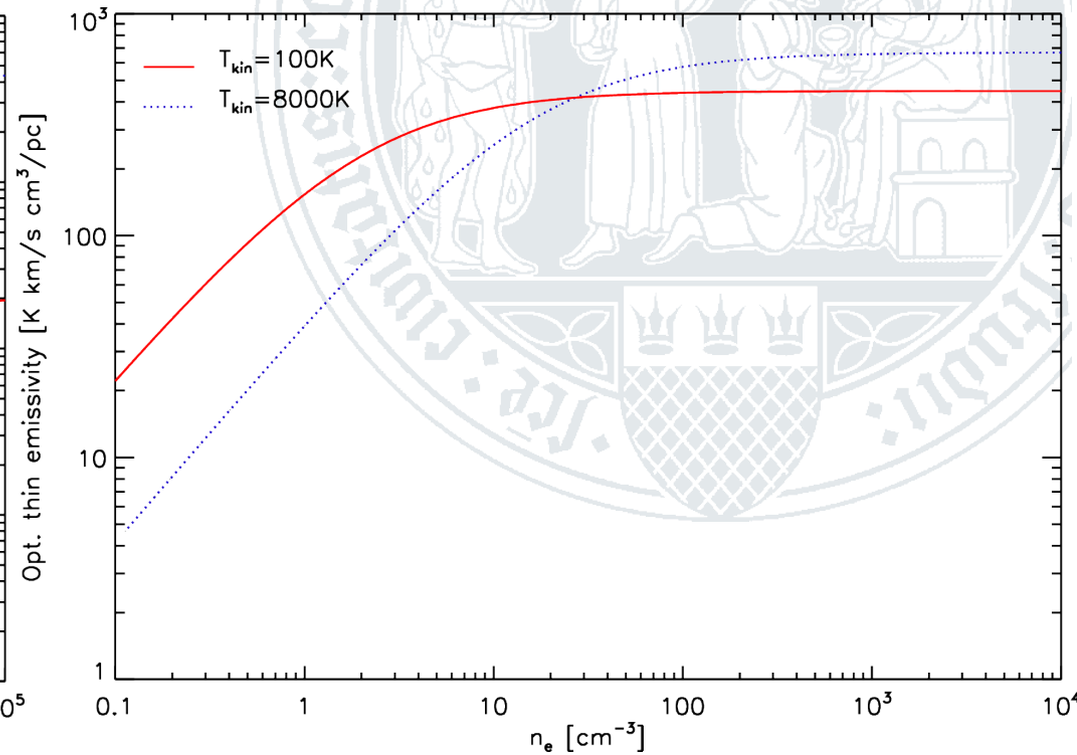
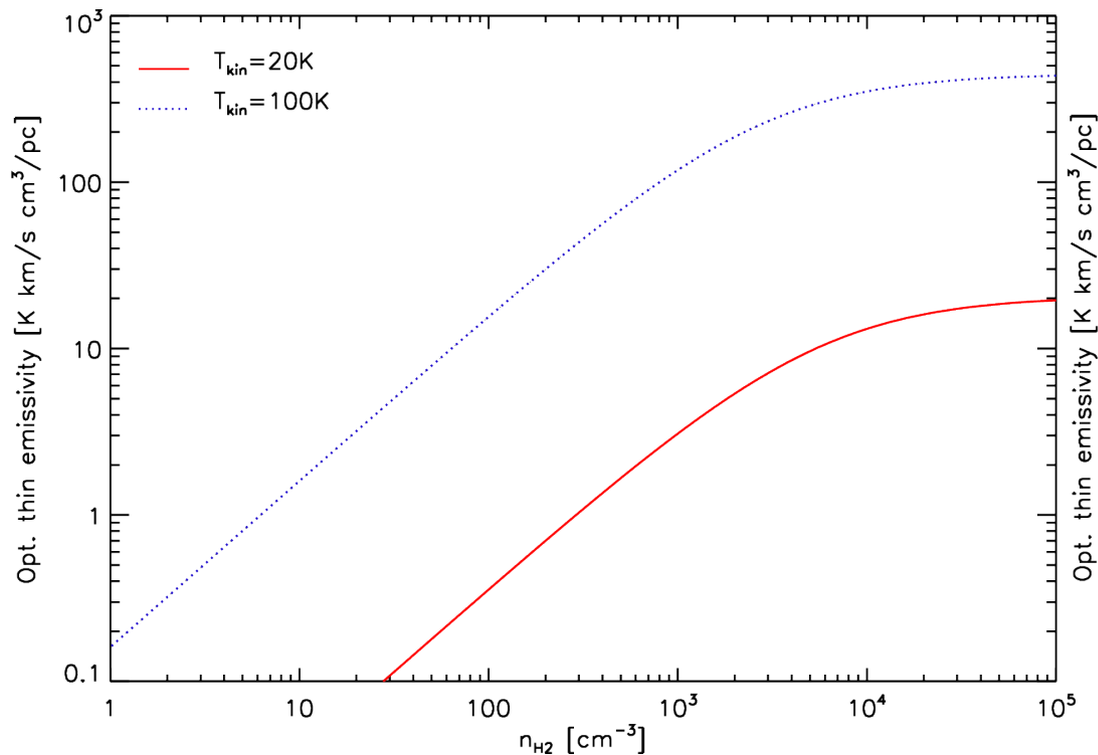
- and 3-level system:

