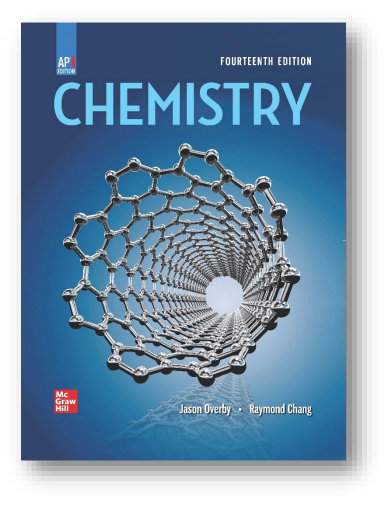
Advanced Placement[®] CORRELATION GUIDE



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| Topic 1.1 Moles and Mo | Topic 1.1 Moles and Molar Mass Enduring Understanding | | |
|--|--|--|--|
| Enduring Understandin | | | |
| SPQ-1 The mole allows | different units to be compared. | | |
| Learning Objective SPQ-1.A Calculate quantities of a substance or its relative number of particles using dimensional analysis and the mole concept. | Essential Knowledge SPQ-1.A.1 One cannot count particles directly while performing laboratory work. Thus, there must be a connection between the masses of substances reacting and the actual number of particles undergoing chemical changes. | Page numbers 80-84, 100-106; 155- 161; 196-199; 736-743 | |
| | SPQ-1.A.2 Avogadro's number ($N_A = 6.022 \times 10^{23}$ mol ⁻¹) provides the connection between the number of moles in a pure sample of a substance and the number of constituent particles (or formula units) of that substance. | 82-85, 100-106; 196- 199 | |
| | SPQ-1.A.3 Expressing the mass of an individual atom or molecule in atomic mass units (amu) is useful because the average mass in amu of one particle (atom or molecule) or formula unit of a substance will always be numerically equal to the molar mass of that substance in grams. Thus, there is a quantitative connection between the mass of a substance and the number of particles that the substance contains. EQN: n = m/M | 80-85 | |

Unit 1: Atomic Structure and Properties

| Topic 1.2 Mass Spectro | scopy of Elements | |
|------------------------|--|------------------|
| Enduring Understandin | g | |
| SPQ-1 The mole allows | different units to be compared. | |
| Learning Objective | Essential Knowledge | Page numbers |
| SPQ-1.B Explain the | SPQ-1.B.1 The mass spectrum of a sample | 80-81, 88-89 |
| quantitative | containing a single element can be used to | |
| relationship between | determine the identity of the isotopes of that | |
| the mass spectrum of | element and the relative abundance of each | |
| an element and the | isotope in nature. | |
| masses of the | SPQ-1.B.2 The average atomic mass of an element | 81, 88-89; AP123 |
| element's isotopes. | can be estimated from the weighted average of the | |
| | isotopic masses using the mass of each isotope and | |
| | its relative abundance. | |

| Topic 1.3 Elemental Co | omposition of Pure Substances | |
|------------------------|---|----------------|
| Enduring Understandi | ng | |
| SPQ-2 Chemical formu | las identify substances by their unique combination of a | atoms. |
| Learning Objective | Essential Knowledge | Page numbers |
| SPQ-2.A Explain the | SPQ-2.A.1 Some pure substances are composed of | 54-60; 494-500 |
| quantitative | individual molecules, while others consist of atoms | |
| relationship between | or ions held together in fixed proportions as | |
| the elemental | described by a formula unit. | |
| composition by mass | SPQ-2.A.2 According to the law of definite | 42 |
| and the empirical | proportions, the ratio of the masses of the | |
| formula of a pure | constituent elements in any pure sample of that | |
| substance. | compound is always the same. | |
| | SPQ-2.A.3 The chemical formula that lists the lowest | 57-58; 93-94 |
| | whole number ratio of atoms of the elements in a | |
| | compound is the empirical formula. | |

| Topic 1.4 Composition | Topic 1.4 Composition of Mixtures | | |
|-----------------------|--|---------------------|--|
| Enduring Understandi | Enduring Understanding | | |
| SPQ-2 Chemical formu | las identify substances by their unique combination of atc | oms. | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SPQ-2.B Explain the | SPQ-2.B.1 While pure substances contain molecules or | 8-10; 150-154; 199- | |
| quantitative | formula units of a single type, mixtures contain | 205; 936-937 | |
| relationship between | molecules or formula units of two or more types, | | |
| the elemental | whose relative proportions can vary | | |
| composition by mass | SPQ-2.B.2 Elemental analysis can be used to | 90-92; 155-157 | |
| and the composition | determine the relative numbers of atoms in a | | |
| of substances in a | substance and to determine its purity | | |
| mixture. | | | |

| Topic 1.5 Atomic Struc | ture and Electron Configuration | | |
|------------------------|---|------------------------------|--|
| Enduring Understandi | Enduring Understanding | | |
| SAP-1 Atoms and mole | cules can be identified by their electron distribution and e | energy. | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SAP-1.A Represent | SAP-1.A.1 The atom is composed of negatively | 43-49; 283-290 | |
| the electron | charged electrons and a positively charged nucleus | | |
| configuration of an | that is made of protons and neutrons. | | |
| element or ions of an | SAP-1.A.2 Coulomb's law is used to calculate the force | 336-350, 375-380 | |
| element using the | between two charged particles. | | |
| Aufbau principle. | SAP-1.A.3 In atoms and ions, the electrons can be thought of as being in "shells (energy levels)" and "subshells (sublevels)," as described by the electron configuration. Inner electrons are called core electrons, and outer electrons are called valence electrons. The electron configuration is explained by quantum mechanics, as delineated in the Aufbau principle and exemplified in the periodic table of the elements. | 290-317; 333-336; 371-372 | |
| | SAP-1.A.4 The relative energy required to remove an electron from different subshells of an atom or ion or from the same subshell in different atoms or ions (ionization energy) can be estimated through a qualitative application of Coulomb's law. This energy is related to the distance from the nucleus and the effective (shield) charge of the nucleus. | 343-347 | |

| Topic 1.6 Photoelectro | n Spectroscopy | | |
|--|---|---|--|
| Enduring Understandi | Enduring Understanding | | |
| SAP-1 Atoms and mole | cules can be identified by their electron distribution and e | energy. | |
| Learning Objective | Essential Knowledge | Page number | |
| SAP-1.B Explain the relationship between the photoelectron spectrum of an atom or ion and: a. The electron configuration of the species. b. The interactions | SAP-1.B.1 The energies of the electrons in a given shell can be measured experimentally with photoelectron spectroscopy (PES). The position of each peak in the PES spectrum is related to the energy required to remove an electron from the corresponding subshell, and the height of each peak is (ideally) proportional to the number of electrons in that subshell. | 281-283 *see additional online activity | |
| between the electrons and the nucleus. | | | |

| Topic 1.7 Periodic Trends | | |
|---------------------------|--|------------------|
| Enduring Understanding | | |
| SAP-2 The periodic tab | le shows patterns in electronic structure and trends in at | omic properties. |
| Learning Objective | Essential Knowledge | Page number |
| SAP-2.A Explain the | SAP-2.A.1 The organization of the periodic table is | 51-52; 329-336 |
| relationship between | based on the recurring properties of the elements | |
| trends in atomic | and explained by the pattern of electron | |
| properties of | configurations and the presence of completely or | |
| elements and | partially filled shells (and subshells) of electrons in | |
| electronic structure | atoms. | |
| and periodicity. | SAP-2.A.2 Trends in atomic properties within the | 336-350 |
| | periodic table (periodicity) can be qualitatively | |
| | understood through the position of the element in | |
| | the periodic table, Coulomb's law, the shell model, | |
| | and the concept of shielding/effective nuclear charge. | |
| | These properties include: | |
| | a. Ionization energy | |
| | b. Atomic and ionic radii | |
| | c. Electron affinity | |
| | d. Electronegativity. | |
| | SAP-2.A.3 The periodicity (in SAP-2.A.2) is useful to | 336-350 |
| | predict/estimate values of properties in the absence | |
| | of data. | |

| Topic 1.8 Valence Elec | trons and Ionic Compounds | |
|-------------------------------|---|----------------------|
| Enduring Understandi | ng | |
| SAP-2 The periodic tab | le shows patterns in electronic structure and trends in a | tomic properties. |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-2.B Explain the | SAP-2.B.1 The likelihood that two elements will form | 54-60; 333, 350-361 |
| relationship between | a chemical bond is determined by the interactions | |
| trends in the | between the valence electrons and nuclei of | |
| reactivity of | elements. | |
| elements and | SAP-2.B.2 Elements in the same column of the | 51-52; 947-952; 989- |
| periodicity. | periodic table tend to form analogous compounds. | 994 |
| | SAP-2.B.3 Typical charges of atoms in ionic | 54-55; 334-336; 372- |
| | compounds are governed by their location on the | 374; 947-952; 989- |
| | periodic table and the number of valence electrons. | 994 |

| Topic 2.1 Types of Chemical Bonds | | |
|-----------------------------------|---|----------------------|
| Enduring Understandin | | |
| SAP-3 Atoms or ions bo | nd due to interactions between them, forming molecule | es. |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-3.A Explain the | SAP-3.A.1 Electronegativity values for the | 383-385 |
| relationship between | representative elements increase going from left to | |
| the type of bonding | right across a period and decrease going down a | |
| and the properties of | group. These trends can be understood qualitatively | |
| the elements | through the electronic structure of the atoms, the | |
| participating in the | shell model, and Coulomb's law. | |
| bond. | SAP-3.A.2 Valence electrons shared between atoms | 54-55; 380-382, 385- |
| | of similar electronegativity constitute a nonpolar | 386; 429-434 |
| | covalent bond. For example, bonds between carbon | |
| | and hydrogen are effectively nonpolar even though | |
| | carbon is slightly more electronegative than | |
| | hydrogen. | |
| | SAP-3.A.3 Valence electrons shared between atoms | 54-55; 383-387; 429- |
| | of unequal electronegativity constitute a polar | 434 |
| | covalent bond. | |
| | a. The atom with a higher electronegativity will | |
| | develop a partial negative charge relative to the | |
| | other atom in the bond. | |
| | b. In single bonds, greater differences in | |
| | electronegativity lead to greater bond dipoles. | |
| | c. All polar bonds have some ionic character, and the | |
| | difference between ionic and covalent bonding is | |
| | not distinct but rather a continuum. | |
| | SAP-3.A.4 The difference in electronegativity is not | 343-347; 383-387 |
| | the only factor in determining if a bond should be | |
| | designated as ionic or covalent. Generally, bonds | |
| | between a metal and nonmetal are ionic, and bonds | |
| | between two nonmetals are covalent. Examination | |
| | of the properties of a compound is the best way to | |
| | characterize the type of bonding. | |
| | SAP-3.A.5 In a metallic solid, the valence electrons | 497-499; 943-944 |
| | from the metal atoms are considered to be | |
| | delocalized and not associated with any individual | |
| | atom. | |

Unit 2: Molecular and Ionic Compound Structure and Properties

| Topic 2.2 Intramolecula | Topic 2.2 Intramolecular Force and Potential Energy | | |
|-------------------------|---|-------------------|--|
| Enduring Understanding | | | |
| SAP-3 Atoms or ions bo | SAP-3 Atoms or ions bond due to interactions between them, forming molecules. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SAP-3.B Represent the | SAP-3.B.1 A graph of potential energy versus the | 383; 434-436 | |
| relationship between | distance between atoms is a useful representation | | |
| potential energy and | for describing the interactions between atoms. Such | | |
| distance between | graphs illustrate both the equilibrium bond length | | |
| atoms, based on | (the separation between atoms at which the | | |
| factors that influence | potential energy is lowest) and the bond energy (the | | |
| the interaction | energy required to separate the atoms). | | |
| strength. | SAP-3.B.2 In a covalent bond, the bond length is | 382-383, 403-408 | |
| | influenced by both the size of the atom's core and | | |
| | the bond order (i.e., single, double, triple). Bonds | | |
| | with a higher order are shorter and have larger bond | | |
| | energies. | | |
| | SAP-3.B.3 Coulomb's law can be used to understand | 259-261; 339-340; | |
| | the strength of interactions between cations and | 375-382 | |
| | anions. | | |
| | a. Because the interaction strength is proportional | | |
| | to the charge on each ion, larger charges lead to | | |
| | stronger interactions. | | |
| | b. Because the interaction strength increases as the | | |
| | distance between the centers of the ions (nuclei) | | |
| | decreases, smaller ions lead to stronger interactions. | | |

| Topic 2.3 Structure of Id | onic Solids | |
|---------------------------|---|--------------|
| Enduring Understandin | g | |
| SAP-3 Atoms or ions bo | nd due to interactions between them, forming molecule | es. |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-3.C Represent an | SAP-3.C.1 The cations and anions in an ionic crystal | 494-496 |
| ionic solid with a | are arranged in a systematic, periodic 3-D array that | |
| particulate model that | maximizes the attractive forces among cations and | |
| is consistent with | anions while minimizing the repulsive forces. | |
| Coulomb's law and | | |
| the properties of the | | |
| constituent ions. | | |

| Topic 2.4 Structure of Metals and Alloys | | |
|--|---|------------------|
| Enduring Understanding | | |
| SAP-3 Atoms or ions bo | nd due to interactions between them, forming molecule | es. |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-3.D Represent a | SAP-3.D.1 Metallic bonding can be represented as | 497-498; 943-944 |
| metallic solid and/or | an array of positive metal ions surrounded by | |
| alloy using a model to | delocalized valence electrons (i.e., a "sea of | |
| show essential | electrons"). | |
| characteristics of the | SAP-3.D.2 Interstitial alloys form between atoms of | 938-941 |
| structure and | different radii, where the smaller atoms fill the | |
| interactions present in | interstitial spaces between the larger atoms (e.g., | |
| the substance. | with steel in which carbon occupies the interstices in | |
| | iron). | |
| | SAP-3.D.3Substitutional alloys form between atoms | 936 |
| | of comparable radius, where one atom substitutes | |
| | for the other in the lattice. (In certain brass alloys, | |
| | other elements, usually zinc, substitute for copper.) | |

