A Dynamic Kinetic C–P Cross–Coupling for the Asymmetric Synthesis of Axially Chiral P,N Ligands

Pedro Ramírez-López, Abel Ros, Beatriz Estepa, Rosario Fernández, Béla Fiser, Enrique Gómez-Bengoa, José M. Lassaletta

ABSTRACT: The Pd-catalyzed enantioselective C–P cross–coupling between racemic, configurationally stable heterobiaryl triflates andtrialkylsilyldiaryl(dialkyl)phosphines has been used for the synthesis of several families of enantiomerically enriched heterobiaryl phosphines including QUINAP, PINAP, and QUINAZOLINAP analogues, which can be performed with good yields and enantioselectivities using JOSIPHOS-type bidentate phosphines. The strategy relies on two key assumptions: (I) the N-atom of the heterocycle is a better ligand than triflate and, upon oxidative addition, it incorporates into the coordination sphere of the Pd II center to form cationic cyclic intermediates, and (II) the geometry of the palladacycle results in a widening of the angles involved in the stabilization of the stereogenic axis, facilitating a fast interconversion of diastereomeric structures and, hence, a dynamic kinetic C–P cross–coupling reaction. These starting hypotheses are supported by experimental and computational studies.

KEYWORDS: P,N-ligands, QUINAP, DYKAT, Asymmetric Catalysis, C–P coupling, Silylphosphines, Heterobiaryls

1. INTRODUCTION

Axially chiral P,N-ligands have found important applications in the field of asymmetric catalysis. Since the pioneering developments by Brown and co–workers on RhI–catalyzed asymmetric hydroboration/oxidation of styrenes,1 the original ligand QUINAP1 and related axially chiral P,N-ligands have found many other applications in asymmetric catalysis, including RhI–catalyzed hydroboration/amination,2 RhI–catalyzed diboration of alkenes,3 CuI–catalyzed conjugate boration,4 AgI–catalyzed 1,3-dipolar cycloadditions,5 Ni0–catalyzed cycloaddition of 1,2,3,4-benzothiatriazine-1,1(2H)-dioxides with alkenes,6 CuI–catalyzed 1,2-addition of alkynes to enamines7 or iminium ions8 or CuI–catalyzed conjugate addition to alkylidene Meldrum’s acids9 (Scheme 1).

In spite of the excellent ligand properties exhibited by QUINAP, its resolution via stoichiometric PdII complexes10 has been a serious drawback that has hampered its application by the chemical industry. Even being commercially available, its high price has probably excluded its structure from being a common candidate in many exploratory screenings at average research laboratories. A second aspect that has retarded the development of applications for axially chiral P,N-ligands has been the lack of structural variability, as for years QUINAP itself and a few analogues differing in the dialkyl phosphino group10 have been the only available option. These problems have motivated the development of alternative axially chiral P,N ligands, on one side, and studies directed to improve the efficiency and economy of the synthetic methods, on the other. Thus, the groups of Guiry, Carreira, Apponik, Chan and others

Scheme 1. Selected Applications of Axially Chiral P,N-Ligands
have approached the problem by introducing alternative ligands such as QUINAZOLINAP II, PINAP III, PyPHOS IV, and StackPHOS V (Figure 1). These ligands have also been successfully applied in asymmetric catalysis, matching or improving in some cases the results collected with QUINAP in previously developed or new catalytic reactions. However, the methods required for their synthesis are still far from practical: in the best cases the synthesis requires resolution of diastereomers by chromatography or crystallization (PINAP), while most of them (QUINAZOLINAP, PyPHOS, and StackPHOS) have to be prepared by crystallization of stoichiometric amounts of Pd(II) complexes.

