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Pincer versus pseudopincer: isomerism in palladium(II) complexes bearing $\kappa^3 C$, S, C ligands \dagger

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In NHC pincer complexes incorporating a hemilabile donor site, there exists an equilibrium between the true pincer form and a pseudopincer coordination isomer. The influence of the NHC

moieties on this isomerism has been studied by DFT calculations.

Tridentate, pincer and pincer-type ligands have gained increasing popularity due to the robustness and thermal stability the rigid, κ^3 -mer or κ^3 -(pseudo)-mer coordination geometry of these ligands confers on their complexes. Transition metal complexes incorporating pincer ligands have been shown to be versatile and useful catalysts in organic synthesis. Beyond catalysis, pincer complexes have also been utilized successfully for the development of new materials.² The fine-tuning of the steric and electronic properties of pincer ligands - be it by modifications in the backbone or changes to the nature of the donor sites - has attracted considerable research interest, in order to provide coordination spheres for each and every application.3 While catalysis often involves the coordination sites on the metal centre not occupied by the pincer ligand, there are also cases in which partial decoordination of a hemilabile donor takes place. The advantage is the rapid creation of a coordinatively and electronically unsaturated metal centre, open for incoming substrates, while retaining the tethered donor functionality in the vicinity in order to allow for rapid recoordination and stabilisation of the active species. As a result, highly active yet stable catalysts are obtained.4 Frequently, the donor functionalities in the side chains of the pincer ligands are hemilabile sites,5 but cases in which the central donor moiety exhibits hemilabile behaviour have also been reported.⁶ However, to control hemilabile behaviour, the

governing geometric and electronic factors still need to be understood. 7

N-heterocyclic carbenes (NHCs) have become ubiquitously employed ligands for transition metal catalysts due to their strong σ -basicity, the robustness of their complexes, and the ease with which they can be synthesized. Their steric bulk and electronic properties can be fine-tuned rapidly by modifying the backbone and side chains, and additional donor functionalities can be introduced in the side chains, which give rise to chelating and pincer-type ligands. Donor moieties based on heteroatoms such as nitrogen, oxygen, phosphorus, and sulfur have been reported. Phosphorus, and sulfur have been reported. Especially the soft thioether functionalities are known to exhibit hemilabile behavior in their complexes.

It has been shown that in the presence of coordinating counteranions, complexes of CSC pincer ligands are isolated either as κ^2 -trans-C,C pseudopincer complexes or in their κ^3 -mer-C,S,C pincer form. The electron-donating abilities of the carbene moieties were suggested to have a pivotal influence on this isomerism. To study this electronic effect, we examined the electronic properties of NHCs with markedly different donor strengths and the preference for either the pincer or the pseudopincer form by means of computational methods.

The relative Gibbs free energies ΔG of a neutral pseudopincer species $\bf A$ and the corresponding cationic pincer $\bf C$ and a bromide ion are difficult to obtain due to the significant impact of solvation on the ion pair. Instead, the Gibbs free energy ΔG_R of the reaction of a pincer and a pseudopincer complex was used as an indication of the preference for the pincer form (Scheme 1). During this fictitious reaction, the number of ionic species remains constant throughout the reaction, and solvation effects as well as the Gibbs free energy of the bromide counteranion can be neglected as they are almost identical for the starting materials and products.

The resulting scale of pincer preference energy is arbitrary in the sense that the absolute values will depend on the choice of the reference system. We decided to use complexes **B** and **D** featuring dichloroimidazolin-2-ylidene moieties as the probes.

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Scheme 1 Homodesmotic pincer/pseudopincer exchange reaction.

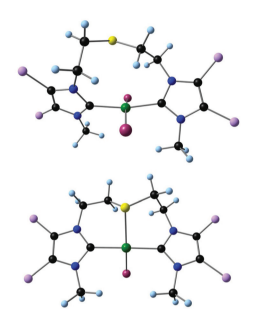


Fig. 1 Optimised pseudopincer and pincer geometries based on the dichloromidazolin-2-ylidene moiety.

