

***Organic Chemistry Principles and Mechanisms 2nd  
edition Karty Test Bank***

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TEST BANK

# Organic Chemistry: Principles and Mechanisms

Second Edition

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# PREFACE

When was the last time you were pleased with the consistency and quality of the assessment supplements that come with introductory texts? If you are like most professors, you probably find that these assessment packages do not always meet your needs. To address this issue, Norton has collaborated with Valerie Shute (Florida State University) and Diego Zapata-Rivera (Educational Testing Services) to develop a methodology for delivering high-quality, valid, and reliable assessment supplements through our Test Banks and extensive suite of support materials.

## WHY A NEW APPROACH?

In evaluating the test banks that accompany introductory texts, we found four substantive problem areas associated with the questions:

1. Test questions were misclassified in terms of type and difficulty.
2. The prevalence of low-level and factual questions misrepresented the goals of the course.
3. Topics were unevenly distributed: Trivial topics were tested via multiple items, while important concepts were not tested at all.
4. Links to course topics were too general, thus preventing diagnostic use of the item information.

## STUDENT COMPETENCIES AND EVIDENCE-CENTERED DESIGN

We first conducted a focus group with the brightest minds in educational testing to create a new model for assessment. A good assessment tool needs to do several things: (a) define what students need to know and the level of knowledge and skills expected; (b) include test items

that assess the material to be learned at the appropriate level; and (c) enable instructors to accurately judge students' mastery of the material based on the assessment outcomes in terms of what they know, what they don't know, and the depth of their knowledge. Accurate assessments of student mastery allow instructors to focus on areas where students need the most help in learning.

## HOW DOES IT WORK?

For each chapter, the learning objectives that students could be expected to master by reading the text are listed. The questions are identified as remembering, understanding, applying, analyzing, evaluating, and creating. This classification is patterned after Bloom's taxonomy of educational objectives. Bloom listed six levels of learning: knowledge (information), comprehension, application, analysis, synthesis, and evaluation. Questions are also posed at three difficulty levels: easy, medium, and difficult. By asking students questions that vary in both type and level of difficulty, instructors can gather different types of evidence, which will allow them to more effectively assess how well students understand specific concepts.

### Six Question Types (classified according to Bloom's taxonomy):

1. *Remembering* questions—Test declarative knowledge, including textbook definitions and relationships between two or more pieces of information. Can students recall or remember the information in the same form it was learned?
2. *Understanding* questions—Pose problems in a context different from the one in which the material was learned, requiring students to draw from their declarative and/or procedural understanding of

important concepts. Can students explain ideas or concepts?

3. *Applying* questions—Ask students to draw from their prior experience and use critical-thinking skills to engage in qualitative reasoning about the real world. Can students use learned information in another task or situation?
4. *Analyzing* questions—Test students' ability to break down information and see how different elements relate to each other and to the whole. Can students distinguish among the different parts?
5. *Evaluating* questions—Ask students to assess information as a whole and frame their own argument. Can students justify a stand or decision?
6. *Creating* questions—Pose questions or objectives that prompt students to put elements they have learned together into a coherent whole to generate new ideas. Can students create a new product or point of view based on data?

#### Three Difficulty Levels:

1. *Easy* questions—Require a basic understanding of the concepts, definitions, and examples.
2. *Medium* questions—Direct students to use critical thinking skills, to demonstrate an understanding of core concepts independent of specific textbook examples, and to connect concepts across chapters.
3. *Difficult* questions—Ask students to synthesize textbook concepts with their own experience, making analytical inferences about topics discussed in the text.

Each question is linked to links to a specific learning objective and is written in clear, concise, and grammatically correct language appropriate for the learning objec-

tive and difficulty level being assessed. Every effort is made to eliminate bias (e.g. race, gender, cultural, ethnic, regional, handicap, age) to focus on the material and to assure validity and reliability.

#### KEY TO THE QUESTION META-DATA

Each question in the Test Bank is tagged with five pieces of information designed to help instructors create the most ideal mix of questions for their quiz or exam. These tags are:

**ANS:** This is the correct answer for each question (or, in the case of some short-answer questions, a possible correct answer to the question).

**DIF:** This is the difficulty assigned to the problem. Problems have been classified as Easy, Medium, or Difficult.

**REF:** This is the section in the textbook from which a question is drawn.

**OBJ:** This is the learning objective that the question is designed to test.

**MSC:** This is the knowledge type (described above) that the question is designed to test.

Test Bank files are available in Word, PDF, and Exam-View® Assessment Suite formats.

Finally, we would like to thank Suzette Mooring of Georgia State University and Joshua Osbourn of West Virginia University, whose careful review improved the accuracy and usefulness of this product.

## Chapter 1: Atomic and Molecular Structure

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### LEARNING OBJECTIVES

Determine the number of valence and/or core electrons for an atom or ion.

Interpret the electron configuration and formal charge for an atom or ion.

Identify forces that are involved in chemical bonding.

Analyze an energy versus internuclear distance diagram to understand the properties of a chemical bond.

Predict the properties of a covalent bond based on known periodic trends, and vice versa.

Assess the validity of a Lewis structure.

Apply knowledge of chemical structure to determine the formal charge of an unknown species.

Compare a series of structures to determine if they are resonance structures.

Determine the molecular formula of an organic compound from a structural drawing or condensed formula.

Master the structural drawing of organic molecules—specifically, Lewis structures and line structures.

Assimilate your knowledge of molecular structure to identify and/or draw organic functional groups.

Identify the key structural features of amino acids, saccharides, and nucleotides.

Deduce and draw the resonance structures that contribute to the resonance hybrid, and vice versa.

Elaborate how an electrostatic potential map correlates to molecular structure and properties.

Predict the ionic or covalent nature of an organic structure from physical property data.

Indicate bond dipoles and lone pairs on an organic structure, and predict how these structural features impact chemical reactivity.

Apply the concept of resonance to predict the outcome of a chemical reaction.

Depict electron delocalization via resonance using appropriate arrow notation.

Recognize and name functional groups within a complex molecule.

Draw a structure of a given molecular formula that contains a specific functional group.

## MULTIPLE CHOICE

1. Which orbital does NOT house core electrons for a bromine atom?
- 1s
  - 4p
  - 3p
  - 2s
  - 3s

ANS: B                      DIF: Easy                      REF: 1.3

OBJ: Determine the number of valence and/or core electrons for an atom or ion.

MSC: Remembering

2. An atom of which element would have an electron configuration of  $1s^2 2s^2 2p^6 3s^2 3p^1$ ?
- Al
  - Ne
  - B
  - Si
  - Na

ANS: A                      DIF: Easy                      REF: 1.3

OBJ: Interpret the electron configuration and formal charge for an atom or ion.

MSC: Understanding

3. Which electron configuration is correct for a carbon atom with a formal charge of -1?
- $1s^2 2s^2 2p^6 3s^1$
  - $1s^2 2s^2 2p^3$
  - $1s^2 2s^2 2p^5$
  - $1s^2 2s^2 2p^6 3s^2 3p^5$
  - $1s^2 2s^2 2p^4$

ANS: B                      DIF: Easy                      REF: 1.3 | 1.9

OBJ: Interpret the electron configuration and formal charge for an atom or ion.

MSC: Understanding

4. Which electron configuration is correct for the carbon of a carbocation?
- $1s^2 2s^2 2p^1$
  - $1s^2 2s^2 2p^3$
  - $1s^2 2s^2 2p^5$
  - $1s^2 2s^2 2p^6 3s^2 3p^5$
  - $1s^2 2s^2 2p^4$

ANS: A                      DIF: Easy                      REF: 1.3 | 1.9

OBJ: Interpret the electron configuration and formal charge for an atom or ion.

MSC: Understanding

5. Which electron configuration is correct for a  $\text{Ca}^{2+}$  ion?
- $1s^2 2s^2 2p^6 3s^2 3p^1$
  - $1s^2 2s^2 2p^6 3s^2 3p^6 4s^2$
  - $1s^2 2s^2 2p^6 3s^2 3p^6$
  - $1s^2 2s^2 2p^6 3s^2 3p^6 4s^2 4p^6$
  - $1s^2 2s^2 2p^6 3s^2$

ANS: C                      DIF: Easy                      REF: 1.3 | 1.9

OBJ: Interpret the electron configuration and formal charge for an atom or ion.

MSC: Applying

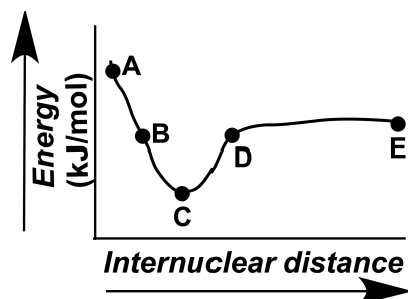
- ANS: D                      DIF: Easy                      REF: 1.3 | 1.9  
OBJ: Determine the number of valence and/or core electrons for an atom or ion.  
MSC: Analyzing

- ANS: D                      DIF: Moderate                      REF: 1.4  
OBJ: Identify forces that are involved in chemical bonding.                      MSC: Remembering

- 

- ANS: E                      DIF: Moderate                      REF: 1.4  
OBJ: Analyze an energy versus internuclear distance diagram to understand the properties of a chemical bond.                      MSC: Analyzing

9. Which point on the following diagram can be extrapolated to identify the length and strength of a chemical bond?



- a. A  
b. B  
c. C  
d. D  
e. E

ANS: C DIF: Difficult REF: 1.4

OBJ: Analyze an energy versus internuclear distance diagram to understand the properties of a chemical bond. MSC: Analyzing

10. A C—O single bond is 143 pm in length, whereas an O—O single bond is 148 pm in length. Which bond is weaker and why?
- The C—O bond is weaker because O is more electronegative than C.
  - The C—O bond is weaker because each O atom is electronegative and pulls the shared electrons toward itself.
  - The O—O bond is weaker because oxygen has *d* orbitals to engage in bonding.
  - The C—O bond is weaker because C is more electronegative than O.
  - The O—O bond is weaker because both oxygens are equally electronegative and pull the shared electrons toward themselves.