Alternative approaches to the synthesis of QUINAP have also been reported. The first practical synthesis avoiding the use of half-equivalents of Pd salts was reported in 2007 by Knochel and co-workers, who exploited the easy chromatographic separation of diastereomeric sulfoxide intermediates, and their easy transformation into enantiopure QUINAP after sulfoxide–lithium exchange, quenching with Ph₂PCl and sulfur, and reduction with Raney-Ni. Two years later, Clayden and co-workers went a step further and, taking advantage of the stereochemical control by the sulfinyl group in heterobiaryl sulfoxides, succeeded in the development of a ‘dynamic thermodynamic resolution’ leading to QUINAP after the functional group transformations mentioned above. This constitutes the first asymmetric synthesis of QUINAP, but still requires the introduction of an enantiopure sulfinyl group as a sacrificial auxiliary.

Therefore, there is still demand of a general methodology for the synthesis of axially chiral heterobiaryls, ideally based on a catalytic asymmetric procedure enabling the introduction of structural variability at both the heterocycle and the diaryl(dialkyl)phosphino group. The direct construction of the stereoaxis by a cross-coupling reaction might appear as the most straightforward approach to these systems. However, in spite of the great progress achieved during the last years in asymmetric Suzuki-Miyaura cross-couplings, the reaction using heterocyclic substrates remains as an unsolved synthetic challenge, presumably due to the interferences caused by the coordination of the heteroatoms located on the substrate, the limitations associated with the availability and poor stability of heteroaromatic organometallics, and the lower configurational stability of the products compared to standard biaryls. In the frame of our research program in asymmetric cross-coupling reactions, we recently developed an alternative methodology for the asymmetric synthesis of axially chiral heterobiaryls consisting of a dynamic asymmetric Suzuki-Miyaura coupling (DYKAT) between racemic, configurationally stable heterobiaryl triflates and arylboroxines (Scheme 2, eq. 1). In a parallel work, we started investigations to apply this strategy for C–P bond forming reactions while, simultaneously, Virgil, Stoltz, et al. also reported the enantioselective synthesis of QUINAP according to a similar procedure (eq. 2). In both cases, a single example (QUINAP itself) was reported, and different mechanisms involved in the dynamic kinetic cross-coupling were invoked. In this paper, an expanded, general procedure for the synthesis of enantiomerically enriched isoquinoline, 3-methylpyridine, quinazoline and phthalazine derivatives is reported (eq. 3), along with experimental and computational support for a mechanism based on the labilization of cyclic, cationic oxidative addition intermediates.

**Dynamic kinetic asymmetric Suzuki-Miyaura coupling (ref. 18)**

Scheme 2. DYKAT Techniques for the Asymmetric Synthesis of Heterobiaryls

**RESULTS AND DISCUSSION**

**Starting Hypothesis and Method Development.** Basically, our strategy is based in two key assumptions: 1) thanks to the poor coordinating ability of the triflate anion, the oxidative addition of Pd⁻⁻ LL’ catalysts [LL’ = chiral ligand(s)] should generate cyclic cationic intermediates OAI and OAI’ incorporating the isoquinoline/pyridine N atom as a ligand (Scheme 3 and 2) a widening of the angles φ and φ' would compromise the configurational stability of the stereogenic axis, facilitating an easy equilibration of atropisomeric intermediates OAI and OAI’. In this scenario, two additional conditions are required to achieve a highly enantioselective dynamic kinetic C–P coupling to products P or P’:

- 3) the transmetalation step from both OAI and OAI’ into intermediates T1 and T2 should be relatively slow with respect to the interconversion between OAI and OAI’ and
- 4) the chiral ligand(s) LL’ should provide a substantial energy gap between the diastereomeric transmetalation transition states.

