GRANDE PRAIRIE REGIONAL COLLEGE DEPARTMENT OF SCIENCE: CHEMISTRY

FORTY-FOURTH SESSION 2009 – 2010

COURSE OUTLINE: ORGANIC CHEMISTRY

CH2610 A2

CH2610 A2: Organic Chemistry I; Prerequisite, CH1010 or CH1030

INSTRUCTOR: Dr. John P. Sloan

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E-mail SLOAN@GPRC.AB.CA

LECTURE: CH2610 T, R 11:30 – 12:50 in J204

ALBERTA TRANSFER CREDIT

(Ref: 2009-20010 Guide to Transfer Credit at Alberta Post-Secondary Institutions)

GPRC: CH2610 (3)

U of Alberta: CHEM 261 (3) or AUCHE 250 (3)

U of Calgary:
CHEM 351 (3)
U of Lethbridge:
CHEM 2500 (3)
Athabasca U:
Canadian UC:
Chem 241 (4)
Concordia UC:
CHEM 261 (3)

COURSE OUTLINE:

LECTURE COMPONENT:

A study of the fundamental principles of the chemistry of carbon compounds. The study is based on a reaction mechanism approach to the functional group chemistry of alkanes, alkenes, alkynes, cycloalkanes, alkyl halides, alcohols and ethers. Topics include: structure and bonding; physical properties; acidity and basicity; conformations of molecules; stereochemistry; addition, elimination and substitution reactions; structure-reactivity relationships; and introduction to methods for structure determination.

A representative selection of molecules found in agricultural, biological, environmental, industrial, medical, and pharmatheutical applications of organic chemistry will be discussed, e.g., molecules found in agrochemicals, fibres, food additives, perfumes, polymers, and prescription drugs.

LABORATORY COMPONENT:

Laboratory Techniques in organic chemistry; preparation of some organic compounds, and; methods of qualitative organic analysis.

TUTORIAL COMPONENT:

Problem solving and discussion sessions with short problem sets for completing and marking during the tutorial. In addition, weekly assignments consisting of 10 questions per assignment will be given. These assignments will consist of exam type questions and do not need to be submitted for marking. Detailed solutions to the assignments will be posted on Blackboard about 1 week after distribution.

NOTES:

- 1. Lectures, Time and Place CH2610 A2 T, R 11:30 - 12:50 in J204
- 2. Laboratory Component, Time and Place CH2610 L1 T 14:30 - 17:20 in J116
- 3. Tutorial Component, Time and Place CH2610 S1 F 8:30 - 9:20 in J204
- 4. Office Hours: Individual and group assistance will normally be available in office J207 during regular college business hours outside of formal class lecture, laboratory and tutorial hours.

TEXT BOOKS AND LABORATORY ITEMS:

The following text book is required:

CH2610

Solomons, T.W.G., and C.B. Fryhle, *Organic Chemistry*, 9th Edition, Wiley, 2008

And

A Three Ring Binder to Hold: Sloan, J.P., *Organic Chemistry Experiments, Chemistry 2610/2630*, Grande Prairie Regional College, 2009/2010.

Molecular Models are highly recommended, namely:

Molecular Model Set for Organic Chemistry, Prentice Hall.

Study Guides, Solutions Manuals, and Wiley Plus are supplementary items, namely:

- 1. Fernandez, J.E., and Solomons, T.W.G., *Study Guide and Solutions Manual to Organic Chemistry*, 9th Edition, 2008;
- 2. Wiley Plus.

Note:

1. All required and supplementary books, molecular structure model sets, safety glasses, and lab coats are available at the College Bookstore. *Organic Chemistry Experiments*, by J.P. Sloan, will be given as handouts in advance of each lab period. These are to be inserted in a three ring binder.

EVALUATION:

Examination Schedule and Composition of the Final Grade:

1.	Midterm Exam # 1, Friday October 9	15%
2.	Midterm Exam # 2, Friday November 13	20%
2.	Final Exam to be scheduled between December 9 - 18	30%
3.	Laboratory	25%
4.	Tutorial Grading Component	10%
	1	00%

The Grades are based on the alpha grading system. The Registrar's Office will convert alpha grades to four-point equivalence for the calculation of grade point averages. Alpha grades, 4-point equivalence, and grade descriptors are as follows:

Alpha	4-Point Equivalence	Descriptor
Grade		
A^{+}	4.0	Excellent
A	4.0	
A-	3.7	Very Good
B+	3.3	First Class Standing
В	3.0	Good
B-	2.7	
C+	2.3	Satisfactory
С	2.0	
C-	1.7	
D+	1.3	Poor*
D	1.0	Minimal Pass*
F	0.0	Failure

^{*} Other post secondary institutions may not award transfer credit for grades of D and D+.

Notes:

1. The Mid-Term Exams will be of 1.5 hours duration and the Final Exam will be of 3 hours duration.

- 2. Between 5 and 15% of exam content will be taken from a combination of weekly assignments, Wiley Plus, and questions in the organic chemistry textbook by Solomons and Fryhle.
- 3. A pass grade is essential for the Laboratory Component.
- 4. The Tutorial Grading Component will contribute to 10% of the final grade and will consist of two components as determined by the Instructor with input from the class. The two components are;
 - 4.1 Printed assignments consisting of ten questions per assignment, and;
 - 4.2 On-line Wiley Plus tests prepared by the Instructor and marked electronically, with feedback to the students.
- 5. Assistance with assignments will be given upon request.
 - 5. Regular attendance in Lecture, Laboratory, and Tutorial Components is a Course Requirement.

Grande Prairie Regional College Calendar 2009 - 2010: Course Description (p 185).

