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Hydroelementation of diynes†

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This review highlights the hydroelementation reactions of conjugated and separated diynes, which depending on the process conditions, catalytic system, as well as the type of reagents, leads to the formation of various products: enynes, dienes, allenes, polymers, or cyclic compounds. The presence of two triple bonds in the diyne structure makes these compounds important reagents but selective product formation is often difficult owing to problems associated with maintaining appropriate reaction regio- and stereoselectivity. Herein we review this topic to gain knowledge on the reactivity of diynes and to systematise the range of information relating to their use in hydroelementation reactions. The review is divided according to the addition of the E–H (E = Mg, B, Al, Si, Ge, Sn, N, P, O, S, Se, Te) bond to the triple bond(s) in the diyne, as well as to the type of the reagent used, and the product formed. Not only are the hydroelementation reactions comprehensively discussed, but the synthetic potential of the obtained products is also presented. The majority of published research is included within this review, illustrating the potential as well as limitations of these processes, with the intent to showcase the power of these transformations and the obtained products in synthesis and materials chemistry.

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^c Cardiff Catalysis Institute, Cardiff University, School of Chemistry, Park Place, Main Building, Cardiff CF10 3AT, Cymru/Wales, UK. E-mail: MelenR@cardiff.ac.uk † This article is dedicated to Prof. Bogdan Marciniec from Adam Mickiewicz University in Poznań (Poland), expert in hydrosilylation reactions, on the occasion of his

80th birthday.



Jędrzej Walkowiak

Jędrzej Walkowiak received his PhD degree (maxima cum laude) in 2009 from Adam Mickiewicz University in Poznań (Poland) with Prof. B. Marciniec. He completed postdoctoral research in Prof. W. Leitner group at the RWTH Aachen in Germany. In 2019, he obtained habilitation in chemical sciences. From 2011 he has been employed at the Center for Advanced Technology, at Adam Mickiewicz University in Poznan, first as an Assistant

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1. Introduction

Hydroelementation reactions are one of the most prominent transformations in organic and organometallic chemistry, to obtain functionalised compounds from the addition of E-H bonds (E = Mg, B, Al, Si, Ge, Sn, N, P, O, S, Se, Te) to unsaturated C-C bonds in olefins (C=C) or alkynes (C=C), $^{1-31}$ C-N bonds in imines $(C=N)^{2,32-36}$ or nitriles $(C=N)^{2,33,34,37}$ and C=O bonds in carbonyl compounds.^{2,32–35,38–47} The processes are mostly catalytic but may also occur as uncatalysed. In both cases, the stereo- and regioselectivity of the reaction depends upon the catalyst, reagent, and reaction conditions.

Hydroelementation of alkynes is perhaps the simplest, most straightforward, and atom economic method for the synthesis of unsaturated organometallic or organometalloid compounds. Over the last few decades, several reviews have been published focusing on this subject.^{14,17,18,22,25,30,48-61} Particularly useful are hydroboration, hydrosilylation, hydroamination, hydrophosphination, and hydrostannation processes, which lead to important building blocks in organic and materials chemistry. Although many different terminal and internal alkynes have been used in these transformations, literature focused on the hydroelementation of conjugated or separated divnes is much rarer and has never been collated in a review before. The more complex structure of diynes together with the possibility to obtain various isomers or different products (e.g., enynes, dienes, allenes, heterocyclic compounds, polymers), as well as the problems with carrying out monohydroelementation or bishydroelementation selectively, define the complexity of these processes (Scheme 1). For example, as was described by Perry et al., the hydrosilylation of conjugated 1,3-diynes may lead to the formation of nine different products.⁶² The difficulties in distinction in the reactivity of both $C \equiv C$ bonds and the potential for overreduction are the most problematic issues reported. Actually, because of the synthetic potential of diyne hydrometallative products in the production of natural compounds, pharmaceuticals, or highly conjugated materials, within the last two decades, the subject is getting more explored.63-71

Working on the hydroboration and hydrosilylation of various unsaturated compounds and especially on the reactivity of conjugated 1,3-divnes in these processes, we have found that literature information is often scattered, with no detailed procedures or much discussion on the process optimisation or methodology.⁷²⁻⁸⁷ Therefore, we have decided to build a comprehensive and critical compendium focused on this subject, which will systemise the existing knowledge on the hydroelementation of diynes in relation to the formation of different products. We will also show the possible applications of the obtained products in the synthesis of fine chemicals and materials. The review is divided into subchapters according to the type of hydroelementation reactions according to the element group of the periodic table of elements: hydromagnesation, hydroboration, hydroalumination, hydrosilylation, hydrogermylation, hydrostannation, hydroamination, hydrophosphination, hydration, hydrothiolation, hydroselenation, and hydrotelluration. Each hydroelementation reaction type is then subdivided into conjugated and separated diynes, and the type of product formed (enynes, dienes, heterocyclic compounds, and polymers). This comprehensive review will be helpful for all advanced researchers and newcomers working on the synthesis of organometallic compounds and their further applications in organic chemistry, synthesis of natural compounds, pharmaceuticals, with the emphasis placed on the process regio- and stereoselectivity.