ANS: E DIF: Moderate REF: 1.4

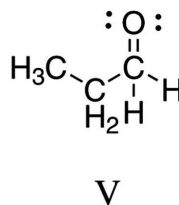
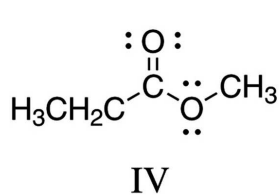
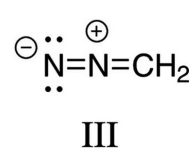
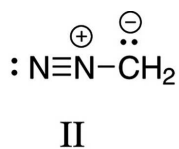
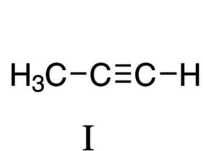
OBJ: Predict the properties of a covalent bond based on known periodic trends, and vice versa. MSC: Evaluating

11. How many total valence electrons are used in the structure of ammonium chloride,  $\text{NH}_4\text{Cl}$ ?
- 13
  - 14
  - 15
  - 16
  - 28

ANS: D DIF: Easy REF: 1.5

OBJ: Determine the number of valence and/or core electrons for an atom or ion. MSC: Analyzing

12. Which of the following Lewis structures violates the octet rule and is therefore incorrect?



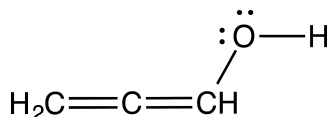
- |                  |                 |
|------------------|-----------------|
| a. Structure I   | d. Structure IV |
| b. Structure II  | e. Structure V  |
| c. Structure III |                 |

ANS: E                      DIF: Moderate              REF: 1.5

OBJ: Assess the validity of a Lewis structure.

MSC: Analyzing

13. Evaluate the following Lewis structure and determine its legitimacy.



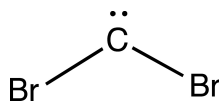
- The structure is legitimate.
- The structure is not legitimate, because the oxygen does not have an octet.
- The structure is not legitimate, because the formal charges are not shown.
- The structure is not legitimate, because the middle carbon lacks an octet.
- The structure is not legitimate, because the leftmost carbon is missing a lone pair.

ANS: A                      DIF: Moderate              REF: 1.5 | 1.6

OBJ: Assess the validity of a Lewis structure.

MSC: Evaluating

14. Consider the interesting structure below, called a dibromocarbene. The carbon of the dibromocarbene has one lone electron pair and two separate covalent bonds to individual bromine atoms. What is the formal charge on the carbon atom of the dibromocarbene?



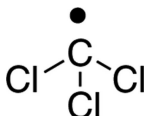
- |       |       |
|-------|-------|
| a. +2 | d. -1 |
| b. +1 | e. -2 |
| c. 0  |       |

ANS: C                      DIF: Easy                      REF: 1.9

OBJ: Apply knowledge of chemical structure to determine the formal charge of an unknown species.

MSC: Understanding

15. Consider the interesting structure below, called a free radical. The carbon of this free radical has one unpaired electron and three separate covalent bonds to individual chlorine atoms. What is the formal charge on the carbon atom?



- a. +2  
b. +1  
c. 0  
d. -1  
e. -2

ANS: C

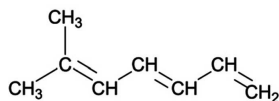
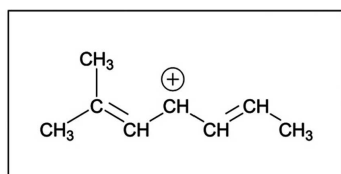
DIF: Moderate

REF: 1.9

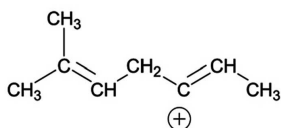
OBJ: Apply knowledge of chemical structure to determine the formal charge of an unknown species.

MSC: Applying

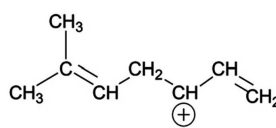
16. Which of the following is a resonance structure of the given molecule?



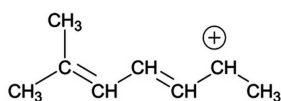
I



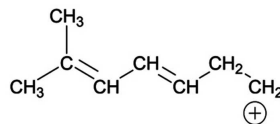
II



III



IV



V

- a. I  
b. II  
c. III  
d. IV  
e. V

ANS: D

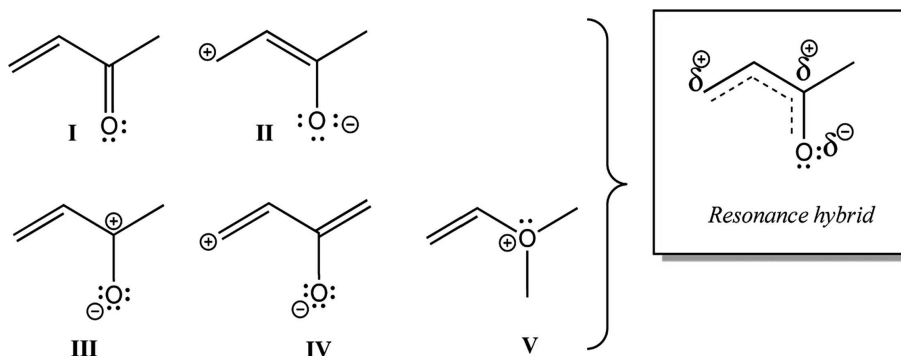
DIF: Moderate

REF: 1.10

OBJ: Compare a series of structures to determine if they are resonance structures.

MSC: Analyzing

17. Which individual structures below could be contributing resonance structures to the given hybrid structure?



- All are contributing structures.
- All but V are contributing structures.
- I, II, and III are contributing structures.
- I, III, and IV are contributing structures.
- Only I and III are contributing structures.

ANS: C                      DIF: Moderate                      REF: 1.10

OBJ: Deduce and draw the resonance structures that contribute to the resonance hybrid, and vice versa.                      MSC: Evaluating

18. When two Lewis structures are related as resonance forms, which of the following are true?
- When compared, the resonance forms have the same atoms connected in the same order.
  - Either individual Lewis structure can be used as an accurate representation of valence electron distribution.
  - Electrons in single bonds may be delocalized in the resonance forms.
  - Electrons in a multiple bond may be delocalized in the resonance forms.
  - A lone pair of electrons on an atom adjacent to a multiple bond can be delocalized in the resonance forms.
- All are true.
  - Only I, II, IV, and V are true.
  - Only I, III, IV, and V are true.
  - Only I, IV, and V are true.
  - Only I and IV are true.

ANS: D                      DIF: Difficult                      REF: 1.10 | 1.11

OBJ: Compare a series of structures to determine if they are resonance structures.

MSC: Evaluating

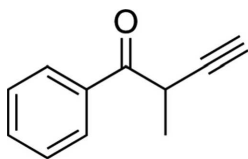
19. Which of following does NOT suggest that another resonance structure exists?
- a lone pair of electrons adjacent to a multiple bond
  - an incomplete octet on an atom adjacent to a multiple bond
  - a lone pair of electrons on an atom adjacent to an atom with an incomplete octet
  - a ring containing an atom with an incomplete octet
  - a ring of alternating single and double bonds

ANS: D                      DIF: Easy                      REF: 1.11

OBJ: Depict electron delocalization via resonance using appropriate arrow notation.

MSC: Remembering

20. How many hydrogen atoms are in the following molecule?



- a. 7  
b. 8  
c. 9
- d. 10  
e. 11

ANS: D

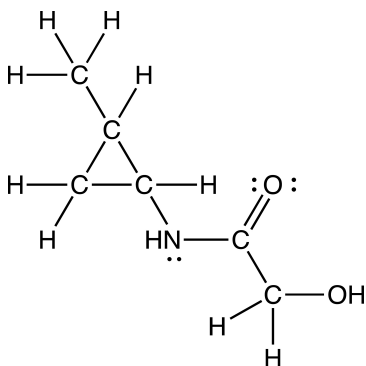
DIF: Easy

REF: 1.12

OBJ: Master the structural drawing of organic molecules—specifically, Lewis structures and line structures.

MSC: Understanding

21. What is the molecular formula of this compound?



- a.  $C_6H_{12}NO_2$   
b.  $C_5H_{12}NO_2$   
c.  $C_5H_{11}NO_2$
- d.  $C_6H_{11}NO_2$   
e.  $C_6H_{13}NO_2$

ANS: D

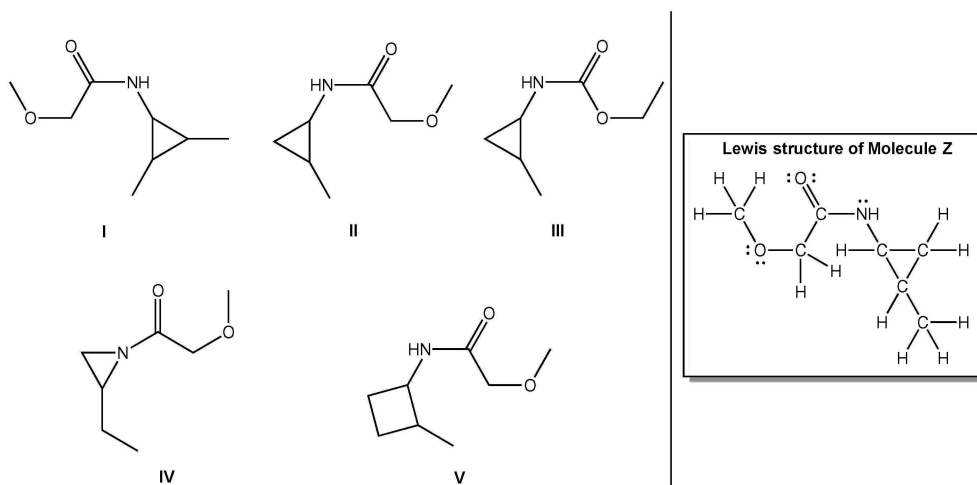
DIF: Easy

REF: 1.12

OBJ: Determine the molecular formula of an organic compound from a structural drawing or condensed formula.

