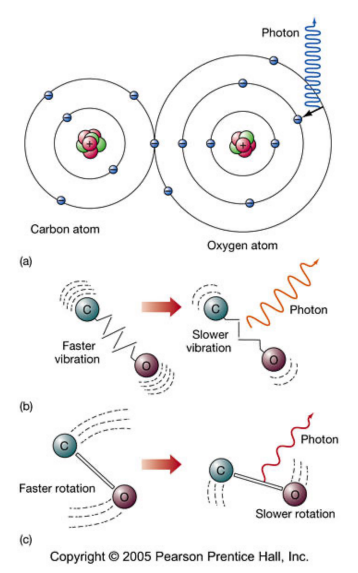
# Astrochemistry (1)

Planets and Astrobiology (2020-2021) G. Vladilo

# Molecular spectra

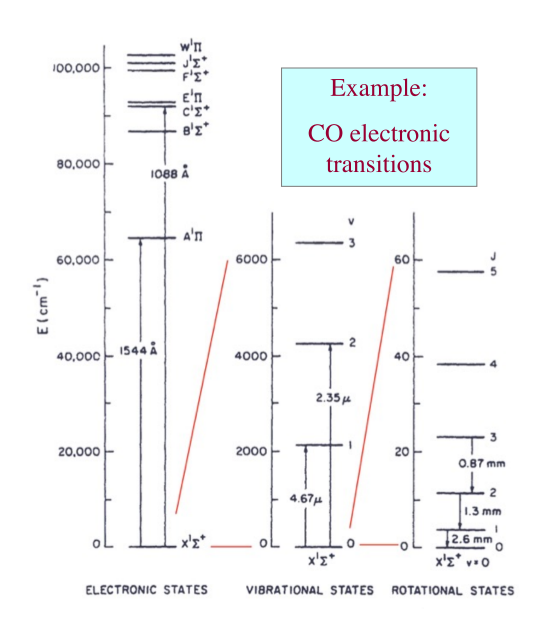
- Molecular spectra are much more complex than atomic spectra
- Molecular lines result from the combination of different types of transitions
  - Electronic
  - Vibrational
  - Rotational

Example in figure: CO molecule



#### Electronic transitions in molecules

- Electronic transitions in molecules are equivalent to atomic transitions
  - Typical energies are in the order of some eV
     They generally lie in the optical/UV spectral range
  - When observed at high spectral resolution, they can be solved in vibrational sub-levels which, in turn, can be solved in rotational sub-levels



## Quantization of vibrational and rotational motions

- Motions of atomic nuclei with respect the molecule's center of mass
- Main physical quantities that describe vibrational and rotational motions:
  - Total angular momentum
     The symbol *J* is used for diatomic molecules
     For polyatomic molecules the treatment is more complex
  - Reduced mass For diatomic molecules  $\mu = m_1 m_2 / (m_1 + m_2)$

#### Vibrational transitions

- Between energy levels that result from the quantization of vibrational modes
  - They are mostly "stretching" modes
    (variations of interatomic distances)
    Stretching is the only possible mode for diatomic molecules
    Complex molecules also have "bending" and "deformation" modes
- The quantum number v indicates the vibrational state of the system (for v=0 the vibrational energy is not null)

$$E^{vib} = \hbar \omega_e (v + \frac{1}{2})$$

$$v = 0, 1, 2, ...$$

As v varies in a given molecule, the vibrational energy levels are equidistant For different molecules the separation of the vibrational levels is a function of the reduced mass  $\mu$ 

For instance, H<sub>2</sub>, HD e D<sub>2</sub> produce different vibrational spectra

### Vibrational transitions

- The typical energies of vibrational modes lie in the range 0.1 e 0.3 eV
  - The corresponding wavelength  $\lambda$  lies in the near IR

CO: 
$$v = 1 - 0$$
 band  $\lambda = 4.67 \ \mu m \iff 2140 \ cm^{-1}$   
 $v = 2 - 0$  band  $\lambda = 2.35 \ \mu m \iff 4250 \ cm^{-1}$   
H<sub>2</sub>:  $v = 1 - 0$  band  $\lambda = 2.40 \ \mu m \iff 4150 \ cm^{-1}$ 

The energy of the transitions scales as  $\lambda^{-1}$  and can be expressed in cm<sup>-1</sup>

