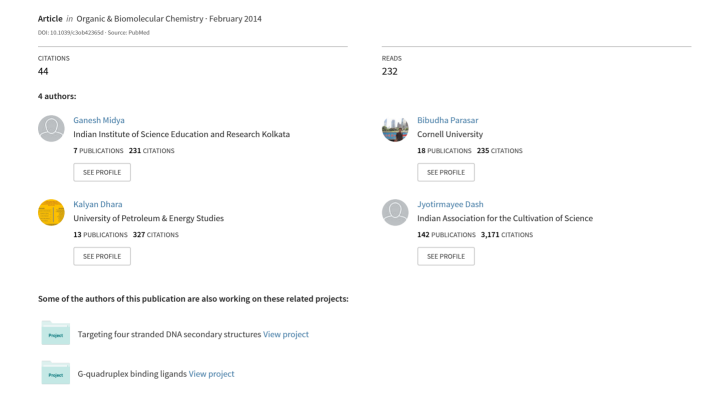
ChemInform Abstract: Ligand Mediated Iron Catalyzed Dimerization of Terminal Aryl Alkynes: Scope and Limitations.



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Ligand mediated iron catalyzed dimerization of terminal aryl alkynes: scope and limitations†‡

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Regioselective dimerization of terminal aryl alkynes to produce conjugated enynes has been achieved using FeCl₃ and KO^tBu in the presence of either DMEDA or dppe. The reaction proceeds smoothly in toluene at 145 °C for 2 h to give the corresponding head-to-head dimers in good to excellent yields (54 to 99%) with high E-selectivity (67: 33 to 83: 17 E/Z). Both strongly electron-donating and electron-with-drawing groups are compatible with this procedure. The bidentate phosphine (dppe) ligand exhibits better catalytic activity than the bidentate amine (DMEDA). The aliphatic acetylene fails to react under this catalytic system which suggests that potassium tertiary butoxide activates the conjugated system of aryl acetylene through cation-pi interaction and pi-pi interaction. A radical inhibitor (galvinoxyl or TEMPO) completely suppresses the reaction. Employing FeCl₂ as a catalyst instead of FeCl₃, only phenyl acetylene afforded the corresponding head to head dimer in good yield. Mechanistic pathways for both FeCl₃ catalyzed dimerization of aryl alkynes and FeCl₂ catalyzed dimerization of phenyl acetylene have been proposed.

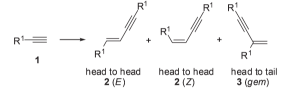
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Introduction

Conjugated enynes are the key structural unit in many natural and bioactive molecules, drug intermediates and organic materials. Conjugated enynes can be used as precursors for the synthesis of polysubstituted benzenes and stereospecific 1,3-dienes and natural products. Thus, the past few decades have witnessed substantial efforts in the direction of developing versatile strategies to synthesize the enyne synthons. Commonly used practical methods in the literature include alkynylation of alkene metal, metal-catalyzed coupling between an alkene and an organometallic alkyne, Sonogashira type coupling between terminal alkyne and vinyl halide and the dimerization of terminal alkynes. The synthesis of enynes by dimerization of terminal alkynes is a straightforward and an atom-economical process (Scheme 1).

However, a highly selective synthesis of conjugated enynes by a dimerization is challenging due to the competitive formation of the other three possible (E), (Z), and gem-enyne isomers (Scheme 1). $^{10-20}$ Catalysts based on transition



Scheme 1 Dimerization of terminal alkyne.

metals^{10–17} (Zr,¹⁰ Hf,¹¹ Re,¹² Ru,¹³ Rh,¹⁴ Ir,¹⁵ Ni¹⁶ and Pd¹⁷), lanthanides,¹⁸ actinides¹⁹ and main group elements²⁰ have been developed for the dimerization of terminal alkynes. Although these methods have been utilized extensively and effectively, some limitations exist because of the difficulties in synthesizing the organometallic substrates, toxicity and the recovery of expensive noble metal catalyst for an industrial scale preparation.

Recently iron-based catalytic systems have attracted significant growing interest because iron salts are readily available, inexpensive, and environmentally benign. ^{22–25} Iron salts have been used as important alternatives to established transition metal-catalyzed carbon–carbon and carbon–hetero atom bond formation reactions. ^{22–25} Realizing the importance of the enyne motif and its potential application we thought it would be useful if a protocol for the facile transformation of alkyne to enyne can be designed using the cheap iron salts as catalysts.

It is noteworthy that iron/copper promoted oxidative homocoupling reaction of terminal alkynes has been reported to give the corresponding diynes (Scheme 2, eqn (ii))²⁶ and an

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[†]This paper is dedicated to Professor Rabindranath Mukherjee on the occasion of his 60th birthday.