| Topic 2.5 Lewis Diagrams | | | |
|---------------------------|--|--------------|--|
| Enduring Understandin | g | | |
| SAP-4 Molecular compo | SAP-4 Molecular compounds are arranged based on Lewis diagrams and Valence Shell Electron Pair | | |
| Repulsion (VSEPR) theory. | | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SAP-4.A Represent a | SAP-4.A.1 Lewis diagrams can be constructed | 388-394 | |
| molecule with a Lewis | according to an established set of principles. | | |
| diagram. | | | |

| Topic 2.6 Resonance and Formal Charge | | | |
|---------------------------------------|--|--------------|--|
| Enduring Understanding | | | |
| SAP-4 Molecular compo | SAP-4 Molecular compounds are arranged based on Lewis diagrams and Valence Shell Electron Pair | | |
| Repulsion (VSEPR) theo | ry. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SAP-4.B Represent a | SAP-4.B.1 In cases where more than one equivalent | 393-394 | |
| molecule with a Lewis | Lewis structure can be constructed, resonance must | | |
| diagram that accounts | be included as a refinement to the Lewis structure. | | |
| for resonance | In many such cases, this refinement is needed to | | |
| between equivalent | provide qualitatively accurate predictions of | | |
| structures or that uses | molecular structure and properties. | | |
| formal charge to | SAP-4.B.2 The octet rule and formal charge can be | 391-394 | |
| select between | used as criteria for determining which of several | | |
| nonequivalent | possible valid Lewis diagrams provides the best | | |
| structures. | model for predicting molecular structure and | | |
| | properties. | | |
| | SAP-4.B.3 As with any model, there are limitations | 397-398 | |
| | to the use of the Lewis structure model, particularly | | |
| | in cases with an odd number of valence electrons. | | |

| Topic 2.7 VSEPR and Bo | nd Hybridization | | |
|------------------------|---|-----------------------|--|
| Enduring Understandin | g | | |
| SAP-4 Molecular compo | ounds are arranged based on Lewis diagrams and Valence | e Shell Electron Pair | |
| Repulsion (VSEPR) theo | Repulsion (VSEPR) theory. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SAP-4.C Based on the | SAP-4.C.1 VSEPR theory uses the Coulombic | 419 | |
| relationship between | repulsion between electrons as a basis for predicting | | |
| Lewis diagrams, | the arrangement of electron pairs around a central | | |
| VSEPR theory, bond | atom. | | |
| orders, and bond | SAP-4.C.2 Both Lewis diagrams and VSEPR theory | 419-428 | |
| polarities: | must be used for predicting electronic and structural | | |
| a. Explain structural | properties of many covalently bonded molecules | | |
| properties of | and polyatomic ions, including the following: | | |
| molecules. | a. Molecular geometry | | |
| b. Explain electron | b. Bond angles | | |
| properties of | c. Relative bond energies based on bond order | | |
| molecules. | d. Relative bond lengths (multiple bonds, effects of | | |
| | atomic radius) | | |
| | e. Presence of a dipole moment | | |
| | f. Hybridization of valence orbitals of the molecule | | |
| | SAP-4.C.3 The terms "hybridization" and "hybrid | 437-445 | |
| | atomic orbital" are used to describe the | | |
| | arrangement of electrons around a central atom. | | |
| | When the central atom is <i>sp</i> hybridized, its ideal | | |
| | bond angles are 180°; for <i>sp</i> ² hybridized atoms the | | |
| | bond angles are 120°; and for <i>sp</i> ³ hybridized atoms | | |
| | the bond angles are 109.5°. | | |
| | SAP-4.C.4 Bond formation is associated with overlap | 449-457 | |
| | between atomic orbitals. In multiple bonds, such | | |
| | overlap leads to the formation of both sigma and pi | | |
| | bonds. The overlap is stronger in sigma than pi | | |
| | bonds, which is reflected in sigma bonds having | | |
| | greater bond energy than pi bonds. The presence of | | |
| | a pi bond also prevents the rotation of the bond and | | |
| | leads to structural isomers. | | |

| Topic 3.1 Intermolecula | r Forces | |
|-------------------------|---|--------------------|
| Enduring Understandin | | |
| SAP-5 Intermolecular fo | rces can explain the physical properties of a material. | |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-5.A Explain the | SAP-5.A.1 London dispersion forces are a result of | 473-476; 1064-1069 |
| relationship between | the Coulombic interactions between temporary, | |
| the chemical | fluctuating dipoles. London dispersion forces are | |
| structures of | often the strongest net intermolecular force | |
| molecules and the | between large molecules. | |
| relative strength of | a. Dispersion forces increase with increasing contact | |
| their intermolecular | area between molecules and with increasing | |
| forces when: | polarizability of the molecules. | |
| a. The molecules are | b. The polarizability of a molecule increases with an | |
| of the same chemical | increasing number of electrons in the molecule; and | |
| species. | the size of the electron cloud. It is enhanced by the | |
| b. The molecules are | presence of pi bonding. | |
| of two different | c. The term "London dispersion forces" should not | |
| chemical species. | be used synonymously with the term "van der Waals | |
| | forces." | |
| | SAP-5.A.2 The dipole moment of a polar molecule | 429-430; 473-476; |
| | leads to additional interactions with other chemical | 530; 1042 |
| | species. | |
| | a. Dipole-induced dipole interactions are present | |
| | between a polar and nonpolar molecule. These | |
| | forces are always attractive. The strength of these | |
| | forces increases with the magnitude of the dipole of | |
| | the polar molecule and with the polarizability of the | |
| | nonpolar molecule. | |
| | b. Dipole-dipole interactions are present between | |
| | polar molecules. The interaction strength depends | |
| | on the magnitudes of the dipoles and their relative | |
| | orientation. Interactions between polar molecules | |
| | are typically greater than those between nonpolar | |
| | molecules of comparable size because these | |
| | interactions act in addition to London dispersion | |
| | forces. | |
| | c. Ion-dipole forces of attraction are present | |
| | between ions and polar molecules. These tend to be | |
| | stronger than dipole-dipole forces. | |

Unit 3: Intermolecular Forces and Properties

| Topic 3.1 Intermolecular Forces (continued) | | |
|---|--|--------------|
| Enduring Understanding | | |
| SAP-5 Intermolecular for | prces can explain the physical properties of a material. | |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-5.A Explain the | SAP-5.A.3 The relative strength and orientation | 473-476 |
| relationship between | dependence of dipole-dipole and ion-dipole forces | |
| the chemical | can be understood qualitatively by considering the | |
| structures of | sign of the partial charges responsible for the | |
| molecules and the | molecular dipole moment, and how these partial | |
| relative strength of | charges interact with an ion or with an adjacent | |
| their intermolecular | dipole. | |
| forces when: | SAP-5.A.4 Hydrogen bonding is a strong type of | 473-476 |
| a. The molecules are | intermolecular interaction that exists when | |
| of the same chemical | hydrogen atoms covalently bonded to the highly | |
| species. | electronegative atoms (N, O, and F) are attracted to | |
| b. The molecules are | the negative end of a dipole formed by the | |
| of two different | electronegative atom (N, O, and F) in a different | |
| chemical species. | molecule, or a different part of the same molecule. | |
| | SAP-5.A.5 In large biomolecules, noncovalent | 1072-1080 |
| | interactions may occur between different molecules | |
| | or between different regions of the same | |
| | large biomolecule. | |

| Topic 3.2 Properties of Solids | | |
|--------------------------------|--|--------------------|
| Enduring Understandin | g | |
| SAP-5 Intermolecular for | prces can explain the physical properties of a material. | |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-5.B Explain the | SAP-5.B.1 Many properties of liquids and solids are | 502-509 |
| relationship among | determined by the strengths and types of | |
| the macroscopic | intermolecular forces present. Because | |
| properties of a | intermolecular interactions are broken when a | |
| substance, the | substance vaporizes, the vapor pressure and boiling | |
| particulate-level | point are directly related to the strength of those | |
| structure of the | interactions. Melting points also tend to correlate | |
| substance, and the | with interaction strength, but because the | |
| interactions between | interactions are only rearranged, in melting, the | |
| these particles. | relations can be more subtle. | |
| | SAP-5.A.2 Particulate-level representations, showing | 126-128; 472-473, |
| | multiple interacting chemical species, are a useful | 477-478, 483; 529; |
| | means to communicate or understand how | 683; 1073-1074, |
| | intermolecular interactions help to establish | 1080 |
| | macroscopic properties. | |

| Topic 3.2 Properties of Solids (continued) | | |
|--|--|--------------------|
| Enduring Understanding | | |
| SAP-5 Intermolecular forces can explain the physical properties of a material. | | |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-5.B Explain the | SAP-5.B.3 Due to strong interactions between ions, | 125-127; 176-177; |
| relationship among | ionic solids tend to have low vapor pressures, high | 375-380, 382; 494- |
| the macroscopic | melting points, and high boiling points. They tend to | 499 |
| properties of a | be brittle due to the repulsion of like charges caused | |
| substance, the | when one layer slides across another layer. They | |
| particulate-level | conduct electricity only when the ions are mobile, as | |
| structure of the | when the ionic solid is melted or dissolved in water | |
| substance, and the | or another solvent. | |
| interactions between | SAP-5.B.4 In covalent network solids, the atoms are | 496-497 |
| these particles. | covalently bonded together into a three-dimensional | |
| | network (e.g., diamond) or layers of two- | |
| | dimensional networks (e.g., graphite). These are | |
| | only formed from nonmetals: elemental (e.g., | |
| | diamond, graphite) or binary compounds of two | |
| | nonmetals (e.g., silicon dioxide and silicon carbide). | |
| | Due to the strong covalent interactions, covalent | |
| | solids have high melting points. Three-dimensional | |
| | network solids are also rigid and hard, because the | |
| | covalent bond angles are fixed. However, graphite is | |
| | soft because adjacent layers can slide past each | |
| | other relatively easily | |
| | SAP-5.B.5 Molecular solids are composed of distinct, | 176-177; 380-383; |
| | individual units of covalently-bonded molecules | 496-497; 1063-1068 |
| | attracted to each other through relatively weak | |
| | intermolecular forces. Molecular solids generally | |
| | have a low melting point because of the relatively | |
| | weak intermolecular forces present between the | |
| | molecules. They do not conduct electricity because | |
| | their valence electrons are tightly held within the | |
| | covalent bonds and the lone pairs of each | |
| | constituent molecule. Molecular solids are | |
| | sometimes composed of very large molecules or | |
| | polymers. | |

| Topic 3.2 Properties of | Topic 3.2 Properties of Solids (continued) | | |
|--------------------------|--|--------------------|--|
| Enduring Understanding | | | |
| SAP-5 Intermolecular for | prces can explain the physical properties of a material. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SAP-5.B Explain the | SAP-5.B.6 Metallic solids are good conductors of | 497-498; 936, 938- | |
| relationship among | electricity and heat, due to the presence of free | 944 | |
| the macroscopic | valence electrons. They also tend to be malleable | | |
| properties of a | and ductile, due to the ease with which the metal | | |
| substance, the | cores can rearrange their structure. In an interstitial | | |
| particulate-level | alloy, interstitial atoms tend to make the lattice | | |
| structure of the | more rigid, decreasing malleability and ductility. | | |
| substance, and the | Alloys typically retain a sea of mobile electrons and | | |
| interactions between | so remain conducting. | | |
| these particles. | SAP-5.B.7 In large biomolecules or polymers, | 1054-1055; 1072- | |
| | noncovalent interactions may occur between | 1075, 1078-1080 | |
| | different molecules or between different regions of | | |
| | the same large biomolecule. The functionality and | | |
| | properties of such molecules depend strongly on the | | |
| | shape of the molecule, which is largely dictated by | | |
| | noncovalent interactions. | | |

| Topic 3.3 Solids, Liquids | s, and Gases | |
|---------------------------|--|----------------------|
| Enduring Understandin | g | |
| SAP-6 Matter exists in t | hree states: solid, liquid, and gas, and their differences a | are influenced by |
| variances in spacing and | l motion of the molecules. | |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-6.A Represent | SAP-6.A.1 Solids can be crystalline, where the | 11-13; 485-491, 494- |
| the differences | particles are arranged in a regular three-dimensional | 498 |
| between solid, liquid, | structure, or they can be amorphous, where the | |
| and gas phases using a | particles do not have a regular, orderly | |
| particulate-level | arrangement. In both cases, the motion of the | |
| model. | individual particles is limited, and the particles do | |
| | not undergo overall translation with respect to each | |
| | other. The structure of the solid is influenced by | |
| | interparticle interactions and the ability of the | |
| | particles to pack together. | |
| | SAP-6.A.2 The constituent particles in liquids are in | 11-13; 479-484 |
| | close contact with each other, and they are | |
| | continually moving and colliding. The arrangement | |
| | and movement of particles are influenced by the | |
| | nature and strength of the forces (e.g., polarity, | |
| | hydrogen bonding, and temperature) between the | |
| | particles. | |

Topic 3.3 Solids, Liquids, and Gases (continued)

Enduring Understanding

SAP-6 Matter exists in three states: solid, liquid, and gas, and their differences are influenced by variances in spacing and motion of the molecules.

| Learning Objective | Essential Knowledge | Page numbers |
|------------------------|--|---------------------|
| SAP-6.A Represent | SAP-6.A.3 The solid and liquid phases for a particular | 11-13; 508-511 |
| the differences | substance typically have similar molar volume | |
| between solid, liquid, | because, in both phases, the constituent particles | |
| and gas phases using a | are in close contact at all times. | |
| particulate-level | SAP-6.A.4 In the gas phase, the particles are in | 11-13; 178, 206-207 |
| model. | constant motion. Their frequencies of collision and | |
| | the average spacing between them are dependent | |
| | on temperature, pressure, and volume. Because of | |
| | this constant motion, and minimal effects of forces | |
| | between particles, a gas has neither a definite | |
| | volume nor a definite shape. | |

| Topic 3.4 Ideal Gas Law | Topic 3.4 Ideal Gas Law | | |
|-------------------------|--|---------------------|--|
| Enduring Understandin | g | | |
| SAP-7 Gas properties ar | e explained macroscopically—using the relationships ar | nong pressure, | |
| volume, temperature, n | noles, gas constant—and molecularly by the motion of t | he gas. | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SAP-7.A Explain the | SAP-7.A.1 The macroscopic properties of ideal gases | 188-199, 214-216; | |
| relationship between | are related through the ideal gas law: | 634-635, 654-655 | |
| the macroscopic | EQN: PV = nRT. | | |
| properties of a sample | SAP-7.A.2 In a sample containing a mixture of ideal | 199-205; 654-655 | |
| of gas or mixture of | gases, the pressure exerted by each component (the | | |
| gases using the ideal | partial pressure) is independent of the other | | |
| gas law. | components. Therefore, the total pressure of the | | |
| | sample is the sum of the partial pressures. | | |
| | EQN: $P_A = P_{total} \times X_A^3$ | | |
| | where XA = moles A/total moles; | | |
| | EQN: $P_{total} = P_A + P_B + P_C +$ | | |
| | SAP-7.A.3 Graphical representations of the | 184, 185, 204, 208, | |
| | relationships between P, V, T, and n are useful to | 214; 797 | |
| | describe gas behavior. | | |

