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# Thermodynamics of the condensation of the $Si_8O_{20}(SnMe_3)_8$ building block with M-X (M = B, Al, Si, P, Ti, V, Zn, Sn, Sb, X = Cl, Me, Et) precursors by **DFT-D3** calculations†

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Synthesis of porous metallosilicate materials from siloxane oligomers is a promising approach for constructing well-defined structures at a molecular level. Here, we use quantum chemistry DFT methods and demonstrate a computationally cheap method for screening potential precursors for synthesizing porous metallosilicates. We estimate the thermodynamic parameters of condensation reactions of the octakis(trimethyltin) spherosilicate Si<sub>8</sub>O<sub>20</sub>(SnMe<sub>3</sub>)<sub>8</sub> (CUBE) building block with metal chlorides and alkyl metals. These reactions represent the initial steps in the non-hydrolytic synthesis of metallosilicate gels containing potentially uniform single-site metal centers. Our main emphasis was on the spontaneity and irreversibility of the condensation and the computational screening of potential metal center sources. The precursors previously reported in successful condensations with CUBE, such as AlCl<sub>3</sub>, [AlCl<sub>4</sub>]<sup>-</sup>, Si-Cl compounds, PCl<sub>3</sub>, TiCl<sub>4</sub>, and VOCl<sub>3</sub>, are shown to undergo sufficiently irreversible reactions, as are the untested precursors BCl<sub>3</sub>, VCl<sub>4</sub>, and POCl<sub>3</sub>. Interestingly, AIMe<sub>3</sub> proves to be twice as exoergic as AICl<sub>3</sub>. The first chloride in Cp<sub>2</sub>TiCl<sub>2</sub> reacts readily, but the second may be partially reversible. SbCl<sub>3</sub> and Ph<sub>3</sub>SbCl<sub>2</sub> are borderline cases, and the reversibility of their condensations might pose a problem. SnCl<sub>4</sub> was found unsuitable as a precursor to stannosilicates. It should be possible to prepare zincosilicates from ZnEt<sub>2</sub>, but not from ZnCl<sub>2</sub>, as the affinity of Zn for Cl<sup>-</sup> is so high that in the presence of a source of Cl<sup>-</sup>, zincosilicate structures will dissolve back to CUBE and ZnCl<sub>2</sub>. The oxophilicity of the metal in the precursor is the main factor in the driving force for the condensation with CUBE. Alkyl metals and lighter elements are more prone to the reaction than the corresponding metal chlorides and heavier analogs. The propensity of [SnMe<sub>3</sub>]<sup>+</sup> to bind to Cl⁻ in preference to CUBE has a supporting effect. At low temperatures, the condensation is slightly disfavored, while at the experimentally used temperature of 100 °C, this process contributes over 20 kJ mol<sup>-1</sup> of the additional driving force and helps to complete the condensation. The reliability of B3LYP-D3 and PBE0-D3, together with the CBS extrapolation scheme, is also evaluated in calculations.

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## Introduction

Porous metallosilicates represent a diverse class of chemically simple yet powerful heterogeneous catalysts - partially hollow extended networks generally composed of [SiO<sub>4</sub>] and [MO<sub>x</sub>] (M = metal) polyhedra. The potentially omnidirectional bonding between the [SiO<sub>4</sub>] tetrahedra provides a chemically and thermally stable backbone that supports porosity and can withstand harsh reaction environments. The silicate networks can accommodate most chemical elements that can form

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stable bonds to oxygen, with the general trend of enhancing the electrophilic character of the resulting metal center, attributable to the negative inductive effect of the oxygen. The applicability in catalysis and viable synthesis routes depends entirely on the target metal, its oxidation state, its chemical environment, and the required porosity of the final material.

A spherosilicate consisting of a double 4-ring core (D4R, Si<sub>8</sub>O<sub>20</sub>) with every vertex functionalized with a suitable reactive group has attracted significant attention as a molecular building block for constructing a whole array of nanoporous materials.2-5 By direct crosslinking of these cubic units into 3-dimensional infinite networks, microporous materials have been obtained, while mesoporous systems can be prepared with templates or long-alkyl chain substituents.

Highly promising possibilities arise in the controlled synthesis of single-site catalysts from the well-behaved

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condensation chemistry of the octakis(trimethyltin) spherosilicate building block Si<sub>8</sub>O<sub>20</sub>(SnMe<sub>3</sub>)<sub>8</sub> (CUBE), first reported by Feher and Weller. 6,7 The trimethyltin functionalization provides high solubility in non-polar solvents and supports metathesis reactions with covalent dand p-block element (M) chlorides and bromides as well as organometallic compounds, forming Me<sub>3</sub>SnCl, Me<sub>3</sub>SnBr, or Me<sub>3</sub>SnR as byproducts, respectively. Multiple silicate Si-O-M linkages are thus created by non-hydrolytic condensation, leading to the formation of gels. The metathesis is analogous to common proton-exchange reactions. However, due to the relatively low reactivity and weak coordinating nature of the [Me<sub>3</sub>SnOSi] moiety, it is highly compatible with other functional groups and ligands.

Increasingly more complex non-hydrolytic sol-gel schemes were reported by Ghosh et al., 8 Clark et al., 9 Lee et al., 10 Clark and Barnes, 11 and Barnes et al., 12 utilizing TiCl4, Cp2TiCl2, ZrCl4, Cp2ZrCl2, VOCl3, WOCl4, and AlCl3 as metal sources and SiCl<sub>4</sub>, HSiCl<sub>3</sub>, MeSiCl<sub>3</sub>, and Me<sub>2</sub>SiCl<sub>2</sub> as inert cross-linkers. This line of research has culminated in a flexible two-step, one-pot procedure (Fig. 1). The CUBE building block is first cross-linked by a limited amount of the metal precursor, providing ideal conditions for M-X group condensation, homogeneous incorporation of the metal sites, and structural relaxation. A true solution of oligomeric species is formed, which are then "knitted" together by further cross-linking with silyl chlorides to produce porous silicate matrices. The solvent and all reaction byproducts are then removed under vacuum to obtain pure materials. Subsequently, Styskalik et al. 13 used the approach to prepare a series of porous Lewis acidic aluminosilicate matrices containing  $[R]^+$   $[AlO_4]^ (R^+$  = lutidinium, NBu<sub>4</sub><sup>+</sup>) and [L-AlO<sub>3</sub>] (L = pyridine, THF) sites starting from AlCl<sub>3</sub> and its coordination compounds in THF. The method was improved by using longer hybrid silyl chlorides (ClMe<sub>2</sub>Si(CH<sub>2</sub>)<sub>n</sub>SiMe<sub>2</sub>Cl, where n = 1-3) for the final cross-linking. This resulted in increased average pore sizes and better diffusion properties, leading to superior activities in Lewis acid catalysis.

Most recently, we used the first step of this procedure to prepare a series of Lewis acidic aluminosilicate oligomers starting from L-AlCl<sub>3</sub> and L-AlMe<sub>3</sub> compounds. We characterized the structure of the aluminosilicate sites by <sup>27</sup>Al MQ/MAS NMR techniques and studied the conditions leading to single-site species.<sup>14</sup>

Since the first applications of CUBE to the synthesis of metallosilicate catalysts, it was presumed that the condensation reactions used to connect the metallosilicate networks are irreversible for all reported precursors. The validity of this assumption has far-reaching implications. The method of sequential additions assumes that the connections and structures formed in one cross-linking step remain unchanged during the next steps. Each subsequent crosslinking step only adds further connectivity through the residual -SnMe3 groups left at the CUBE vertices in the structures. oligomeric Under such conditions, rearrangement or metal migration can occur, and the structures formed should be amorphous and purely statistical, allowing for the use of mean-field concepts, such as average connectivity and Vegard's law. Irreversibility should also result in better decoupling between the chemistries of the precursors used in the sequential crosslinking steps, and it would imply that the order of the steps strictly matters. This work aims to answer the question of reversibility for a broader spectrum of interesting metal site precursors using quantum chemistry DFT computational methods.