Our initial ligand screening was performed using the coupling of triflate 1A and tBuMe₃SiPPh₂ as a model reaction, with CsF as the base, dry THF at 55 °C as the solvent and 5 mol% of the base.
Pd\((\text{dba})_3\)/10 mol% ligand as the catalyst system (Scheme 4). The use of silylphosphine reagents,\(^{24}\) whose reactivities can be tuned by adjusting the steric and/or electronic properties of the silyl group, was envisaged as a potentially useful method to modulate the rate of release of the phosphine fragment; according with our strategy, a relatively slow transmetallation step is necessary to facilitate the equilibration of the oxidative addition intermediates. Bishydrazone and phosphino–hydrazine ligands L\(1\) and L\(2\), which showed very good enantioselectivities and activities in Suzuki–Miyaura cross-couplings,\(^{19}\) and a privileged ligand such as the phosphino oxazoline L\(3\), were employed as candidates and tested in the model reaction. These ligands provided moderate to excellent conversions into the desired product 3Aa, although low enantioselectivities were observed. Motivated by the excellent performance of these type of ligands in dynamic asymmetric (DYKAT) Suzuki–Miyaura couplings,\(^{25}\) we also examined TADDOL-derived phosphoramidite ligand L\(4\) and related binaphthol-derivatives L\(5\) and L\(6\). L\(5\)-L\(6\) afforded 3Aa in >90% conversions but with low enantioselectivities. Axially chiral, commercially available ligands such as BINAP, MeO-MOP, and SEGPHOS L\(7\)-L\(9\) also provided high catalytic activity but only in the last case a low yet significant 63:37 er was observed. Additionally, a kind of ‘autocatalytic’ reaction promoting QUINAP L\(6\) played during the model reaction was also considered. Using commercially available \((S)\)-QUINAP as the ligand, a moderate ~65% conversion was observed after 20 hours and the product 3Aa was obtained with 67:33 er, thus demonstrating that QUINAP itself can be used for its own synthesis, although in an inefficient manner. P-stereogenic ligands L\(10\)-L\(11\) and Josiphos-type ligands with planar chirality L\(12\)-L\(18\) were also tested in the model reaction, yielding the desired product 3Aa with moderate to excellent conversions. The reactions with ligands L\(12\) and L\(16\) proved to be the most selective (er 77:23 and 78.5:21.5, respectively). Due to the higher activity of L\(12\) (~58 and 34% conversions after 6 h for L\(12\) and L\(16\), respectively), it was selected for the following optimization studies, devoted to explore the effect of the phosphorous reagent and the Pd precatalyst. Silylphosphines with different steric and electronic properties of the silyl fragment, as well as other phosphine sources such as HPPh\(6\) and KPPh\(7\) were tested in the model reaction (1A—3Aa), using 2 equiv of CsF, dry THF as the solvent and 10 mol% L\(12\)/5 mol% Pd\((\text{dba})_3\), as the catalyst (Table 1, entries 1-7). The size of the trialkysilyl

\[\text{Scheme 3. Starting Mechanistic Hypothesis}\]

\[\text{Scheme 4. Ligand Screening}^{a}\]

\(^{a}\) Reactions performed at 0.1 mmol scale in dry THF (2 mL). Conversions estimated by \(^1\)H NMR. Enantiomeric ratios were determined by HPLC on chiral stationary phases after oxidation of 3Aa to the corresponding phosphine oxide. \(^{b}\) Reaction time was 38 h. \(^{c}\) Reaction time was 6 h.
Table 1. Condition Optimization for the Synthesis of QUINAP