CH2610 3(3-1-3)UT, 105 Hours, Organic Chemistry I

The correlation of structure and bonding in carbon compounds with the physical properties and chemical reactivity of organic molecules. Discussion will be based on functional groups with emphasis on hydrocarbons and derivatives that contain halogens, oxygen, sulphur and the hydroxyl group. Introduction to stereochemistry, three dimensional structure, reaction mechanisms, especially addition to double bonds, nucleophilic substitution and elimination reactions, and methods of structure determination. The study covers the functional group chemistry of alkanes, alkenes, alkynes, alcohols, ethers and sulfides.

Prerequisites: CH1010 or CH1030

Notes: Credit will be granted for only one of CH1610 or CH2610

Transfer: UA, UC, UL, AU, AF, CU, CUC, KUC

CHEMISTRY 2610: READING, STUDYING, AND PRACTICE PROBLEMS

All references are to T.W.G. Solomons and C.B. Fryhle, *Organic Chemistry*, 9th Edition, Wiley, 2008.

FALL SEMESTER

Weeks of

Sept 3 & 7: THE BASICS: Bonding and Molecular Structure

Molecular Graphic: Glycine, an organic molecule found in space

Sect # Page # Read and Study Chapter 1 "We are Star Dust"

1.1	2	Organic Chemistry and Life
1.2	3	The Structural Theory of Organic Chemistry
1.3	4	Isomers: The Importance of Structural Formulas
1.4	5	Chemical Bonds: The Octet Rule
1.5	7	Writing Lewis Structures
1.6	9	Exceptions to the Octet Rule
1.7	10	Formal Charge
1.8	13	Resonance Theory
1.8A	15	Summary of Rules for Resonance
1.9	18	Quantum Mechanics and Atomic Structure
1.10	20	Atomic Orbitals and Electron Configuration:
1.10A	21	Aufbau Principle; the Pauli Exclusion Principle; Hund's Rule
1.11	21	Molecular Orbitals: Bonding and Antibonding
1.12	24	The Structure of Methane and Ethane: sp ³ Hybridization;
1.12A	24	The Structure of Methane
1.12B	27	The Structure of Ethane
1.13	28	The Structure of Ethene (Ethylene): sp ² Hybridization
1.13A	31	Restricted Rotation and the Double Bond
1.13B	32	Cis-Trans Isomers
1.14	33	The Structure of Ethyne (Acetylene): sp Hybridization
1.14A	34	Bond Lengths of Ethyne, Ethene, and Ethane
1.15	35	A Summary of Important Concepts that Come from Quantum Mechanics
1.16	36	Molecular Geometry: The Valence Shell Electron-Pair Repulsion (VSEPR) Model.
1.16A	-F 37	Molecular Geometry: VSEPR Models for Methane, Ammonia, Water, Boron Trifluoride,
		Berylium Hydride and Carbon Dioxide
1.17	39	Representation of Structural Formulas: Dash; Condensed; Bond Line; and the Three
		Dimensional Wedge, Dash, Line Representation
1.18	44	Applications of Basic Principles: Opposite Charges Attract; Like Charges repel; Nature
		Tends Towards States of Lower Potential Energy; Orbital Overlap Stabilizes Molecules
	45	Key Terms and Concepts
	46	Concept Map

Practice Problems: You are encouraged to work all of the in-chapter problems, and you are required to complete the assignments given in-seminar-class and from Wiley Plus. Routinely doing problems in organic chemistry leads to understanding of the theory, and good grades in organic chemistry.

In the words of Solomons and Fryhle:

"One way to check your progress is to work each of the in-chapter problems when you come to it. These problems have been written just for this purpose and are designed to help you decide whether or not you understand the material that has just been explained."

And, in the words of Wade:

"It's easy to fool yourself into thinking you understand organic chemistry when you actually do not. As you read through this book, all the facts and ideas may make sense, yet you have not learned to combine and use those facts and ideas. An examination is a painful time to learn that you do not really understand the material.

The best way to understand organic chemistry is to use it. You will certainly need to read and reread all the material in the chapter, but this level of understanding is just the beginning. Problems are provided so you can work with the ideas, applying them to new compounds and new reactions that you have never seen before. By working problems, you force yourself to use the material and fill in the gaps in your understanding. You also increase your level of self-confidence and your ability to do well on exams".

Problems: In-Chapter 1.1 to 1.15

- 47 End of Chapter 1.16 to 1.38
- 50 Learning Group Problem

Week of Sept 14: REPRESENTATIVE CARBON COMPOUNDS: Functional Groups, Intermolecular Forces, and Infrared (IR) Spectroscopy

- 51 Structure and Function: Organic Chemistry, Nanotechnology, and Bioengineering
- 2.1 52 Carbon-Carbon Covalent Bonds
- 2.2 52 Hydrocarbons: Representative, Alkanes, Alkenes, Alkynes, and Aromatic Compounds
- 2.3 55 Polar Covalent Bonds
- 2.4 56 Polar and Nonpolar Molecules
- 2.4A 58 Dipole Moments in Alkenes
- 2.5 59 Functional Groups
- 2.5A 59 Alkyl Groups and the Symbol R
- 2.5B 60 Phenyl and Benzyl Groups
- 2.6 60 Alkyl Halides or Haloalkanes
- 2.7 61 Alcohols, including Classification as Primary, Secondary and Tertiary (1E, 2E, 3E)
- 2.8 63 Ethers
- 2.9 Amines, including Classification as Primary, Secondary and Tertiary
- 2.10 65 Aldehydes and Ketones
- 2.11 65 Carboxylic Acids, Esters, and Amides
- 2.12 67 Nitriles
- 2.13 68 Summary of Important Families of Organic Compounds
- 2.14 68 Physical Properties and Molecular Structure with emphasis on Intermolecular Interactions, namely:
- 2.14A 69 Ion-Ion Forces in ionic compounds, e.g. sodium acetate, sodium chloride