This system was the most weakly donating ligand under scrutiny, and the respective complexes are known to prefer the pseudopincer form. A negative value for $\Delta G_{\rm R}$ will thus signify a higher preference for the pincer form in comparison to the reference system. Both the pincer and pseudopincer forms of all examined complexes were optimised at the B3LYP/cc-pVDZ level of theory, ^{14–16} and their respective Gibbs free energies were calculated at the same level of theory (see ESI† for computational details).

To avoid the influence of variations in steric bulk associated with changing the substituents at nitrogen and to limit the computational cost, NHC side chains were simplified to sterically unassuming methyl groups, and the effect of five vastly different NHC backbones (*cf.* Fig. 2) was examined, two of which have already been studied experimentally.¹³

The optimised structures are comparable to the reported molecular structures. All complexes have a distorted square planar coordination geometry of the palladium center (Fig. 1). In the pincer complexes, the metallacycles adopt a distorted boat conformation and the planes defined by the NHC rings are almost coplanar, while they showed a twist of 26–43° in the corresponding pseudopincer forms. From these optimized structures, $\Delta G_{\rm R}$ can be obtained.

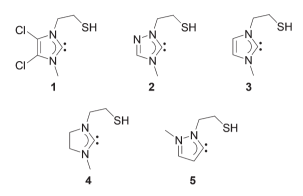


Fig. 2 Truncated ligands with different NHC moieties.

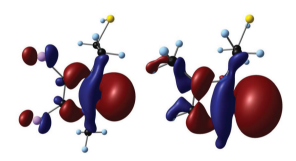


Fig. 3 σ-HOMO of NHCs 1 (HOMO–1) and 5 (HOMO).

No experimental donor strength values are available for the ligands in the calculated complexes. However, it has been shown that the Kohn–Sham eigenvalue of the σ -lone pair orbital $\varepsilon(\sigma$ -HOMO) correlates well with the donor strength. To limit the computational cost, orbital energies were calculated for truncated ligands at the B3LYP/auc-cc-pVTZ//B3LYP/cc-pVDZ level (Fig. 2). 14,16,18 In all cases, the highest orbital with σ -symmetry was found to be the lone pair at $C_{carbene}$ (Fig. 3).

Orbital energies for the σ -lone pair orbital were found to range from -6.56 to -5.52 eV, and this confirmed the expected increase in donor strength from 1 to 5 (Table 1). The preference for the pincer form, represented by $\Delta G_{\rm R}$ of the test reaction, increased in the same order, and Fig. 4 shows the strong correlation between donor strength and $\Delta G_{\rm R}$ ($R^2 = 0.9931$).

It is apparent that more strongly donating carbene moieties in the CSC ligand favour the pincer form. The cationic pincer complexes are better stabilised by stronger NHCs than the **Table 1** $\varepsilon(\sigma\text{-HOMO})$ of the truncated ligand and $\Delta G_{\mathbb{R}}$

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| NHC moiety | ε (σ-HOMO) [eV] | $\Delta G_{ m R}$ [kJ mol ⁻¹] |
|----------------------------|-----------------------------|---|
| Cl ₂ -imidazole | -6.56 | 0.0 |
| 1,2,4-Triazole | -6.52 | -5.4 |
| Imidazole | -6.12 | -26.5 |
| Imidazoline | -5.94 | -32.8 |
| Pyrazole | -5.52 | -57.0 |
| | | |

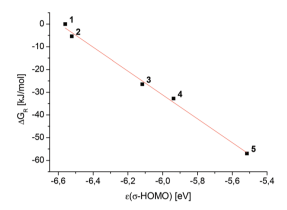


Fig. 4 Correlation of $\varepsilon(\sigma\text{-HOMO})$ [eV] and ΔG_R [kJ mol⁻¹].

neutral pseudopincer form, and this difference in stabilization increases with the donor strength. Additionally, entropy and solvation also favour the formation of the ionic pincer over the pseudopincer form. On the other hand, the neutral pseudopincer complexes are preferred if the NHC moieties transfer less electron density towards the metal atom due to the fact that bromide is a better ligand than the thioether.