2. Hydromagnesation

Hydromagnesation of conjugated 1,4-diaryl-1,3-diynes 2.1.

Hydromagnesation of alkynes is just limited to a few examples, which were performed in the presence of nickel, titanium, or



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applications of main group chemistry in organic synthesis.

diverse aspects of main group reactivity and catalysis, including the



E = Mg, B, Al, Si, Ge, Sn, N, P, O, S, Se, Te

Scheme 1 Possible products obtained from the hydroelementation of conjugated 1,3-diynes and separated 1,*n*-diynes. Only selected products and their isomers are presented.



iron complexes.88-95 Hydromagnesation of conjugated and separated divnes was reported only by Nakamura et al., who synthesised alkenylmagnesium compounds using EtMgBr 2 as a hydrogen source and FeCl₂ 3 as a catalyst.⁹⁶ The reaction occurred with 1,2-diarylalkynes and 1,3-diynes 1a-e with high (Z)-selectivity, in short reaction time (15 min), at room temperature. For the reaction 5 mol% of an iron catalyst was used. Under the applied reaction conditions, the alkenes were unreactive, so no overreduction was observed. Moreover, only one $C \equiv C$ bond in the diynes 1a-e was converted to the magnesium derivative 4 or 5 (regioisomers), which then was treated with an electrophile (HCl, D₂O, allyl bromide, or DMF) giving products with H, D, allyl or CHO groups respectively 6, 7 (Scheme 2 and Table 1). Other iron complexes (e.g., FeCl₃ 8, $Fe(acac)_2$ 9, $Fe(acac)_3$ 10) were also active in this transformation but gave products with lower yields. Primary alkyl magnesium derivatives were also active in this reaction (except for bulky isobutylmagnesium bromide), as well as secondary alkyl Grignard compounds (with cyclohexyl or cyclopentyl groups). The magnesium compound had to be used in a 2.0 to 2.5-fold excess to obtain high conversion of alkyne or diyne, but the necessity of

 Table 1
 Hydromagnesation of conjugated 1,3-diynes 1a-e with ethylmagnesium bromide 2 followed by the electrophilic substitution

Entry	$R^1 = R^2$	1	Electrophile	6- <i>E</i> / <i>Z</i>	6:7	Isol. yield of 6 [%]
1	Ph	1a	D^+	14:86	>99:1	63
2	Ph	1a	DMF	97:3	>99:1	50
3	$4-FC_6H_4$	1b	D^+	11:89	97:3	63
4	4-MeC ₆ H ₄	1c	D^+	25:75	97:3	55
5	4-MeOC ₆ H ₄	1d	D^+	22:78	97:3	65
6	$R^{1} \neq R^{2}$ $R^{1} = Me_{3}Si, R^{2} = Ph$	1e	DMF	17:83	98:2	55

this excess was not clear. The source of hydrogen was from the magnesium compound, which was determined from the reaction of 1,2-diphenylethyne with deuterated d_5 -ethylmagnesium bromide. Both diynes with electron-donating and electron-withdrawing groups attached to aryl rings were transformed into enynes with high regio- and stereoselectivity. The stereoselectivity was slightly lower for electron-rich reagents **1c** and **1d** (Table 1, entries 4 and 5). When the unsymmetrically substituted diyne with phenyl and trimethylsilyl groups **1e**



Scheme 3 Selective hydromagnesation of unsymmetrically substituted 1,4-diethynylbenzene **11** with ethylmagnesium bromide **2** catalysed by FeCl₂ **3**.

was used, the addition of magnesium compound 2 occurred at the $C \equiv C$ bond to which the phenyl ring was attached (Table 1, entry 6). Moreover, the lack of reactivity of alkyl-substituted alkynes permitted the selective functionalisation of 1-(3,3-dimethylbut-1-yn-1-yl)-4-(phenylethynyl)benzene **11**, which reacted at the diarylalkyne site. Only a small amount (7%) of diene was formed in the reaction mixture as a side product (Scheme 3). As it was shown for hydromagnesation of alkynes, the catalytic system tolerates many functional groups (*e.g.*, halogens, amines, phenoxide, alkenes).

