

48th International Chemistry Olympiad

Theoretical Problems

28 July 2016 Tbilisi, Georgia

Instructions

- Begin only when the START command is given. You have 5 hours to work on the problems.
- Use only the pen and calculator provided.
- The problem booklet has <mark>25</mark> pages, the answer sheet is <mark>29</mark> pages.
- Make sure that your code is on every page of the answer sheet.
- Questions are identical in the problem text and on the answer sheets.
- All results must be written in the appropriate boxes on the answer sheets. Anything written elsewhere will not be graded. Use the reverse of the problem pages if you need scratch paper.
- Write relevant calculations in the appropriate boxes when necessary. If you provide only correct end results for complicated questions, you will receive no score.
- Raise your hand if you need a restroom break.
- When you have finished the examination, put your answer sheets into the envelope provided. Do not seal the envelope.
- You can keep the problem booklet.
- You must stop your work immediately when the STOP command is given. A delay in doing this may lead to cancellation of your exam.
- Do not leave your seat until permitted by the supervisors.
- The official English version of this examination is available on request only for clarification.

Constants and formulae

Avogadro constant:	$N_{\rm A} = 6.022 \cdot 10^{23} {\rm mol}^{-1}$	Zero of the Celsius scale:	273.15 K
Gas constant:	$R = 8.314 \text{ J K}^{-1} \text{ mol}^{-1}$	Faraday constant:	F = 96485 C mol⁻¹
Ideal gas equation:	pV = nRT	Gibbs energy:	G = H - TS
$\Delta_r G^o = -RT lnK = -r$	nFE ^o _{cell}	Ionic product of water at 298.15K :	$K_w = 10^{-14}$
Nernst equation:	$E = E^{o} + \frac{RT}{nF} \ln \frac{c_{ox}}{c_{red}} = E^{o}$ or $E = E^{o} - \frac{RT}{nF} \ln Q = E$	$\frac{1}{r^o} + \frac{0.059 \text{ V}}{n} \log \frac{c_{ox}}{c_{red}}$ $\frac{1}{r^o} - \frac{0.059 \text{ V}}{n} \log Q$	<u>.</u>
Beer-Lambert law:	$A = \log \frac{I_0}{I} = \varepsilon c l$		

Consider all gases ideal throughout the exam.

Periodic table with relative atomic masses

1																	18
1																l	2
H	~											40		45	4.0	47	He
1.008	2	1										13	14	15	16	17	4.003
3	4											5	6	7	8	9	10
LI	Ве											B	C	IN	O	F	INE
6.94	9.01	-										10.81	12.01	14.01	16.00	19.00	20.18
	12											13	14 C:	15 D	16 C		18 A m
INa	IVIG	З	Λ	5	6	7	8	٩	10	11	12		30	P	32.06		
22.99	24.30	21	-	22	24	7	26	27	20	20	12	20.90	20.09	30.97	32.00	35.45	39.95
19		Ŝ	Ti	23	$\hat{\mathbf{C}}_{\mathbf{r}}$	Mn	Ē		20 NIi		- 30 7n	Go			34 So	30 Dr	30 Kr
IN 39.10	40.08	44 96	47 87	V 50.94	52.00	54 94	1 C	58.93	1 NI 58 69	63 55	65.38	69.72	72.63	74 92	78 97	79 90	1 NI 83 80
37		39	40	41	42	43	44	45	46	47	48	49	50	51	52	53	54
Rh	Sr	Ŷ	7r	Nb	Mo	Tr	Ru	Rh	Рď	DΑ	Сd	In	Sn	Sh	Te	Ĩ	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	Ha	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		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	Ρm	Sm	Eu	Gd	Tb	Dy	Ho	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	۱h	Ра	U	Np	Pu	Am	Cm	Bk	Ct	Es	⊦⊢m	Md	No	Lr
			-	232.0	231.0	238.0	-	-	-	-	-	-	-	-	-	-	-

Problem 1 5% o

5% of the total

Nitrogen trifluoride is a surprisingly stable compound that was first prepared in the melt electrolysis of a mixture of ammonium fluoride and hydrogen fluoride.

1.1. <u>On which electrode</u> does nitrogen trifluoride form? <u>Write</u> a balanced chemical equation for the electrode half reaction for the formation of NF₃.

Interestingly the related fluoroamine (NH₂F) and difluoroamine (NHF₂) are very unstable materials; decomposition of either pure substance can even be explosive. This is dangerous as they are formed in the electrolysis as side products.

1.2. Which of NF₃, NHF₂ or NH₂F compound is expected to condense at the lowest temperature?

The N-F bond lengths in these molecules were determined to be 136, 140 and 142 pm. The change in the bond lengths can be explained with a simple electrostatic model taking into account the partial charges on the atoms.

1.3. <u>Assign</u> the N-F bond lengths (136, 140, 142 pm) to the molecules.

When NHF₂ is bubbled through a solution of KF in HF, a binary nitrogen – fluorine compound can be obtained as a mixture of two geometric isomers.

1.4. <u>Write</u> a balanced chemical equation for the formation of the binary nitrogen-fluorine compound.

Tetrafluoroammonium ion (NF $_4$ ⁺) and its corresponding salt can form from NF $_3$ with elementary fluorine in the presence of an appropriate reagent.

1.5. <u>Propose</u> a suitable reagent and <u>write</u> a balanced chemical equation for the reaction.

 NF_{4^+} ions form stable salts with a number of anions. These are very sensitive to humidity, because NF_{4^+} ion hydrolyzes forming NF_3 and O_2 . Interestingly nitrogen trifluoride always forms quantitatively, while the quantity of oxygen is often less than expected due to side reactions.

1.6. <u>Write</u> a balanced chemical equation for the hydrolysis of NF₄⁺ <u>Write</u> a balanced chemical equation for a possible side reaction that can decrease the theoretically expected O₂:NF₃ mole ratio.

