# THEORETICAL EXAM



Making science together!

2019-07-26





MINISTÈRE DE L'ÉDUCATION NATIONALE ET DE LA JEUNESSE

MINISTÈRE DE L'ENSEIGNEMENT SUPÉRIEUR, DE LA RECHERCHE ET DE L'INNOVATION

#### **General instructions**

- This theoretical exam booklet contains 61 pages.
- You may begin writing as soon as the **Start** command is given.
- You have 5 hours to complete the exam.
- All results and answers must be written clearly in pen in their respective designated areas on the exam papers. Answers written outside the answer boxes will **not** be graded.
- If you need scratch paper, use the backside of the exam sheets. Remember that nothing outside the designated areas will be graded.
- Use only the pen and calculator provided.
- The official English version of the exam booklet is available upon request and serves for clarification only.
- If you need to leave the exam room (i.e. to use the toilet or have a snack), wave the corresponding IChO card. An exam supervisor will come to accompany you.
- For multiple-choice questions: if you want to change your answer, fill the answer box completely and then make a new empty answer box next to it.
- The supervisor will give a warning when there is only 30 minutes left before the **Stop** command.
- You must stop working **immediately** when the **Stop** command is announced. Failure to stop writing by ½ minute or longer will lead to nullification of your theoretical exam.
- After the Stop command has been given, place your exam booklet back in your exam envelope, then wait at your seat. The exam supervisor will come to seal the envelope in front of you and collect it.

#### **GOOD LUCK!**

# **Table of Contents**

This theoretical exam is composed of 9 independent problems, as follows. Their relative weight is indicated in parenthesis.

Problem T1: Infinite well and butadiene	(6%)	p. 8
Problem T2: Hydrogen production by water-splitting	(7%)	p. 13
Problem T3: About silver chloride	(5%)	p. 19
Problem T4: From black powder to the discovery of iodine	(7%)	p. 24
Problem T5: Complexes for the formation of nanomachines	(8%)	p. 31
Problem T6: Characterization of a block-copolymer	(8%)	p. 40
Problem T7: Ring motion in a [2]catenane	(6%)	p. 48
Problem T8: Identification and synthesis of inositols	(6%)	p. 53
Problem T9: Synthesis of levobupivacaine	(7%)	p. 58

#### Physical constants and equations

In these tasks, we assume the activities of all aqueous species to be well approximated by their respective concentration in mol L<sup>-1</sup>. To further simplify formulas and expressions, the standard concentration  $c^{\circ} = 1 \text{ mol } L^{-1}$  is omitted.

Avogadro's constant:

Universal gas constant:

Standard pressure:

Atmospheric pressure:

Zero of the Celsius scale:

Faraday constant:

Watt:

Kilowatt hour:

Planck constant:

Speed of light in vacuum:

Elementary charge:

Electron-volt

Electrical power:

Power efficiency:

Planck-Einstein relation:

Ideal gas equation:

Gibbs free energy:

Reaction quotient Q for a reaction a A(aq) + b B(aq) = c C(aq) + d D(aq):

Henderson-Hasselbalch equation:

Nernst–Peterson equation:

where O is the reaction quotient of the reduction half-reaction

Beer-Lambert law:

Rate laws in integrated form:

- Zero order:
- First order:
- Second order:

Half-life for a first order process:

Number average molar mass  $M_n$ :

Mass average molar mass  $M_w$ :

Polydispersity index  $I_n$ :

$$N_{\rm A} = 6.022 \cdot 10^{23} \text{ mol}^{-1}$$
  
 $R = 8.314 \text{ J mol}^{-1} \text{ K}^{-1}$   
 $p^{\circ} = 1 \text{ bar} = 10^{5} \text{ Pa}$   
 $P_{\rm atm} = 1 \text{ atm} = 1.013 \text{ bar} = 1.013 \cdot 10^{5} \text{ Pa}$   
 $273.15 \text{ K}$ 

$$F = 9.6485 \cdot 10^4 \text{ C mol}^{-1}$$
  
1 W = 1 J s<sup>-1</sup>

1 kWh = 
$$3.6 \cdot 10^6$$
 J  
 $h = 6.6261 \cdot 10^{-34}$  J s

$$c = 2.998 \cdot 10^8 \,\mathrm{m \, s}^{-1}$$

$$e = 1.6022 \cdot 10^{-19} \text{ C}$$
  
1 eV = 1.6022 \cdot 10^{-19} J

$$P = \Delta E \times I$$

$$\eta = P_{\text{obtained}}/P_{\text{applied}}$$

$$E = hc/\lambda = h \nu$$

$$pV = nRT$$

$$G = H - TS$$

$$\Delta_{\rm r}G^{\circ} = -RT \ln K^{\circ}$$
  
$$\Delta_{\rm r}G^{\circ} = -n F E_{\rm cell}^{\circ}$$

$$\Delta_{\rm r}G = \Delta_{\rm r}G^{\circ} + RT \ln O$$

$$Q = \frac{[\mathbf{C}]^{c}[\mathbf{D}]^{d}}{[\mathbf{A}]^{a}[\mathbf{B}]^{b}}$$

$$[\mathbf{A}]^{-1}$$

$$pH = pK_a + \log \frac{[A^-]}{[AH]}$$

$$E = E^{o} - \frac{RT}{zF} \ln Q$$

at 
$$T = 298 \text{ K}, \frac{RT}{F} \ln 10 \approx 0.059 \text{ V}$$

$$A = \varepsilon lc$$

$$[\mathbf{A}] = [\mathbf{A}]_0 - kt$$

$$\ln[\mathbf{A}] = \ln[\mathbf{A}]_0 - kt$$

$$1/[\mathbf{A}] = 1/[\mathbf{A}]_0 + kt$$

$$\frac{\ln 2}{k}$$

$$M_{\rm n} = \frac{\sum_{\rm i} N_{\rm i} M_{\rm i}}{\sum_{\rm i} N_{\rm i}}$$

$$M_{\rm w} = \frac{\sum_{\rm i} N_{\rm i} M_{\rm i}^2}{\sum_{\rm i} N_{\rm i} M_{\rm i}}$$
$$I_{\rm p} = \frac{M_{\rm w}}{M_{\rm p}}$$

$$I_{\rm p} = \frac{M_{\rm w}}{M_{\rm n}}$$

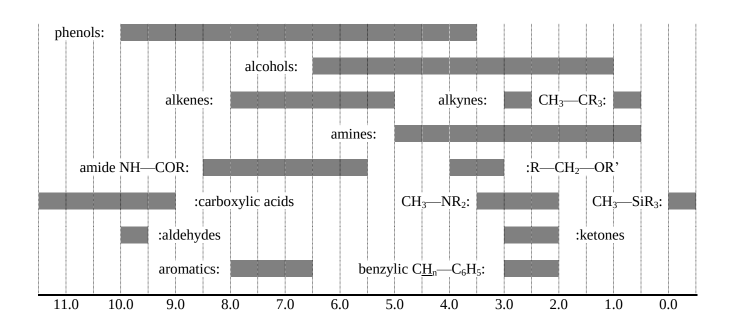
#### Periodic table

1																	18
1 <b>H</b> 1.008	2											13	14	15	16	17	2 He 4.003
3	4											5	6	7	8	9	10
Li	Be											В	С	N	0	F	Ne
6.94	9.01											10.81	12.01	14.01	16.00	19.00	20.18
11	12											13	14	15	16	17	18
Na	Mg	3	4	5	6	7	8	9	10	11	12	ΑI	Si	Р	S	CI	Ar
22.99	24.31											26.98	28.09	30.97	32.06	35.45	39.95
19	20	21	22	23	24	25	26	27	28	29	30	31	32	33	34	35	36
K	Ca	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr
39.10	40.08	44.96	47.87	50.94	52.00	54.94	55.85	58.93	58.69	63.55	65.38	69.72	72.63	74.92	78.97	79.90	83.80
37	38	39	40	41	42	43	44	45	46	47	48	49	50	51	52	53	54
Rb	Sr	Υ	Zr	Nb	Мо	Тс	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Te	ı	Xe
85.47	87.62	88.91	91.22	92.91	95.95	-	101.1	102.9	106.4	107.9	112.4	114.8	118.7	121.8	127.6	126.9	131.3
55	56		72	73	74	75	76	77	78	79	80	81	82	83	84	85	86
Cs	Ba	57-71	Hf	Та	W	Re	Os	Ir	Pt	Au	Hg	TI	Pb	Bi	Po	At	Rn
132.9	137.3		178.5	180.9	183.8	186.2	190.2	192.2	195.1	197.0	200.6	204.4	207.2	209.0	-	-	-
87	88	00	104	105	106	107	108	109	110	111	112	113	114	115	116	117	118
Fr	Ra	89- 103	Rf	Db	Sg	Bh	Hs	Mt	Ds	Rg	Cn	Nh	FI	Мс	Lv	Ts	Og
-	-		-	-	-	-	-			-	-	-	-	-	-	-	-

57	58	59	60	61	62	63	64	65	66	67	68	69	70	71
La	Ce	Pr	Nd	Pm	Sm	Eu	Gd	Tb	Dy	Но	Er	Tm	Yb	Lu
138.9	140.1	140.9	144.2	-	150.4	152.0	157.3	158.9	162.5	164.9	167.3	168.9	173.0	175.0
89	90	91	92	93	94	95	96	97	98	99	100	101	102	103
Ac	Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	Lr
-	232.0	231.0	238.0	-	-	-	-	-	-	-	-	-	-	-



<sup>1</sup>H NMR
Chemical shifts of hydrogen (in ppm / TMS)



