Research article

The self–indicating preparation of bromoferrocenes from stannylferrocenes and an improved synthesis of di-iodoferrocene

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ABSTRACT

A simple method for the preparation of bromoferrocenes from stannylferrocenes is described: the preparation of 1,1′-dibromoferrrocene is used as an example. Stannylferrocenes are reacted with bromine directly in dichloromethane to give bromoferrocenes in a self-indicating titration reaction. Also, an enhanced method of removal of the highly soluble organotin by-products formed in the preparation of 1,1′-diiodoferrocene from 1,1′-tri-n-butylstannylferrocene is reported allowing the preparation of large-scale quantities of 1,1′-diiodoferrocene, which is one of the most important starting materials in ferrocene chemistry.

1. Introduction

Halogenoferrocenes are particularly useful synthons in the preparation of a broad range of useful ferrocenes such as ferrocenylenamines [1, 2], diisocyanoferoferrocenes [3], trifluoroferrocenes [4], and of course in the metathesis reaction to lithioferrocenes [5, 6] (although lithioferrocenes to halogenoferrocenes is more common) which lead to hundreds of ferrocene-based compounds. Of the halogenoferoferrocenes arguably the most important are iodo- and bromo-ferrocenes because they are more reactive in standard organic coupling reactions, [7, 8]. There has been much debate on the synthesis of iodoferrocenes concerning the most useful and viable synthetic methodology some of which are summarised for just one representative compound, 1,1′-diiodoferrocene, Figure 1 [9, 10, 11, 12, 13, 14, 15, 16, 17].

All the synthetic methods shown in Figure 1, use iodine as a synthon with a ferrocene-metal or ferrocene-metalloid compound although clearly there are other quenching reagents such as iodine monobromide, iodine monochloride, diiodotetrafluoroethane (route A (ii), yields 90–95%)*, and N-iodosuccinimide (35–40%)* which can replace iodine (60–65%)*("yields quoted are those we have routinely obtained based on many synthetic attempts). The very first preparations of iodoferrocenes were made using the reactions of ferrocenyl mercuric chlorides (route E, Figure 1) which requires the preparation of the poorly soluble bis mercury salt which is difficult to crystallise and the reaction of lithioferrocenes with iodine (route A(i), Figure 1) which were reported by Rausch and co-workers, [9, 10]. Additionally, ferrocene based organometallic reagents such as Grignard reagents or ferrocenyloboronic acids may be directly reacted with iodine (routes B and D, Figure 1). Both these routes require the formation of a precursor compound which is itself difficult to obtain pure without product loss. The synthetic route A (ii) using di-iodotetrafluoroethane was by far the best synthetic route as a cheap supply of this quenching reagent was available in the early 1990’s but the industrial synthesis was halted. The reaction with iodine on large scale worked reasonably well although it was imperfect; nevertheless, we and others used the lithiation route routinely in the early 1980s although many by-products such as 1,1′′′-di-iodobiferrocene, 7, Figure 2, which results from metathesis, were formed, [14]. Of course in some instances the metathesis reaction itself can be useful, for example, when [Fe(CO) 2CpI] is used to quench 1,1′-dilithioferrocene to give the iron-substituted iodobiferrocene, 8, in good yields (20–25%) together with [Fe(CO) 2Cp] 2 and small quantities of other iodoferrocenes.

The separation and purification of iodoferrocenes was carried out using column chromatography. This served well as an undergraduate teaching exercise to clean up the crude product but later a commercial temperature gradient sublimation apparatus was used to purify the iodoferrocenes. In response to the poor-quality synthesis we developed a new synthesis which used stannylferrocenes as intermediates, Figure 1, C [16]. This synthetic method is a cheap and high-yielding but it does suffer from one major disadvantage which is the product is difficult to crystallise, i.e. it has a low melting point and is non-polar, so it is