<table>
<thead>
<tr>
<th>MPPH₂: L</th>
<th>[Pd]</th>
<th>t (°C)</th>
<th>Conv (%)</th>
<th>er (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>L12</td>
<td>Pd₂(dba)₃</td>
<td>55</td>
</tr>
<tr>
<td>2</td>
<td>4</td>
<td>L12</td>
<td>Pd₂(dba)₃</td>
<td>55</td>
</tr>
<tr>
<td>3</td>
<td>5a</td>
<td>L12</td>
<td>Pd₂(dba)₃</td>
<td>55</td>
</tr>
<tr>
<td>4</td>
<td>5a</td>
<td>L12</td>
<td>Pd₂(dba)₃</td>
<td>40</td>
</tr>
<tr>
<td>5</td>
<td>6</td>
<td>L12</td>
<td>Pd₂(dba)₃</td>
<td>55</td>
</tr>
<tr>
<td>6</td>
<td>7</td>
<td>L12</td>
<td>Pd₂(dba)₃</td>
<td>-20</td>
</tr>
<tr>
<td>7</td>
<td>7°</td>
<td>L12</td>
<td>Pd₂(dba)₃</td>
<td>40</td>
</tr>
<tr>
<td>8</td>
<td>5a</td>
<td>L12</td>
<td>Pd₂(dba)₃</td>
<td>40</td>
</tr>
<tr>
<td>9</td>
<td>5a</td>
<td>L15</td>
<td>Pd₂(dba)₃</td>
<td>40</td>
</tr>
<tr>
<td>10</td>
<td>5a</td>
<td>L16</td>
<td>Pd₂(dba)₃</td>
<td>40</td>
</tr>
<tr>
<td>11</td>
<td>5a</td>
<td>L17</td>
<td>Pd₂(dba)₃</td>
<td>40</td>
</tr>
<tr>
<td>12</td>
<td>5a</td>
<td>L12</td>
<td>Pd₂(dba)₃</td>
<td>40</td>
</tr>
<tr>
<td>13</td>
<td>5a</td>
<td>L12</td>
<td>Pd(OAc)₂</td>
<td>40</td>
</tr>
<tr>
<td>14</td>
<td>5a</td>
<td>L16</td>
<td>Pd₂(dba)₃</td>
<td>40</td>
</tr>
<tr>
<td>15</td>
<td>5a</td>
<td>L12°</td>
<td>Pd₂(dba)₃</td>
<td>40</td>
</tr>
<tr>
<td>16</td>
<td>5a</td>
<td>L12°</td>
<td>Pd₂(dba)₃</td>
<td>40</td>
</tr>
</tbody>
</table>

*Reactions performed on a 0.1 mmol scale using anhydrous THF (2 mL/0.1 mmol 1A), 2 equiv phosphine source and 2 equiv of CsF. Conversions were estimated by 1H-NMR and refer to the consumption of the starting triflate. Enantiomeric excesses were determined by HPLC on chiral stationary phases, after oxidation of 3Aa to the corresponding phosphine oxide. Commercially available Me₅SiPPh₂ 5a from Aldrich containing 8% of HPPH₂ 6, was employed. A 0.2 M solution in THF of KPPH₂ 7 was slowly added (0.1 mL/h) at 40 °C using a syringe pump, for a 10 hour-period. Synthetic, HPPh₂-free Me₅SiPPh₂ 5a was employed. 20 mol% of L12 was used. Freshly prepared triflate 1A was used.

Further efforts were directed to extend the scope of the methodology to related systems. To this aim, silylphosphines 5a–e were synthesized by a modified protocol of the described procedure and reacted with heterobiaryl triflates 1A–D or nonaflates 8C–D in the presence of Pd(dba)₂/L12 as the catalytic system and CsF as the base. Heterobiaryl phosphines 3 were obtained in excellent yields and good enantiomeric excesses (Table 2). Silylphosphines

Table 2. Dynamic Kinetic Asymmetric C–P Couplings: Scope
excellent yields and enantioselectivities (er 85.5:14.5-95:5). Additionally, the heterobiaryl dialklyphosphine 3Be could also be obtained in 93% yield and a good 85:15 er when L16 was used as the ligand. Reproducibility problems were observed when triflate 1C was employed as the substrate, due presumably to some undetected impurity. In this case, we resorted to using nonaflate 8C, which allowed to obtain the desired products 3Ca-3Cd in excellent yields and enantioselectivities of 79-90%. Dialkyl heterobiarylphosphine 3Ce was obtained in 72% yield and a moderate 72.5:27.5 er. Phthalazine-derived triflate 1D was also tested in the C-P coupling reaction, giving the corresponding biaryl- phosphines 3Da-3Db in 71 and 73% yield, and 85.5:14.5 and 91:9 er, respectively. The enantimeric purity of products 3 could be increased by crystallization in some cases. As representative examples, QUINAP itself (3Aa) and 3Ca were obtained in 99.5:0.5 er after a single recrystallization or washing with cold acetone, respectively. In the case of 3Ac, the crystallization of small amount of the racemate served to increase the optical purity of the remaining mother liquors (up to 97.5:2.5 er after a single crystallization).