### H-H coupling constants (in Hz)

Hydrogen type	$ J_{ m ab} $ (Hz)
$R_2CH_aH_b$	4-20
$R_2H_aC$ — $CR_2H_b$	2-12 if free rotation: 6-8 ax-ax (cyclohexane): 8-12 ax-eq or eq-eq (cyclohexane): 2-5
$R_2H_aC$ — $CR_2$ — $CR_2H_b$	if free rotation: < 0.1 otherwise (rigid): 1-8
RH <sub>a</sub> C=CRH <sub>b</sub>	cis: 7-12 trans: 12-18
$R_2C=CH_aH_b$	0.5-3
H <sub>a</sub> (CO)—CR <sub>2</sub> H <sub>b</sub>	1-3
RH <sub>a</sub> C=CR—CR <sub>2</sub> H <sub>b</sub>	0.5-2.5

eq = equatorial, ax = axial

## IR spectroscopy table

Vibrational mode	$\sigma$ (cm <sup>-1</sup> )	Intensity
alcohol O—H (stretching)	3600-3200	strong
carboxylic acid O—H (stretching)	3600-2500	strong
N—H (stretching)	3500-3350	strong
	3300	strong
≡C—H (stretching)	3100-3000	strong weak
=C—H (stretching)		*** - **
C—H (stretching)	2950-2840	weak
–(CO)—H (stretching)	2900-2800	weak
C≡N (stretching)	2250	strong
C≡C (stretching)	2260-2100	variable
aldehyde C=O (stretching)	1740-1720	strong
anhydride C=O (stretching)	1840-1800; 1780-1740	weak; strong
ester C=O (stretching)	1750-1720	, ,
,	1730-1720	strong
ketone C=O (stretching)		strong
amide C=O (stretching)	1700-1500	strong
alkene C=C (stretching)	1680-1600	weak
aromatic C=C (stretching)	1600-1400	weak
CH <sub>2</sub> (bending)	1480-1440	medium
CH <sub>3</sub> (bending)	1465-1440; 1390-1365	medium
C—O—C (stretching)	1250-1050	strong
C—OH (stretching)		strong
NO <sub>2</sub> (stretching)	1200-1020	strong
1102 (succining)	1600-1500; 1400-1300	strong

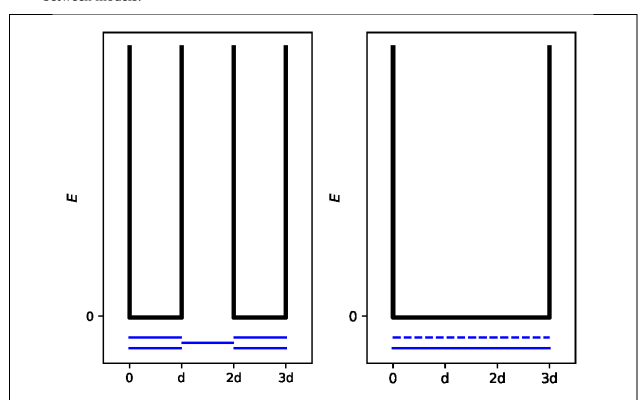
Problem	Question	1	2	3	4	5	6	7	8	9	10	11	Total
T1	Points	3	4	4	2	3	2	2	4.5	2.5	3	3	33
6%	Score												

#### Problem T1: Infinite well and butadiene

The buta-1,3-diene molecule is often written  $CH_2$ =CH-CH= $CH_2$ , with alternating single and double bonds. Nevertheless, its chemical reactivity is not consistent with this description and the  $\pi$  electrons are better described by a distribution along the three bonds:

This system can be modeled as a 1D box (*i.e.* infinite well) where the electrons are free. The energy of an electron in an infinite well of length L is:  $E_n = \frac{n^2 h^2}{8m_e L^2}$ , where n is a **non-zero** positive integer.

1. Two different models are studied. <u>Sketch</u> at least the three lowest-energy levels  $E_n$  <u>for each</u> <u>model</u> in the respective diagrams, showing how the relative energy levels differ within and between models.



**Model 1** (« **localized** »): The  $\pi$  electrons are localized on the extremal bonds and evolve into two separate infinite potential wells of length d.

Model 2 (« delocalized »): The  $\pi$  electrons are delocalized on the whole molecule and evolve into a single infinite potential well of length 3d.

2.	<u>Place</u> the $\pi$ electrons for model 1 in the previous diagrams and <u>express</u> the total energy of the $\pi$ system in model 1, as a function of $h$ , $m_{\rm e}$ and $d$ .
E(	1) =
3.	<u>Place</u> the $\pi$ electrons for model 2 in the previous diagrams and <u>express</u> the total energy of the $\pi$ system in model 2, as a function of $h$ , $m_{\rm e}$ and $d$ .
<i>E</i> (	(2) =
	e conjugation energy is the total energy of the actual $\pi$ system, minus the sum of the energies of ylene molecules involving the same number of electrons.
4.	<b>Express</b> the conjugation energy $\Delta E_c$ of butadiene, as a function of $h$ , $m_e$ and $d$ .
	7
$\Delta E$	$E_{\rm c} =$
Mo	odels 1 and 2 are too simplistic. A new model will be detailed in the following.
5.	<u>Draw</u> three other resonance structures of butadiene using Lewis notation.
	$H_2C$ $CH_2$

To take into account the size of carbon atoms, model 2 is now modified into model 3, as follows:

- The new length of the well is L and is located between the abscissa 0 and L;
- The carbon atoms are located at the abscissas L/8; 3L/8; 5L/8 and 7L/8.

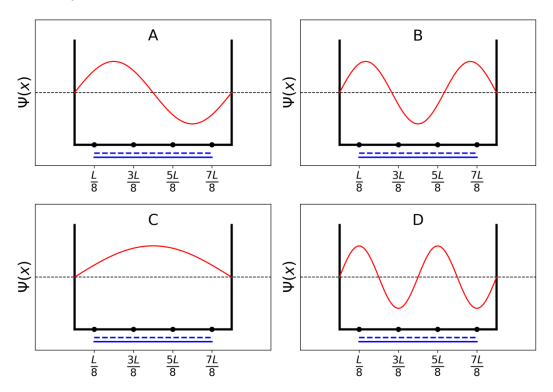
For each level n, the  $\pi$  wavefunction is:

$$\psi_{\rm n}(x) = \sqrt{\frac{2}{L}} \sin\left(\frac{n\pi x}{L}\right)$$

and the  $\pi$  electron density for a system with  $N \pi$  electrons is:

$$\rho(x) = 2 \sum_{i=1}^{N/2} |\psi_i(x)|^2$$

The four  $\pi$  wavefunctions, which correspond to the molecular orbitals of the  $\pi$  system, are depicted below (**arbitrary order**).



6. **Arrange** the energies of the four  $\pi$  wavefunctions ( $E_A$ ,  $E_B$ ,  $E_C$  and  $E_D$ ).

< < <

7. Give the labels (A, B, C or D) of the orbitals that are filled with electrons in butadiene.

8. Within model 3, **give** the values of the  $\pi$  wavefunctions  $\psi_n$  for occupied levels at positions 0, L/4 and L/2, for n = 1 and n = 2, as a function of L.

 $\psi_1(0) =$ 

 $\psi_1\left(\frac{L}{4}\right) =$ 

 $\psi_1\left(\frac{L}{2}\right) =$ 

 $\psi_2(0) =$ 

 $\psi_2\left(\frac{L}{4}\right) =$ 

 $\psi_2\left(\frac{L}{2}\right) =$ 

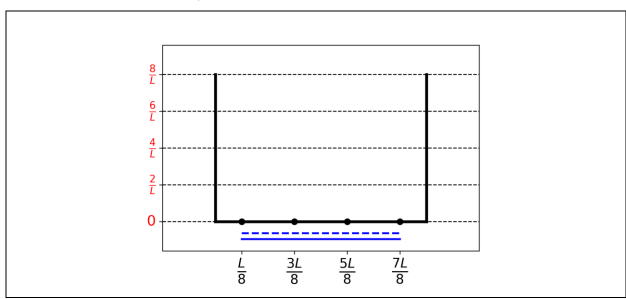
9. Within model 3, give the value of the  $\pi$  electron density at positions 0, L/4 and L/2.

 $\rho(0) =$ 

$$\rho\left(\frac{L}{4}\right) =$$

$$\rho\left(\frac{L}{2}\right) =$$

10. **Draw** the  $\pi$  electron density between 0 and L.



11. <u>Arrange</u> the following CC bonds (B1, B2, ..., B5) in order of increasing length, using the symbols = or <:

B1: C1C2 in the butadiene molecule

B2: C2C3 in the butadiene molecule

B3: C3C4 in the butadiene molecule

B4: CC in the ethane molecule

B5: CC in the ethene molecule

Problem	Question	1	2	3	4	5	6	7	8	9	10	Total
T2	Points	1	4	2	3	3	6	4	1	8	2	34
7%	Score											

# Problem T2: Hydrogen production by water-splitting

#### Data:

Compound	$H_2(g)$	H <sub>2</sub> O(1)	H <sub>2</sub> O(g)	O <sub>2</sub> (g)
$\Delta_{\rm f} H^{\circ} ({\rm kJ~mol}^{-1})$	0	-285.8	-241.8	0
$S_{\rm m}^{\circ} (\mathrm{J}  \mathrm{mol}^{-1}  \mathrm{K}^{-1})$	130.6	69.9	188.7	205.2

Molecular hydrogen  $(H_2)$  can be used as an alternative to carbon dioxide-emitting fuels. Hence, lowering the cost and the environmental impact of its production is a major challenge. In this field, water-splitting is a promising candidate technology.

1.	<u>Write down</u> the balanced equation of liquid water splitting reaction. <u>Use 1 as the stoichiometric coefficient of water.</u>
2.	Using only the provided thermodynamic data, <b>justify numerically</b> whether this reaction is thermodynamically favorable at 298 K.
Cal	lculations:
Is t	the reaction thermodynamically favorable?
	□ Yes □ No

Water splitting can be performed electrochemically using two electrodes in an acidic water bath, connected by a generator (*Fig. 1*). Gas bubbles are formed at both electrodes.