## Computational methods

The input geometries were drawn and preoptimized by molecular mechanics in Avogadro 1.2.0 software. 15 Further calculations were performed using the GAMESS 2019(R2) suite.16,17 The structures were optimized in delocalized internal coordinates<sup>18</sup> by the B3LYP<sup>19,20</sup> and PBE0<sup>21,22</sup> exchange-correlation functionals (DFT grid: 96 radial/302 angular points) with the correlation-consistent valence triple- $\zeta$  cc-pVTZ (default)<sup>23–25</sup> and the augmented small-core pseudopotential-based aug-cc-pVTZ-PP (Sb, Sn) basis sets<sup>26,27</sup> in vacuum. Grimme's empirical D3 dispersion correction with Becke-Johnson damping<sup>28-30</sup> was used in all calculations. The following convergence criteria were used: 10<sup>-5</sup> for the largest absolute change in the density matrix and 5·10<sup>-5</sup> Hartree Bohr<sup>-1</sup> for the largest component of the geometry gradient. Vibrational analyses with harmonic oscillator approximation were used to confirm the energyminimum character of the obtained stationary point geometries and to calculate the thermodynamic corrections to the electronic energy at 0, 273.15, 298.15, and 373.15 K (ideal gas at 101.325 kPa and rigid rotor approximations).

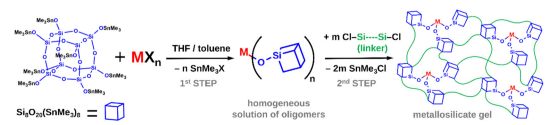


Fig. 1 General reaction scheme for the two-step one-pot synthesis of metallosilicate gels from CUBE

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The structures were first optimized by B3LYP-D3, and the resulting geometries were used as inputs for the optimizations by PBE0-D3 to minimize the conformational variations between the two optimized geometries. Additional single-point energy calculations using the double- and quadruple- $\zeta$  cc-pVxZ + aug-cc-pVxZ-PP (Sn) (x = D, Q) basis sets were conducted with the corresponding method. The exponential complete basis set limit (CBS) extrapolation scheme developed by Halkier et al.31 was then used to obtain refined electronic energies  $(E_{CBS})$  with decreased effects of the basis set superposition error (BSSE) and the fitting parameters B and  $\alpha$  according to eqn (1)-(3). The choice of temperatures was aimed to provide thermodynamic data ( $\Delta H$ ,  $\Delta S$ , and  $\Delta G$ ) at two standard temperatures as well as a third set at a reasonable high-temperature limit (100 °C) for the practical use of the condensation reaction, given the thermal stability of CUBE (decomposes above ~120 °C). All the produced data is available in the ESI† (Tables S1-S46 for thermodynamic data by reaction; Tables S47 and S48 for the

electronic energy and thermal corrections by structure; Tables

S49-S57 for equilibrium geometries by structure).

$$E_{\rm CBS} = \frac{E_{\rm DZ} E_{\rm QZ} - E_{\rm TZ}^2}{E_{\rm DZ} + E_{\rm OZ} - 2E_{\rm TZ}} \tag{1}$$

$$B = \frac{(E_{\rm DZ} - E_{\rm TZ})^4}{(E_{\rm DZ} + E_{\rm QZ} - 2E_{\rm TZ})(E_{\rm TZ} - E_{\rm QZ})^2}$$
(2)

$$\alpha = \ln \left( \frac{E_{\rm DZ} - E_{\rm TZ}}{E_{\rm TZ} - E_{\rm OZ}} \right) \tag{3}$$

## Results and discussion

#### **Reaction spontaneity**

The reversibility of the condensation reaction of the  $Si_8O_{20}(SnMe_3)_8$  building block (CUBE) with precursors of interest was characterized through the calculation of the Gibbs energy change for model reactions with truncated molecular representations of the polymeric products (metallosilicate networks). A single reactive group (corner) of the CUBE was modeled by either  $Me_3SnOSiH_3$  (model A, reactions (1A)–(26A)) for all condensation steps with  $MX_n$  precursors (x = 0, ..., n-1) and by  $Me_3SnOSi(OSiH_3)_3$  (model B, reactions (1B)–(20B)) for the condensation of only the first X group (X = Cl, Me, Et).

This aimed to provide both precise thermodynamic data for the initial condensation reaction (the first substitution on the metal) and to get a sense of the level of precursor deactivation with subsequent condensation steps all the way to the completely condensed metallosilicate sites. Of course, this must be viewed as an approximation that reflects mainly the electronic effects on the central atom (metal) and excludes any steric repulsion between multiple connected CUBEs.

The driving force for the condensation reaction may be viewed as a sum of two contributions (positive or negative) – the exchange of the reactive group M–X for the silicate moiety M–O–Si on the metal (different for each precursor) and the exchange of the silicate moiety Sn–O–Si for Sn–X on the trimethyltin group (intrinsic driving force, common to all precursors). Reactions (26A) and (20B) model the latter process for precursors with Cl<sup>-</sup> as a leaving group to assess how much of the driving force (and more importantly, with what sign) comes intrinsically from the chemistry of the trimethyltin group.

As there is no sharp distinction between reversible and irreversible reactions, in the following discussion, as a rule of thumb, reactions with  $\Delta G \leq -30$  kJ mol<sup>-1</sup> shall be considered sufficiently irreversible to prevent any measurable extent of network equilibration over the reaction times of interest ( $\leq 1$  week).

#### Reactions A.

py-BCl<sub>3-x</sub>(OSiH<sub>3</sub>)<sub>x</sub> + Me<sub>3</sub>SnOSiH<sub>3</sub> 
$$\rightarrow$$
 py-BCl<sub>2-x</sub>(OSiH<sub>3</sub>)<sub>1+x</sub>  
+ SnMe<sub>3</sub>Cl (where:  $x = 0$ -2) (1A)

py-AlMe<sub>3-x</sub>(OSiH<sub>3</sub>)<sub>x</sub> + Me<sub>3</sub>SnOSiH<sub>3</sub> 
$$\rightarrow$$
 py-AlMe<sub>2-x</sub>(OSiH<sub>3</sub>)<sub>1+x</sub>  
+ SnMe<sub>4</sub> (where:  $x = 0-2$ ) (2A)

py-AlMe<sub>3-x</sub>(OSiH<sub>3</sub>)<sub>x</sub>(THF) + Me<sub>3</sub>SnOSiH<sub>3</sub> 
$$\rightarrow$$
 py-AlMe<sub>2-x</sub>(OSiH<sub>3</sub>)<sub>1+x</sub>(THF)  
+ SnMe<sub>4</sub> (where:  $x = 0-2$ ) (3A)

py-AlCl<sub>3-x</sub>(OSiH<sub>3</sub>)<sub>x</sub> + Me<sub>3</sub>SnOSiH<sub>3</sub> 
$$\rightarrow$$
 py-AlCl<sub>2-x</sub>(OSiH<sub>3</sub>)<sub>1+x</sub>  
+ SnMe<sub>3</sub>Cl (where:  $x = 0-2$ ) (4A)

py-AlCl<sub>3-x</sub>(OSiH<sub>3</sub>)<sub>x</sub>(THF) + Me<sub>3</sub>SnOSiH<sub>3</sub> 
$$\rightarrow$$
 py-AlCl<sub>2-x</sub>(OSiH<sub>3</sub>)<sub>1+x</sub>(THF)  
+ SnMe<sub>3</sub>Cl (where:  $x = 0-2$ ) (5A)