2.14B 2.14C 2.14D 2.14E 2.14F 2.14G	70 71 73 74	Dipole-Dipole Forces resulting from permanent dipoles, e.g. acetone, chloromethane Hydrogen Bonds van der Waals Forces, or London forces or dispersion forces, e.g. methane Solubilities Guidelines for Water Solubility Intermolecular Forces in Biochemistry, and Organic Templates Engineered to Mimic Bone
2.15	75	Growth Summary of Attractive Floating Forese
2.152.16	75 76	Summary of Attractive Electric Forces Infrared Spectroscopy: An Instrumental Method for Detecting Functional Groups
2.16A		Infrared Spectroscopy. An instrumental Method for Detecting Functional Groups Infrared Spectra of Hydrocarbons
2.16B		IR Spectra of Some Functional Groups Containing Heteroatoms including Carbonyl Functional Groups of Aldehydes, Ketones, Esters, Carboxylic Acids and Amides, plus Alcohols, Phenols and Amines
2.17	84	Applications of Basic principles: Polar Bonds are Caused by Electronegativity Differences; Opposite Charges Attract; Molecular Structure Determines Properties
	85	Key Terms and Concepts
	86	Concept Map
Proble	ms: 87 90	In-Chapter 2.1 to 2.19 End of Chapter 2.20 to 2.48 Learning Group Problem

Week of Sept 21: AN INTRODUCTION TO ORGANIC REACTIONS: ACIDS AND BASES IN ORGANIC CHEMISTRY

		ACIDS AND DASES IN ORGANIC CHEMISTRY
		Read and Study Chapter 3
	91	Diamox, a drug that prevents altitude sickness
	91	Shuttling the Protons, or, from the Lewis and Sloan perspective, Shuttling the Electrons
3.1	92	Reactions and their Mechanisms - Substitution, Addition, Elimination and Rearrangement
		Reactions
3.1A	92	Homolysis and Heterolysis of Covalent Bonds, and Introduction to the Use of Curved
		Arrows
3.2	94	Acids and Bases
3.2A	94	The BrNnsted-Lowry Definition of Acids and Bases
3.2B	95	The Lewis Definition of Acids and Bases
3.2C	96	Opposite Charges Attract
	97	The Chemistry of HOMOs and LUMOs in Reactions
3.3	97	Heterolysis of Bonds to Carbon - Carbocations and Carbanions
3.4	98	The Use of Curved Arrows in Illustrating Reactions
3.5	100	The Strength of Acids and Bases, K _a and pK _a
3.5A	100	The Acidity Constant, K _a
3.5B	100	Acidity and pK _a
	101	Table 3.1: Relative Strength of Selected Acids and Their Conjugate Bases
3.5C	102	Predicting the Strength of Bases
		the Stronger the Acid, the Weaker the Conjugate Base
3.6	103	Predicting the Outcome of Acid-Base Reactions
3.6A	104	Water Solubility as a Result of Salt Formation

3.7	105	The Relationship between Structure and Acidity, i.e. Structural Effects on Acidity and
		Basicity, namely:
		1. Size Effect, acidity increases upon descending a column in the Periodic Table,
		H-I is a stronger acid than H-F; the acidity order is: H-I > H-Br > H-Cl > H-F
		2. Electronegativity Effect, acidity increases from left to right in the Periodic Table,
		H-F is a stronger acid than CH_4 ; the acidity order is: $HF > H_20 > NH_3 > CH_4$
3.7A	107	3. The Effect of Hybridization, more s-character means the anion has lower energy, is more stable, and is a weaker base
3.7B	108	4. Inductive Effects, from polarization by electron attracting and electron withdrawing
		groups
3.8	108	Energy Changes; higher potential and kinetic energy implies less stable, lower energy
		implies more stable
3.8A	109	Potential Energy and Covalent Bonds, exothermic reactions give out heat, endothermic
		reactions absorb heat
3.9	110	The Relationship Between the Equilibrium Constant and the Standard Free-Energy
		Change, ΔG^{o} ; a negative value favours products at equilibrium
3.10	111	The Acidity of Carboxylic Acids, with explanations arising from Resonance Effects and
		Inductive Effects
3.10A	112	The Effect of Delocalization: An Explanation based on Resonance Effects, due to
		resonance stabilization of the carboxylate anion
3.10B	113	An Explanation based on Inductive Effects, due to inductive withdrawal of electronic
		charge by –O and -C=O in carboxylate anions
3.10C	114	Summary of a Comparison of Conjugate Acid-Base Strengths
3.10D	114	Inductive Effects of Other Groups
3.11	115	The Effect of Solvent on Acidity - Protic Solvents
3.12	116	Organic Compounds as Bases
3.13	117	A Mechanism for an Organic Reaction
	118	The Chemistry of carbonic Anhydrase
3.14	119	Acid and Base in Nonaqueous Solutions
3.15	120	Acid-Base Reactions, and Synthesis of Deuterium- and Tritium-Labelled Compounds
3.16	121	Applications of Basic Principles: Electronegativity Differences Polarize Bonds; Polarized
		Bonds Underlie Inductive Effects; Opposite Charges Attract; Nature Prefers States of
		Lower Potential Energy; Resonance Effects Can Stabilize Molecules and Ions
	122	Key Terms and Concepts
	123	Concept Map
Proble	ms:	In-Chapter 3.1 to 3.14
	124	End of Chapter 3.15 to 3.42