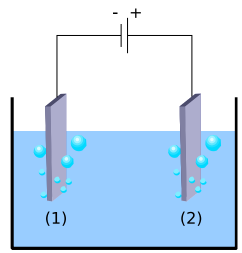


Fig. 1 – Water-splitting electrochemical cell.

3. Write down the balanced electrochemical half reactions occurring at each electrode.

On electrode (1):	
On electrode (2):	

4. Using only the provided thermodynamic data (or question 2), <u>derive</u> the condition on the applied voltage between electrodes ( $\Delta E_{\rm applied}$ ) and compare it to the value for the process to be thermodynamically favorable at 298 K ( $\Delta E_{\rm th}$ ), when all reactants and products are in their standard state. <u>Tick</u> the right condition and <u>give</u> the numerical value with 3 decimal places.

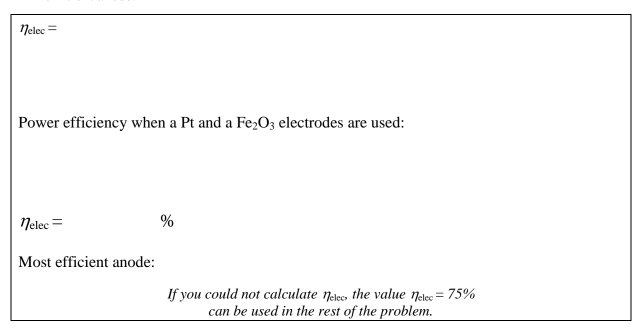
Calculation:	
$\Box$ $\Delta E$ $-\Delta E$	
$\Box  \Delta E_{\text{applied}} = \Delta E_{\text{th}}$	where AE V
$\square$ $\Delta E_{\text{applied}} > \Delta E_{\text{th}}$	where $\Delta E_{\mathrm{th}} = $ V
$\square$ $\Delta E_{\text{applied}} < \Delta E_{\text{th}}$	(give the result with 3 decimal places)
	If you could not calculate $\Delta E_{th}$ , the value 1.200 V
	can be used in the rest of the problem.

Experimentally, a higher voltage is needed to observe water splitting. For a given Pt cathode, the minimum voltage necessary to observe water splitting,  $\Delta E_{\min}$ , depends on the nature of the anode, as displayed in the table below:

Anode	$\Delta E_{\min}(V)$
$IrO_x$	1.6
$NiO_x$	1.7
$CoO_x$	1.7
$Fe_2O_3$	1.9

The difference between  $\Delta E_{min}$  and  $\Delta E_{th}$  is responsible for losses in the device.

5. <u>Give</u> the expression of the device power efficiency  $\eta_{elec}$  (fraction of the power used for water splitting) as a function of  $\Delta E_{th}$  and  $\Delta E_{min}$ . Assuming an identical current value *I*, <u>calculate</u> the water electrolysis power efficiency when a Pt cathode and a Fe<sub>2</sub>O<sub>3</sub> anode are used. <u>Give</u> the most efficient anode.



An alternative to water electrolysis is direct photocatalytic water-splitting. It uses a semiconductor that can be activated by absorbing light.

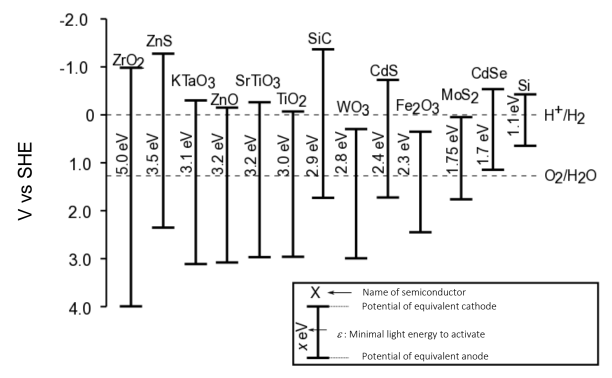


Fig. 2 – Activation condition and equivalent electrode potentials of different semiconductors. Dashed lines correspond to water oxidation and reduction potentials. SHE = Standard Hydrogen Electrode

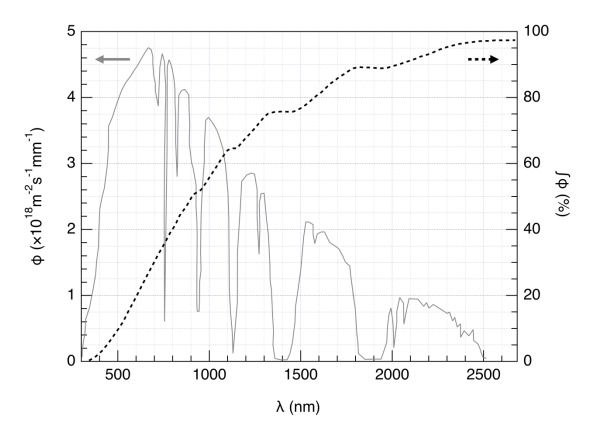


Fig. 3 – Left axis: Spectral distribution of the solar photon flux ( $\phi$ ). The photon flux is the number of photons per unit area per unit time arriving on the semiconductor. Right axis and dashed line: cumulative photon flux (i.e. fraction of the photon flux with smaller wavelength).

6. **Estimate** the fraction of the solar photon flux that can activate the following semiconductors: TiO<sub>2</sub>, CdS, Si. **State** explicitly the equations and units used for the computation.

Explanation / calculation:	

	Approximate fraction	
TiO <sub>2</sub>		%
CdS		%
Si		%

The activation of the semi-conductor results in a modification of the surface potentials, so that it can be seen as two electrodes of different potentials.

7.		Fig. 2, <b>choose</b> the semiconducts of anode and cathode for water		list that, once activated,
	ZrO <sub>2</sub>	□ZnO	□ TiO <sub>2</sub>	□ WO <sub>3</sub>
	CdS	$\square$ Fe <sub>2</sub> O <sub>3</sub>	□ CdSe	□ Si
8.	· · · · · · · · · · · · · · · · · · ·	ductor that, used as both cathod upon a given solar shining.	e and anode, is expected	to be the most efficient
$p_{ati}$ $S =$	m was recently studices 16 mm <sup>2</sup> surface, suction.	and $O_2$ when a semiconductor is it ed. Using an incident power light the production of $V = 0.37$ cm <sup>2</sup> wer efficiency $\eta_{\text{direct}}$ of the conve	at of $P = 1.0 \text{ kW m}^{-2}$ and of $H_2(g)$ was measur	a photoelectrode with a
	lculation:	, paneer		

$\eta_{ m direct}$ =	%	
	If you could not calculate $\eta_{\text{direct}}$ , the value $\eta_{\text{direct}} = 10\%$ can be used in the rest of the problem.	

Two modes of converting solar energy to hydrogen can thus be compared: direct photocatalysis, and indirect photo-electrolysis combining a photovoltaic panel with an electrolyzer. The efficiency of photovoltaic panels on the market is around  $\eta_{panels} = 20\%$ .

10. <u>Compare</u> the power efficiencies of the two modes,  $\eta_{\text{direct}}$  and  $\eta_{\text{indirect}}$ , using Fe<sub>2</sub>O<sub>3</sub> and Pt electrodes for the electrolysis.

Calculation:		
	_	
$\square$ $\eta_{ m direct} > \eta_{ m indirect}$	$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \eta_{ m indirect} $	$\square$ $\eta_{ m direct} < \eta_{ m indirect}$

Problem	Question	1	2	3	4	5	6	7	8	9	10	11	12	Total
T3	Points	1	3	3	3	4	2	7	2	2	3	4	6	40
5%	Score													

#### Problem T3: About silver chloride

#### Data at 298 K:

 $pK_{s1}(AgCl) = 9.7; pK_{s2}(Ag_2CrO_4) = 12$ 

Formation constant of the complex  $[Ag(NH_3)_n]^+$ :  $\beta_n = 10^{7.2}$ 

Potentials against the standard hydrogen electrode (SHE):

Standard potential of  $Ag^{+}|Ag(s)$ :  $E^{\circ}(Ag^{+}|Ag(s)) = 0.80 \text{ V}$ 

Apparent potential of  $O_2(aq)|HO^-(aq)$  (in seawater):  $E'(O_2(aq)|HO^-(aq)) = 0.75 \text{ V}$ 

#### Part A: Quotes from a chemistry lesson by Louis Joseph Gay-Lussac

The following quotes from a chemistry lesson by Louis Joseph Gay-Lussac (French chemist and physicist, 1778–1850) deal with some properties of silver chloride.

**Quote A:** "I will now talk about silver chloride, a milk-white solid. It is easily obtained by pouring hydrochloric acid into an aqueous solution of silver nitrate."

Quote B: "This salt has no taste since it is insoluble."

**Quote C:** "This compound is completely insoluble in alcohol and even in acids, except in concentrated hydrochloric acid which dissolves it readily."

Quote D: "On the other hand, silver chloride is highly soluble in aqueous solution of ammonia."

Quote E: "Then, we can make silver chloride appear again by adding an acid which reacts with ammonia."

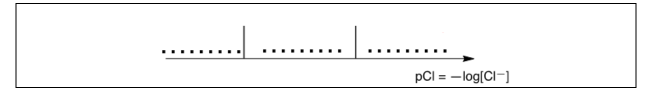
**Quote F:** "If you take a bowl made of silver to evaporate salty seawater, you will get impure sodium chloride, mixed with a milk-white solid."

2. **Quote B:** Calculate the solubility s of AgCl(s) in water at 298 K in mol L<sup>-1</sup>.

1. **Quote A: Write** the balanced chemical equation for synthesis AgCl(s).