$$[AlCl4-x(OSiH3)x]- + Me3SnOSiH3 \rightarrow [AlCl3-x(OSiH3)1+x]- + SnMe3Cl (where:  $x = 0-3$ ) (6A)$$

$$SiCl_{4-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow SiCl_{3-x}(OSiH_3)_{1+x} + SnMe_3Cl \text{ (where: } x = 0-3)$$
 (7A)

$$Me_3SiCl + Me_3SnOSiH_3 \rightarrow Me_3SiOSiH_3 + SnMe_3Cl$$
 (8A)

$$PhMe_2SiCl + Me_3SnOSiH_3 \rightarrow PhMe_2SiOSiH_3 + SnMe_3Cl$$
 (9A)

$$PCl_{3-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow PCl_{2-x}(OSiH_3)_{1+x} + SnMe_3Cl \text{ (where: } x = 0-2)$$
 (10A)

$$POCl_{3-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow POCl_{2-x}(OSiH_3)_{1+x} + SnMe_3Cl \text{ (where: } x = 0-2\text{)}$$
 (11A)

$$TiCl_{4-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow TiCl_{3-x}(OSiH_3)_{1+x} + SnMe_3Cl \text{ (where: } x = 0-3\text{)}$$
 (12A)

$$Cp_2TiCl_{2-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow Cp_2TiCl_{1-x}(OSiH_3)_{1+x} + SnMe_3Cl \text{ (where: } x = 0-1\text{)}$$
 (13A)

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(16A)

$$VCl_{4-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow VCl_{3-x}(OSiH_3)_{1+x}$$

$$+ SnMe_3Cl \text{ (where: } x = 0-3)$$

$$VOCl_{3-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow VOCl_{2-x}(OSiH_3)_{1+x}$$

+ SnMe<sub>3</sub>Cl (where: 
$$x = 0-2$$
) (15A)  

$$ZnEt_{2-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow ZnEt_{1-x}(OSiH_3)_{1+x}$$

$$ZnEt_{2-x}(OSiH_3)_x(THF) + Me_3SnOSiH_3 \rightarrow ZnEt_{1-x}(OSiH_3)_{1+x}(THF) + SnMe_3Et \text{ (where: } x = 0-1\text{)}$$
 (17A)

+ SnMe<sub>3</sub>Et (where: x = 0-1)

$$ZnEt_{2-x}(OSiH_3)_x(THF)_2 + Me_3SnOSiH_3 \rightarrow ZnEt_{1-x}(OSiH_3)_{1+x}(THF)_2 + SnMe_3Et \text{ (where: } x = 0-1\text{)}$$
 (18A)

$$ZnCl_{2-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow ZnCl_{1-x}(OSiH_3)_{1+x} + SnMe_3Cl \text{ (where: } x = 0-1\text{)}$$
 (19A)

$$ZnCl_{2-x}(OSiH_3)_x(THF) + Me_3SnOSiH_3 \rightarrow ZnCl_{1-x}(OSiH_3)_{1+x}(THF) + SnMe_3Cl \text{ (where: } x = 0-1)$$
 (20A)

$$ZnCl_{2-x}(OSiH_3)_x(THF)_2 + Me_3SnOSiH_3 \rightarrow ZnCl_{1-x}(OSiH_3)_{1+x}(THF)_2 + SnMe_3Cl \text{ (where: } x = 0-1\text{)}$$
 (21A)

$$[ZnCl_{3-x}(OSiH_3)_x]^- + Me_3SnOSiH_3 \rightarrow [ZnCl_{2-x}(OSiH_3)_{1+x}]^- + SnMe_3Cl \text{ (where: } x = 0-1\text{)}$$
 (22A)

$$SnCl_{4-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow SnCl_{3-x}(OSiH_3)_{1+x} + SnMe_3Cl \text{ (where: } x = 0-3)$$
 (23A)

$$SbCl_{3-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow SbCl_{2-x}(OSiH_3)_{1+x} + SnMe_3Cl \text{ (where: } x = 0-2\text{)}$$
 (24A)

$$Ph_3SbCl_{2-x}(OSiH_3)_x + Me_3SnOSiH_3 \rightarrow Ph_3SbCl_{1-x}(OSiH_3)_{1+x} + SnMe_3Cl \text{ (where: } x = 0-1)$$
 (25A)

$$Cl^- + Me_3SnOSiH_3 \rightarrow H_3SiO^- + SnMe_3Cl$$
 (26A)

## Reactions B.

$$\begin{array}{c} py\text{-BCl}_3 + Me_3SnOSi(OSiH_3)_3 \rightarrow py\text{-BCl}_2OSi(OSiH_3)_3 \\ + SnMe_3Cl \end{array} \tag{1B}$$

py-AlMe<sub>3</sub> + Me<sub>3</sub>SnOSi(OSiH<sub>3</sub>)<sub>3</sub> 
$$\rightarrow$$
 py-AlMe<sub>2</sub>OSi(OSiH<sub>3</sub>)<sub>3</sub>  
+ SnMe<sub>4</sub> (2B)

$$\begin{aligned} & py\text{-AlCl}_3 + Me_3SnOSi(OSiH_3)_3 \rightarrow py\text{-AlCl}_2OSi(OSiH_3)_3 \\ & + SnMe_3Cl \end{aligned} \tag{3B}$$

$$\begin{aligned} & \left[ AlCl_4 \right]^- + Me_3SnOSi(OSiH_3)_3 \rightarrow \left[ AlCl_3OSi(OSiH_3)_3 \right]^- \\ & + SnMe_3Cl \end{aligned} \tag{4B}$$

 $SiCl_4 + Me_3SnOSi(OSiH_3)_3 \rightarrow SiCl_3OSi(OSiH_3)_3 + SnMe_3Cl(5B)$ 

$$Me_3SiCl + Me_3SnOSi(OSiH_3)_3 \rightarrow Me_3SiOSi(OSiH_3)_3 + SnMe_3Cl$$
 (6B)

$$PhMe_{2}SiCl + Me_{3}SnOSi(OSiH_{3})_{3} \rightarrow PhMe_{2}SiOSi(OSiH_{3})_{3} + SnMe_{3}Cl$$
(7B)

$$PCl_3 + Me_3SnOSi(OSiH_3)_3 \rightarrow PCl_2OSi(OSiH_3)_3 + SnMe_3Cl$$
 (8B)

$$\begin{aligned} \text{POCl}_3 + \text{Me}_3 \text{SnOSi}(\text{OSiH}_3)_3 &\rightarrow \text{POCl}_2 \text{OSi}(\text{OSiH}_3)_3 \\ + \text{SnMe}_3 \text{Cl} \end{aligned} \tag{9B}$$

$$TiCl_4 + Me_3SnOSi(OSiH_3)_3 \rightarrow TiCl_3OSi(OSiH_3)_3 + SnMe_3Cl$$
 (10B)

$$\begin{aligned} \text{Cp}_2\text{TiCl}_2 + \text{Me}_3\text{SnOSi}(\text{OSiH}_3)_3 &\rightarrow \text{Cp}_2\text{TiClOSi}(\text{OSiH}_3)_3 \\ &+ \text{SnMe}_3\text{Cl} \end{aligned} \tag{11B}$$

$$VCl_4 + Me_3SnOSi(OSiH_3)_3 \rightarrow VCl_3OSi(OSiH_3)_3 + SnMe_3Cl(12B)$$

$$VOCl_3 + Me_3SnOSi(OSiH_3)_3 \rightarrow VOCl_2OSi(OSiH_3)_3 + SnMe_3Cl$$
 (13B)

$$ZnEt_2(THF)_2 + Me_3SnOSi(OSiH_3)_3 \rightarrow ZnEtOSi(OSiH_3)_3(THF)_2 + SnMe_3Et$$
 (14B)