3. Hydroboration

Organoboron compounds constitute important building blocks in the synthesis of structurally advanced organic and organometallic compounds due to their versatile reactivity in many catalytic and noncatalytic couplings and deborylation reactions, together with their low toxicity and moderate stability. There are numerous papers focused on the synthesis and applications of organoboron compounds, especially arylboronic acids and vinyl boranes.^{97–110} The hydroboration reaction is still the most important and useful transformation in the synthesis of boranes because of its straightforward procedure, 100% atom economy, the possibility to control process regio- and stereoselectivity by the application of a catalyst or modification of the reagent steric properties or process conditions. The hydroboration of monoalkynes furnishing important alkenylborane building blocks is well established in the literature and has been discussed in several reviews.^{1,2,4,7,10,12,48} Hydroboration of conjugated and separated diynes, because of the increased complexity of their structure, is much more challenging in the case of selectivity control. Moreover, the possibility for carrying out mono-, bis-hydroboration, polyaddition reactions and cyclisation reactions with these reagents creates the possibility to obtain various products, which have been used in the synthesis of natural compounds, pharmaceuticals (*e.g.*, anticancer rizoxin D, cytotoxic nannocystin Ax, ivorenolides),^{111–119} dyes,¹²⁰ π -conjugated compounds, or heterocycles.^{121–123} The information in this section is divided according to the type of reagent used: conjugated or separated diynes, as well as the formation of different products: enynes, dienes, heterocyclic compounds, and polymers.

3.1. Hydroboration of conjugated 1,3-diynes

Hydroboration of conjugated 1,3-diynes is the simplest procedure for the synthesis of boryl-substituted 1,3-enynes or bisboryl-substituted 1,3-dienes, but the transition metalcatalysed selective addition of the B–H bond to the C \equiv C bond is limited only to three recently published examples.^{72,124,125}

The first selective hydroboration of 1,3-diynes was reported by Zweifel and Ponso. Noncatatalytic reduction of alkylsubstituted buta-1,3-diynes 13a-c was carried out using disiamylborane (bis(3-methyl-2-butyl)borane) 14 or less bulky dicyclohexylborane 15. The monohydroboration of dodeca-5,7-diyne 13a, 2,7-dimethylocta-3,5-diyne 13b and 2,2,7,7tetramethyl-octa-3,5-diyne 13c with disiamylborane 14 (used in 1.1-fold excess) occurred with high regio- and stereoselectivity at 0-5 °C within 3 h. Protonolysis of the enynylborane intermediate 16a (obtained in the reaction of 13a with 14) with acetic acid at 55-60 °C for 5 h furnished (Z)-5-dodecen-7-yne 17a (83%), (Z,Z)-dodeca-5,7-diene (7%), and only traces of unreacted diyne 13a. Bulkier diyne 13c yielded the bishydroboration products only in trace amounts. Oxidation of the monohydroboration products with NaOH/H2O2 (30%) afforded the acetylenic ketones **18a–c** in high yields >70% (Scheme 4).

The boryl group was attached to the external carbon atom, which was proved by protonolysis with deuterated acetic acid. NMR analysis indicated that deuterium was attached to the internal carbon atom (95(\pm 3%)). Synthesis of (*Z*,*Z*)-bisborylated dienes was more effective when less hindered borane **15** was



Scheme 4 Non-catalytic hydroboration of symmetrical dialkyl-substituted 1,3-diynes 13a-c with disiamylborane 14 and dicyclohexylborane 15 followed by protonolysis or oxidation towards enynes 17a-c or α,β -acetylenic ketones 18a-c.



Scheme 5 Non-catalytic hydroboration of 1,4-dialkyl 1,3-diynes **13a-d** with boranes **14** and **15** towards (*Z*,*Z*)-dienes: (a) two-step method when **14** was used; (b) one-step procedure for **15**.



Scheme 6 Functionalisation of silyl-substituted 1,3-diynes 20a-d with boranes 14 and 15, and further synthesis of silyl-substituted enynes 22, terminal enynes 23, and silyl-functionalised ketones 24.

used and could also be reacted with the non-symmetrically substituted diyne *e.g.*, 2-methyldodeca-3,5-diyne **13d**. In the case of hindered diynes **13b** and **13c**, the reduction was carried stepwise using borane **14** to obtain (*Z*,*Z*)-diene, by the hydroboration protonolysis of the first triple bond and the subsequent repetition of these processes. The application of **14** instead of **15** was caused due to the fact that dienes **19** formed in the reaction with **15** have a similar boiling point to the side product cyclohexanol, making the distillation method ineffective for their separation (Scheme 5). The analogous experiment with deuterium labeling proved that the second boryl group is attached to the external carbon atom of the side of the firstly attached boryl group.¹²⁶

