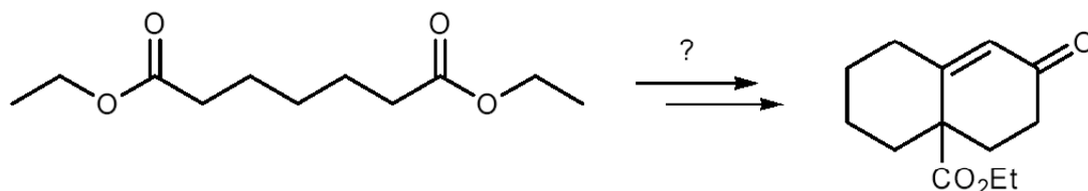


Problem set 2 (for discussion on May 5, 2010)

Exercise 1

Show detailed mechanism



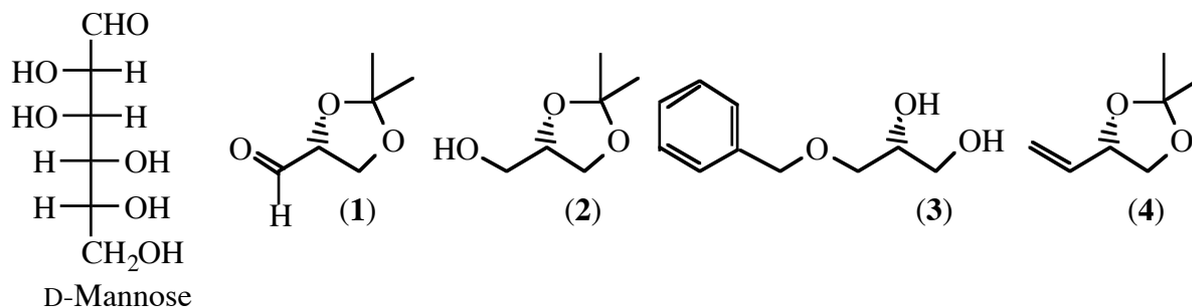
Exercise 2

Propose a synthesis of racemic phenylalanin (Phe) from benzene. Show how the dipeptide Phe-Gly could be made using appropriate protecting groups and reagents.

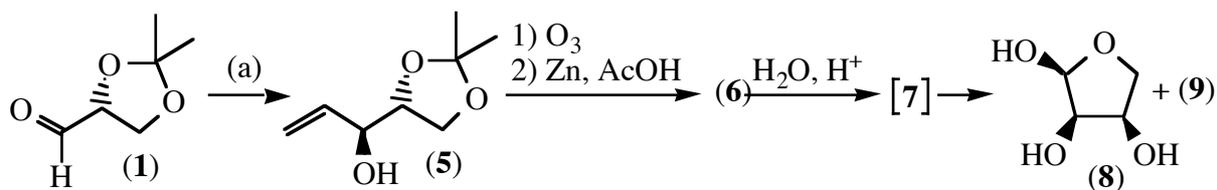
Exercise 3

a) D-mannose is shown below

- D-Mannose is an aldohexose. Explain.
- Draw L-mannose.
- D-Mannose can exist as both α - and β -D-mannopyranose. Draw the β -form.



- The chiral aldehyde **1** can be synthesized from D-mannose. How can compounds **2** – **4** be synthesized from **1**? (Show also synthetic intermediates)
- Compound **8** is available from **1**. Show reagents used in (a) and the structures of intermediates **6** and **7**. Compound **7** is not a stable intermediate and will go directly to **8** and **9** under these reaction conditions. **9** is an isomer of **8**. Show structure of **9**, What kind of isomers are **8** and **9**? In the first step, the yield of **5** is modest. An other compound with the same molecular formula as **5** is formed in almost equal amounts, explain.

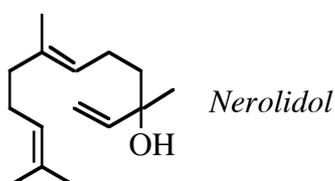


Exercise 4

Draw a Fischer projection of (2*R*,3*S*,4*R*,5*S*,6*S*)-2,3,4,5,6,7-hexahydroxyheptanal. Is this a D- or L sugar? The heptose is subjected to a Wohl degradation. Suggest a mechanism for the reaction and draw the Fischer projection of the product. What will be the more stable chair conformation of the pyranose form of the product? What will be the major pyranose anomer?

Exercise 5 (Try to answer the question without looking up the structure of γ -bisabolene)

- The natural product γ -bisabolene has the molecular formula $C_{15}H_{24}$. Catalytic hydrogenation (H_2 -gas, Pt-cat., AcOH) of γ -bisabolene gives comp. **A**, $C_{15}H_{30}$. How many unsaturations are there in γ -bisabolene, and how many of these are rings.
- In cyclohexane, γ -Bisabolene may be reduced to **B**, $C_{15}H_{28}$. Ozonolysis of **B** gives 6-methyl-2-heptanone and 4-methyl cyclohexanone. Draw **B**.
- Ozonolysis of γ -bisabolene followed by oxidative work up gives among other things acetone and 4-oxopentanoic acid. Formic acid is not a product. Draw possible structures of γ -bisabolene
- γ -Bisabolene can be formed from the natural product nerolidol. Suggest a mechanism and the structure for γ -bisabolene



Exercise 6

Peptide **X** is widely distributed in the body. You have the following information about **X**:

- Vigorous acidic hydrolysis gives Arg, Glu (2), Gly, Leu, Lys, Met, Phe (2), Pro (2)
- Enzymatic hydrolysis gives Arg, Gln (2), Gly, Leu, Lys, Met, Phe (2), Pro (2)
- When **X** is treated with phenylisothiocyanate, phenylthiohydantoin derivatives derived from Arg, Pro, Lys and Pro can be obtained in that order
- Incubation of **X** with chymotrypsin gives peptides **A** and **B**.
- Peptide **A** contains Arg, Gln (2), Lys, Phe, Pro (2). Degradation with Edman's reagent gives the same phenylthiohydantoin derivatives derived from intact **X**. Carboxypeptidase releases first Phe, then Gln
- Peptide **B** reacts with phenylisothiocyanate to give phenylthiohydantoin derivatives derived from Phe, Gly and Leu in that order
- Peptide **X** is strongly basic, pI above 8.9. No amino acids are released when **X** is incubated with carboxypeptidase.

- If **X** is treated with 0.03 M HCL at 110 °C, for 8-12 h (a method that cleaves carboxylic acid amides bonds but leave most peptide bonds untouched) peptide **C** is formed. Peptide **C** reacts with carboxypeptidase to give Gly, Met, Leu and Phe. The order in which these AA were released is not known
- a) What is the N-terminal AA in **X**?
 - b) What is the C-terminal AA in **X**?
 - c) What is the sequence of AAs in peptide **A**?
 - d) What is the sequence of AAs in peptide **B**?
 - e) What is the complete structure of peptide **B**?
 - f) What is the sequence of AAs in peptide **C**?
 - g) What is the structure of **X**?