$$ZnCl_2(THF)_2 + Me_3SnOSi(OSiH_3)_3 \rightarrow ZnClOSi(OSiH_3)_3(THF)_2 + SnMe_3Cl$$
 (15B)

$$\begin{aligned} & \left[ ZnCl_{3} \right]^{-} + Me_{3}SnOSi(OSiH_{3})_{3} \rightarrow \left[ ZnCl_{2}OSi(OSiH_{3})_{3} \right]^{-} \\ & + SnMe_{3}Cl \end{aligned} \tag{16B}$$

$$SnCl_4 + Me_3SnOSi(OSiH_3)_3 \rightarrow SnCl_3OSi(OSiH_3)_3 + SnMe_3Cl$$
 (17B)

$$SbCl_3 + Me_3SnOSi(OSiH_3)_3 \rightarrow SbCl_2OSi(OSiH_3)_3 + SnMe_3Cl$$
 (18B)

$$Ph_3SbCl_2 + Me_3SnOSi(OSiH_3)_3 \rightarrow Ph_3SbClOSi(OSiH_3)_3 + SnMe_3Cl$$
 (19B)

$$Cl^- + Me_3SnOSi(OSiH_3)_3 \rightarrow (H_3SiO)_3SiO^- + SnMe_3Cl$$
 (20B)

Fig. 2 displays the values of the Gibbs energy change for the first step of the condensation of all the precursors of interest at 298.15 K as calculated by B3LYP-D3 and PBE0-D3 using both molecular models A and B. Fig. 3 and 4 show the comparisons of all the condensation steps (all steps represented by model A and the first step represented by model B) as calculated by the two DFT functionals, respectively. The results show that the reaction is sufficiently spontaneous and irreversible for most precursors, especially those reported to be utilized experimentally, thereby generally supporting the conclusions of Clark, Lee, Barnes, and Styskalik, 10-13 and the validity of the method of sequential additions.

Fig. S1 and S2 (ESI $\dagger$ ) show the corresponding values of the Gibbs energy change for the first step of the condensation at 0 and 373.15 K, respectively. The negligible differences between the two figures indicate that the effects of

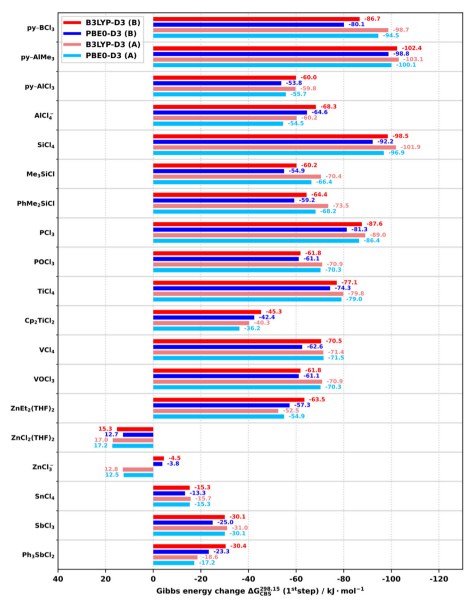


Fig. 2 Gibbs energy change of the first step of the condensation of studied precursors with models A (light) and B (dark) of CUBE at 298.15 K, as calculated by B3LYP-D3/CBS (red) and PBE0-D3/CBS (blue).

temperature do not affect the interpretation of the results in any significant way, and, therefore, the following discussion will deal primarily with the Gibbs energy change at 298.15 K ( $\Delta G^{298.15}$ ).

 $PCl_3$  was the very first precursor reported in reactions with CUBE by Weller and Feher, and their experiments indicated a fast, highly exothermic reaction (10A)/(8B). Our calculations concur with the predicted  $\Delta G^{298.15} < -80$  kJ mol<sup>-1</sup> for the first step and approximately equal to -70 kJ mol<sup>-1</sup> for the third step (electronic effects-only approximation), making the condensation perfectly irreversible. Although the reported products were only partially condensed and inhomogeneous, this was likely only due to the crude reaction conditions ( $PCl_3$ -rich stoichiometry, poor solvent, and fast mixing at room temperature), and if appropriately conducted,  $PCl_3$ 

should yield silicophosphate sites with the general formula  $[P(OSi\equiv)_3].$  Porous materials containing such sites would be interesting as strong Lewis bases and redox catalysts, with the added benefit of utilizing a light, abundant, non-toxic p-block element. Analogous to  $PCl_3$ , silicophosphate sites with the general formula  $[PO(OSi\equiv)_3]$  should be available through the condensation of  $POCl_3$  (11A)/(9B). Calculations predict irreversible condensation  $(\Delta G^{298.15} < -60 \text{ kJ mol}^{-1})$  with very little change in the subsequent condensation steps.

The condensation reactions of CUBE with VCl<sub>4</sub> (14A)/(12B) and especially VOCl<sub>3</sub> (15A)/(13B) were reported initially by Ghosh *et al.*<sup>8</sup> and later extensively studied by Lee *et al.*<sup>10</sup> In the first condensation step, both precursors exhibit thermodynamic characteristics nearly identical to POCl<sub>3</sub>, however, compared to VOCl<sub>3</sub>, VCl<sub>4</sub> is significantly deactivated

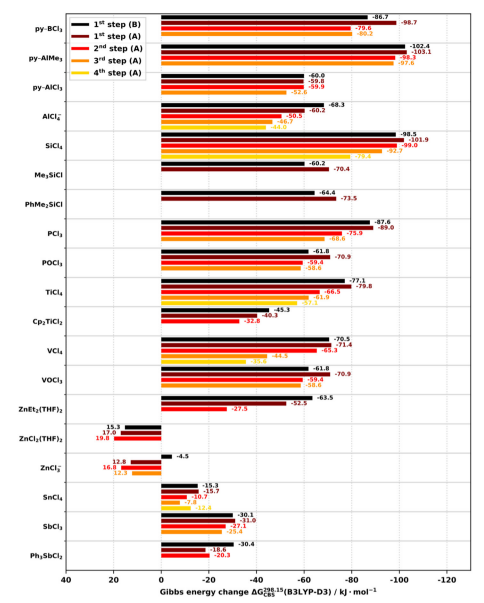


Fig. 3 Gibbs energy change of the condensation of selected precursors with models A and B of CUBE at 298.15 K, as calculated by B3LYP-D3/ CBS.

as the condensation progresses, decreasing  $\Delta G^{298.15}$  to mere -35.6/-33.8 kJ mol<sup>-1</sup> by B3LYP-D3 and PBE0-D3, respectively.

The reaction of TiCl<sub>4</sub> (12A)/(10B) was reported (along with ZrCl<sub>4</sub> and SiCl<sub>4</sub>) by Barnes et al., 12 who successfully used multistep template-free procedures to obtain micromesoporous titano- and zirconosilicates. Calculations showed that the condensation of TiCl<sub>4</sub> is highly exoergic with  $\Delta G^{298.15}$  ranging from nearly -80 kJ mol<sup>-1</sup> for the first step to -57 kJ mol<sup>-1</sup> for the last step, making it irreversible. Reactions of CUBE with Cp2TiCl2 were reported by Clark et al.,9 who obtained a molecular substitution product of CUBE Si<sub>8</sub>O<sub>20</sub>(Cp<sub>2</sub>TiCl)<sub>8</sub> in high yield. In harmony with this experimental observation, our calculations show that the condensation of this precursor is nearly half as energetic as TiCl<sub>4</sub>, with  $\Delta G^{298.15}$  ranging from over -40 kJ mol<sup>-1</sup> to

approximately -32 kJ mol<sup>-1</sup>. Since the steric repulsion between the cyclopentadienyl rings and the CUBE(s) is completely neglected by model A (and the results given by model B for the first condensation step do not significantly differ), this deactivation compared to TiCl4 is attributed in large part to electronic effects - the electron-donating ability of the Cp rings. The Cp rings will impose additional steric repulsion and, especially in the case of the second condensation, a large additional penalty can be expected, explaining the apparently high activation barrier of the reaction and the tendency for stepwise substitution.

