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Intramolecular C–H insertion vs. Friedel–Crafts coupling induced by silyl cation-promoted C–F activation†

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Silyl cation-promoted aryl C–F activation can lead to formal C–H activation and the formation of new C(ar)–C(alkyl) bonds. Deuterium-labeling experiments suggest an insertion of a phenyl cation into the C–H bond. From competition experiments, a relation between reaction rate and C–H bond strength could be established. Mechanistic parallels are drawn to the Mascarelli reaction.

Introduction

Silyl cations activate even highly stable C–F bonds.^{1,2} Prominent cases include hydrodefluorinations of fluoroalkanes, by simple $3-5$ as well as bridging $6,7$ cationic silanes, and catalytic intramolecular coupling of a fluoroarene with a non-functionalized arene.^{8,9} Specifically, triisopropylsilylium hexachlorocarba-closo-dodecaborate (1), induces the cleavage of an aryl carbon–fluorine bond allowing a proximal aryl moiety to attack the incipient phenyl cation. If attack by the adjacent arene generates too much strain, as in the formation of a 4-membered ring, the reaction is either very slow or not observed; however, in such cases the presence of an alternative nucleophile, such as the C–H sigma bond of a proximal methyl group, can create a competitive reaction path to capture the phenyl cation (Scheme 1). Exactly this C–F/C–H activation couple forms the basis of this report. RESEARCH ARTICLE

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Scheme 1 Friedel–Crafts type arylation (right) and C–H insertion, involving an alkyl group (left).

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Reaction of 2-fluoro-2′-methylbiphenyl (2) with 5 mol% of $\left[\begin{matrix} \text{Br}_3\text{Si} \end{matrix}\right]$ CHB₁₁H₅Cl₆ (1) and stoichiometric dimethyldimesitylsilane (DMDMS) illustrates the fundamental C–F/C–H activation couple (Scheme 2); wherein 1 serves as catalytic initiator and DMDMS serves as Brønsted base and pro-Lewis acid catalyst. Notably, C–F activation of 2 does not lead to the highly strained biphenylene, but rather to fluorene (3).

Analogous diazoniumbiphenyls exhibit similar reactivity under either thermal or acidic conditions, as observed by Mascarelli in 1932.^{10,11} Recently, studies on Brønsted acidmediated intramolecular cyclization of biaryl triazenes for the synthesis of fluorenes showed similar results.¹² Decomposition of these diazonium salts to form arene carbonium ions has precedence.¹³

Speculations on the mechanism of the Mascarelli and analogous reactions are plentiful with one of the most persistent theories being the shift of a hydride from the methyl group to the phenyl cation, resulting in a benzyl cation, followed by intramolecular electrophilic aromatic substitution (Scheme 3). Alternative mechanisms for the Mascarelli reaction include intermediates such as a pentacoordinated carbocation¹⁴ or a Zwitterion, 15 postulated to be formed by abstraction of a benzylic proton; however no credible experimental evidence has been provided.

Scheme 2 Transformation of 2-fluoro-2'-methylbiphenyl to give fluorene using silyl cation 1 as initiator.

Scheme 3 Postulated mechanisms for the Mascarelli reaction.^{10,11,14,15}

Results and discussion

Evidence for a hydride shift was observed in the denitrogenation of 2-diazonium-N,N-dibenzylbenzamides;¹⁶ the resulting iminium ion does not undergo ring closure, but leads to products of hydrolysis. The phthalimidine originating from intramolecular cyclization, was proposed to be generated from the initial phenyl cation by an insertion mechanism.¹⁷ A hydride shift-derived mechanism was further inferred from the observation of a 2-phenylbenzyl alcohol, generated likely from the reaction of the 2-phenylbenzyl cation with water; 18 but, only 8% of alcohol was isolated. Notably, the benzyl cation generated by protonation of 2-biphenylmethanol with Amberlyst-15 in benzene showed only coupling with solvent; 14 and, heating in cyclohexane produced fluorene, but very slowly compared to the known rate of the Mascarelli and related reactions.

Thermal decomposition of the 2′-trifluoromethylbiphenyl-2 yldiazonium salt generates a difluorobenzylcation, which cyclizes in hexafluorobenzene under oxygen atmosphere.¹⁹ An example relying on niobium-mediated C–F activation was also reported, where 2-trifluoromethylbiphenyl was transformed to 9,9-difluorofluorene.^{20,21}

On the basis of the assumption that silylium-activated fluoroarenes react in a similar fashion as diazonium compounds, the deuterated analog $2-d_3$ was synthesized to observe the fate of the methyl hydrogens. MS and NMR analysis showed that most of the deuterium atoms were still present in the product (ca. 85%). The deuterium distribution within the product was determined by ${}^{2}H$ and ${}^{1}H$ NMR spectroscopy to be 60% @ C1, 18% @ C2, 2% @ C3, 5% @ C4, and ca. 15% deuterium loss.22 Thus, products from deuteration at C1, or C2, plus deuterium loss account for 90+ % of the product formation. Very little deuterium ends up at C4, inconsistent with the expectation of a hydride abstraction mechanism. In contrast, an insertion of the phenyl cation into the carbon deuterium bond followed by elimination or migration could account for this product distribution by the following rationale (Scheme 4).

Intermediate 1, IM1, (a Wheland cation) can either eliminate to yield fluorene possessing no deuterium atom at the arene, or undergo a deuteride shift to form intermediate 2, IM2. IM2 could rapidly transform by another migration into the much more stable intermediate 3, IM3, from which the

Scheme 4 Deuterium distribution in 3 produced from $2-d₃$; postulated intermediates and their M06-2X/Def2-TZVPP(chlorobenzene) computed energetics.

majority of elimination occurs. Taking into account that eliminations and migrations occur more readily for hydrogen than deuterium, one would predict a pronounced deuteration at position 1, but also substantial deuteration at position 2.