In a more formalistic approach, the [PdBr(NHC)₂]⁺ fragment can be understood as a Lewis acid interacting with either the thioether moiety or a bromido ligand as a Lewis base. The internal charge redistribution in the Lewis acid upon charge transfer from the Lewis base is energetically more favourable with more weakly donating NHC moieties, which allows the coordination of the more strongly donating bromido ligand.¹⁹

In summary, the relationship between donor strength and pincer/pseudopincer isomerism has been unambiguously demonstrated to be in good agreement with experimental observations. Calculated or experimentally determined $\varepsilon(\sigma\text{-HOMO})$ values can thus serve as an indication of the hemilability of the central donor site in similar complexes, allowing for a more rational ligand design and the synthesis of tailor-made pincer complexes.

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Notes and references

- 1 The Chemistry of Pincer Compounds, ed. D. Morales-Morales and C. Jensen, Elsevier, Amsterdam; G. Rivera and R. H. Crabtree, N-Heterocyclic Carbenes in Synthesis, ed. S. P. Nolan, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim, ch. 9, p. 223; D. Pugh and A. A. Danopoulos, Coord. Chem. Rev., 2007, 251, 610; N. Selander and K. J. Szabó, Chem. Rev., 2011, 111, 2048; G. van Koten, J. Organomet. Chem., 2013, 730, 156; K. J. Szabó, Top. Organomet. Chem., 2013, 40, 203.
- 2 M. Albrecht and G. van Koten, *Angew. Chem., Int. Ed.*, 2001, 40, 3750; J. L. Hawk and S. L. Craig, *Top. Organomet. Chem.*, 2013, 40, 319.
- 3 G. van Koten, *Pure Appl. Chem.*, 1989, 61, 1681; I. Moreno, R. SanMartin, B. Inés, F. Churruca and E. Domínguez, *Inorg. Chim. Acta*, 2010, 363, 1903; D. Gelman and S. Musa, *ACS Catal.*, 2012, 2, 2456; D. M. Roddick, *Top. Organomet. Chem.*, 2013, 40, 49.
- 4 J. C. Jeffrey and T. B. Rauchfuss, *Inorg. Chem.*, 1979, 18, 2658; C. S. Slone, D. A. Weinberger and C. A. Mirkin, *Prog. Inorg. Chem.*, 1999, 48, 233; P. Braunstein and F. Naud, *Angew. Chem., Int. Ed.*, 2001, 40, 680; H. V. Huynh, C. H. Yeo and G. K. Tan, *Chem. Commun.*, 2006, 3833.
- 5 L. Canovese, F. Visentin, G. Chessa, P. Uguagliati, C. Santo, G. Bandoli and L. Maini, *Organometallics*, 2003, 22, 3230; M. Bassetti, A. Capone and M. Salamone, *Organometallics*, 2004, 23, 247; D. Vuzman, E. Poverenov, L. J. W. Shimon, Y. Diskin-Posner and D. Milstein, *Organometallics*, 2008, 27, 2627; G. R. Fulmer, W. Kaminsky, R. A. Kemp and K. I. Goldberg, *Organometallics*, 2011, 30, 1627; A. J. Ruddy, S. J. Mitton, R. McDonald and L. Turculet, *Chem. Commun.*, 2012, 48, 1159.
- 6 G. L. Moxham, H. E. Randell-Sly, S. K. Brayshaw, R. L. Woodward, A. S. Weller and M. C. Willis, *Angew. Chem., Int. Ed.*, 2006, 45, 7618; C. Zhao, M. C. Jennings and R. J. Puddephatt, *Dalton Trans.*, 2008, 1243; G. L. Moxham, H. Randell-Sly, S. K. Brayshaw, A. S. Weller and M. C. Willis, *Chem. – Eur. J.*, 2008, 14, 8383.
- 7 E. Poverenov, M. Gandelman, L. J. W. Shimon, H. Rozenberg, Y. Ben-David and D. Milstein, *Organometallics*, 2005, 24, 1082; R. Lindner, B. van den Bosch, M. Lutz, J. N. H. Reek and J. I. van der Vlugt, *Organometallics*, 2011, 30, 499.
- 8 W. A. Herrmann and C. Köcher, Angew. Chem., Int. Ed. Engl., 1997, 36, 2162; D. Bourissou, O. Guerret, F. P. Gabbaï and G. Bertrand, Chem. Rev., 2000, 100, 39;
 W. A. Herrmann, Angew. Chem., Int. Ed., 2002, 41, 1290;
 C. M. Crudden and D. P. Allen, Coord. Chem. Rev., 2004, 248, 2247; F. E. Hahn and M. C. Jahnke, Angew. Chem., Int. Ed., 2008, 47, 3122; L. Benhamou, E. Chardon, G. Lavigne, S. Bellemin-Laponnaz and V. César, Chem. Rev., 2011, 111, 2705.
- S. Díez-González and S. P. Nolan, Coord. Chem. Rev., 2007,
 251, 874; T. Dröge and F. Glorius, Angew. Chem., Int. Ed.,
 2010, 49, 6940.