[‡] Electronic supplementary information (ESI) available: Analytical data and NMR spectra of all synthetic compounds. See DOI: 10.1039/c3ob42365d

Scheme 2 Dimerization and trimerization of terminal alkyne using iron salts

iron salt/organolanthanide-based bimetallic catalytic system has been efficiently used for cyclotrimerization of terminal alkynes (Scheme 2, eqn (iii)).27 We have recently presented the first report of an iron catalyzed highly regioselective protocol for the dimerization of terminal alkynes using N^1 , N^2 -dimethylethane-1,2-diamine (DMEDA) as a ligand in the presence of KO^tBu (Scheme 2, eqn (i)).²⁸

Recently Bolm and Buchwald reported that FeCl3 catalyzed reactions may be influenced by trace amounts of copper impurities.²⁹ However, iron/copper promoted oxidative homo-coupling reaction of terminal alkynes has been reported to give the corresponding diynes (Scheme 2, eqn (ii)).26 Herein we explore various aspects of this head to head dimerization of terminal aryl alkynes, from the effect of electronic and steric properties of ligands, to the scope and limitations to mechanistic manifolds.

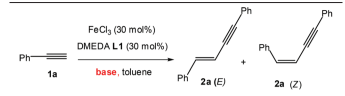
Results and discussion

In our previous work, 28 we employed 30 mol% of FeCl₃ in the presence of 30 mol% of DMEDA and 3 equivalents of KO^tBu for the dimerization of terminal alkynes. The reaction was performed in toluene at 145 °C for 2 h (Table 1). To improve the yield and selectivity, it was decided to study the effect of bases first keeping other parameters fixed.

Screening of bases

Bases tend to affect the dimerization reactions to a major extent. In the absence of a base, no reaction took place (Table 1, entry 1). The use of cesium chloride dramatically improved the yield to 48% (entry 2) with a good selectivity (89:11). When potassium carbonate (K₂CO₃) was employed as the base, the reaction took even longer time (72 h) without an appreciable increase in yield of the dimers with decreased stereoselectivity (entry 3). Potassium phosphate (K₃PO₄) after 3 days gave E/Z selectivity comparable to that of the cesium carbonate (Cs₂CO₃); however, the yield of 2a was very low (35%, entry 4). The reaction was sluggish in the presence of 3 equivalents of lithium tertiary butoxide (LiO^tBu) and sodium tertiary butoxide (NaO^tBu) (entries 5 and 6). Surprisingly, when 2 equivalents of potassium tertiary butoxide (KO^tBu) were used

Table 1 Screening of bases for iron catalyzed dimerization of phenyl acetylene 1a



Base (equiv.)	Temp. (°C), time (h)	Ratio ^a of 2a (E/Z)	Yield ^b (%) of $2a (E/Z)$
No base	145. 48	No reaction	No base
	,		48
	,	67:33	50
$K_3PO_4(3)$,	86:14	35
LiO^tBu (3)	145, 2	Not determined	<5
$NaO^tBu(3)$	145, 2	Not determined	< 5
$KO^t Bu(2)$	145, 2	78:22	65
$KO^t Bu (3)$	145, 2	78:22	73
$KO^t Bu (3)$	65, 15	75:25	51
$KO^t Bu (3)$	145, 7	60:40	67
$KO^t Bu (0.4)$	145, 15	55:45	63
DABCO (3)	145, 72	No reaction	
$Et_3N(3)$	145, 72	No reaction	
$\mathrm{Et_3N}^c$	130, 72	No reaction	
	(equiv.) No base Cs ₂ CO ₃ (3) K ₂ CO ₃ (3) K ₃ PO ₄ (3) LiO'Bu (3) NaO'Bu (3) KO'Bu (2) KO'Bu (3) KO'Bu (3) KO'Bu (3) KO'Bu (3) KO'Bu (3) EO'Bu (0.4) DABCO (3) Et ₃ N (3)	No base 145, 48 Cs ₂ CO ₃ (3) 145, 48 K ₂ CO ₃ (3) 145, 72 K ₃ PO ₄ (3) 145, 72 LiO'Bu (3) 145, 2 NaO'Bu (3) 145, 2 KO'Bu (2) 145, 2 KO'Bu (3) 145, 2 KO'Bu (3) 145, 2 KO'Bu (3) 145, 2 KO'Bu (3) 145, 7 KO'Bu (3) 145, 7 KO'Bu (0.4) 145, 15 DABCO (3) 145, 72 Et ₃ N (3) 145, 72	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

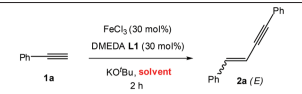
^a E: Z ratios were determined by ¹H NMR analysis of the crude reaction mixture. b Isolated yields after chromatography. C Using Et3N as a solvent.