Conversion factor:  $1 \text{ cm}^{-1} = 1.24 \text{ x } 10^{-4} \text{ eV}$ 

- Every chemical group has a characteristic vibrational energy
  - Examples: C-H stretch, C $\equiv$ H stretch, CH<sub>2</sub> angle bending

<u>Different</u> complex molecules that have in common a chemical group will show the <u>same</u> vibrational transitions characteristic of that group

Problem of identification of complex molecules

### Rotational transitions

- Result from the quantization of the molecular rotational energies
  - The rotation can take place around the main axis of inertia
  - For complex molecules it could be a form of internal rotation
- Rotational transitions have energies in the order of  $\sim 10^{-3} \text{ eV}$ 
  - The wavelength  $\lambda$  generally lies in the millimetric or sub-millimetric spectral bands
- Rotational states are identified with the quantum number *J*

Example: CO

CO 
$$J=1-0$$
  $\nu=115$  GHz  $\Leftrightarrow \lambda=2.6$  mm  $J=2-1$   $\nu=230$  GHz  $\Leftrightarrow \lambda=1.3$  mm  $J=3-2$   $\nu=345$  GHz  $\Leftrightarrow \lambda=0.87$  mm

### Roto-vibrational transitions

- Vibrational transitions can be decomposed in rotational levels, the combination of the two giving rise to a roto-vibrational band
- Selection rules:
  - There are no restrictions on the variation of the vibrational state,  $\Delta v$
  - Variations of angular momentum are constrained by the selection rules  $\Delta J = 0, \pm 1$

Depending on the value  $\Delta J$  the bands are called as follows:

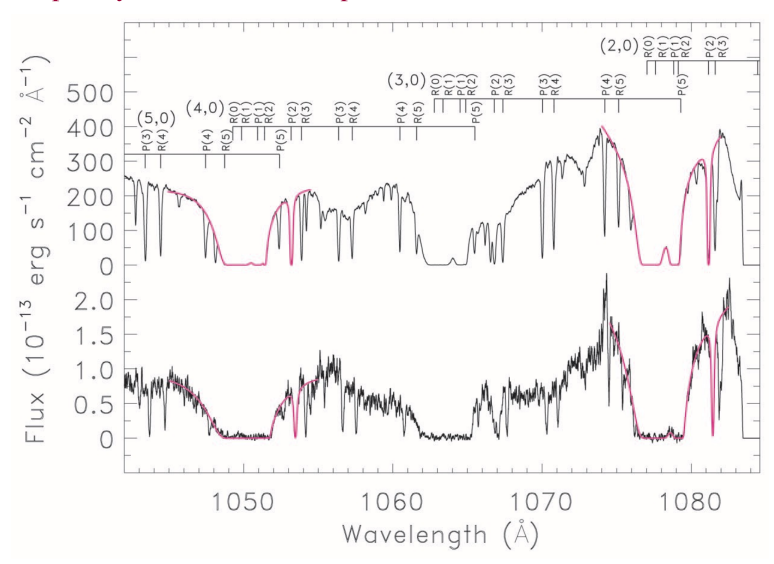
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\Delta J = -1 "P branch"

\Delta J = 0 "Q branch" (but J=0 \rightarrow J=0 is forbidden)