Calculation:		
	s =	$mol L^{-1}$

3. **Quote C:** In a highly concentrated solution of chloride ions, a well-defined complex of stoichiometry 1:2 is formed. On the following qualitative axis (with pCl increasing from left to right), **place** in each domain the silver-containing species that is predominant (or exists, for solids). pCl values at frontiers are not expected.



**Quote D:** When ammonia is added to silver chloride, a well-defined complex of stoichiometry n is formed.

4. <u>Write</u> the balanced equation corresponding to the synthesis of the complex  $[Ag(NH_3)_n]^+$  from silver chloride and <u>calculate</u> the corresponding equilibrium constant.

Equation:	
Calculation:	
	•
	K =
	If you could not calculate $K$ , the following value can be used in the rest of the problem: $K = 10^{-3}$

5. Ammonia is added to 0.1 mol of silver chloride in 1 L of water until the last grain of solid disappears. At this moment,  $[NH_3] = 1.78 \text{ mol } L^{-1}$ . **Determine** the stoichiometry of the complex neglecting dilution effects.

n =	

6.	<u>Write</u> the balanced chemical equation corresponding to <b>quote E</b> .
7.	Assuming that seawater is slightly basic and rich in dioxygen, and that silver metal can reduce dioxygen in such conditions, <b>write</b> a balanced chemical equation corresponding to the formation of the solid mentioned in <b>quote F.</b> Use 1 as the stoichiometric coefficient of dioxygen. <b>Calculate</b> its equilibrium constant at 298 K.
Equ	uation:
Cal	culation:
	K =
	n –

#### Part B: The Mohr method

The Mohr method is based on the colorimetric titration of Cl<sup>-</sup> by Ag<sup>+</sup> in the presence of potassium chromate  $(2K^+, CrO_4^{2^-})$ . Three drops (~ 0.5 mL) of  $7.76 \cdot 10^{-3}$  mol L<sup>-1</sup> K<sub>2</sub>CrO<sub>4</sub> solution are added to V<sub>0</sub> = 20.00 mL of a sodium chloride solution of unknown concentration  $C_{Cl}$ . This solution is then titrated by silver nitrate  $(Ag^+, NO_3^-)$  with  $C_{Ag} = 0.050$  mol L<sup>-1</sup>, which immediately leads to the formation of solid **A**. A red precipitate (solid **B**) appears at  $V_{Ag} = 4.30$  mL.

8.	Write the balanced equations of the two reactions	-	ent. <u>Calculate</u> the
	corresponding equilibrium constants.		
		$K^{\circ}{}_{1} =$	
		$K^{\circ}{}_{2} =$	
9.	<u>Identify</u> the solids.		
	Solid A:		
	Solid <b>B</b> :		
10	. <u>Calculate</u> the unknown concentration of chloride	ions ( $C_{\rm Cl}$ ) in the sodium chloric	de solution.
Ca	lculation:		
		$C_{\mathrm{Cl}} =$	$mol L^{-1}$
	If you could not calculate $C_{Cl}$ , the can be used in the rest		

11. <u>Calculate</u> the minimal volume $V_{Ag}(min)$ for which AgCl(s) precipitates.					
Calculation:					
	$V_{ m Ag}({ m min}) =$	mI	_		
	·				
			es) when silver chromate beg		
	Jusury wny CrO <sub>4</sub> is a g	good indicator for Monr	titration by comparing two valu	es.	
Calculation:					
		$[Cl^-]_{res} =$	$mol L^{-1}$		
			mor L		
$CrO_4^{2-}$ is a goo	d titration endpoint ind	licator because:			

Problem	Question	1	2	3	4	5	6	7	8	Total
T4	Points	6	9	8	5	6	2	2	12	50
7%	Score									

# Problem T4: From gunpowder to the discovery of iodine

In the 19<sup>th</sup> century, the French entrepreneur B. Courtois specialized in the production of nitrate  $\mathbf{A}$  ( $\mathbf{M}_{\mathbf{A}}(\mathrm{NO}_3)_m$ ), used for gunpowder. Initially imported from Asia,  $\mathbf{A}$  was later produced from nitrate  $\mathbf{B}$  ( $\mathbf{M}_{\mathbf{B}}(\mathrm{NO}_3)_n$ ) using exchange reaction with compound  $\mathbf{C}$ , obtained from algae.

1.	. <u>Find</u> the formulas of nitrates $A$ and $B$ knowing that they are anhydrous salts of alkali or alkaline-earth metal ( $M_A$ and $M_B$ ). One of the nitrates contains no more than 1 w% of non-metallic impurities while the other contains $9 \pm 3$ w% of impurities. The content of metals $M_A$ and $M_B$ in the samples is 38.4 w% and 22.4 w% respectively. <u>Support</u> your answer with calculations.						
	<b>A</b> :		and <b>B</b> :				

the sample (containing only nitrites, NO<sub>2</sub><sup>-</sup>) was constant. The only gaseous product was dioxygen: 60.48 L at 0 °C at 1 atm (dioxygen can be considered as an ideal gas). 2. Calculate the composition (in w%) of mixture E considering that it contained only compounds A and **B** and no other impurities, and that **C** was taken in pure anhydrous state. w% of **A**: and of **B**:

To obtain **A**, 262.2 g of solid compound **C** were added to the solution containing 442.8 g of **B**. **B** is known to be in excess. As a result, 190.0 g of white precipitate **D** were formed and removed by filtration. The filtrate was evaporated, and the obtained solid mixture **E** was heated until the mass of

3.	<b>Determine</b> the formulas of condetween <b>B</b> and <b>C</b> .	ompounds <b>C</b> and <b>D</b> and <u>wri</u>	<b>te</b> the balanced equation for the reaction
	C:		and <b>D</b> :
Re	action between <b>B</b> and <b>C</b> :		

In 1811, when working with algae ashes, Courtois observed that copper vessels were worn out faster than usual. While he was studying this phenomenon, his cat entered the laboratory and spilled the solution of concentrated sulfuric acid on the dry algae ashes: violet vapors instantly came out of the vessel (1, sulfuric acid is the oxidizing agent): iodine ( $I_2$ ) had just been discovered! Iodine was the cause of the copper corrosion (2). However, because of the medicinal applications of iodine, Courtois opened a new manufacture to produce it by reaction of algae ashes with chlorine (3). Nowadays, iodine is prepared from the set of reactants ( $NO_3^-$ ,  $I^-$ ,  $H^+$ ) (4) or ( $IO_3^-$ ,  $I^-$ ,  $H^+$ ) (5).

4. **Write** the balanced chemical equation for reactions **1–5**.

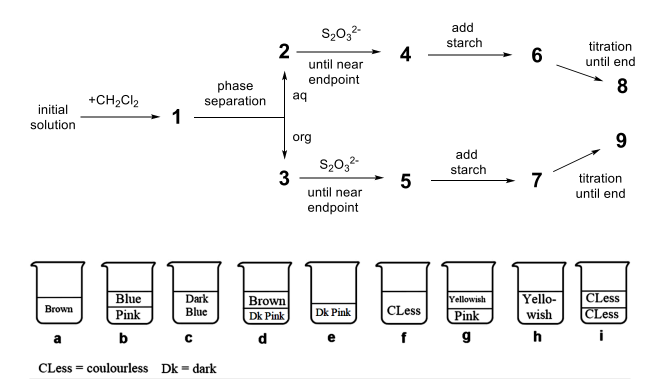
1		
2		
3		
4		
5		

The solubility of iodine is very low in water but significantly increases when iodide ions are added. Together they form ions such as triiodide,  $I_3^-$ :

$$I^{-}(aq) + I_{2}(aq) = I_{3}^{-}(aq)$$
 (6)

Equilibrium (6) can be studied through the extraction of  $I_2$  with dichloromethane. Indeed,  $I^-$  and  $I_3^-$  do not dissolve in organic solvents but  $I_2$  does and, when extracted, it is 15 times more concentrated in dichloromethane than in water.

The following experiment was performed. To prepare the initial solution, a few crystals of solid iodine were dissolved in 50.0 mL of an aqueous solution of potassium iodide (0.1112 g). Then, 50.0 mL of dichloromethane were added, and the mixture was vigorously shaken until equilibration. After phase separation, the organic phase was titrated by 16.20 mL while the aqueous phase was titrated by 8.00 mL of the standard aqueous solution of sodium thiosulfate pentahydrate (14.9080 g in 1.000 L of solution) in the presence of starch. The process is schematically represented on the next page:



5. <u>Find</u> the correspondence between the stages on the scheme (1–9) and the schematic pictures representing them (a–i).

Stages	Picture
1	
2	
3	
4	
5	
6	
7	
8	
9	

6.	<u>Write</u> balanced equations for the two possible chemical reactions in the aqueous phase during the titration involving iodine species and sodium thiosulfate.
7.	<u>Calculate</u> the mass of iodine used to prepare the initial solution.
	$m(I_2) = g$

8.	3. <u>Calculate</u> the equilibrium constant $K^{\circ}$ for equilibrium	juilibrium of reaction (6).	
			<i>K</i> ° =
			<b>11</b> –

Problem	Question	1	2	3	4	5	6	7	8	9	10	11	12	Total
T5	Points	3	4	4	2	5	5	4	3	5	2	2	2	41
8%	Score													

# Problem T5: Azobenzene – $\beta$ -cyclodextrin complexes for the formation of nanomachines

Nanomachines are molecular assemblies that enable the transformation of an energy source into a nano-movement for applications such as drug delivery. Numerous nanomachines make use of the isomerization of azo compounds (R-N=N-R') upon irradiation.

1. <u>Draw</u> the stereoisomers of azobenzene ( $H_5C_6-N=N-C_6H_5$ ) and <u>draw</u> a line between the two carbon atoms that are the furthest apart. <u>Compare</u> these two distances ( $d_{trans}$  and  $d_{cis}$ ).