Topic 3.5 Kinetic Molecular Theory

| Topic 3.5 Kinetic Molec | | |
|--|---|------------------|
| Enduring Understanding | | |
| SAP-7 Gas properties are explained macroscopically—using the relationships among pressure, | | |
| volume, temperature, n | noles, gas constant—and molecularly by the motion of t | he gas. |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-7.B Explain the | SAP-7.B.1 The kinetic molecular theory (KMT) | 205-216; 593-596 |
| relationship between | relates the macroscopic properties of gases to | |
| the motion of | motions of the particles in the gas. The Maxwell- | |
| particles and the | Boltzmann distribution describes the distribution of | |
| macroscopic | the kinetic energies of particles at a given | |
| properties of gases | temperature. | |
| with: | SAP-7.B.2 All the particles in a sample of matter are | 205-206 |
| a. The kinetic | in continuous, random motion. The average kinetic | |
| molecular theory | energy of a particle is related to its average velocity | |
| (KMT). | by the equation: | |
| b. A particulate | EQN: KE = $\frac{1}{2}$ mv ² | |
| model. | SAP-7.B.3 The Kelvin temperature of a sample of | 184-186; 593-596 |
| c. A graphical | matter is proportional to the average kinetic energy | |
| representation. | of the particles in the sample. | |
| | SAP-7.B.4 The Maxwell-Boltzmann distribution | 207-209 |
| | provides a graphical representation of the energies/ | |
| | velocities of particles at a given temperature. | |

| Topic 3.6 Deviation from | m Ideal Gas Law | |
|--------------------------|---|------------------|
| Enduring Understandin | g | |
| SAP-7 Gas properties ar | e explained macroscopically—using the relationships a | mong pressure, |
| volume, temperature, n | noles, gas constant—and molecularly by the motion of | the gas. |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-7.C Explain the | SAP-7.C.1 The ideal gas law does not explain the | 188-189; 214-216 |
| relationship among | actual behavior of real gases. Deviations from the | |
| non-ideal behaviors of | ideal gas law may result from interparticle | |
| gases, interparticle | attractions among gas molecules, particularly at | |
| forces, and/or | conditions that are close to those resulting in | |
| volumes. | condensation. Deviations may also arise from | |
| | particle volumes, particularly at extremely high | |
| | pressures. | |

| Topic 3.7 Solutions and | Topic 3.7 Solutions and Mixtures | | |
|-------------------------|---|------------------------|--|
| Enduring Understandin | g | | |
| SPQ-3 Interactions betw | veen intermolecular forces influence the solubility and s | eparation of mixtures. | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SPQ-3.A Calculate the | SPQ-3.A.1 Solutions, also sometimes called | 8-9; 125; 199-201, | |
| number of solute | homogeneous mixtures, can be solids, liquids, or | 202-203; 528-531, | |
| particles, volume, or | gases. In a solution, the macroscopic properties do | 554-557; 753-758 | |
| molarity of solutions. | not vary throughout the sample. In a heterogeneous | | |
| | mixture, the macroscopic properties depend on | | |
| | location in the mixture. | | |
| | Essential Knowledge | 150-154, 157-162; | |
| | SPQ-3.A.2 Solution composition can be expressed in | 532-536 | |
| | a variety of ways; molarity is the most common | | |
| | method used in the laboratory. | | |
| | EQN: $M = n_{solute} / L_{solution}$ | | |

| Topic 3.8 Representations of Solutions | | | |
|--|---|-------------------------|--|
| Enduring Understandin | Enduring Understanding | | |
| SPQ-3 Interactions betw | veen intermolecular forces influence the solubility and | separation of mixtures. | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SPQ-3.B Using | SPQ-3.B.1 Particulate representations of solutions | 126-128, 139-140, | |
| particulate models for | communicate the structure and properties of | 153; 199; 529, 539, | |
| mixtures: | solutions, by illustration of the relative | 553, 556-557; 683, | |
| a. Represent | concentrations of the components in the solution | 701 | |
| interactions between | and drawings that show interactions among the | | |
| components. | components. | | |
| b. Represent | | | |
| concentrations of | | | |
| components. | | | |

| Topic 3.9 Separation of Solutions and Mixtures Chromatography | | | |
|---|---|------------------------|--|
| Enduring Understanding | Enduring Understanding | | |
| SPQ-3 Interactions betw | veen intermolecular forces influence the solubility and s | eparation of mixtures. | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SPQ-3.C Explain the | SPQ-3.C.1 The components of a liquid solution | 359; 429-433 | |
| relationship between | cannot be separated by filtration. They can, | *See additional | |
| the solubility of ionic | however, be separated using processes that take | online activity; | |
| and molecular | advantage of differences in the intermolecular | 544; 1054-1056 | |
| compounds in | interactions of the components. | | |
| aqueous and | a. Chromatography (paper, thin-layer, and column) | | |
| nonaqueous solvents, | separates chemical species by taking advantage of | | |
| and the | the differential strength of intermolecular | | |
| intermolecular | interactions between and among the components of | | |
| interactions between | the solution (the mobile phase) and with the surface | | |
| particles. | components of the stationary phase. | | |
| | b. Distillation separates chemical species by taking | | |
| | advantage of the differential strength of | | |
| | intermolecular interactions between and among the | | |
| | components and the effects these interactions have | | |
| | on the vapor pressures of the components in the | | |
| | mixture. | | |

| Topic 3.10 Solubility | Topic 3.10 Solubility | | |
|-------------------------|---|-------------------------|--|
| Enduring Understandin | g | | |
| SPQ-3 Interactions betw | veen intermolecular forces influence the solubility and | separation of mixtures. | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SPQ-3.C Explain the | SPQ-3.C.2 Substances with similar intermolecular | 128; 528-531,, 537- | |
| relationship between | interactions tend to be miscible or soluble in one | 541; 747-748, 755, | |
| the solubility of ionic | another | 759-764 | |
| and molecular | | | |
| compounds in | | | |
| aqueous and | | | |
| nonaqueous solvents, | | | |
| and the | | | |
| intermolecular | | | |
| interactions between | | | |
| particles. | | | |

Topic 3.11 Spectroscopy and the Electromagnetic Spectrum

Enduring Understanding

SAP-8 Spectroscopy can determine the structure and concentration in a mixture of a chemical species.

| Learning Objective | Essential Knowledge | Page numbers |
|------------------------|---|------------------|
| SAP-8.A Explain the | SAP-8.A.1 Differences in absorption or emission of | 275-290; 917-918 |
| relationship between | photons in different spectral regions are related to | |
| a region of the | the different types of molecular motion or electronic | |
| electromagnetic | transition: | |
| spectrum and the | a. Microwave radiation is associated with transitions | |
| types of molecular or | in molecular rotational levels. | |
| electronic transitions | b. Infrared radiation is associated with transitions in | |
| associated with that | molecular vibrational levels. | |
| region. | c. Ultraviolet/visible radiation is associated with | |
| | transitions in electronic energy levels. | |
| | | |

| Topic 3.12 Photoelectri | Topic 3.12 Photoelectric Effect | | |
|--|---|---------------------|--|
| Enduring Understandin | Enduring Understanding | | |
| SAP-8 Spectroscopy car species. | n determine the structure and concentration in a mixtur | e of a chemical | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SAP-8.B Explain the | SAP-8.B.1 When a photon is absorbed (or emitted) | 274, 275-290; 1015- | |
| properties of an | by an atom or molecule, the energy of the species is | 1016 | |
| absorbed or emitted | increased (or decreased) by an amount equal to the | | |
| photon in relationship | energy of the photon. | | |
| to an electronic | SAP-8.B.2 The wavelength of the electromagnetic | 274, 278-280; 1015- | |
| transition in an atom | wave is related to its frequency and the speed of | 1017 | |
| or molecule. | light by the equation: | | |
| | EQN: $c = \lambda v$. | | |
| | The energy of a photon is related to the frequency | | |
| | of the electromagnetic wave through Planck's | | |
| | equation ($E = hv$). | | |

Topic 3.13 Beer-Lambert Law

| Enduring Understandin | g | |
|--|---|---------------------|
| SAP-8 Spectroscopy ca | n determine the structure and concentration in a mixtur | re of a chemical |
| species. | | |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-8.C Explain the amount of light absorbed by a solution of molecules or ions in relationship to the concentration, path length, and molar absorptivity. | SAP-8.C.1 The Beer-Lambert law relates the absorption of light by a solution to three variables according to the equation: EQN: A = ε bc. The molar absorptivity ε describes how intensely a sample of molecules or ions absorbs light of a specific wavelength. The path length b and concentration c are proportional to the number of absorbing species. | 149; 570; 1015-1017 |
| molar absorptivity. | SAP-8.C.2 In most experiments the path length and wavelength of light are held constant. In such cases, the absorbance is proportional only to the concentration of absorbing molecules or ions. | 532-536; 1015-1017 |

Unit 4: Chemical Reactions

Topic 4.1 Introduction for Reactions

| Topic 4.1 Indioduction for Reactions | | | |
|---|--|---|--|
| Enduring Understandin | Enduring Understanding | | |
| TRA-1 A substance that | TRA-1 A substance that changes its properties, or that changes into a different substance, can be | | |
| represented by chemica | al equations. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-1.A Identify evidence of chemical and physical changes | TRA-1.A.1 A physical change occurs when a substance undergoes a change in properties but not a change in composition. Changes in the phase | 13-14; 472, 502-516 | |
| in matter | of a substance (solid, liquid, gas) or formation/ separation of mixtures of substances are common physical changes. | | |
| | TRA-1.A.2 A chemical change occurs when substances are transformed into new substances, typically with different compositions. Production of heat or light, formation of a gas, formation of a precipitate, and/or color change provide possible evidence that a chemical change has occurred. | 13-14; 95; 127, 132, 136-138 139, 144-148, 155-157; 232-234 | |

| Topic 4.2 Net Ionic Equations | | | |
|-------------------------------|--|-----------------------|--|
| Enduring Understanding | | | |
| TRA-1 A substance that | TRA-1 A substance that changes its properties, or that changes into a different substance, can be | | |
| represented by chemica | l equations. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-1.B Represent | TRA-1.B.1 All physical and chemical processes can | 95-100; 129-132; 816- | |
| changes in matter | be represented symbolically by balanced equations. | 818 | |
| with a balanced | TRA-1.B.2 Chemical equations represent chemical | 95-100; 129-132; 372- | |
| chemical or net ionic | changes. These changes are the result of a | 373, 380-381; 816-818 | |
| equation: | rearrangement of atoms into new combinations; | | |
| a. For physical | thus, any representation of a chemical change must | | |
| changes. | contain equal numbers of atoms of every element | | |
| b. For given | before and after the change occurred. Equations | | |
| information about the | thus demonstrate that mass is conserved in | | |
| identity of the | chemical reactions. | | |
| reactants and/or | TRA-1.B.3 Balanced molecular, complete ionic, and | 95-100; 129-132; 816- | |
| product. | net ionic equations are differing symbolic forms | 818 | |
| c. For ions in a given | used to represent a chemical reaction. The form | | |
| chemical reaction. | used to represent the reaction depends on the | | |
| | context in which it is to be used. | | |

| Topic 4.3 Representations of Reactions | | |
|--|--|-------------------------|
| Enduring Understanding | | |
| TRA-1 A substance that changes its properties, or that changes into a different substance, can be | | |
| represented by chemica | al equations. | |
| Learning Objective | Essential Knowledge | Page numbers |
| TRA-1.C Represent a | TRA-1.C.1 Balanced chemical equations in their | 3, 14; 41, 42; 96, 100, |
| given chemical | various forms can be translated into symbolic | 107, 116, 117, 118, |
| reaction or physical | particulate representations. | 120; 126, 135; 187; |
| process with a | | 243, 246, 261, 267; |
| consistent | | 378, 398, 402, 406, |
| particulate model. | | 407, 414; 529; 569, |
| | | 572, 576, 578, 582, |
| | | 584, 586, 588, 595, |
| | | 596, 597, 601, 603, |
| | | 604, 609, 613, 618, |
| | | 619, 620, 621, 622, |
| | | 623; 631, 637, 638, |
| | | 639, 646, 647, 651; |
| | | 654, 658, 660, 662, |
| | | 667, 668; 675, 683; |
| | | 769, 770; 783, 789, |
| | | 791, 808, 811, 813; |
| | | 820; 877; 1072, 1077 |

| Topic 4.4 Physical and Chemical Changes | | |
|---|---|--------------------|
| Enduring Understanding | | |
| TRA-1 A substance that changes its properties, or that changes into a different substance, can be | | |
| represented by chemica | l equations. | |
| Learning Objective | Essential Knowledge | 13-14; 95-97; 127- |
| TRA-1.D Explain the | TRA-1.D.1 Processes that involve the breaking | 130; 253-254; 372- |
| relationship between | and/or formation of chemical bonds are typically | 373, 380-382; 472, |
| macroscopic | classified as chemical processes. Processes that | 502-516 |
| characteristics and | involve only changes in intermolecular interactions, | |
| bond interactions for: | such as phase changes, are typically classified as | |
| a. Chemical processes. | physical processes. | |
| b. Physical processes. | TRA-1.D.2 Sometimes physical processes involve | 125-127; 259- 262; |
| | the breaking of chemical bonds. For example, | 375-380; 529-531 |
| | plausible arguments could be made for the | |
| | dissolution of a salt in water, as either a physical or | |
| | chemical process, involves breaking of ionic bonds, | |
| | and the formation of ion-dipole interactions | |
| | between ions and solvent. | |