The stabilities of IM1 and IM2 were calculated to be very similar $(\Delta E_{\text{[IM2-IM1]}} = -2.1 \text{ kJ mol}^{-1})$, in contrast to IM3, which is much lower in energy $(\Delta E_{\text{[IM3–IM1]}} = -34.3 \text{ kJ mol}^{-1})$.²³ Transition state $TS(1,2)$, is calculated to be rate determining over TS(2,3) $\left[E_{TS(1,2)} = 25.7 \text{ kJ mol}^{-1} \text{ vs. } E_{TS(2,3)} = 21.7 \text{ kJ mol}^{-1} \right]$ for the hydride migration IM1–IM2–IM3. This computational data leads to a model in which the initially formed IM1 partitions with preference for intramolecular migration over abstraction of D⁺. IM2 is fleeting and further hydride migration occurs to IM3 without appreciable elimination. A Hammond Postulate analysis allows this to be modeled by basic principles of physical organic chemistry (Scheme 5).

Intramolecular hydride transfer to form a benzyl cation followed by ring closure would place substantial deuteration at C4, which is not observed. Additionally, abstraction of fluoride from 2-fluoromethylbiphenyl (4) yielded primarily 5 and 6 (coupling with mesitylene from DMDMS^{24}) and traces of 7 but no fluorene, further demonstrating the preference of the benzyl cation for intermolecular coupling (Scheme 6). Only trace amounts of two intermolecular coupling isomers with DMDMS could be observed for the reaction of 2. Thus, intramolecular hydride transfer seems mechanistically insignificant in the formation of fluorene from 2.

C–F activation of 2 led exclusively to 3 by benzyl C–H insertion. In contrast, for substrates 8a–8c where benzyl C–H insertion and Friedel–Crafts reaction pathways are probable, a competitive distribution of products can be observed (Scheme 7). Whereas the substrate possessing the stronger benzylic C–H bond (8a, toluene-like: 375 kJ mol⁻¹) prefers for-

Scheme 5 Calculated reaction energy profile (M06-2X/Def2-TZVPP in chlorobenzene)²³ for hydride migration from $IM1-M2-IM3$ [TS(1,2) rate determining]. (a) Projection of hydride position in TS(1,2) and TS(2,3) onto the carbon–carbon bond length as an estimate of late/early character; (b) triangle defined by carbon and hydrogen positions in TS(1,2) and TS(2,3) to show late/early TS assignment.

Scheme 6 Product distribution from abstraction of fluoride from 4.

Scheme 7 Product distributions from C–F activation 8a–c.

mation of the fluoranthene derivative 9a over 10a, with a ratio of 1.6 : 1.0, the substrates featuring the weaker benzylic C–H bond (8b & 8c, cumene-like: 348 kJ mol⁻¹) rather form products of the insertion $10b$ and $10c$ $(1.0:1.9$ and $1.0:2.1$, respectively).

Scheme 8 Non-benzylic C–H bond insertion from C–F activation of 11.

Computational studies predict an activation energy for C–F activation of ~100 kJ mol⁻¹, which is consistent with the reaction proceeding at 90 °C over 2 h. Calculated transition states leading to products 9a and 10a are very close in energy $(\Delta E_{TS}(9a-10a) = -1.2 \text{ kJ mol}^{-1}$ [$\varepsilon = 5.7 \text{ chlorobenzene}$]).¹⁵ Experimental product ratios also suggest that the transition states are very close in energy $(\Delta E_{TS((9a-10a)} = +1.5 \text{ kJ mol}^{-1})$. The difference between theory and experiment (2.7 kJ mol⁻¹ in chlorobenzene) is well within the limits of reliability – emphasizing the importance of small energy differences for product ratios in the range of 1:2 to 2:1 (ca. 4 kJ mol⁻¹). Computations suggest that changes in dielectric constant can influence the product selectivity $(\Delta E_{TS(9a-10a)} = +1.5 \text{ kJ mol}^{-1} [\varepsilon = 1$ vacuum]) but, they do not change the general course and nature of the reaction.

Noteworthy is that only a small fraction of the difference in C–H bond strength shows up in the difference in transition state energy ($ca. 5-10\%$). Invoking the Hammond postulate again here, one would conclude that the reaction product selectivity step has an early transition state, further supporting a direct C–H insertion mechanism. An early C–H insertion mechanism would predict the reaction to be relative promiscuous with regard to C–H insertion chemistry. Indeed, the reactivity is not restricted to weak benzylic CH groups. Applying the reaction conditions on 2-fluoro-2′-tert-butylbiphenyl (11) also yields ring-closed product (12) (Scheme 8). Despite the large difference in C–H bond strength for a primary aliphatic CH_3 group compared to a benzylic CH_3 group $(\Delta E \sim 40 - 50 \text{ kJ mol}^{-1})$, the formation of the six-membered ring, aliphatic insertion product, is only slightly slower than that of the model compound. 25 This result further supports the early transition state, direct insertion model for this reaction.

Conclusions and outlook

Parallel to the Mascarelli reaction for diazonium salts, phenyl cations generated from arylfluorides by C–F bond activation with cationic silyl Lewis acids can undergo intramolecular arene-coupling and C–H insertion. Deuterium labeling studies suggest a direct C–H insertion mechanism. Competition experiments showed only a slight dependence of aryl–aryl coupling vs. C–H insertion based on the C–H bond strength, consistent with an early-transition-state to a direct-insertion mechanism. The tandem activation of two generally stable functional groups (Ar–F, alkyl–H) provides a new reactivity pattern in cationic silyl Lewis acids.

Experimental and computational details

Synthesis and characterization data, together with computational details, are available in the ESI.†

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