Tetrafluoroammonium salts were investigated for use as solid rocket fuels, because NF₃ and F₂ are released from them on heating. One of them has a fluorine content of 65.6 m/m%, all of which is converted into NF₃ and F₂ upon decomposition. During the decomposition 2.5 times as many moles of F₂ are formed as of NF₃.

1.7. <u>Determine</u> the formula of the salt in question.

8% of the total

One of the first materials used in solid state electronics was red copper(I) oxide. Interest is renewed nowadays because it could be a non-toxic and cheap component of solar cells.



The two figures above depict the cubic unit cell of the Cu_2O crystal. The lattice constant of the structure is 427.0 pm.

2.1.1. <u>Which of the atoms (A or B) is copper?</u> <u>Which basic structure (primitive cubic, face centered cubic, body centered cubic, bo</u>

diamond) is formed by the A atoms and <u>which structure</u> is formed by the B atoms?

What are the coordination numbers of the atoms?

- 2.1.2. <u>Calculate</u> the smallest 0-0, Cu-0 and Cu-Cu distances in the structure?
- **2.1.3.** <u>What</u> is the density of pure copper(I) oxide?

A common defect in this crystal is some copper atoms missing with the oxygen lattice unchanged. The composition of one such crystal sample was studied, and 0.2% of all copper atoms were found to be in oxidation state +2.

2.2. What percentage of normal copper sites are empty in the crystal sample? What is x in the empirical formula Cu_{2-x}O of the crystal?

Copper(I) oxide is insoluble in water. It is stable in dry air, but humidity in the air catalyzes a transformation (Reaction 1)

When copper(I) oxide is dissolved in dilute sulfuric acid, a blue solution containing a precipitate is formed without evolution of a gas (Reaction 2). When hot, concentrated sulfuric acid is used, no precipitate remains, but an odorous gas forms (Reaction 3). The same gas forms when the precipitate from reaction 2 is dissolved in hot concentrated sulfuric acid.

2.3. <u>Write</u> balanced chemical equations for reactions (1-3).

Copper (I) oxide can be produced in a number of ways. Heating copper in air is a common method in the synthesis of semiconductor Cu_2O . In a pure oxygen atmosphere, the three species containing copper (Cu(s), $Cu_2O(s)$ or CuO(s)) can potentially interconvert.

Suppose that the $\Delta_f H^0$ and S^0 data given for 10^5 Pa are independent of temperature:

	$\Delta_{\rm f} H^{\rm o}$ / kJ mol ⁻¹	<i>S</i> °/ J mol ⁻¹ K ⁻¹
Cu(s)	0	65
02(g)	0	244
CuO(s)	-156	103
$Cu_2O(s)$	-170	180

2.4. <u>Determine</u> the temperature ranges, if any, of thermodynamic stability of copper and its oxides between 500 and 1500 K in a 10⁵ Pa oxygen atmosphere.

Important data are given for 298 K. Use this temperature in the following calculations:

 $\begin{array}{ll} K_{\rm sp}({\rm Cu}({\rm OH})_2) = 2 \cdot 10^{-19} \\ {\rm Cu}_2{\rm O}({\rm s}) + {\rm H}_2{\rm O}({\rm l}) + 2{\rm e}^- \longrightarrow 2{\rm Cu}({\rm s}) + 2{\rm OH}^-({\rm aq}) \\ {\rm Cu}^{2+}({\rm aq}) + {\rm e}^- \longrightarrow {\rm Cu}^+({\rm aq}) \\ {\rm Cu}^{2+}({\rm aq}) + 2{\rm e}^- \longrightarrow {\rm Cu}({\rm s}) \\ \end{array} \qquad \begin{array}{ll} E^{\rm o} = -0.360 \ {\rm V} \\ E^{\rm o} = +0.159 \ {\rm V} \\ E^{\rm o} = +0.337 \ {\rm V} \end{array}$

One possibility for producing Cu₂O is the anodic oxidation of copper. Electrolysis of an aqueous basic solution (e.g. NaOH) with a copper anode and platinum cathode can lead to formation of copper(I) oxide on the anode.

2.5. <u>Write</u> the half reaction equations for the electrode processes during the anodic production of Cu₂O in NaOH solution with a platinum cathode and copper anode.

Electrolytic reduction of copper(II) ions in solution is another possibility.

2.6.1. Write the half reaction equation of the cathode process giving Cu₂O in acidic medium.

Let us use 0.100 mol dm $^{\rm -3}$ Cu $^{\rm 2+}$ solution and carry out electrolysis with platinum electrodes.

2.6.2. What is the maximum pH of this solution at which the concentration of copper(II) can be maintained at 0.100 mol dm⁻³?

If the pH is too low, reduction to metallic copper is preferred to the formation of copper(I) oxide.

2.6.3. What is the minimum pH at which the cathodic production of Cu_2O in a 0.100 mol dm⁻³ Cu^{2+} solution is still possible?

9% of the total

Iodine deficiency is of special concern in Georgia because it occupies a region where iodine is scarce in soil and water. Iodine deficiency can be effectively and inexpensively prevented if salt for human consumption is fortified with small amounts of iodine. Methods for analyzing salt for iodine content are thus important. Current regulations in Georgia are that iodized salt must contain between 25-55 ppm iodine (1 ppm = 1 mg iodine/kg salt).

Most salt is iodized by fortification with potassium iodate (KIO₃). Iodate content can be determined in salt samples using iodometric titration. In a typical procedure, 10.000 g of an iodized salt sample is dissolved in 100 cm³ of 1.0 mol/dm³ aqueous HCl to which 1.0 g KI has been added. The solution is then titrated with 0.00235 mol/dm³ aqueous sodium thiosulfate solution to a starch endpoint; this requires 7.50 cm³ of titrant.