$$n_2 = \frac{g_2 n_0}{g_0} \frac{C_{12}C_{01} + C_{12}C_{10} + C_{02}C_{12}}{C_{21}C_{10} + C_{20}C_{10} + C_{21}C_{12}} + \frac{C_{20}A_{20}}{A_{10}(C_{21} + C_{20}) + A_{21}C_{10} + A_{10}A_{21}}$$



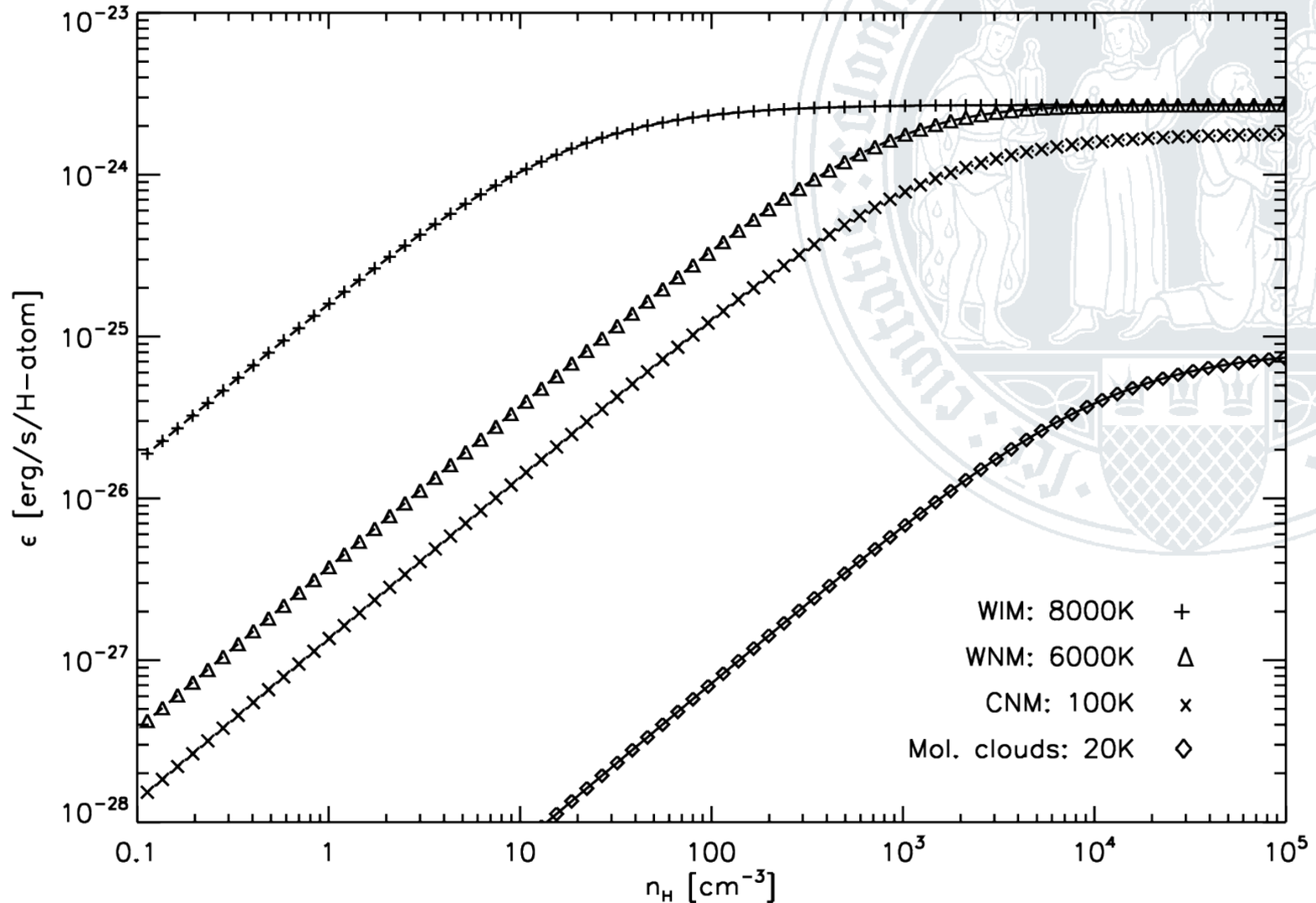
[CII] emissivity:

- For collisions with H₂ (left) and e⁻ (right)



- Transition from subthermal excitation at low densities to LTE at high densities
- At high densities only T_{kin} counts
- Critical density for electron-collisions lower by factor 1000