MSC: Understanding

22. Which line structure is correct for Molecule Z?

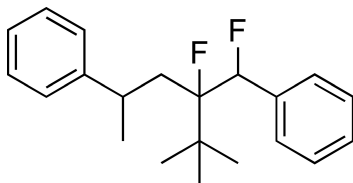


- a. I  
b. II  
c. III  
d. IV  
e. V

ANS: B DIF: Moderate REF: 1.12

OBJ: Master the structural drawing of organic molecules—specifically, Lewis structures and line structures. MSC: Understanding

23. In sum, how many total hydrogen atoms are directly connected to the benzene rings found in the following molecule?



- a. 8  
b. 9  
c. 10  
d. 11  
e. 12

ANS: C DIF: Moderate REF: 1.12

OBJ: Assimilate your knowledge of molecular structure to identify and/or draw organic functional groups. MSC: Understanding

24. Which condensed formula contains an aldehyde functional group?

- a.  $\text{CH}_3\text{COCH}_3$   
b.  $\text{CH}_3\text{CO}_2\text{CH}_2\text{CH}_3$   
c.  $\text{CH}_3\text{CH}_2\text{CH}_2\text{COOH}$   
d.  $\text{CH}_3\text{CH}_2\text{OH}$   
e.  $\text{CH}_3\text{CHO}$

ANS: E DIF: Moderate REF: 1.12 | 1.13

OBJ: Assimilate your knowledge of molecular structure to identify and/or draw organic functional groups. MSC: Applying

25. Which condensed formula contains a carboxylic acid?
- $\text{CH}_3\text{COCH}_3$
  - $\text{CH}_3\text{CO}_2\text{CH}_2\text{CH}_3$
  - $\text{CH}_3\text{CH}_2\text{CH}_2\text{COOH}$
  - $\text{CH}_3\text{CH}_2\text{OH}$
  - $\text{CH}_3\text{CHO}$

ANS: C      DIF: Moderate      REF: 1.12 | 1.13

OBJ: Assimilate your knowledge of molecular structure to identify and/or draw organic functional groups.  
MSC: Applying

26. Which condensed formula contains a ketone?
- $\text{CH}_3\text{COCH}_3$
  - $\text{CH}_3\text{CO}_2\text{CH}_2\text{CH}_3$
  - $\text{CH}_3\text{CH}_2\text{CH}_2\text{COOH}$
  - $\text{CH}_3\text{CH}_2\text{OH}$
  - $\text{CH}_3\text{CHO}$

ANS: A      DIF: Moderate      REF: 1.12 | 1.13

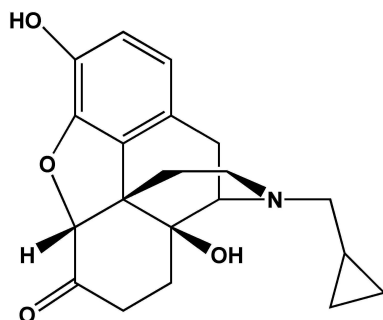
OBJ: Assimilate your knowledge of molecular structure to identify and/or draw organic functional groups.  
MSC: Applying

27. Which condensed formula contains an ester?
- $(\text{CH}_3\text{CH}_2)_2\text{O}$
  - $\text{CH}_3\text{CO}_2\text{CH}_2\text{CH}_3$
  - $\text{CH}_3\text{CH}_2\text{CH}_2\text{COOH}$
  - $\text{CH}_3\text{CH}_2\text{OH}$
  - $\text{CH}_3\text{CHO}$

ANS: B      DIF: Moderate      REF: 1.12 | 1.13

OBJ: Assimilate your knowledge of molecular structure to identify and/or draw organic functional groups.  
MSC: Applying

28. Naltrexone is an FDA-approved treatment for alcoholism that targets the mu opioid receptor. Name four functional groups that are present in naltrexone.

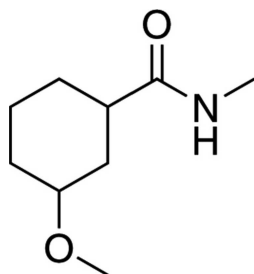


- amine, phenol, aldehyde, ether
- amine, phenol, amide, alcohol
- amine, phenol, ketone, alcohol
- ether, ketone, amide, alcohol
- ketone, phenol, alcohol, ester

ANS: C      DIF: Moderate      REF: 1.13

OBJ: Assimilate your knowledge of molecular structure to identify and/or draw organic functional groups.  
MSC: Applying

29. Which functional group is present in the following molecule?



- a. amine
- b. alcohol
- c. ketone
- d. ester
- e. ether

ANS: E                      DIF: Moderate                      REF: 1.13  
OBJ: Recognize and name functional groups within a complex molecule.  
MSC: Analyzing

30. Which of the following  $\alpha$ -amino acids possesses two hydrogen atoms adjacent to the carboxylic acid?

- a. serine
- b. phenylalanine
- c. glycine
- d. tryptophan
- e. lysine

ANS: C                      DIF: Easy                      REF: 1.14  
OBJ: Identify the key structural features of amino acids, saccharides, and nucleotides.  
MSC: Remembering

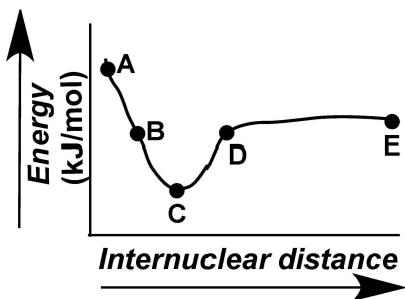
31. For which of the following  $\alpha$ -amino acids can resonance forms be drawn for its side chain?

- a. alanine
- b. methionine
- c. glycine
- d. proline
- e. histidine

ANS: E                      DIF: Moderate                      REF: 1.14  
OBJ: Identify the key structural features of amino acids, saccharides, and nucleotides.  
MSC: Remembering

## SHORT ANSWER

1. The evolution of a chemical bond can be tracked by plotting energy versus internuclear distance, as shown in the figure here. Describe what is occurring at the atomic level for points A–E on the graph.



ANS:

**POINT A.** Point of highest energy, because the two positive nuclei are pushed close together. Like charges repel.

**POINT B.** The positively charged nuclei are farther apart; thus, this data point occurs at lower energy than Point A.

**POINT C.** Point of lowest energy; this is the optimum internuclear distance to form a chemical bond.

**POINT D.** Nuclei are farther apart and weakly bonded to one another.

**POINT E.** Nuclei are functioning independently, and the internuclear distance is too great for a chemical bond to form.

DIF: Moderate REF: 1.4

OBJ: Analyze an energy versus internuclear distance diagram to understand the properties of a chemical bond. MSC: Understanding

2. Which would you expect to be a stronger bond: C–Si or C–C? Explain your response.

ANS:

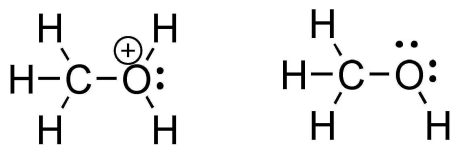
A C–Si single bond is weaker and longer than a C–C single bond. Because silicon is larger and is polarizable (since it is in period 3), its electrons do not have as great an attraction to the nucleus, and the bond between Si and C is thus longer.

DIF: Difficult REF: 1.4

OBJ: Predict the properties of a covalent bond based on known periodic trends, and vice versa.

MSC: Analyzing

3. Oxygen is an important heteroatom found in many organic molecules. Consider methanol and its protonated derivative, shown below. How does an oxygen with a positive charge, called an oxonium species, influence the magnitude of the partial positive charge on the carbon atom? Which oxygen-carbon bond do you think is more difficult to break? Explain.



ANS:

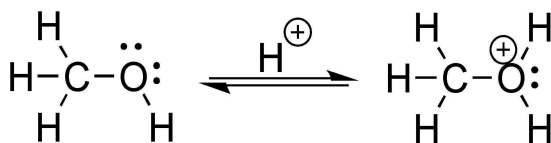
The C–O bond in the oxonium species would have the greatest bond dipole. Consequently, this bond would be weaker and easier to break than the C–O bond of methanol.

DIF: Difficult REF: 1.4 | 1.5 | 1.9

OBJ: Predict the properties of a covalent bond based on known periodic trends, and vice versa.

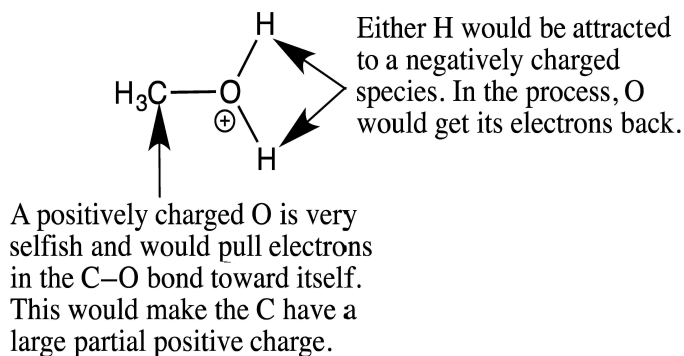
MSC: Evaluating

4. Using a chemical reaction to convert an alcohol to an oxonium species is a highly valued tool in every organic chemist's arsenal (see the following figure). Although this reaction was not covered in the current chapter, it will be discussed in due course. Reconsider your response to the preceding question. Can you identify the two different atoms of the oxonium species to which a negatively charged species might be most attracted? Explain. Hint: It is not the oxygen with the positive charge.



ANS:

Oxygen is an electronegative atom that is relatively unstable owning a positive charge; thus, it wants to engage in a process that will make it more stable.

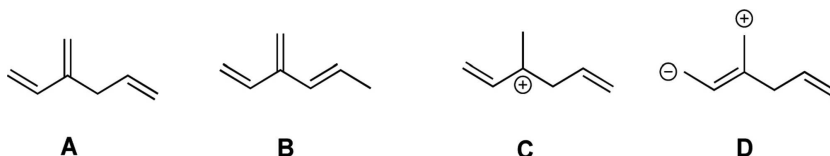


DIF: Difficult REF: 1.4 | 1.5 | 1.9

OBJ: Predict the properties of a covalent bond based on known periodic trends, and vice versa.

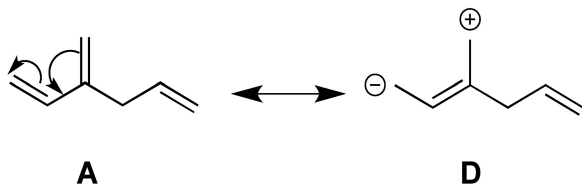
MSC: Evaluating

5. Compare Structure A with Structures B, C, and D. Is Structure B, C, or D a resonance structure of A? Justify your response. For any structures that are resonance forms, use curved arrows to show how the resonance forms are interconverted.

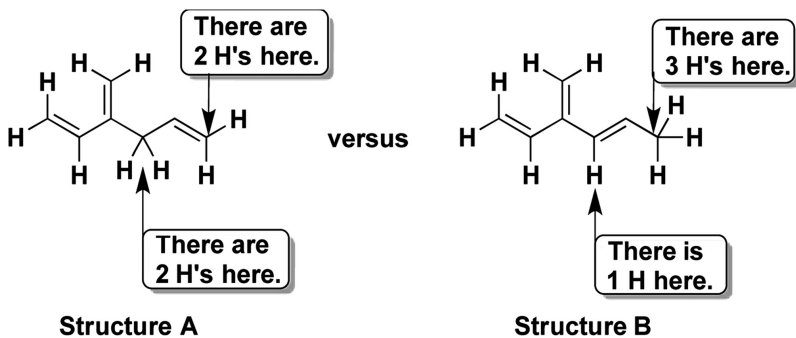


ANS:

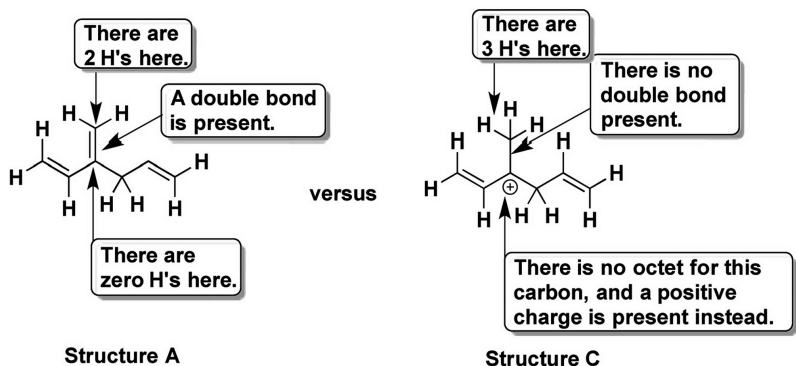
Structure D is a resonance structure of Structure A.



