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Rapid Sonogashira cross-coupling of iodoferrocenes and the unexpected cyclo-oligomerization of 4-ethynylphenylthioacetate

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A systematic study into the Sonogashira cross-coupling of 1,1'-diiodoferrocene (fcI₂) confirms that the Pd(0)/P(^tBu)₃ system provides a remarkable rate increase over Pd(0)/(PPh₃)₂. Attempts to couple 4-ethynylphenylthioacetate ¹⁰ (2) with fcI₂ instead produced a novel cyclic trimer of the

former, from *syn* addition of S–Ac across C≡C.

Whilst large quantities of pure iodo-¹ and 1,1'-diiodoferrocene² can now easily be obtained, the full synthetic exploitation of ¹⁵ these useful starting materials may only be realized through optimizing onward reaction conditions (enabling high product yields). Towards this end, increasing the typically low/moderate

reactivity of iodoferrocenes³ (versus aryliodides/bromides) under Sonogashira cross-coupling conditions was considered a primary ²⁰ target. Convenient and widely applicable, this reaction (from Fc–I containing materials) has been used to construct compounds for

- containing materials) has been used to construct compounds for molecular⁴ and organic⁵ electronics, the study of intramolecular electron transfer,⁶ photo-⁷ and electro-chemical sensing,⁸ catalysis (pincer complexes),⁹ and artificial bio-receptors.¹⁰ It is worth ²⁵ noting that just two years after the seminal 1975 papers
- concerning anyl iodides by Heck,¹¹ Cassar¹² and Sonogashira *et al.*,¹³ cross-coupling of iodoferrocenes was under investigation.¹⁴

In the reports referenced above, syntheses have usually employed the convenient $PdCl_2(PPh_3)_2$ precatalyst in DIPA/THF.

³⁰ However, variances in substrate structure and reagent stoichiometry make it difficult to ratify the superiority of any particular set of conditions. Of particular significance, one paper¹⁵ describes unsuccessful attempts with Buchwald and Fu's Pd(PhCN)₂Cl₂/P(^tBu)₃ combination – utilized to rapidly cross-

³⁵ couple electron-rich aryl-bromides at room temperature.¹⁶ It appeared that normally outstanding catalytic systems might offer little or no benefit over the use of PPh₃-ligated complexes in Sonogashira reactions with iodoferrocenes.

We were accordingly motivated to explore the reaction ⁴⁰ between fcI₂ and phenylacetylene as a model in an attempt to optimise the Sonogashira cross coupling of iodoferrocenes and terminal alkynes in general (Scheme 1). Concentration, temperature, phosphine (14 examples), solvent (3 examples), time and the phenylacetylene/fcI₂ ratio were systematically varied to ⁴⁵ examine their individual effects. Product yields were determined

via ¹H NMR spectroscopy of crude reaction mixtures. As shown in Fig. S-2, overall reaction yields were found to

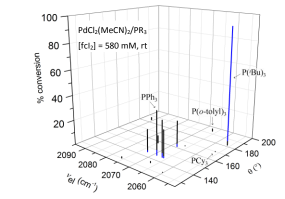
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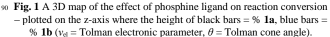
increase substantially with substrate concentration (using PdCl₂(PPh₃)₂ at 80°C). The effect of phosphine ligand on reaction ⁵⁰ yields was subsequently explored by employing PdCl₂(MeCN)₂ *at room temperature* – PR₃ was added separately to form PdCl₂(PR₃)₂ *in situ*. Though PPh₃ was found to be the best ligand within its local steric/electronic landscape (10% yield **1b**, Fig. 1), the PdCl₂(MeCN)₂/P('Bu)₃ combination provided an exceptional ⁵⁵ rate improvement (93% yield **1b**). Furthermore, whilst all reactions were run for 20 h, with P('Bu)₃ a large exotherm, and rapid and complete precipitate formation was observed after ~15 min. Such observations eliminate any previous doubt¹⁵ as to the superior reactivity of bulky, electron-rich phosphines¹⁶⁻¹⁷ in this ⁶⁰ context.