SiCl<sub>4</sub> (7A)/(5B), Me<sub>3</sub>SiCl (8A)/(6B), and PhMe<sub>2</sub>SiCl (9A)/(7B) are relevant as inert cross-linking and capping agents. Alternatively, a precursor with the general formula RSiCl<sub>3</sub>, where R is a functional moiety, could be used for embedding.

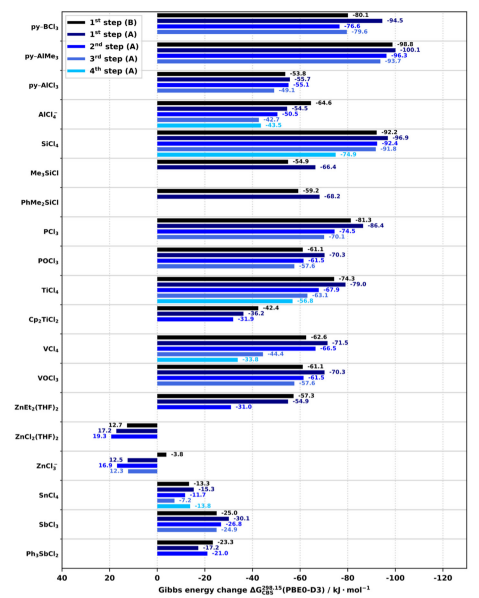


Fig. 4 Gibbs energy change of the condensation of selected precursors with models A and B of CUBE at 298.15 K, as calculated by PBE0-D3/CBS.

Clark and Barnes<sup>11</sup> studied various cross-linking procedures by SiCl<sub>4</sub>, HSiCl<sub>3</sub>, and Me<sub>2</sub>SiCl<sub>2</sub>. Styskalik et al. 13 used ditopic linkers with general formulas  $ClMe_2Si(CH_2)_nSiMe_2Cl$  (n = 1-3) as well as the very similar Me<sub>3</sub>SiCl directly as a capping agent to remove the residual -SnMe3 groups left in the prepared metallosilicate matrices. The results show that SiCl4 is one of the most exoergic precursors with estimated  $\Delta G^{298.15}$  ranging from -102 to -75 kJ mol<sup>-1</sup> while the condensation of Me<sub>3</sub>SiCl is moderated by the positive inductive effect of the methyl substituents to -60.2/-54.9 kJ mol<sup>-1</sup> by B3LYP-D3 and PBE0-D3, respectively. Surprisingly, the condensation of PhMe<sub>2</sub>SiCl is predicted to be even slightly more energetic than that of Me<sub>3</sub>SiCl, despite our expectations that a positive mesomeric effect of the phenyl substituent would decrease the reactivity even further. The reactivity of compounds such as HSiCl3 and Me<sub>2</sub>SiCl<sub>2</sub> is expected to range somewhere between the

extremes of SiCl<sub>4</sub> and Me<sub>3</sub>SiCl. In conclusion, there is no doubt about the irreversibility of the condensation reaction for any Si-Cl precursors.

The condensation of CUBE with py-AlCl<sub>3</sub> (4A)/(3B) and [AlCl<sub>4</sub>] (6A)/(4B) was described by Styskalik et al. 13 with reports of full condensation and undisturbed coordination environment around Al even after subsequent cross-linking by the Si-Cl linkers. We have recently employed both precursors, along with py-AlMe<sub>3</sub> (2A)/(2B) experimentally. 14 The calculations show that while condensation for all three precursors is certainly irreversible, there are vast differences between py-AlCl<sub>3</sub> and py-AlMe<sub>3</sub>. While the former is a moderately exoergic precursor (-70 to -50 kJ mol<sup>-1</sup>), the latter is the most energetic in the whole series (around -100 kJ mol<sup>-1</sup>). This agrees with our experimental observations of the relative reactivity of the two precursors.<sup>14</sup> In our experience,

py–AlMe $_3$  is much more sensitive to concentration, proper cooling (–80 °C), and slow addition speed. Mismanagement of these conditions leads to the formation of particulate precipitates. This is attributed to the emergence of molecular hotspots – a behavior typical of highly exothermic, low activation barrier reactions. [AlCl $_4$ ] $^-$  is comparable to py–AlCl $_3$  with a greater degree of deactivation in the subsequent condensation steps.

Given the rather hard character of the Al-based Lewis acids, precursors based on B and Zn were screened as softer alternatives. Borosilicate Lewis acids differ from aluminosilicate counterparts by the tendency of the [BO<sub>3</sub>] moiety to assume planar geometry with  $\pi$ -electron delocalization across the whole moiety. This results in much weaker, more reversible coordination to ligands and the fact that, in contrast to [Al(OSi≡)<sub>3</sub>], [B(OSi≡)<sub>3</sub>] is a well-defined and stable species.<sup>32</sup> To facilitate maximally relevant comparison, py-BCl<sub>3</sub> (1A)/(1B) was chosen as the starting precursor, and the results show that its condensation with CUBE is highly exoergic (-99 to -77 kJ mol<sup>-1</sup>) with very little deactivation in subsequent steps. Therefore, B-Cl compounds, in general, are predicted to be perspective, irreversibly reacting precursors for the synthesis of CUBE-based borosilicate materials. The possibility of maintaining an undisturbed coordination environment throughout the condensation is, however, questionable given the aforementioned general lability of neutral ligands on [BO<sub>3</sub>]

ZnCl2 represents another alternative and it is often used in synthetic organic chemistry as a milder analog to AlCl3 in reactions, such as the Friedel-Crafts methods.<sup>33</sup> In analogy to Al, we also considered the use of the commercially available ZnEt<sub>2</sub>. Due to the weaker coordination to Zn, no special ligands were assumed, but rather the precursors were studied primarily in the state of saturation (coordination number 4) by THF - the aprotic solvent of choice to dissolve both ZnCl2 and CUBE. The results show that the condensation of ZnEt<sub>2</sub>(THF)<sub>2</sub> (18A)/(14B) is moderately energetic (-64 to -28 kJ mol<sup>-1</sup>) and thus quite irreversible, albeit with significant deactivation between the first and the second steps. In contrast, the condensation of ZnCl<sub>2</sub>(THF)<sub>2</sub> (21A)/(15B) is a clear example of the limits of the CUBE-based synthetic approach as the predicted  $\Delta G^{298.15}$  is positive (+13 to +20 kJ mol<sup>-1</sup>). Therefore, the equilibrium is shifted towards CUBE + ZnCl2, and the condensation is not spontaneous. Moreover, this also implies, that even if [Zn(OSi≡)<sub>2</sub>] sites were generated from ZnEt<sub>2</sub>, the method of sequential additions could not be used because as soon as any SnMe<sub>3</sub>Cl was generated by subsequent steps, such as the inert cross-linking by Si-Cl compounds, the equilibrium of the CUBE/ZnCl2 system would lead to the dissolution of the Zn-O-Si bonds and leaching out ZnCl<sub>2</sub>. The presence of Cl<sup>-</sup> donors, such as SnMe<sub>3</sub>Cl, may also give rise to the anionic species [ZnCl<sub>3</sub>] (22A)/(16B), which, according to the results, should behave in the same manner as ZnCl2, although model B predicts the condensation of the first Zn-Cl to be slightly spontaneous.

SbCl<sub>3</sub> (24A)/(18B) was chosen as a softer isoelectronic analog to the Lewis basic PCl<sub>3</sub>. The results show that its condensation thermodynamics are right on the established border of absolute irreversibility (-30 to -25 kJ mol<sup>-1</sup>).

