- **A. Draw the two chair** conformations for each of the following di-substituted cyclohexanes. Circle the more stable one.
 - For convenience, you may abbreviate the substituents (Me, Et, Pr, Bu, iPr, tBu, or the like rather than drawing out methyl, ethyl, propyl, butyl, isopropyl, t-butyl....)
 - Assume that a halogen, OH, or NH₂ is smaller than a CH₃ or any other alkyl group.
 - Remember to draw in the hydrogens on each of the "substituted" carbons
- 1. Cis-2-bromo-1-methylcyclohexane

2. Cis-3-isopropyl-1-methylcyclohexane

3. Cis -4-ethyl-1-hydroxycyclohexane

4. trans-2-butyl-1-isopropylcyclohexane

- 5. trans-3-t-butyl-1-methylcyclohexane
- 6. trans -4-chloro-1-propylcyclohexane

B. For each of the following, do two things:

- A. draw the most stable chair form for the more stable stereoisomer for the molecule
- B. identify whether the more stable stereoisomer is cis or trans.
- 7. 1-butyl-2-methylcyclohexane
- 8. 3-t-butyl-1-methylcyclohexane
- 9. 1,4-diethylcyclohexane

C. For each of the following, do two things:

- A. draw the most stable chair form
- B. identify whether the more stable stereoisomer would be the cis or the trans stereoisomer
- 10. Cis-2-chloro-1-ethylcyclohexane
- 11. trans-3-butyl-1-isopropylcyclohexane

12. trans -4-hydroxy-1-t-butylcyclohexane

Organic Chemistry I Jasperse Newman Projections and Cyclohexane Chairs. Steps.

Steps for processing a di-substituted cyclohexane chair:

Summary: Draw chairs; install sticks; install substituents appropriately

- 1. Draw both "right-" and "left-handed" chairs
- 2. Draw in "axial" sticks on the relevant carbons; then draw in "equatorial" sticks on the relevant carbons
 - Use the left-most carbon for your first substituted carbon
- 3. On the left-most carbon, put your first substituent in on both chairs.
 - It should be equatorial in the "right-handed" chair, and axial in the other.
- 4. Use "upper/downer" logic to decide whether the second substituent belongs eq or ax on the first chair (then make it the opposite on the second chair)
 - Draw in the H's on the relavent carbons
- 5. Are the two substituents eq/eq, eq/ax, or ax/ax? This will help recognize relative stability
- 6. If one subst. is forced axial, the preferred chair has the bigger subst. equatorial
- 7. The best cis vs trans isomer has both substituents equatorial.
- 8. Note: To draw and identify the best cis versus trans, just draw a chair with both groups equatorial, and then identify whether that is cis or trans

Steps for Drawing the Best Newman projection

Summary: Draw staggered sticks; install substituents appropriately

- 1. Draw a staggered Newman projection, with three sticks on the "back" carbon and three on the "front". Have a stick up on the back carbon, and one down on the front.
- 2. Draw your biggest substituent on the back carbon on the "up" stick
- 3. Draw your biggest substituent on the front on the "down" "anti" stick
- 4. Fill in the other two back attachments on the other two back-carbon sticks.
- 5. Fill in the other two front attachments on the other two front-carbon sticks.

Steps for Drawing the Worst Newman projection

Summary: Draw eclipsed sticks; install substituents appropriately

- 1. Draw an eclipsed Newman projection, with three sticks on the "back" carbon and three on the "front". Have a stick up on both the back and front carbons.
- 2. Draw your biggest substituent on the back carbon on the "up" stick
- 3. Draw your biggest substituent on the front on the "up" "totally eclipsed" stick
- 4. Fill in the other back and front attachments.

Note: The more severe the eclipsing in the "worst" projection, the greater the rotation barrier

Tips for creating a Newman Projection Energy Diagram

- 1. Use the "worst" (totally eclipsed version) as 0° and 360°.
- 2. 120° and 240° will be the other "eclipsed" conformations => energy crests.
- 3. 60°, 180°, and 300° will be the staggered conformations => energy valleys
- 4. 60° and 300° will be the other two staggered conformations (gauche) => energy valleys.
- 5. To compared the relative energies of the eclipsed crests, evaluate the sizes of the eclipsing substituents (when two non-hydrogens eclipse) and
- 6. To compare the relative energies of the staggered valleys, evaluate the number/severity of gauche interactions