(entry 7), within 2 hours the reaction afforded the dimer 2a in moderately good yield (65%) with high selectivity (78:22). Use of 3 equivalents of KO^tBu was then used to provide the dimer 2a in good yield (73%) with identical selectivity (entry 8). However, when the reaction was carried out at lower temperature (65 °C), even after 15 h the yield could not reach up to the expectation, although the selectivity was still high (entry 9). After screening of different bases (entries 1-7), potassium tertbutoxide (3 equiv.) was found to be the most effective (entry 7). Our hypothesis to further increase the yield by keeping the reaction for a longer time failed, as both the yield and selectivity of product 2a was decreased with further increase of the reaction time (entry 10). The reaction was found to proceed using 40 mol% KO^tBu (entry 11) to give 2a in 63% yield. No product was detected using nitrogen containing bases like DABCO (entry 12) and Et₃N (entries 13 and 14).

Solvent optimization

Different solvents were then screened with the aim of further improving the reaction outcome (Table 2). Solvents such as THF (entry 1), DMF (entry 2), DMSO (entry 3) and dioxane (entry 4) gave the desired product 2a in low yields with moderate stereoselectivity, whereas no product was detected using CH₂Cl₂ (entry 5) and MeCN (entry 6). Among the studied aromatic solvents (entries 7-9), toluene gave the best results with facile dimerization of 1 with high yield and regioselectivity. Lower yield of the product 2a was obtained using 1,2-dichlorobenzene (entry 8) and xylene (entry 9), albeit a nearly similar

Table 2 Screening of solvents for iron catalyzed dimerization of phenyl acetylene ${\bf 1a}^a$



Entry	Solvent	Temp. (°C)	Ratio ^a	Yield ^b (%) of $2a (E/Z)$
1	THF	85	61:29	67
2	DMF	145	75:25	55
3	DMSO	145	74:26	50
4	Dioxane	145	72:28	45
5	CH_2Cl_2	85	No :	reaction
6	MeCN	145	No :	reaction
7	Toluene	145	78:22	73
8	1,2-Dichlorobenzene	145	76:24	65
9	Xylene	145	75:25	63
10	Neat	145	71:29	45

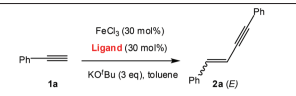
 $^aE:Z$ ratios were determined by 1H NMR analysis of a crude reaction mixture. b Isolated yields after chromatography.

selectivity to toluene as a solvent. Under solvent free conditions the dimer 2a was obtained in 45% yield (entry 10).

Choice of the ligand

With the aim of improving the reaction outcome, various ligands were examined in the iron catalyzed dimerization of phenyl acetylene (Table 3). Surprisingly only iron could drive the selectivity as in the absence of any ligand (entry 1, Table 3), an E-selective dimerization was observed with low yield (42%). We initially screened the nitrogen based ligands (entries 2-8). In all the cases, the use of bulkier ligands resulted in lower yields. When TMEDA (L2) was used instead of DMEDA (L1), the yield slightly decreased (60%). When a chiral ligand cyclohexane-1,2-diamine was used (L3), the selectivity was not at all affected, although the yield was slightly lower, possibly owing to its bulk (entry 4). The product 2a was isolated in moderate yields when aromatic ligands such as 2,2'-bipyridine, L4 (entry 5), and 1,10-phenanthroline, L5 (entry 6), were used. We thought that perhaps the use of the aromatic ligand would facilitate the reaction as the reagents have aromatic rings. But unfortunately, the yield did not improve, rather it decreased even more, presumably because of its higher bulk. 8-Hydroxyquinoline, L6, gave compound 2a with a slightly better selectivity (entry 7); however, the yield was very low (58%). Use of the bathophenanthroline ligand L7 also failed to give good yield (entry 8). When triphenylphosphine, L8, was used as the ligand, the product was obtained in 50% yield with a slightly decreased selectivity (entry 9). Using the bidentate ligand 1,1-bis(diphenylphosphino)methane (dppm) L9, the enyne was isolated in a good yield (85%). Notably, when the phosphene analogue of DMEDA, i.e. 1,1-bis(diphenylphosphino)ethane (dppe) L10, was used, the

Table 3 Iron catalyzed dimerization of phenyl acetylene 1a using different ligands^a