- **3.1.1.** <u>Write</u> a balanced net ionic equation for the reaction when iodate reacts with excess iodide in acidic solution.
- **3.1.2.** <u>Write</u> a balanced net ionic equation for the reaction taking place during the titration with thiosulfate.
- **3.1.3.** <u>Calculate</u> the iodization level, in ppm, of this salt sample.

A less common agent for iodizing salt is potassium iodide, which cannot be easily measured by iodometric titration.

One possible method for analyzing iodide in the presence of chloride is potentiometric titration. However, this method is not very precise in the presence of large amounts of chloride.

In this method, a silver wire is immersed in the solution (containing iodide and chloride) to be analyzed and silver ion is gradually added to the solution. The potential of the silver wire is measured relative to a reference electrode consisting of a silver wire in a 1.000 mol/dm³ solution of AgNO₃. The measured potentials are negative and the absolute values of these potentials are reported. The solution to be analyzed has a volume of 1.000 dm³ (which you may assume does not change as silver ion is added), and $T = 25.0^{\circ}$ C.

The results of this experiment are governed by three equilibria: the solubility of AgI(s) $[K_{spI}]$ and AgCl(s) $[K_{spCl}]$ and the formation of AgCl₂-(aq) $[K_f]$. (Iodide also forms complex ions with silver but this may be neglected at the very low concentrations of iodide present in this experiment).

$AgI(s) \rightleftharpoons Ag^{+}(aq) + I^{-}(aq)$	KspI
$AgCl(s) \rightleftharpoons Ag^{+}(aq) + Cl^{-}(aq)$	KspCl
$Ag^{+}(aq) + 2 Cl^{-}(aq) \rightleftharpoons AgCl_{2}^{-}(aq)$	Kf

Below are shown the results of two experiments measuring the observed potential as a function of added number of moles of silver ion. Experiment **A** (solid circles) was carried out with 1.000 dm³ of solution containing $1.00 \cdot 10^{-5}$ mol/dm³ iodide and no chloride ion. Experiment **B** (open circles) was done using 1.000 dm³ of solution containing $1.00 \cdot 10^{-5}$ mol/dm³ iodide and $1.00 \cdot 10^{-1}$ mol/dm³ chloride.



- **3.2.1.** Select an appropriate data point from the experiments and use it to <u>calculate</u> the solubility product of AgI (K_{spl}).
- **3.2.2.** Select an appropriate data point from the experiments and use it to <u>calculate</u> the solubility product of AgCl (K_{spCl}).
- **3.2.3.** Select an appropriate data point from the experiments and use it to <u>calculate</u> $K_{\rm f}$. You may need to use values of $K_{\rm spl}$ or $K_{\rm spCl}$ to do this calculation. If you were unable to carry out the calculations in **3.2.1**. or **3.2.2**., you may use the arbitrary values of $K_{\rm spl} = 1.00 \cdot 10^{-15}$ and $K_{\rm spCl} = 1.00 \cdot 10^{-9}$ without penalty.

An analytical method that is more practical, because it is not sensitive to the presence of chloride, uses the Sandell-Kolthoff reaction. This is the reaction of H_3AsO_3 with Ce(IV) to give Ce(III) in acidic solution, which is strongly catalyzed by iodide ion.

3.3.1. <u>Write</u> balanced net ionic equations for the reaction of cerium(IV) with H₃AsO₃ in acidic solution, as well as reactions of cerium(IV) with a species containing the element iodine and H₃AsO₃ with a species containing the element iodine, that could reasonably account for the catalysis of the net reaction by iodide.

The reaction of Ce(IV) with H_3AsO_3 can be monitored by measuring the absorbance at 405 nm, as Ce(IV) is orange and absorbs significantly at 405 nm, while the other reactants and products are colorless and do not absorb appreciably. Three runs were carried out, all in 0.50 mol/dm³ H_2SO_4 at 25.0°C using the following initial concentrations:

Run	[H3AsO3]0,	[Ce(IV)]0,	[I-]0,
	mol dm ⁻³	mol dm ⁻³	mol dm ⁻³
1	0.01250	0.00120	$1.43 \cdot 10^{-6}$
2	0.00625	0.00060	$1.43 \cdot 10^{-6}$
3	0.01250	0.00120	7.16.10-7

An analyst initiated the reactions by mixing the reagents in a cuvette. After a short variable delay absorbance measurements were started, with the first measurement recorded at t=0 s. The data obtained are shown below:



t c	A_{405}	A_{405}	A_{405}
ι, s	Run 1	Run 2	Run 3
0	0.621	0.287	0.818
20	0.348	0.149	0.608
40	0.198	0.083	0.455
60	0.113	0.046	0.340
80	0.064	0.025	0.254
100	0.037	0.014	0.191

Under these conditions (0.5 mol/dm 3 H₂SO₄, 25.0°C), the rate law for the reaction can be written as

Rate = $k[H_3AsO_3]^m[Ce(IV)]^n[I^-]^p$

where *m*, *n*, and *p* are integers.

3.3.2. Determine the values of *m*, *n*, and *p* and <u>calculate</u> the value of *k* (be sure to specify its units).

A 1.000 g sample of iodized salt is dissolved in water to give 10.00 cm^3 of solution. A 0.0500 cm^3 aliquot of this solution is added to a mixture of 1.000 cm^3 0.025 mol/dm^3 H₃AsO₃ in 0.5 mol/dm³ H₂SO₄ and 0.800 cm³ 0.5 mol/dm³ H₂SO₄. To this mixture is added 0.200 cm³ 0.0120 mol/dm³ Ce(NH₄)₂(NO₃)₆ in 0.5 mol/dm³ H₂SO₄ and the absorbance at 405 nm is measured as a function of time at 25.0°C:



t, s	A_{405}
0	0.756
20	0.531
40	0.373
60	0.262
80	0.185
100	0.129

3.3.3. <u>Calculate</u> the iodization level, in ppm, of this salt sample.