Week of Sept 28: NOMENCLATURE AND CONFORMATIONS OF ALKANES AND CYCLOALKANES

Read and Study Chapter 4

Learning Group Problem

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4.1

- To be Flexible or Inflexible Molecular Structure Makes the Difference
 Introduction to Alkanes and Cycloalkanes
- 4.1A 130 Sources of Alkanes: Petroleum
 - 130 The Chemistry of Petroleum Refining

	131	Typical Fractions Obtained by Distillation of Petroleum
4.2	132	Shapes of Alkanes
	133	Tables 4.1: Physical Constants of Hexane Isomers
	134	Table 4.2: Number of Alkane Isomers
	135	Table 4.3: The Unbranched Alkanes
4.3	134	IUPAC Nomenclature of Alkanes, Alkyl Halides and Alcohols
4.3A	135	Nomenclature of Unbranched Alkyl Groups
4.3B	135	Nomenclature of Branched-Chain Alkanes
4.3C	137	Nomenclature of Branched Alkyl Groups
4.3D	138	Classification of Hydrogen Atoms,
		as Primary (1°), Secondary (2°), and Tertiary (3°)
4.3E	139	Nomenclature of Alkyl Halides
4.3F	139	Nomenclature of Alcohols
4.4	141	Nomenclature of Cycloalkanes
4.4A	141	Monocyclic Compounds
4.4B	142	Bicyclic Compounds
4.5	143	Nomenclature of Alkenes and Cycloalkenes
4.6	145	Nomenclature of Alkynes
4.7	146	Physical Properties of Alkanes and Cycloalkanes
	148	The Chemistry of Pheromones: Communication by Means of Chemicals
4.8	148	Sigma (Φ) Bonds and Bond Rotation
4.9	151	Conformational Analysis of Butane
4.10	153	The Relative Stability of Cycloalkanes: Ring Strain
4.10A		Heats of Combustion
4.10B		Heats of Combustion of Cycloalkanes
	154	Table 4.5: Heats of Combustion and Ring Strain of Cycloalkanes
4.11	155	The origin of Ring Stain in Cyclopropane and Cyclobutane: Angle Strain and Torsional
.,,,	100	Strain
4.11A	155	Cyclopropane
4.11B		Cyclobutane
4.11C		Cyclopentane
4.12		Conformations of Cyclohexane
4.12A		Conformations of Higher Cycloalkanes
.,,_,,	159	The Chemistry of Nanoscale Motors and Molecular Switches
4.13	160	Substituted Cyclohexanes, Axial and Equatorial Hydrogen Atoms
4.14	163	Disubstituted Cyclohexanes, Cis-Trans Isomerism
4.14A		Cis-Trans Isomerism and Conformational Structures
4.15	166	Bicyclic and Polycyclic Alkanes
	167	The Chemistry of Elemental Carbon
4.16	168	Chemical Reactions of Alkanes
4.17	168	Synthesis of Alkanes and Cycloalkanes
4.17A		Hydrogenation of Alkenes and Alkynes
4.18	169	Structural Information from Molecular Formulas and the Index of Hydrogen Deficiency
4.18A		Compounds Containing Halogens, Oxygen, or Nitrogen
4.19	171	¹³ C NMR Spectroscopy- A Practical Introduction
4.19A		One Signal for each Unique Carbon
4.19B		Chemical Shift – Location of the Signal Depends on Electronic Environment
4.19C		Using ¹³ C NMR to Elucidate Structure
	175	Application of Basic Principles: Nature Prefers States of Lower Energy;

- **Key Terms and Concepts** 176
- Concept Maps 177

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Concept Map

Problems:

- In-Chapter 4.1 to 4.21 End of Chapter 4.22 to 4.54 178
- Learning Group Problems 180

STEREOCHEMISTRY: CHIRAL MOLECULES Week of Oct 5:

		Read and Study Chapter 5
	181	The Handedness of Life
5.1	182	The Biological Significance of Chirality
5.2	183	Isomerism, Constitutional Isomers and Stereoisomers
5.3	184	Enantiomers and Chiral Molecules
5.4	187	More about the Biological Importance of Chirality
5.5	188	The Historical Origin of Stereochemistry
5.6	189	Tests for Chirality, Planes of Symmetry and Points of Symmetry
5.7	190	Nomenclature of Enantiomers: The R-S System
5.8	194	Properties of Enantiomers, Optical Activity
5.8A	195	Plane-Polarized Light
5.8B	195	The Polarimeter
5.8C	195	Specific Rotation
5.9	198	The Origin of Optical Activity
5.9A	199	Racemic Forms
5.9B	199	Racemic Forms and Enantiomeric Excess
5.10	200	The Synthesis of Chiral Molecules
5.10A	200	Racemic Forms
5.10B	201	Stereoselective Synthesis
5.11	202	Chiral Drugs
	203	The Chemistry of: Selective Binding of Drug Enantiomers to Left- and Right-Hand Coiled
		DNA
5.12	203	Molecules with More Than One Chirality Centree
5.12A		Meso Compounds
5.12B		Naming Compounds with More than One Chirality Centre
5.13	207	Fischer Projection Formulas
5.14	209	Stereoisomerism of Cyclic Compounds
5.14A		Cyclohexane Derivatives
5.15	211	Relating Configurations Through Reactions in Which No Bonds to the Chirality Centre are
		Broken
5.15A		Relative and Absolute Configurations
5.16	213	Separation of Enantiomers: Resolution
5.16A		Pasteur's Method for Separating Enantiomers
5.16B		Current Methods for Resolution of Enantiomers
5.17	214	Compounds with Chirality Centres Other than Carbon
5.18	215	Chiral Molecules that do not Possess a Chirality Centre (a Tetrahedral Atom with Four
		Different Groups Attached)
	216	Key Terms and Concepts