Entry	Ligand (30 mol%)	Ratio ^a (E/Z)	Yield b (%) of 2a
1	No ligand	77:23	42
2	N H L1	78:22	73
3	N L2	75:25	60
4	$ \begin{array}{c} \stackrel{\text{NH}_2}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}{\overset{\text{NH}_2}{\overset{\text{NH}_2}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}{\overset{\text{NH}_2}{\overset{\text{NH}_2}{\overset{\text{NH}_2}{\overset{\text{NH}_2}{\overset{\text{NH}_2}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}{\overset{\text{NH}_2}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}{\overset{\text{NH}_2}}}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}}}{\overset{\text{NH}_2}}{\overset{\text{NH}_2}}}{\overset{\text{NH}_2}}{\overset{N}}}{\overset{N}}}}}}}}}}}}}}}}}}}}}}}}$	77:23	68
5	N L4	75:25	45
6	N L5	76:24	43
7	OH L6	83:17	58
8	Ph Ph L7	77:23	60
9	PPh ₃ L8	71:29	50
10	Ph ₂ P PPh ₂ L9	80:20	85
11	Ph ₂ P PPh ₂ L10	81:19	90
12	PPh ₂ PPh ₂ L11	79:21	70

 $[^]a$ Ratios were determined by 1 H NMR analysis of the unpurified reaction mixture. b Isolated yields after chromatography.

best reaction conditions were achieved with the highest yield (90%) and good selectivity (81:19).

When ligand bulk was increased by introducing 2,2'-bis-(diphenylphosphino)-1,1'-binaphthyl (BINAP) L11, the reaction

yield decreased. All the observations point to the following conclusions. (1) The ligand has a primary role in improving the yield. (2) The selectivity is primarily decided by the catalyst. On the basis of selectivity, reaction times and yields, the best result was achieved using 30 mol% FeCl₃, 30 mol% dppe, and 3 equiv. of KO^tBu in toluene at 145 °C for 2 h.

Substrate scope and limitations

We found that a series of aryl acetylenes 1 smoothly underwent dimerization to produce the head-to-head dimers 2 in good to excellent yield (Table 4). Both strongly electron-donating and electron-withdrawing groups were compatible with this procedure. The yields of the dimerization ranged from 54 to 99% and the ratio of E and Z head-to-head dimers varied from 67:33 to 83:17 in favor of E isomer. Whenever the yields were compared in the presence of DMEDA (L1) and dppe (L10), the yield was the same or higher for L10 over L1. For example, the dimerization of 4-fluorophenylacetylene 1b in toluene gave a 67:33 mixture of the E and Z head-to-head dimers in the presence of L1 with 71% yield (entry 4), whereas in the presence of L10, the yield improved to 80% (entry 5). The 4-bromo phenylacetylene 1c was successfully reacted under the effect of DMEDA (L1) conditions (entry 7) to give the corresponding product 2c in 67% yield. When dimerization of aryl acetylene 1d was carried out in the absence of the ligand, the corresponding product 2d was obtained in 29% yield as a nearly 1:1 ratio of E/Z isomers (data not shown in the table), ²⁸ while in the presence of ligand L1 (entry 8), the product 2d was obtained in an improved yield (55%) and selectivity (E/Z = 67:33). And when ligand L10 was used, the reaction yield even improved (60%, entry 9), which confirms the key role of the ligand in iron catalyzed dimerization.

When m-fluoro 1e was employed (entry 11) under the same conditions, the dimer 2e was obtained in moderate yield (55%). The product 2e was obtained in improved yield (65%), when ligand L10 was used instead of L1 (entry 12). But 1-ethynyl-3,5-difluorobenzene (1f) was converted to the corresponding dimers with a slightly higher yield (59%) than the meta derivative 1e with an E/Z ratio of 3:1 in the presence of L1 (entry 13). When ligand L1 was employed, the ortho fluoro derivative 1g furnished good yield (65%) of the dimerized product 2g (entry 14). The order was 4-fluoro > 4-bromo > 2-fluoro > 3,5-difluoro > 3-fluoro. The orthotrifluoromethyl derivative 1h gave the corresponding dimer in moderate yield (54%, entry 15).

Then a set of methyl substituents were analyzed under the dimerization conditions to get further insight into the reaction. Methyl derivative at para positions of phenyl acetylenes 1i accelerated the reaction (entries 16 and 17) as evidenced by quantitative yield (99%) irrespective of the ligands. The meta analogue 1j resulted in the corresponding dimer in moderate yield (73%) in the presence of L1 (entry 18). So, the order is 4-methyl > 3-methyl. However, a reduced yield (57-58%) was observed with the tert-butyl group at the para position of phenyl acetylene (1k) with a similar 2:1 stereoselectivity in the presence of either L1 or L10 (entries 19 and 20).