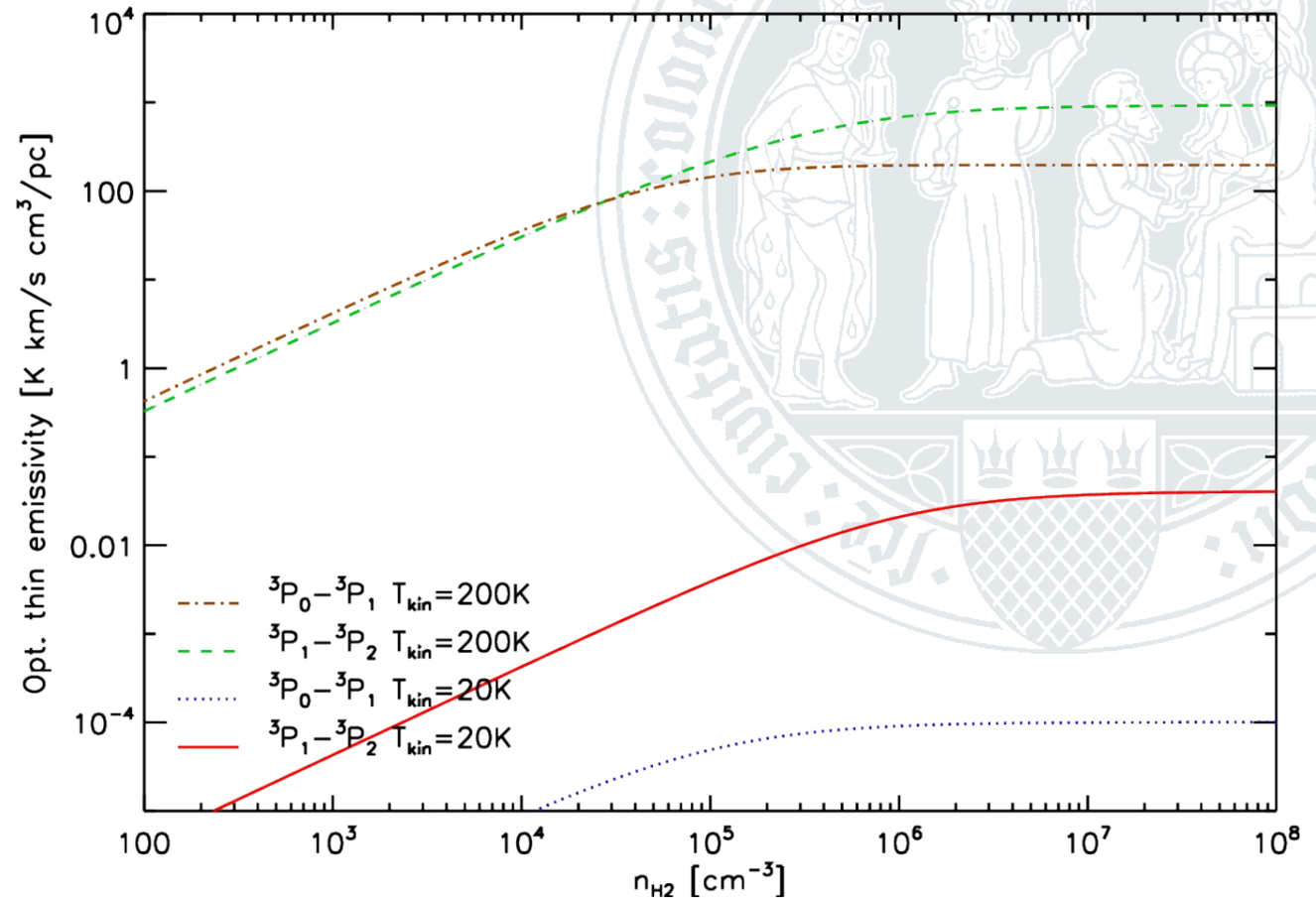
[CII] emissivity:



- Combination of higher emissivity above 91.2K and lower critical density for excitation through electron-collisions

[OI] emissivity:

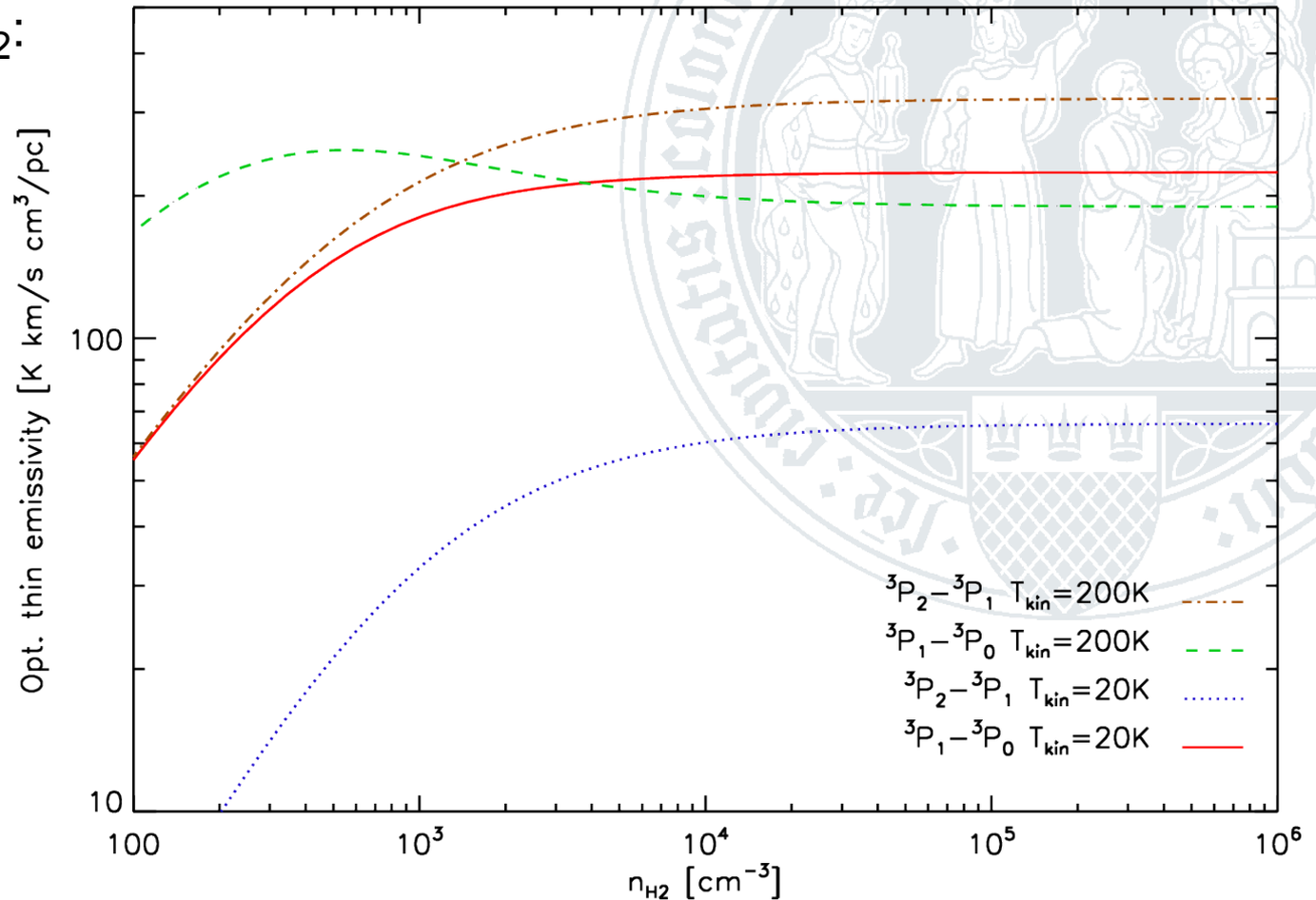
- Excitation through H₂



- Very simple: $\epsilon_\nu \propto n_{H_2}$ for subthermal excitation, monotonic transition to LTE
- High densities needed for thermal excitation
- Significant emission only for temperatures above 150K

[CI] emissivity:

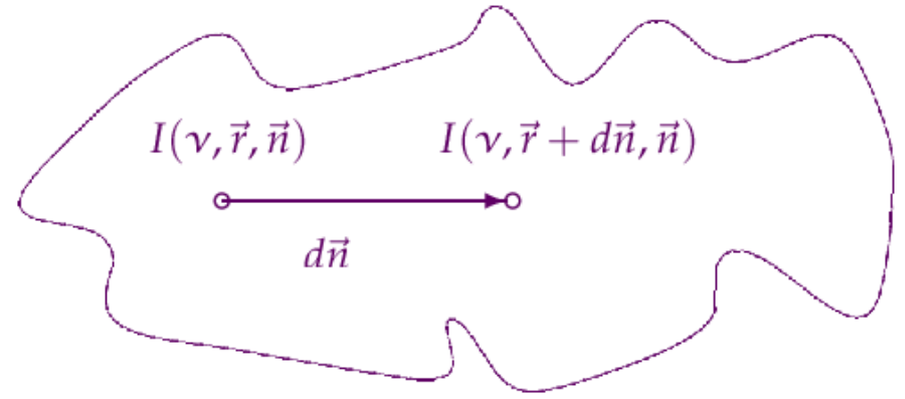
- Excitation through H₂:



- Partially non-monotonic behaviour for ground-state transition
- Much lower densities and temperatures needed for excitation
- Qualitatively same for [NII]

