Direct Lithiation of (Cyclobutadiene) tricarbonyliron and (**(Trimethylsilyl) c yclobutadiene) tricarbonyliron with sec-Butyllithium: Selective** *para* **Metalation**

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(Cyc1obutadiene)tricarbonyliron (la) is metalated on treatment with sec-BuLi in **THF,** as could be shown by reaction of the product **(1ithiocyclobutadiene)tricarbonyliron** (lb) with a variety of electrophiles (inter *alia* trimethylchlorosilane, methyl disulfide, methyl iodide, and diiodoethane); the respective adducts were obtained in good yields. Similarly, metalation of **((trimethylsily1)cyclobutadiene)tricarbonyliron** (IC) affords **(3-lithio-l-(trimethylsilyl)cyclobutadiene)tricarbonyliron** (3a), which has also been trapped with standard electrophiles. **No** products of ortho metalation were observed.

Introduction

The chemistry of cyclobutadiene' has challenged both experimentalists and theoreticians due to its importance with regard to the concept of (anti)aromaticity.^{1b} The extreme reactivity of cyclobutadiene does not allow isolation of this compound under ordinary conditions.2 Stabilization can be attained by capture through a macrocyclic host^{2a} or complexation with an 14-electron organotransition-metal fragment to yield stable organometallic compounds,2b of which the most famous are **(cyc1obutadiene)tricarbonyliron** (la13 and cobalt complex **2.4** Free cyclobutadiene is liberated by oxidative decomposition of $1a^{2b}$ by cerium(IV) salts; $1a$ can be considered a storage form of cyclobutadiene. Substituted cyclobutadiene complexes undergo this reaction as well.2b

The synthesis of substituted **(cyc1obutadiene)tricarbo**nyliron complexes has been achieved by preparation of suitable dihalocyclobutene precursors⁵ in multistep sequences, by electrophilic substitution of la under Lewis acid catalysis, thereby exploiting its metalloaromatic character,^{5b,6} or most elegantly by dimerization of alkynes under the influence of a transition-metal fragment.⁷

Direct lithiation on the other hand is an important and often very efficient way of functionalization for organic

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and organometallic entities.8 **A** prerequisite for the success of this reaction is the sufficient activation of the **C-H** bond involved and the stability of the substrate to strongly basic reaction conditions.

In contrast to the well-developed field of lithiated cyclopentadienyl complexes? the chemistry of lithiated **organotransition-metal-bonded** cyclobutadienes has attracted much less attention. Early attempts to deprotonate la by either MeLi or n-BuLi afforded the ketones **lj,k** instead.1° Up to now there have been only three examples of lithiated cyclobutadiene complexes. One is prepared by metal halogen exchange of the iodide 1d¹⁰ (which itself is not easily available $6a$) with methyllithium. In the two other cases the organolithiums were obtained either by tin lithium exchange of a cobalt cyclobutadiene complex¹¹ or by direct deprotonation of 2 with *n*-butyllithium in tetramethylethylenediamine.⁴ In the latter case the organolithiums were quenched by carbon dioxide. Metalation of both rings is a side reaction, and the yield of the desired mono acid (30%) is not satisfactory.

To find conditions for direct lithiation of la would be a desirable process, considering the synthetic potential of organolithiums!

Results

Reaction of 1 with sec-butyllithium in tetrahydrofuran at **-78** "C for **15** min (conditions described by Stille for lithiation of **cyclopentadienylmanganese** tricarbonyle) afforded a dark brown but clear solution of lb. Reaction of lb with trimethylchlorosilane, 1,2-diiodoethane, methyl disulfide, trimethyltin chloride, methyl iodide, hexachloroethane, and diphenylchlorophosphane yielded the corresponding heterosubstituted **(cyc1obutadiene)tricarbo**nyliron complexes **IC-i** in yields between **58** and **67%** (Table I). **A** notable exception is chloride **lh,** which was isolated in **26%** yield only. Compounds ld-g have not been described in the literature.