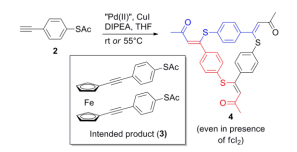
In comparison, other variables provided only small changes in yields. It was additionally noted that hydrodehalogenation reactions (Fc–I \rightarrow Fc–H) occur under these conditions (limiting yields in some cases), attributed to adventitious water. Full ⁶⁵ experimental details may be found in the supporting information.[†] Intrigued as to why previous reactions using PdCl₂(PhCN)₂/P([']Bu)₃ had proven unsuccessful, we noted that in these attempts substrates had featured thioacetyl and pyridyl

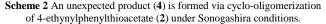
functionalities.¹⁵ This prompted investigation into Fe + (2 eq. per Cp-I) $(2 \text{$

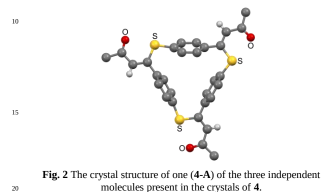
75 Scheme 1 Model reaction used to study the Sonogashira cross-coupling of iodoferrocenes and terminal alkynes.

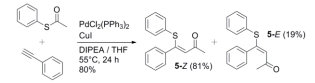












25 Scheme 3 Monofunctional analogues of 4-ethynylphenylthioacetate react under Sonogashira conditions to form the addition product 5.

the apparently straightforward, though as yet unreported, Sonogashira cross-coupling reaction between 4ethynylphenylthioacetate (**2**) and fcI₂. The intended product, 1,1'-³⁰ bis[(4-thioacetylbenzene)ethynyl]ferrocene (**3**), has previously been prepared via Stille coupling.¹⁸

Remarkably, all attempts to synthesise **3** via Sonogashira coupling failed – ¹H NMR spectroscopy of the crude product mixture indicated that the fcI₂ had not appreciably reacted even ³⁵ after 24 h (PdCl₂(PPh₃)₂, 55°C). However, upon closer inspection it became apparent that resonances attributable to **2** had disappeared, and new peaks were observed at approximately δ 2.3, 6.4 and 7.0 ppm (intensities 3:1:4). Column chromatography led to the isolation of a bright yellow solid, the identity of which

- ⁴⁰ was unambiguously confirmed by X-ray crystallography[‡] as that of the novel cyclic trimer (**4**, 24%; Scheme 2 and Fig. 2). A material with matching spectral features was produced using $PdCl_2(MeCN)_2/P(^{t}Bu)_3$ in an analogous procedure at room temperature.
- ⁴⁵ Overall this reaction may be described as cyclooligomerization of **2**, generating **4** via the intermolecular *syn* addition of acetyl and thiolate moieties to C≡C (accompanied by cleavage of the S–Ac bond, and reduction to C=C). Whereas **4** comprises all-Z linkages, reaction between monofunctional
- ⁵⁰ analogues of **2** (S-phenylthioacetate and phenylacetylene) yielded the known compound **5**¹⁹ in 80% yield as a mixture of *Z* (81%) and *E* (19%) isomers (Scheme 3). Sequential elimination of components showed that $PdCl_2(PPh_3)_2$, CuI and DIPEA were all

necessary for the reaction to occur in THF at a reasonable rate ⁵⁵ (proceeding sluggishly in the absence of CuI, and not at all without PdCl₂(PPh₃)₂); fcI₂ is not required in any case. Based on studies by other groups,²⁰ we propose a reaction mechanism such as that shown in Scheme 4. (Tokuyama *et al.* previously isolated a series of 1-alkynyl ketones via an analogous process, using an ⁶⁰ excess of CuI to trap the thiolate anion.^{20a} Under more forcing conditions, Minami *et al.* used a similar Pd/Cu catalysed reaction to prepare 2,3-dihydrothiopyran-4-one derivatives, isolating and reacting key intermediates to demonstrate the reaction pathway.^{20b}) In the absence of Cu(I), it is speculated that an ⁶⁵ alkyne insertion step may occur in place of transmetallation.²¹

It was of additional interest to identify where Fc–I sits within the well-established aryl halides/triflate rate series,²² and the observation that **4** forms even in the presence of fcI₂ is particularly revealing. From the above discussion it is certainly 70 evident that $k_{(Ar-I)} >> k_{(Fc-I)}$, as aryl iodides will rapidly crosscouple with terminal alkynes under most circumstances (resulting in high to quantitative yields, even at room temperature).¹³ Noting also that aryl bromides are readily cross-coupled in the presence of thioacetate moieties,²³ it is inferred that rates of 75 oxidative addition to Pd(0) follow the series: $k_{(Ar-I)} > k_{(Ar-OTf)} > k_{(Ar-Br)} > k_{(S-Ac)} > k_{(Fc-I)}$ (assuming oxidative addition to Pd(0) is always the rate-limiting step).