Problems: In-Chapter 5.1 to 5.29

- 218 End of Chapter 5.30 to 5.44
- 220 Learning Group Problems

Additional Problems - The CD accompanying the text book includes a set of computer

molecular model stereochemistry exercises that are keyed to the text

Weeks of Oct 12 & 19: IONIC REACTIONS: Nucleophilic Substitution and Elimination Reactions of Alkyl Halides

	221	Breaking Bacteria Cell Walls With Organic Chemistry
6.1	222	Organic Halides
	222	Table 6.1: Carbon-Halogen Bond lengths and Bond Strengths
6.1A	223	Physical Properties of Organic Halides
	223	Table 6.2: Organic Halides
6.2	224	Nucleophilic Substitution Reactions
6.3	224	Nucleophiles
6.4	225	Leaving Groups
6.5	226	Kinetics of a Nucleophilic Substitution Reaction -
		a Substitution Nucleophilic Bimolecular (S _N 2) Reaction
6.6	227	A Mechanism for the S_N 2 Reaction
6.7	228	Transition State Theory: Free-Energy Diagrams
6.8	229	The Stereochemistry of S _N 2 Reactions
6.9	235	The Reaction of Tert-Butyl Chloride with Hydroxide Ion: An S _N 1 Reaction
6.9A	235	Multistep Reactions and the Rate-Determining Step
6.10	236	A Mechanism for the S_N1 Reaction
6.11	237	Carbocations
6.11A	237	The Structure of Carbocations
6.11B	238	The Relative Stabilities of Carbocations
6.12	239	The Stereochemistry of S _N 1 Reactions
6.12A	239	Reactions That Involve Racemization
6.12B	240	Solvolysis – Cleavage of the Solvent by the Nucleophile
6.13	241	Factor's Affecting the Rates of S _N 1 and S _N 2 Reactions
6.13A	241	The Effect of the Structure of the Substrate
	241	Table 6.4: Relative Rates of Reactions of Alkyl Halides in S _N 2 Reactions
	243	S _N 1 Reactions and the Hammond-Leffler Postulate
6.13B	244	The Effect of the Concentration and the Strength of the Nucleophile
	244	Nucleophilicity versus Basicity
6.13C	245	Solvent Effects on S _N 2 Reactions: Polar Protic and Aprotic Solvents
6.13D	247	Solvent Effects on S _N 1 Reactions: The Ionizing Ability of the Solvent
	247	Table 6.5: Dielectric Constants of Common Solvents
6.13E	247	The Nature of the Leaving Group
	249	Summary of $S_N 1$ versus $S_N 2$
	249	Table 6.6: Factors Favouring S_N1 versus S_N2 Reactions
6.14	250	Organic Synthesis - Functional Group Transformations Using S _N 2 Reactions
	251	The Chemistry of Biological Methylation: A Biological Nucleophilic Substitution
		Reaction

6.14A	252	The Unreactivity of Vinylic and Phenyl Halides
6.15	253	Elimination Reactions of Alkyl Halides
6.15A	253	Dehydrohalogenation (loss of H-X)
6.15B	254	Bases Used in Dehydrohalogenation
6.15C	255	Mechanisms of Dehydrohalogenation: E2 and E1 Mechanisms
6.16	255	The Elimination-Bimolecular (E2) Reaction
6.17	256	The Elimination-Unimolecular (E1) Reaction
6.18	257	Substitution versus Elimination
6.18A	257	S _N 2 versus E2
6.18B	259	Tertiary Halides: S _N 1 versus E1
6.19	260	Overall Summary
	260	Table 6.7: Overall Summary of S _N 1, S _N 2, E1 and E2 Reactions
	261	Summary and Review Tools
	262	Key Terms and Concepts
Proble	ms:	In-Chapter 6.1 to 6.12
	252	End of Chapter 6.13 to 6.48
	268	Learning Group Problems

Week of Oct 26: ALKENES AND ALKYNES I: Properties and Synthesis. Elimination Reactions of Alkyl Halides

	269	Cell Membrane Fluidity
7.1	270	Introduction
7.1A	270	Physical Properties of Alkenes and Alkynes
7.2	270	The (E) - (Z) System for Designating Alkene Diastereomers
7.3	272	Relative Stabilities of Alkenes
7.3A	272	Heat of Reaction
	272	Figure 7.2: Order of Stability of Alkenes from Heats of Hydrogenation
7.3B	273	Overall Relative Stabilities of Alkenes
7.4	274	Cycloalkenes
7.5	274	Synthesis of Alkenes via Elimination Reactions
7.6	275	Dehydrohalogenation of Alkyl Halides
7.6A	275	Zaitsev's Rule: Formation of the Most Substituted Alkene is Favoured with a Small Base
7.6B	277	Formation of the Less Substituted Alkene Using a Bulky Base
7.6C	278	The Stereochemistry of E2 Reactions: The Orientation of Groups in the Transition State
7.7	280	Acid-Catalyzed Dehydration of Alcohols
7.7A	281	Mechanism for Dehydration of Secondary and Tertiary Alcohols: An E1 Reaction
7.7B	282	Carbocation Stability and the Transition State
7.7C	284	A Mechanism for Dehydration of Primary Alcohols: An E2 Reaction
7.8	285	Carbocation Stability and the Occurrence of Molecular Rearrangements
7.8A	285	Rearrangements During Dehydration of Secondary Alcohols
7.8B	287	Rearrangement after Dehydration of a Primary Alcohol
7.9	288	Synthesis of Alkynes by Elimination Reactions:
		Dehydrohalogenation of vic-Dibromides
7.10	290	The Acidity of Terminal Alkynes
7.11	290	Replacement of the Acetylenic Hydrogen Atom of Terminal Alkynes