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*⁴*The entries for **lj-l** are only to list the substituents.

When silane IC was subjected to the same metalation conditions with sec-BuLi in THF, a clear yellow solution of 3a formed. Functionalization of the organolithium compound with methyl iodide resulted in isolation of a yellow oil, whose proton NMR spectrum shows three singlets at δ 4.05, 1.79, and 0.10 in the ratio 2:3:9. The carbon NMR shows six signals at 6 215.1 **(s),** 93.9 **(s),** 69.4 (d), 61.2 **(s),** 13.7 (q), and -0.7 **(9).** The signal at 6 215.1 is attributed to the carbonyl groups, while three resonances at 93.9, 69.4, and 61.2 are assigned to the complexed cyclobutadienyl ring. The remaining two signals are due to the methyl and the trimethylsilyl group, respectively. These spectroscopic data strongly suggest that the two substituents are placed para to each other, leaving a plane of symmetry in 3b intact. Reaction of 3awith diiodoethane or methyl disulfide gives rise to the isolation of compounds 3c and 3d. Their NMR spectroscopic data are in full accordance with the above para assignment. In the case of **(bis(trimethylsily1)cyclobutadiene)tricarbonyliron (38)** (obtained by functionalization of 3a with trimethylchlorosilane in 46% yield) the para substitution pattern can only be concluded from the other functionalization experiments. To strengthen the assumption of para metalation, le was deprotonated and the corresponding metalated species were functionalized with trimethylchlorosilane. An inseparable mixture of the ortho and para isomers 3d and 4d is formed in the ratio of ca. 1:2:3. The major component has the same NMR spectroscopic characteristics as the independently prepared para compound 3d. In the case of the ortho-substituted 4d the two ring protons appear as two singlets at δ 4.16 and 4.64, respectively; for the cyclobutadiene ring of 4d the carbon NMR spectrum shows also four different signals of correct multiplicity, in contrast to the spectrum for the parasubstituted 3d, where only three signals are observed.

When 3b was subjected to the above-described metalation conditions and functionalized with methyl disulfide, a single compound was isolated in 16% yield **after** chromatography involving heavy losses. Its proton NMR spectrum shows four singlets at δ 4.17, 3.06, 2.22, and 0.10 in the ratio of 2:2:3:9. Moreover, the carbon NMR spectrum shows seven signals at δ 214.6 (s), 91.7 (s), 68.5 (d), 63.9 **(s),** 31.6 (t), 16.0 (q), and-0.8 (q). These spectral data rule out this compound being (l-(trimethylsilyl)-2- **(methylthio)-3-methylcyclobutadiene)** tricarbonyliron **(5):** the proton-coupled carbon NMR spectrum shows the presence of a methylene group and one instead of two methyl groups. The carbon NMR spectrum of **5** would be expected to show signals for three and not for only two quarternary carbon atoms **as** observed. Additionally, the integration of the proton NMR spectrum is at odds with the proposed structure. Instead, structure 3f is in full accordance with the spectral data.

In order to make sure that the methyl position is not generally more acidic than the ring protons in compounds such as li, we carried out a deprotonation reaction of this substrate under the above-described conditions and functionalized with methyl disulfide. We isolated an inseparable mixture which contained two main compounds, the ortho- and para-disubstituted complexes 3g and 4g, and a small (less than 5 *5%* by proton NMR integration) amount of other compounds which might have resulted by deprotonation of the methyl group. The selectivity of the ortho position over the para position is in this case ca. 2.51, corrected by the statistical bias.

Discussion

Attempts to deprotonate $1a$ with n -alkyllithium reagents such as n-BuLi or MeLi resulted in the formation of ketones 1*i* and $1k^{10}$ The use of sec-BuLi at -78 °C in THF fortuitously suppresses this undesired side reaction. Instead, clean lithiation of the complexed cyclobutadiene ring is achieved. A wide variety of heterosubstituted complexes lc-i can be conveniently synthesized by this method. These compounds are otherwise not easily prepared. Especially, the hitherto unknown alkynylsubstituted **(cyc1obutadiene)tricarbonylirons** should be accessible from iodide^{6a} 1d, which we can now obtain in gram quantities.