| Topic 4.5 Stoichiometry | | |
|--|--|-----------------------|
| Enduring Understanding | | |
| SPQ-4 When a substance changes into a new substance, or when its properties change, no mass is | | |
| lost or gained. | | |
| Learning Objective | Essential Knowledge | Page numbers |
| SPQ-4.A Explain | SPQ-4.A.1 Because atoms must be conserved | 100-112; 150-163 |
| changes in the | during a chemical process, it is possible to calculate | |
| amounts of reactants | product amounts by using known reactant | |
| and products based | amounts, or to calculate reactant amounts given | |
| on the balanced | known product amounts. | |
| reaction equation for | SPQ-4.A.2 Coefficients of balanced chemical | 95-112; 129-132; 849- |
| a chemical process. | equations contain information regarding the | 851 |
| | proportionality of the amounts of substances | |
| | involved in the reaction. These values can be used | |
| | in chemical calculations involving the mole concept. | |
| | SPQ-4.A.3 Stoichiometric calculations can be | 150-163; 196-199 |
| | combined with the ideal gas law and calculations | |
| | involving molarity to quantitatively study gases and | |
| | solutions. | |

| Topic 4.6 Introduction to Titration | | |
|-------------------------------------|---|----------------------|
| Enduring Understanding | | |
| SPQ-4 When a substance | e changes into a new substance, or when its propertie | s change, no mass is |
| lost or gained. | | |
| Learning Objective | Essential Knowledge | Page numbers |
| SPQ-4.B Identify the | SPQ-4.B.1 Titrations may be used to determine the | 157-163; 736-743 |
| equivalence point in a | concentration of an analyte in solution. The titrant | |
| titration based on the | has a known concentration of a species that reacts | |
| amounts of the titrant | specifically and quantitatively with the analyte. The | |
| and analyte, assuming | equivalence point of the titration occurs when the | |
| the titration reaction | analyte is totally consumed by the reacting species | |
| goes to completion. | in the titrant. The equivalence point is often | |
| | indicated by a change in a property (such as color) | |
| | that occurs when the equivalence point is reached. | |
| | This observable event is called the endpoint of the | |
| | titration. | |

Topic 4.7 Types of Chemical Reactions

Enduring Understanding

TRA-2 When a substance changes into a new substance, or when its properties change, no mass is lost or gained.

| Learning Objective | Essential Knowledge | Page number |
|------------------------|--|---------------------|
| | • | - |
| TRA-2.A Identify a | TRA-2.A.1 Acid-base reactions involve transfer of | 132-138; 673-676; |
| reaction as acid-base, | one or more protons between chemical species. | 736-743 |
| oxidation-reduction, | TRA-2.A.2 Oxidation-reduction reactions involve | 138-150, 816-819 |
| or precipitation. | transfer of one or more electrons between chemical | |
| | species, as indicated by changes in oxidation | |
| | numbers of the involved species. Combustion is an | |
| | important subclass of oxidation-reduction reactions, | |
| | in which a species reacts with oxygen gas. In the | |
| | case of hydrocarbons, carbon dioxide and water are | |
| | products of complete combustion. | |
| | TRA-2.A.3 In a redox reaction, electrons are | 138-141; 822-827 |
| | transferred from the species that is oxidized to the | |
| | species that is reduced. | |
| | TRA-2.A.4 Oxidation numbers may be assigned to | 138-141; 822-827 |
| | each of the atoms in the reactants and products; this | |
| | is often an effective way to identify the oxidized and | |
| | reduced species in a redox reaction. | |
| | TRA-2.A.5 Precipitation reactions frequently involve | 127-132, 155-157; |
| | mixing ions in aqueous solution to produce an | 51528-531; 746-748, |
| | insoluble or sparingly soluble ionic compound. All | 764-767 |
| | sodium, potassium, ammonium, and nitrate salts are | |
| | soluble in water. | |

Topic 4.8 Introduction to Acid-Base Reactions

Enduring Understanding

TRA-2 When a substance changes into a new substance, or when its properties change, no mass is lost or gained.

| Learning Objective | Essential Knowledge | Page numbers |
|------------------------|---|-------------------|
| TRA-2.B Identify | TRA-2.B.1 By definition, a Brønsted-Lowry acid is a | 133-136; 673-674 |
| species as Brønsted- | proton donor and a Brønsted-Lowry base is a | |
| Lowry acids, bases, | proton acceptor. | |
| and/or conjugate | TRA-2.B.2 Only in aqueous solutions, water plays | 133-136; 673-676 |
| acid-base pairs, based | an important role in many acid-base reactions, | |
| on proton-transfer | as its molecular structure allows it to accept | |
| involving those | protons from and donate protons to | |
| species. | dissolved species. | |
| | TRA-2.B.3 When an acid or base ionizes in water, | 133-136; 673-676, |
| | the conjugate acid-base pairs can be identified and | 682-686 |
| | their relative strengths compared. | |

| Topic 4.9 Oxidation-Reduction (Redox) Reactions | | | |
|---|--|-------------------|--|
| Enduring Understanding | | | |
| TRA-2 When a substance changes into a new substance, or when its properties change, no mass is | | | |
| lost or gained. | | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-2.C Represent a | TRA-2.C.1 Balanced chemical equations for redox | 138-150; 816-819, | |
| balanced redox | reactions can be constructed from half-reactions | 846-849 | |
| reaction equation | | | |
| using half-reactions. | | | |

Unit 5: Kinetics

| Topic 5.1 Reaction Rates | | |
|--------------------------|--|--------------------|
| Enduring Understanding | | |
| TRA-3 Some reactions h | appen quickly, while others happen more slowly and d | lepend on reactant |
| concentrations and tem | perature. | |
| Learning Objective | Essential Knowledge | Page numbers |
| TRA-3.A Explain the | TRA-3.A.1 The kinetics of a chemical reaction is | 568-571 |
| relationship between | defined as the rate at which an amount of reactants | |
| the rate of a chemical | is converted to products per unit of time. | |
| reaction and | TRA-3.A.2 The rates of change of reactant and | 573-575 |
| experimental | product concentrations are determined by | |
| parameters. | the stoichiometry in the balanced | |
| | chemical equation. | |
| | TRA-3.A.3 The rate of a reaction is influenced by | 570-600, 607-614 |
| | reactant concentrations, temperature, surface | |
| | area, catalysts, and other environmental factors. | |

| Topic 5.2 Introduction to Rate Law | | | |
|------------------------------------|---|------------------|--|
| Enduring Understanding | | | |
| TRA-3 Some reactions h | TRA-3 Some reactions happen quickly, while others happen more slowly and depend on reactant | | |
| concentrations and tem | iperature. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-3.B Represent | TRA-3.B.1 Experimental methods can be used to | 569-573, 576-580 | |
| experimental data | monitor the amounts of reactants and/or products | | |
| with a consistent rate | of a reaction and to determine the rate of | | |
| law expression. | the reaction. | | |
| | TRA-3.B.2 The rate law expresses the rate of a | 576-592 | |
| | reaction as proportional to the concentration of | | |
| | each reactant raised to a power. | | |
| | TRA-3.B.3 The power of each reactant in the rate | 579-592 | |
| | law is the order of the reaction with respect to that | | |
| | reactant. The sum of the powers of the reactant | | |
| | concentrations in the rate law is the overall order | | |
| | of the reaction. | | |
| | TRA-3.B.4 The proportionality constant in the rate | 572, 580-592 | |
| | law is called the rate constant. The value of this | | |
| | constant is temperature dependent and the units | | |
| | reflect the overall reaction order. | | |
| | TRA-3.B.5 Comparing initial rates of a reaction is a | 570-575 | |
| | method to determine the order with respect to | | |
| | each reactant. | | |

| Topic 5.3 Concentration | Topic 5.3 Concentration Changes Over Time | | |
|---|--|----------------------|--|
| Enduring Understanding | | | |
| TRA-3 Some reactions happen quickly, while others happen more slowly and depend on reactant | | | |
| concentrations and tem | perature. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-3.C Identify the | TRA-3.C.1 The order of a reaction can be inferred | 568-580, 592 | |
| rate law expression of | from a graph of concentration of reactant versus | | |
| a chemical reaction | time. | | |
| using data that show | TRA-3.C.2 If a reaction is first order with respect to | 580-592 <i>,</i> 579 | |
| how the | a reactant being monitored, a plot of the natural | | |
| concentrations of | log (In) of the reactant concentration as a function | | |
| reaction species | of time will be linear. | | |
| change over time. | TRA-3.C.3 If a reaction is second order with respect | 588-592 | |
| | to a reactant being monitored, a plot of the | | |
| | reciprocal of the concentration of that reactant | | |
| | versus time will be linear. | | |
| | TRA-3.C.4 The slopes of the concentration versus | 580-592 | |
| | time data for zeroth, first, and second order | | |
| | reactions can be used to determine the rate | | |
| | constant for the reaction. | | |
| | Zeroth order: EQN: $[A]_t - [A]_0 = -kt$ | | |
| | First order: EQN: $ln[A]_t - ln[A]_0 = -kt$ | | |
| | Second order: EQN: $1/[A]_t - 1/[A]_0 = kt$ | | |
| | TRA-3.C.5 Half-life is a critical parameter for first | 580-592; 874-877 | |
| | order reactions because the half-life is constant and | | |
| | related to the rate constant for the reaction by the | | |
| | equation: | | |
| | EQN: $t_{1/2} = 0.693/k$. | | |
| | TRA-3.C.6 Radioactive decay processes provide an | 874-877 | |
| | important illustration of first order kinetics | | |

| Topic 5.4 Elementary R | Topic 5.4 Elementary Reactions Enduring Understanding TRA-4 There is a relationship between the speed of a reaction and the collision frequency of particle | | |
|------------------------------|---|--------------|--|
| Enduring Understandin | | | |
| TRA-4 There is a relatio | | | |
| collisions. | | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-4.A Represent an | TRA-4.A.1 The rate law of an elementary reaction | 600-607 | |
| elementary reaction | can be inferred from the stoichiometry of the | | |
| as a rate law | molecules participating in a collision. | | |
| expression using | TRA-4.A.2 Elementary reactions involving the | 600-601 | |
| stoichiometry. | simultaneous collision of three or more particles | | |
| | are rare. | | |

Topic 5.5 Collision Model

| Enduring Understandin | lg | |
|--------------------------|---|-------------------------|
| TRA-4 There is a relatio | nship between the speed of a reaction and the collisior | n frequency of particle |
| collisions. | | |
| Learning Objective | Essential Knowledge | Page numbers |
| TRA-4.B Explain the | TRA-4.B.1 For an elementary reaction to | 592-594, 600-607 |
| relationship between | successfully produce products, reactants must | |
| the rate of an | successfully collide to initiate bond-breaking and | |
| elementary reaction | bond-making events. | |
| and the frequency, | TRA-4.B.2 In most reactions, only a small fraction of | 592-607 |
| energy, | the collisions leads to a reaction. Successful | |
| and orientation of | collisions have both sufficient energy to overcome | |
| molecular collisions. | energy barriers and orientations that allow the | |
| | bonds to rearrange in the required manner. | |
| | TRA-4.B.3 The Maxwell-Boltzmann distribution | 207-209; 592-594, |
| | curve describes the distribution of particle | 607-614 |
| | energies; this distribution can be used to gain a | |
| | qualitative estimate of the fraction of collisions | |
| | with sufficient energy to lead to a reaction, and also | |
| | how that fraction depends on temperature. | |

| Topic 5.6 Reaction Energy Profile | | | |
|---|--|--------------|--|
| Enduring Understandin | g | | |
| TRA-4 There is a relationship between the speed of a reaction and the collision frequency of particle | | | |
| collisions. | - | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-4.C Represent | TRA-4.C.1 Elementary reactions typically involve | 600 | |
| the activation energy | the breaking of some bonds and the forming of | | |
| and overall energy | new ones. | | |
| change in an | TRA-4.C.2 The reaction coordinate is the axis along | 568-572 | |
| elementary reaction | which the complex set of motions involved in | | |
| using a reaction | rearranging reactants to form products can be | | |
| energy profile. | plotted. | | |
| | TRA-4.C.3 The energy profile gives the energy along | 592-600 | |
| | the reaction coordinate, which typically proceeds | | |
| | from reactants, through a transition state, to | | |
| | products. The energy difference between the | | |
| | reactants and the transition state is the activation | | |
| | energy for the forward reaction. | | |
| | TRA-4.C.4 The Arrhenius equation relates the | 592-600 | |
| | temperature dependence of the rate of an | | |
| | elementary reaction to the activation energy | | |
| | needed by molecular collisions to reach the | | |
| | transition state. | | |

| Topic 5.7 Introduction | to Reaction Mechanisms | |
|------------------------|--|---------------------|
| Enduring Understandir | ng | |
| TRA-5 Many chemical r | eactions occur through a series of elementary reaction | s. These elementary |
| reactions when combin | ed form a chemical equation. | |
| Learning Objective | Essential Knowledge | Page numbers |
| TRA-5.A Identify the | TRA-5.A.1 A reaction mechanism consists of a | 600-614 |
| components of a | series of elementary reactions, or steps, that occur | |
| reaction mechanism. | in sequence. The components may include | |
| | reactants, intermediates, products, and catalysts. | |
| | TRA-5.A.2 The elementary steps when combined | 600 |
| | should align with the overall balanced equation of a | |
| | chemical reaction. | |
| | TRA-5.A.3 A reaction intermediate is produced by | 600-601 |
| | some elementary steps and consumed by others, | |
| | such that it is present only while a reaction is | |
| | occurring. | |
| | TRA-5.A.4 Experimental detection of a reaction | 600-607 |
| | intermediate is a common way to build evidence in | |
| | support of one reaction mechanism over an | |
| | alternative mechanism. | |

| Topic 5.8 Reaction Mec | hanism and Rate Law | |
|-------------------------|--|---------------------|
| Enduring Understandin | g | |
| TRA-5 Many chemical re | eactions occur through a series of elementary reaction | s. These elementary |
| reactions when combine | ed form a chemical equation. | |
| Learning Objective | Essential Knowledge | Page numbers |
| TRA-5.B Identify the | TRA-5.B.1 For reaction mechanisms in which each | 600-607 |
| rate law for a reaction | elementary step is irreversible, or in which the first | |
| from a mechanism in | step is rate limiting, the rate law of the reaction is | |
| which the first step is | set by the molecularity of the slowest elementary | |
| rate limiting. | step (i.e., the rate-limiting step). | |