General case

- To derive physical parameters, the full radiative transfer problem needs to be solved



$$\vec{n} \cdot \text{grad} I_{ul} = -\kappa_{ul} I_{ul} + \epsilon_{ul}$$

$$\kappa_{ul} = \frac{h\nu_{ul}}{c} [n_l B_{lu} - n_u B_{ul}] \Phi(\nu)$$

$$\epsilon_{ul} = \frac{h\nu_{ul}}{c} n_u A_{ul} \Phi(\nu)$$

$$n_i \left(\sum_{j<i} A_{ij} + \sum_{j\neq i} B_{ij} u_{ij} + \sum_{j\neq i} C_{ij} \right) = \sum_{j>i} A_{ji} n_j + \sum_{j\neq i} B_{ji} u_{ij} n_j + \sum_{j\neq i} C_{ji} n_j$$

$$u_{ul} = \frac{1}{c} \int_{4\pi} d\vec{n} \int_0^\infty d\nu \Phi(\nu) I_{ul}$$

- Practical way out: **Local approximation – escape probability**

Probability that a line-photon escapes from the cloud β :

- Same probability that a background photon enters the cloud
- Consequence:
 - Radiation field given by “trapped” radiation field and external field:

$$u = (1 - \beta)B_\nu(T_{ex}) + \beta B_\nu(T_{bg})$$

- “Cooling of the system” not through all spontaneously emitted photons, but only through those photons also escaping from the cloud

- Modified Einstein-A coefficient: $A_{ul} \times \beta$

- Analytic solution for 2-level system:

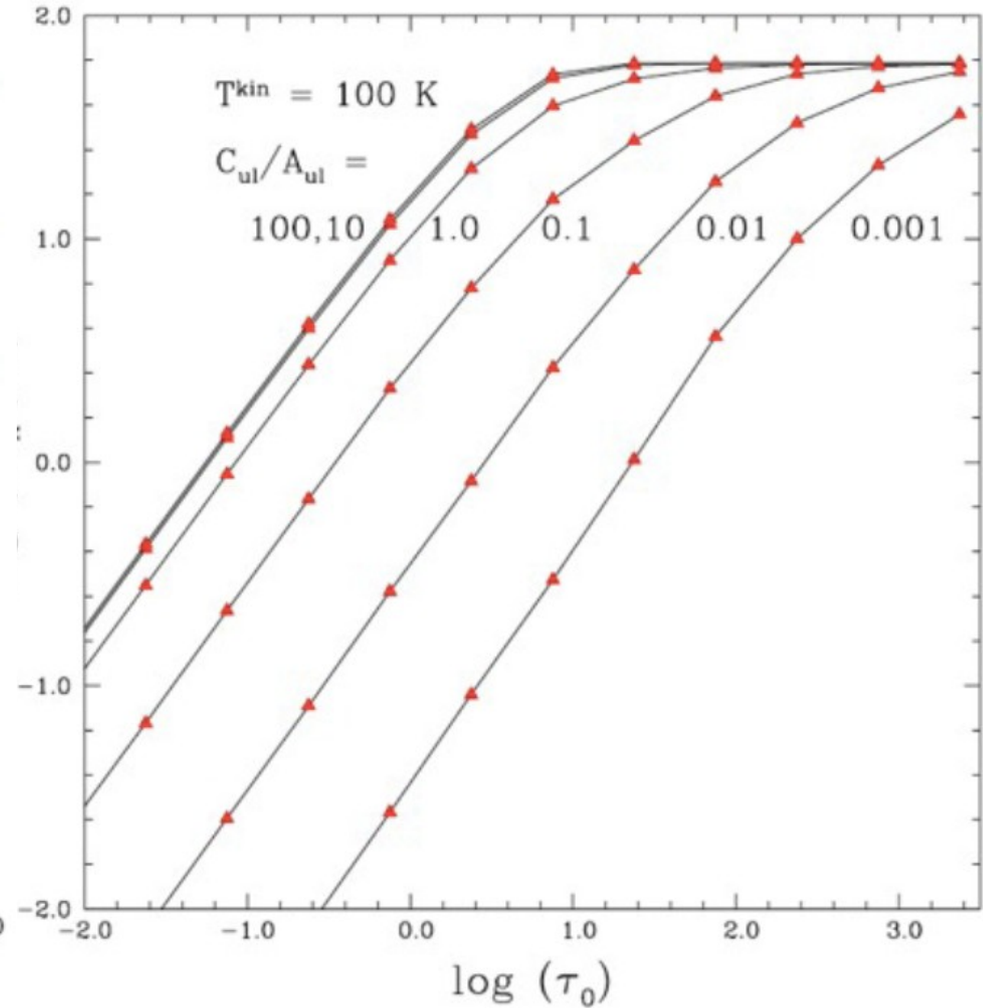
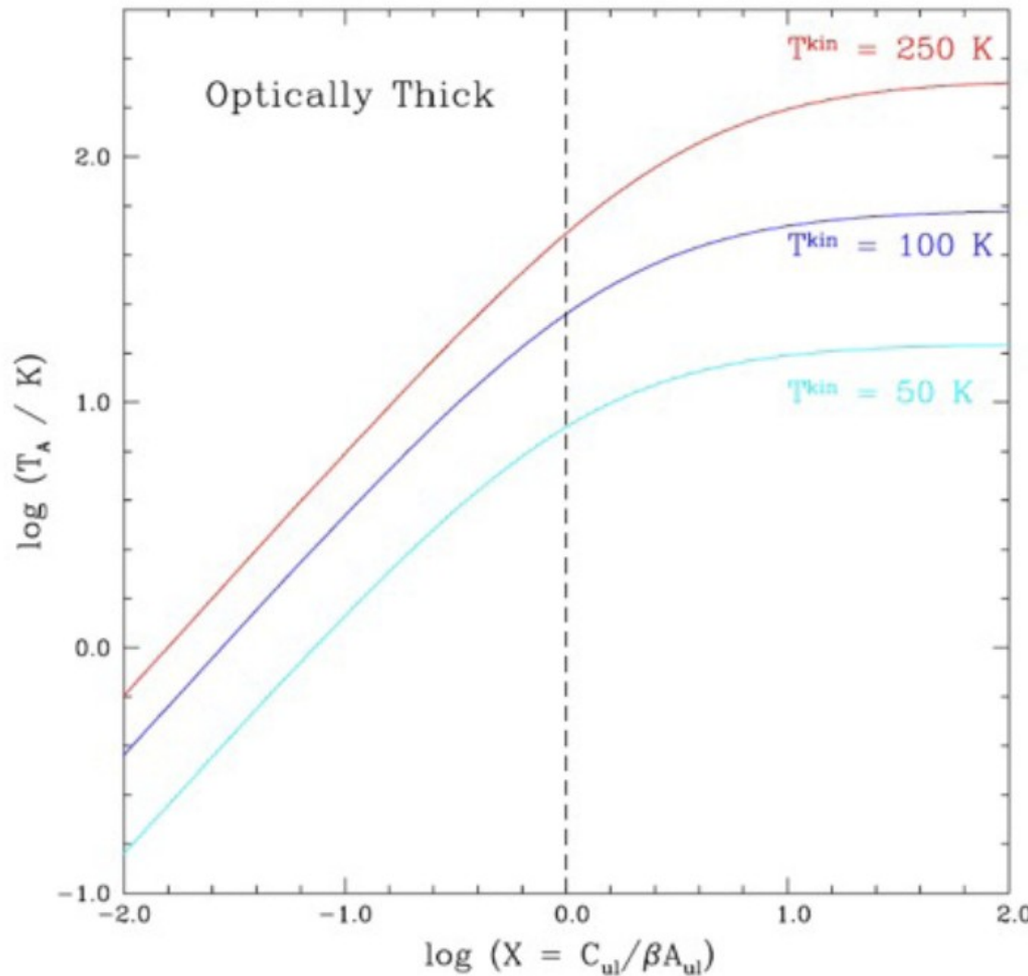
$$n_u = \frac{n_l g_u}{g_l} \frac{\beta A_{ul} G + C_{ul} \exp\left(-\frac{h\nu_{ul}}{kT_{kin}}\right)}{\beta A_{ul} (1 + G) + C_{ul}}$$