In the case of silane 1c a pronounced para selectivity of the metalation reaction is observed. It is known that silicon substituents stabilize adjacent negative charges,¹² while at the β -position a positive charge is heavily stabilized13 by the overlap of the carbon-silicon bond with the empty π orbital of the positive charge. The enhanced electron density of the carbon-silicon bond, on the other hand, destabilizes a negative charge β to the silicon substituent. This is probably more pronounced due to the forced syn-periplanar arrangement in the situation under consideration. The large steric demand of the trimethylsilyl group additionally enhances the para selectivity. 1,3-Heterodisubstituted (cyclobutadiene)tricarbonylirons are otherwise difficult to obtain;^{2b,5b,6b} an exception is the low-yield dimerization of a 1-silyl-2-amino-

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substituted alkyne by iron pentacarbonyl.¹⁴ Therefore, the method described here offers an access to *para***trimethylsilyl-substituted (cyc1obutadiene)tricarbonyl**irons **(3).**

In the case of the methyl- and the methylthic-substituted cyclobutadiene complexes **li** and **le,** deprotonation occurs at both *ortho* and *para* positions and the small differences in regioselectivity obtained render these reactions unattractive for preparation of disubstituted (cyclobutadiene)tricarbonylirons. In **li** we could not detect metalation of the methyl substituent.

However, the regioselectivity changes in the case **of 3b,** whose most acidic proton is not located on the cyclobutadiene ring but at the attached methyl group. It is not unreasonable to assume that the increased steric bulk of the two substituents suppresses the ring metalation completely and instead leads to the metalation of the methyl protons. In a similar case, the attempted functionalization of **116b** with n-butyllithium merely leads to isolation of starting material. The reason for the difference in behavior **of 11** compared to that of **3b** is not clear.

Experimental Section

General Procedure. A flame-dried and nitrogen-flushed **100** mL Schlenk flask with magnetic stirbar is charged with **50** mL of anhydrous **THF** (freshly distilled from benzophenonepotassium) and the specified amount of (cyclobutadiene)tricarbonyliron complex. After the mixture is cooled to **-78** "C, **1.1** equiv of an **1.4** mol/L solution of sec-BuLi in pentane/ cyclohexane is added by syringe. After **15** min of stirring at this temperature a dark clear solution has formed. An excess of the electrophile is added by syringe, and the cooling bath is removed. The reaction mixture is stirred for **1** h. After aqueous workup the residues are distilled at **0.001** mmHg and subsequently chromatographed over silica with pentane as eluent to give the desired compound as brown-yellow oils.

Tricarbonyl[(1-4-η)-(trimethylsilyl)-1,3-cyclobutadiene]**iron10 (IC). la (500** mg, **2.60** mmol), sec-BuLi **(2.00** mL, **2.80** mmol), and trimethylchlorosilane **(326** mg, **3.00** mmol) are treated according to the general procedure. Distillation at **20** "C/O.001 mmHg and chromatography yields **403** mg **(59** %) of the known **IC.**

Tricarbonyl[(1-4- η)-iodo-1,3-cyclobutadiene]iron^{6a} (1d). **la (2.00** g, **10.4** mmol), sec-BuLi **(8.20** mL, **11.5** mmol), and diiodoethane **(3.24** g, **11.5** mmol), after distillation at **20** "C/O.OOl mmHg, yield **2.23** g **(67.3%)** of the known **Id.**