| Topic 5.9 Steady-State | Topic 5.9 Steady-State Approximation | | |
|---|---|---------------------|--|
| Enduring Understandin | g | | |
| TRA-5 Many chemical re | eactions occur through a series of elementary reactions | s. These elementary | |
| reactions when combined form a chemical equation. | | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-5.C Identify the | TRA-5.C.1 If the first elementary reaction is not rate | 600-607 | |
| rate law for a reaction | limiting, approximations (such as steady state) | | |
| from a mechanism | must be made to determine a rate law expression. | | |
| in which the first step | | | |
| is not rate limiting. | | | |

| Topic 5.10 Multistep Reaction Energy Profile | | |
|--|--|-----------------------|
| Enduring Understandin | g | |
| TRA-5 Many chemical re | eactions occur through a series of elementary reactions | s. These elementary |
| reactions when combine | ed form a chemical equation. | |
| Learning Objective | Essential Knowledge | Page numbers |
| TRA-5.D Represent the activation energy and overall energy change in a multistep reaction with a reaction energy profile. | TRA-5.D.1 Knowledge of the energetics of each elementary reaction in a mechanism allows for the construction of an energy profile for a multistep reaction. | 592-593, 601-605, 608 |

| Topic 5.11 Catalysis | | |
|--------------------------|---|---------------------|
| Enduring Understanding | g | |
| ENE-1 Many chemical re | eactions occur through a series of elementary reactions | s. These elementary |
| reactions when combine | ed form a chemical equation. | |
| Learning Objective | Essential Knowledge | Page numbers |
| ENE-1.A Explain the | ENE-1.A.1 In order for a catalyst to increase the | 607-614; 656 |
| relationship between | rate of a reaction, the addition of the catalyst must | |
| the effect of a catalyst | increase the number of effective collisions and/ or | |
| on a reaction | provide a reaction path with a lower activation | |
| and changes in the | energy relative to the original reaction coordinate. | |
| reaction mechanism. | ENE-1.A.2 In a reaction mechanism containing a | 607-608 |
| | catalyst, the net concentration of the catalyst is | |
| | constant. However, the catalyst will frequently be | |
| | consumed in the rate-determining step of the | |
| | reaction, only to be regenerated in a subsequent | |
| | step in the mechanism. | |
| | ENE-1.A.3 Some catalysts accelerate a reaction by | 607-614 |
| | binding to the reactant(s). The reactants are either | |
| | oriented more favorably or react with lower | |
| | activation energy. There is often a new reaction | |
| | intermediate in which the catalyst is bound to the | |
| | reactant(s). Many enzymes function in this manner. | |
| | ENE-1.A.4 Some catalysts involve covalent bonding | 607-614 |
| | between the catalyst and the reactant(s). An | |
| | example is acid-base catalysis, in which a reactant | |
| | or intermediate either gains or loses a proton. This | |
| | introduces a new reaction intermediate and new | |
| | elementary reactions involving that intermediate. | |
| | ENE-1.A.5 In surface catalysis, a reactant or | 607-610 |
| | intermediate binds to, or forms a covalent bond | |
| | with, the surface. This introduces elementary | |
| | reactions involving these new bound | |
| | reaction intermediate(s). | |

Unit 6: Thermodynamics

| Topic 6.1 Endothermic | and Exothermic Processes | | |
|-------------------------|---|-----------------------|--|
| Enduring Understanding | | | |
| ENE-2 Changes in a subs | ENE-2 Changes in a substance's properties or change into a different substance requires an exchange | | |
| of energy. | | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| ENE-2.A Explain the | ENE-2.A.1 Temperature changes in a system | 11; 232-234, 236-239, | |
| relationship between | indicate energy changes. | 259-262 | |
| experimental | ENE-2.A.2 Energy changes in a system can be | 11; 232-234 | |
| observations and | described as endothermic and exothermic | | |
| energy changes | processes such as the heating or cooling of a | | |
| associated with a | substance, phase changes, or chemical | | |
| chemical or physical | transformations. | | |
| transformation. | ENE-2.A.3 When a chemical reaction occurs, the | 11; 232-239, 246-249 | |
| | energy of the system either decreases (exothermic | | |
| | reaction), increases (endothermic reaction), or | | |
| | remains the same. For exothermic reactions, the | | |
| | energy lost by the reacting species (system) is | | |
| | gained by the surroundings, as heat transfer from | | |
| | or work done by the system. Likewise, for | | |
| | endothermic reactions, the system gains energy | | |
| | from the surroundings by heat transfer to or work | | |
| | done on the system. | | |
| | ENE-2.A.4 The formation of a solution may be an | 259-262 | |
| | exothermic or endothermic process, depending on | | |
| | the relative strengths of | | |
| | intermolecular/interparticle interactions before | | |
| | and after the dissolution process. | | |

| Topic 6.2 Energy Diagrams | | |
|---|---|---------------|
| Enduring Understanding | ng | |
| ENE-2 Changes in a substance's properties or change into a different substance requires an exchange | | |
| of energy. | | |
| Learning Objective | Essential Knowledge | Page numbers |
| ENE-2.B Represent a | ENE-2.B.1 A physical or chemical process can be | 233, 242, 259 |
| chemical or physical | described with an energy diagram that shows the | |
| transformation with | endothermic or exothermic nature of that process. | |
| an energy diagram. | | |

Topic 6.3 Heat Transfer and Thermal Equilibrium

Enduring Understanding

ENE-2 Changes in a substance's properties or change into a different substance requires an exchange of energy.

| Learning Objective | Essential Knowledge | Page numbers |
|-----------------------|---|-------------------|
| ENE-2.C Explain the | ENE-2.C.1 A physical or chemical process can be | 203-206; 231-232 |
| relationship between | described with an energy diagram that shows the | |
| the transfer of | endothermic or exothermic nature of that process. | |
| thermal energy and | ENE-2.C.2 Collisions between particles in thermal | 203-206; 232-234, |
| molecular collisions. | contact can result in the transfer of energy. This | 238-243 |
| | process is called "heat transfer," "heat exchange," | |
| | or "transfer of energy as heat." | |
| | ENE-2.C.3 Eventually, thermal equilibrium is | 231-232, 235-236 |
| | reached as the particles continue to collide. At | |
| | thermal equilibrium, the average kinetic energy of | |
| | both bodies is the same, and hence, their | |
| | temperatures are the same. | |

| Topic 6.4 Heat Capacity | and Calorimetry | |
|--------------------------|---|------------------------|
| Enduring Understanding | - | |
| ENE-2 Changes in a subst | tance's properties or change into a different substanc | e requires an exchange |
| of energy. | | |
| Learning Objective | Essential Knowledge | Page numbers |
| ENE-2.D Calculate the | ENE-2.D.1 The heating of a cool body by a warmer | 247-249 |
| heat q absorbed or | body is an important form of energy transfer | |
| released by a system | between two systems. The amount of heat | |
| undergoing heating/ | transferred between two bodies may be | |
| cooling based on the | quantified by the heat transfer equation: | |
| amount of the | EQN: q = mc∆T. | |
| substance, the heat | Calorimetry experiments are used to measure the | |
| capacity, and the | transfer of heat. | |
| change in temperature. | ENE-2.D.2 The first law of thermodynamics states | 235 |
| | that energy is conserved in chemical and | |
| | physical processes. | |
| | ENE-2.D.3 The transfer of a given amount of | 247 |
| | thermal energy will not produce the same | |
| | temperature change in equal masses of matter | |
| | with differing specific heat capacities. | |
| | ENE-2.D.4 Heating a system increases the energy | 235-242 |
| | of the system, while cooling a system decreases | |
| | the energy of the system. | |
| | ENE-2.D.5 The specific heat capacity of a | 247-250 |
| | substance and the molar heat capacity are both | |
| | used in energy calculations. | |
| | ENE-2.D.6 Chemical systems change their energy | 231-234, 240-244; |
| | through three main processes: heating/cooling, | 502-511 |
| | phase transitions, and chemical reactions. | |

Topic 6.5 Energy of Phase Changes

Enduring Understanding

ENE-2 Changes in a substance's properties or change into a different substance requires an exchange of energy.

| Learning Objective | Essential Knowledge | Page numbers |
|-------------------------|--|--------------|
| ENE-2.E Explain | ENE-2.E.1 Energy must be transferred to a system | 502-511 |
| changes in the heat q | to cause a substance to melt (or boil). The energy | |
| absorbed or released | of the system therefore increases as the system | |
| by a system | undergoes a solid-to-liquid (or liquid-to-gas) phase | |
| undergoing a phase | transition. Likewise, a system releases energy | |
| transition based on the | when it freezes (or condenses). The energy of the | |
| amount of the | system decreases as the system undergoes a | |
| substance in moles and | liquid-to-solid (or gas-to-liquid) phase transition. | |
| the molar enthalpy of | The temperature of a pure substance remains | |
| the phase transition. | constant during a phase change. | |
| | ENE-2.E.2 The energy absorbed during a phase | 502-511 |
| | change is equal to the energy released during a | |
| | complementary phase change in the opposite | |
| | direction. For example, the molar heat of | |
| | condensation of a substance is equal to the | |
| | negative of its molar heat of vaporization. | |

| Topic 6.6 Introduction to | Topic 6.6 Introduction to Enthalpy of Reaction | | |
|---------------------------|--|------------------------|--|
| Enduring Understanding | | | |
| ENE-2 Changes in a subst | ance's properties or change into a different substanc | e requires an exchange | |
| of energy. | | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| ENE-2.F Calculate the | ENE-2.F.1 The enthalpy change of a reaction gives | 242-243 | |
| heat q absorbed or | the amount of heat energy released (for negative | | |
| released by a system | values) or absorbed (for positive values) by a | | |
| undergoing a chemical | chemical reaction at constant pressure. | | |
| reaction in relationship | | | |
| to the amount of the | | | |
| reacting substance in | | | |
| moles and the molar | | | |
| enthalpy of reaction. | | | |

| Topic 6.7 Bond Enthalpies | | |
|---|--|------------------|
| Enduring Understanding | | |
| ENE-3 The energy exchar | nged in a chemical transformation is required to break | and form bonds. |
| Learning Objective | Essential Knowledge | Page numbers |
| ENE-3.A Calculate the enthalpy change of a reaction based on the | ENE-3.A.1 During a chemical reaction, bonds are broken and/or formed, and these events change the potential energy of the system. | 231-232; 403-408 |
| average bond energies of bonds broken and formed in the reaction. | ENE-3.A.2 The average energy required to break all of the bonds in the reactant molecules can be estimated by adding up the average bond energies of all the bonds in the reactant molecules. Likewise, the average energy released in forming the bonds in the product molecules can be estimated. If the energy released is greater than the energy required, the reaction is exothermic. If the energy required is greater than the energy released, the reaction is endothermic. | 358-361; 403-408 |

| Topic 6.8 Enthalpy of Formation | | | |
|---|---|--------------|--|
| Enduring Understanding | | | |
| ENE-3 The energy exchanged in a chemical transformation is required to break and form bonds. | | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| ENE-3.B Calculate the enthalpy change for a chemical or physical process based on the standard enthalpies of formation. | ENE-3.B.1 Tables of standard enthalpies of formation can be used to calculate the standard enthalpies of reactions. EQN: $\Delta H^{\circ}_{reaction} = \Sigma \Delta H^{\circ}_{f products} - \Sigma \Delta H^{\circ}_{f reactants}$ | 253-260 | |

| Topic 6.9 Hess's Law | | | | |
|--|---|------------------|--|--|
| Enduring Understandin | g | | | |
| ENE-3 The energy exchanged in a chemical transformation is required to break and form bonds. | | | | |
| Learning Objective | Essential Knowledge | Page numbers | | |
| ENE-3.C Represent a | ENE-3.C.1 Although the concept of "state function" | 253-260 | | |
| chemical or physical | is not required for the course, two principles of | | | |
| process as a sequence | Hess's law should be understood. First, when a | | | |
| of steps. | reaction is reversed, the enthalpy change stays | | | |
| | constant in magnitude but becomes reversed in | | | |
| | mathematical sign. Second, when two (or more) | | | |
| | reactions are added to obtain an overall reaction, | | | |
| | the individual enthalpy changes of each reaction | | | |
| | are added to obtain the net enthalpy of the overall | | | |
| | reaction. | | | |
| ENE-3.D Explain the | ENE-3.D.1 When the products of a reaction are at a | 231-234; 655-656 | | |
| relationship between | different temperature than their surroundings, they | | | |
| the enthalpy of a | exchange energy with the surroundings to reach | | | |
| chemical or physical | thermal equilibrium. Thermal energy is transferred | | | |
| process and the sum | to the surroundings from the products of an | | | |
| of the enthalpies of | exothermic reaction. Thermal energy is transferred | | | |
| the individual steps. | from the surroundings to the products of an | | | |
| | endothermic reaction. | | | |

Unit 7: Equilibrium

| Topic 7.1 | Introduction | to | Equilibrium |
|-----------|--------------|----|-------------|
|-----------|--------------|----|-------------|

| Enduring | Understanding |
|----------|---------------|
|----------|---------------|

TRA-6 Some reactions can occur in both forward and reverse directions, sometimes proceeding in each direction simultaneously

| Learning Objective | Essential Knowledge | Page numbers |
|------------------------|---|---------------------|
| TRA-6.A Explain the | TRA-6.A.1 Many observable processes are | 125-129, 133-134, |
| relationship between | reversible. Examples include evaporation and | 136-138; 208-210; |
| the occurrence of a | condensation of water, absorption and desorption | 502-508; 538-541; |
| reversible chemical or | of a gas, or dissolution and precipitation of a salt. | 673-674; 746-753, |
| physical process, and | Some important reversible chemical processes | 753-754; 816-8819 |
| the establishment of | include the transfer of protons in acid-base | |
| equilibrium, to | reactions and the transfer of electrons in redox | |
| experimental | reactions. | |
| observations. | TRA-6.A.2 When equilibrium is reached, no | 630-633, 647-648 |
| | observable changes occur in the system. Reactants | |
| | and products are simultaneously present, and the | |
| | concentrations or partial pressures of all species | |
| | remain constant. | |
| | TRA-6.A.3 The equilibrium state is dynamic. The | 503; 630-633 |
| | forward and reverse processes continue to occur at | |
| | equal rates, resulting in no net observable change. | |
| | TRA-6.A.4 Graphs of concentration, partial | 569, 571, 572, 573; |
| | pressure, or rate of reaction versus time for simple | 631, 653, 659 |
| | chemical reactions can be used to understand the | |
| | establishment of chemical equilibrium. | |