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difficult to remove the tin by-products and trace ferrocene. Despite this difficulty the method has been well used, [18, 19, 20, 21]. In the original work the tin by-products were removed by adding fluoride, which turns them into a polymeric material which could be filtered off. However, the large-scale use of fluoride is not ideal because of its high toxicity. Two papers describe the clean-up methods [22, 23] for removal of trace ferrocene by selective oxidation which is a particularly elegant solution to the first problem. Given the inherent importance of iodoferrocenes it was decided to re-examine the second problem, i.e. the removal of the by-product, tri-\(n\)-butylstannyl iodide, on scales greater than 5g. Corey had proposed a method to remove similar alkyltins from organic reaction mixtures using mixed metal salts [24] but this turned out to be synthetically difficult because of the nature of the reagents and scales involved. We now report a solution to this problem which is simple, effective and the synthetic method has been developed into an alternative synthesis of bromoferrocenes too. It should be noted that (trimethylstannyl)ferrocenes also react with iodine to give iodoferrocenes and may be less problematic in the workup because of their reduced solubility, however we have focussed on the use of more readily available low cost \(n\)-butyltins to produce an efficient synthesis.

2. Results and discussion

The preparation of tri (\(n\)-butylstannyl)ferrocenes from the reaction of lithioferrocenes with chloro-\(n\)-butyltin is a routine procedure which can be carried out on very large scale in a broad range of dry solvents [25]. Following an exhaustive review of the literature where tin halides were unwanted by-products (and many trials) it was decided to attempt the use of aqueous sodium hydroxide or ammonia solutions to remove the tin containing compounds. (Supplementary Figures S1 and S2) This initially gave limited success but subsequently we developed the method using biphasic conditions to obtain pure iodoferrocenes. The method is summarised in Figure 3.

![Figure 1. General synthetic routes to diiodoferrocene, A-E.](image1)

![Figure 2. Biferrocenes 7 and 8 obtained with iodine in coupling reaction of 1,1'-dilithioferrocenes.](image2)

![Figure 3. Schematic of synthetic steps in 1,1'-diiodoferroocene preparation and purification. The progress of iodination reaction, step 1, may be followed by NMR (Figure S3).](image3)
1,1′-Bis-tri-n-butylstannylferrocene, which may be prepared and used in situ, (1H NMR spectrum shown in Figure S4) was reacted with iodine in dichloromethane and the product was vigorously stirred overnight with aqueous ammonia with the addition of some methanol to enhance the reagent miscibility. A fine white precipitate removed. It is known that trimethyltin halides react with NaOH to form the corresponding tin hydrides and also that trimethyl tin iodide reacts with trimethyl tin hydride to give a white crystalline solid, [(Me3SnOH)2SnMe3] + 1−. A similar chemistry is expected in this case where methyl substituents are replaced with n-Bu groups. The reason ammonium hydroxide is more efficient in the clean-up process than using aqueous sodium hydroxide is almost certainly a case of better miscibility of the reaction solvents. It is possible that bis-(tri-n-butyltin)oxides, such as (n-Bu3Sn)2O are also formed. Thus, the filtered solution was added to alumina and the solvent was removed to give a free-flowing sample. This solid was loaded onto a standard alumina column and the product was eluted with petrol as a yellow orange band to give pure 1,1′-diiodoferrocene. (1H NMR, Figure S5). The remaining tin containing by-products adhere to the column and are apparent as a pale green and deep green bands allowing the elution of pure di-iodoferrocene with petrol (column photograph, Figure S6). The deep green band was eluted with copious amounts of diethyl ether however it turns yellow instantly on contact with air and after solvent removal it was clear that it contained only trace amounts of iodoferrocenes. We assume it is an oxidised ferrocene which accounts for the adherence to the column and are apparent as a pale green and deep green bands allowing the elution of pure di-iodoferrocene with petrol (column photograph, Figure S6). The deep green band was eluted with copious amounts of diethyl ether however it turns yellow instantly on contact with air and after solvent removal it was clear that it contained only trace amounts of iodoferrocenes. We assume it is an oxidised ferrocene which accounts for the intense deep green colour. The reason for the adherence to the column is assumed to be that the remaining by-products are complex alkyltin hydrides as described earlier. The reaction of 1,1′-bis-(tri-n-butylstannyl)ferrocenes with iodine monobromide likewise produces 1,1′-diiodoferrocenes, again relatively cleanly with the formation of the by-product tri-n-butylstannylbromide. This was not too surprising – the only surprise was that the reaction was so clean with minimum halide scrambling. The reaction is faster than that of the corresponding reaction with iodine. The latter observation led us to examine the reaction of bromine with 1,1′-bis-(tri-n-butylstannyl)ferrocene, which proceeds even more rapidly and thus can be carried out as a titration reaction in dichloromethane: this effectively removes the possibility of oxidation of ferrocenes with excess bromine, Figure 4.