The same group reported that for unsymmetrically substituted 1,3-diynes with silyl and alkyl or cycloalkyl groups attached to the opposite sides of **20a-d**, monohydroboration predominantly occurred at the $C \equiv C$ bond without the silyl group. Application of the more hindered borane **14**, as well as the structure of silyl groups (trimethyl, (*tert*-butyl)dimethyl, dimethylthexyl) influenced the reaction selectivity. For symmetrical 1,4-bis(trimethylsilyl)-1,3-butadiyne, the addition of sterically hindered **14** occurred at positions C^1 and C^2 in the ratio 26:74. The high regioselectivity of the reaction towards the product with borane at C^2 position was obtained when the bulky dimethylthexylsilyl substituent was attached in **20c** and $20d.^{127}$ The obtained products were further transformed into silyl-functionalised ketones 24 or enynes 22–23 with the above-described procedure (Scheme 6).¹²⁶

These alkylboryl-substituted enynes (16, 21) are difficult to handle due to their low stability, therefore the hydroboration of diynes with alkoxyboranes (e.g., pinacolborane 25 or cateholborane 26) is much more desirable. Moreover, alkoxyboranes are easy to use and non-flammable. However, due to the lower acidity of the B-H bond in comparison to alkylboranes, the addition of alkoxyboranes to unsaturated $C \equiv C$ bonds requires the application of a catalyst to accelerate the process. Relating to this, our recent paper focused on the selective monohydroboration of 1,4-diaryl-buta-1,3-diynes 1a-d, 27a-c with pinacolborane 25 in the presence of $Ru(CO)Cl(H)(PPh_3)_3$ 28. Compound 28 has previously been described as an active catalyst in hydroboration of terminal- or internal monoalkynes in conventional and novel, green reaction media (supercritical CO2 (scCO₂), ionic liquids (ILs), polyethylene glycol, (PEG)).^{72,75,76,128,129} The reaction proceeded effectively for various diynes possessing electron-withdrawing or electron-donating substituents on the aryl ring, as well as for heterocyclic 1,4-di(thiophen-3-yl)buta-1,3-diyne 27c. Alkyl-substituted diynes yielded boryl-substituted enynes by *cis*-addition of borane to the $C \equiv C$ bond, but the postreaction mixture also consisted of other monoborylated enynes, bisborylfunctionalised dienes, and some undefined products. Thus, the electronic properties of diynes have an important influence on



Ru(CO)Cl(H)(PPh₃)₃ catalyst 28

the process regio- and stereoselectivity. Under the optimised reaction conditions (3 mol% of Ru(CO)Cl(H)(PPh₃)₃ **28**, 60 °C, 24 h, with a small excess of borane **25** (1.2 equiv.)), the boryl-substituted enynes **29** were obtained with high yields (85–97%). Due to their instability during purification by column chromatography, the products were directly transformed to the corresponding stable trifluoroborane salts **31** with KHF₂ **30** furnishing the desired products with 75–84% yield (Scheme 7).

The regioselectivity of the process was confirmed by NOESY and X-ray diffraction analysis for product **29** obtained in the hydroboration of **27b** with **25** (Fig. 1). The borane **25** was added to the $C \equiv C$ bond in a *syn*-manner according to the anti-Markownikow rule, with the boron group attached to the less shielded internal carbon atom in **29**. We have also proposed the mechanism of the process according to the stoichiometric reactions monitored by ¹H NMR and 1D selective gradient NOESY. The process initiates from the insertion of diyne **1a–d** or **27a–c** into the Ru–H bond of catalyst **28** forming but-3-en-1-yn-3-yl complex **32**. The addition of borane **25** then leads to a σ -bond metathesis between Ru–C and B–H (**33**), followed by the elimination of the product **29** and regeneration of the initial Ru-hydride complex **28** (Scheme 8 and Fig. 1).

Fig. 1 Molecular structure of 2-(4,4,5,5,-tetramethyl-1,3,2-dioxaborolanyl)-1,4-bis(4-*tert*-butylphenyl)but-1-en-3-yne **29** obtained in the hydroboration of **27b** with **25** in the presence of catalyst **28**.⁷²



Scheme 8 Proposed catalytic cycle for the hydroboration of 1,3diynes 1a-d, 27a-c with pinacolborane 25 in the presence of $[Ru(CO)Cl(H)(PPh_3)_3]$ 28.



Scheme 9 Suzuki–Miyaura coupling of enynes 29a and 31a with iodobenzene 34.

The utility of the resulting boryl-substituted 1,4-diaryl-but-1en-3-ynes was presented in the Suzuki coupling reaction of pinacoloborane derivative **29a** and trifluoroborate salt **31a** with iodobenzene **34** using 5 mol% of Pd(PPh₃)₄ **35**. The reaction occurred with the retention of the configuration and (*Z*)-1,2,4triphenylbut-1-en-3-yne **36a–b** was formed with high yields 71% and 78%, respectively (Scheme 9).⁷²