Topic 7.2 Direction of Reversible Reactions

TRA-6 Some reactions can occur in both forward and reverse directions, sometimes proceeding in each direction simultaneously

| Learning Objective | Essential Knowledge | Page numbers |
|------------------------|--|------------------|
| TRA-6.B Explain the | TRA-6.B.1 If the rate of the forward reaction is | 630-633, 644-646 |
| relationship between | greater than the reverse reaction, then there is a | |
| the direction in which | net conversion of reactants to products. If the rate | |
| a reversible reaction | of the reverse reaction is greater than that of the | |
| proceeds and the | forward reaction, then there is a net conversion of | |
| relative rates of the | products to reactants. An equilibrium state is | |
| forward and reverse | reached when these rates are equal. | |
| reactions. | | |

| Topic 7.3 Reaction Quo | Topic 7.3 Reaction Quotient and Equilibrium Constant | | |
|--|---|------------------|--|
| Enduring Understandin | g | | |
| TRA-7 A system at equil | ibrium depends on the relationships between concent | rations, partial | |
| pressures of chemical sp | pecies, and equilibrium constant K. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-7.A Represent the reaction quotient Q_c or Q_p , for a reversible reaction, and the corresponding equilibrium expressions $K_c = Q_c$ or $K_p = Q_p$ | TRA-7.A.1 The reaction quotient Q_c describes the relative concentrations of reaction species at any time. For gas phase reactions, the reaction quotient may instead be written in terms of pressures as Q_p . The reaction quotient tends toward the equilibrium constant such that at equilibrium $K_c = Q_c$ and $K_p = Q_p$. | 647-651 | |
| | TRA-7.A.2 The reaction quotient does not include substances whose concentrations (or partial pressures) are independent of the amount, such as for solids and pure liquids. | 633-651 | |

| Topic 7.4 Calculating th | Topic 7.4 Calculating the Equilibrium Constant | | |
|----------------------------------|--|-----------------------|--|
| Enduring Understandin | g | | |
| TRA-7 A system at equil | ibrium depends on the relationships between concent | rations, partial | |
| pressures of chemical sp | pecies, and equilibrium constant K. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-7.B Calculate K _c | TRA-7.B.1 Equilibrium constants can be determined | 630-644; 691-692; 749 | |
| or K_p based on | from experimental measurements of the | | |
| experimental | concentrations or partial pressures of the reactants | | |
| observations of | and products at equilibrium. | | |
| concentrations or | | | |
| pressures at | | | |
| equilibrium. | | | |

| Topic 7.5 Magnitude of | the Equilibrium Constant | |
|-------------------------|--|--------------------|
| Enduring Understandin | g | |
| TRA-7 A system at equil | ibrium depends on the relationships between concer | ntrations, partial |
| pressures of chemical s | pecies, and equilibrium constant K. | |
| Learning Objective | Essential Knowledge | Page numbers |
| TRA-7.C Explain the | TRA-7.C.1 Some equilibrium reactions have very | 632, 644-646 |
| relationship between | large K values and proceed essentially to | |
| very large or very | completion. Others have very small K values and | |
| small values of K and | barely proceed at all. | |
| the relative | | |
| concentrations of | | |
| chemical species | | |
| at equilibrium. | | |

| Topic 7.6 Magnitude of | the Equilibrium Constant | | | |
|--------------------------|---|------------------|--|--|
| Enduring Understandin | g | | | |
| TRA-7 A system at equil | ibrium depends on the relationships between concent | rations, partial | | |
| pressures of chemical sp | pecies, and equilibrium constant K. | | | |
| Learning Objective | Essential Knowledge | Page numbers | | |
| TRA-7.D Represent a | TRA-7.D.1 When a reaction is reversed, K is | 642 | | |
| multistep process | inverted. | | | |
| with an overall | TRA-7.D.2 When the stoichiometric coefficients of | 642-643 | | |
| equilibrium | a reaction are multiplied by a factor c, K is raised to | | | |
| expression, using the | the power c. | | | |
| constituent K | constituent K | | | |
| expressions for each | TRA-7.D.3 When reactions are added together, the | 642-643 | | |
| individual reaction. | K of the resulting overall reaction is the product of | | | |
| | the K's for the reactions that were summed. | | | |
| | | | | |
| | TRA-7.D.4 Since the expressions for K and Q have | 646-651 | | |
| | identical mathematical forms, all valid algebraic | | | |
| | manipulations of K also apply to Q. | | | |
| | | | | |

| Topic 7.7 Calculating E | Topic 7.7 Calculating Equilibrium Concentrations | | |
|-------------------------|---|-------------------|--|
| Enduring Understandi | Enduring Understanding | | |
| TRA-7 A system at equi | ilibrium depends on the relationships between concent | rations, partial | |
| pressures of chemical s | species, and equilibrium constant K. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-7.E Identify the | TRA-7.E.1 The concentrations or partial pressures of | 630-633, 637-638, | |
| concentrations or | species at equilibrium can be predicted given the | 647-651; 676-682, | |
| partial pressures of | balanced reaction, initial concentrations, and the | 687-701, 705-710; | |
| chemical species at | appropriate K. | 726-743, 746-755 | |
| equilibrium based on | | | |
| the initial conditions | | | |
| and the equilibrium | | | |
| constant. | | | |

| Topic 7.8 Representati | ons of Equilibrium | | |
|---|---|--|--|
| Enduring Understandi | ng | | |
| TRA-7 A system at equ | TRA-7 A system at equilibrium depends on the relationships between concentrations, partial | | |
| pressures of chemical s | pressures of chemical species, and equilibrium constant K. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-7.F Represent a system undergoing a reversible reaction with a | TRA-7.F.1 Particulate representations can be used to describe the relative numbers of reactant and product particles present prior to and at equilibrium, and the value of the equilibrium constant. | 639, 651, 654, 662, 665, 668; 683, 701, 710; 753 | |
| particulate model. | · | | |

| Topic 7.9 Introduction | Topic 7.9 Introduction to Le Châtelier's Principle | | |
|------------------------|--|-------------------|--|
| Enduring Understanding | | | |
| TRA-8 Systems at equil | ibrium respond to external stresses to offset the effect | of the stress. | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-8.A Identify the | TRA 8.A.1 Le Châtelier's principle can be used to | 651-658 | |
| response of a system | predict the response of a system to stresses such as | | |
| at equilibrium to an | addition or removal of a chemical species, change in | | |
| external stress, using | temperature, change in volume/ pressure of a gas- | | |
| Le Châtelier's | phase system, or dilution of a reaction system. | | |
| principle. | TRA 8.A.2 Le Châtelier's principle can be used to | 651-658; 729-743, | |
| | predict the effect that a stress will have on | 755-758 | |
| | experimentally measurable properties such as pH, | | |
| | temperature, and color of a solution. | | |

| Topic 7.10 Reaction Qu | Topic 7.10 Reaction Quotient and Le Châtelier's Principle | | |
|--|--|----------------|--|
| Enduring Understandi | Enduring Understanding | | |
| TRA-8 Systems at equil | ibrium respond to external stresses to offset the effect | of the stress. | |
| Learning Objective | Essential Knowledge | Page numbers | |
| TRA-8.B Explain the relationships between Q, K, and the direction in which a reversible | TRA 8.B.1 A disturbance to a system at equilibrium causes Q to differ from K, thereby taking the system out of equilibrium. The system responds by bringing Q back into agreement with K, thereby establishing a new equilibrium state. | 647-651 | |
| reaction will proceed to reach equilibrium. | TRA 8.B.2 Some stresses, such as changes in concentration, cause a change in Q only. A change in temperature causes a change in K. In either case, the concentrations or partial pressures of species redistribute to bring Q and K back into equality. | 647-658 | |

| Topic 7.11 Introduction | Topic 7.11 Introduction to Solubility Equilibria | | |
|-------------------------------|--|-----------------------|--|
| Enduring Understandi | Enduring Understanding | | |
| SPQ-5 The dissolution | of a salt is a reversible process that can be influenced by | environmental factors | |
| such as pH or other dis | solved ions. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SPQ-5.A Calculate | SPQ-5.A.1 The dissolution of a salt is a reversible | 746-753 | |
| the solubility of a salt | process whose extent can be described by K_{sp} , the | | |
| based on the value of | solubility-product constant. | | |
| K _{sp} for the salt. | SPQ-5.A.2 The solubility of a substance can be | 746-764 | |
| | calculated from the K _{sp} for the dissolution process. | | |
| | This relationship can also be used to predict the | | |
| | relative solubility of different substances. | | |
| | SPQ-5.A.3 The solubility rules (see TRA-2.A.5) can be | 737-741 | |
| | quantitatively related to K_{sp} , in which K_{sp} values >1 | | |
| | correspond to soluble salts | | |

Topic 7.12 Common-Ion Effect

| Enduring Understanding | | |
|---|--|--------------|
| SPQ-5 The dissolution of a salt is a reversible process that can be influenced by environmental factors | | |
| such as pH or other dis | solved ions. | |
| Learning Objective | Essential Knowledge | Page numbers |
| SPQ-5.B Identify the | SPQ-5.B.1 The solubility of a salt is reduced when it | 755-758 |
| solubility of a salt, | is dissolved into a solution that already contains one | |
| and/or the value of | of the ions present in the salt. The impact of this | |
| K _{sp} for the salt, based | "common-ion effect" on solubility can be | |
| on the concentration | understood qualitatively using Le Châtelier's | |
| of a common ion | principle or calculated from the K _{sp} for the | |
| already present | dissolution process. | |
| in solution. | | |

| Topic 7.13 pH and Solubility | | | |
|-------------------------------------|---|-----------------------|--|
| Enduring Understandi | ng | | |
| SPQ-5 The dissolution | of a salt is a reversible process that can be influenced by | environmental factors | |
| such as pH or other dissolved ions. | | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SPQ-5.C Identify the | SPQ-5.C.1 The solubility of a salt is pH sensitive | 757-758 | |
| qualitative effect of | when one of the constituent ions is a weak acid or | | |
| changes in pH on the | base. These effects can be understood qualitatively | | |
| solubility of a salt. | using Le Châtelier's principle | | |

| Topic 7.14 Free Energy | Topic 7.14 Free Energy of Dissolution | | |
|--------------------------|--|-------------------|--|
| Enduring Understanding | | | |
| SPQ-5 The dissolution of | SPQ-5 The dissolution of a salt is a reversible process that can be influenced by environmental factors | | |
| such as pH or other dis | solved ions. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SPQ-5.D Explain the | SPQ-5.D.1 The free energy change (△G°) for | 259-262; 529-531; | |
| relationship between | dissolution of a substance reflects a number of | 782-784, 801-803 | |
| the solubility of a salt | factors: the breaking of the intermolecular | | |
| and changes in the | interactions that hold the solid together, the | | |
| enthalpy and entropy | reorganization of the solvent around the dissolved | | |
| that occur in the | species, and the interaction of the dissolved species | | |
| dissolution process. | with the solvent. It is possible to estimate the sign | | |
| | and relative magnitude of the enthalpic and entropic | | |
| | contributions to each of these factors. However, | | |
| | making predictions for the total change in free | | |
| | energy of dissolution can be challenging due to the | | |
| | cancellations among the free energies associated | | |
| | with the three factors cited. | | |

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Unit 8: Acids and Bases

| Topic 8.1 Introduction | Topic 8.1 Introduction to Acids and Bases | | |
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| Enduring Understandi | ng | | |
| SAP-9 The chemistry o | f acids and bases involves reversible proton-transfer rea | ctions, with equilibrium | |
| concentrations being r | elated to the strength of the acids and bases involved. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| SAP-9.A Calculate | SAP-9.A.1 The concentrations of hydronium ion and | 132-133; 677-682; | |
| the values of pH and | hydroxide ion are often reported as pH and pOH, | 757-758 | |
| pOH, based on K _w | respectively. | | |
| and the | EQN: $pH = -log[H_3O^+]$ | | |
| concentration of all | EQN: $pOH = -log[OH^{-}]$ | | |
| species present in a | The terms "hydrogen ion" and "hydronium ion" and | | |
| neutral solution of | the symbols $H^+(aq)$ and $H_3O^+(aq)$ are often used | | |
| water. | interchangeably for the aqueous ion of hydrogen. | | |
| | Hydronium ion and H₃O⁺ (aq) are preferred, but H⁺ | | |
| | (aq) is also accepted on the AP Exam. | | |
| | SAP-9.A.2 Water autoionizes with an equilibrium | 674-676 | |
| | constant Kw. | | |
| | EQN: $K_w = [H_3O^+][OH^-] = 1.0 \times 10^{-14} \text{ at } 25^{\circ}C$ | | |
| | SAP-9.A.3 In pure water, pH = pOH is called a neutral | 677-682 | |
| | solution. At 25°C, pK _w = 14.0 and thus pH = pOH = | | |
| | 7.0. | | |
| | EQN: pK _w = 14 = pH + pOH at 25°C | | |
| | SAP-9.A.4 The value of Kw is temperature | 675 | |
| | dependent, so the pH of pure, neutral water will | | |
| | deviate from 7.0 at temperatures other than 25°C. | | |

Topic 8.2 Introduction to Acids and Bases

Enduring Understanding

SAP-9 The chemistry of acids and bases involves reversible proton-transfer reactions, with equilibrium concentrations being related to the strength of the acids and bases involved.