8% of the total

Application of kinetic studies in water treatment

Industrial waste is a major cause of water pollution and kinetic studies are carried out in a laboratory to design effluent treatment. 1,4-dioxane, more commonly known as dioxane (C₄H₈O₂), an industrial solvent and by-product, is a significant water contaminant. It can be oxidised to hazard free chemicals using oxidants such as peroxodisulfate, ozone or hydrogen peroxide.

The data obtained in the kinetic study of oxidation of dioxane with potassium peroxodisulfate ($K_2S_2O_8$) as oxidant and AgNO₃ as catalyst at *T* = 303.15 K are given below. The reaction was monitored by the estimation of unreacted peroxodisulfate. The concentration of AgNO₃ used in this study was $1.00 \cdot 10^{-3}$ mmol·dm⁻³.

Trial	Dioxane	$K_2S_2O_8$	Initial rate
	mmol·dm ⁻³	mmol·dm ⁻³	mmol·dm ⁻³ ·min ⁻¹
1	0.0100	2.50	1.661.10-2
2	0.0100	5.10	3.380.10-2
3	0.00500	13.8	9.200·10 ⁻²
4	0.0110	13.8	9.201·10 ⁻²

In many countries the accepted maximum level of dioxane in drinking water is specified as 0.35 $\mu g \ dm^{-3}.$

A water sample contains an initial dioxane concentration of 40.00 μ g dm⁻³. Assume that 1 mol dioxane requires 1 mol of peroxodisulfate for oxidation. The concentration of AgNO₃ used in this study was 1.00·10⁻³ mmol·dm⁻³.

4.1.1. <u>Calculate</u> the time in minutes the oxidation process has to continue in order to reach the accepted level of dioxane at 303.15 K if the initial concentration of $K_2S_2O_8$ is $5.0 \cdot 10^{-6}$ mol dm⁻³. Assume that the rate law obtained from the data above is valid under these conditions.

Various mechanisms have been proposed for the peroxodisulfate oxidation of dioxane. Misra and Ghosh (1963) proposed the following mechanism:

$$S_{2}O_{8}^{2-} + Ag^{+} \stackrel{k_{1}}{\rightleftharpoons} Ag^{3+} + 2SO_{4}^{2-}$$

$$k_{2}$$

$$Ag^{3+} + D (dioxane) \stackrel{k_{3}}{\longrightarrow} D' (dioxane oxidised) + 2H^{+} + Ag^{+}$$

4.1.2. Assuming Ag(III) to be in steady state, <u>deduce</u> the rate equation for the oxidation of dioxane.

4.1.3. Which of the following is/are correct?

A) The rate equation based on the mechanism given in **4.1.2**, at very high concentrations of dioxane, is consistent with the experimental data in **4.1.1**. B) The rate equation based on the mechanism given in **4.1.2**, at very low concentrations of dioxane, is consistent with the experimental data in **4.1.1**. C) The units of the observed rate constant are dm³·mol⁻¹·s⁻¹ at very high concentrations of dioxane.

D) The units of the observed rate constant are $dm^3 \cdot mol^{-1} \cdot s^{-1}$ at very low concentrations of dioxane.

Degradation of pharmaceutical products - a kinetic overview

Kinetic studies are important in deciding the shelf life of a pharmaceutical product. Several chemical reactions can affect the shelf life of pharmaceutical products and the rates of these reactions depend on conditions such as pH, temperature, humidity.

Lysine acetylsalicylate (LAS) is prescribed as a pain killer and anti-inflammatory drug under the brand name Aspegic. LAS on hydrolysis forms lysine salicylate and acetic acid.



Hydrolysis of LAS can proceed via three different pathways (a) acid catalysed, (b) uncatalysed and (c) base catalysed.

If [LAS] denotes the concentration of LAS at time '*t*', the overall rate of the hydrolysis reaction can be written as

$$-\frac{d[LAS]}{dt} = k_{H}[LAS][H^{+}] + k_{0}[LAS] + k_{OH}[LAS][OH^{-}]$$

where $k_{\rm H}$, k_0 and $k_{\rm OH}$ are the rate constants of the acid catalysed, uncatalysed and base catalysed pathways of hydrolysis, respectively. The overall rate constant is defined as:

$$-\frac{d[\text{LAS}]}{dt} = k_{\text{obs}}[\text{LAS}]$$

4.2.1. <u>Write</u> an expression for k_{obs} in terms of k_H , k_0 , k_{OH} and $[H^+]$.

Hydrolysis of LAS was carried out at 298.15 K at various pH values (0.50 to 13.0). A very low initial concentration of LAS ensured that the pH did not change during the course of the reaction.

The following graph shows the pH dependence of the hydrolysis of LAS.



- 4.2.2. Which of the following is/are correct?
 A) k_{obs} ≈ k₀ at pH = 12
 B) k_{obs} ≈ k₀ at pH = 5.0
 C) The rate of the reaction increases when the pH is changed from 0.50 to 1.0.
 D) The rate of the reaction increases when the pH is changed from 10 to 12.
- **4.2.3.** Using the diagram and the data given below, <u>calculate</u> *k*_H, *k*₀ and *k*_{OH}. Make sure to specify the units.

рН	1.300	5.300	12.180
$\log (k_{\rm obs}/{\rm min}^{-1})$	-3.886	-4.000	-1.726

Acetylsalicylic acid, more commonly known as aspirin is a medicine often used for reducing fever, pain and inflammation. Like LAS, the hydrolysis of aspirin can also take different pathways depending on the pH. The pH rate profile of aspirin hydrolysis at **333.15 K** is given below:



The following are possible reactions for the hydrolysis of aspirin. Depending on the pH, one or more of these reactions will predominate.