Structure B is not a resonance structure of Structure A, because a C–H single bond changes position.



Structure C is not a resonance structure of Structure A, because a double bond from Structure A is replaced with a C–H bond and a positive charge. Furthermore, the molecular formulas of A and C are different.



DIF: Moderate      REF: 1.4 | 1.5 | 1.9 | 1.10

OBJ: Compare a series of structures to determine if they are resonance structures.

MSC: Evaluating

6. Draw a Lewis structure of thionyl chloride ( $\text{SOCl}_2$ ), showing all lone pairs. Show each bond dipole using a dipole arrow.

ANS:



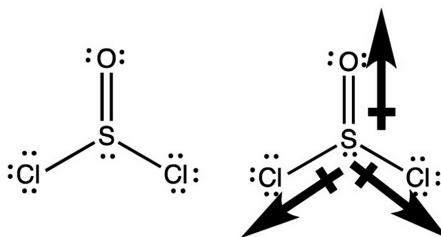
Electron count

S = group 6  $\times$  1 = 6 valence e $^-$

O = group 6  $\times$  1 = 6 valence e $^-$

Cl = group 7  $\times$  2 = 14 valence e $^-$

-----  
26 total valence electrons

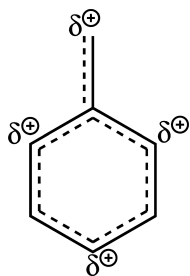


DIF: Moderate      REF: 1.5 | 1.9

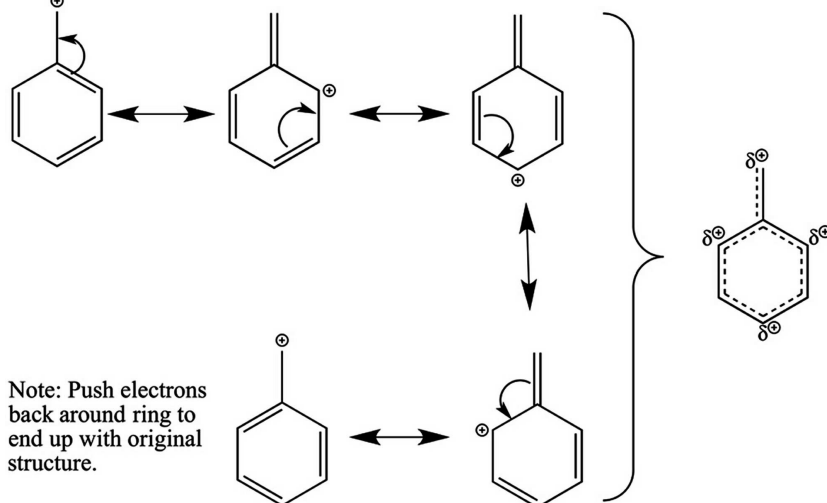
OBJ: Assimilate your knowledge of molecular structure to identify and/or draw organic functional groups.

MSC: Creating

7. Using line structures, draw the individual resonance contributors from the resonance hybrid structure given here.



ANS:



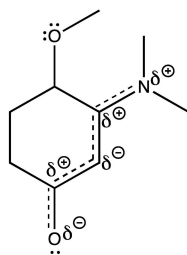
**NOTICE THAT POSITIVE CHARGES ARE ON THE SPECIFIC ATOMS WHERE THE PARTIAL CHARGES WERE LOCATED IN THE HYBRID.**

DIF: Difficult REF: 1.5 | 1.9 | 1.10 | 1.12

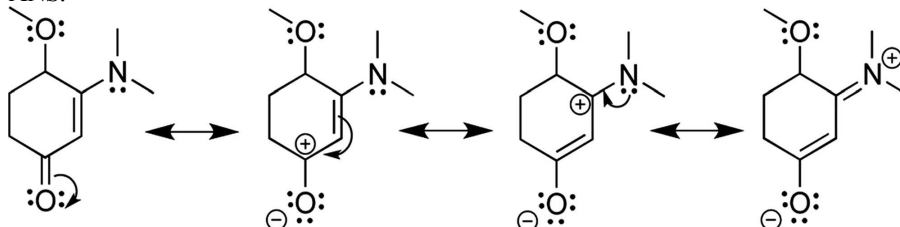
OBJ: Deduce and draw the resonance structures that contribute to the resonance hybrid, and vice versa.

MSC: Creating

8. Using line structures, deduce individual resonance contributors from the resonance hybrid structure given here.



ANS:



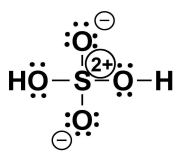
DIF: Difficult

REF: 1.5 | 1.9 | 1.10 | 1.12

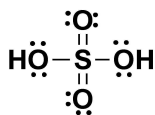
OBJ: Deduce and draw the resonance structures that contribute to the resonance hybrid, and vice versa.

MSC: Creating

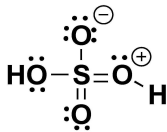
9. Are the following Lewis structures for sulfuric acid related as resonance structures? Explain.



Structure I



Structure II



Structure III

ANS:

Yes, they are all resonance structures. The structures differ in the position of lone pairs and  $\pi$  bonds. Atoms are in the same position and no sigma bonds have been broken. The structures have different potential energies due to structural and electrostatic differences.

DIF: Moderate

REF: 1.5 | 1.9 | 1.10

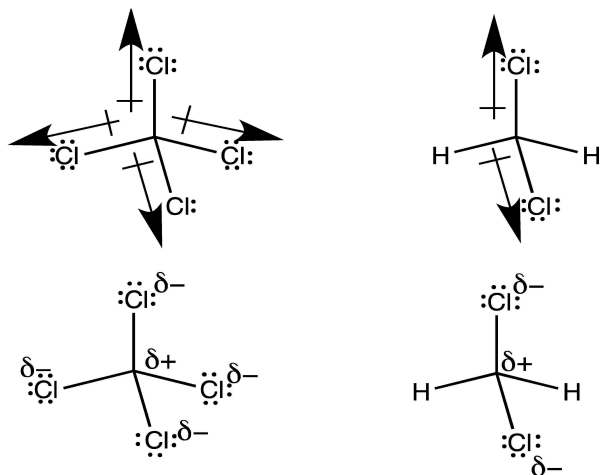
OBJ: Compare a series of structures to determine if they are resonance structures.

MSC: Analyzing

10. Consider two solvents that are commonly used for organic chemistry reactions:  $\text{CH}_2\text{Cl}_2$  and  $\text{CCl}_4$ . It is interesting that studies have shown that one of these solvents is polar and one is nonpolar. Draw valid Lewis structures for these two molecules. Show bond dipoles and both partial charges (use  $\delta^+$  and  $\delta^-$ ). How would the electrostatic potential maps for the two molecules be similar? How would they be different?

ANS:

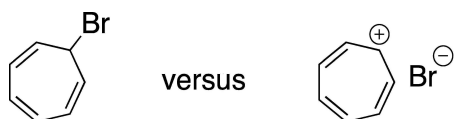
Although  $\text{CCl}_4$  has individual bond dipoles, the vector sum of the bond dipoles is zero. This means that  $\text{CCl}_4$  is a nonpolar molecule.  $\text{CH}_2\text{Cl}_2$  has two bond dipoles that do not cancel; thus,  $\text{CH}_2\text{Cl}_2$  is a polar molecule. The electrostatic potential map of  $\text{CH}_2\text{Cl}_2$  would show that the electron density in each of the C–Cl bonds is shifted toward the Cl atom.



DIF: Moderate REF: 1.7

OBJ: Elaborate how an electrostatic potential map correlates to molecular structure and properties. MSC: Analyzing

11. A compound with a molecular formula of  $\text{C}_7\text{H}_7\text{Br}$  has a melting point of  $203^\circ\text{C}$ . The compound is soluble in water but not in diethyl ether. Based on your knowledge of organic structure, is the compound most stable in its ionic or covalent form? Justify your answer.



ANS:

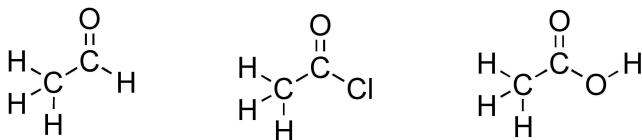
The high melting point of  $+200$  degrees and the water solubility of the seven-carbon molecule suggest that the compound is ionic. Additionally, because the compound is insoluble in ether, a charged hydrocarbon salt is most likely the most stable structure.

DIF: Difficult REF: 1.7 | 1.8

OBJ: Predict the ionic or covalent nature of an organic structure from physical property data.

MSC: Evaluating

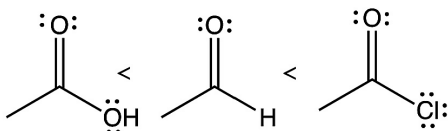
12. A carbonyl, the  $C=O$  unit, is a component of many important functional groups. Consider the Lewis structures below. Convert the Lewis structures to line structures, showing all lone pairs. Rank the structures for increasing partial positive charge. Predict which carbonyl carbon should have greatest partial positive charge, assuming that the chlorine lone pairs do not engage in resonance. Explain your answer.



ANS:

The point of difference in the three structures is the substitution on the carbon of the carbonyl. Any property that influences electron distribution will influence the magnitude of the positive charge. Effects are either electron withdrawing (e.g., an electronegative atom) or electron donating (e.g., compare the Pauling electronegativity values and resonance potential of each atom). The carbonyl carbon of the acyl chloride has the greatest partial positive charge. There are three lone pairs on Cl and two on the O of the OH. Because one of the oxygen lone pairs can be delocalized through resonance, electron density is added to the  $C=O$ , and the partial positive charge of the carbonyl carbon is reduced. Given that Cl does not typically engage in resonance, then the electron withdrawing nature of the electronegative Cl predominates.

**INCREASING  $\delta^+$  CHARGE ON C of  $C=O$**



Note: Hydrogen atoms bonded to carbon atoms are not typically drawn on line structures. The aldehyde H is shown here for emphasis.

