

A Level Chemistry B (Salters) H433/02 Scientific literacy in chemistry

Sample Question Paper

Date - Morning/Afternoon

Time allowed: 2 hours 15 minutes



You must have:

- the Advance Notice (inserted)
- the Data Sheet for Chemistry B (Salters)

You may use:

· a scientific calculator



First name	
Last name	
Centre number	Candidate number

INSTRUCTIONS

- Use black ink. You may use an HB pencil for graphs and diagrams.
- Complete the boxes above with your name, centre number and candidate number.
- Answer all the questions.
- · Write your answer to each question in the space provided.
- Additional paper may be used if required but you must clearly show your candidate number, centre number and question number(s).
- Do **not** write in the bar codes.

INFORMATION

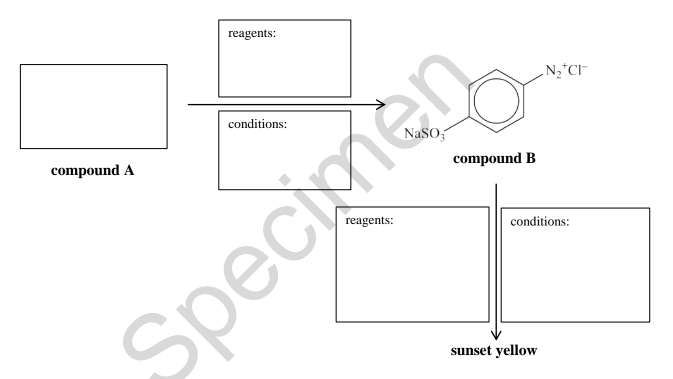
- The total mark for this paper is 100.
- The marks for each question are shown in brackets [].
- Quality of extended responses will be assessed in questions marked with an asterisk (*).
- This document consists of 24 pages.

[5]

Answer all the questions.

1 The azo dye 'sunset yellow' is a food colouring, though its use is being phased out.

(a) This dye can be made by a synthetic route that is shown, in outline, below.



Complete the diagram by drawing the structure of compound A, and giving the reagents and conditions for the reactions.

(b) The groups shown as ' SO_3Na ' aid the solubility of the dye.

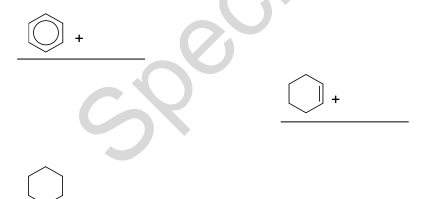
Explain how the SO ₃ Na groups do this, in terms of bonds broken and made.	
[4	

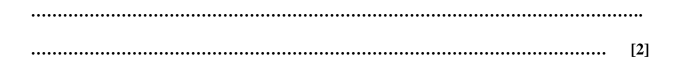
(c)	Sunset yellow absorbs blue light of wavelength 450 nm.
	Calculate the energy change of this absorption in kJ mol ⁻¹
	Show all your working.

energy change =	 kJ mol ⁻¹	[3]
onergy onange	 The state of the s	[C]

(d) Enthalpy changes of hydrogenation show that benzene is not best represented as:Complete and explain the energy level diagram below to show why this is so.







[2]

(e) A compound similar to compound **B** decomposes in dilute aqueous solution.

$$C_6H_5N_2C1 + H_2O \rightarrow N_2 + HC1 + C_6H_5OH$$

Some chemists investigate the rate of this decomposition. They collect the nitrogen gas in a graduated syringe at different initial concentrations of the $C_6H_5N_2Cl$ solution. They time how long it takes for $50~\text{cm}^3$ of nitrogen to be collected. The volume of solution used in each experiment is $100~\text{cm}^3$.

(i) Two of the chemists' results are given in the table below.

Initial concentration / mol dm ⁻³	0.020	0.040
Initial rate / mol dm ⁻³ s ⁻¹	8.70×10^{-7}	1.74×10^{-6}

Use these data to calculate the rate constant for the reaction. Include units in your answer.

Assume that the reaction is zero order with respect to water.

Show all your working.

	rate = units
(ii)	The chemists could have measured the time to produce a much larger volume of nitrogen.
	Suggest why it would have been inappropriate to measure the time to collect larger volumes of gas, particularly in the experiments with lower concentrations of $C_6H_5N_2Cl$.