- with $G = \frac{1}{\exp\left(\frac{h\nu_{ul}}{kT_{bg}}\right) - 1}$

Goldsmith et al. (2013)



Result

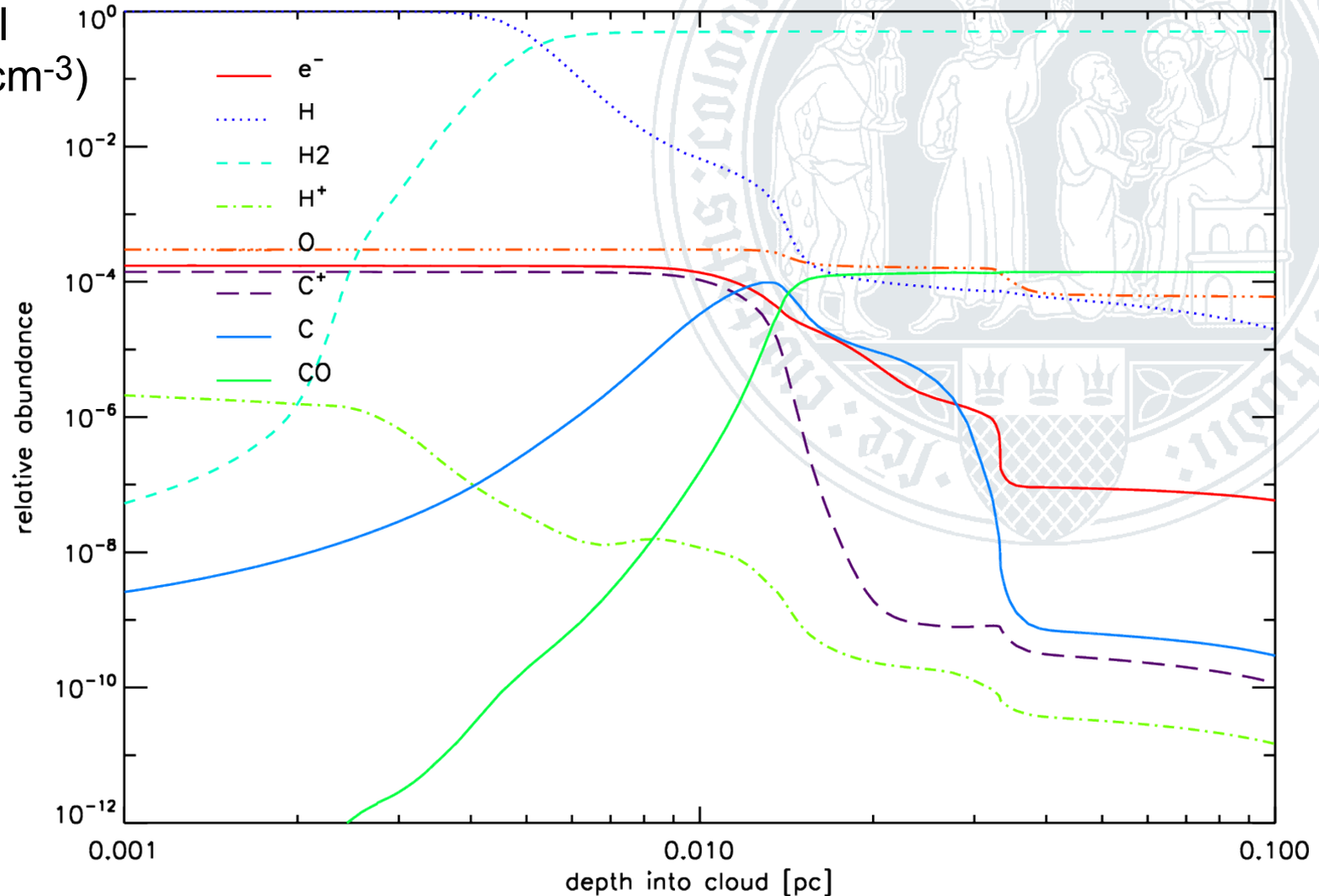


- Transition to LTE shifted to lower densities
- Emissivity remains proportional to column density in subthermal range

Goldsmith et al. (2013)

Combination with model for chemical abundance

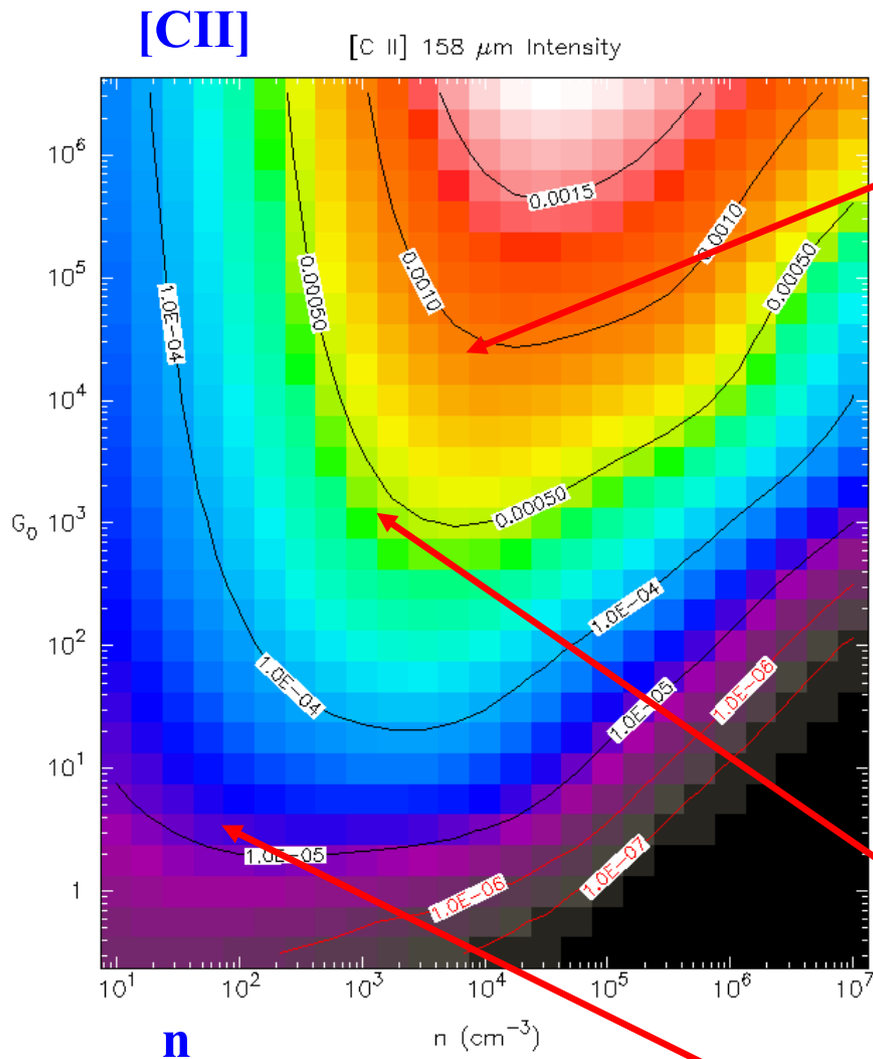
- e.g. PDR model
($\chi=10^4$, $n=10^5 \text{ cm}^{-3}$)



- Convolution of abundance profile with density and temperature for given chemical model needed.