The para SMe substituted phenyl acetylene 11 afforded lower yield (60%) of the corresponding dimers 21 in the presence of L1 (entry 21); however, the yield improved (68%) upon employing L10 (entry 22). The methoxy substituent behaved similar to that of the methyl substituted derivatives. 4-Methoxy phenylacetylenes 1m furnished the corresponding dimer 2m in near quantitative yield (E/Z = 2.5:1, entries 23 and 24). Its ortho analogue 10 also resulted in the dimer 20 with a high yield (entries 28 and 29). For para and ortho methoxy derivatives, the yields were similar for both the ligands. However, when the meta derivative 1n was employed, the yield was reduced dramatically to 57% in the presence of L1 (entry 25). The yield was improved to 63% using L10 (entry 26). The yield in the case of 3,5-dimethoxy derivative 1p (entry 30) was lower than that of the 3,5-difluoro derivative 1f in the presence of L1 (entry 13). So, for the methoxy family, the order is 4-methoxy > 2-methoxy \gg 3-methoxy > 3,5-dimethoxy. The para SMe substituted phenyl acetylene 11 afforded a lower yield (60%) of the corresponding dimers 2l in the presence of L1 (entry 21); however, the yield improved (68%) upon employing L10 (entry 22).

The reaction of 2-ethynyl-6-methoxynaphthalene (1q) afforded the desired products in 65% yield with high selectivity (entry 31). Similarly, bulky alkyne 9-ethynylanthracene 1r reacted to afford the enynes in moderate yield (57%) and decreased selectivity (entry 32). Notably, 4-N,N-dimethyl derivative 1s did not proceed when L1 was employed (entry 33); however, it went smoothly in the presence of ligand L10 to give moderate yield (56%, entry 34). Heteroaromatic thiophene alkyne 1t in the presence of ligand L10 gave moderate yield with high E selectivity (entry 35). The reaction of ester 1u (entry 36) and the amide 1v (entry 37) at the para position did not take place in the presence of either of the ligands. Most surprisingly, under the reaction conditions employed, aliphatic alkyne 1w did not yield the corresponding dimer at all (entry 38). When iron(II) chloride was employed as a catalyst instead of iron(III) chloride, only phenyl acetylene reacted to give the dimer in acceptable yields (Table 4, entry 3). Interestingly, none of the phenyl acetylene derivatives 1 could successfully undergo the dimerization reaction using FeCl2 either using L1 or L2. Since the result was independent of the type of the group, i.e. electron donating or withdrawing, the above phenomenon was solely attributed to the steric factor. Notably, the phosphene ligand dppe L10 proved to be more effective than DMEDA L1, even for the iron(II) chloride catalyst. The phenyl acetylene derivatives could undergo dimerization reaction in the presence of dppe as the ligand, although the yield was low (10-23%, entries 6, 10, and 27).

Radical quenching experiment

When 0.5 equiv. of galvinoxyl (radical inhibitor) was added to the reaction system, the yield of 2a significantly dropped to 30% (Table 5, entry 1). The reaction completely stopped in the presence of 1 equiv. of galvinoxyl (entry 2). Similarly in the presence of 2,2,6,6-tetramethylpiperidine 1-oxyl (TEMPO), no dimer 2a was isolated (entry 3). These results indicate that the

 Table 4
 Iron catalyzed dimerization of aryl acetylenes 1^a

Entry	Alkyne 1a	Product	Yield ^a (%), E/Z^b
1 2 3	1a	2a	L1, 73% yield, 78: 22 <i>E/Z</i> L10, 90% yield, 81: 19 <i>E/Z</i> L1, 63% yield, 67: 33 <i>E/Z^c</i>
4 5 6	F—————————————————————————————————————	F—————————————————————————————————————	L1 , 71% yield, 67: 33 <i>E/Z</i> L10 , 80% yield, 71: 29 <i>E/Z</i> L10 , 20% yield, 71: 29 <i>E/Z</i>
7	Br————————————————————————————————————	Br————————————————————————————————————	L1 , 67% yield, 67:33 <i>E/Z</i>
8 9 10	F ₃ C————————————————————————————————————	F_3C \longrightarrow $2d$ \longrightarrow CF_3	L1 , 55% yield, 67 : 33 <i>E/Z</i> L10 , 60% yield, 70 : 30 <i>E/Z</i> L10 , 10% yield, 70 : 30 <i>E/Z</i> ^c
11 12	F 1e	F Ze	L1 , 55% yield, 73 : 27 <i>E/Z</i> L10 , 65% yield, 73 : 27 <i>E/Z</i>
13	F 1f	F F F	L1 , 59% yield, 75 : 25 <i>E/Z</i>
14	7 == 1g	Zg F	L1 , 65% yield, 83 : 17 <i>E/Z</i>
15	CF ₃	2h F ₃ C	L1 , 54% yield, 68:32 <i>E/Z</i>
16 17	Me————————————————————————————————————	Me————————————————————————————————————	L1 , 99% yield, 73 : 27 <i>E/Z</i> L1 , 99% yield, 73 : 27 <i>E/Z</i>
18	Me	Me	L1 , 73% yield, 67 : 33 <i>E/Z</i>