| Learning Objective | Essential Knowledge | Page numbers |
|-----------------------|---|------------------|
| SAP-9.B Calculate pH | SAP-9.B.1 Molecules of a strong acid (e.g., HCl, HBr, | 134-135; 683-686 |
| and pOH based on | HI, HClO ₄ , H_2SO_4 , and HNO ₃) will completely ionize | |
| concentrations of all | in aqueous solution to produce hydronium ions. As | |
| species in a solution | such, the concentration of H_3O^+ in a strong acid | |
| of a strong acid or a | solution is equal to the initial concentration of the | |
| strong base | strong acid, and thus the pH of the strong acid | |
| | solution is easily calculated. | |
| | SAP-9.B.2 When dissolved in solution, strong bases | 134-135; 683-686 |
| | (e.g., group I and II hydroxides) completely | |
| | dissociate to produce hydroxide ions. As such, the | |
| | concentration of OH- in a strong base solution is | |
| | equal to the initial concentration of the strong base, | |
| | and thus the pOH (and pH) of the strong base | |
| | solution is easily calculated. | |

| Topic 8.3 Weak Acid a | nd Base Equilibria | |
|-----------------------|---|---------------------------|
| Enduring Understandi | ng | |
| SAP-9 The chemistry o | f acids and bases involves reversible proton-transfer rea | actions, with equilibrium |
| | elated to the strength of the acids and bases involved. | |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-9.C Explain the | SAP-9.C.1 Weak acids react with water to produce | 687, 701-715 |
| relationship among | hydronium ions. However, molecules of a weak acid | |
| pH, pOH, and | will only partially ionize in this way. In other words, | |
| concentrations of all | only a small percentage of the molecules of a weak | |
| species in a solution | acid are ionized in a solution. Thus, the | |
| of a monoprotic | concentration of H3 O+ is much less than the initial | |
| weak acid or weak | concentration of the molecular acid, and the vast | |
| base. | majority of the acid molecules remain un-ionized. | |
| | SAP-9.C.2 A solution of a weak acid involves | 687-693, 697-701; |
| | equilibrium between an un-ionized acid and its | 726-729 |
| | conjugate base. The equilibrium constant for this | |
| | reaction is Ka , often reported as pKa . The pH of a | |
| | weak acid solution can be determined from the | |
| | initial acid concentration and the pKa . | |
| | SAP-9.C.3 Weak bases react with water to produce | 693-694, 705-706, |
| | hydroxide ions in solution. However, ordinarily just a | 710-715 |
| | small percentage of the molecules of a weak base in | |
| | solution will ionize in this way. Thus, the | |
| | concentration of OH- in the solution does not equal | |
| | the initial concentration of the base, and the vast | |
| | majority of the base molecules remain un-ionized. | |
| | SAP-9.C.4 A solution of a weak base involves | 693-697 |
| | equilibrium between an un-ionized base and its | |
| | conjugate acid. The equilibrium constant for this | |
| | reaction is Kb , often reported as pKb . The pH of a | |
| | weak base solution can be determined from the | |
| | initial base concentration and the pKb . | |
| | SAP-9.C.5 The percent ionization of a weak acid (or | 692-693 |
| | base) can be calculated from its pKa (pKb) and the | |
| | initial concentration of the acid (base). | |

| Topic 8.4 Acid-Base Reactions and Buffers | | | | |
|---|---|--------------------------|--|--|
| Enduring Understandi | | | | |
| - | f acids and bases involves reversible proton-transfer rea | ctions, with equilibrium | | |
| | concentrations being related to the strength of the acids and bases involved. | | | |
| Learning Objective | Page numbers | | | |
| SAP-9.D Explain the | Essential Knowledge SAP-9.D.1 When a strong acid and a strong base are | 136-137; 736-743 | | |
| relationship among | mixed, they react quantitatively in a reaction | , | | |
| the concentrations of | represented by the equation: | | | |
| major species in a | H^+ (aq) + OH ⁻ (aq) $\rightarrow H_2O(I)$. | | | |
| mixture of weak and | The pH of the resulting solution may be determined | | | |
| strong acids | from the concentration of excess reagent. | | | |
| and bases. | SAP-9.D.2 When a weak acid and a strong base are | 738-741 | | |
| | mixed, they react quantitatively in a reaction | | | |
| | represented by the equation: | | | |
| | $HA(aq) + OH^{-}(aq) \Box A^{-}(aq) H_2O(I).$ | | | |
| | If the weak acid is in excess, then a buffer solution is | | | |
| | formed, and the pH can be determined from the | | | |
| | Henderson-Hasselbalch (H-H) equation (see SAP- | | | |
| | 10.C.1). If the strong base is in excess, then the pH | | | |
| | can be determined from the moles of excess | | | |
| | hydroxide ion and the total volume of solution. If | | | |
| | , they are equimolar, then the (slightly basic) pH can | | | |
| | be determined from the equilibrium represented by | | | |
| | the equation: | | | |
| | A⁻(aq) + H₂O(I) □ HA(aq) + OH⁻ (aq). | | | |
| | SAP-9.D.3 When a weak base and a strong acid are | 741-743 | | |
| | mixed, they will react quantitatively in a reaction | | | |
| | represented by the equation: | | | |
| | B(aq) + H₃O⁺ (aq) □ HB⁺(aq) + H₂O(I). | | | |
| | If the weak base is in excess, then a buffer solution is | | | |
| | formed, and the pH can be determined from the H-H | | | |
| | equation. If the strong acid is in excess, then the pH | | | |
| | can be determined from the moles of excess | | | |
| | hydronium ion and the total volume of solution. If | | | |
| | they are equimolar, then the (slightly acidic) pH can | | | |
| | be determined from the equilibrium represented by | | | |
| | the equation: | | | |
| | $HB^{+}(aq) + H_2O(I) \square B(aq) + H_3O^{+}(aq).$ | | | |
| | SAP-9.D.4 When a weak acid and a weak base are | 736 | | |
| | mixed, they will react to an equilibrium state whose | | | |
| | reaction may be represented by the equation: | | | |
| | $HA(aq) + B(aq) \rightleftharpoons A^{-}(aq) + HB^{+}(aq).$ | | | |
| | $HA(aq) + B(aq) \rightleftharpoons A^{-}(aq) + HB^{+}(aq).$ | | | |

Topic 8.5 Acid-Base Titrations

| Enduring Understandi | ng | |
|------------------------|---|---------------------------|
| SAP-9 The chemistry o | f acids and bases involves reversible proton-transfer rea | actions, with equilibrium |
| concentrations being r | elated to the strength of the acids and bases involved. | |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-9.E Explain | SAP-9.E.1 An acid-base reaction can be carried out | 157-160; 736-743 |
| results from the | under controlled conditions in a titration. A titration | |
| titration of a mono- | curve, plotting pH against the volume of titrant | |
| or polyprotic acid or | added, is useful for summarizing results from | |
| base solution, in | a titration | |
| relation to the | SAP-9.E.2 At the equivalence point, the number of | 157-160; 736-743 |
| properties of the | moles of titrant added is equal to the number of | |
| solution and its | moles of analyte originally present. This relationship | |
| components. | can be used to obtain the concentration of the | |
| | analyte. This is the case for titrations of strong | |
| | acids/bases and weak acids/bases. | |
| | SAP-9.E.3 For titrations of weak acids/bases, it is | 769, 772 |
| | useful to consider the point halfway to the | |
| | equivalence point, that is, the half-equivalence | |
| | point. At this point, there are equal concentrations | |
| | of each species in the conjugate acid-base pair, for | |
| | example, for a weak acid $[HA] = [A^{-}]$. Because pH = | |
| | pK _a when the conjugate acid and base have equal | |
| | concentrations, the pKa can be determined from the | |
| | pH at the half-equivalence point in a titration. | |
| | SAP-9.D.4 For polyprotic acids, titration curves can | 697-701; 775 |
| | be used to determine the number of acidic protons. | |
| | In doing so, the major species present at any point | |
| | along the curve can be identified, along with the pK _a | |
| | associated with each proton in a weak polyprotic | |
| | acid. | |

| Topic 8.6 Molecular Str | ucture of Acids and Bases | |
|-------------------------|---|---------------------------|
| Enduring Understandin | g | |
| SAP-9 The chemistry of | acids and bases involves reversible proton-transfer rea | actions, with equilibrium |
| concentrations being re | lated to the strength of the acids and bases involved. | |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-9.F Explain the | SAP-9.F.1 The protons on a molecule that will | 683-713; 1050-1051 |
| relationship between | participate in acid-base reactions, and the relative | |
| the strength of an acid | strength of these protons, can be inferred from the | |
| or base and the | molecular structure. | |
| structure of the | a. Strong acids (such as HCl, HBr, HI, HClO4, H2 SO4 | |
| molecule or ion. | , and HNO3) have very weak conjugate bases that | |
| | are stabilized by electronegativity, inductive | |
| | effects, resonance, or some combination thereof. | |
| | b. Carboxylic acids are one common class of weak | |
| | acid. | |
| | c. Strong bases (such as group I and II hydroxides) | |
| | have very weak conjugate acids. | |
| | d. Common weak bases include nitrogenous bases | |
| | such as ammonia as well as carboxylate ions. | |
| | e. Electronegative elements tend to stabilize the | |
| | conjugate base relative to the conjugate acid, and | |
| | so increase acid strength. | |

| Topic 8.7 pH and pK _a | | |
|-----------------------------------|--|-----------------------|
| Enduring Understandin | g | |
| SAP-10 A buffered solut | ion resists changes to its pH when small amounts of ac | id or base are added. |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-10.A Explain the | SAP-10.A.1 The protonation state of an acid or base | 687-697 |
| relationship between | (i.e., the relative concentrations of HA and A^-) can | |
| the predominant form | be predicted by comparing the pH of a solution to | |
| of a weak acid or base | the pK_a of the acid in that solution. When solution | |
| in solution at a given | pH < acid pK _a , the acid form has a higher | |
| pH and the pK _a of the | concentration than the base form. When solution | |
| conjugate acid or the | pH > acid pK _a , the base form has a higher | |
| pK_{b} of the | concentration than the acid form. | |
| conjugate base. | SAP-10.A.2 Acid-base indicators are substances | 743-746 |
| | that exhibit different properties (such as color) in | |
| | their protonated versus deprotonated state, | |
| | making that property respond to the pH of | |
| | a solution. | |

| Topic 8.8 Properties of Buffers | | |
|--|---|-------------------------|
| Enduring Understanding | | |
| SAP-10 A buffered solut | ion resists changes to its pH when small amounts of | acid or base are added. |
| Learning Objective | Essential Knowledge | Page numbers |
| SAP-10.B Explain the relationship between the ability of a buffer to stabilize pH and the reactions that occur when an acid or a base is added to a buffered solution. | SAP-10.B.1 A buffer solution contains a large concentration of both members in a conjugate acid-base pair. The conjugate acid reacts with added base and the conjugate base reacts with added acid. These reactions are responsible for the ability of a buffer to stabilize pH. | 729-730 |

| Topic 8.9 Henderson-H | Topic 8.9 Henderson-Hasselbalch Equation | | | |
|------------------------|---|------------------------|--|--|
| Enduring Understandin | Enduring Understanding | | | |
| SAP-10 A buffered solu | tion resists changes to its pH when small amounts of a | cid or base are added. | | |
| Learning Objective | Essential Knowledge Page numbers | | | |
| SAP-10.C Identify the | SAP-10.C.1 The pH of the buffer is related to the | 726-729 | | |
| pH of a buffer | pK _a of the acid and the concentration ratio of the | | | |
| solution based on the | conjugate acid-base pair. This relation is a | | | |
| identity and | consequence of the equilibrium expression | | | |
| concentrations of the | associated with the dissociation of a weak acid, and | | | |
| conjugate acid-base | is described by the Henderson-Hasselbalch | | | |
| pair used to create | equation. Adding small amounts of acid or base to a | | | |
| the buffer. | buffered solution does not significantly change the | | | |
| | ratio of [A ⁻]/[HA] and thus does not significantly | | | |
| | change the solution pH. The change in pH on | | | |
| | addition of acid or base to a buffered solution is | | | |
| | therefore much less than it would have been in the | | | |
| | absence of the buffer. | | | |

| Topic 8.10 Buffer Capa | Topic 8.10 Buffer Capacity | | |
|------------------------|---|------------------------|--|
| Enduring Understandi | Enduring Understanding | | |
| SAP-10 A buffered solu | ition resists changes to its pH when small amounts of a | cid or base are added. | |
| Learning Objective | Essential Knowledge Page numbers | | |
| SAP-10.D Explain the | SAP-10.D.1 Increasing the concentration of the | 726-729 | |
| relationship between | buffer components (while keeping the ratio of | | |
| the buffer capacity of | these concentrations constant) keeps the pH of the | | |
| a solution and the | buffer the same but increases the capacity of the | | |
| relative | buffer to neutralize added acid or base. | | |
| concentrations of the | SAP-10.D.2 When a buffer has more conjugate acid | 729-736 | |
| conjugate acid and | than base, it has a greater buffer capacity for | | |
| conjugate base | addition of added base than acid. When a buffer | | |
| components of | has more conjugate base than acid, it has a greater | | |
| the solution. | buffer capacity for addition of added acid than | | |
| | base. | | |

| Unit 9: | Applications | of Thermody | vnamics |
|------------------------|--------------|-------------|---------|
| O me 3 . | Applications | | / names |

| Topic 9.1 Introduction to Entropy | | | |
|-----------------------------------|---|------------------|--|
| Enduring Understanding | | | |
| ENE-4 Some chemical of | or physical processes cannot occur without interventio | n. | |
| Learning Objective | Essential Knowledge | Page numbers | |
| ENE-4.A Identify the | ENE-4.A.1 Entropy increases when matter becomes | 780-786 | |
| sign and | more dispersed. For example, the phase change | | |
| relative magnitude of | from solid to liquid or from liquid to gas results in a | | |
| the entropy change | dispersal of matter as the individual particles | | |
| associated with | become freer to move and generally occupy a | | |
| chemical or physical | larger volume. Similarly, for a gas, the entropy | | |
| processes. | increases when there is an increase in volume (at | | |
| | constant temperature), and the gas molecules are | | |
| | able to move within a larger space. For reactions | | |
| | involving gas-phase reactants or products, the | | |
| | entropy generally increases when the total number | | |
| | of moles of gas-phase products is greater than the | | |
| | total number of moles of gas-phase reactants. | | |
| | ENE-4.A.2 Entropy increases when energy is | 205-209; 780-786 | |
| | dispersed. According to kinetic molecular theory | | |
| | (KMT), the distribution of kinetic energy among the | | |
| | particles of a gas broadens as the temperature | | |
| | increases. As a result, the entropy of the system | | |
| | increases with an increase in temperature. | | |

| Topic 9.2 Absolute Entropy and Entropy Change | | | | |
|---|---|---------|--|--|
| Enduring Understandi | | | | |
| • | ENE-4 Some chemical or physical processes cannot occur without intervention. | | | |
| Learning Objective | Essential Knowledge Page numbers | | | |
| ENE-4.B Calculate | ENE-4.B.1 The entropy change for a process can be | 786-791 | | |
| the entropy change | calculated from the absolute entropies of the | | | |
| for a chemical or | species involved before and after the | | | |
| physical process | process occurs. | | | |
| based on the | EQN: ΔS° reaction = ΣS° products - ΣS° reactants | | | |
| absolute entropies of | | | | |
| the species involved | | | | |
| in the process. | | | | |