\Delta J = +1 "R branch"
```

## H<sub>2</sub> absorption lines

• Observations with high resolution spectroscopy show the extreme complexity of the molecular spectrum



# H<sub>2</sub> emission

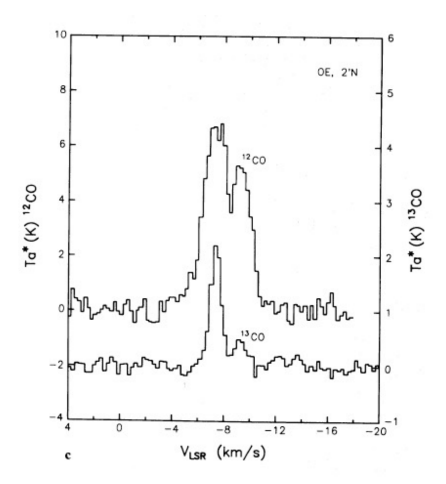
- Molecular hydrogen is a symmetric molecule
  - The electric dipole moment is zero

Rotational transitions are forbidden, even though electronic transitions are allowed

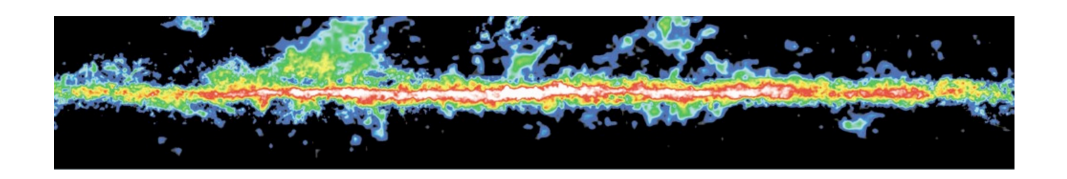
- Quadrupole transitions are possible, but very weak
  - Their energies are much higher than the typical rotational energies
  - Can only be observed in relatively warm regions
     Thanks to the abundance of H<sub>2</sub> some of these transitions have been observed in the mid IR
- Due to the lack of H<sub>2</sub> emissions it is hard to map the distribution of molecular hydrogen in galaxies
  - UV absorptions require (rare) bright background sources and are not suited to map the molecular gas, especially in dust-rich regions, where the background sources are obscured

## Molecular emission spectra in the millimetric band

- CO emission lines
  - J = 1-0 at 2.6 mm
- Used as a tracer of H<sub>2</sub>
  - CO is relatively abundant and has rotational transitions (not a symmetric molecule)
  - the dipole moment is relatively small and the molecule can be easily excited also in gas with relatively low density
- If the emission is too strong one can use less abundant istotopes, such as <sup>13</sup>CO, to obtain optically thin emissions



# Maps of the Galactic distribution of CO Tracer of molecular gas in the Galaxy



#### Chemical bonds of interstellar molecules

- Atoms in interstellar molecules are held together by covalent bonds
  - Superposition of the atomic orbitals
  - Sharing of the eletrons in external shells
- Typical energies of covalent bonds
  - $\sim 100 \text{ kcal/mol} \Rightarrow \sim 4 \text{ eV}$
- In the harsh conditions of the ISM the molecules can be easily dissociated
  - Kinetic temperatures in excess of ~10⁴ K would dissociate molecules by collisions
  - Photons with energies up to 13.6 eV can penetrate HI regions and photodissociate molecules
  - Interstellar molecules can survive in cold regions protected by the interstellar radiation field

#### Interstellar chemical reactions

#### Formation of bonds

- Radiative associationBetween neutrals and ions
- Reactions on the surface of dust grains