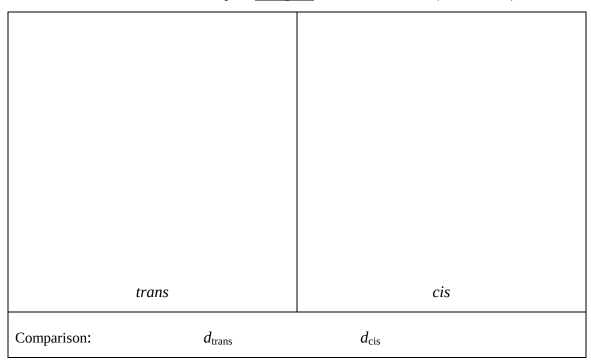


Fig. 1 – Possible reactants for the synthesis of M.

2. **M** can be synthesized in two steps from simple reactants (*Fig. 1*). <u>Choose</u> among the suggested reactants (**N** to **Q**) the ones that can provide **M** with very high regioselectivity. Sodium nitrite (NaNO<sub>2</sub>) in cold aqueous hydrochloric acid is used as reagent for the first step of the synthesis.

Reactants:	and

#### Determination of the association constant $K_t$

β-cyclodextrin ( $\mathbf{C}$ , Fig. 2) is a cyclic heptamer of glucose, which can form inclusion complexes with azo compounds. In tasks 3 to 6, we will determine by spectroscopy the association constant  $K_t$ , corresponding to the formation of the inclusion complex  $CM_{trans}$  as depicted in Fig. 2.

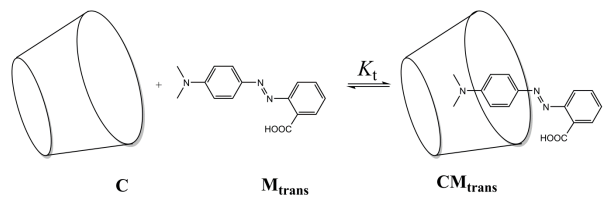


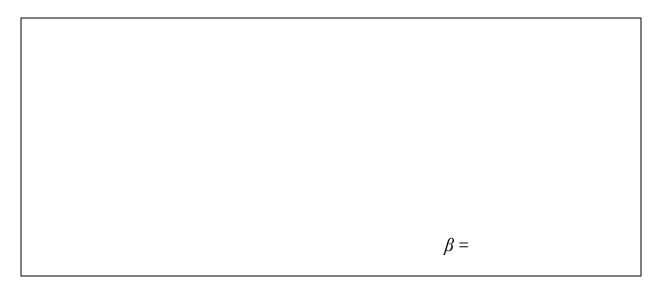
Fig. 2 – Formation of the  $CM_{trans}$  inclusion complex.

Several solutions are prepared by mixing C and  $M_{trans}$  in different proportions to reach initial concentrations  $[C]_0$  and  $[M_{trans}]_0$ . While  $[M_{trans}]_0$  is identical for all solutions,  $[C]_0$  varies. We follow, at a fixed wavelength, the evolution of the difference in absorbance ( $\Delta A$ ) between the absorbance of each solution and the pure  $M_{trans}$  solution. We note the molar absorption coefficients of  $CM_{trans}$  and  $M_{trans}$ , respectively. L is the path length of the beam through the sample. The absorbance of C ( $\varepsilon_C$ ) is negligible.

3. **Prove** that  $\Delta A = \alpha \cdot [\mathbf{CM_{trans}}]$  and **express**  $\alpha$  in terms of known constant(s).

Proof:			

lpha =
4. <u>Prove</u> that, when $C$ is in large excess with respect to $M_{trans}$ (i.e. $[C]_0 >> [M_{trans}]_0$ ), the concentration of $C$ may be considered as constant, $[C] \simeq [C]_0$ .
Proof:
5. <b>Prove</b> that, when <b>C</b> is in large excess with respect to $\mathbf{M}_{\text{trans}}$ ( <i>i.e.</i> $[\mathbf{C}]_0 >> [\mathbf{M}_{\text{trans}}]_0$ ), $\Delta A = \alpha \cdot \frac{\beta \cdot [\mathbf{C}]_0}{1 + K_t \cdot [\mathbf{C}]_0}$ and $\underline{\text{express}} \beta$ in terms of constant(s) and initial concentration(s).
Proof:



6. **Determine**  $K_t$  using the following experimental curve (*Fig. 3*).

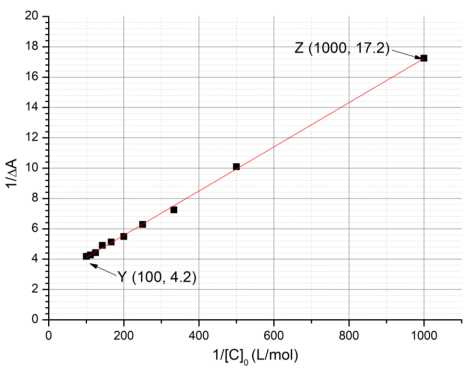
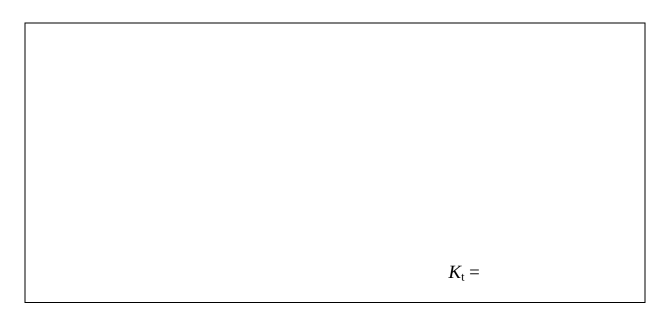


Fig. 3 – Evolution of  $1/\Delta A$  as a function of  $1/[C]_0$ .

Calculations:		



#### Determination of the association constant $K_c$

In tasks 7 to 9, we will determine by kinetic studies the association constant  $K_c$ , corresponding to the formation of the inclusion complex with  $\mathbf{M}_{cis}$ ,  $\mathbf{CM}_{cis}$ . A sample containing only  $\mathbf{M}_{trans}$  is irradiated, thus producing  $\mathbf{M}_{cis}$  with a known concentration  $[\mathbf{M}_{cis}]_0$ .  $\mathbf{M}_{cis}$  (free or within the inclusion complex) then thermally isomerizes into  $\mathbf{M}_{trans}$ . In the absence of  $\mathbf{C}$ , the isomerization follows a first order kinetics with a rate constant  $k_1$ . All complexation equilibria are faster than the isomerization processes. The kinetic scheme corresponding to this experiment is provided in Fig. 4.

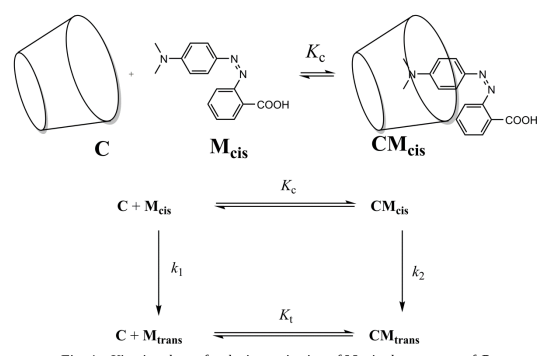


Fig. 4 – Kinetic scheme for the isomerization of  $M_{cis}$  in the presence of C.

The rate of disappearance r for the total amount of  $\mathbf{M}_{cis}$  (free and complexed) is defined as  $r = k_1[\mathbf{M}_{cis}] + k_2[\mathbf{C}\mathbf{M}_{cis}]$ 

Experimentally, r follows an apparent first order kinetics with an apparent rate constant  $k_{obs}$ :

$$r = k_{\text{obs}}([\mathbf{M_{cis}}] + [\mathbf{CM_{cis}}])$$

7. **Prove** that  $k_{\text{obs}} = \frac{\gamma + \delta \cdot k_2[C]}{1 + K_C[C]}$  and **express**  $\gamma$  and  $\delta$  in terms of known constant(s).

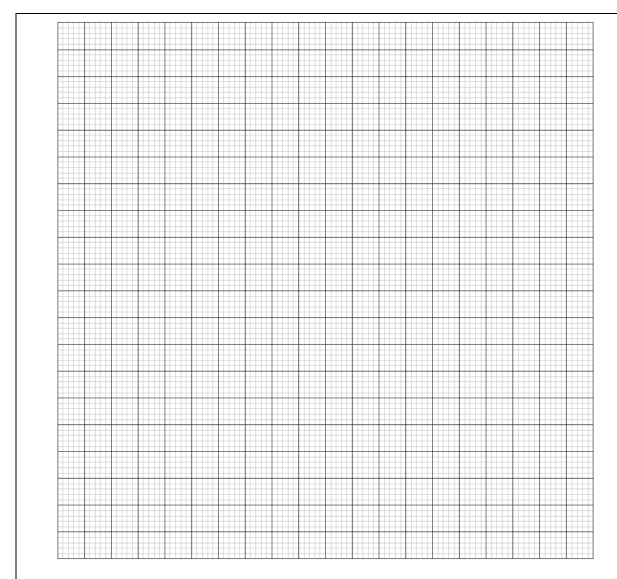
	=([-]		
Proof:			
	$\gamma =$	and	$\delta$ =

8. Choose in which condition(s) the half-life  $t_{1/2}$  corresponding to  $k_{\rm obs}$  can be expressed as  $t_{1/2} = \frac{\ln 2}{\gamma} \ (1 + K_{\rm c}[{\bf C}]_0)$  given that  $[{\bf C}]_0 >> [{\bf M}_{\rm cis}]_0$ . Mathematically justify your answer.