Ge *et al.* reported an interesting method for the synthesis of boryl-substituted enynes from unsymmetrical and symmetrical 1,3-diynes **1c**, **1e**, **37a-t** in the presence of a cobalt catalyst

generated from inexpensive and stable Co(acac)₂ 38 and bidentate phosphine ligands 39a-c. The authors showed that the regioselectivity of the process was dependent on the bidentate phosphine ligand used. When Co(acac)₂/xantphos 38/39a was used, enynes with boron groups attached to the internal carbon atom were formed 40. However, when applying dppf 39b as a ligand, the opposite regioselectivity was observed, furnishing the product functionalised with borane at the external position 41. The pinacolborane derivatives were further transformed to more stable 1,8-diaminonaphthalene boronates 43 and 44 respectively, with 1,8-diaminonaphthalene 42, which were easier to isolate (Scheme 10).¹²⁵ To find the answer to this different reactivity pathway, the authors carried out a reaction with deuterated DBpin 45 (Scheme 11a). The formation of products with different regioselectivity was also confirmed using 1D NOE and 2D HMBC NMR correlations. For the Co(acac)₂/xantphos 38/39a catalyst, the reaction occurred through the formation of Co-H intermediate 48, while for Co(acac)₂/dppf 38/39b the process proceeded through the Co-borane species 50 (Scheme 11b). For both catalytic systems, several borylsubstituted envnes 40 and 41 were formed using an equimolar amount of reagents. When silyl groups were attached to one alkyne, the borane was added to the second triple bond with different (aryl, heteroaryl, or alkyl) substituents. The products were obtained with excellent regioselectivity and yield. For alkylsubstituted diynes, better selectivity was obtained when L-N₃ 39c was used as a ligand instead of xphos 39a. The catalytic systems tolerate a lot of functional groups in the divne structure. No significant changes in their reactivity were observed (Scheme 10).¹²⁵ The utility of the resulting boryl-functionalised 1,3-envnes was presented in the bromodeborylation reaction with CuBr₂ 52, as well as the Pd-catalysed Suzuki-Miyaura and Hiyama coupling reactions (Scheme 12).

Applying a CuCl 55/P(p-Tol)₃/NaOt-Bu catalytic system, it was possible to carry out formal hydroboration of symmetrical and unsymmetrical 1,3-diynes 1a-b, 1d-e, 13a, 13c, 27a, and 60a-e with bis(pinacolato)diboron 61 and methanol as a proton source. The process was carried out under strictly assigned conditions (11 °C for 6 h) with 5 mol% of CuCl 55, 6 mol% of phosphine, and 10 mol% of the base. Reactions with weaker donating phosphines (P(OEt)₃ or PPh₃) resulted in lower yields and longer reaction times, with some exceptions. For diynes with tert-butyl 13c, 4-methoxyphenyl 1d, cyclohexyl 60a groups, P(OEt)₃ was applied, while for 4-fluorophenyl-substituted divne 1b, PPh₃ was successfully used. The boryl group was attached at the external carbon position of the 1,3-diyne and only for sterically hindered 2,2,7,7-tetramethyl-octa-3,5-diyne 13c the regioselectivity was reversed and the borane was bonded to the less shielded internal carbon atom. When hydroboration of unsymmetrically functionalised diynes with aryl and alkyl groups in the terminal position was carried out, the reaction occurred at a more accessible $C \equiv C$ bond. Divnes bearing a silvl group were characterised by their strong directing effect, where the functionalisation proceeded at the triple bond situated further from the silvl group.¹²⁴ The same observation was noticed in the noncatalytic hydroboration of silyl-substituted diynes.¹²⁷ The resulting pinacolborane derivatives **62** were then transformed to their potassium trifluoroborate analogs **63** with KHF₂ **30**. When unsymmetrical 1,4-diaryl-diyne was used, a 1:1 mixture of regioisomers was formed due to the similar reactivity of both $C \equiv C$ bonds (Scheme 13). Using this catalytic system, different regioisomers were formed with the boryl group attached to the internal carbon bond in comparison to the previously described works on noncatalytic or Ru–H catalysed hydroboration.^{72,126,130} The obtained products were subsequently derivatised by Suzuki-Miyaura coupling with iodobenzene **34** using PdCl₂ **64**/dppf **39b** as a catalyst and KOH as a base, as well as deborylated to enynes with acetic acids. Moreover, it was possible to carry out selective desilylation of silylboryl-substituted enynes with K₂CO₃/MeOH while Bpin remained unreactive under applied process conditions.