Ph<sub>3</sub>SbCl<sub>2</sub> (25A)/(19B) was selected as an interesting antimony(v) precursor for computational screening because the obvious candidate, SbCl5, is unsuitable for several reasons. First, our recent work showed that fitting four CUBEs around an Al3+ center is strongly disfavored.14 It is doubtful that fitting five CUBEs around the 11% larger Sb5+ is possible. Second, SbCl<sub>5</sub> is a strong oxidizing agent, and third, the extreme stability of the [SbCl<sub>6</sub>] species implies a side reaction between unconsumed SbCl<sub>5</sub> and the evolving SnMe<sub>3</sub>Cl byproduct, producing ionic compounds, such as [SnMe<sub>3</sub>][SbCl<sub>6</sub>]. In contrast, Ph<sub>3</sub>SbCl<sub>2</sub> was expected to be much less energetic, and there is no doubt that two CUBEs can fit around Sb at 180° apart with the [Ph<sub>3</sub>Sb] moiety in an equatorial plane between them. Our calculations showed that this precursor is even slightly less exoergic than SbCl<sub>3</sub>, with  $\Delta G^{298.15}$  of at most -30 and as little as -17 kJ mol<sup>-1</sup>, making its irreversibility questionable.

Finally, Clark and Barnes<sup>11</sup> mentioned using  $SnCl_4$  to obtain stannosilicates (23A)/(17B). The possibility for an irreversible exchange of  $Cl^-$  between different tin(rv) species appeared highly questionable to us. Our calculations show that although the condensation reaction is still spontaneous in this case, it is hardly irreversible with  $\Delta G^{298.15}$  of at most -16 and as little as -7 kJ  $mol^{-1}$ . A facile ligand exchange with  $SnMe_3Cl$ , leading to a mixture of  $SnMe_2Cl_2$  and  $SnMeCl_3$  is also a possibility. Therefore,  $SnCl_4$  is likely not a useful precursor, and this type of condensation reaction is generally not well suited for the synthesis of stannosilicates.

Examination of the results for (26A)/(20B) (Table 1) shows that at low temperatures, the driving force for the condensation of CUBE with M-Cl compounds comes exclusively from the preferences of the metal site precursors. Thermal corrections, however, have the effect of favoring the condensation, contributing over -20 kJ mol<sup>-1</sup> at 373.15 K. Since, in this case, there is a large disparity between model A (26A) and B (20B), the latter is taken as more accurate and should be used for interpretation. This situation is favorable with respect to moderating the reactivity of highly energetic precursors, as cooling not only decreases the kinetic rates but also decreases the thermodynamic driving force.

#### **Routes to Al- and Zn-silicates**

The model A of CUBE was used to study the condensation with py–AlMe $_3$ /py–AlCl $_3$  and ZnEt $_2$ /ZnCl $_2$  with respect to the effects of the coordination of THF (one of the two most successful solvents for the CUBE-based systems) as both metals can exist in multiple coordination states.

For py-AlMe<sub>3</sub>/py-AlCl<sub>3</sub>, pentacoordinate states (5C) with THF donors in axial positions were assumed: py-AlMe<sub>3</sub>(THF) (3A) and py-AlCl<sub>3</sub>(THF) (5A). To visualize the interaction of the condensation with coordination, these two reactions,

Table 1 Gibbs energy change for the exchange of [SnMe<sub>3</sub>]<sup>+</sup> between the models A and B of CUBE and Cl<sup>-</sup>

Reaction	Method	$\Delta E \text{ (kJ mol}^{-1}\text{)}$	$\Delta G^0$ (kJ mol <sup>-1</sup> )	$\Delta G_{\mathrm{CBS}}^{273.15} \mathrm{~kJ~mol}^{-1}$	$G_{\rm CBS}^{298.15}  ({\rm kJ \ mol}^{-1})$	$\Delta G_{\rm CBS}^{373.15}  ({\rm kJ \ mol}^{-1})$
(26A)	B3LYP-D3	52.5	45.4	32.9	31.8	28.9
	PBE0-D3	47.3	40.5	28.1	27.2	24.3
(20B)	B3LYP-D3	12.9	6.0	-15.5	-17.3	-22.8
, ,	PBE0-D3	6.1	0.2	-20.5	-21.3	-27.4

along with (2A) and (4A), were added up to form a system with total composition py-AlMe<sub>3</sub> + py-AlCl<sub>3</sub> + 3 Me<sub>3</sub>SnOSiH<sub>3</sub> + THF, where either of the Al precursors can undergo condensation (evolving SnMe4 or SnMe3Cl, respectively) and both Al educt and Al product can coordinate to THF. The total Gibbs free energy of the system in various states is then displayed in Fig. 5 and 6 for 0 and 298.15 K, respectively. Starting from the left and right, the initial precursors py-AlMe<sub>3</sub>/py-AlCl<sub>3</sub> (denoted as states Me<sub>3</sub> and Cl<sub>3</sub>) can undergo three condensation steps towards the center of the diagrams. The energy scale is referenced with respect to the final state with the tetracoordinate (4C) aluminosilicate product py-Al(OSiH<sub>3</sub>)<sub>3</sub>, which is common to both precursors (states denoted as O). In parallel, states containing the pentacoordinate analogs (with additional THF at Al) are visualized in lighter colors (5C).

The results show that while the additional coordination of THF is favored at 0 K (for py-AlCl<sub>3</sub> and py-Al(OSiH<sub>3</sub>)<sub>3</sub> by 50 kJ mol<sup>-1</sup>), it is disfavored at 298.15 K for all states. This agrees with our recent experimental observations, where no evidence for 5- or 6-coordinated Al species was observed in <sup>27</sup>Al TQ/MAS NMR spectra of dried products prepared in THF. While the results are completely realistic for the initial precursors, model A falls short of representing the steric demands of CUBE, therefore, in reality, the coordination of THF to py-AlMe<sub>3-x</sub>(OCUBE)<sub>x</sub> (x = 1-3,

equivalent to states Me2, Me1, and O in Fig. 5 and 6) and py-AlCl<sub>3-x</sub>(OCUBE)<sub>x</sub> (x = 1-3, equivalent to states Cl<sub>2</sub>, Cl<sub>1</sub>, and O) can be expected to be progressively more disfavored with the increasing number of CUBEs in the coordination sphere of Al, even at 0 K. The interaction of the incoming CUBE with the coordinated THF will likely result in an increased reaction barrier. The comparison of the energy difference between the 4-(4C) and 5-coordinated (5C) states (equivalent to the Gibbs energy change of coordination of THF to the 4-coordinated Al species) shows that the dative coordination is most favored by py-AlCl<sub>3</sub> and least by py-AlMe<sub>3</sub>. The final aluminosilicate site is comparable to py-AlCl<sub>3</sub> at low temperatures and it assumes a behavior intermediate between the two precursors at higher temperatures. This has the minor effect of generally decreasing  $\Delta G$  of the condensation steps starting from py-AlCl<sub>3</sub>(THF) and increasing it for py-AlMe<sub>3</sub>(THF).

The conclusion is that although electronically favored, the coordination of weak ligands to Al precursors, resulting in pentacoordinate states, is a minor effect, which does not significantly affect the thermodynamics of the condensation, but it may significantly increase its activation barriers if the reactions are conducted at low temperatures. In any case, if pentacoordinate aluminosilicate species remain in the reaction mixtures upon reaching room temperature, the coordination is lost with heating and solvent removal.

