



A better model for the mechanism of Lithal (LAH) reduction of cinnamaldehyde?

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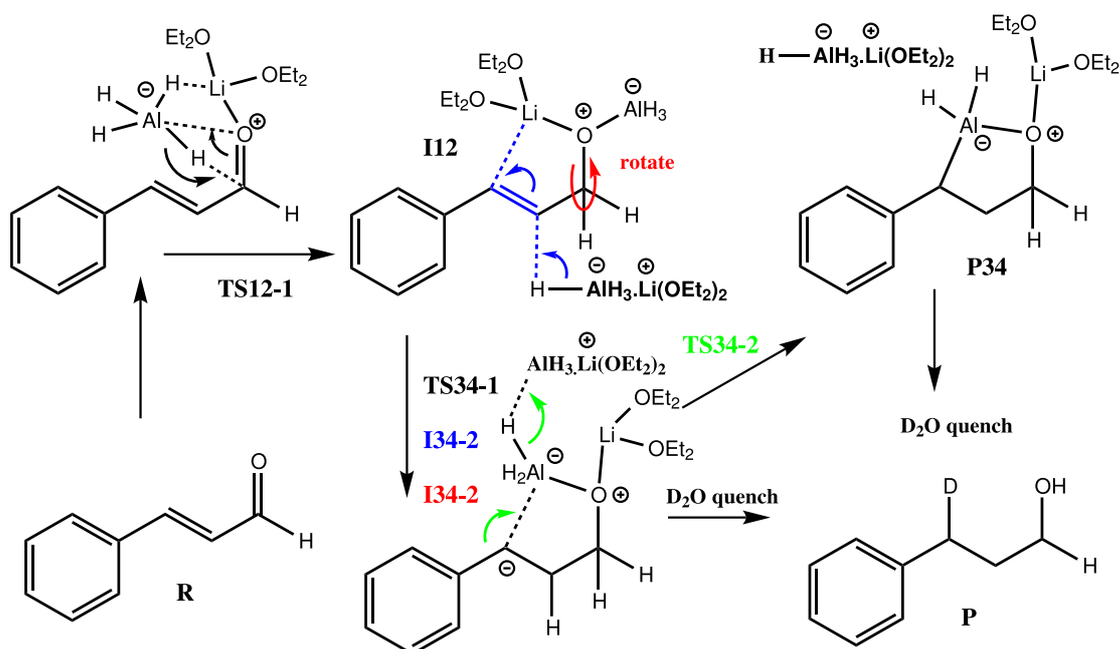
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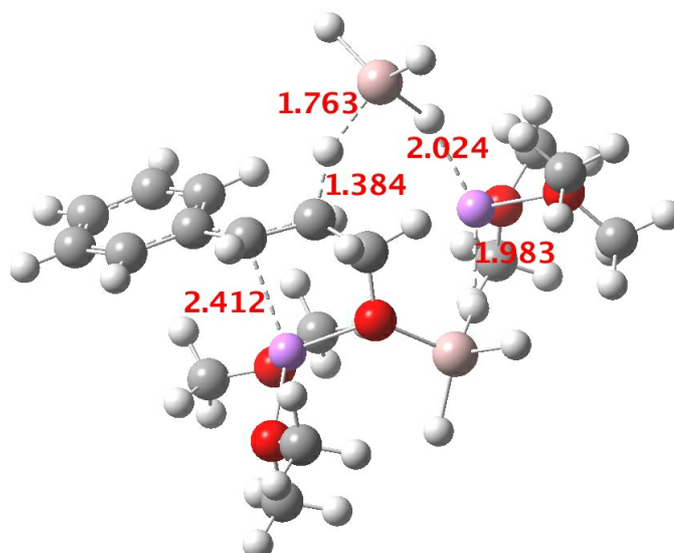
Previously on this blog: modelling the reduction of cinnamaldehyde using one molecule of lithal shows easy reduction of the carbonyl but a high barrier at the next stage, the reduction of the double bond. Here is a quantum energetic exploration of what might happen when a second LAH is added to the brew (the usual ωB97XD/6-311+G(d,p)/SCRF=diethyl ether).