Very recently, Taniguchi and co-workers reported the first trans-hydroboration of 1,3-diyne derivatives 60a and 65a-l under radical conditions in the presence of AIBN (2,2'azobis(isobutyronitrile)) 67 or ACCN ((1,1'-azobis(cyclohexane-1-carbonitrile))) 68 as an azo initiator and tert-dodecanethiol 69 (TDT) as a polarity-reversal catalyst. The addition of an Nheterocyclic carbene borane 66 to symmetrical and unsymmetrical 1,3-diynes gave (E)-alkynylalkenyl boranes 70 in high selectivity (E/Z = 95/5) and good isolated yields (51-77%). Interestingly, a 4-fold excess of the borane 66 with diynes 65a-l and 60a caused the formation of bisadducts in inconsiderable amounts. The protocol was suitable for 1,3-diynes with different substituents (n-alkyl, c-alkyl, propargyl ether, silyl) however, hydroboration of 1,4-diphenylbuta-1,3-diyne 1a under the standard conditions gave a mixture of undefined products due to different rate of hydrogen atom transfer for aryl and alkyl or silvl substituted 1,3-diynes. It is worth noting that obtained NHC-based boryl functionalised enynes 70, in contrast to pinacolborane-based enynes, are bench-stable compounds and can be easily purified by silica-column chromatography (Scheme 14). The authors proposed the mechanism of this antiselective hydroboration of 1,3-diynes which in the first step involved the thermal decomposition of azo initiator AIBN 67 or ACCN 68 and formation of thivl radical 71 from thiol 69. The abstraction of the hydrogen atom from NHC-borane 66 by the radical 71 yielded NHC-boryl radical 72. Subsequently, the regioselective addition of boryl-radical 72 to 1,3-diyne 60a, 65a-l gave alkenyl radical 73 conjugated to the alkyne moiety. The presence of thiol TDT 69 promoted the hydrogen atom transfer step and formation of thiyl radical 71 which abstracted hydrogen atom from NHC-borane 66 and closed radical chain. The authors suggested that the bisadducts 74 would not be efficiently formed since the presence of electron-rich NHCborylalkenyl group of 73 would cause polarity mismatching during the addition of nucleophilic NHC-boryl radical 72 (Scheme 15). 70a could be easily converted to the corresponding (Z)-1-aryl-1,3-enyne derivatives 77a and 77b through a one-pot procedure involving chlorination and hydrolysis of the boron moiety with N-chlorosuccinimide (NCS) 78 and water followed by a Suzuki-Miyaura coupling with aryl iodides 75 and 76 (Scheme 16).¹³¹



Scheme 10 Selective hydroboration of conjugated dignes 1c, 1e and 37a-t using $Co(acac)_2$ 38 as a catalyst and different ligands: path A – xantphos 39a or L-N₃ 39c; path B – dppf 39b.



Scheme 11 (a) Mechanistic study on the hydroboration of conjugated diynes with **38/39a** and **38/39b** with DBpin **45**; (b) proposed catalytic cycles for Co-catalysed regiodivergent hydroboration of 1,3-diynes.

3.2. Hydroboration of separated 1,n-diynes

The hydroboration of separated 1,*n*-diynes leads to the formation of various products from boryl-substituted enynes and bisboryl-substituted dienes, towards cyclic products, which are important building blocks in the synthesis of natural compounds, that can be used in *e.g.*, Diels–Alder reactions. The selectivity of the hydroboration processes depends on the catalyst type and reagent structure. Their choice is essential to direct the desired course of the reaction. In this section, the synthesis of molecular boryl-derivatives is presented, while the formation of macromolecular compounds is described in Section 3.3.

Selective hydroboration of the separated 1,6-diynes, 1-pinacolboryl-hepta-1,6-diyne **79a** or 1-pinacolboryl-octa-1,7-diyne **79b**, occurred at the terminal $C \equiv C$ bond with the *syn*-addition of pinacolborane **25** in the presence of 5 mol% Cp₂ZrHCl **80** as a catalyst (according to Wang's procedure)¹³² or using HBBr₂–SMe₂ **81**, which was further transformed with pinacol **82** to generate stable boryl derivatives **83a–b** in 94–95% yield. The obtained diborylenynes **83a–b** were further cyclised using Cp₂ZrCl₂ **84** and *n*-BuLi **85** followed by treatment with anhydrous HCl in diethyl ether. The resulting products **87a–b** possessing boryl groups attached to C_{sp^2} and C_{sp^3} were used in Suzuki coupling reactions with iodoarenes in typical conditions, applying the commonly used Pd(PPh₃)₄ **35** as a catalyst. Here the reaction occurred exclusively on C_{sp^2} -B bond, because of its much higher reactivity in this coupling reaction (Scheme 17).¹³⁰

Wang's procedure was also used for the bishydroboration of aminodiyne **90** with pinacolborane **25**. (*E*,*E*)-Bis(vinylboronate ester) **91** was obtained in 54% yield. When the modified Srebrnik procedure was used, the same product was formed but with opposite (*Z*,*Z*)-stereoselectivity **93**.¹³³ Here reaction of diyne **90** with *n*-BuLi **85** (2 equiv.) in Et₂O at -78 °C, which was then transferred to a solution of 2 equiv. of PINBOP **92**, followed by the addition of HCl. The crude bis-alkynyl-Bpin was then added to a solution of Schwartz reagent Cp₂ZrHCl **80**,



Scheme 12 Transformation of borylsilyl-substituted enynes 41 in bromodeborylation, Suzuki–Miyaura and Hiyama coupling reactions.