7.12	292	Alkylation of Alkynide Anions: Some General Principles of Structure and Reactivity
		Illustrated
7.13	292	Hydrogenation of Alkenes
	293	The Chemistry of Hydrogenation in the Food Industry
7.14	294	Hydrogenation: The Function of the Catalyst
7.14A	295	Syn and Anti Additions
	295	The Chemistry of Homogeneous Asymmetric Catalytic Hydrogenation:
		Examples Involving L-DOPA, (S)-Naproxen, and Aspartame
7.15	297	Hydrogenation of Alkynes
7.15A	297	Syn Addition of Hydrogen: Synthesis of cis-Alkenes
7.15B	297	Anti Addition of Hydrogen: Synthesis of trans-Alkenes
7.16	298	An Introduction to organic Synthesis
7.16A	298	Why do Organic Synthesis?
7.16B	299	Retrosynthetic Analysis – Planning an Organic Synthesis
7.16C	300	Identifying Precursors
	302	The Chemistry of – From the Inorganic to the Organic
7.16D	302	Raison d'Etre
	303	Summary and Review Tools
	304	Summary of Methods for the Preparation of Alkenes and Alkynes;
		1. Dehydrohalogenation of Alkyl Halides (Section 7.6, p 275)
		2. Dehydration of Alcohols (Sections 7.7 & 7.8, p 280)
		3. Hydrogenation of Alkynes (Section 7.15, p 297)
		(4. Dehydrohalogenation of vic-Dihalides x 2 (Section 7.9, p 288))
	305	Summary and Review Tools
	306	Key Terms and Concepts
Proble	ms:	In-Chapter 7.1 to 7.17
	306	End of Chapter 7.18 to 7.46
	310	Learning Group Problems

Week of Nov 2: ALKENES AND ALKYNES II: Addition Reactions.

	311	The Sea: A Treasure of Biologically Active Natural Products
8.1	312	Introduction: Addition to Alkenes
8.1A	313	Understanding Additions to Alkenes
8.2	314	Electrophilic Addition of Hydrogen Halides to Alkenes:
		Mechanism and Markovnikov's Rule
8.2A	316	Theoretical Explanation of Markovnikov's Rule
8.2B	318	Modern Statement of Markovnokov's Rule
8.2C	319	Regioselective Reactions
8.2D	319	An Exception to Markovnikov's Rule
8.3	319	Stereochemistry of the Ionic Addition to an Alkene
8.4	320	Addition of Sulfuric Acid to Alkenes
8.4A	320	Alcohols from Alkyl Hydrogen Sulfates
8.5	321	Addition of Water to Alkenes: Acid Catalyzed Hydration
8.5A	321	Mechanism for Acid-Catalyzed Hydration
8.5B	322	Rearrangements

8.6	323	Alcohols from Alkenes through Oxymercuration-Demercuration: Markovnikov Addition
8.6A	323	Regioselectivity of Oxymercuration-Demercuration
8.6B	324	Rearrangements Seldom Occur in Oxymeercuration-Demercuration
8.6C	324	Mechanisms of Oxymercuration
8.7	326	Alcohols from Alkenes through Hydroboration-Oxidation:
		Anti-Markovnikov Syn Hydration
8.8	326	Hydroboration: Synthesis of Alkylboranes
8.8A	327	Mechanism of Hydroboration
8.8B	328	Stereochemistry of Hydroboration
8.9	329	Oxidation and Hydrolysis of Alkyl Boranes
8.9A	330	Regiochemistry and Stereochemistry of Alkyl Boranes:
		Oxidation and Hydrolysis
8.10	331	Summary of Alkene Hydration Methods
8.11	331	Proponolysis of Alkyl Boranes
8.12	332	Addition of Bromine and Chlorine to Alkenes
8.12A		Mechanism of Halogen Addition
8.13	334	Stereochemistry of the Addition of Halogens to Alkenes
8.13A		Stereospecific Reactions
8.14	337	Halohydrin Formation
8.15	338	Divalent Carbon Compounds: Carbenes
8.15A		Structure and Reactions of Methylene
8.15B		Reactions of Other Carbenes: Dihalocarbenes
8.15C		
		Carbenoids: The Simmons-Smith Cyclopropane Synthesis
8.16	340	Oxidation of Alkenes: Syn 1,2-Dihydroxylation
8.16A		Mechanisms for Syn Dihydroxylations of Alkenes
0.17	342	The Chemistry of Catalytic Asymmetric Dihydroxylations
8.17	343	Oxidative Cleavage of Alkenes
8.17A		Cleavage with Hot Basic Potassium Permanganate
8.17B		Cleavage with Ozone
8.18	345	Addition of Bromine and Chlorine to Alkynes
8.19	346	Addition of Hydrogen Halides to Alkynes
8.20	347	Oxidative Cleavage of Alkynes
8.21	347	Synthetic Strategies Revisited, including:
		1. Construction of the Carbon Skeleton
		2. Functional Group Interconversions
		3. Control of Regiochemistry and
		4. Control of Stereochemistry
8.21A	347	Retroactive Analysis
8.21B	348	Disconnections, Synthons, and Synthetic Equivalents
8.21C	349	Stereochemical Considerations
	350	The Chemistry of Cholesterol Biosynthesis: Elegant and Familiar Reactions in Nature
	354	Summary and Review Tools:
	354	Mechanism Review: Summary of Alkene Addition Reactions
	355	Synthetic Connections of Alkynes and Alkenes: II
	356	Key Terms and Concepts
Duckle.		In Chanter 9.1 to 9.26