	Very slow isomerization of $\mathbf{M}_{\mathbf{cis}}$ within cyclodextrin
	Very slow isomerization of free $\mathbf{M}_{\mathbf{cis}}$
	CM <sub>cis</sub> very stable
	CM <sub>trans</sub> very stable
Desof	
Proof:	

9. Assuming the condition(s) in task 8 satisfied, <u>determine</u>  $K_c$  by linear regression using the data below. You may use a calculator or plot a graph.

$[\mathbf{C}]_0 \text{ (mol } \mathbf{L}^{-1})$	$t_{1/2}$ (s)	$[\mathbf{C}]_0 \text{ (mol } \mathbf{L}^{-1})$	$t_{1/2}$ (s)
0	3.0	$3.0 \cdot 10^{-3}$	5.9
$1.0 \cdot 10^{-4}$	3.2	$5.0 \cdot 10^{-3}$	7.7
$5.0 \cdot 10^{-4}$	3.6	$7.5 \cdot 10^{-3}$	9.9
$1.0 \cdot 10^{-3}$	4.1	$1.0 \cdot 10^{-2}$	12.6



Equation of the linear regression:

 $K_{\rm c} =$ 

### **Formation of nanomachines**

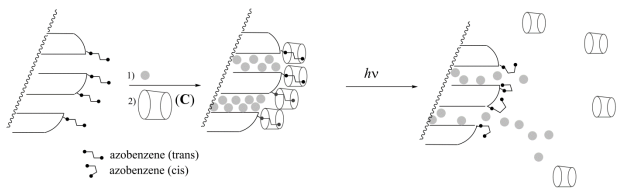


Fig. 5 – Cleavage of an azobenzene-cyclodextrin inclusion complex induced by a light-triggered isomerization, which allows delivery of a dye (grey circles).

Another azobenzene compound (for which  $K_c \ll K_t$ ), initially in the *trans* form, is covalently grafted on silica (*Fig. 5*). The silica pores are filled with a dye (rhodamine B, grey circles in *Fig. 5*). Upon addition of **C**, an inclusion complex is formed, which blocks the pores and prevents the release of the dye.

10. <u>Choose</u> the most appropriate condition (choose only one) so that the pores are initially blocked in the presence of **C**, and the dye can be released upon irradiation.

$K_{\rm t} >> 1$
$K_{\rm t} >> 1$ and $K_{\rm c} << 1$
$K_{\rm t} / K_{\rm c} << 1$
$K_{\rm t} >> 1$ and $K_{\rm c} >> 1$
$K_{\rm c} \ll 1$

This azobenzene-silica powder loaded with a dye is placed in the corner of a cuvette (Fig. 6) so that this powder cannot move into the solution. The powder is irradiated at a wavelength  $\lambda_1$  to trigger the release of the dye from the pores (Fig. 5). To monitor this release, we measure the absorbance of the solution at wavelength  $\lambda_2$ .

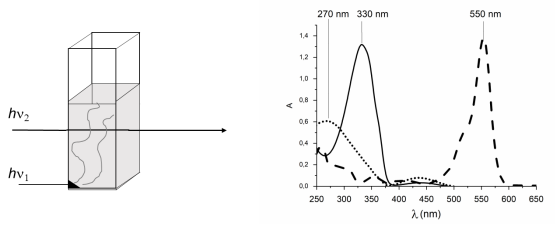


Fig. 6 – Left: experimental setup used to monitor the release of the dye; right: absorption spectra of trans-azobenzene (full line), cis-azobenzene (dotted line) and rhodamine B (dashed line).

### 11. **Determine** $\lambda_1$ .

$$\lambda_1 = nm$$

### 12. **Determine** $\lambda_2$ .

 $\lambda_2$  = nm

Problem	Question	1	2	3	4	5	6	7	8	9	Total
Т6	Points	4	4	5	3	10	2	9	6	5	48
8%	Score										

# Problem T6: Characterization of a block-copolymer

Block-copolymers, obtained by linking different polymers (blocks), have unique properties, such as the ability to self-assemble. In this problem, the synthesis and characterization of such a macromolecule are studied.

### Study of the first block

$$H_2N$$

$$\begin{array}{c}
O \\
 & \\
1
\end{array}$$
och<sub>3</sub>

In this first part, we will study the water soluble homopolymer  ${\bf 1}$  ( $\alpha$ -methoxy- $\omega$ -aminopolyethyleneglycol).

The <sup>1</sup>H NMR spectrum of **1** (DMSO- $d_6$ , 60 °C, 500 MHz) includes the following signals:

Index	δ (ppm)	Peak Area
a	2.7*	0.6
b	3.3	0.9
c	3.4	0.6
d	~ 3.5	133.7

Table 1, \*in the presence of  $D_2O$ , the signal at 2.7 ppm disappears.

1. Match the <sup>1</sup>H NMR signals (a, b, c, d) from *Table 1* with each of the corresponding protons.

2.	<b>Express</b> the average degree of polymerization $n$ as a function of the area $A_{\text{OC2H4}}$ of the NMR peak
	of the repeating unit and the area $A_{\text{OCH3}}$ of the NMR peak of the methyl end group. Calculate $n$ .

n =

If you could not calculate n, the value n = 100 can be used in the rest of the problem.

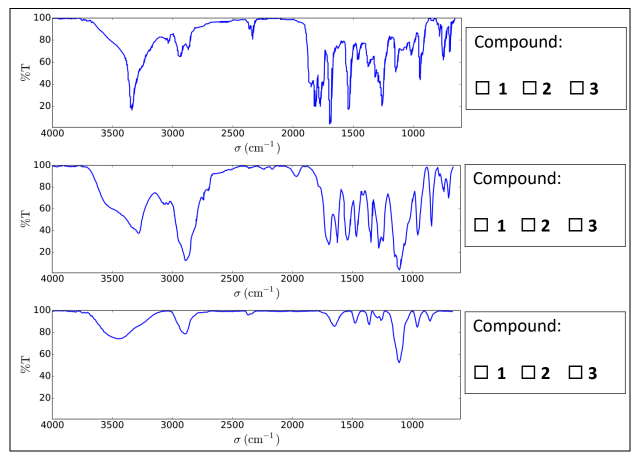
## Study of a diblock-copolymer

The synthesis of the second block of the copolymer is performed through the reaction of 1 with 2 ( $\epsilon$ -(benzyloxycarbonyl)-lysine N-carboxyanhydride). This yields the block-copolymer 3.

3. <u>Draw</u> the reaction intermediate that is formed in the first step of the addition of 1 to 2. The second step of the mechanism leads to the formation of a gas molecule, G. <u>Draw</u> its structure.

**G**:

4. Infrared (IR) measurements are performed to characterize the compounds. <u>Match</u> the three IR spectra with compounds 1, 2 and 3.



5. The <sup>1</sup>H NMR spectrum of copolymer **3** (in DMSO- $d_6$ , at 60 °C, 500 MHz) is reported in Fig. 1. Using some or all of the NMR signals, the areas of which are reported in *Table 2*, <u>calculate</u> its number average molar mass  $M_n$ , considering n from question 2. For your calculations, <u>draw</u> a circle around the group(s) of atoms you used and <u>give</u> their corresponding symbol(s)  $(\alpha, \beta, ...)$ .

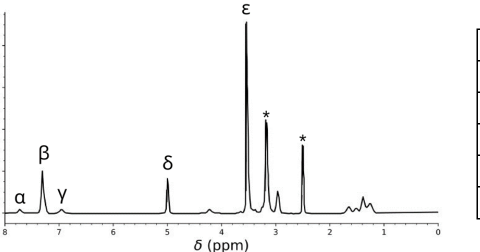


Fig. 1 – signals marked with \* correspond to the solvent and water.

Tal	ole 2
Peak	Area
α	22.4
В	119
γ	23.8
δ	47.6
3	622

$$H \longrightarrow \begin{pmatrix} 0 \\ N \\ H \end{pmatrix}_{m} \longrightarrow \begin{pmatrix} 0 \\ 0 \\ 0 \end{pmatrix}$$

 $M_{\rm n} = {\rm kg \ mol}^{-1}$  (Express your answer with two decimal places)

This reaction of **1** with **2** yielded the copolymers **3a** after 20 h, **3b** after 25 h and **3c** after 30 h of reaction at 40 °C. Results of size-exclusion chromatography (SEC) experiments are presented in Fig. 2.

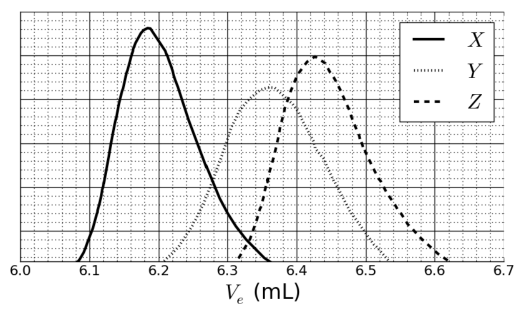


Fig. 2 – SEC chromatograms of 3a, 3b and 3c as a function of the elution volume,  $V_e$ .

6. Match the signals in Fig. 2 with the copolymers 3a, 3b and 3c.

3a:	$\square X$	$\square Y$	$\square Z$	
3a: 3b:	$\square X$	$\square Y$	$\square Z$	
3c:	$\square X$	$\square Y$	$\square Z$	

In order to calibrate the chromatogram, a mixture of standard polymers of known masses (3, 30, 130, 700 and  $7000 \text{ kg mol}^{-1}$ ) has been studied (*Fig. 3*).

The log value of the molar mass is a linear function of the elution volume,  $V_{\rm e}$ .

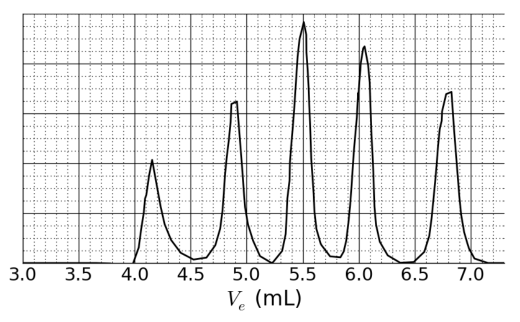


Fig. 3-SEC chromatogram of the mixture of standards.