Scheme 13 Monohydroboration of symmetrical and nonsymmetrical conjugated diynes with bis(pinacolato)diboron 61 and methanol as a proton source catalysed by a CuCl 55/P(p-Tol)₃/NaOt-Bu system.

and the obtained zirconocene was then hydrolysed to the bisborylated diene **93** in 49% yield (Scheme 18).¹³⁴ Wang's and Srebrnik's procedures were also applied for the formation of other (*E*,*E*)- or (*Z*,*Z*)-vinyl boronate esters with moderate yields, which were further cyclised using PdCl₂(PPh₃)₂ **94** to various cyclic polyenes with controlled (*E*,*E*), (*Z*,*Z*) or (*E*,*Z*) selectivities **95–104** (Schemes 18 and 19).¹³⁴

Non-catalytic hydroboration of diyne **105** with sterically hindered di(iso-pinocampheyl)borane (l Ipc₂BH) **106** was carried out in THF at 0 °C yielded the desired selective hydroboration of the Me-substituted alkyne. Addition of bromodienoate **107**, Pd(PPh₃)₄ **35**, and TlOEt to the product provided the targeted cross-coupling product **109** in 83% yield with excellent regioselectivity (>95:5) (Scheme 20). This method was developed and used as a part of the synthesis of the natural compound *Apoptolidin An* isolated from actinomycete identified as *Nocardiopsis sp*, which possesses cytotoxic properties. Interestingly, the application of the less hindered pinacolborane 25, cateholborane 26, or dicyclohexylborane 15, led to the mixture of isomers in the reaction.

Hydroboration of chiral binol derived diynes **110a–e** was carried out with Piers borane (HB(C_6F_5)₂) **111** in mesitylene to **112a–e**. After 5 min, tri(*tert*-butyl)phosphine **113** was added to **112** to generate a frustrated Lewis pair *in situ*. This system was used for the enantioselective reduction of enol silyl ethers **114a–u** under 40 bars of H₂. After workup with TBAF





(tetrabutylammonium fluoride) **108**, chiral secondary alcohols **115a–u**, with excellent yields and enantioselectivities (87–99% ee) were obtained. The catalytic alkenylborane activity **112a–e** (Lewis acidity) was tuned by conjugation of the system as well as the type of electron-rich or deficient substituents attached to the binaphthyl ring (Scheme 21).¹³⁵

Ruthenium catalysts have not only been used in the hydroboration of conjugated 1,3-diynes⁷² but also in the hydroboration of separated diynes **116a–c**.¹³⁶ In this case, application of ruthenium hydride pincer complex [Ru(*t*-BuPNP)(H)₂(H₂)] **117** (PNP = 1,3-bis(di-*tert*-butyl-phosphinomethyl)pyridine) permitted anti-Markownikow *trans*-hydroboration leading to (*Z*)-vinyl boranates **118** under mild reaction conditions (r.t., 24 h, toluene). Generally, the system was active in the reaction with various terminal alkynes, but very good yields and selectivities were also obtained in bishydroboration of hepta-1,6-diyne **116a**, deca-1,9-diyne **116b** and 1,4-diethynylbenzene **116c** (Scheme 22).

The complex $[Ru(PNP)(H){(\mu-H)_2Bpin}]$ **119**, which is formed in the reaction of 117 with pinacolborane 25, with the simultaneous evolution of H₂, was found to be the catalyst for this transformation which was structuraly characterised (Scheme 23). Based on stoichiometric reactions, DFT calculations, and catalytic transformation with deuterated d1-phenylacetylene, the mechanism of this transformation was determined. 119 is generated by the formation of the ruthenium hydride complex with a covalent bond Ru–B through σ -bond metathesis. 119 is subsequently further substituted with an alkyne generating complex **120.** Then dihydrogen migration led to η^1 -vinylidene complex 121. Complex 122 is then formed by the coupling between borane and vinylidene ligands. Coordination of pinacolborane 25 followed by σ -bond metathesis releases product 124 and generates complex 125. The addition of the next alkyne molecule regenerates complex 120 closing the catalytic cycle. The (Z)-stereochemistry of the product is determined in the reaction



Scheme 15 Proposed mechanism of radical *trans*-hydroboration of 1,3-diynes 60a and 65a–l with an N-heterocyclic carbene borane 66.