Tricarbonyl[(l-4-~)-(methylthio)-l,3-cyclobutadiene] iron (1e). 1a (265 mg, 1.38 mmol), sec-BuLi(1.10 mL, 1.54 mmol), and methyl disulfide **(1.00** g, **10.62** mmol), after distillation at **20** "C/0.005 mmHg, yield **201** mg **(61** %) of the analytically pure **le.** IR (KBr, cm-'): **v 2925,2043 (CO),1964** (CO), **1434,1424,1342, 1317.** 'H NMR (CDCl3): 6 **2.21** *(8,* **3** H), **4.13** *(8,* **1 H), 4.23 (8, ²** H). '3C NMR (CDC13): 6 **18.38 (q),60.45** (d), **65.25** (d), **84.34 (81, 213.77 (e).** Mass (EI; *m/z* (relative intensity, *7%)):* **238** (M+, **531, ²¹⁰**(M+ - CO, **loo), 182** (M+ - **2C0, 161, 154** (M+ - **3CO), ¹²¹ (48).** Anal. Calcd: C, **40.36;** H, **2.54.** Found: C, **40.00;** H, **2.64.**

Tricarbonyl[(1-4- η)-(trimethylstannyl)-1,3-cyclobutadi**eneliron (If). la (312** mg, **1.63** mmol), sec-BuLi **(1.28** mL, **1.80** mmol), and trimethyltin chloride (400 mg, **2.01** mmol), after distillation at **40** 0C/0.005 mmHg, yield **346** mg **(60%)** of the sensitive **If.** IR (KBr, cm-1): **v 2986,2919,2040** (CO), **1960** (CO), **1256, 1193, 992, 927, 824. ¹H NMR (CDCl₃): δ 0.20 (s, 9 H), 3.97** *(8,* **2** H), **4.42** *(8,* **1** H). "C NMR (CDC13): 6 **-8.71** (q), **65.81 (s), 70.72** (d), **71.36** (d), **215.09** (9).

Tricarbonyl[$(1-4-\eta)$ -(diphenylphosphino)-1,3-cyclobuta**dieneliron (le). la (208** mg, **1.08** mmol), sec-BuLi **(0.86** mL,

Bunz

1.20 mmol), and chlorodiphenylphosphine **(265** mg, **1.20** mmol), after chromatography (pentane/methylene chloride, **211,** yield 237 mg (58%) of the analytically pure 1g. IR (KBr, cm^{-1}) : ν **3071, 3056, 2048** (CO), **1975** (CO), **1907, 1479, 1434, 1093.** 'H NMR (CDCl3): 6 **4.36** (d, *JPH* = **0.64** Hz, **2** H), **4.49** (d, *JPH* **6.54** Hz, **1** H) **7.3-7.6** (m, **10** H). 13C('H) NMR (CDC13): 6 **63.70,68.86, 77.00,77.43,137.57,133.10,132.87,128.90,128.41,213.11.** Mass (EI; *m/z* (relative intensity, %)I: **376** (M+, **0.5), 348** (M+ - CO, **58), 292** (M+ - **3C0, 100), 291 (61), 188 (48), 183 (51).** Anal. Calcd C, **60.67;** H, **3.48.** Found C, **60.44;** H, **3.63.**

 $Tricarbonyl[(1-4-*n*)-chloro-1,3-cyclobutadiene]iron¹⁵ (1 h).$ **la** *(500* mg, **2.61** mmol), sec-BuLi **(1.90** mL, **2.66** mmol), and hexachloroethane **(660** mg, **2.79** mmol), after distillation at **20** $^{\circ}$ C/0.01 mmHg, yield 152 mg (26%) of the known 1h.