(iii)	The progress of the reaction can also be followed with a pH meter.
	What is the pH of the solution once 50 cm ³ of nitrogen have been collected?
	Give your answer to two decimal places.
	pH = [2]
(iv)	Besides gas collection and measuring pH, suggest one other method with which the chemists could measure the progress of the reaction.
	[1]
	QUESTION 1 CONTINUES ON PAGE 6

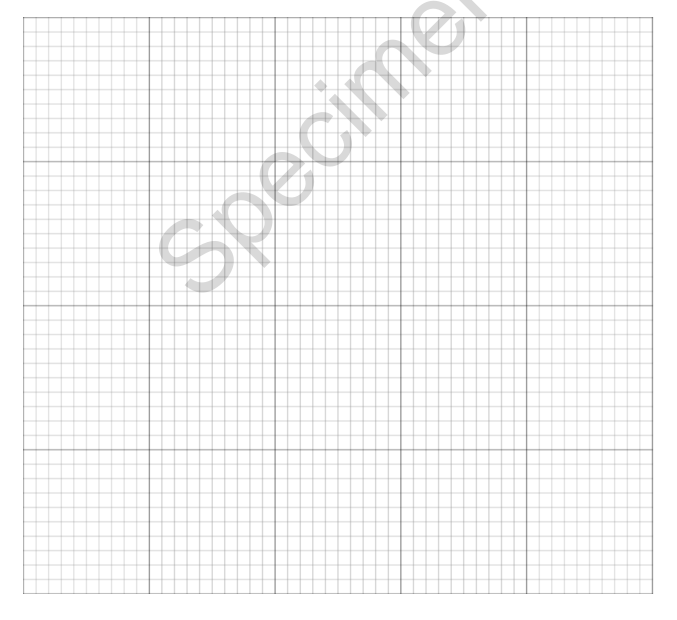
(v) The chemists carried out the reaction at several temperatures and obtained the results in the left-hand two columns of **Table 1.1**.

Table 1.1

Temperature, T / K	rate constant, <i>k</i> / s ⁻¹	1/T / K ⁻¹	$\ln k$
278	1.5×10^{-6}	3.60×10^{-3}	-13.41
298	4.1×10^{-5}		-10.10
308	2.0×10^{-4}	3.25×10^{-3}	
323	1.4×10^{-3}	3.10×10^{-3}	-6.57

Calculate the missing values in **Table 1.1** and plot a graph of $\ln k$ against 1/T on the graph paper below.

Use your graph to calculate a value for the activation enthalpy of the reaction.



7

 $activation\ enthalpy = \dots \qquad kJ\ mol^{-1}$

[5]



2	condiox	crete (kide fi	contains calcium hydroxide, Ca(OH) ₂ , and is alkaline when damp. The steel reinforcements do not corrode in an alkaline environment. When the calcium hydroxide reacts with carbon com the air to form calcium carbonate, 'carbonatation' occurs and the pH falls. The increased in cause the steel to corrode.	
	(a)	(i)	Write the equation for carbonatation.	
				[1]
		(ii)	Calculate the mass of calcium carbonate that is formed when 15 cm ³ of carbon dioxide (measured at room temperature and pressure) react with excess calcium hydroxide.	
			Give your answer to an appropriate number of significant figures.	
	(b)	Carl	$mass = \dots \qquad g$ conatation produces calcium carbonate. The carbonate ion is alkaline in aqueous solution.	[2]
		(i)	Write an equation for the reaction that occurs between the carbonate ion and water.	[1]
		(ii)	Explain how the carbonatation reaction in (a)(i) accounts for the fall in pH.	· ··
				[2]

(c)	(i)	Calculate the pH of a 0.0020 mol dm ⁻³ solution of calcium hydroxide.	
		Show all your working.	
		pH = [3]]
	(ii)	Calculate the pH of a 0.0010 mol dm ⁻³ solution of carbon dioxide.	
		$K_{\rm a} = 4.5 \times 10^{-7} \; {\rm mol \; dm^{-3}}$	
		pH = [2]]
	(iii)	Explain the approximation that is made in performing the calculation in (ii) when $[H^+]$ is low.	
]

(d)	The	rusting of steel reinforcements can affect the strength of the structure.
	(i)	Write the half-equations (as equilibria) for the formation of hydroxide ions and Fe^{2+} ions in the first stage of rusting.