- I. $CH_3COOC_6H_4COOH + H_3O^+ \rightarrow HOC_6H_4COOH + CH_3COOH + H^+$
- II. $CH_3COOC_6H_4COOH + H_2O \rightarrow HOC_6H_4COOH + CH_3COOH$
- III. $CH_3COOC_6H_4COOH + OH^- \rightarrow HOC_6H_4COOH + CH_3COO^-$
- IV. $CH_3COOC_6H_4COO^- + H_3O^+ \longrightarrow HOC_6H_4COOH + CH_3COOH$
- V. $CH_3COOC_6H_4COO^- + H_2O \rightarrow HOC_6H_4COOH + CH_3COO^-$
- VI. $CH_3COOC_6H_4COO^- + OH^- \rightarrow HOC_6H_4COO^- + CH_3COO^-$

4.3.1. Using the pH-rate profile diagram and the reactions given above, <u>state</u> which of the following statements is/are correct. (pK_a of aspirin = 3.57 at 333.15 K)

a) In the region C-D, reaction IV is predominant

b) In the region C-D, reaction V is predominant

c) In the region D-E reaction VI is predominant

d) In the region A-B, reaction II is predominant

The plot of k_{obs} vs pH for the hydrolysis of aspirin has been confirmed to show a minimum at a particular pH. At **290.15 K** the following rate constants for reactions I, II and III were determined:

<i>k</i> н (reaction I)	k_0 (reaction II)	<i>k</i> он (reaction III)
1.99 dm ³ ·mol ⁻¹ ·day ⁻¹	2.29·10 ⁻⁴ day ⁻¹	$3.18\cdot10^9 \mathrm{dm^3\cdot mol^{-1}\cdot day^{-1}}$

The ionic product of water at 290.15 K can be taken as $1.0 \cdot 10^{-14}$.

4.3.2. Assuming that only reactions I, II and III occur, <u>calculate</u> the value of the pH at the minimum.

8% of the total

5500 years ago in ancient Egypt people learned for the first time how to synthesize a blue pigment. Now we know this pigment as Egyptian blue. About 2000 years later in ancient China another pigment was widely used, which is now referred to as Chinese blue. The two pigments are similar in structure, but have different elemental compositions.

The ancient method of preparation for these pigments can be easily reproduced in a modern laboratory.

When considering the amounts, assume that all of the compounds in this task are pure, and the yields are quantitative.

To make **Egyptian blue**, one should heat 10.0 g of mineral **A** with 21.7 g of SiO₂ and 9.05 g of mineral **B** at 800–900°C for a prolonged time. 16.7 dm³ of a mixture of two gaseous products are released (the volume is measured at 850°C and $1.013 \cdot 10^5$ Pa (1.013 bar) pressure. In result, 34.0 g of the pigment was obtained. No other products are formed. As the gas mixture is cooled, one component of the mixture condenses. As the remaining gas is further cooled to 0°C, the gaseous volume reduces to 3.04 dm³.

- **5.1.1.** <u>Find</u> the mass of the gaseous mixture formed upon heating of **A** with **B** and SiO₂.
- **5.1.2.** <u>Determine</u> the quantitative composition of this gas mixture.

When 10.0 g of mineral **A** is heated with 21.7 g of SiO₂ in the absence of **B**, it forms 8.34 dm³ of gaseous products (measured at 850°C and $1.013 \cdot 10^5$ Pa = 1.013 bar pressure). Mineral **A** contains only one metal.

5.1.3. <u>Calculate</u> the molar mass and determine the formula of mineral **B**. Hint: it is an ionic solid insoluble in water and containing no water of crystallization.

In order to obtain **Chinese blue**, one should take 17.8 g of mineral **C** instead of mineral **B** (keeping the amounts of mineral **A** and SiO_2 same as for Egyptian blue), and run the reaction at higher temperatures. Besides the pigment, the same gaseous products in the same quantities are formed as in the preparation of Egyptian blue.

5.1.4. <u>Determine</u> the formula of mineral **C**.

- **5.1.5.** <u>Determine</u> the formulae of Egyptian blue and Chinese blue.
- **5.1.6.** <u>Determine</u> the formula of mineral **A**.

Elemental analysis of some samples of Chinese blue shows traces of sulfur. This led to a conclusion that those were synthesized using another common mineral instead of **C**.

- **5.2.1.** <u>Suggest</u> a formula for the mineral used in place of **C**.
- **5.2.2.** <u>Could</u> the temperature of synthesis of Chinese blue be decreased if this mineral is used instead of **C**?

If during the synthesis of Chinese blue we take a smaller amount of silica than in the process above, we will obtain a purple pigment: Chinese violet. It was used, in particular, for coloring the famous Terracotta army soldiers.

5.3. <u>Write down</u> the formula of a binary compound that forms under the conditions required for Chinese violet and is responsible for the change of the color.

Problem 6 7% of the total

Although there is currently no known cure for Alzheimer's disease, there are medications available to manage the neurodegenerative disorder. Among these are acetylcholinesterase inhibitors, of which galantamine **1** is an example. This molecule can be isolated from the Caucasian snowdrop, a plant native to Georgia; however, the large amounts needed for therapy require a synthetic route. Shown below is the route used to prepare galantamine industrially.



Notes about the synthesis:

- ¹H NMR of **A** indicates 2 aromatic protons in a *para* arrangement.
- **C** is labile in aqueous conditions, so it is not isolated, but rather reacted immediately with NaBH₄ to convert it to **D**.
- **6.1.1.** <u>Suggest</u> structures for **A**, **B**, **C**, **D**, **F**, and **G**. None of the reactions except for the final transformation with L-selectride are stereoselective. Therefore, stereochemistry does not need to be indicated in your answers.

6.1.2. <u>Give</u> the formula for a possible reagent, **X**, to convert compound **D** to **E**.

The optical rotation of the material obtained by resolution was -400° cm² g⁻¹, while that of the enantiomerically pure compound is -415° cm² g⁻¹ when measured under the same conditions. You may assume that the only optical impurity is the other enantiomer.

One way of describing optical purity is enantiomeric excess (*ee*). It is defined as the difference in the percentages of the enantiomers in a mixture. For example in a mixture of 70% R and 30% S, the *ee* is 40%.