10 D. J. Nielsen, K. J. Cavell, B. W. Skelton and A. H. White, Organometallics, 2006, 25, 4850; F. E. Hahn, M. C. Jahnke and T. Pape, Organometallics, 2006, 25, 5927; S. Warsink, P. Hauwert, M. A. Siegler, A. L. Spek and C. J. Elsevier, Appl. Organomet. Chem., 2009, 23, 225; J. DePasquale, M. Kumar, M. Zeller and E. T. Papish, Organometallics, 2013, 32, 966; H. V. Huynh and C.-S. Lee, Dalton Trans., 2013, 42, 6803; X. Liu and P. Braunstein, Inorg. Chem., 2013, 52, 7367.

Communication

- 11 M. Bierenstiel and E. D. Cross, *Coord. Chem. Rev.*, 2011, 255, 574; D. Yuan and H. V. Huynh, *Molecules*, 2012, 17, 2491.
- S. J. Roseblade, A. Ros, D. Monge, M. Alcarazo, E. Álvarez, J. M. Lassaletta and R. Fernández, *Organometallics*, 2007, 26, 2570; C. Fliedel, G. Schnee and P. Braunstein, *Dalton Trans.*, 2009, 2474; H. V. Huynh, C. H. Yeo and Y. X. Chew, *Organometallics*, 2010, 29, 1479; C. Fliedel and P. Braunstein, *Organometallics*, 2010, 29, 5614; L. Canovese, F. Visentin, C. Levi, C. Santo and V. Bertolasi, *J. Organomet. Chem.*, 2013, 732, 27; D. Krishnan, S. A. Pullarkat, M. Wu, Y. Li and P.-H. Leung, *Chem. Eur. J.*, 2013, 19, 5468.

- 13 H. V. Huynh, D. Yuan and Y. Han, *Dalton Trans.*, 2009, 7262; D. Yuan, H. Tang, L. Xiao and H. V. Huynh, *Dalton Trans.*, 2011, 40, 8788.
- 14 A. D. Becke, J. Chem. Phys., 1993, 98, 5648; A. D. Becke, Phys. Rev. A, 1988, 38, 3098; C. Lee, W. Yang and R. G. Parr, Phys. Rev. B: Condens. Matter, 1988, 37, 785.
- K. A. Peterson, D. Figgen, E. Goll, H. Stoll and M. Dolg, J. Chem. Phys., 2003, 119, 11113; K. A. Peterson and C. Puzzarini, Theor. Chem. Acc., 2005, 114, 283; K. A. Peterson, D. Figgen, M. Dolg and H. Stoll, J. Chem. Phys., 2007, 126, 124101.
- 16 T. H. Dunning Jr., J. Chem. Phys., 1989, 90, 1007; D. E. Woon and T. H. Dunning Jr., J. Chem. Phys., 1993, 98, 1358.
- 17 R. Tonner, G. Heydenrych and G. Frenking, *Chem. Asian J.*, 2007, 2, 1555; R. Tonner, G. Heydenrych and G. Frenking, *ChemPhysChem*, 2008, 9, 1474; J. C. Bernhammer, G. Frison and H. V. Huynh, *Chem. Eur. J.*, 2013, 19, 12892.
- 18 R. A. Kendall and T. H. Dunning Jr., J. Chem. Phys., 1992, 96, 6796.
- 19 S. E. Denmark and G. L. Beutner, Angew. Chem., Int. Ed., 2008, 47, 1560.