When a bromine solution in dichloromethane is added to a dichloromethane solution of stannyferrocenes a deep green/black spot on the reaction mixture produces an intense dark green colour. The reaction of 1,1′-diiodoferrocene with diethyl ether (100mL) and the organic layer was separated. A further extraction was carried out as a self-indicating titration. The product 1,1′-dibromoferrocene, 9, (1H NMR spectrum, Figure S7) does not suffer from the purification problems of 1,1′-diiodoferrocene because it is so easy to crystallise from methanol or hexanes. As noted earlier the use of trimethyltin chloride as an alternative quenching reagent gives less soluble reaction products however the low cost of tri-n-butyltin chloride means that on large scale this is the reagent of choice and of course this clean-up method will also ameliorate the purification step with this reagent. Clearly the use of a surfactant as a phase transfer reagent could further enhance the removal of the tin containing by-products by increased solvent miscibility, but this was not found to be necessary in this case.

3. **Summary**

An improved method of producing pure 1,1′-diiodoferrocene on a scale up to 100g has been developed. Adaptation of this method can be used for the formation of bromoferrrocenes. Although the clean-up method has focussed on diiodoferrocene it will have broad applicability towards other reactions containing alkyltin halide by-products.

4. **Experimental section**

All reagents used were of reagent grade. Note: Rubber gloves should be worn when working with alkyltin compounds because of their toxicity.

4.1. **1,1′-diodoferrocene preparation**

The general method given is suitable for scales between 5g and 50g ferrocene; the following is an example on 100mmol scale. All reactions are carried out under an inert atmosphere whereas workup methods are carried out under normal laboratory conditions. When using aqueous ammonium hydroxide it is particularly important to work in fume hood, neutralising any excess before disposal.

1,1′-Dilithioferrocene/TMEDA was prepared in hexanes as follows: a solution of n-butyllithium in hexanes (88 mL of a 2.5M solution) was added to a slurry of ferrocene (18.6g, 100mmol) and excess TMEDA (12.0g, 103 mmol) in hexanes (500mL). The mixture was stirred overnight, (min 12h, and it is important that stirring does not stop and a heavy-duty magnetic stirrer at slow speed is always used). The reaction flask is immersed in an external cooling bath of acetone and the mixture was allowed to warm to room temperature without removal of the cooling bath over a period of 2–3h. before being re-cooled to ca -30°C. It was then hydrolysed slowly by the gradual addition of water (300mL) initially dropwise, increasing volumes as the hydrolysis proceeded. The organic layer was separated by addition of further diether ether (100mL) and diethyl ether (100mL) hexanes is then added dropwise over a period of 20 min to the reaction mixture while the cooling bath is maintained at -75°C. The mixture was allowed to warm to room temperature without removal of the cooling bath over a period of 2–3h. before being re-cooled to ca -30°C. It was then hydrolysed slowly by the gradual addition of water (300mL) initially dropwise, increasing volumes as the hydrolysis proceeded. The organic layer was separated by addition of further diether ether (100mL) and the organic layer was separated. A further extraction was performed on the aqueous layer with diethyl ether 200mL and the organic layers were combined vacuum filtered through a loose plug of celite (in a 15cm Buchner funnel) and dried over stirred anhydrous magnesium sulfate. Volatiles and solvent were subsequently removed on a rotary evaporator to furnish a thick amber colored oil of the crude product 1,1′-bis-(tri-n-butylstannyl)ferrocene. This oil was dissolved in dichloromethane (250 mL) and this was treated incrementally with excess freshly ground iodine (2.2 equiv. l2). [slow addition is recommended as an exotherm occurs]. The slurry was stirred in the dark

![Figure 4. Simple synthesis of 1,1′-dibromoferrocene from the 1,1′-tri-(n-butyl-stannyl)ferrocene using bromine.](image-url)
It is noted that if bromine is added one drop at a time to the centre of an unstirred solution of bi-tri-n-butylstannylferrocene then the disappearance of the blue/black spot that forms in the centre is an excellent teaching demonstration.

Declarations

Author contribution statement

Ian R. Butler: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Wrote the paper.

Yassin T. H. Mehdar, Zahraa S. Al-Taie: Performed the experiments.

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Competing interest statement

The authors declare no conflict of interest.

Additional information

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