6.2.1. <u>What</u> is the enantiomeric excess of the resolved compound as prepared by the industrial route?

L-selectride is a commercial reagent that performs the final reaction stereoselectively.



- **6.3.1.** <u>Give</u> the formula for compound **Y** to carry out the first step of the route.
- 6.3.2. <u>Suggest</u> structures for J and K.

8% of the total

This question looks at the synthesis of *dolasetron mesylate*, **Z** (shown right), a drug sold under the tradename *Anzemet* and used to treat post-operative nausea and vomiting.

The synthesis begins as shown below.



LiH (2.5 eqiv) NaOH(aq) Β Α 2. HCl(aq), Δ C_2H_5O C_2H_5 CI CI (1.1 eqiv) SOCl₂ E1 + E2 HIO₄(aq) OsO₄(aq) C₂H₅OH O_3 (+ reductive workup) G D С C 55.81% H 7.03% O 37.17% HIO₄(aq) F1 + F2он $(in CH_2Cl_2)$ ċι

First cyclic compound **A** is made, which contains C, H, and O only. Compound **G** is achiral and can be prepared directly from **D** using ozone under reductive conditions, or via stereoisomers **E1** and **E2** using OsO₄, or via stereoisomers **F1** and **F2** using the peracid shown.

- **7.1.** <u>Determine</u> the empirical formula of **G** from the percentage masses given.
- **7.2.** <u>Give</u> the structures of **A**, **B**, **C**, **D**, **E1**, **E2**, **F1**, **F2** and **G**.

Compound **G** is used in the next stage of the synthesis, under buffered conditions, to form **H** (as a mixture of two achiral diastereoisomers). Reduction of **H** with NaBH₄ gives alcohol **I** (as a mixture of four achiral diastereoisomers). **I** reacts with acidified dihydropyran to form **J** (as a mixture of even more diastereoisomers). **J** is then treated first with *t*-butoxide base, then refluxed with acid before finally extracting under weakly basic conditions to form **K** as a mix of two diastereomers, **K1** (major product) and **K2** (minor product). These could be separated, and **K1** was used in the final stages of the synthesis.



- **7.3.1.** <u>Give</u> the structures of **H**, **I**, and **J**. There is no need to show the different diastereoisomers formed.
- **7.3.2.** Give the structures of diastereoisomers **K1**, and **K2**.

In the final stage of the synthesis, **L** and **M** react to form intermediate **N**. **N** then reacts with **K1** to form, after extraction, the neutral amine which gives the target compound upon protonation with CH_3SO_3H .



7.4. <u>Give</u> the structure of **N**.

7% of the total

An exotic, but biologically relevant sugar analogue can be prepared from D-glucose in the following manner. Heating a mixture of D-glucose and acetone with a few drops of concentrated acid results in the formation of a diacetonide **A**. Then **A** can be hydrolyzed selectively to **B**.



8.1. <u>Give</u> your chosen answers to the questions on the answer sheet.

B is treated with sodium metaperiodate to get **C**. **C** is then reacted with an aqueous solution of NaCN, then heated with 10% NaOH solution to get a mixture of two diastereomeric compounds D_1 and D_2 . These compounds can be separated by column chromatography.



Reaction of D_1 with LiAlH₄ followed by heating with 1M HCl solution gives sugar **F** that is the hydrolysis product of the most abundant natural polysaccharide.

	LiAlH ₄			1M HCl/H ₂ O		
\mathbf{D}_1		\longrightarrow	E		\rightarrow	F

- B.2.1. Draw the structures of C, D₁, D₂, E and F including stereochemical information. Show F as the more stable 6-membered ring containing isomer using the ring skeleton. Indicate with a wavy line if absolute chirality around a carbon is not known.
- **8.2.2.** <u>Give</u> your chosen answer to the question on the answer sheet.

Neutralization of D_2 with HCl followed by heating in toluene results in dehydration and formation of **G**, which has a tricyclic structure in water-free solvents. Boiling **G** in 1M HCl solution gives **H** (C₆H₁₀O₇), which is a natural sugar derivative containing a 6 membered ring. **H** is a building block of heparin, an anticoagulant polysaccharide produced by our bodies.

	1. Equimolar HCl		
	2. Heat, toluene	1M HCl/H ₂ O	
D ₂	\longrightarrow G	\longrightarrow	Н
	-H ₂ O		

- B.3.1. <u>Draw</u> the structure of **G** including the stereochemistry. <u>Draw</u> **H** as the more stable 6-membered ring containing isomer using the ring skeleton. <u>Indicate</u> with a wavy line if absolute chirality around a carbon is not known.
- **8.3.2.** <u>Give</u> your chosen answer to the question on the answer sheet.
- **8.3.3.** <u>Give</u> your answer to the question on the answer sheet.



48th International Chemistry Olympiad

Theoretical Problems Answer sheets

28 July 2016 Tbilisi, Georgia

Problem 1 5% of the total

1.1.	1.2.	1.3.	1.4.	1.5.	1.6.	1.7.	Sum
3	1	2	2	2	4	3	17

1.1. <u>On which electrode</u> does nitrogen trifluoride form?

Cathode Anode

<u>Write a balanced chemical equation for the electrode half reaction for the formation of NF3.</u>

1.2.	<u>Which of NF₃, NHF₂ or NH₂F compound is expected to condense at the lowest</u>
	temperature?

NF3

NHF2 NH2F

1.3. <u>Assign</u> the N-F bond lengths (136, 140, 142 pm) to the molecules.

Molecule	NH ₂ F	NHF ₂	NF3
N-F bond			
length, pm			

- **1.4.** <u>Write</u> a balanced chemical equation for the formation of the binary nitrogen fluorine compound.
- **1.5.** <u>Propose</u> a suitable reagent for the formation of NF₄⁺ and <u>write</u> a balanced chemical equation for the reaction.