	FeCl ₃ (30 mol%)	R	R
R	Ligand (30 mol%)		
1	KO ^t Bu (3 equiv) toluene 145 °C, 2 h	R 2 (E)	2 (Z)

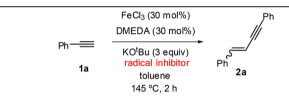
Entry	Alkyne 1a	Product	Yield ^a (%), E/Z^b
19 20)—————————————————————————————————————	<u> </u>	L1 , 57% yield, 73 : 27, <i>E/Z</i> L10 , 58% yield, 75 : 25 <i>E/Z</i>
21 22	MeS————————————————————————————————————	MeS — SMe	L1 , 60% yield, 75 : 25 <i>E/Z</i> L10 , 68% yield, 77 : 23 <i>E/Z</i>
23 24	MeO ————————————————————————————————————	MeOOMe	L1 , 97% yield, 70 : 30 <i>E/Z</i> L10 , 98% yield, 75 : 25 <i>E/Z</i>
25 26 27	MeO	MeO OMe	L1, 57% yield, 75:25, <i>E/Z</i> L10, 63% yield, 77:23, <i>E/Z</i> L10, 23% yield, 75:25, <i>E/Z</i>
28 29	OMe 10	OMe 20 MeO	L1, 91% yield, 71 : 29, <i>E/Z</i> L10, 93% yield, 75 : 25, <i>E/Z</i>
30	MeO	MeO OMe OMe	L1 , 55% yield, 78 : 22 <i>E/Z</i>
31	MeO————	MeO Zq OMe	L1 , 65% yield, 78:22 <i>E/Z</i>
32		2r	L1 , 57% yield, 69:31 <i>E/Z</i>
33 34	$ \begin{array}{c} \text{1r} \\ \text{Me}_2\text{N} \longrightarrow \\ \text{1s} \end{array} $	Me_2N NMe_2 NMe_2	L1 , no reaction L10 , 56% yield, 77 : 28 <i>E/Z</i>
35	S	S Zt	L10 , 57% yield, 71 : 29 <i>E/Z</i>

Table 4 (Contd.)

Entry	Alkyne 1a	Product	Yield ^a (%), E/Z^b
36	MeO_2C	No reaction	
37	Bn-NH	No reaction	
38	C ₅ H ₁₂ -=== 1w	No reaction	

^a Ratios were determined by ¹H NMR analysis of unpurified reaction mixture. ^b Isolated yields after chromatography. ^c Using FeCl₂ (99.99%) as the catalyst.

Table 5 Effect of radical inhibitor



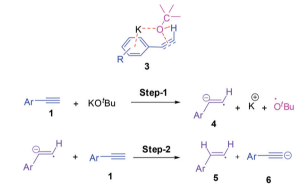
Entry	Additive/equiv.	Ratio of $2^a (E/Z)$	Yield (%) of 2^b
1	Galvinoxyl/0.5	77:23	30
2	Galvinoxyl/1	No	reaction
3	Tempo/1	No	reaction

 $[^]a$ Ratios were determined by 1 H NMR analysis of the unpurified reaction mixture. b Isolated yields after chromatography.

radical process may be a possible pathway in this reaction, unlike the protocol reported previously by others in the presence of other catalysts. On the basis of all the above observations, a plausible mechanism is proposed.

Mechanism

We propose that the potassium tertiary butoxide interacts with the conjugated system of aryl acetylene through cation-pi interaction and pi-pi interaction as shown in the transition state 3 to promote the reactivity of the inactive aryl acetylene (Scheme 3). Subsequently, the increased reactivity of the transition state 3 results in two important species, one tertiary butoxide radical and another radical anion 4. The radical anion 4 then reacts with the second molecule of aryl acetylene 1 and abstracts the acidic hydrogen to give the corresponding alkene radical 5 and the aryl acetylide 6 (Scheme 3). The reaction yields employing bases of similar structures validate the



Scheme 3 First two steps of the iron dimerization reaction.