Applications of Thermodynamics

| Topic 9.3 Gibbs Free Er | nergy and Thermodynamic Favorability | |
|------------------------------------|---|--------------|
| Enduring Understandi | ng | |
| ENE-4 Some chemical of | or physical processes cannot occur without intervention | |
| Learning Objective | Essential Knowledge | Page numbers |
| ENE-4.C Explain | ENE-4.C.1 The Gibbs free energy change for a | 794-795 |
| whether a physical or | chemical process in which all the reactants and | |
| chemical process is | products are present in a standard state (as pure | |
| thermodynamically | substances, as solutions of 1.0 M concentration, or | |
| favored based on an | as gases at a pressure of 1.0 atm (or 1.0 bar)) is | |
| evaluation of ΔG° . | given the symbol ΔG° . | |
| | ENE-4.C.2 The standard Gibbs free energy change | 792-794 |
| | for a chemical or physical process is a measure of | |
| | thermodynamic favorability. Historically, the term | |
| | "spontaneous" has been used to describe processes | |
| | for which ΔG° < 0. The phrase "thermodynamically | |
| | favored" is preferred instead so that common | |
| | misunderstandings (equating "spontaneous" with | |
| | "suddenly" or "without cause") can be avoided. | |
| | When ΔG° < 0 for the process, it is said to be | |
| | thermodynamically favored. | |
| | ENE-4.C.3 The standard Gibbs free energy change | 792-795 |
| | for a physical or chemical process may also | |
| | be determined from the standard Gibbs free energy | |
| | of formation of the reactants and products. | |
| | EQN: $\Delta G^{\circ}_{reaction} = \Sigma \Delta G^{\circ}_{f products} - \Sigma \Delta G^{\circ}_{f reactants}$ | |
| | ENE-4.C.4 In some cases, it is necessary to consider | 796-799 |
| | both enthalpy and entropy to determine if a process | |
| | will be thermodynamically favored. The freezing of | |
| | water and the dissolution of sodium nitrate are | |
| | examples of such phenomena. | |

| Topic 9.3 Gibbs Free Er | nergy and T | hermodynar | nic Favorability | (continued) | |
|------------------------------------|---------------|---------------------------|--------------------------|-------------------------|--------------|
| Enduring Understandi | ng | | | | |
| ENE-4 Some chemical of | or physical p | processes car | nnot occur with | out intervention | l <u>.</u> |
| Learning Objective | Essential | Knowledge | | | Page numbers |
| ENE-4.C Explain | ENE-4.C.5 | Knowing the | e values of ∆Ho | and ∆So for a | 796-799 |
| whether a physical or | process at | a given tem | perature allows | ∆Go to be | |
| chemical process is | calculated | directly. | | | |
| thermodynamically | EQN: ∆G° | = ΔH° – Τ ΔS [°] | | | |
| favored based on an | ENE-4.C.6 | In general, t | he temperature | conditions for | 795-796 |
| evaluation of ΔG° . | a process | to be thermo | odynamically fav | vored (∆Go < | |
| | - | | m the signs of <i>L</i> | LHo and ΔSo as | |
| | shown in t | the table belo | ow: | | |
| | ΔH° | ΔS° | Symbols | $\Delta G^{\circ} < 0,$ | |
| | | | | favored | |
| | | | | at: | |
| | < 0 | > 0 | <> | all T | |
| | >0 | < 0 | >< | no T | |
| | >0 | > 0 | >> | high T | |
| | < 0 | < 0 | < < | low T | |
| | | | | | |
| | In cases w | here $\Delta Ho > 0$ |) and $\Delta So < 0, n$ | o calculation | |
| | of ∆Go is ı | necessary to | determine that | the process is | |
| | thermody | namically un | favored (∆Go > | 0). | |

| Topic 9.4 Thermodyna | amic and Kinetic Control | | |
|----------------------|---|-------------------|--|
| Enduring Understand | ing | | |
| ENE-4 Some chemical | or physical processes cannot occur without intervention | l. | |
| Learning Objective | Essential Knowledge Page numbers | | |
| ENE-4.D Explain, in | ENE-4.D.1 Many processes that are | 778-779, 792-794, | |
| terms of kinetics, | thermodynamically favored do not occur to any | 795-796 | |
| why a | measurable extent, or they occur at extremely slow | | |
| thermodynamically | rates. | | |
| favored reaction | ENE-4.D.2 Processes that are thermodynamically | 593-596; 778-779 | |
| might not occur at a | favored, but do not proceed at a measurable rate, | | |
| measurable rate. | are under "kinetic control." High activation energy is | | |
| | a common reason for a process to be under kinetic | | |
| | control. The fact that a process does not proceed at | | |
| | a noticeable rate does not mean that the chemical | | |
| | system is at equilibrium. If a process is known to be | | |
| | thermodynamically favored, and yet does not occur | | |
| | at a measurable rate, it is reasonable to conclude | | |
| | that the process is under kinetic control. | | |

Topic 9.5 Free Energy and Equilibrium

| Enduring Understandi | ng | |
|-------------------------------------|--|----------------------|
| ENE-5 The relationship | between ΔG° and K can be used to determine favorabil | ity of a chemical or |
| physical transformation | n. | |
| Learning Objective | Essential Knowledge | Page numbers |
| ENE-5.A Explain | ENE-5.A.1 The phrase "thermodynamically favored" | 801-803 |
| whether a process is | $(\Delta G^{\circ} < 0)$ means that the products are favored at | |
| thermodynamically | equilibrium (K > 1). | |
| favored using the | ENE-5.A.2 The equilibrium constant is related to free | 801-803 |
| relationships | energy by the equations | |
| between K, ΔG° , and | EQN: $K = e - \Delta G^{\circ}/RT$ and | |
| Т. | EQN: ΔG° = -RT ln K. | |
| | ENE-5.A.3 Connections between K and ΔG° can be | 801-803 |
| | made qualitatively through estimation. When ΔG° is | |
| | near zero, the equilibrium constant will be close to | |
| | 1. When ΔG° is much larger or much smaller than | |
| | RT, the value of K deviates strongly from 1. | |
| | ENE-5.A.4 Processes with $\Delta G^{\circ} < 0$ favor products | 801-803 |
| | (i.e., $K > 1$) and those with $\Delta G^{\circ} > 0$ favor reactants | |
| | (i.e., K < 1). | |

| Topic 9.6 Coupled Reactions | | |
|--|--|-----------------------|
| Enduring Understanding | | |
| ENE-5 The relationship | between ΔG° and K can be used to determine favorabil | lity of a chemical or |
| physical transformatio | n | |
| Learning Objective | Essential Knowledge | Page numbers |
| ENE-5.B Explain the relationship between external sources of energy or coupled reactions and their ability to drive | ENE-5.B.1 An external source of energy can be used to make a thermodynamically unfavorable process occur. Examples include: a. Electrical energy to drive an electrolytic cell or charge a battery. b. Light to drive the overall conversion of carbon dioxide to glucose in photosynthesis | 801-803, 804-805 |
| thermodynamically unfavorable processes. | ENE-5.B.2 A desired product can be formed by coupling a thermodynamically unfavorable reaction that produces that product to a favorable reaction (e.g., the conversion of ATP to ADP in biological systems). In the coupled system, the individual reactions share one or more common intermediates. The sum of the individual reactions produces an overall reaction that achieves the desired outcome and has $\Delta G^{\circ} < 0$. | 804-805 |

| Topic 9.7 Galvanic (Vol | taic) and Electrolytic Cells | | |
|---|---|---------------------|--|
| Enduring Understandi | ng | | |
| ENE-6 Electrical energy can be generated by chemical reactions. | | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| ENE-6.A Explain the | ENE-6.A.1 Each component of an electrochemical | 819-827 | |
| relationship between | cell (electrodes, solutions in the half-cells, salt | | |
| the physical | bridge, voltage/current measuring device) plays a | | |
| components of an | specific role in the overall functioning of the cell. The | | |
| electrochemical cell | operational characteristics of the cell (galvanic vs. | | |
| and the overall | electrolytic, direction of electron flow, reactions | | |
| operational | occurring in each half-cell, change in electrode mass, | | |
| principles of the cell. | evolution of a gas at an electrode, ion flow through | | |
| | the salt bridge) can be described at both the | | |
| | macroscopic and particulate levels. | | |
| | ENE-6.A.2 Galvanic, sometimes called voltaic, cells | 820, 823, 836, 837, | |
| | involve a thermodynamically favored reaction, | 838, 840, 841, 843, | |
| | whereas electrolytic cells involve a | 846, 847 | |
| | thermodynamically unfavored reaction. Visual | | |
| | representations of galvanic and electrolytic cells are | | |
| | tools of analysis to identify where half-reactions | | |
| | occur and in what direction current flows. | | |
| | ENE-6.A.3 For all electrochemical cells, oxidation | 819-821, | |
| | occurs at the anode and reduction occurs at the | 836-840, 846-851 | |
| | cathode. | | |

| Topic 9.8 Cell Potential and Free Energy | | | |
|--|---|--------------|--|
| Enduring Understandi | ng | | |
| ENE-6 Electrical energy | ENE-6 Electrical energy can be generated by chemical reactions. | | |
| Learning Objective | Essential Knowledge | Page numbers | |
| ENE-6.B Explain the | ENE-6.B.1 Electrochemistry encompasses the study | 816, 827-830 | |
| relationship between | of redox reactions that occur within electrochemical | | |
| the physical | cells. The reactions are either thermodynamically | | |
| components of an | favored (resulting in a positive voltage) or | | |
| electrochemical cell | thermodynamically unfavored (resulting in a | | |
| and the overall | negative voltage and requiring an externally applied | | |
| operational | potential for the reaction to proceed). | | |
| principles of the cell. | ENE-6.B.2 The standard cell potential of | 822-827 | |
| | electrochemical cells can be calculated by identifying | | |
| | the oxidation and reduction half-reactions and their | | |
| | respective standard reduction potentials. | | |
| | ENE-6.B.3 ΔG° (standard Gibbs free energy change) | 827-830 | |
| | is proportional to the negative of the cell potential | | |
| | for the redox reaction from which it is constructed. | | |
| | Thus, a cell with a positive E° involves a | | |
| | thermodynamically favored reaction, and a cell with | | |
| | a negative E° involves a thermodynamically | | |
| | unfavored reaction. EQN: $\Delta G^{\circ} = -nFE^{\circ}$ | | |

| Topic 9.9 Cell Potentia | l Under Nonstandard Conditions | |
|-------------------------|---|---------|
| Enduring Understandi | ng | |
| ENE-6 Electrical energy | can be generated by chemical reactions. | |
| Learning Objective | rning Objective Essential Knowledge | |
| ENE-6.C Explain the | ENE-6.C.1 In a real system under nonstandard | 822-835 |
| relationship between | conditions, the cell potential will vary depending on | |
| deviations from | the concentrations of the active species. The cell | |
| standard cell | potential is a driving force toward equilibrium; the | |
| conditions and | farther the reaction is from equilibrium, the greater | |
| changes in the cell | the magnitude of the cell potential. | |
| potential. | ENE-6.C.2 Equilibrium arguments such as Le | 827-835 |
| | Châtelier's principle do not apply to electrochemical | |
| | systems, because the systems are not in equilibrium. | |
| | ENE-6.C.3 The standard cell potential Eo | 827-835 |
| | corresponds to the standard conditions of Q = 1. As | |
| | the system approaches equilibrium, the magnitude | |
| | (i.e., absolute value) of the cell potential decreases, | |
| | reaching zero at equilibrium (when Q = K). | |
| | Deviations from standard conditions that take the | |
| | cell further from equilibrium than Q = 1 will increase | |
| | the magnitude of the cell potential relative to Eo . | |
| | Deviations from standard conditions that take the | |
| | cell closer to equilibrium than Q = 1 will decrease | |
| | the magnitude of the cell potential relative to Eo . In | |
| | concentration cells, the direction of spontaneous | |
| | electron flow can be determined by considering the | |
| | direction needed to reach equilibrium. | |
| | ENE-6.C.4 Algorithmic calculations using the Nernst | 831-835 |
| | equation are insufficient to demonstrate an | |
| | understanding of electrochemical cells under | |
| | nonstandard conditions. However, students should | |
| | qualitatively understand the effects of concentration | |
| | on cell potential and use conceptual reasoning, | |
| | including the qualitative use of the Nernst equation: | |
| | EQN: E = Eo - (RT/nF) ln Q | |
| | to solve problems. | |

| Topic 9.10 Electrolysis and Faraday's Law | | | | |
|---|--------------------------------------|------------------|--|--|
| Enduring Understanding | | | | |
| ENE-6 Electrical energy can be | generated by chemical reactions. | | | |
| Learning Objective | Essential Knowledge | Page numbers | | |
| ENE-6.D Calculate the | ENE-6.D.1 Faraday's laws can be used | 827-830, 846-851 | | |
| amount of charge flow based | to determine the stoichiometry of | | | |
| on changes in the amounts | the redox reaction occurring in an | | | |
| of reactants and products in | electrochemical cell with respect to | | | |
| an electrochemical cell. | the following: | | | |
| | a. Number of electrons transferred | | | |
| | b. Mass of material deposited on or | | | |
| | removed from an electrode | | | |
| | c. Current | | | |
| | d. Time elapsed | | | |
| | e. Charge of ionic species | | | |
| | | | | |
| | EQN: $I = q/t$ | | | |

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