	[2]
(ii)	Fe(OH) ₂ is oxidised by O ₂ to give rust, Fe ₂ O ₃ .
	Use this information and your answer from part (i) to suggest why iron is less likely to rust in alkaline conditions.
	[3]

(e) The indicator phenolphthalein can be used to detect carbonatation in concrete. Phenolphthalein has a variety of related structures, depending on the pH.

Species	C	D	E	F
Structure	НОСООН	НООН	O CO ₂	-о о- он он
pН	<0	0-8.2	8.2-12.0	>12.0
Conditions	strongly acidic	acidic or near- neutral	alkaline	strongly alkaline
Colour	orange	colourless	pink	colourless

(i)	Give the colours of phenolphthalein for normal concrete and for concrete that has undergone carbonatation.
	normal:
	carbonatation: [1]
(ii)	An acid is titrated with a strong alkali using phenolphthalein until the pink colour just persists. If the solution is then allowed to stand in the titration flask it slowly goes colourless.
	Explain what is happening.
	[2]

3 The compound 'GMV' is a potential 'green fuel' as it can be made from glucose and can be blended with petrol.

(a) (i) Write an equation, using molecular formulae, for the complete combustion of GMV to carbon dioxide and water.

[2]

[2]

(ii) Calculate the ratio of the masses of carbon dioxide produced by burning equal masses of GMV and hexane.

Show all your working.

(iii) A scientist says that the result in (ii) is an indication of the relative effects on the environment when GMV and hexane are used as vehicle fuels.

Discuss this statement.

b)	GMV can be made in the laboratory by reacting a single organic compound with concentrated sulfuric acid.
	Give the structure of this organic compound and state how it forms GMV.

QUESTION 3 CONTINUES ON PAGE 14

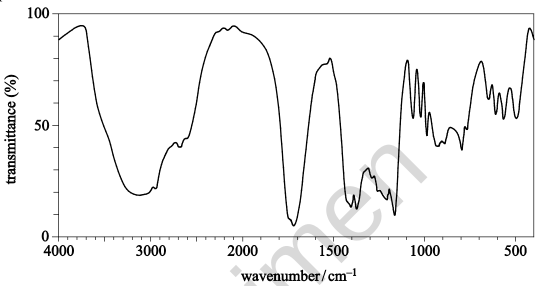
(c)* Compound G is an intermediate in the manufacture of GMV from glucose.

Compound G has the following composition by mass:

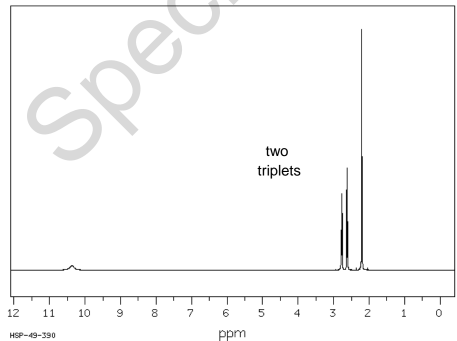
C, 51.72%; H, 6.90%; O, 41.38%

Compound **G** has IR and ¹H NMR spectra as shown below.

IR spectrum:



¹H NMR spectrum:



4	When brass objects are used in the presence of horses, the brass tends to corrode. This is because
	ammonia from horse urine reacts with copper in the brass to form the soluble compound [Cu(NH ₃) ₄]SO ₄ .

(a) Draw a diagram to show the square planar shape of the complex ion $[Cu(NH_3)_4]^{2+}$ and give the co-ordination number.

On your diagram name the type of bond joining atoms to the metal ion.

[2]

(b) This complex ion forms when ammonia solution is added to copper(II) sulfate solution. The equation for the reaction can be represented as below.

$$Cu^{2+}(aq) + 4NH_3(aq) \rightleftharpoons [Cu(NH_3)_4]^{2+}(aq)$$
 Equation 4.1

(i) High concentrations of ammonia in horse's urine can be a sign of infection. Adding a fixed amount of copper(II) sulfate solution to a given volume of urine produces the deep blue/purple ion, $[Cu(NH_3)_4]^{2+}(aq)$ in **equation 4.1**.

Suggest how the production of this ion can be used as a measure of the ammonia content in the horse's urine.



(ii) Write an equation for the equilibrium constant, K_c , of the reaction in equation 4.1.