In-Chapter 8.1 to 8.26 End of Chapter 8.27 to 8.68 Learning Group Problems. Problems: 356

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Week of Nov 9: RADICAL REACTIONS

	427	Radicals in Biology, Medicine, and Industry
10.1	428	Introduction
10.1A	428	Production of Radicals
10.1B	428	Reactions of Radicals
10.2	429	Homolytic Bond Dissociation Energies
10.2A	430	Homolytic Bond Dissociation Energies and Heats of Reaction
	430	Table 10.1 Single-Bond Homolytic Dissociation Energies DH° at 25° C
10.2B	431	Homolytic Bond Dissociation Energies and the Relative Stabilities of Radicals
10.3	433	The Reactions of Alkanes with Halogens
10.3A	433	Multiple Substitution Reactions versus Selectivity
10.4	435	Chlorination of Methane: Mechanism of Reaction
		1. Chain Initiation
		2. Chain Propagation
		3. Chain Termination
10.5	437	Chlorination of Methane: Energy Changes
10.5A	438	The Overall Free-Energy Change
10.5B	439	Activation Energies
10.5C	441	Reaction of Methane with other Halogens
10.6	443	Halogenation of Higher Alkanes
10.6A	445	Selectivity of Bromine, and Selectivity versus Reactivity
10.7	446	The Geometry of Alkyl Radicals
10.8	446	Reactions that Generate Tetrahedral Chirality Centres
10.8A	447	Generation of a Second Chirality Centre in a Radical Halogenation
10.9	449	Radical Addition to Alkenes:
		The Anti-Markovnikov Addition of Hydrogen Bromide
10.9A		Summary of Markovnikov versus Anti-Markovnikov Addition of HBr to Alkenes
10.10		Radical Polymerization of Alkenes: Chain Growth Polymers
	451	Radical Polymerization of Ethene
	453	Other Common Chain-Growth Polymers
10.11		Other Important Radical Reactions
10.11A		Molecular Oxygen and Super Oxide
10.11E		Nitric Oxide
10.110		Combustion of Alkanes
10.11E		Autoxidation
	458	The Chemistry of Antioxidants
	459	The Chemistry of Ozone Depletion and Chlorofluorocarbons (CFCs)
	460	Concept Map: Mechanism Review of Radical Reactions
	461	Key Terms and Concepts
	464	Special Topic A: Chain-Growth Polymers
Proble	ms:	In-Chapter 10.1 to 10.22
	461	End of Chapter 10.23 to 10.34
	463	Learning Group Problems.

Week of Nov 16: ALCOHOLS AND ETHERS.

	469	Molecular Hosts
11.1	470	Structure and Nomenclature
11.1A	471	Nomenclature of Alcohols
11.1B	472	Nomenclature of Ethers
11.2	472	Physical Properties of Alcohols and Ethers
	473	Tables 11.1 and 11.2: Physical Properties of Ethers and Alcohols
11.3	474	Important Alcohols and Ethers
11.3A-	D 474	Methanol, Ethylene Glycol, Diethyl Ether
11.4	476	Synthesis of Alcohols from Alkenes
	476	1. Acid-Catalyzed Hydration of Alkenes
	477	2. Oxymercuration-Demercuration
	477	3. Hydroboration-Oxidation
11.5	478	Reactions of Alcohols
11.6	479	Alcohols as Acids
11.7	480	Conversion of Alcohols into Alkyl Halides
11.8	480	Alkyl Halides from the Reactions of Alcohols with Hydrogen Halides
11.8A		Mechanisms of the Reactions of Alcohols with HX
11.9	483	Alkyl Halides from the Reactions of Alcohols with PBr ₃ or SOCl ₂
11.10	484	Tosylates, Mesylates and Triflates:
		Leaving Group Derivatives of Alcohols
	487	The Chemistry of Alkyl Phosphates
11.11		Synthesis of Ethers
11.11 <i>A</i>		Ethers by Intermolecular Dehydration of Alcohols
11.11E		The Williamson Synthesis of Ethers
11.11C		Synthesis of Ethers by Alkoxymercuration-Demercuration
11.11E		tert-Butyl Ethers by Alkylation of Alcohols: Protecting Groups
11.11E		Silyl Ether Protecting Groups
11.12		Reactions of Ethers: Ether Cleavage by Strong Acids
11.13		Epoxides
11.13A		Synthesis of Epoxides: Epoxidation of Alkenes
11.13E		Stereochemistry of Epoxidation
	494	The Chemistry of The Sharpless Asymmetric Epoxidation
11.14	496	Reactions of Epoxides:
		1. Acid Catalyzed Ring Opening
	400	2. Base Catalyzed Ring Opening
	498	The Chemistry of Epoxides, Carcinogens, and Biological Oxidation
11.14A		Polyethers from Epoxides
11.15	500	Anti 1,2-Dihydroxylation of Alkenes via Epoxides
1116	502	The Chemistry of Environmentally Friendly Alkene Oxidation Methods
11.16	503	Crown Ethers: Nucleophilic Substitution Reactions in Relatively Nonpolar Aprotic
11 1 / 4	504	Solvents by Phase-Transfer Catalysis
11.16A		Crown Ethers
11.16E		Transport Antibiotics and Crown Ethers
11.17	506	Summary of Reactions of Alkenes, Alcohols and Ethers

- 11.17A 506 Alkenes in Synthesis
 - 507 Key Terms and Concepts.
 - 508 Summary and Review Tool: Some Synthetic Connections of

Alkynes, Alcohols, Alkyl Halides and Ethers

Problems: In-Chapter 11.1 to 11.24

- 509 End of Chapter 11.25 to 11.51
- 512 Learning Group Problems.