PDR emission:



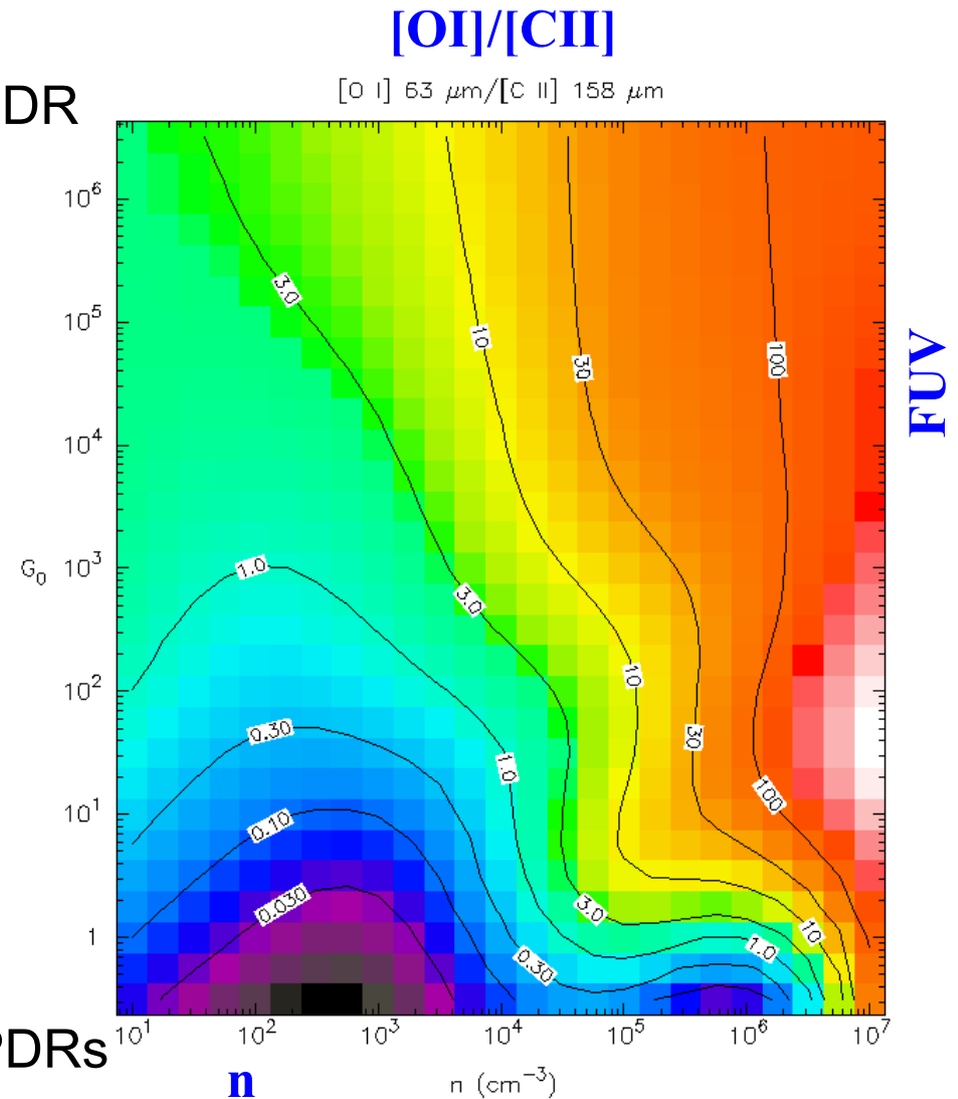
Orion PDR

FUV

Classic PDRs

Diffuse Gas

→ see talk by M. Röllig



FUV

Kaufman et al. 1999

Fine structure transitions are simple!

- Just a few numbers to describe them
- **But always think of the dominant collision partners**

	OIII	NII	CII	OI	CI
Line frequencies [GHz]	3393.0 5785.9	1461.3 2459.4	1900.5	4745.8 2060.0	492.2 809.3
Formation energy [eV]	35.1eV (IP)	14.5eV (IP)	11.2eV (IP)	11.1eV (CO diss.)	11.1eV (CO diss.)
Main collision partners	e ⁻	e ⁻	e ⁻ H H ₂	H ₂	H ₂
Typical critical densities [cm ⁻³]	500 4000	200 100	10 3000 5000	5 10 ⁵ 8 10 ⁴	1000 1500
High-temperature LTE emissivities [K km/s cm ³ pc ⁻¹]		550 940	699	1990 670	182 345