Also between neutrals example: H<sub>2</sub> formation

#### Destruction of bonds

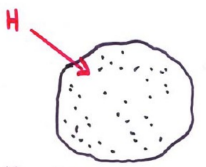
- Photo-dissociation
- Dissociative recombination

$$X^{+} + Y \rightarrow XY^{+} + h\nu$$
$$X + Y:g \rightarrow XY + g$$

$$XY + h\nu \rightarrow X + Y$$
  
 $XY^+ + e \rightarrow X + Y$ 

# H<sub>2</sub> formation on dust grains

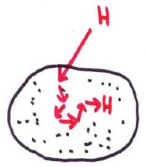
Credits: van Dishoeck



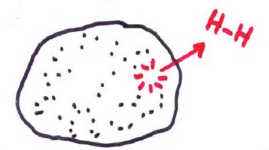
1. H collides with grain



2. It explores grain until either 1 (encounter with another II)
I immobilized at enhanced binding site



.3+4. second H atom collides with grain: explores surface and encounters first H atom



5. Hz formation on surface Hz ejected from surface

### Interstellar molecules

- About two hundreds gas-phase molecular species have been detected so far
- Besides simple molecules with a few atoms, also complex molecules with a relatively large number of atoms have been detected

http://www.astro.uni-koeln.de/cdms/molecules

Different types of molecules are observed in different types of interstellar or circumstellar regions

Some of them are only observed in dense molecular clouds

Symmetric molecules are harder to detect: they could be more abundant than what observed

 TABLE 1
 Interstellar and circumstellar molecules as compiled by Al Wootten (see text)

					Number of Atoms					
2	3	4	5	6	7	8	9	10	11	13
$\overline{H_2}$	C <sub>3</sub>	c-C <sub>3</sub> H	C <sub>5</sub>	C <sub>5</sub> H	C <sub>6</sub> H	CH <sub>3</sub> C <sub>3</sub> N	CH <sub>3</sub> C <sub>4</sub> H	CH <sub>3</sub> C <sub>5</sub> N?	HC <sub>9</sub> N	HC <sub>11</sub> N
A1F	$C_2H$	1-C <sub>3</sub> H	$C_4H$	$1-H_2C_4$	CH <sub>2</sub> CHCN	HCOOCH3	CH <sub>3</sub> CH <sub>2</sub> CN	(CH <sub>3</sub> ) <sub>2</sub> CO		
A1C1	$C_2O$	$C_3N$	C <sub>4</sub> Si	$C_2H_4$	$CH_3C_2H$	CH <sub>3</sub> COOH?	$(CH_3)_2O$	NH <sub>2</sub> CH <sub>2</sub> COOH?		
$C_2$	$C_2S$	C <sub>3</sub> O	$1-C_3H_2$	CH <sub>3</sub> CN	HC <sub>5</sub> N	$C_7H$	CH <sub>3</sub> CH <sub>2</sub> OH			
CH	$CH_2$	C <sub>3</sub> S	$c-C_3H_2$	CH <sub>3</sub> NC	HCOCH <sub>3</sub>	$H_2C_6$	HC <sub>7</sub> N			
CH <sup>+</sup>	HCN	$C_2H_2$	CH <sub>2</sub> CN	CH <sub>3</sub> OH	NH <sub>2</sub> CH <sub>3</sub>		C <sub>8</sub> H			
CN	HCO	$CH_2D+?$	$CH_4$	CH <sub>3</sub> SH	$c-C_2H_4O$					
CO	HCO+	HCCN	$HC_3N$	HC <sub>3</sub> NH <sup>+</sup>						