[1]

	(iii)	A solution initially contains 0.10 mol dm^{-3} of Cu(II) ions and 0.40 mol dm^{-3} of ammonia. A equilibrium, the concentration of $[\text{Cu}(\text{NH}_3)_4]^{2+}$ formed is x .	t
		Write an expression for the equilibrium constant K_c in terms of x .	
			[2]
	(iv)	Two students wanted to measure the concentration of ammonia in the complex solution at equilibrium. One suggested that they titrate samples of the complex solution with standard sulfuric(VI) acid. The other said the results would be inaccurate.	
		Comment on who is correct and give a reason for your answer.	
			••
			••
			[2]
(c)	Cop	per(II) ions also form a complex with the bidentate ligand NH ₂ -CH ₂ -CH ₂ -NH ₂ .	
	Sug	gest the formula of the complex ion formed with this ligand and give the co-ordination number	er.
	•••••		•••
	••••		[2]
(d)	Cop	per(II) ions react with iodide ions in aqueous solution to give CuI and I ₂ .	
	(i)	Write an equation for the reaction.	
			[1]
	(ii)	Give the systematic name for CuI.	
			[1]

[2]

5	This	s ques	tion refers to the Article 'Sniffing out an Explanation' which comes as an insert in this paper	er.
	(a)	(i)	Draw, in the space below, the skeletal formula of the structure you would expect to obtain when ambrinol (Fig. 2) reacts with bromine.	
				[1]
		(ii)	How many peaks would you expect in the ¹³ C NMR spectrum of ambrinol?	
				[1]
		(iii)	Give the number of chiral centres in ambrinol.	
			Explain how you arrived at your answer.	
				• • • • •
				[2]
		(iv)	Discuss whether or not you would expect the enantiomers of ambrinol to smell the same.	
				••••
				••••

(b)		cribe chemical tests, resulting in colour changes, by which you could distinguish between illin, guaiacol and benzaldehyde (Fig. 3).	
	Incl	ude a positive reaction for each substance.	
	••••		•••
	••••		•••
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	••••	•••••••••••••••••••••••••••••••••••••••	•••
	••••		[4]
(c)	Exp	plain the significance of the graph (shown in Fig. 3) to Turin's theory.	
	Giv	e the cause of the peak just above 3000 cm ⁻¹ .	
	••••		••••
	••••		••••
	••••		••••
	••••		[3]
(d)	Thi	s part of the question explores Turin's use of isotopes to support his theory.	
	(i)	State the meaning of the term <i>isotopes</i> , using the terms <i>atomic number</i> and <i>mass number</i> .	
		Give the number of neutrons in an atom of deuterium and an atom of ¹³ C.	
		meaning:	••••
			••••
		number of neutrons:	••••
			[2]

(ii)	A sample of carbon dioxide contains 98.89% ¹² C, the rest being ¹³ C.
	Calculate the $A_{\rm r}$ value for the carbon in the carbon dioxide to two decimal places.
	4 [1]
	$A_{\rm r} = \dots \qquad [1]$
(iii)	Draw the skeletal formula of 'acetophenone'.
	State the difference between this molecule and 'acetophenone- d_8 ' (page 5).
	[2]

(iv)*	Discuss examples from the article that support Turin's theory and not the 'lock and key' theory
	••••••••••••••••••••••••••••••••••••
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A Level Chemistry B (Salters) H433/02 Scientific literacy in chemistry Sample Advance Notice Article

For issue on or after: Date/Year





NOTES FOR GUIDANCE (CANDIDATES)

- 1. This leaflet contains an article which is needed in preparation for a question in the externally assessed examination H433/02 Scientific literacy in chemistry.
- 2. You will need to read the article carefully and also have covered the learning outcomes for A Level in Chemistry B (Salters). The examination paper will contain questions on the article. You will be expected to apply your knowledge and understanding of the work covered in A Level in Chemistry B (Salters) to answer this question. There are 20–25 marks available on the question paper for this question.
- 3. You can seek advice from your teacher about the content of the article and you can discuss it with others in your class. You may also investigate the topic yourself using any resources available to you.
- **4.** You will not be able to bring your copy of the article, or other materials, into the examination. The examination paper will contain a fresh copy of the article as an insert.
- **5.** You will not have time to read this article for the first time in the examination if you are to complete the examination paper within the specified time. However, you should refer to the article when answering the questions.

This document consists of 8 pages. Any blank pages are indicated.