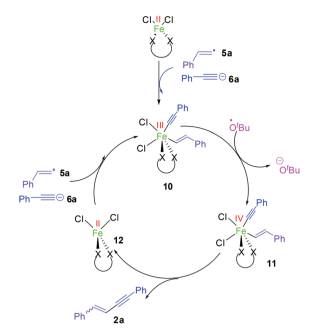
formation of transition state 3 (Table 1). In addition to potassium tertiary butoxide, potassium carbonate (K-O-), cesium carbonate (Cs-O-) and potassium phosphate (K-O-) also proceed through the mechanism to give the corresponding product. So, bases sharing bond connectivity similar to potassium tertiary butoxide undergo the reaction. Notably, NaO^tBu and LiO'Bu did not yield anything, which can be explained as follows. Both the phenyl acetylene and toluene have an aromatic ring; however, toluene being the solvent molecule has a much higher probability of forming cation-pi interaction with the cations. Smaller cations such as lithium and sodium have the highest efficiency of forming cation-pi interaction;³⁰ also they are most easily solvated. So, the interplay between cationpi interaction energy and desolvation energy decides the most effective catalyst among all. The high desolvation energy restricts their binding to the benzene ring of the phenyl acetylene, which explains the inability of the tertiary butoxide salt of lithium and sodium to push the reaction forward. Organic

Scheme 4 Iron(III) cycle for the dimerization of aryl acetylene.

bases having structures different from the above class like DABCO and Et₃N did not yield anything.

Notably, only aryl acetylene gave the enyne product and the aliphatic acetylene did not. We think that the transition state 3 solely decides which reactant to favor. As the observations suggest, the cation-pi interaction plays an important role in stabilizing the transition state. In the case of aliphatic acetylene, the cation-pi interaction is absent, which then inhibits further progress of the reaction.

In the presence of iron(III) chloride, the outcome of Step 2, i.e. the alkene radical 5 and aryl acetylide 6 attack to form the reactive species 7 (Scheme 4). Subsequently, this tetravalent hexacoordinated species 7 undergoes reductive elimination to give the Fe(II) species 8 and the desired enyne product 2. The tertiary butoxide radical further abstracts an electron from the divalent iron center 8 to generate the corresponding Fe(III) species 9 and the alkene radical 5. The aryl acetylide 6 then attacks the tetra-coordinated species 9 regenerating 7 to complete the cycle. The yield is based on the hypothesis that more the nucleophilicity of the acetylide, more the probability of attacking the Fe(III) center to regenerate the catalyst, and hence the better is the yield. So, in accordance with our proposition, phenyl acetylene bearing ortho or para electron donating groups almost gives the corresponding product in a quantitative amount. The phosphene ligand gave better results for alkyne containing electron withdrawing groups, which also corroborates our hypothesis. Electron withdrawing groups at ortho and para positions decrease its ability to attack and coordinate to the metal center. The strong pi-acceptor ligand dppe compensates for it. Being attached to the metal center, it increases the effective local charge (oxidation state) of the metal, to which the radical 5 and anion 6 components containing electron withdrawing groups at ortho and para positions can



Scheme 5 Iron(II) cycle for the dimerization of phenyl acetylene 1a.

efficiently attack, and the better is the yield. Thus, the ligand plays a crucial role in carrying out the reaction.

The inability of iron(II) chloride to dimerize derivatives of phenyl acetylene also fits our model (Scheme 5). In the case of Fe(II) chloride, the alkene radical 5a and the aryl acetylide 6a attack to form the Fe(III) species 10. The generated tertiary butoxide radical subsequently abstracts an electron from the Fe(III) species 10, thus resulting in the formation of the tetravalent iron species 11. Reductive elimination from 11 gives the enyne product 2a. The alkene radical 5a and aryl acetylide 6a subsequently attacks the tetra-coordinated species regenerating 10 to complete the cycle.

Then the question arises: why does only phenyl acetylene give the envne product in the presence of FeCl2 but its derivatives do not? The answer behind this is the steric crowding. In the case of the Fe(III) catalytic cycle (Scheme 4) the tertiary butoxide radical abstracts an electron from a tetra-coordinated species 8, whereas in the case of the Fe(II) catalytic cycle the tertiary butoxide radical has to abstract the electron from a hexacoordinated species 10 (Scheme 5).

Whenever we employ phenyl acetylene derivatives (Scheme 6), we effectively increase the steric bulk of the complex (irrespective of the electron donating or withdrawing groups), which makes it even more difficult or impossible for the tertiary butoxide radical to abstract the electron from 13 to form 15. As a result, the Fe(II) species interacts with the alkene radical to generate a Fe(III) species 13, which on reductive elimination has to form a Fe(1) species 14, which is neither stable nor favorable. When the phosphene ligand was used instead of DMEDA, low yield was observed. In the case of dppe L10, either the alkene radical or aryl acetylide interacts with the benzene ring of the ligand via weak pi-pi stacking, which brings either of the groups closer to the ligand, finally

Scheme 6 Schematic diagram for the two unlikely options for the dimerization of phenyl acetylene derivatives in the iron(II) cycle.

resulting in the availability of space for tertiary butoxide radical to attack the Fe center and abstract the electron. As a result, the reaction proceeds, but with a low yield.