**Mechanism and Computational Studies.** In the preliminary investigations mentioned above, two different mechanisms were postulated to explain the observed dynamic asymmetric cross-coupling. On one side, our group assumed that the nitrogen atom of the isoquinoline coordinates to the Pd center, leading to diastereomeric, cyclic OAI and OAI’ intermediates, displacing the poorly coordinating triflate anion. The geometry of these OA intermediates suggests that the equilibration proceeds via a transition state TS_{enol-OAI} in which the angles $\phi_1$ and $\phi_2$ are just slightly wider to allow hydrogen atoms H(8) and H(8') to reach coplanarity with the stereogenic axis (Scheme 5, path a). On the other hand, Virgil, Stoltz et al. suggested that unsaturated T-shaped OAI, and OAI’ intermediates equilibrate via a square planar transition state TS_{enol-OAI’s} stabilized by an agostic Pd–H(8) interaction (path b).

---

* Reactions performed on a 0.1 mmol scale. Synthetic and phosphine–free silylphosphines 5a-e were used in all cases. Isolated yields and er’s determined by HPLC on chiral stationary phases are shown. * Triflate 1 was used as starting material. * Er after recrystallization (toluene/CH2Cl2). * Er of mother liquor after crystallization of minor amounts of racemate (n-hehane/AcOEt). * Air-sensitive compounds: fast flash chromatography under nitrogen was required for purification. * L16 was used instead of L12. * Nonaflate 8 was used as starting material. * Reaction time 40 hours. * Er after washing with cold acetone.

5b-d, which bear electron withdrawing and electron-donating groups on the aryl fragment, afforded the corresponding chiral phosphines 3Ab-3Ad in 74-88% isolated yields and er’s from 78:22 to 92:8. Pyridine-derived phosphines 3Ba-Bd were also obtained in excellent yields and enantioselectivities (er 85.5:14.5-95:5). Additionally, the heterobiaryl dialklyphosphine 3Be could also be obtained in 93% yield and a good 85:15 er when L16 was used as the ligand. Reproducibility problems were observed when triflate 1C was employed as the substrate, due presumably to some undetected impurity. In this case, we resorted to using nonaflate 8C, which allowed to obtain the desired products 3Ca-3Cd in excellent yields and enantioselectivities of 79-90%. Dialkyl heterobiarylphosphine 3Ce was obtained in 72% yield and a moderate 72.5:27.5 er. Phthalazine-derived triflate 1D was also tested in the C-P coupling reaction, giving the corresponding biaryl- phosphines 3Da-3Db in 71 and 73% yield, and 85.5:14.5 and 91:9 er, respectively. The enantimeric purity of products 3 could be increased by crystallization in some cases. As representative examples, QUINAP itself (3Aa) and 3Ca were obtained in 99.5:0.5 er after a single recrystallization or washing with cold acetone, respectively. In the case of 3Ac, the crystallization of small amount of the racemate served to increase the optical purity of the remaining mother liquors (up to 97.5:2.5 er after a single crystallization).

**Mechanism and Computational Studies.** In the preliminary investigations mentioned above, two different mechanisms were postulated to explain the observed dynamic asymmetric cross-coupling. On one side, our group assumed that the nitrogen atom of the isoquinoline coordinates to the Pd center, leading to diastereomeric, cyclic OAI and OAI’ intermediates, displacing the poorly coordinating triflate anion. The geometry of these OA intermediates suggests that the equilibration proceeds via a transition state TS_{enol-OAI} in which the angles $\phi_1$ and $\phi_2$ are just slightly wider to allow hydrogen atoms H(8) and H(8') to reach coplanarity with the stereogenic axis (Scheme 5, path a). On the other hand, Virgil, Stoltz et al. suggested that unsaturated T-shaped OAI, and OAI’ intermediates equilibrate via a square planar transition state TS_{enol-OAI’s} stabilized by an agostic Pd–H(8) interaction (path b).