Conclusions

A systematic study into the Sonogashira cross-coupling of iodoferrocenes indicates that yields are maximised by employing high reagent concentrations and reaction temperatures. Though not immediately apparent given prior reports utilising iodoferrocenes,¹⁵ we have shown that superior reactivity *can* be obtained in this context using the PdCl₂(MeCN)₂/P([']Bu)₃ sc combination. Utilisation of such conditions may prove invaluable when attempting cross-couplings of iodoferrocenes at room temperature or lower concentrations (Fig. S-2), for example when working with temperature-sensitive substrates or small quantities of advanced intermediates (following multi-step syntheses).

It was further demonstrated that the cross-coupling of terminal alkynes and iodoferrocenes is impracticable in the presence of thioacetate moieties. This is presumably due to a competing Pdcatalysed reaction between the thioacetate group and terminal alkyne(s), effectively blocking oxidative addition of Fc–I to

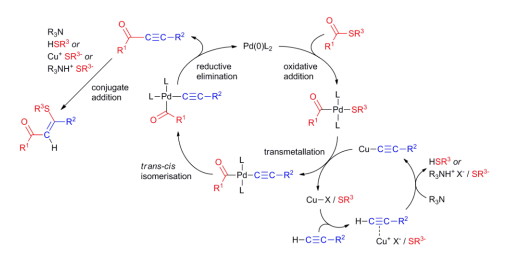
Pd(0). Whilst the bifunctional ligand 2 yielded an isolable cyclic trimer (4), it is considered that systems of higher complexity have previously formed unexpected, potentially polymeric, product mixtures under Sonogashira conditions. Future work in our laboratories will explore these Fc–I and S–Ac based catalytic processes in more detail, the latter being important for biological applications (e.g. functionalization of SAc-terminated bioconjugates,²⁴ modification of biologically-relevant small

molecules²⁵) and the syntheses of novel chelates, redox-active materials and conducting polymers with β -thioketone linkages.

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Notes and references

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Scheme 4 A proposed reaction mechanism: producing cross-coupled acetyl-alkyne and thiolate products that subsequently react under basic conditions to produce $R^1C(O)CH=C(R^2)S(R^3)$ -type compounds (interpreted from work by Tokuyama and Minami *et al.*²⁰).

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† Electronic Supplementary Information (ESI) available: detailed experimental procedures, tables of data for all catalytic runs, discussion of hydrodehalogenation reactions, ¹H/¹³C{¹H} NMR spectra, ²⁰ crystallographic information. See DOI: 10.1039/b000000x/

- ^{*} Crystal data for 4: C₃₀H₂4O₃S₃·CH₂Cl₂ M = 613.60, monoclinic, Cc (no. 9), a = 24.8640(4), b = 24.0808(3), c = 15.1854(3) Å, $\beta = 96.5311(18)^\circ$, V = 9033.2(3) Å³, Z = 12 [3 independent molecules], $D_c = 1.354$ g cm⁻³, μ(Mo-Ka) = 0.455 mm⁻¹, T = 173 K, yellow blocks, Oxford
- ²⁵ Diffraction Xcalibur 3 diffractometer; 22101 independent measured reflections ($R_{int} = 0.0304$), F^2 refinement,^[26] $R_1(obs) = 0.0489$, $wR_2(all) =$ 0.1470, 18011 independent observed absorption-corrected reflections [$|F_o|$ $> 4\sigma(|F_o|)$, $2\theta_{max} = 65^\circ$], 1099 parameters. The absolute structure of **4** was determined by a combination of *R*-factor tests [$R_1^+ = 0.0489$, $R_1^- =$ 30 0.0498] and by use of the Flack parameter [$x^+ = 0.00(4)$, $x^- = 1.01(4)$]. CCDC 927952.
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