Conclusions

In summary, we have described novel iron catalyzed regio- and stereoselective dimerization of terminal aryl alkynes to give the corresponding head-to-head dimers in moderate to excellent yields. Phenyl acetylene bearing ortho or para electron donating groups almost gave the corresponding product in quantitative yield. Mechanistic evidence suggests that the transformation proceeds through the activation of aryl acetylene via cation-pi interaction and pi-pi interaction with potassium tertiary butoxide to produce alkene radical and aryl acetylide. In the presence of iron(III) chloride, the alkene radical and aryl acetylide attacks to form the reactive tetravalent hexacoordinated species which then undergoes reductive elimination to give the desired enyne product 2. Notably, the phosphene analogue dppe gave marginally better results as compared to DMEDA. We further identified that FeCl2 could only promote the dimerization of phenyl acetylene. This novel catalytic system provides an alternative to toxic and expensive transition metals for a variety of conjugated enyne compounds.

Experimental

General information

All experiments were carried out under an inert atmosphere of argon in flame-dried microwave vials. Solvents were dried using standard procedures. Starting materials were obtained from commercial suppliers and used as received. Products were purified by flash chromatography on silica gel (100-200 mesh). NMR spectra were recorded in CDCl₃. ¹H NMR spectra were recorded at 500 MHz and 400 MHz instruments at 278 K. Signals are quoted as δ values in ppm using residual protonated solvent signals as the internal standard (CDCl₃: δ 7.26 ppm). Data are reported as follows: chemical shift, integration, multiplicity (s = singlet, d = doublet, t = triplet, q = quartet, m = multiplet), and coupling constants (Hz). ¹³C NMR spectra were recorded on either a JEOL-400 (100 MHz) or a Brüker AVANCE 500 MHz (125 MHz) with complete proton decoupling. Chemical shifts (δ) are reported in ppm downfield from tetramethylsilane with the solvent as the internal reference (CDCl₃: δ 77.26 ppm). Infrared (FTIR) spectra were recorded by the KBr disk and KBr plate techniques for solid and liquid samples, $\nu_{\rm max}~{\rm cm}^{-1}$. HRMS analyses were performed by +ve mode electrospray ionization.

General procedure for dimerization of aryl acetylenes 1

To a 10 mL oven dried microwave vial were added KO^tBu (3 equiv.), anhydrous toluene (4 mL mmol⁻¹) and anhydrous FeCl₃ (0.3 equiv.). Then, the resulting mixture was submitted to argon/vacuum cycles stirring under an atmosphere of argon. Subsequently DMEDA L1 (0.3 equiv.) or dppe L10 (0.3 equiv.) followed by aryl acetylene 1 (1 equiv.) were added. Then, the resulting mixture was stirred at 145 °C for 2 h. After cooling, the solvent was evaporated and the crude residue was purified by flash column chromatography on silica gel using distilled *n*-hexanes–EtOAc (100:0 to 90:10) as an eluent to give the desired enynes 2. The ¹H and ¹³C NMR spectroscopic data of the enyne compounds 2a–g,²⁸ 2h,^{13b} 2i–j,²⁸ 2k (E),^{13f} 2k (Z),³¹ 2m–2q,²⁸ 2r (E),^{17d} 2r (Z),³² 2s (E),^{14b} 2s (Z),³¹ 2t (E),¹⁷ⁱ and 2t (Z)^{18d} matched with those reported in the literature (see ESI‡).

(4,4'-(But-1-en-3-yne-1,4-diyl)bis(4,1-phenylene))bis (methylsulfane) 2l

Light yellow oil; ESI-MS: calcd for $C_{18}H_{16}S_2$ [M]⁺: 296.0693; found, 296.0767. (*E*)-isomer (2**I**): FT-IR (neat): 3401, 3016, 2925, 2853, 2395, 2193, 1934, 1607, 1581, 1486, 1268, 1178, 1152, 1075; ¹H NMR (400 MHz): 7.38–7.32 (m, 4H), 7.21–7.17 (m, 4H), 6.97 (d, J = 15.9 Hz, 1H), 6.33 (d, J = 16.5 Hz, 1H), 2.49 (s, 6H); ¹³C NMR (100 MHz): 140.4, 139.3, 139.2, 133.1, 131.7, 126.6, 126.3, 125.8, 119.6, 107.2, 91.7, 89.1. (*Z*)-isomer (2**I**): data from a mixture of E/Z = 84 : 16; ¹H NMR (400 MHz): δ = 7.85 (d, J = 8.5 Hz, 2H), 7.41–7.33 (m, 2H, merged with *trans* isomer), 7.23–7.17 (m, 4H, merged with *trans* isomer), 6.63 (d, J = 12.2 Hz, 1H), 5.86 (d, J = 11.6 Hz, 1H), 2.51, (s, 3H), 2.50 (s, 3H).

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