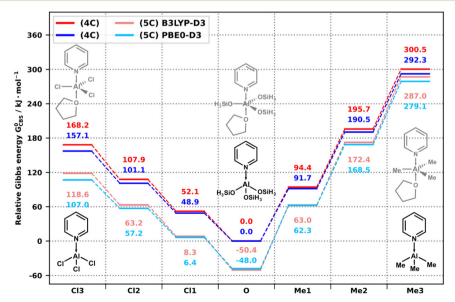


Fig. 5 Relative Gibbs energy of the condensation of py-AlCl<sub>3</sub>/py-AlMe<sub>3</sub> (dark, 4C) and py-AlCl<sub>3</sub>(THF)/py-AlMe<sub>3</sub>(THF) (light, 5C) with model A of CUBE at 0 K, as calculated by B3LYP-D3/CBS (red) and PBE0-D3/CBS (blue).

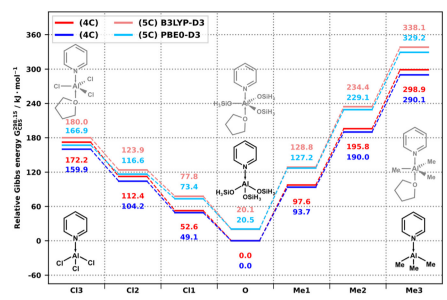
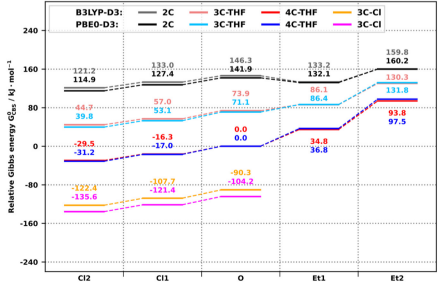


Fig. 6 Relative Gibbs energy of the condensation of py-AlCl<sub>3</sub>/py-AlMe<sub>3</sub> (dark, 4C) and py-AlCl<sub>3</sub>(THF)/py-AlMe<sub>3</sub>(THF) (light, 5C) with model A of CUBE at 298.15 K, as calculated by B3LYP-D3/CBS (red) and PBE0-D3/CBS (blue).

In the case of ZnEt<sub>2</sub>/ZnCl<sub>2</sub> precursors, 2-, 3-, and 4-coordinated states were studied, where 0 (2C), 1 (3C-THF), or 2 (4C-THF) molecules of THF can coordinate to both precursors. For ZnCl<sub>2</sub>, the coordination of a free Cl<sup>-</sup>, resulting in [ZnCl<sub>3</sub>] (3C-Cl), was also envisioned. Thus, reactions (16A)-(22A) were added together to form a system with total composition ZnEt2 + ZnCl<sub>2</sub> + 2 Me<sub>3</sub>SnOSiH<sub>3</sub> + 2 THF + Cl, and the total Gibbs free energy was plotted in Fig. 7-9, for 0, 298.15, and 373.15 K, respectively. Much like in the case of Al above, starting from left and right, the original precursors, denoted as states Cl<sub>2</sub> and Et<sub>2</sub>, can undergo two steps of condensation towards the common zincosilicate products in the center - Zn(OSiH<sub>3</sub>)<sub>2</sub>(THF)<sub>x</sub> for  $ZnEt_2(THF)_x/ZnCl_2(THF)_x$  precursors and  $[ZnCl(OSiH_3)_2]^-$  for [ZnCl<sub>3</sub>]. Likewise, vertical transitions among 2C, 3C-THF, and 4C-THF series involve association or dissociation of THF, while a transition between any one of these states and the 3C-Cl series involves the replacement of any coordinated THF for a single Cl or vice versa. The energy scale is referenced with respect to the state with the most stable uncharged zincosilicate species Zn(OSiH<sub>3</sub>)<sub>2</sub>(THF)<sub>2</sub>.

The results show that the whole dataset exhibits the same monotonic trend hinted at in the previous sections by



 $\textbf{Fig. 7} \quad \text{Relative Gibbs energy of the condensation of } ZnCl_2/ZnEt_2 \text{ (black/gray, 2C), } ZnCl_2(THF)/ZnEt_2(THF) \text{ (light, 3C-THF), } ZnCl_2(THF)_2/ZnEt_2/ZnEt_2/ZnE$ (dark, 4C-THF), and [ZnCl<sub>3</sub>] (orange/pink, 3C-Cl) with model A of CUBE at 0 K, as calculated by B3LYP-D3/CBS (gray/red/orange) and PBE0-D3/ CBS (black/blue/pink).

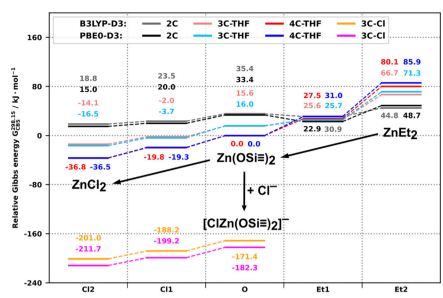


Fig. 8 Relative Gibbs energy of the condensation of  $ZnCl_2/ZnEt_2$  (black/gray, 2C),  $ZnCl_2(THF)/ZnEt_2(THF)$  (light, 3C-THF),  $ZnCl_2(THF)/ZnEt_2(THF)$  (dark, 4C-THF), and  $[ZnCl_3]^-$  (orange/pink, 3C-Cl) with model A of CUBE at 298.15 K, as calculated by B3LYP-D3/CBS (gray/red/orange) and PBE0-D3/CBS (black/blue/pink).

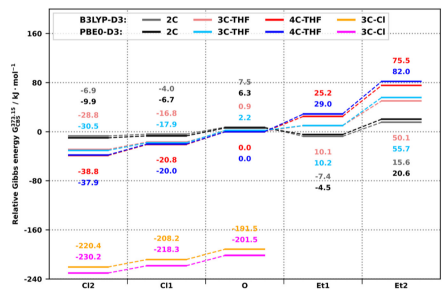


Fig. 9 Relative Gibbs energy of the condensation of ZnCl<sub>2</sub>/ZnEt<sub>2</sub> (black/gray, 2C), ZnCl<sub>2</sub>(THF)/ZnEt<sub>2</sub>(THF) (light, 3C-THF), ZnCl<sub>2</sub>(THF)<sub>2</sub>/ZnEt<sub>2</sub>(THF)<sub>2</sub> (dark, 4C-THF), and [ZnCl<sub>3</sub>]<sup>-</sup> (orange/pink, 3C-Cl) with model A of CUBE at 373.15 K, as calculated by B3LYP-D3/CBS (gray/red/orange) and PBE0-D3/CBS (black/blue/pink).

reactions (18A)/(14B) and (21A)/(15B), where alkyl zinc precursors tend to condense with CUBE to zincosilicates, but these tend to decompose in the presence of SnMe<sub>3</sub>Cl to produce ZnCl<sub>2</sub> and CUBE. The only anomaly is in the case of uncoordinated ZnEt<sub>2</sub>, where the first condensation is spontaneous but the second is not, unless additional stabilization is provided by at least one ligand – a possible sign of the presence of hyperconjugation effects between Zn and Et, which is therefore energetically expensive to lose. This would also explain the much milder  $\Delta G$  of condensation

from uncoordinated states  $Et_2$  to  $Et_1$  compared to their counterparts with coordinated THF. Analogously to the case of Al, the Zn–Cl structures show greater affinity towards THF coordination compared to their Zn–Et counterparts. At 0 K, coordination of THF is universally preferred, but with increasing temperature, the states with higher coordination numbers are gradually more disfavored to the point where at 373.15 K even the coordination to the zincosilicate site  $Zn(OSiH_3)_2$  is in equilibrium. The association of a free  $Cl^-$  ion with the Zn site is already strongly preferred over the

Table 2 Summary of the thermodynamic parameters for the first step condensation of all the studied precursors and their comparison to the properties