7. Based on the SEC curves in *Fig. 2 and 3*, <u>determine</u>  $V_e$  of the polymer that corresponds to curve X and use it to <u>estimate</u> the degree of polymerization m of its second block. <u>Detail</u> your calculation; you may use a calculator or plot a graph.

## **Triblock copolymer synthesis**

For biological applications, involving the formation of micelles, a triblock copolymer 9 can be synthesized through the introduction of a middle block, **B**, using monomer 5.

$$\begin{array}{c|c}
CI & CI \\
MsCI: & O=S=O \\
CH_3
\end{array}$$

8. **Draw** the structures of **5**, **7** and **8**.

5 (no other products than 6: A-B is obtained)

7 (a gas is formed in the final step)

8

9. Amphiphilic block copolymers, such as **9: A-B-C**, can be used for medical applications, as they self-assemble into micelles in water (pH = 7), which can be used as drug carriers. <u>Assign</u> each block of the copolymer to a property. <u>Draw</u> a scheme of the micelle with only 4 polymer chains.

<b>A</b> :	☐ hydrophobic	□ hydrophilic	
<b>B</b> :	☐ hydrophobic	□ hydrophilic	
<b>C</b> :	☐ hydrophobic	□ hydrophilic	
	<b>A W</b>	В —	C
3	<i>&gt;</i>		
my 3	}	<del></del>	
	W/\		

## Problem T7: Ring motion in a [2]catenane

Problem	Question	1	2	3	4	5	6	7	8	9	10	11	Total
<b>T7</b>	Points	4	12	2	2	2	5	5	8	4	5	5	54
6%	Score												

In 2016, the Nobel Prize in Chemistry was awarded to J.-P. Sauvage, Sir J. F. Stoddart and B. L. Feringa "for the design and synthesis of molecular machines". An example of these is [2]catenane, a molecule consisting of two interlocked rings. In this system, one macrocycle contains a single phenanthroline (bidentate) ligand and the second contains two ligands: a phenanthroline and a terpyridine (tridentate) ligand. A copper ion is coordinated by one ligand from each macrocycle. Depending on the oxidation state of the copper (+I or +II), two configurations are obtained (*Fig. 1*).

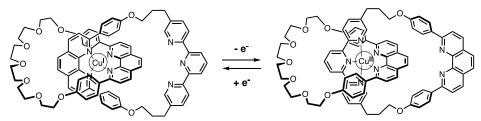


Fig. 1 – Multi-stability of a ring in a [2]catenane.

The synthesis of the macrocycle is the following:

$$\begin{array}{c} \text{Br} \\ \text{C} \\ \text{C} \\ \text{CPquiv.} \end{array} \end{array} \longrightarrow \begin{array}{c} \text{LDA} \\ \text{Q2 equiv.} \end{array} \longrightarrow \begin{array}{c} \text{C} \\ \text{Pr} \\ \text{Q2 equiv.} \end{array} \longrightarrow \begin{array}{c} \text{C} \\ \text{Pr} \\ \text{Q2 equiv.} \end{array} \longrightarrow \begin{array}{c} \text{C} \\ \text{Pr} \\ \text{C2 equiv.} \end{array} \longrightarrow \begin{array}{c} \text{C} \\ \text{C3 equiv.} \end{array} \longrightarrow \begin{array}{c} \text{C3 equiv.} \\ \text{C4 equiv.} \end{array} \longrightarrow \begin{array}{c} \text{C4 equiv.} \\ \text{C4 equiv.} \end{array} \longrightarrow \begin{array}{c} \text{C4 equiv.} \\ \text{C4 equiv.} \end{array} \longrightarrow \begin{array}{c} \text{C4 equiv.} \\ \text{C5 equiv.} \end{array} \longrightarrow \begin{array}{c} \text{C6 equiv.} \\ \text{C7 equiv.} \end{array} \longrightarrow \begin{array}{c} \text{C7 equiv.} \\ \text{C9 equiv.} \end{array} \longrightarrow \begin{array}{c} \text{C9 equiv.$$

#### 1. **<u>Draw</u>** the structure of **B**.

B

2.	<u>Draw</u> the structures of <b>E</b> , <b>F</b> and <b>G</b> .
E	
F	
G	
3.	Out of the following the reaction conditions, $\underline{choose}$ which one(s) can produce <b>E</b> from <b>D</b> :
П	$H^+$ , $H_2O$
	$OH^-, H_2O$
	NaBH <sub>4</sub> , CH <sub>3</sub> OH
	H <sub>2</sub> , Pd/C, THF
4.	In the synthetic strategy, MsCl is used to obtain:
	a leaving group
	a protecting group
	a deactivating group
Ш	a directing group
5.	${f G}$ is obtained by the reaction between ${f F}$ and LiBr in acetone. This reaction is:
	electrophilic aromatic substitution
	nucleophilic aromatic substitution
	$S_N 1$

6.	$\underline{\mathbf{Draw}}$ the transition state of the rate-determining step of the reaction $\mathbf{F} \to \mathbf{G}$ , showing the 3D
	geometry. Depict only one reaction center. The main carbon chain can be represented as an R
	group.

Transition state:			

The synthesis of [2]catenane **L** uses the template effect of a copper complex:

7. <u>Write</u> the full electron configuration of Cu(0) in its ground state. Give the oxidation state of Cu in complex **J** and write the electron configuration of Cu in the free ion corresponding to **J**.

Electron configuration of Cu(0):
Oxidation state of Cu in <b>J</b> :
Oxidation state of Cu in §.
Electron configuration of Co. in I.
Electron configuration of Cu in <b>J</b> :

8. <u>Select</u> the geometry of the copper ion in **L**. Assuming an ideal geometry of the ligands around the copper center, <u>draw</u> the electronic levels of the *d* orbitals subject to the crystal field. <u>Fill</u> the orbital diagram. <u>Give</u> the maximum value of the spin (*S*) for this complex.

The geometry of Cu in <b>L</b> is:
□ Octahedral
☐ Tetrahedral
□ Square planar
☐ Trigonal bipyramid
Splitting and filling of <i>d</i> orbitals:
C C
S =

9. Out of the following compounds, <u>choose</u> the one(s) that can remove the copper ion in L to obtain the free [2]catenane:

$$\begin{array}{c|c} \square & CH_3CN \\ \square & NH_4PF_6 \\ \square & KCN \\ \square & tren \end{array}$$

In [2]catenane **L**, the copper ion can exist in two oxidation states (+I) or (+II), and each of them exhibits a different coordination sphere (tetra- or penta-coordinated, respectively).

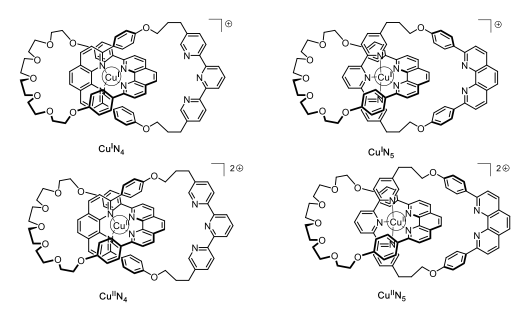


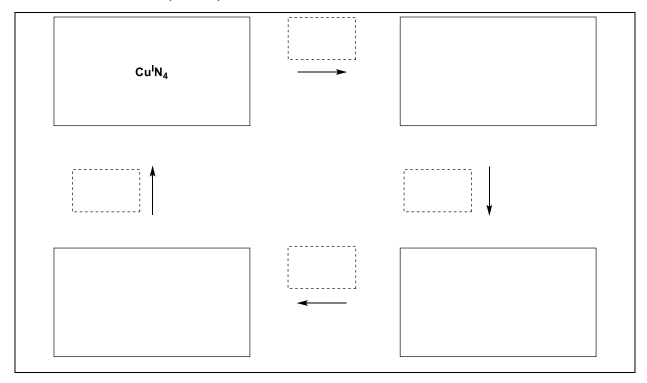
Fig. 2 – [2] catenane L states

The stability of Cu(I) complexes can be inferred by comparing their electronic structures to that of a noble gas.

10. **Fill** in the blanks with a number or a tick:

The Cu <sup>I</sup> N <sub>4</sub> complex has electrons in the coordination sphere of the metal.
The Cu <sup>I</sup> N <sub>5</sub> complex has electrons in the coordination sphere of the metal.
The $Cu^IN_4$ complex is $\square$ more / $\square$ less stable than the $Cu^IN_5$ complex.

11. <u>Fill</u> in the solid boxes with the designation of the involved complexes in *Fig.* 2 and <u>complete</u> the sequence to achieve electrochemical control of the system using the following notation for the dashed boxes: (rotation);  $+ e^-$ ;  $- e^-$ .



Problem	Question	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	Total
Т8	Points	2	6	2	2	11	2	4	3	4	2	6	8	2	6	4	64
6%	Score																

# **Problem T8: Identification and synthesis of inositols**

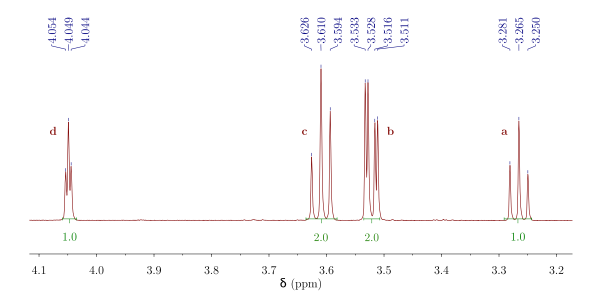
In this problem, we define " $\underline{3D}$  structure" and " $\underline{perspective}$  formula" as indicated for  $\beta$ -glucose in the following figure.