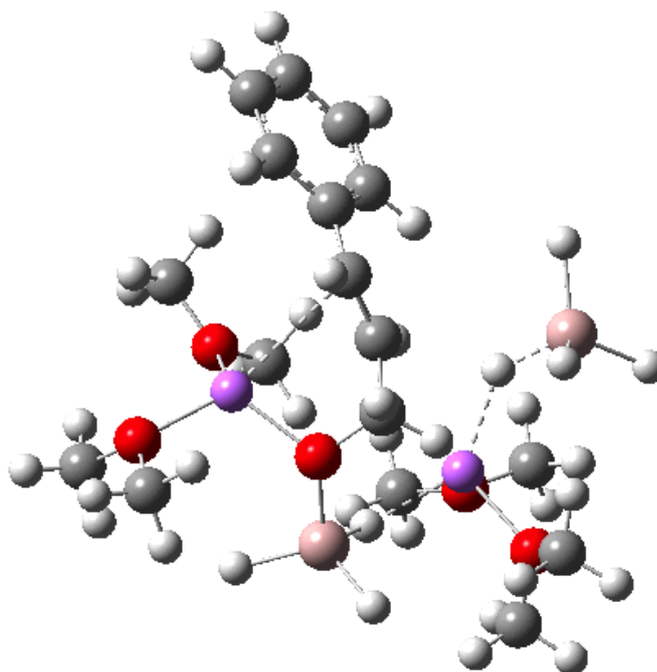
In a [comment](#) at the end of the first post on this theme, I had noted some crystal structures containing in effect $H_xAl.Li(OR)_y$ units ($x=3,4$; $y=0-3$), noting the variety of structural motifs. The current exploration does not even attempt to cover this range of possibilities, but it is informed by the types of weak interaction that these structures reveal. I will nevertheless accept that whatever pathway is revealed here is likely to represent an energetic upper bound and recognise that lower energy pathways may well exist but are yet to be explored.

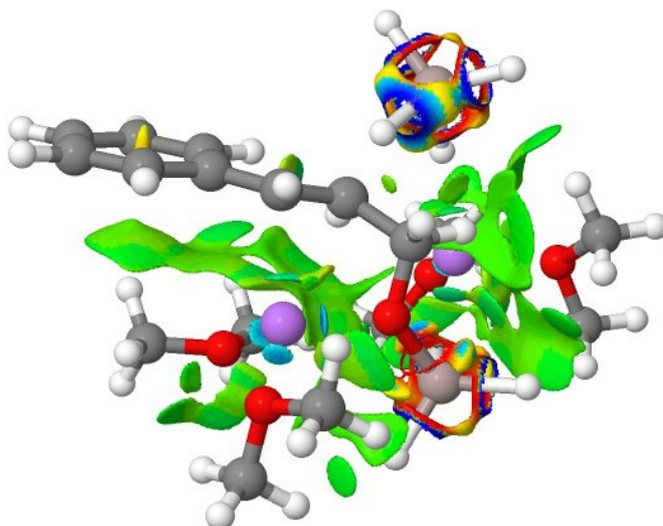
1. At the **I12** stage, a second $AlH_4^-.Li(OMe)_2$ is added and hydride transfer occurs *antiperiplanar* across the C=C bond (**TS34-1**). The computed free energy barrier ΔG_{298}^\ddagger is ~ 24 kcal/mol. The

magnitude of this barrier corresponds to a relatively slow reaction occurring around room temperatures or slightly higher.