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Sniffing out an explanation

By Josh Howgego from The Mole, Issue 6, November 2012. Published by the Royal Society of Chemistry.

It's strange to think that no one knows exactly how smell works. Josh Howgego explains the chemistry behind the puzzle.

In laboratories up and down the country there are chemists who have the misfortune to work with thiols. Some of these whiffy molecules have a stench so pungent, so disgusting, that they have been known to empty entire buildings when a few drops are spilled in the wrong place.

But why do thiols smell so bad? It doesn't seem to make sense when you consider their structures (Fig. 1). Thiols are the sulfur analogues of alcohols – they look very similar, so what makes their odours different?

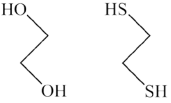


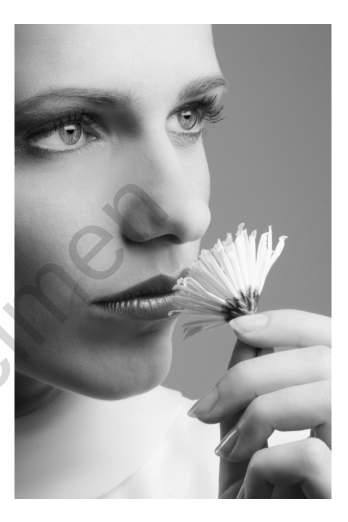
Fig. 1: Ethylene glycol (left) doesn't have much of a smell, but ethane dithiol (right) gives off one of the strongest and most unpleasant stenches known.

The answer is surprisingly complicated, and scientists are still arguing about it. It's strange to think that smell remains a mysterious process.

You would think that vision – which we understand quite well – would be a much more complicated sense than smell. But actually our eyes only have to detect one thing: photons (albeit with different wavelengths). Whereas our noses are chemical sensors: capable of identifying thousands of different molecules.

Smells of the past

Smell has an emotional power over us that the other senses can't emulate. The scent of cut grass on a sunny day can transport us back to memories of childhood, and we are beginning to understand why smell can be so evocative. The brain's olfactory bulb – the first port of call for impulses



from the nose – has a direct neural link to the limbic system, the region of the brain that processes memory and emotions. In contrast the major nerves from all the other senses are routed *via* the thalamus for pre-processing before the information is fully interpreted. That might be because humans have evolved to use mainly sight and touch for day to day activities (and so these inputs need careful attention). Leaving our sense of smell as a more instinctive and subtle guide.

Many animals use smell as a means of communication too. The chemicals that animals secrete as messages are called pheromones. They can be warnings of danger, or even hints that the

sender is on the lookout for a mate. There is little evidence to suggest that humans use pheromones but scientists think our ancestors may have done because there are parts of our nose which are no longer connected up to the brain. These may once have been part of the vomeronasal organ (the part of the nose which detects pheromones), but we have lost the use of it during evolution.

Getting a whiff of the truth

But how do our noses detect chemicals anyway? Early in the 20th century scientists thought that olfactory receptors worked much like any other enzyme or protein: they have a well-defined three-dimensional structure and a specifically shaped cleft in the middle. Only molecules with a complementary shape are able to fit inside the cleft, and so the idea became known as the 'lock and key' theory, because of the shape-dependent way the receptor and small molecule fit together.

You can imagine the nose like a dungeon with hundreds of locked doors leading to the brain (an analogy that works quite well, given the dark and slimy nasal environment). When a molecule wafts into the chamber it can only unlock a specific door, and the passage it activates tells the brain the identity of the intruder.

But alas, the concept of olfactory receptors being like locks has seemed over-simplified for decades. Molecules that have similar shapes would be expected to open the same locks, but as we've already seen (Fig. 1), structurally related molecules can smell utterly different. On the other hand, it's not clear how timberol, cedramber and ambrinol (Fig. 2) could unlock the same smell receptors when they have such dissimilar structures, yet all three smell very alike.

Shaking possible answers out

Luca Turin, a Lebanon-born biophysicist fascinated with the mechanisms of smell, came up with an innovative theory to explain all this in 1996. It's not just the molecule's shape that's important, according to Turin, but also how it vibrates once inside the receptor's binding pocket. If it oscillates in just the right way it encourages an electron to 'jump' from one part of the receptor

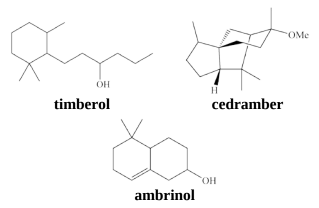


Fig. 2: Three structurally different molecules which all smell very similar

protein to another (in a process called quantum tunnelling), and this sets off a nerve impulse.