Inositols are cyclohexane-1,2,3,4,5,6-hexols. Some of these 6-membered carbocycles, in particular *myo*-inositol, are involved in a number of biological processes.

## Structure of myo-inositol

1. <u>Draw</u> the structural formula of inositols, without stereochemical details.						
This family of molecules contains 9 different stereoisomers, including enantiomers.						
2. <u>Draw</u> all 3D structures of the stereoisomers that are optically active.						

The structure of a specific inositol, called myo-inositol, is studied here. Only one of its chair conformers is predominant and its structure can be deduced from its  $^{1}H$  NMR spectrum. The spectrum below was obtained at 600 MHz in  $D_{2}O$ . No other signal from that compound was observed in the spectrum. The integration is indicated on the spectrum below each signal.



3. <u>Give</u> the molecular formula of the predominant compound derived from *myo*-inositol in this sample that is consistent with the number of protons observed in the <sup>1</sup>H NMR spectrum.

4. Based on the number and integrations of the proton signals, **give** the number of symmetry plane(s) that exist(s) in this molecule.

5. <u>Complete</u> the following perspective drawing of the most stable conformation of *myo*-inositol. Then <u>label</u> each hydrogen with the corresponding letter (**a**, **b**, **c** or **d**) according to the NMR spectrum above. Proton **a** must be on carbon **a** on the following representation. <u>Draw</u> its 3D structure.

## Synthesis of inositols

For medicinal applications, it is useful to synthesize some inositol phosphates on a large scale. We will study the synthesis of inositol 2 from bromodiol 1.

6. Choose the correct structural relationship(s) between 2 and 3.

☐ enantiomers		
□ epimers		
☐ diastereomers		
□ atropoisomers		

Inositol 2 can be obtained from compound 1 in 7 steps.

7.	<u>Draw</u> the 3D structure of 4.	
4		
8.	below the structure of 1-bromo-1,3-cyclohexa	e bond with the highest electron density. Consider adiene, which is a substructure of 4. Encircle the On separate structures, represent all the electronic
	Br	
9.	<u>Draw</u> the 3D structure of the major diastereom	er <b>5</b> .
5		
10	<ul> <li>Give the total number of stereoisomers of 5 enantiopure compound 1.</li> </ul>	possibly obtained by this synthesis, starting from
11	. For the step $5 \to 6$ , another product with produced. $\underline{\mathbf{Draw}}$ the 3D structures of $6$ and $6$ .	the same molecular formula, denoted 6', can be
6		6'

12. <u>Draw</u> the 3D structures of major diastereomers 8 and 9.						
8	9					
13. <b>Select</b> the right set(s) of conditions <b>A</b> to obtain	n <b>2</b> .					
<ul> <li>□ H<sub>2</sub>, Pd/C</li> <li>□ K<sub>2</sub>CO<sub>3</sub>, HF</li> <li>□ HCOOH, H<sub>2</sub>O</li> <li>□ BF<sub>3</sub>·OEt<sub>2</sub></li> </ul>						
obtained. Considering that the stereoselectiv	1, in addition to 2, another stereoisomer would be ity of the reactions that take place in the synthesis ups involve the same number of equivalents as for 2, d give its relationship with 2.					
□ enantiomers						
<ul><li>□ epimers</li><li>□ diastereoisomers</li><li>□ atropoisomers</li></ul>						
15. During the synthesis of <b>2</b> from <b>1</b> , <b>choose</b> the r	emoval step(s) of protecting or directing groups.					
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$						
$ \begin{array}{ccc} \square & 5 \to 6 \\ \square & 6 \to 7 \\ \square & 7 \to 8 \end{array} $						
$ \begin{array}{c} \square & 8 \rightarrow 9 \\ \square & 9 \rightarrow 2 \end{array} $						

Problem	Question	1	2	3	4	5	6	7	8	9	10	11	12	13	Total
Т9	Points	2	2	4	3	2	17	1	1	2	4	2	2	2	44
7%	Score														

# **Problem T9: Synthesis of levobupivacaine**

### Part I.

The local anesthetic bupivacaine (marketed as Marcaine) is on the World Health Organization List of Essential Medicines. Although the drug is currently used as a racemic mixture, it was demonstrated that one enantiomer of bupivacaine, levobupivacaine, is less cardiotoxic and, therefore, safer than the racemate. Levobupivacaine can be synthesized from the natural amino acid L-lysine.

$$CI^ H_3N$$
 $O$ 

L-Lysine hydrochloride

1. <u>Assign</u> the absolute configuration of the stereogenic center in L-lysine hydrochloride and <u>justify</u> your answer by assigning the substituents their order of priority.

Configuration:	Priority 1 > 2 > 3 > 4:
$\square R$	$NH_3^+_{Cl}$ $NH_3^+$ $COO^ H$
$\square S$	

2. The prefix L in L-lysine refers to relative configuration. **Choose** all correct statements:

- ☐ All natural L-amino acids are levorotatory.
- □ Natural L-amino acids can be levorotatory or dextrorotatory.
- $\square$  All natural L-amino acids are (S).
- $\square$  All natural L-amino acids are (R).

Often, we want only one of the amino groups in L-lysine to react. A  $Cu^{2+}$  salt with excess aqueous hydroxide can selectively mask the reactivity of one of the amino groups. After the complex is formed, only the non-complexed  $NH_2$  group is available to react.

3. Considering that L-lysine acts as a bidentate ligand and that two L-lysine molecules coordinate to one Cu<sup>2+</sup> ion in the presence of aqueous hydroxide, <u>draw</u> the structure of the intermediate complex.

Complex:			

Fortunately, in the synthesis of levobupivacaine shown below, the same amino group reacts even without  $Cu^{2+}$  salt.

$$\begin{array}{c} \text{CI} \\ \text{H}_{3} \text{N} \\ \text{L-Lysine} \\ \text{hydrochloride} \\ \text{NaNO}_{2}, \text{NaOAc} \\ \text{AcOH} \\ \text{C}_{16} \text{H}_{21} \text{NO}_{6} \\ \text{C}_{2} \text{H}_{20} \text{N}_{2} \text{O}_{4} \\ \text{D} \\ \text{DCC} \\ \text{DCC} \\ \text{PhCHO} \\ \text{Alignment of the ph of the ph$$

From this point on, you can use the abbreviations proposed in the scheme above.

4. **Draw** the structure of compound **A**, including the appropriate stereochemistry.

		,			3
A					
5	Transformation of L-lysine into	A is (choo	se all annronri	ate answer(s))·	
<i>J</i> .	Transformation of E-tystic into	A 13 ( <u>choo</u>	<u>зе</u> ан арргоргі	ate answer(s)).	
	an enantioselective reaction.				
	1				
	a regioselective reaction.				

6. <b><u>Draw</u></b> the structures of compounds <b>B</b> - <b>F</b> , including the appropriate stereochemistry.					
$\mathbf{B} \ \mathrm{C}_{14} \mathrm{H}_{20} \mathrm{N}_2 \mathrm{O}_4$	$\mathbf{C}  \mathbf{C}_{16} \mathbf{H}_{21} \mathbf{NO}_{6}$				
D	$\mathbf{E}  \mathbf{C}_{29} \mathbf{H}_{34} \mathbf{N}_2 \mathbf{O}_6 \mathbf{S}$				
F C <sub>21</sub> H <sub>28</sub> N <sub>2</sub> O <sub>4</sub> S					
7. What is the role of DCC in the transformation	$\mathbf{C} \to \mathbf{D}$ ?				
<ul> <li>□ Protecting group for the amino group.</li> <li>□ Protecting group for the hydroxy group.</li> <li>□ Activating agent for the amide bond formation.</li> </ul>					
8. TsCl is used in the synthesis to enable:					
<ul> <li>□ Nucleophilic substitution of an amino group.</li> <li>□ Electrophilic substitution of an amino group.</li> <li>□ Nucleophilic substitution of a hydroxy group.</li> <li>□ Electrophilic substitution of a hydroxy group.</li> </ul>					

9. <b>Mark</b> all possible reagents which could be used a	as reagent H:				
□ diluted HCl	□ Zn/HCl				
$\square$ K <sub>2</sub> CO <sub>3</sub>	$\Box$ H <sub>2</sub> SO <sub>4</sub>				
☐ diluted KMnO <sub>4</sub>	☐ diluted NaOH				
$\square$ SOCl <sub>2</sub>	$\square$ PCl <sub>5</sub>				
10. <u>Draw</u> the structure of levobupivacaine, including the appropriate stereochemistry.					
Levobupivacaine C <sub>18</sub> H <sub>28</sub> N <sub>2</sub> O					
<b>-</b>					
<b>Part II.</b> The synthesis of levobupivacaine requires the use of enantiomerically pure L-lysine. A common method to confirm the enantiomeric purity of aminoacids is their transformation into amides using Mosher's acid (see the structure of the (S) isomer below).					
_0 C	F <sub>3</sub>				
HO					
∬ (S) `					
0					
(S)-Mosher	's acid				
11. <u>Draw</u> the structure of the amide formed when the α-amino group of L-lysine is derivatized with (S)-Mosher's acid. Clearly show the stereochemistry of each chiral center.					
12. <u>How many products</u> will be formed from racemic lysine and ( <i>S</i> )-Mosher's acid (consider that only the α-amino group of lysine is derivatized)?					
☐ Two diastereoisomers.					
☐ Four diastereoisomers.					
☐ A racemic mixture of two enantiomers.					
☐ Four compounds: two enantiomers and two diaste	reoisomers.				
13. <u>Choose</u> the method(s) which can be used to quallysine after its derivatization with (S)-Mosher's ac	* * *				
☐ NMR spectroscopy.					
☐ Liquid chromatography.					
☐ Mass spectrometry.					
iviass spectrometry.					