DIF: Difficult REF: 1.7 | 1.10

OBJ: Indicate bond dipoles and lone pairs on an organic structure, and predict how these structural features impact chemical reactivity. MSC: Evaluating

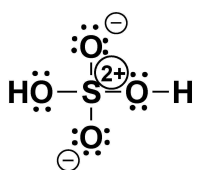
13. (a) Sulfuric acid,  $\text{H}_2\text{SO}_4$ , is an important strong oxo acid in organic chemistry. Propose Lewis structures for sulfuric acid, showing all lone pairs and formal charges, using the following guidelines:

In Structure I, one atom has a +2 formal charge. Two other atoms each possess a charge of -1. All atoms in Structure II are completely neutral. In Structure III, one atom has a +1 charge and another a -1 charge.

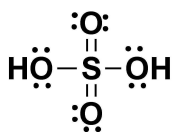
(b) Which Lewis structure is most stable? Why?

ANS:

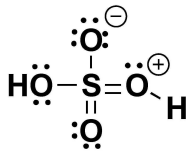
The neutral structure is most stable. The number of covalent bonds is maximized, and charges are minimized (e.g., there are no individual formal charges on atoms).



**Structure I**



**Structure II**



**Structure III**

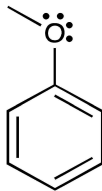
DIF: Easy

REF: 1.9 | 1.10

OBJ: Master the structural drawing of organic molecules—specifically, Lewis structures and line structures.

MSC: Analyzing | Creating

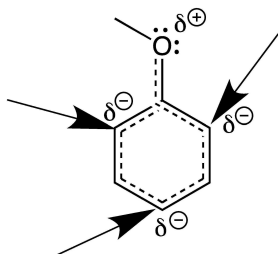
14. To which carbon atoms in anisole would a positively charged species, called “E+,” bond? Explain your answer using the concept of resonance. Hint: Refer to your response from question 16 for further insight.



**anisole**

ANS:

Opposites attract. It is reasonable to predict that any positively charged species would be attracted to the carbon atoms with a partial negative charge. These partially negative carbon atoms are highlighted with an arrow in the resonance hybrid.

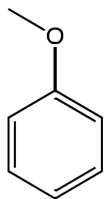


DIF: Moderate REF: 1.9 | 1.10

OBJ: Apply the concept of resonance to predict the outcome of a chemical reaction.

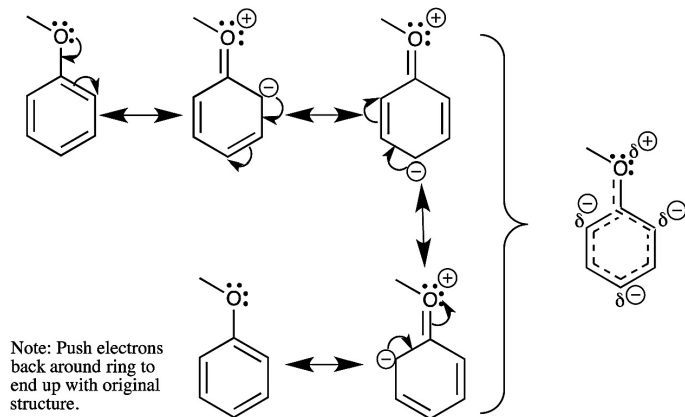
MSC: Analyzing | Evaluating

15. Draw all possible resonance forms for anisole using appropriate arrow notation. Which resonance structure is most stable? Which is least stable? Draw the resonance hybrid for anisole, indicating all partial charges.



ANS:

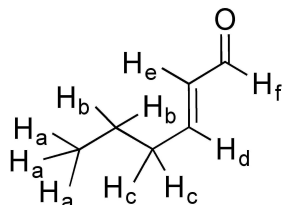
The neutral structure is most stable. The two structures with the partial negative charge closest to the oxygen are equal in energy. These structures are also more stable than the structure with the charges farther apart; thus, they contribute a greater degree to the hybrid.



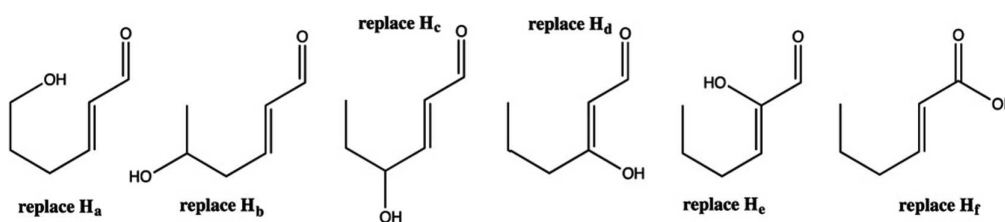
DIF: Moderate REF: 1.9 | 1.10

OBJ: Deduce and draw the resonance structures that contribute to the resonance hybrid, and vice versa. MSC: Creating

16. The six unique hydrogen atoms in the molecule below are labeled a–f. Suppose we individually replace each of these unique hydrogen atoms with a hydroxyl group (–OH) and draw a new molecule of formula  $C_6H_{10}O_2$ . (In the cases of  $H_a$ ,  $H_b$ , and  $H_c$ , substitute only one designated H with an OH.) Which of the new –OH groups would have localized lone pairs on oxygen? Which of the –OH groups would have a delocalized lone pair? Finally, for each molecule, identify the new functional group that was created.



ANS:



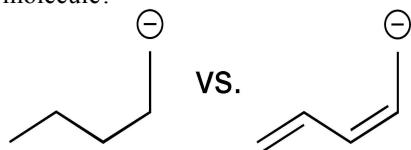
- \*  $H_a$  replacement by OH yields an alcohol that is primary, meaning the carbon connected to the oxygen of the OH has one hydrocarbon connected. The oxygen's lone pairs are localized.
- \*  $H_b$  replacement by OH yields an alcohol that is secondary, meaning the carbon connected to the oxygen of the OH has two hydrocarbons connected. The oxygen's lone pairs are localized.
- \*  $H_c$  replacement by OH yields an allylic alcohol that is also secondary, meaning the carbon connected to the oxygen of the OH has two hydrocarbons connected. The oxygen's lone pairs are localized.
- \*  $H_d$  replacement by OH yields an alcohol that is referred to as vinylic, meaning the oxygen of the alcohol is directly connected to an alkene. The "OH" + "alkene" unit is also referred to as an enol, which is a functional group that will be studied extensively in future chapters. One oxygen lone pair is delocalized through resonance; thus, resonance forms can be drawn.
- \*  $H_e$  replacement by OH yields an alcohol that is referred to as vinylic, meaning the oxygen of the alcohol is directly connected to an alkene. The "OH" + "alkene" unit is also referred to as an enol, which is a functional group that will be studied extensively in future chapters. This OH is also special because it is on a carbon that is next to (or alpha to) the carbonyl  $C=O$ . One oxygen lone pair from the OH is delocalized through resonance; thus, resonance forms can be drawn.
- \*  $H_f$  replacement by OH yields a carboxylic acid, meaning the oxygen of the alcohol is directly connected to a carbonyl,  $C=O$ . One of the oxygen lone pairs from the OH is delocalized through resonance; thus, resonance forms can be drawn.

DIF: Difficult      REF: 1.9 | 1.11

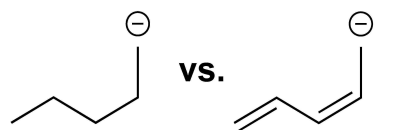
OBJ: Assimilate your knowledge of molecular structure to identify and/or draw organic functional groups.

MSC: Applying | Analyzing

17. Predict which carbon atom should have the greatest negative charge density. Explain your answer. What would the electrostatic potential map show for this carbon, in comparison with others in the molecule?



ANS:



**Negative  
charged  
localized**

**Negative  
charge  
delocalized**

The electrostatic potential map for the molecule with the localized charge would have a focused area of red (e.g., negative charge) on the rightmost carbon. On the other hand the rightmost molecule with delocalized electrons would have a broad area of light red across all carbons to indicate that the charge is spread out.

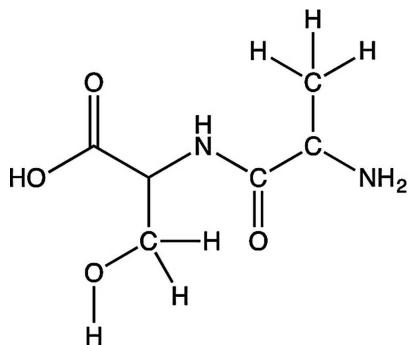
DIF: Moderate

REF: 1.10

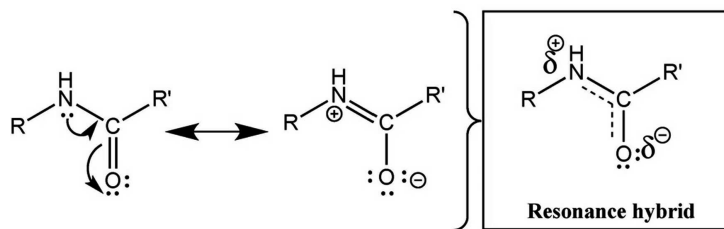
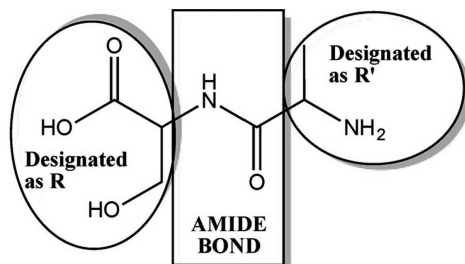
OBJ: Elaborate how an electrostatic potential map correlates to molecular structure and properties.

MSC: Evaluating

18. Peptide bonds are the building blocks of proteins. Consider a peptide bond formed from the amino acids alanine and serine, shown below. Draw the resonance forms and the resonance hybrid for the amide bond of the dipeptide. Use appropriate arrow notation.



ANS:



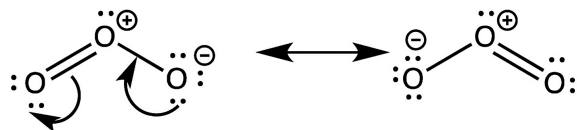
DIF: Moderate REF: 1.10 | 1.11

OBJ: Deduce and draw the resonance structures that contribute to the resonance hybrid, and vice versa.

MSC: Creating

19. Global warming and ozone depletion both have far-reaching environmental consequences and are linked by human cause. Propose two reasonable structures of ozone,  $O_3$ , that differ in electron delocalization and do not contain a ring. Show how these two structures are interconverted using appropriate arrow notation. Explain the relationship between these molecules.