Tricarbonyl[(1-4-₇)-methyl-1,3-cyclobutadiene]iron¹⁶(1i). **la** *(500* mg, **2.61** mmol), sec-BuLi **(1.90** mL, **2.66** mmol), and methyl iodide **(452** mg, **3.18** mmol), after distillation at **20** "C/ **0,001** mmHg, yield **309** mg **(58%) of** the known **li.**

Tricarbonyl[(1-4-r))-3-methyl-l-(trimethylsilyl)-1,3 cyclobutadiene]iron (3b). IC (362 mg, **1.37** mmol), sec-BuLi (LlOmL, **1.54** mmol), and iodomethane (1.00g, **7.05** mmol), after distillation at $20 °C/0.05 mmHg$ and subsequent chromatography, yield 239 mg (63%) of analytically pure 3b. IR (KBr, cm^{-1}) : ν **3085,2959,2925,2900,2040** (Co), **1963** (CO), **1451,1337,1251, 1054, 1014, 841.** ¹**H** NMR (CDCl₃): δ 0.10 (s, 9 **H**), 1.79 (s, 3 **H**), **4.05 (~,2** H). I3C NMR (CDCl3): 6 **-0.70** (q), **13.66** (q), **61.20 (s),** *76)):* **278** (M+, **91,250** (M+ - CO, **16), 222** (M+ - **2C0, 24), 194 93.91 (s), 69.38** (d), **215.14** *(8).* Mass (EI; *m/z* (relative intensity, (M+ - **3C0, 581, 154 (100).** Anal. Calcd: C, **47.50;** H, **5.07.** Found: C, **47.47;** H, **5.28.**

Tricarbonyl[(1-4-9)- l-iodo-3-(trimet hylsilyl)-1,3-cyclobutadieneliron (3c). IC (507 mg, **1.92** mmol), sec-BuLi **(1.57** mL, **2.20** mmol), and diiodoethane **(620** mg, **2.20** mmol), after distillation at 40 °C/0.001 mmHg and subsequent chromatography, yield **422** mg **(56%)** of **3c.** IR (KBr, cm-l): *Y* **3110,2958, 2048** (CO), **1974** (CO), **1308,1252,1012,843.** 'H NMR (CDCl3): δ -0.05 (s, 9 H), 3.81 (s, 2 H). ¹³C NMR (C₆D₆): δ -1.06 (q), **31.29 (s), 65.77 (s), 74.13** (d), **214.45** *(8).* Mass (EI; *m/z* (relative intensity, $\%$)): 390 (M⁺, 5), 362 (M⁺ ~ CO, 16), 334 (M⁺ - 2CO, **23), 306** (M+ - 3C0, **E), 266 (12).**

Tricarbonyl[(1-4-9)- 1-(methylthio)3- (trimethylsily1)- 1,3-cyclobutadiene]iron (3d). IC (236 mg, **0.893** mmol), sec-BuLi **(0.650** mL, **0.910** mmol), and methyl disulfide **(500** mg, **5.31** mmol) yield **199** mg **(72** %) **of** crude product and after chromatography yield **92** mg **(33%)** of analytically pure material. IR (KBr, cm-l): **v 2957,2926,2039** (CO), **1965** (CO) **1962** (CO), **1435, 1425, 1251, 843.** ¹**H** NMR (CDCl₃): δ 0.09 **(s, 9 H), 2.20 (s, 3 H) 4.28 (8,2** H). "C NMR (CDCl3): 6 **-0.81** (q), **15.88** (q), **60.89 (s),** %)): **310** (M+, **19), 282** (M+ - CO, **84), 254** (Mt - **2CO,59), 226 97.17 (s), 67.67** (d), **214.16** (9). Mass (EI; *m/z* (relative intensity, (M+ - **3C0,49), 168 (77), 154 (100).** Anal. Calcd: C, **42.59;** H, **4.55.** Found: C, **42.67;** H, **4.53.**

Tricarbonyl[(1-4-9)- 1,3-bis(trimet hylsilyl)-l,3-cyclobutadieneliron (3e). IC (244 mg, **0.924** mmol), sec-BuLi **(0.700** mL, **0.980** mmol), and trimethylchlorosilane **(200** mg, **1.84** mmol), after distillation at **0.01** mmHg/3O "C and subsequent chromatography, yield **143** mg **(46%)** of **3e.** IR (KBr, cm-'): **v 3120, 2960,2038** (CO), **1966** (CO), **1251,839.** 'H NMR (CDCl3): 6 **0.05** $(s, 18 H), 3.74 (s, 2 H).$ ¹³C NMR $(C_6D_6): \delta -1.12 (q), 73.39 (d),$ **77.96 (s), 215.81** *(8).* Mass (EI; *m/z* (relative intensity, %)I: **336** (M+, **23), 308** (M+ - CO, **32), 280** (M+ - **2C0,43), 252** (Mt - 3C0, **loo), 154 (72).** Anal. Calcd: C, **46.25;** H, **5.99.** Found C, **46.35;** H, **6.08.**