1.6. <u>Write</u> a balanced chemical equation for the hydrolysis of NF₄⁺

<u>Write</u> a balanced chemical equation for a possible side reaction that can decrease the theoretically expected O_2 :NF₃ mole ratio.

1.7. <u>Determine</u> the formula of the salt in question.

Your work:

Formula:

8% of the total

2.1.1	2.1.2	2.1.3	2.2	2.3	2.4	2.5	2.6.1	2.6.2	2.6.3	Sum
5	3	2	2	3	6	2	1	3	6	33

2.1.1. <u>Which</u> of the atoms (A or B) is copper?

B:

Cu:

<u>Which basic structure</u> (primitive cubic, face centered cubic, body centered cubic, diamond) is formed by the A atoms and <u>which structure</u> is formed by the B atoms?

	pr. cubic	fcc	bcc	diamond
А				
В				

<u>Calculate</u> the coordination numbers of the atoms?

Δ.		
41.		

2.1.2. <u>What</u> are the smallest 0-0, Cu-0 and Cu-Cu distances in the structure?

Your work: O-O: Cu-O: Cu-Cu:

2.1.3. <u>What</u> is the density of pure copper(I) oxide?

Your work:

density:

2.2. What percentage of normal copper sites are empty in the crystal sample?

Your work:

percentage:

What is *x* in the empirical formula $Cu_{2-x}O$ of the crystal?

Your work:

x:

2.3. <u>Write</u> balanced chemical equations for reactions (1-3).

Reaction 1:

Reaction 2:

Reaction 3:

2.4. Determine the temperature ranges, if any, of thermodynamic stability of copper and its oxides between 500 and 1500 K in a 10⁵ Pa oxygen atmosphere.

Your work:

Temperature range between 500- 1500K	The most stable form (Cu, Cu2O or CuO)

2.5. <u>Write</u> the half reaction equations for the electrode processes during the anodic production of Cu₂O in NaOH solution with a platinum cathode and copper anode.

Cathode:			
Anode:			

- **2.6.1.** <u>Write</u> the half reaction equation of the cathode process giving Cu₂O in acidic medium.
- **2.6.2.** What is the maximum pH at which the concentration of copper(II) can be maintained at 0.100 mol dm⁻³?

Your work:

maximum pH:

2.6.3. What is the minimum pH at which the cathodic production of Cu_2O in a 0.100 mol dm⁻³ Cu²⁺ solution is still possible?

Your work:

minimum pH:

Problem 3 9% of the total

3.1.1.	3.1.2.	3.1.3.	3.2.1.	3.2.2.	3.2.3.	3.3.1.	3.3.2.	3.3.3.	Sum
2	2	6	5	4	7	6	11	5	48

- **3.1.1.** <u>Write</u> a balanced net ionic equation for the reaction when iodate reacts with excess iodide in acidic solution.
- **3.1.2.** <u>Write</u> a balanced net ionic equation for the reaction taking place during the titration with thiosulfate.

3.1.3. <u>Calculate</u> the iodization level, in ppm, of the salt sample.

Your work:

ppm iodine =

3.2.1. Select an appropriate data point from the experiments and use it to calculate the solubility product of AgI (*K*_{spl}).

Your work:		
Ksnl:		

3.2.2. Select an appropriate data point from the experiments and use it to calculate the solubility product of AgCl (K_{spCl}).

Your work:

K_{spCl}:

3.2.3. Select an appropriate data point from the experiments and use it to calculate $K_{\rm f}$. You may need to use values of $K_{\rm spl}$ or $K_{\rm spCl}$ to do this calculation. If you were unable to carry out the calculations in **3.2.1**. or **3.2.2**., you may use the arbitrary values of $K_{\rm spl} = 1.00 \cdot 10^{-15}$ and $K_{\rm spCl} = 1.00 \cdot 10^{-9}$ without penalty.

values of Kspi – 1.00 10	without penalty.	
Your work:		
Kf:		

3.3.1. <u>Write</u> balanced net ionic equations for the reaction of cerium(IV) with H₃AsO₃ in acidic solution, as well as reactions of cerium(IV) with a species containing the element iodine and H₃AsO₃ with a species containing the element iodine, that could reasonably account for the catalysis of the net reaction by iodide.

Net reaction of cerium(IV) with H₃AsO₃ in acidic solution:

Reaction of cerium(IV) with an iodine-containing species:

Reaction of H_3AsO_3 with an iodine-containing species:

3.3.2. Determine the integer values of *m*, *n*, and *p* and <u>calculate</u> the value of *k* (be sure to <u>specify</u> its units).

Your work:

m = n = p = k =

3.3.3. <u>Calculate</u> the iodization level, in ppm, of the salt sample.

Your work:

ppm I =

8% of the total

4.1.1	4.1.2	4.1.3	4.2.1	4.2.2	4.2.3	4.3.1	4.3.2	Sum
5	3	2	1	2	6	2	4	25

4.1.1. <u>Calculate</u> the time in minutes the oxidation process has to continue in order to reach the accepted level of dioxane at 303.15 K if the initial concentration of $K_2S_2O_8$ is $5.0 \cdot 10^{-6}$ mol/dm³. Assume that the rate law obtained from the data above is valid under these conditions.

Your work:

Oxidation time:

4.1.2. Assuming Ag(III) to be in steady state, deduce the rate equation for the oxidation of dioxane.

Your work:

4.1.3.

4.2.1. Write an expression for k_{obs} in terms of k_H , k_0 , k_{OH} and $[H^+]$.

Your work:

- 4.2.2.
- **4.2.3.** Using the diagram and the data in the table, <u>calculate</u> $k_{\rm H}$, k_0 and $k_{\rm OH}$. Make sure to specify the units.

Your work:

7	7	7	
K0 <u>:</u>	<i>К</i> он <u>:</u>	Кн:	
	—		
4.3.1.			

4.3.2. Assuming that only reactions I, II and III occur, <u>calculate</u> the value of the pH at the minimum of k_{obs} .