ANS:



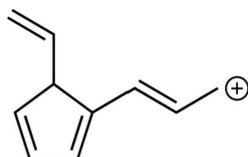
Use 18 valence electrons (O is in group VI;  $6 \times 3 = 18$ ) when drawing the structure. These two Lewis structures are resonance forms because they have the same atomic positions. The structures differ in the placement of electrons in multiple bonds or lone pairs.

DIF: Moderate REF: 1.10 | 1.11

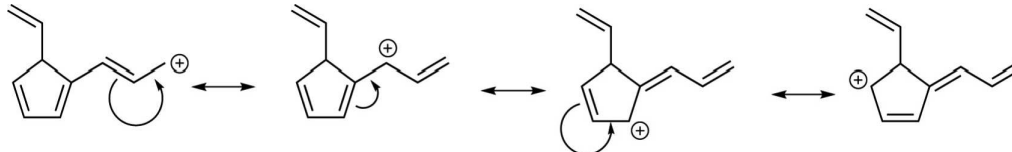
OBJ: Depict electron delocalization via resonance using appropriate arrow notation.

MSC: Creating

20. Draw three resonance structures for the following molecule. Use arrows to show interconversion between each structure.



ANS:

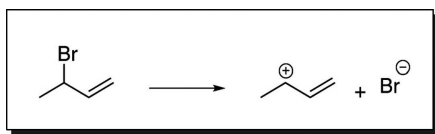


DIF: Moderate REF: 1.11

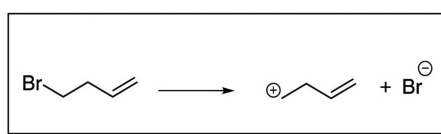
OBJ: Depict electron delocalization via resonance using appropriate arrow notation.

MSC: Applying

21. Which of the following is more likely to occur? Explain your answer referencing product stability.



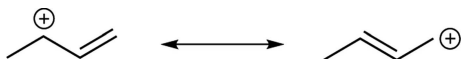
Option 1



Option 2

ANS:

Option 1 is more likely since the product is resonance stabilized. The resonance delocalizes charge and makes the product more stable. Since the product is more stable, option 1 is more likely to occur.

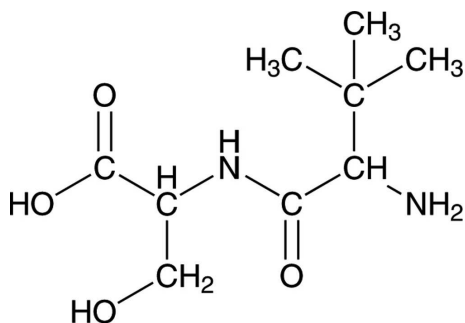


DIF: Difficult REF: 1.11 | 1.12

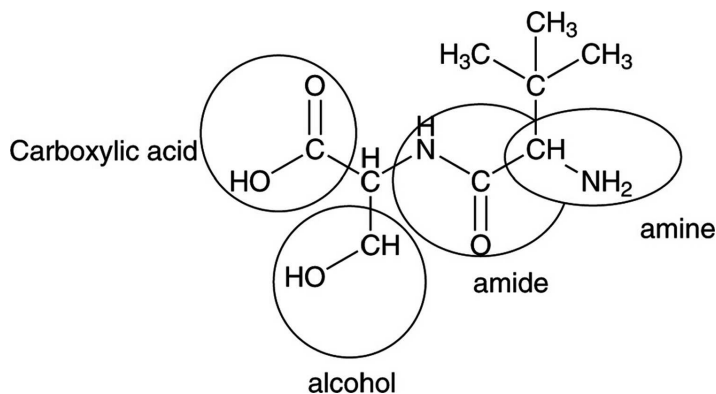
OBJ: Apply the concept of resonance to predict the outcome of a chemical reaction.

MSC: Evaluating

22. Identify the functional groups present in the structure of the dipeptide shown below. In your answer, do not use the word "alkane."



ANS:



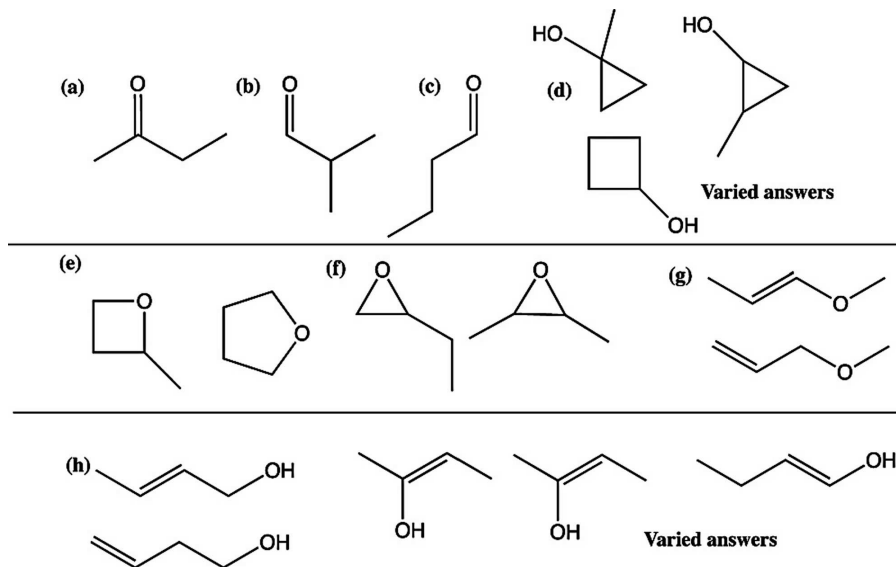
DIF: Moderate REF: 1.13

OBJ: Recognize and name functional groups within a complex molecule.

MSC: Analyzing

23. The way atoms are connected to each other in an organic structure determines the chemical behavior of the structure. Using line structures, propose individual molecules with the formula  $C_4H_8O$  that contain the following functional groups:
- (a) a ketone
  - (b) an aldehyde with a branched alkane
  - (c) an aldehyde with an unbranched alkane
  - (d) a cycloalkane
  - (e) a cyclic ether that is NOT an epoxide
  - (f) an epoxide
  - (g) an acyclic ether
  - (h) an alkene and an alcohol

ANS:

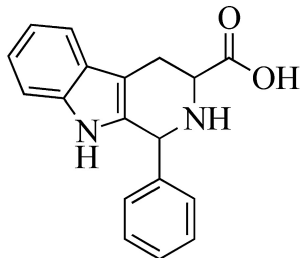


DIF: Difficult REF: 1.13

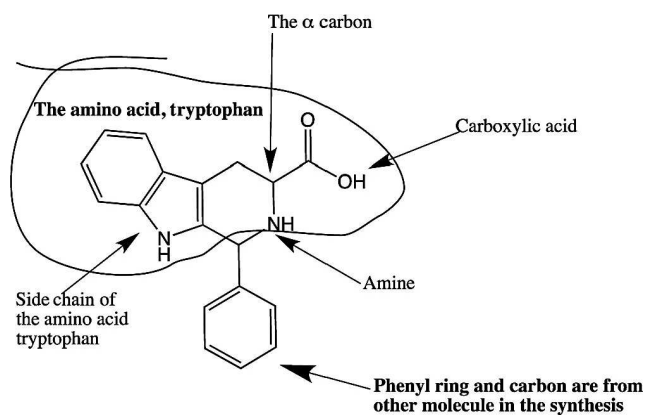
OBJ: Draw a structure of a given molecular formula that contains a specific functional group.

MSC: Creating

24. Amino acids are important building blocks in chemistry. Consider the following structure, which can be formed from reaction of an  $\alpha$ -amino acid with another compound. Identify and circle the  $\alpha$ -amino acid that is used in the synthesis. Clearly label the amino acid side chain, the alpha carbon, the amine, and the carboxylic acid within the larger structure. What parts of the larger molecule do not come from the amino acid?



ANS:

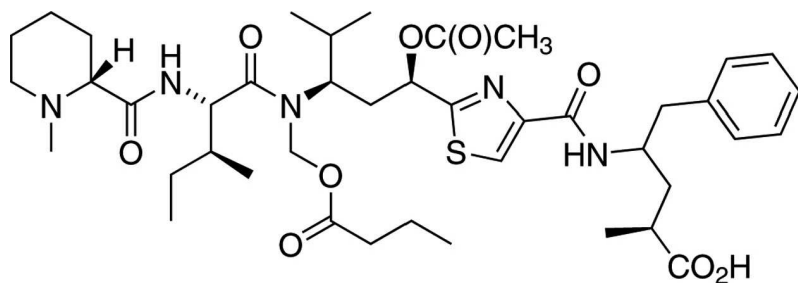


DIF: Difficult REF: 1.13 | 1.14

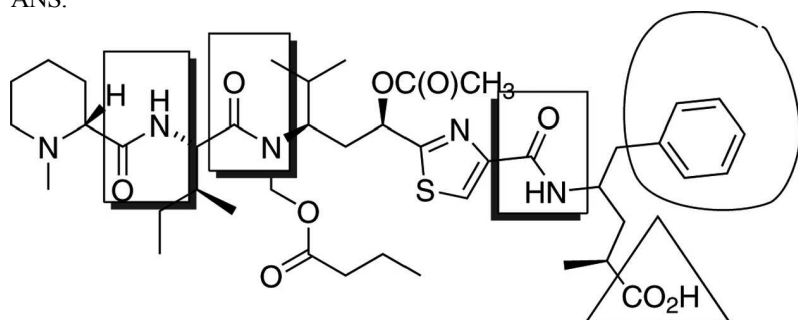
OBJ: Assimilate your knowledge of molecular structure to identify and/or draw organic functional groups.

MSC: Evaluating

25. Tubulysin D is a potent cytotoxic compound that interferes with mitosis. Identify the following structural features of tubulysin D.
- Place a box around any amide found in the molecule.
  - Circle the aryl group(s).
  - Place a triangle around the carboxylic acid(s).



ANS:

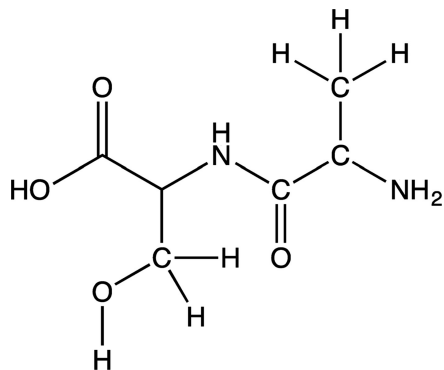


DIF: Moderate REF: 1.14

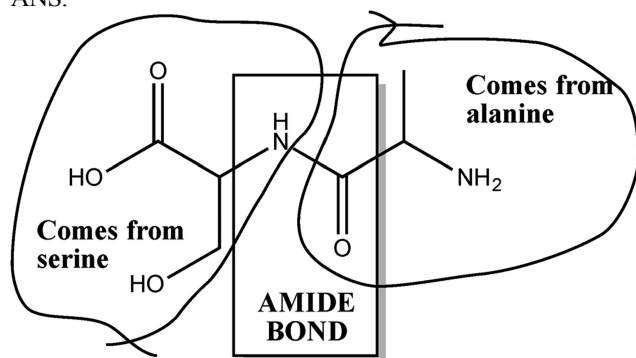
OBJ: Identify the key structural features of amino acids, saccharides, and nucleotides.