CO+	HCS+	HCNH+	HC <sub>2</sub> NC	HC <sub>2</sub> CHO						
CP	HOC+	HNCO	HCOOH	NH <sub>2</sub> CHO	Interstellar molecules					
CSi	$H_2O$	HNCS	H <sub>2</sub> CHN	C <sub>5</sub> N						
HC1	$H_2S$	HOCO+	$H_2C_2O$		liste	ed accord	ling to th	e number c	of atoi	ms
KC1	HNC	H <sub>2</sub> CO	H <sub>2</sub> NCN			Ehronfre	and & C	harnley (200	10)	
NH	HNO	$H_2CN$	$HNC_3$			Ememi		narmey (200	<i>(</i> 0)	
NO	MgCN	H <sub>2</sub> CS	SiH <sub>4</sub>							
NS	MgNC	$H_3O^+$	$H_2COH^+$							
NaC1	$N_2H^+$	$NH_3$								
OH	$N_2O$	SiC <sub>3</sub>								
PN	NaCN		Λ 11 .			0 10000 10	umahan a	fatama	04000	
SO	ocs		All molecules with a large number of atoms are organic							
SO+	SO <sub>2</sub>		A tos	na with	1000,000	mia ahu	ndonoo	ro only dat	actad	in
SiN	c-SiC <sub>2</sub>		Atoms with low cosmic abundance are only detected in							
SiO	$CO_2$		sma	ll molec	cules					
SiS	$NH_2$		Siliu.		0.000					
CS	$H_3^+$									

Note that observations suggest the presence of large PAHs and fullerenes in the interstellar gas (Tielens et al 1999, Foing & Ehrenfreund 1997).

#### Small interstellar molecules

- Found in *diffuse* molecular clouds
  - molecular clouds with relatively low extinction
- Large molecules are absent in *diffuse* clouds because of:
  - physical conditions
     diffuse clouds are less
     protected from interstellar
     radiation field than denser
     molecular clouds
  - observational limitations

diffuse clouds have relatively low column densities and this fact makes hard to detect large molecules, characterized by a low abundance

Table 2 Molecules detected in diffuse molecular clouds

Weight	Species	Method	Target	N(X)/N <sub>H</sub>
2	H <sub>2</sub>	UV	ζOph	0.56
3	HD	UV	ζOph	4.5 (-7)
3	H <sub>3</sub> +	IR	ζPer	5.1 (-8)
13	СН	Optical	ζOph	1.5 (-9)
13	CH+	Optical	ζOph	2.4 (-8)
14	<sup>13</sup> CH <sup>+</sup>	Optical	ζOph	3.5 (-10)
15	NH	Optical	ζOph	6.2 (-10)
17	OH	UV	ζOph	3.3 (-8)
24	C <sub>2</sub>	Optical	ζOph	1.3 (-8)
25	C <sub>2</sub> H	mm abs.	BL Lac	1.8 (-8)
26	CN	Optical	ζOph	1.9 (-9)
27	HCN	mm abs.	BL Lac	2.6 (-9)
27	HNC	mm abs.	BL Lac	4.4 (-10)
28	N <sub>2</sub>	UV	HD 124314	3.1 (-8)
28	CO	UV	X Per	6.4 (-6)
29	HCO+	mm abs.	BL Lac	1.5 (-9)
29	HOC+	mm abs.	BL Lac	2.2 (-11)
29	<sup>13</sup> CO	UV	X Per	8.9 (-8)
29	C <sup>17</sup> O	UV	X Per	7.4 (-10):
30	C18O	UV	X Per	2.1 (-9):
30	H <sub>2</sub> CO	mm abs.	BL Lac	3.7 (-9)
36	C <sub>3</sub>	Optical	ζOph	1.1 (-9)