Turin's theory can potentially explain the different scents of molecules with the same shape but different atoms. The vibrations of any given bond are decided not only by geometry, but also by the mass of the atoms at either end. Sulfur and oxygen are different sizes, so thiol and alcohol functional groups have different vibrational signatures.

Like all good theories, Turin's made some predictions about smell that could be tested. One was that even subtle differences in atomic mass in similar molecules should result in odour variations. He stated, for example, that acetophenone and its deuterated analogue (acetophenone-d₈) smell a little different (see 'Subtle vibrations').

In 2001 scientists from Canada set out to check if this 'isotopes smell different' rule would hold for a similar compound, benzaldehyde. They found that trained noses could distinguish regular and deuterated benzaldehyde quite easily. The two compounds smell similar (like 'bitter almonds'), but not quite the same.

Interestingly, when they tried varying the isotope of carbon in the molecule (from regular ¹²C to ¹³C) people could detect a difference less often. That adds up, because the percentage mass difference between ¹²C and ¹³C is small, and would understandably affect the overall vibrations of the molecule less than the doubling of mass incurred when ¹H is substituted with ²H. The results seemed to support Turin's theory nicely.

But unfortunately this was not to be a case where the scientists all happily agreed with one another. In 2004 another team of scientists looked at Turin's claim about acetophenone and acetophenone-d₈, and disagreed with him: finding that, in general, their test subjects couldn't distinguish the two molecules.

Adding up to a big stink

They also looked at another prediction of Turin's theory – that molecules can add up to produce smells which none of the individual molecules give off in isolation. The classic example is a 1:1 mixture of benzaldehyde and guaiacol. Looking at these molecules (Fig. 3), it is obvious that a mixture would contain the same functional groups as vanillin overall (aldehyde, methyl ether and phenol), but they wouldn't add together to make something the same shape as vanillin (it would be nearly twice as big). Turin says that the 1:1 mixture does smell of vanilla though, adding that 'the illusion is not perfect, but it is striking, because the vanilla character is absent from the components'.

The scientists then set out to test Turin's reports themselves. They asked a group of test subjects to rate vials containing guaiacol, benzaldehyde, and a 1:1 mixture of the two on a scale from 1 (no vanilla

character) to 13 (extremely vanilla). They tried the test at various concentrations, but at no point did the subjects find the 1:1 mixture more vanilla-like than either of the separate components. This apparently disproved another of Turin's findings. The team ended their report of the study saying that 'molecular vibrations alone cannot explain the perceived smell of an odorous chemical'.



Vanillin is extracted from vanilla pods – but can its smell be recreated from a mix of two other compounds?

To date, scientists still don't agree on why similarly shaped molecules can smell so different, but we are certainly closer to the answer than we were before Turin's hypothesis. His ideas may not be fully correct, but they have prompted people to ask some interesting questions that may eventually lead us to the answers.

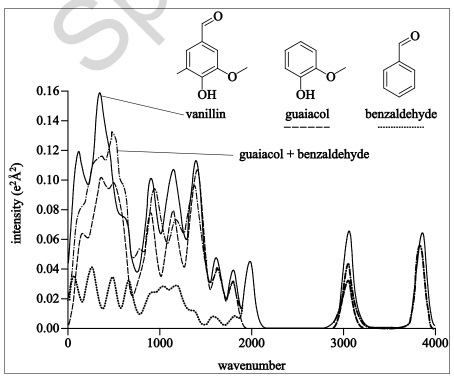


Fig. 3 H433/02

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Subtle vibrations

The mass of the two atoms at either end of a chemical bond helps determine the frequency at which it vibrates. It's a little like a guitar string; if the atoms are heavier, they pull the 'string' taut and the vibration occurs at a higher pitch.

An easy way to change the mass of atoms without changing the shape of the parent molecule is to use isotopes; atoms which have the same number of protons but a different number of neutrons. Turin stated in the 1990s that acetophenone-d₈, which has all its protons (¹H) replaced with deuterium (²H), smells slightly different from the natural, all-hydrogen analogue.

END OF ADVANCE NOTICE ARTICLE