MSC: Analyzing

26. Consider a peptide bond formed from the amino acids alanine and serine. Redraw the dipeptide from the figure below using a line structure. Once this structure has been drawn, place a box around the amide bond. Circle the parts of the dipeptide that originate from serine and alanine.



ANS:

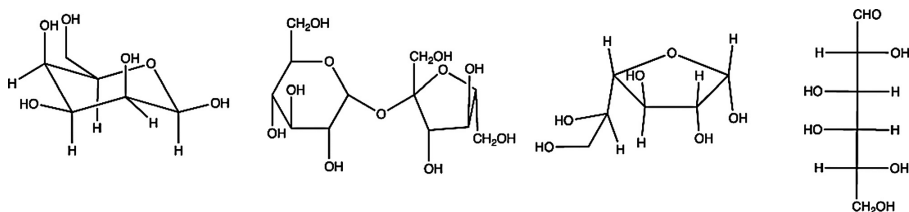


DIF: Moderate REF: 1.14

OBJ: Identify the key structural features of amino acids, saccharides, and nucleotides.

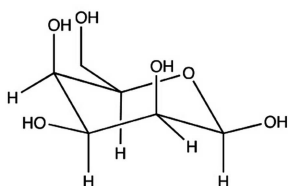
MSC: Understanding

27. Identify the molecules below that can be labeled as a monosaccharide.



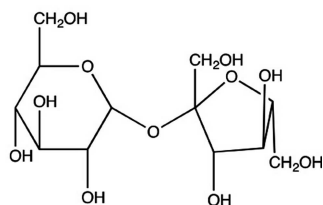
ANS:

**Cyclic  
monosaccharide**



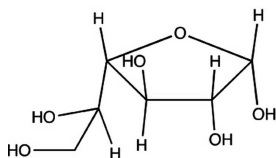
Chemical formula:  $C_6H_{12}O_6$

**Disaccharide**



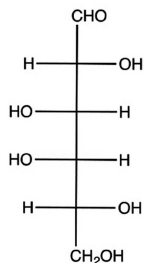
Chemical formula:  $C_{12}H_{22}O_{11}$

**Cyclic  
monosaccharide**



Chemical formula:  $C_6H_{12}O_6$

**Acyclic  
monosaccharide**



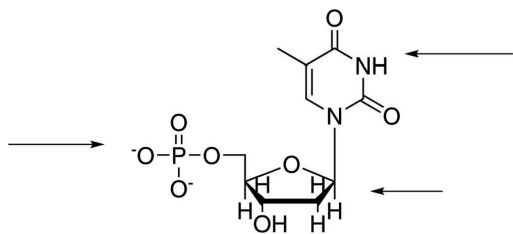
Chemical formula:  $C_6H_{12}O_6$

DIF: Moderate REF: 1.14

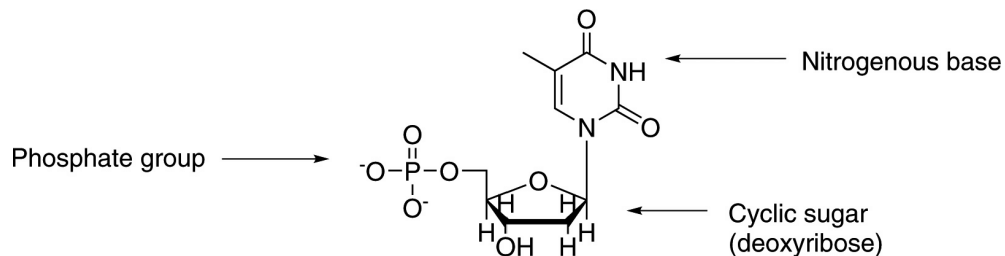
OBJ: Identify the key structural features of amino acids, saccharides, and nucleotides.

MSC: Understanding

28. Label the three distinct components in the following DNA nucleotide.



ANS:



DIF: Moderate REF: 1.14

OBJ: Identify the key structural features of amino acids, saccharides, and nucleotides

MSC: Understanding

29. How many hydrogen atoms are there for each oxygen atom in a carbohydrate?

ANS:

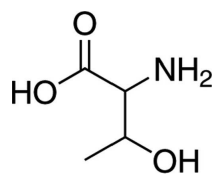
two

DIF: Easy REF: 1.14

OBJ: Identify the key structural features of amino acids, saccharides, and nucleotides.

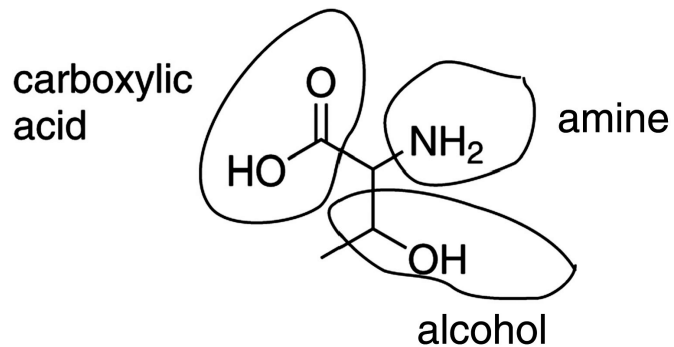
MSC: Remembering

30. Circle and identify three functional groups in threonine.



threonine

ANS:



DIF: Moderate REF: 1.14

OBJ: Identify the key structural features of amino acids, saccharides, and nucleotides.

MSC: Analyzing

## **Interchapter A: Nomenclature 1: The Basic System for Naming Simple Organic Compounds: Alkanes, Haloalkanes, Nitroalkanes, Cycloalkanes, and Ethers**

### **LEARNING OBJECTIVES**

Name alkanes and cycloalkanes using IUPAC nomenclature.

Name haloalkanes and nitroalkanes using IUPAC nomenclature.

Name ethers using IUPAC nomenclature.

Draw ethers when given an IUPAC name.

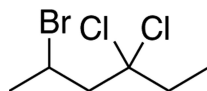
Classify carbon types.

Identify and name trivial alkyl groups.

Draw alkanes and cycloalkanes when given an IUPAC name.

## MULTIPLE CHOICE

1. What is the IUPAC name for the following molecule?



- a. 3,3-dichloro-5-bromohexane      d. 3,3-chloro-5-bromohexane  
b. 3-chloro-3-chloro-5-bromohexane      e. 4,4-dichloro-2-bromohexane  
c. 2-bromo-4,4-dichlorohexane

ANS: C

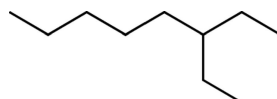
DIF: Easy

REF: A.3

OBJ: Name haloalkanes and nitroalkanes using IUPAC nomenclature.

MSC: Analyzing

2. What is the IUPAC name for the following molecule?



- a. 3-ethyloctane      d. 6,6-diethylpentane  
b. 3-pentylpentane      e. none of the above  
c. 6-ethyloctane

ANS: A

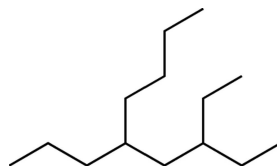
DIF: Easy

REF: A.4

OBJ: Name alkanes and cycloalkanes using IUPAC nomenclature.

MSC: Analyzing

3. What is the IUPAC name for the following molecule?



- a. 4-butyl-6-ethyloctane      d. 3-ethyl-5-propylnonane  
b. 3-ethyl-5-butyloctane      e. 5-propyl-7-ethyldecane  
c. 5-propyl-7-ethylnonane

ANS: D

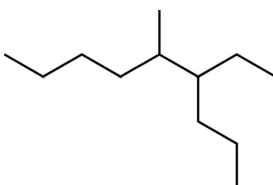
DIF: Moderate

REF: A.4

OBJ: Name alkanes and cycloalkanes using IUPAC nomenclature.

MSC: Analyzing

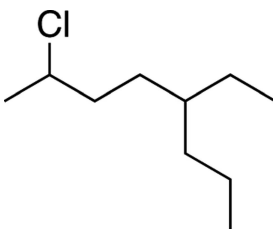
4. What is the IUPAC name for the following molecule?



- a. 5-methyl-6-propyloctane
- b. 3-propyl-4-methyloctane
- c. 2-butyl-3-propylpentane
- d. 4-ethyl-5-methylnonane
- e. 5-methyl-6-ethylnonane

ANS: D      DIF: Moderate      REF: A.4  
OBJ: Name alkanes and cycloalkanes using IUPAC nomenclature.  
MSC: Analyzing

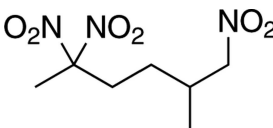
5. What is the IUPAC name for the following molecule?



- a. 3-propyl-6-chloroheptane
- b. 2-chloro-5-propylheptane
- c. 2-chloro-5-ethylheptane
- d. 4-ethyl-7-chlorooctane
- e. 2-chloro-5-ethyloctane

ANS: E      DIF: Easy      REF: A.4  
OBJ: Name haloalkanes and nitroalkanes using IUPAC nomenclature.  
MSC: Analyzing

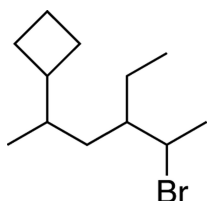
6. What is the IUPAC name for the following molecule?



- a. 1,6-trinitro-2-methylhexane
- b. 2-methyl-1,5,5-trinitrohexane
- c. 1,5,5-trinitro-2-methylhexane
- d. 2,2,6-trinitro-5-methylhexane
- e. 5-methyl-2,2,6-trinitrohexane

ANS: B      DIF: Moderate      REF: A.4  
OBJ: Name haloalkanes and nitroalkanes using IUPAC nomenclature.  
MSC: Analyzing

7. What is the IUPAC name for the following molecule?



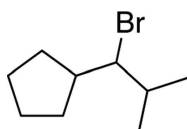
- a. 5-bromo-2-cyclobutyl-4-ethylhexane
- b. 2-cyclobutyl-5-bromo -4-ethylhexane
- c. 2-bromo-5-cyclobutyl-3-ethylheptane
- d. 2-bromo-5-cyclobutyl-3-ethylhexane
- e. 2-bromo-3-ethyl-5-cyclobutylhexane

ANS: D                      DIF: Moderate                      REF: A.5

OBJ: Name haloalkanes and nitroalkanes using IUPAC nomenclature.

