Supplementary data for the article:

Malenov, D. P.; Hall, M. B.; Zarić, S. D. Influence of Metal Ion on Chelate-Aryl Stacking Interactions. International Journal of Quantum Chemistry 2018, 118 (16). https://doi.org/10.1002/qua. 25629

AB(-)



Figure S1. Additional model systems for calculations of chelate-aryl potential energy surfaces


Figure S2. Potential energy curves for chelate-aryl stacking in model system B, calculated at M06-D3/def2-TZVP level. The calculations were performed on a series of geometries by changing the normal distances for a range of offset values; the curves present the energies of strongest interactions at all offset values.


Figure S3. Potential energy curves for chelate-aryl stacking in model system AB, calculated at M06-D3/def2-TZVP level. The calculations were performed on a series of geometries by changing the normal distances for a range of offset values; the curves present the energies of strongest interactions at all offset values.

$A B \min 1$


AB $\min 2$


Figure S4. The parallel displaced energy minima on potential energy curvea $B$ and $A B$ for chelate-aryl stacking

Table S1. M06-D3/def2-TZVP interaction energies and correlation energies calculated as the difference between MP2 and HF energies with cc-pVTZ basis set for the minima on $B$ and AB potential energy curves for chelate-aryl stacking interactions

|  | interaction energy <br> M06-D3/def2-TZVP[kcal/mol] |  |  | correlation energy <br> (MP2 -HF )/cc-pVTZ [kcal/mol] |  |  |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| geometry | Ni | Pd | Pt | Ni | Pd | Pt |
| B min | -4.85 | -4.90 | -4.83 | -7.52 | -8.31 | -8.46 |
| $\mathrm{AB} \min 1$ | -4.42 | -4.70 | -4.80 | -6.96 | -7.55 | -7.66 |
| $\mathrm{AB} \min 2$ | -5.70 | -5.58 | -5.02 | -7.63 | -7.52 | -8.66 |

Optimal normal distances for chelate-aryl stacking


Figure S5. Optimal normal distances for chelate-aryl stacking