Tricarbonyl[(1-4-η)-3-((methylthio)methyl)-1-(trimeth**ylsilyl)-1,3-~yclobutadiene]iron** *(3f).* **3b (211** mg, **0.759mmol),** sec-BuLi **(0.571** mL, 0.800 mmol), and methyl disulfide **(200** mg, **2.12** mmol), after chromatographywith a **1:1** methylene chloride/ pentane mixture, yield **39** mg **(16%) of** analytically pure **3f.** IR

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Selective para Metalation

(KBr, cm-l): *v* 2957, 2919, 2040 (CO), 1965 (CO), 1438, 1337, 1251, 1012. ¹H NMR (CDCl₃): δ 0.10 (s, 9 H), 2.22 (s, 3 H), 3.06 **(s,** 2 H), 4.17 *(8,* 2 H). 1% NMR (CDCl3): **6** -0.83 **(q),** 16.04 **(q),** 31.60 (t), 63.94 **(a),** 68.52 (d), 91.66 **(s),** 214.60 **(8).** Mass **(EI;** *mlz* (relative intensity, *5%)):* 324 (M+, 3), 296 (M+ - CO, 72), 277 Calcd: C, 44.44; H, 4.97. Found: C, 44.62; H, 5.02. (loo), 268 (M+ - 2C0,44), 240 (M+ - 3C0,25), 224 (68). **Anal.**

Metalation of le. le (113 mg, 0.475 mmol), sec-BuLi (0.360 mL,0.504mmol),and trimethylchlorosilane (105mg,0.966mmol), after distillation at 30 °C/0.001 mmHg, yield 51 mg (35%) of a 2.3:l mixture of **3d** and **tricarbonyl[(1-4-r))-l-(methylthio)- 2-(trimethylsilyl)-l,3-cyclobutadiene]iron (4d) as an** inseparable mixture. **3d was** identified according to ita **known** NMR data (vide *supra).* **4d** 1H NMR (CDCla) 6 0.18 (s,9 H), 2.21 **(8,** 3 H), 4.16, 4.64 (2 **S,** 2 H); '3C NMR (CDCl3) **6** -0.64 **(q),** 19.00 **(q),** 65.26 **(s),** 65.60 (d), 72.25 (d), 88.84 **(8)** 214.17 *(8).*

Metalation of li. li (309 mg, 1.50 mmol), sec-BuLi (1.20 mL, 1.68 mmol), and methyl disulfide (212 mg, 2.25 mmol), after distillation at 40 °C/0.01 mmHg, yield 117 mg (31%) of an inseparable mixture of **tricarbonyl**[(1-4- η)-2-methyl-1-**(methylthio)-1,3-cyclobutadiene]iron (4g)** and **tricarbonyl-** [$(1-4-\eta)$ -3-methyl-1-(methylthio)-1,3-cyclobutadiene]iron (3g) in a 5.1:1 ratio. **4g**: ¹H NMR (CDCl₃) δ 1.84 (s, 3 H), 2.23 (s, 3 61.52 (d), 63.01 (d), 63.90 **(s),** 88.18 **(s),** 214.11 *(8).* **3g:** 1H NMR H), 4.15,4.19 (2 **S,** 2 H); '3C NMR (CDCl3) 6 11.63 **(q),** 19.26 **(q),** (CDCla) 6 1.79 **(8,** 3 H), 2.19 *(8,* 3 H), 4.30 **(s,** 2 H).

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