---

**Scheme 5. Proposed Mechanisms for the Epimerization of Diastereomeric Oxidative Addition Intermediates.**

In this last proposal, the presence of a coordinating nitrogen atom on the heteroaryl moiety does not play any role and the DYKAT process should work with triflate 9. However, no C–P
coupling products was observed from the reaction of 9 with 5a, which afforded an incomplete conversion into hydrolysis product 10 after overnight heating at 40 °C (Scheme 6).

![Scheme 6. Experiments with Triflate 9 and Nonafate 11.](image)

This experiment reveals that the presence of a coordinating isoquinolyl/pyridyl nitrogen is not only necessary to favor the formation of the cationic and configurationally labile palladacycle, but also to facilitate the chelate-assisted oxidative addition of the racemic triflate/nonafate to the Pd0 center. Similarly, the reaction of NOBIN-derived nonafate 11 with 5a afforded a low conversion into hydrolysis product 12, even at a higher temperature (60 °C), indicating that the formation of a five-membered cationic palladacycle is essential to reactivity.

In order to gain further insight into the mechanism of the reaction, we decided to set out an in depth DFT computational investigation of the process. According to the aforementioned data, the optimal experimental conditions involved the use of aromatic phosphine (PPh3) as a model for the incoming nucleophilic phosphide bonding interaction, is similar (ca. 2.1 Å) in the ground states of irradiation/epimerization process, like the reduction of the q1 angle to 45° (Scheme 8). Meanwhile, after the oxidative addition of Pd(0) to the C-OTf bond, the chelation complex R-int1-PMe3 presents drastic geometrical changes in the right direction to facilitate the rotation/epimerization process, like the reduction of the q1 angle to 45° (Scheme 8). Furthermore, the N-Pd distance, which is a favorable, bonding interaction, is similar (ca. 2.1 Å) in the ground states of R- or S-int1-PMe3 and during the transition state for the rotation (Ts1). Thus, the energy barrier is dramatically reduced to 18.7 kcal/mol, explaining the fast interconversion of both enantiomeric complexes in the experimental conditions. These results are the confirmation that the palladacyclic intermediates can easily racemize whereas the initial triflates can not.

We found computational evidence supporting our working hypothesis based on the following results. The interconversion of the two enantiomeric triflates R-1A and S-1A is predicted to be an extremely slow process in the absence of palladium, since its activation energy is as high as 29.3 kcal/mol (Tsrot-1A-anti, Scheme 8). There is a clear difficulty in the substrate to adopt the necessary conformation for the rotation. In the ground state, the angle formed by the planes of the two aromatic rings θ, is ca. 90°, but it must approach to 0° during the transition state Tsrot-1A-syn (34.9 kcal/mol, Scheme 8), with the accompanying energetic cost. The rotation is easier through the anti approach, but still too high to be feasible at room temperature (29.3 kcal/mol, Tsrot-1A-anti). Meanwhile, after the oxidative addition of Pd(0) to the C-OTf bond, the chelation complex R-int1-PMe3 presents drastic geometrical changes in the right direction to facilitate the rotation/epimerization process, like the reduction of the q1 angle to 45° (Scheme 8). Furthermore, the N-Pd distance, which is a favorable, bonding interaction, is similar (ca. 2.1 Å) in the ground states of R- or S-int1-PMe3 and during the transition state for the rotation (Tsrot-int1). Thus, the energy barrier is dramatically reduced to 18.7 kcal/mol, explaining the fast interconversion of both enantiomeric complexes in the experimental conditions. These results are the confirmation that the palladacyclic intermediates can easily racemize whereas the initial triflates can not.
implies that 

