Alkali-Metal Silyl Zincates: Synthesis, Structure and Reactivity

S. Belrhomari¹, E. Hevia¹*

¹Universität Bern, Department für Chemie, Biochemie und Pharmazie, Switzerland

The high natural abundance of silicon and its non-toxic nature are just some of this element's advantages, strengthened by the popular use of organosilicon compounds within cross-coupling reactions.^[1] Therefore, there stands a natural drive to easily access such useful organosilicon building blocks. While numerous pathways for the silylation of organic molecules have been designed using electrophilic silicon species (mainly using transition metals and/or harsh conditions), the use of nucleophilic silicon reagents is a lesser-employed alternative.^[2] Current methods that have emerged within the past decade include nucleophilic addition or substitution reactions *via* silylmetalation,^[3a] silylboration,^[3b] defluorosilylation^[3c] or nucleophilic silyl reactions *via* silylmetalation,^[3a] silylboration,^[3b] substitution^[3d] using monometallic-silyl reagents. Despite these advances, a gap in the knowledge still exists with regards to (i) fundamental characterisation and understanding of these bimetallic reagents employed and (ii) the *modus operandi* which they follow.

In an effort to bridge this gap, this work exploits the high reactivity of alkali-metals (e.g. K) combined with the better selectivity of a lower-polarity metal (Zn) as a platform for accessing novel, nucleophilic silyl complexes. The results presented include X-ray crystallographic characterisation of these bimetallic silyl reagents, and our preliminary understandings surrounding their nucleophilic capabilities in the direct silylation of haloarenes or the silylzincation of phenylacetylene.

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