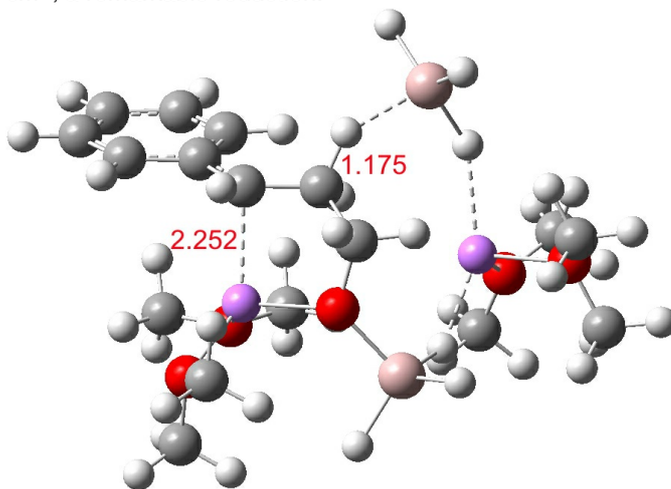
TS. Click for 3D





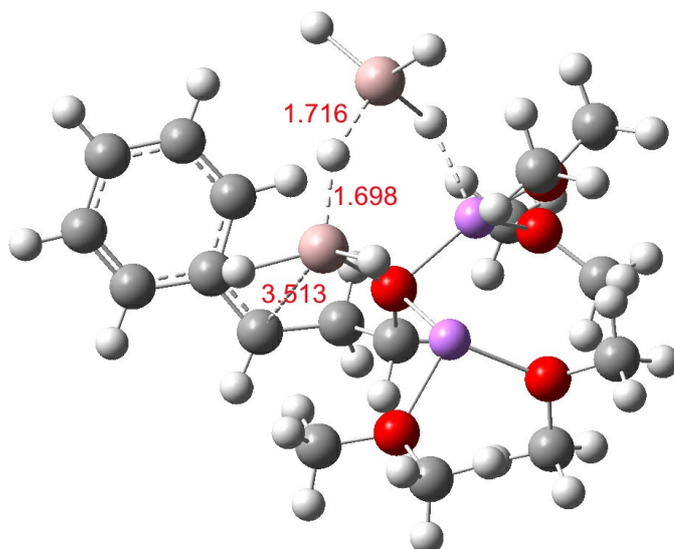
NCI Isosurface (green regions are dispersion stabilizing) Click for 3D

- A transient shallow intermediate **I34-1** is formed in which the benzylic anion is stabilised by an adjacent solvated Li centre. The energy of this species (Table below) needs some explanation.[‡] Can its free energy really be 1.5 kcal/mol higher than that of the preceding transition state? Yes, because its entropy is lower! The transition state is located on a total energy surface, which does not include thermal and entropic corrections; these are always applied AFTER the stationary points are located. If one inspects these total energies, **I34-1** emerges as 1.2 kcal/mol lower than the preceding transition state. This sort of result serves to remind us of the dynamic nature of a potential energy surface, and that static energies may on occasion lead to odd results. Its geometry is shown below, and this too has an interesting feature. The C-H bond just created from the LAH is *antiperiplanar* to the benzylic anion (locked *anti* by the Li) and the resulting stereoelectronic effect reduces its C-H calculated^[1] stretching wavenumber from the normal value of $\sim 3100\text{ cm}^{-1}$ to 2231 cm^{-1} , a remarkable reduction.



I34-1. Click for 3D

- The C-O-AlH₃.Li(OMe)₂ ligand now needs to rotate to **I34-2** so that metal exchange on the benzylic carbon can occur, with Al displacing Li at that position. As with **I34-1**, the free energy of this species is actually slightly higher than that of **TS34-1**. Two AlH₃ groups now exist at this stage (each of them formed by hydride donation as part of the reduction process, see below). A hydride transfer metathesis between them (H₂Al-H-Al₃ is actually a stable bridged species) will generate an AlH₂ as part of the 5-ring aluminate ester in **P34** and regenerate a molecule of LAH. Transition states for these processes (*i.e.* TS34-2) proved difficult to locate;[‡] it may be that the ligand rotation and the hydride metathesis are part of the same concerted process but that is not proven yet.



I34-2. Click for 3D

- The final product prior to hydrolysis is appropriately exoenergetic.
- I would also remark that many aspects of this reaction remain unexplored. For example, AlH_4 can deliver up to four hydrides, becoming progressively substituted as $\text{Al}(\text{OR})_n\text{H}_y$ and in the process losing $\text{Al-H}\dots\text{Li}$ weak interactions. What influence this has on the barriers remains unknown.

Species **Relative ΔG_{298} , kcal/mol** **FAIR Data-**

DOI

| | | |
|--------|-------------------|-----|
| I12 | 0.0 | [2] |
| TS34-1 | 24.1 | [3] |
| I34-1 | 25.5 [†] | [1] |
| I34-2 | 25.0 [†] | [4] |
| P34 | -8.8 | [5] |

In summary, the first step in the reduction of cinnamaldehyde to cinnamyl alcohol requires just one molecule of “ LiAlH_4 ” as reductant and has a very low barrier to reaction. To construct a reasonable model to account for the slower further reduction of the $\text{C}=\text{C}$ bond requires adding a further LiAlH_4 , the key feature being the availability of a lithium centre to stabilise out the forming benzylic carbanion. No doubt even better models might include the effects of adding *e.g.* a third molecule of LAH, and a much more extensive exploration of the various conformational options. But I think the present model might be good enough to augment the apparently relatively limited mechanistic speculations found in text books on the topic.

[†]You sometimes see this phrase in articles reporting transition state location. What it means is that I tried a half-dozen what I thought were reasonable possibilities, and none of them satisfactorily converged. This semi-random exploration of the potential energy surface revealed a very flat energy potential, with lots of conformational possibilities. At this point, you have to decide whether it is worth the time to continue hunting.

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