**Figure 2. Structures of the Two Most Stable Palladacyclic Intermediates.**

The use of an achiral phosphine (PMe₃) as a first approximation implies that **R-int1-PMe₃** and **S-int1-PMe₃** are isoenergetic (Scheme 8), but the computation of the experimental chiral ligand L12 makes the corresponding complexes **R-int1-L12** and **S-int1-L12** diastereomeric (Figure 2), differentiating their energies. Both complexes can still interconvert, and the equilibrium is clearly shifted towards the pro-R complex **R-int1-L12**, which is 4.3 kcal/mol lower in energy than its pro-S counterpart. The two complexes in Figure 2 are actually the lowest in energy intermediates of a total of eight possible diastereomeric species, the rest of them being remarkably higher in energy (between 5.3 to 13.4 kcal/mol) than **R-int1-L12**. Four of these species are the result of the fact that the palladium center is not forming a perfect square planar structure. The two cycles around the palladium atom are not coplanar, and form a pseudo-tetrahedral geometry with two different configurations. Also, the relative disposition of the two phosphorous atoms in the asymmetric di-phosphine (L12) with respect to the C and N atoms of the aromatic rings leads to the formation of the other four diastereoisomers.

It is important to note that the overall lowest in energy intermediate (**R-int1-L12**) corresponds to the *minor* experimental enantiomer R. However, this fact is irrelevant in the present Curtin-Hammett conditions, since all possible isomers are involved in a fast equilibrium, and the final outcome of the reaction is determined by the relative activation energy of the different transmetalation transition states. At this point, attention must be drawn to the fact that the transmetalation step is not in fact a *classical* transmetalation, since it actually consists on an isoquinoline/Me₃SiPPh₂ ligand exchange, with subsequent abstraction of the silyl moiety by the CsF salt (Scheme 9). This system is difficult to compute, but a slightly altered model was envisioned, consisting on the replacement of the actual Me₃SiPPh₂ ligand by PPh₃ (int2-Si vs int2-Ph). This alternative has the great advantage that triphenylphosphine is conformationally much simpler than Me₃SiPPh₂, while maintaining a similar steric hindrance.

**Scheme 9. Two-step Transmetalation Process, and the Computed PPh₃ Model**

Thus, all possible transition structures for the approach of the nucleophilic phosphine were computed, maintaining in all cases similar distances for the coordination of the phosphine to palladium and decoordination of the isoquinoline-nitrogen from the metal (Figure 3). As mentioned before, there are eight main isomers for the int-1 type complexes, and for each of them we found at least two different transition structures, depending on the departing trajectory followed by the isoquinoline. Compare for example the two lowest in energy structures for the *S* enantiomer (**S-TS2-a, S-TS2-b**), in which the isoquinoline ligand leaves the palladium sphere towards the lower or upper face, respectively. All of the possible alternatives were computed, resulting in a large range of activation energies (ca. 10 kcal/mol), but only the most stable ones.
are shown in Figure 3. To our delight, the most favored approach (S-TS2-a) is in agreement with the formation of the experimental major S enantiomer, and the difference with the lowest pro-R structure (R-TS2-a) is 4.2 kcal/mol, large enough to explain a high selectivity in the process. Even more, the second favored structure also corresponds to the S-enantiomer (S-TS2-b, +0.9 kcal/mol). Taking the results of Schemes 8 and Figure 3 together, we are facing the typical situation where the minor, less stable isomer (S-int1-L12) reacts faster than the major unreactive one (R-int1-L12), nicely explaining the experimental stereoselectivity results (Figure 4). It is also important to note that the computed activation energy of the (S)-TS2-a is $\Delta G^\ddagger = 25.2$ kcal/mol from the separate PPh$_3$ and (R)-int1-L12, and can be safely considered the rate limiting step, as it is much larger than either isomerization, or reductive elimination.