M	Site precursor	TD parameters (1st step, model B, kJ mol <sup>-1</sup> )						Metal properties	
		B3LYP-D3		PBE0-D3		Δ (B3LYP-PBE0)		Pauling	
		$\Delta E$	$\Delta G^{298.15}$	$\Delta E$	$\Delta G^{298.15}$	$\Delta\Delta E$	$\Delta\Delta G$	electronegativity	Oxophilicity <sup>34</sup>
В	py-BCl <sub>3</sub>	-88.83	-86.69	-82.42	-80.09	-6.41	-6.60	2.04	1.0
Al	py-AlMe <sub>3</sub>	-113.71	-102.43	-108.44	-98.82	-5.27	-3.62	1.61	0.8
	py-AlCl <sub>3</sub>	-62.45	-59.96	-56.29	-53.76	-6.16	-6.19		
	[AlCl <sub>4</sub> ]	-65.58	-68.33	-62.40	-64.57	-3.18	-3.76		
Si	$SiCl_4$	-89.87	-98.52	-83.95	-92.18	-5.92	-6.35	1.90	0.8
	Me <sub>3</sub> SiCl	-59.37	-60.17	-55.03	-54.95	-4.34	-5.23		
P	$PCl_3$	-80.61	-87.60	-75.65	-81.30	-4.96	-6.31	2.19	0.7
	POCl <sub>3</sub>	-50.28	-61.83	-48.55	-61.14	-1.72	-0.69		
Ti	$TiCl_4$	-62.31	-77.07	-60.88	-74.29	-1.43	-2.77	1.54	1.0
	$Cp_2TiCl_2$	-43.11	-45.32	-37.05	-42.41	-6.06	-2.92		
V	$VCl_4$	-58.35	-70.53	-56.24	-62.56	-2.11	-7.97	1.63	0.8
	VOCl <sub>3</sub>	-50.28	-61.83	-48.55	-61.14	-1.72	-0.69		
Zn	$ZnEt_2(THF)_2$	-66.60	-63.48	-69.00	-57.27	2.39	-6.21	1.65	0.2
	$ZnCl_2(THF)_2$	8.95	15.28	9.55	12.72	-0.60	2.56		
	$[ZnCl_3]^-$	-0.04	-4.47	-0.48	-3.82	0.44	-0.65		
Sn	SnCl <sub>4</sub>	-8.67	-15.31	-6.93	-13.33	-1.74	-1.97	1.96	0.4
Sb	$SbCl_3$	-25.57	-30.07	-23.54	-24.95	-2.04	-5.12	1.82	0.3
	Ph <sub>3</sub> SbCl <sub>2</sub>	-32.34	-30.41	-27.04	-23.34	-5.30	-7.08		

coordination of two THF molecules by 90.3/104.2 kJ mol<sup>-1</sup> at 0 K, by B3LYP-D3 and PBE0-D3, respectively, and it only increases with temperature. Therefore, the Zn/CUBE system will aggressively abstract Cl from available sources, and heating will only thermodynamically promote the process.

Summarized, the coordination of THF has very little effect on the thermodynamics of the condensation of ZnCl<sub>2</sub>, while it has a harmonizing effect on ZnEt2, where it increases the driving force and evens out the differences between the first and the second step of the condensation pathway. The coordination of a free CI ion always dominates over ligands such as THF.

#### General trends

The computed data show several trends across the spectrum of different elements, oxidation states, and computational methods (Table 2). First, the exoergicity of the condensation decreases from light elements down in the groups, which is the primary ordering of Table 2, and it is a natural result of the decreasing ionization energies and electron affinities of heavier elements. There is very little correspondence between the values of the thermodynamic parameters and the Pauling electronegativities of the elements. However, there is a crude correspondence with the oxophilicity scale developed by Early d-block elements are oxophilic, and coincidentally, all of the reported experimentally used transition metal precursors fall into groups 4-6 (Ti to W). Our calculations showed that Zn (group 12) represents the other edge of this presumed "CUBE gap" in the middle of the periodic table. Also, in general, precursors suitable for condensation with CUBE exhibit the covalent character of their bonding to O and the leaving group (Cl, Me, Et).

Second, the condensation is generally more spontaneous and energetic for alkyl metals than for the corresponding metal chlorides. Third, the presence of electron-donating groups on the central metal decreases the driving force (increases  $\Delta G$ ) for the condensation.

Concerning computational characteristics, most of the differences in the driving force come from electronic effects (comparison of  $\Delta E$  to  $\Delta G^{298.15}$ ). The match between the datasets calculated by B3LYP-D3 and PBE0-D3 is remarkable, with the latter predicting systematically higher  $\Delta G^{298.15}$ . The differences are attributable to both electronic energy and thermodynamic corrections. Examination of the CBS extrapolation parameters B (eqn (2)) and  $\alpha$  (eqn (3)) for both datasets (ESI,† Table S47) shows that both parameters assume systematically lower values with PBE0-D3 compared to B3LYP-D3. This means that the CBS convergence for PBE0-D3 is flatter, and the functional is less sensitive to the size of the basis set.

## Conclusions

This study used quantum chemistry DFT methods to estimate the thermodynamic parameters of model reactions in vacuum, representing the steps of the real-world, experimentally reported condensation reactions of the CUBE building block with metal chlorides and alkyl metals conducted in aprotic solvents. The main concerns were the irreversibility of the condensation of the already experimentally reported precursors and the computational screening of other potential precursors for future use.

Our investigation found that the previously reported and experimentally used precursors, AlCl<sub>3</sub>, [AlCl<sub>4</sub>], Si-Cl compounds, PCl<sub>3</sub>, TiCl<sub>4</sub>, and VOCl<sub>3</sub>, should undergo sufficiently irreversible condensation reactions with CUBE.

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The untested precursors BCl<sub>3</sub>, VCl<sub>4</sub>, and POCl<sub>3</sub> are predicted to behave similarly. AlMe3 is almost twice as exoergic as AlCl<sub>3</sub>. Cp<sub>2</sub>TiCl<sub>2</sub> reacts readily in the first step, but the second step may be partially reversible. SbCl3 and Ph3SbCl2 are borderline cases; all steps are similar and spontaneous, but reversibility might pose a problem. SnCl<sub>4</sub> was found to be unsuitable as it is near equilibrium. Zn was found to be an interesting case. While it is possible to prepare zincosilicates from ZnEt2, it is not feasible to use ZnCl2. Moreover, the equilibrium is shifted towards ZnCl2, and the affinity of Zn for Cl is so high that we predict that once a source of Cl, such as SnMe<sub>3</sub>Cl, is introduced, even zincosilicate structures prepared from ZnEt2 will inevitably dissolve back to CUBE and ZnCl<sub>2</sub>.

The condensation of CUBE, in general, appears to be driven mainly by the oxophilicity of the metal in the precursor, which is the primary factor limiting the use of certain elements with CUBE. The driving force is higher for alkyl metals than for the corresponding metal chlorides, and it also decreases for heavy elements. The preference of [SnMe<sub>3</sub>]<sup>+</sup> to bind to Cl<sup>-</sup> over CUBE has a supporting effect, where at low temperatures, the condensation is slightly disfavored, while at high temperatures (100 °C), this process contributes over 20 kJ mol<sup>-1</sup> of the additional driving force, helping to complete the condensation.

This study demonstrated a sound, computationally cheap method for the screening of potential precursors for the CUBE-based synthesis of porous metallosilicates. It also showed that the behavior of B3LYP-D3 and PBE0-D3, together with the CBS extrapolation scheme, is stable and reliable enough that in the future, calculations by a single method can be trusted.

# Data availability

The data supporting this article has been included as part of the ESI.†

## Author contributions

MK: methodology; computation; writing - original draft preparation; visualization; data curation. HS: methodology; writing - review and editing; AS: writing - review and editing; JP: conceptualization; writing - review and editing; funding acquisition; supervision.

## Conflicts of interest

There are no conflicts to declare.

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