Your work:

pH at minimum:

Problem 5 8% of the total

5.1.1.	5.1.2.	5.1.3.	5.1.4.	5.1.5.	5.1.6.	5.2.1.	5.2.2.	5.3.	Sum
1	3	2	2	5	2	1	1	2	19

5.1.1. <u>Find</u> the mass of the gaseous mixture formed upon heating of **A** with **B** and SiO₂.

Your work:

5.1.2. <u>Determine</u> the quantitative composition of this gas mixture.

Your work:

The gaseous mixture formed at 850°C contains mol of and mol of

5.1.3. <u>Calculate</u> the molar mass and determine the formula of mineral **B**. Hint: it is an ionic solid insoluble in water and containing no water of crystallization.

Your work:

B is:

5.1.4. <u>Determine</u> the formula of mineral **C**.

Your work:

C is:

5.1.5. <u>Determine</u> the formulae of Egyptian blue and Chinese blue.

Your work:

Egyptian blue is:

Chinese blue is:

5.1.6. <u>Determine</u> the formula of mineral **A**.

Your work:

A is:

5.2.1. <u>Suggest</u> the formula of the mineral used in place of **C**.

5.2.2. <u>Could</u> the temperature of synthesis of Chinese blue be decreased if this mineral is used instead of **C**?

<u> </u>	
Yes	

No

5.3. <u>Write down</u> a formula of a binary compound that forms under the conditions required for Chinese violet and is responsible for the change of the color.

Problem 6 7% of the total

6.1.1.	6.1.2	6.2.1.	6.2.2.	6.2.3.	6.3.1	6.3.2.	Sum
24	4	3	6	2	2	8	49

6.1.1. <u>Suggest</u> structures for **A**, **B**, **C**, **D**, **F**, and **G**. None of the reactions except for the final transformation with L-selectride are stereoselective. Therefore, stereochemistry does not need to be indicated in your answers.

Α	В
C	D
P	6
r	G

6.1.2. <u>Give</u> the formula for a possible reagent, **X**, to convert compound **D** to **E**.

X

6.2.1. <u>What</u> is the enantiomeric excess of the resolved compound prepared by the industrial route?

ее:	

6.2.2. <u>Assign</u> the labelled stereocentres (α, β, γ) in (-)-**1** as *R* or *S*.

α	β	γ

6.2.3. <u>Give</u> the formula for a reagent that carries out the same reaction as L-selectride, the conversion of **H** to **1**. You need not worry about stereoselectivity.

6.3.1. <u>Give</u> the formula for compound **Y** to carry out the first step of the route.

6.3.2. <u>Suggest</u> structures for **J** and **K**.

Y

J	К

8% of the total

7.1.	7.2.	7.3.1.	7.3.2.	7.4.	Sum
2	36	16	8	4	66

7.1. <u>Determine</u> the empirical formula of **G**.

Your work:

7.2. <u>Give</u> the structures of A, B, C, D, E1, E2, F1, F2 and G.

Α	В
C	D
F1 and F2	



7.3.1. <u>Give</u> the structures of **H**, **I**, and **J**. There is no need to show the different diastereoisomers formed.



7.3.2 Give the structures of diastereoisomers **K1**, and **K2**.

K1		
K2		

7.4. <u>Give</u> the structure of **N**.

Ν

J

7% of the total

8.1.	8.2.1.	8.2.2.	8.3.1.	8.3.2.	8.3.3.	Sum
4	10	1	4	1	2	22

8.1.1. Which of the following sentences is true?

 \Box **A** is an α isomer. \Box **A** is a β isomer. \Box **A** is neither α nor β .

 \Box **A** is a mixture of α and β isomers.

8.1.2. Which of the following sentences is true?

 \square We can get product **A** only if we use α-D glucose as starting material. \square We can get product **A** only if we use β-D glucose as starting material. \square We can get product **A** either from α- or from β-D glucose as starting material.

8.1.3. Which one of these reagents can be utilized as **X** for the selective hydrolysis of **A**?

□ 50% acetic acid
□ 6M HCl in water
□ 6M HCl in acetic acid

concentrated H₂SO₄
 1M NaOH in water

8.1.4. Which is the stereochemically correct structure for compound **B**?



 B.2.1. Draw the structures of C, D₁, D₂, E and F including stereochemical information. Show F as the more stable 6-membered ring containing isomer using the ring skeleton. Indicate with a wavy line if absolute chirality around a carbon is not known.



- **8.2.2.** The reaction sequence from glucose to **F** does not seem to be useful. In some cases, however, this is the most economical way to produce **F**. <u>In which case</u>?
 - \square ¹³C labelling at carbon 6 of **F**
 - \square ¹³C labelling at carbon 5 of **F**
 - \square ¹³C labelling at carbon 1 of **F**
 - \square ¹⁵O labelling at glycosidic OH of **F**
 - \Box synthesis of an uncommon isomer of ${\bf F}$

 B.3.1. <u>Draw</u> the structure of **G** including stereochemistry. <u>Draw</u> **H** as the more stable 6-membered ring containing isomer using the ring skeleton. <u>Indicate</u> with a wavy line if absolute chirality around a carbon is not known.



8.3.2. How are the rings of **G** fused together?

- □ both junctions cis
- □ one cis and one trans junction
- \Box both junctions trans

8.3.3. Which of the following is true for **H**? (You can choose more than one option.)

- □ **H** is a reducing sugar (reacts with Fehling's reagent)
- □ **H** is an aldaric acid (dicarboxylic derivative of an aldose)
- \Box **H** is an aldonic acid (α -carboxyl derivative of an aldose)
- \Box **H** is a uronic acid (ω -carboxyl derivative of an aldose)
- □ **H** is a ketose
- \Box **H** is a lactone
- \Box **H** is a D sugar
- \Box **H** is achiral
- \Box **H** is a meso compound