The final C-P reductive elimination step was also computed (Scheme 10), affording interesting data. First, the activation barrier for TS3 is very low comparing to the rest of energies found in this study ($\Delta G^\ddagger = 12.5$ kcal/mol), making the whole process after transmetalation irreversible. Secondly, the rotation barrier for the int3-type intermediates and for the final QUINAP-type products raise to 34.7 and 32.5 kcal/mol, respectively. Racemization processes at the final stages of the reaction are, thus, unfeasible, confirming that the formation of palladacycles by chelation with the nitrogen of the isoquinoline (like in int1) is a mandatory condition to allow the epimerization of the substrates.

\[
\Delta G^\ddagger = +4.2 \\
\Delta G^\ddagger = +4.3
\]

\[
\Delta G^\ddagger = +12.5 \\
\Delta G^\ddagger = +9.3
\]

Finally, we also checked the possible involvement of the palladium center in a Pd-H agostic bond (int4, Figure 5), which has been proposed by Virgil and Stolz$^{21}$ to explain the epimerization of the oxidative addition intermediates without the intervention of the nitrogen atom. Our calculations show that the model complex int4 is remarkably more unstable ($\Delta G^\ddagger = +9.3$ kcal/mol) than the simple int1-type palladacycle. This large energy difference is enough to completely discard the participation of int4 in the mechanism.

Figure 3. Most Stable Transition States for the Approach of PPh$_3$ to the Pd center, computed at M06/6-31+G(d,p)/(defpcm,THF)//B3LYP/6-31G(d,p)

Figure 4. Energy diagram for the transmetallation step.

CONCLUSION

In summary, the dynamic kinetic asymmetric C–P coupling between heterobiaryl triflates or nonaflates and trimethylsilylphosphines appears as an efficient, general methodology for the asymmetric synthesis of QUINAP, PyPHOS, QUINAZOLINAP, and PINAP analogues. The collected experimental evidences and the results of the performed computational study allow to propose a mechanism based on the formation of cationic oxidative addition intermediates that, under the reaction conditions, undergo a fast interconversion. Coordination of the isoquinoline N atom to Pd is essential to facilitate this process. The calculations also show that the energy requirements for a dynamic kinetic process are met, since the fast equilibrating palladacyclic intermediates evolve through diastereomeric transmetalation steps of very large energy...
difference. The easiness of the final reductive elimination ensures the irreversibility of the process.

ASSOCIATED CONTENT
Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemarxiv.xxx

Experimental procedures, optimization studies, characterisation data, NMR spectra for new compounds, and HPLC traces (PDF). Complete description of the computed structures and energies (PDF).

ACKNOWLEDGMENT

AUTHOR INFORMATION

Corresponding Authors

*abel.ros@uji.csic.es
*ffernan@us.es
*enrique.gomez@ehu.es,
*jmlassa@uji.csic.es

REFERENCES

The authors declare no competing financial interest.

ACKNOWLEDGMENT

The authors thank for technical and human support provided by IZOSGI SGiker of UPV/EHU, European funding [ERDF, ESF and FP7 (ITN ECHONET network MCTIN-2012-316379)], Ministerio de Economía y Competitividad (grants CTQ2013-48164-C2-1-P and CTQ2013-48164-C2-2-P, contract RYC-2013-12585 for AR), and the Junta de Andalucía (Grant 2012/FQM 1078).


27. The S configuration was assigned by chemical correlation. See supporting information for details.

28. Triflates were stored in the fridge for months to prevent the formation of triflic acid traces, affording reproducible enantiomeric excesses throughout this study without the need of further purifications.


32. See Supporting Information for Computational Details.


34. We assume that the barriers of isomerization or reductive elimination will not be strongly affected if the real ligand L12 is used instead of the model bis-PMe₃ surrogate.
racemic \text{OTf} + \text{Me}_3\text{SiPR}_2 \xrightarrow{\text{Pd}\text{LL}^+ \text{(cat)}} \text{high yields up to 95.5:4.5 er} \quad \phi_1, \phi_2 > 120^\circ