36	HCl	UV	ζOph	1.9 (-10)
38	C <sub>3</sub> H <sub>2</sub>	mm abs.	BL Lac	6.4 (-10)
44	CS	mm abs.	BL Lac	1.6 (-9)
64	SO <sub>2</sub>	mm abs.	BL Lac	≤8.2 (−10)

# Complex interstellar molecules

("complex" for interstellar standards, not for chemists)

- Complex interstellar molecules are hydrocarbons
- They are found in:
  - star-forming regions
  - circumstellar envelopes of evolved, late-type stars in the Asymptotic giant branch (AGB)
  - dense clouds in the direction of the Galactic center

Herbst & van Dishoeck (2009)

#### Examples of interstellar hydrocarbons

Table 1 Complex organic interstellar molecules (≥ 6 atoms)

Species	Name	Source	
Hydrocarbons		0	
C <sub>2</sub> H <sub>4</sub>	Ethene	circ	
HC <sub>4</sub> H	Butadiyne	circ	
H <sub>2</sub> C <sub>4</sub>	Butatrienylidene	circ, cc, lc	
C5H	Pentadiynyl	circ, cc	
CH <sub>3</sub> C <sub>2</sub> H	Propyne	cc, lc	
C <sub>6</sub> H	Hexatriynyl	circ, cc, lc	
C <sub>6</sub> H <sup>-</sup>	Hexatriynyl ion	circ, cc, lc	
$H_2C_6$	Hexapentaenylidene	circ, cc, lc	
HC <sub>6</sub> H	Triacetylene	circ	
C <sub>7</sub> H	Heptatriynyl	circ, cc	
CH <sub>3</sub> C <sub>4</sub> H	Methyldiacetylene	cc	
CH₃CHCH₂	Propylene	cc	
C <sub>8</sub> H	Octatetraynyl	circ, cc	
$C_8H^-$	Octatetraynyl ion	circ, cc	
CH₃C <sub>6</sub> H	Methyltriacetylene	cc	
C <sub>6</sub> H <sub>6</sub>	Benzene	circ	

Abbreviations: circ, circumstellar envelope around evolved star/protoplanetary nebula; cc, cold cloud core; hc, hot core/corino; lc, lukewarm corino; gc, galactic center cloud; of, outflow. Not all of these molecules fulfill the strict criteria for identification listed in Section 3.3.

# Saturation of interstellar organic molecules

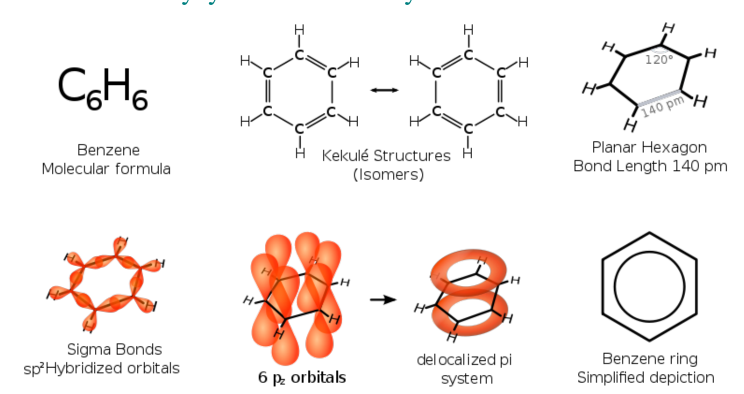
- Saturated hydrocarbons
  - The chain of carbon atoms are held by single bonds
  - The remaining carbon bonds are saturated with hydrogen atoms
- Interstellar organic molecules are usually not saturated
  - Example of <u>saturated</u> molecule <u>not detected</u> in the ISM Cyclohexane, C<sub>6</sub>H<sub>12</sub>

Cyclohexane

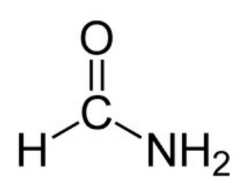
 Example of <u>unsaturated</u> molecule <u>detected</u> in the ISM Benzene, C<sub>6</sub>H<sub>6</sub>

#### Benzene

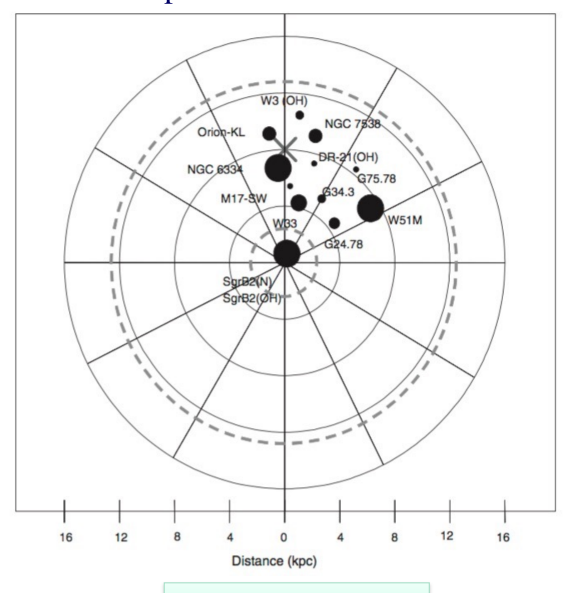
- Aromatic ring
  - Stable electronic structure that results from the superposition of atomic orbitals; the electrons are delocalized and shared by all atoms