MSC: Analyzing

8. What is the IUPAC name for the following molecule?



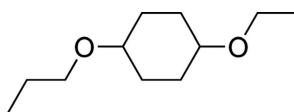
- a. 1-bromo-1-cyclopentyl-2-methylpropane
- b. 1-cyclopentyl-2-bromo-3-methylpropane
- c. 1-cyclopentyl-2-bromo-3-methylbutane
- d. 2-methyl-3-bromo-4-cyclopentylbutane
- e. (1-bromo-2-methylpropyl)cyclopentane

ANS: E                      DIF: Difficult                      REF: A.5

OBJ: Name haloalkanes and nitroalkanes using IUPAC nomenclature.

MSC: Analyzing

9. What is the IUPAC name for the following molecule?



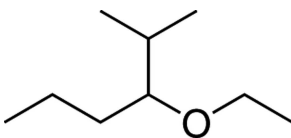
- a. propyloxy ethoxy cyclohexane ether
- b. 1,4-dipropoxyether
- c. 4-propoxycyclohexane-1ethoxycyclohexane
- d. 1-ethoxy-4-propoxycyclohexane
- e. none of the above

ANS: D                      DIF: Moderate                      REF: A.6

OBJ: Name ethers using IUPAC nomenclature.

MSC: Analyzing

10. What is the IUPAC name for the following molecule?



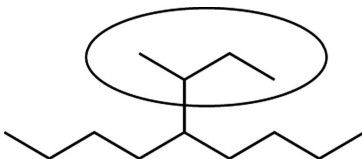
- a. 1-isopropyl-1ethoxybutane
- b. 1-ethoxy-1-isopropylbutane
- c. ethyl-2-isopropylether
- d. 3-ethoxy-2-methylhexane
- e. 2-methyl-3-ethoxyhexane

ANS: D      DIF: Moderate      REF: A.6

OBJ: Name ethers using IUPAC nomenclature.

MSC: Analyzing

11. What is the trivial name for the circled alkyl substituent?



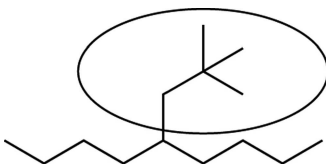
- a. *n*-butyl
- b. 2-butyl
- c. *sec*-butyl
- d. *tert*-butyl
- e. isobutyl

ANS: C      DIF: Easy      REF: A.7

OBJ: Identify and name trivial alkyl groups.

MSC: Remembering

12. What is the trivial name for the circled alkyl substituent?



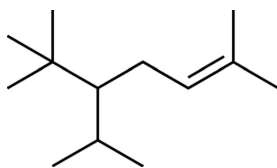
- a. *tert*-butyl
- b. *sec*-butyl
- c. isobutyl
- d. isopentyl
- e. neopentyl

ANS: E      DIF: Easy      REF: A.7

OBJ: Identify and name trivial alkyl groups.

MSC: Remembering

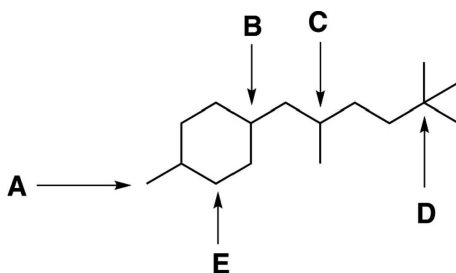
13. How many 4° carbons are in the following molecule?



- a. 1  
b. 2  
c. 3  
d. 4  
e. 5

ANS: A DIF: Moderate REF: A.7 OBJ: Classify carbon types.  
MSC: Remembering

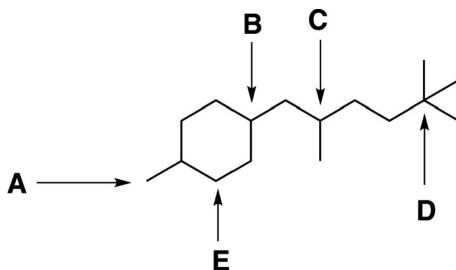
14. Which of the labeled carbon atoms can be classified as a 2° carbon?



- a. A  
b. B  
c. C  
d. D  
e. E

ANS: E DIF: Easy REF: A.7 OBJ: Classify carbon types.  
MSC: Remembering

15. Which of the labeled carbon atoms can be classified as a 1° carbon?

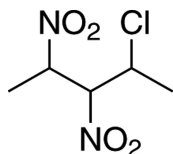


- a. A  
b. B  
c. C  
d. D  
e. E

ANS: A DIF: Easy REF: A.7 OBJ: Classify carbon types.  
MSC: Remembering

## SHORT ANSWER

1. What is the IUPAC name for the following molecule?



ANS:

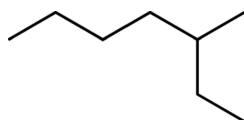
2-chloro-3,4-dinitropentane

DIF: Moderate REF: A.3

OBJ: Name haloalkanes and nitroalkanes using IUPAC nomenclature.

MSC: Applying

2. What is the IUPAC name for the following molecule?



ANS:

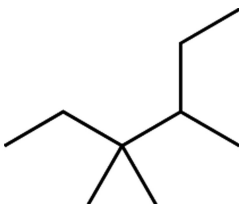
3-methylheptane

DIF: Easy REF: A.4

OBJ: Name alkanes and cycloalkanes using IUPAC nomenclature.

MSC: Applying

3. What is the IUPAC name for the following molecule?



ANS:

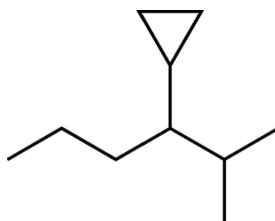
3,3,4-trimethylhexane

DIF: Moderate REF: A.4

OBJ: Name alkanes and cycloalkanes using IUPAC nomenclature.

MSC: Applying

4. What is the IUPAC name for the following molecule?



ANS:

3-cyclopropyl-2-methylhexane

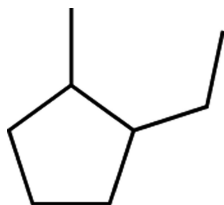
DIF: Easy

REF: A.5

OBJ: Name alkanes and cycloalkanes using IUPAC nomenclature.

MSC: Applying

5. What is the IUPAC name for the following molecule?



ANS:

1-ethyl-2-methylcyclopentane

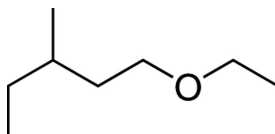
DIF: Easy

REF: A.5

OBJ: Name alkanes and cycloalkanes using IUPAC nomenclature.

MSC: Applying

6. What is the IUPAC name for the following molecule?



ANS:

1-ethoxy-3-methylpentane

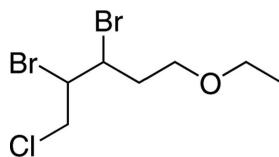
DIF: Easy

REF: A.6

OBJ: Name ethers using IUPAC nomenclature.

MSC: Applying

7. What is the IUPAC name for the following molecule?



ANS:

2,3-dibromo-1-chloro-5-ethoxypentane

DIF: Moderate

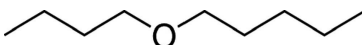
REF: A.6

OBJ: Name ethers using IUPAC nomenclature.

MSC: Applying

8. Draw 1-butoxypentane.

ANS:



DIF: Easy

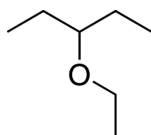
REF: A.6

OBJ: Draw ethers when given an IUPAC name.

MSC: Applying

9. Draw 3-ethoxypentane.

ANS:



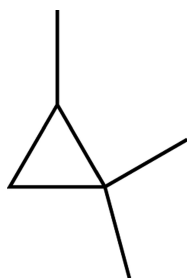
DIF: Moderate

REF: A.6

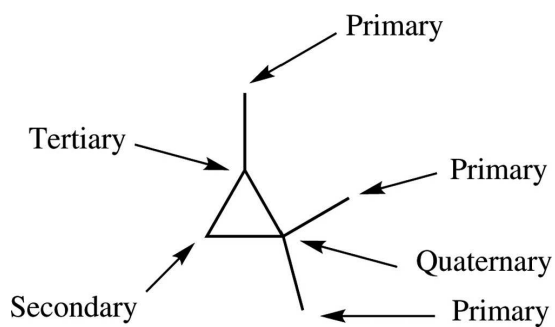
OBJ: Draw ethers when given an IUPAC name.

MSC: Applying

10. Label each carbon in the following molecule as primary, secondary, tertiary, or quaternary.



ANS:



DIF: Easy

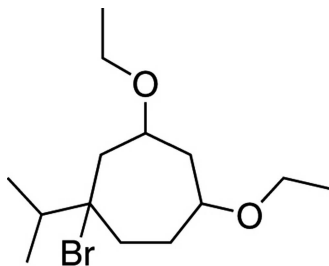
REF: A.7

OBJ: Classify carbon types.

MSC: Remembering

11. Draw 1-bromo-3,5-diethoxy-1-isopropylcycloheptane.

ANS:



DIF: Difficult

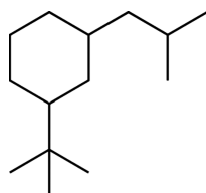
REF: A.7

OBJ: Draw ethers when given an IUPAC name.

MSC: Applying

12. Draw 1-(*tert*-butyl)-3-isobutylcyclohexane.

ANS:

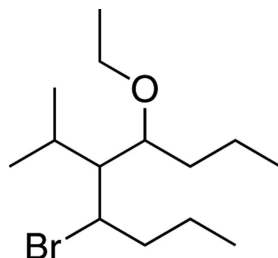


DIF: Moderate REF: A.7

OBJ: Draw alkanes and cycloalkanes when given an IUPAC name.

MSC: Applying

13. What is the IUPAC name for the following molecule?



ANS:

4-bromo-6-ethoxy-5-isopropylnonane

DIF: Difficult

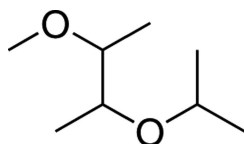
REF: A.7

OBJ: Name ethers using IUPAC nomenclature.

MSC: Applying

14. Draw 2-isopropoxy-3-methoxybutane.

ANS:



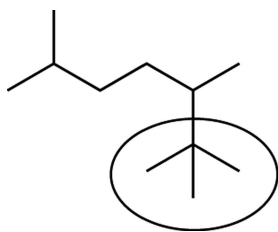
DIF: Moderate

REF: A.7

OBJ: Draw ethers when given an IUPAC name.

MSC: Applying

15. Give the trivial name for the circled alkyl group.



ANS:

*t*-butyl or *tert*-butyl

DIF: Easy

REF: A.7

OBJ: Identify and name trivial alkyl groups.

MSC: Remembering