Week of Nov 23: ALCOHOLS FROM CARBONYL COMPOUNDS: OXIDATION-REDUCTION AND ORGANOMETALLIC COMPOUNDS.

	AND ORGANOMETALLIC COMPOUNDS.
	Read and Study Chapter 12
513	The Two Aspects of the Coenzyme NADH
12.1 514	Introduction
12.1A 514	Structure of the Carbonyl Group
12.1B 515	Reactions of Carbonyl Compounds with Nucleophiles
12.2 515	Oxidation-Reduction Reactions in Organic Chemistry
12.2A 516	Oxidation States of Organic Chemistry
12.3 517	Alcohols by Reduction of Carbonyl Compounds
12.3A 517	Lithium Aluminum Hydride Reductions of Carbonyl Compounds
12.3B 518	Sodium Borohydride Reductions of Carbonyl Compounds
12.3C 519	Overall Summary of LiAlH ₄ and NaBH ₄ Reactivity
519	The Chemistry of Alcohol Dehydrogenase
520	The Chemistry of Stereoselective Reductions of Carbonyl Groups
12.4 521	Oxidation of Alcohols
12.4A 521	Oxidation of Primary Alcohols to Aldehydes: RCH ₂ OH to RCHO
12.4B 522	Oxidation of Primary Alcohols to Carboxylic Acids: RCH ₂ OH to RCO ₂ H
12.4C 522	Oxidation of Secondary Alcohols to Ketones: RCH(OH)R' to RCOR'
12.4D 523	Mechanism of Chromate Oxidations
12.4E 525	A Chemical Test for Primary and Secondary Alcohols
12.4F 525	Spectroscopic Evidence for Alcohols
12.5 526	Organometallic Compounds
12.6 526	Preparation of Organo Lithium and Organo Magnesium Compounds
12.6A 526	Organolithium Compounds
12.6B 527	Grignard Reagents
12.7 528	Reactions of Organolithium and Organomagnesium Compounds
12.7A 528	Reactions with Compounds Containing Acidic Hydrogen Atoms
12.7B 529	Reactions of Grignard Reagents with Oxiranes (Epoxides)
12.7C 530	Reactions of Grignard Reagents with Carbonyl Compounds
12.8 531	Alcohols from Grignard Reagents: Reaction of Grignard Reagents with:
	1. Formaldehyde to Give Primary Alcohols
	2. Other Aldehydes to Give Secondary Alcohols
	3. Ketones to Give Tertiary Alcohols
12.04 522	4. Esters with 2 x RMgX to Give Tertiary Alcohols
12.8A 532	Planning a Grignard Synthesis
12.8B 536	Restrictions on the Use of Grignard Reagents
12.8C 537	The Use of Lithium Reagents
12.8D 537	The Use of Sodium Alkynides
12.9 539	Protecting Groups

- 540 Summary of Reactions
- 541 Synthetic Connections of Alcohols and Carbonyl Compounds
- 541 Key Terms and Concepts

Problems: In-Chapter 12.1 to 12.10

- 542 End of Chapter 12.11 to 12.29
- 545 Learning Group Problems.
- 546 First Review Problem Set 1 to 25.

Week of Nov 30: CONJUGATED UNSATURATED SYSTEMS.

Read and Study Chapter 13.

550	Molecules With the Nobel Prize in Their Synthetic Lineage
13.1 551	Introduction
13.2 551	Allylic Substitution and the Allyl Radical
13.2A 552	Allylic Chlorination (High Temperature)
13.2B 554	Allylic Bromination with N-Bromosuccinimide (Low Conc. of Br ₂)
13.3 555	The Stability of the Allyl Radical
13.3A 555	Molecular Orbital Description of the Allyl Radical
13.3B 557	Resonance Description of the Allyl radical
13.4 558	The Allyl Carbocation
13.5 559	Summary of Rules for Resonance
13.5A 560	Rules for Writing Resonance Structures
13.5B 561	Estimating the Relative Stability of Resonance Structures
13.6 563	Alkadienes and Polyunsaturated Hydrocarbons
13.7 564	1,3-Butadiene: Electron Delocalization
13.7A 564	Bond Lengths of 1,3-Bitadiene
13.7B 565	Conformations of 1,3-Butadiene, s-cis and s-trans
13.7C 565	Molecular Orbitals of 1,3-Butadiene
13.8 566	The Stability of Conjugated Dienes
13.9 568	Ultraviolet-Visible Spectroscopy
13.9A 568	The Electromagnetic Spectrum
13.9B 569	UV-Vis Spectrophotometers
13.9C 571	Absorption Maxima for Nonconjugated and Conjugated Dienes
573	The Chemistry of The Photochemistry of Vision
13.9D 576	Analytical Uses of UV-Vis Spectroscopy
13.10 576	Electrophilic Attack on Conjugated Dienes: 1,4-Electrophilic Addition
13.10A 578	Kinetic Control versus Thermodynamic Control of a Chemical Reaction
13.11 580	The Diels-Alder Reaction: 1,4-Cycloaddition of Dienes
13.11A 581	Factors Favoring the Diels-Alder Reaction
13.11B 582	Stereochemistry of the Diels-Alder Reaction
13.11C 584	Molecular Orbital Considerations That Favor an Endo Transition State
586	The Chemistry of Asymmetric and Intramolecular Diels-Alder Reactions
588	Concept Map
589	Key Terms and Concepts.
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Problems: In-Chapter 13.1 to 13.15

End of Chapter 13.16 to 13.46 Learning Group Problems. 589

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December 7: Review Class, e.g. review of a Practice Final Exam.

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