- Plays an important role in astrochemistry
  - Starting point for the formation of complex aromatic compounds
     PAHs=Polycyclic Aromatic Hydrocarbon



# Formamide An interstellar molecule of prebiotic interest



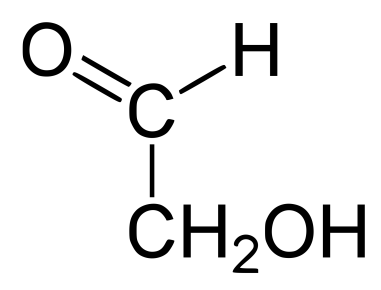
Detected multiple rotational transitions in the sub-millimetric spectral range in molecular clouds at different locations in the Galaxy



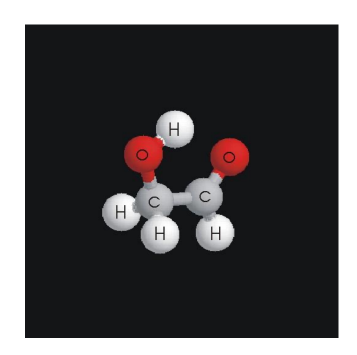
Adande et al. (2013)

## Complex organic molecules in the interstellar medim

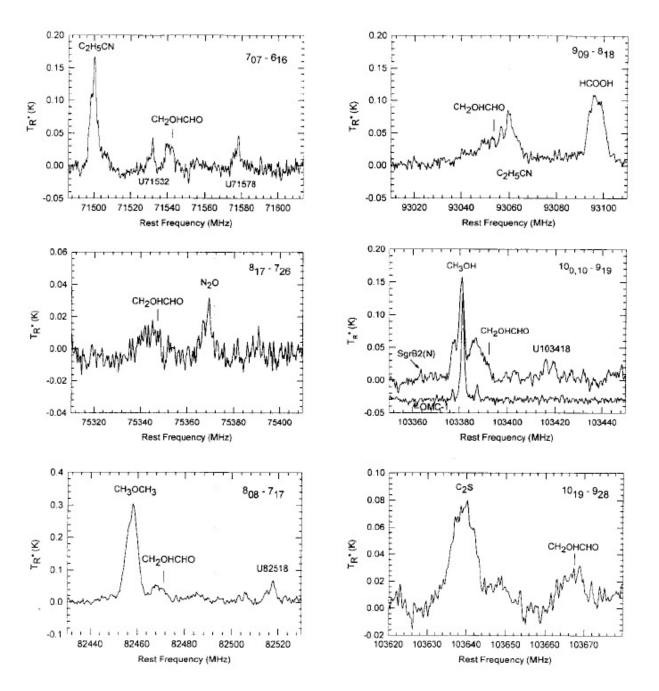
- Glycolaldehyde (C<sub>2</sub>H<sub>4</sub>O<sub>2</sub>)
  - Simplest sugar
  - First intermediate product of the reaction which starts with formaldehyde (H<sub>2</sub>CO) and leads to the formation of various sugars and finally of ribose, one of the DNA building blocks



Generic formula for sugars  $C_n(H_2O)_n$ 

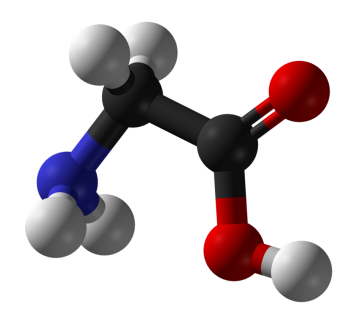


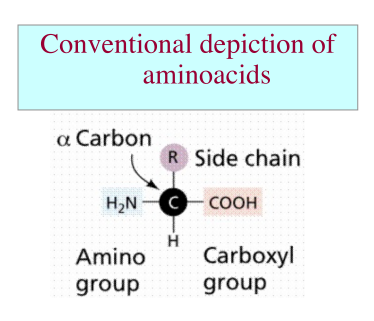
- Detection of glycolaldehyde (CH<sub>2</sub>OHCHO)
  - First detection of interstellar sugar
  - Detected in the millimetric band towards Sagittarius B2(N), a source in the direction of the Galactic center (Hollis et al. 2000)
  - Also observed by ALMA around a young, solar-type star



# Complex organic molecules in the interstellar space The case of glycine

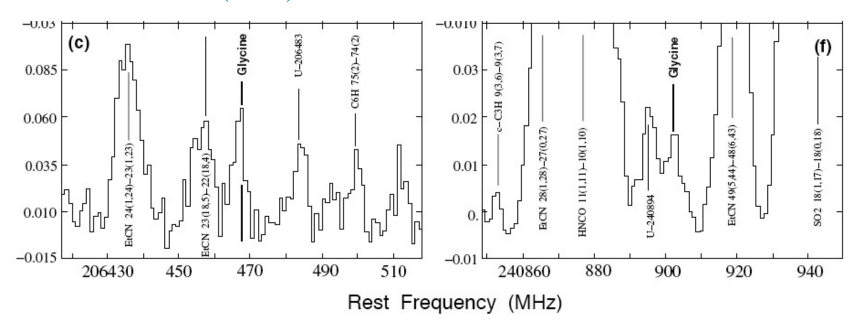
- Glycine is the simplest aminoacid found in biological proteins (NH<sub>2</sub>CH<sub>2</sub>COOH)
  - Its existence in the interstellar space would demonstrate the existence of chemical pathways potentially able to synthesise basic ingredients of life molecules in the interstellar space
  - The "lateral group" R is simply a hydrogen atom





## Tentative evidence for interstellar glycine

- Glycine (NH<sub>2</sub>CH<sub>2</sub>COOH)
  - Several emission lines attributed to interstellar glycine have been reported
     Kuan et al. (2003)



 The identification is not confirmed by a subsequent analysis performed by testing a larger number of lines expected for glycine

Snyder et al. (2005)

# Which is the maximum complexity of interstellar organic molecules in the gas phase?

As molecular complexity increases, the identification of the molecule tends to become uncertain

Gas-phase molecules with a high number of atoms might exist in the interstellar medium, but it is difficult to prove their existence