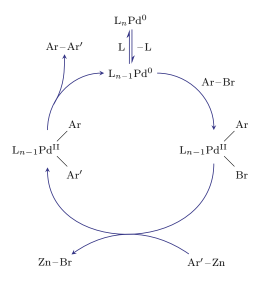
Homework 3 CHEM 20200

- 5/3: 1. Address the following questions.
 - a) Draw a typical mechanism for a Pd catalyzed cross-coupling reaction of an aryl bromide and an aryl zinc reagent.

Answer.



b) Show how this mechanism may differ with a Mizoroki-Heck type reaction involving the coupling of an aryl bromide with an olefin. What additive might be necessary to drive this reaction?

Answer.

With this mechanism, the activation and oxidative addition steps are the same, but then things start to differ. Instead of a transmetallation step, we now have a ligand substitution followed by a 1,2-migratory insertion. Now things start to look a bit similar again as we kick the product out, but here we have a regular elimination step, whereas before we had reductive elimination. Finally, we need one additional last reductive elimination/ligand addition step to regenerate the catalyst. Note that the eliminated acid is neutralized by the added base.

As to the second part of the question, said base is the necessary additive.

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2. A simplified catalytic cycle for hydrocyanation of propylene by NiL_4 ($L = P(OR)_3$) is drawn below. Classify each mechanistic step in the cycle (I-V) using the classifications discussed in class.

Answer.

I Activation (by dissociation).

II Oxidative addition.

III Ligand substitution.

IV 1,2-migratory insertion; ligand addition.

V Reductive elimination.

3. Pd-catalyzed cross couplings with aliphatic groups are typically difficult, especially with bulky alkylhalides. Ni catalysts are typically more effective for these transformations. Starting with a generically ligated Ni^{I} amide complex, $\mathrm{L}_{n}\mathrm{Ni-NPh_{2}}$, show a mechanism for the cross coupling between $\mathrm{LiNPh_{2}}$ and tert-butyliodide that explains this discrepancy.

Answer. I propose the following radical mechanism:

$$\begin{split} t\text{-BuI} + \mathbf{L}_{n} \mathbf{Ni^{I}} - \mathbf{NPh_{2}} & \longrightarrow \mathbf{L}_{n} \mathbf{Ni^{II}} (\mathbf{NPh_{2}}) (\mathbf{I}t\text{-Bu} \cdot) \\ & \longrightarrow \mathbf{L}_{n} \mathbf{Ni^{II}} (\mathbf{NPh_{2}}) (\mathbf{I}) + t\text{-Bu} \cdot \\ & \longrightarrow t\text{-Bu} - \mathbf{NPh_{2}} + \mathbf{L}_{n} \mathbf{NiI} \\ & \frac{\mathbf{LiNPh_{2}}}{-\mathbf{LiI}} t\text{-Bu} - \mathbf{NPh_{2}} + \mathbf{L}_{n} \mathbf{Ni^{I}} - \mathbf{NPh_{2}} \end{split}$$

First off, aliphatic cross couplings are rare because they depend on alkyl electrophiles, and β -H elimination can be a problem. As such, if they do occur, they generally proceed through a radical mechanism. Thus, since first-row transition metals are more likely to participate in one-electron redox chemistry, the first-row metal nickel is favored as a catalyst over the second-row metal palladium.

Homework 3

4. Show generic reactions for the following olefin metathesis processes:

a) Cross metathesis.

Answer.

$$R \longrightarrow R' \longrightarrow R' + M$$

b) Ring-closing metathesis.

Answer.

$$X \longrightarrow X + //$$

c) Ring-opening metathesis polymerization.

Answer.

$$\left\langle \begin{array}{c} \\ \end{array} \right\rangle \longrightarrow \left(\begin{array}{c} \\ \end{array} \right)_{r}$$

Note that although a six-membered ring is pictured, this reaction can proceed with rings containing any number of carbons. Something similar can also happen with alkyne rings. \Box

d) What byproduct is common to many of these reactions?

Answer. Ethylene gas. \Box

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