Asymmetric Nickel-Catalyzed Hydrocarbamoylation of Alkenes

**Proposed mechanism:**

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\begin{align*}
&\text{1} \quad \text{Ni(cod)}_2 (5 \text{ mol}\%), \text{ 2} (5 \text{ mol}\%), \text{ Ph}_3 \text{P} (5 \text{ mol}\%), \text{ Me}_3 \text{Al} (40 \text{ mol}\%) \\
&\text{PhMe, 40 °C, 12 h} \\
&0.1 \text{ mmol scale} \\
&\text{up to 98% yield} \\
&\text{dr up to >20:1} \\
&\text{er up to 97.5:2.5}
\end{align*}
\]

**Significance:** Secondary phosphine oxides (SPO's) have demonstrated to be versatile and robust pre-ligands in a variety of transition-metal-catalyzed transformations due to their stability to air and moisture (see Review below). The authors report the development of a heterobimetallic catalyst system consisting of aluminum and nickel scaffolded on a chiral SPO tether, which enables the enantioselective intramolecular hydrocarbamoylation of alkenes. This method provides access to enantio-enriched pyrrolidones.

**Comment:** The reaction is believed to initiate via tautomeration of pentavalent phosphine oxide 2 to its trivalent phosphinous acid form, 2', which can react with Me_3Al and form complex A. Coordination to nickel(0) generates the active heterobimetallic catalyst B, which complexes to substrate 1a via the Lewis acidic aluminum center, activating the formyl group towards insertion. Following oxidative addition of nickel into the C–H bond, migratory insertion of the pendent olefin and reductive elimination regenerate the catalyst and furnish pyrrolidone 3a.