Scheme 16 Derivatisation of 70a in Pd-catalysed Suzuki-Miyaura coupling.

sequence from 120 to 122, presumably reflecting steric interactions in the formation of complex $121.^{136}$

Applying the low-valent Co catalyst 126, generated in situ from CoCl₂/phenanthroline, TBAF 108, and pinacolborane 25, it was possible to carry out the cyclisation/hydroboration of 1,6divnes 116a, 127a-r, and 129 yielding cyclic 1,3-dienylborons 128. No other work has been reported on this type of cyclisation of diynes, although analogous systems with envnes and dienes have been published.¹³⁷⁻¹⁴² This reaction was observed to be more effective in dilute solutions. Different ligands and activators, e.g., TMSCH₂Li, KOAc, t-BuOK could be used, but TBAF 108 and 126 were the most efficient. It was found that the system was active using various 1,6-divnes 116a and 127a-r with different substituents, e.g., ketone, amide, nitrile, or sulfone. Not only C-tethered but also N- and O-tethered 1,6diynes were reactive in this transformation furnishing heterocyclic compounds 128a-s. Interestingly the reaction with 1,7divne failed in most cases with only one example using 4,4,5,5tetraester 129 which underwent cyclisation/hydroboration to the six-membered ring product 130 (Scheme 24).

The mechanism of this transformation was proposed according to the stoichiometric reactions and experiments with DBpin **45**. The reaction is initiated by the formation of low-valent Co complex **131** in the reaction of L-CoCl₂ **126** with HBpin **25** (DBpin **45**) and TBAF **108**, which reduces the Co(π) to Co(0). In the next step coordination of diyne **116a**, or **127a-r** to **132**, occurs followed by the oxidative cyclisation to form a five-membered cobalt-containing cyclic intermediate **133**. **133** then undergos σ -bond metathesis with **25** (transition state **134**) to give intermediate **135**. Reductive elimination of the product **128a-s** from **135** regenerates the low-valent cobalt species **131** (Scheme 25).¹²³ The utility of **1**,3-dienylborones **128a-s** as building blocks was tested for **128a** in Diels–Alder, oxidation, chlorodeborylation, and Suzuki–Miyaura coupling reactions (Scheme 26).



Scheme 17 Hydroboration of 1,*n*-diynes **79a-b** to borylated enynes according to Wang's procedure followed by the cyclisation and deborylation processes.



Scheme 18 Synthesis of macrocyclic dienes 95 and 96 applying the hydroboration reaction of aminodiyne 90 under Wang's and Srebrnik's procedures.



Scheme 19 Synthesis of macrocyclic dienes 97-103 and 104 based on the hydroboration/Pd-catalysed cyclisation of 1,*n*-diynes.

Taniguchi *et al.* reported another type of hydroboration of benzo[3,4]cyclo-dec-3-ene-1,5-diynes **144a–m** which, following a borylative radical cyclisation, permitted the formation of 5-borylated 6,7,8,9-tetra-hydrobenzo[a]azulenes products **145a–m**.^{143,144} The boryl radicals were formed from N-heterocyclic carbene–boranes **146** with radical initiators, of which di*tert*-butyl hyponitrite (TBHN) **147** was the most effective. The homolytic bond dissociation energies of N-heterocyclic carbene boranes are much lower than those which possess typical boryl hydrides, and therefore these compounds might be used as precursors for rather stable boryl radicals **148**. The reaction



Scheme 20 Towards the synthesis of natural compound Apoptolidin An. Hydroboration of diyne 105 with ${}^{\rm l}{\rm Ipc_2BH}$ 106 followed by the Suzuki–Miyaura coupling with 107 to give 109.

occurred according to a radical chain mechanism by the intramolecular addition of the boryl radical followed by the cyclisation process. Under the optimised reaction conditions, 100 °C, trifluoromethylbenzene, 0.4 equiv. of TBHN 147, and 5-fold excess of NHC-borane 146 products were obtained with moderate yields, which slightly varies depending on the substituents in both reagents: borane 146 and divne 144 (Scheme 27). The obtained borylated compound 145a was subsequently transformed in a one-pot cascade reaction into deborylated products 150a-c and 151a-c in the following reactions with N-chlorosuccinimide (NCS) 78 and then a Suzuki-Miyaura coupling with aryl iodides. Depending on the reaction conditions (path A or B), a different distribution of products was observed (Scheme 28).^{143,144} The mechanism of this transformation was discussed based on the formation of borepin 156 from cyclic diyne 152 and NHC-boryl radical 148, whose formation was initiated by the homolysis of the di-tert-butyl peroxide activator. The obtained radical 148 was added to diyne 152 to form alkenyl radical 153, which was transformed to the hydroboration product by a



Scheme 21 Asymmetric hydrogenation of silyl enol ethers **114a–u** using frustrated Lewis pairs catalysts based on alkenyl boronates **112a–e** formed *in situ* in the hydroboration of diynes **110a–e** with Piers borane **111**.



Scheme 22 Hydroboration of terminal separated diynes **116a–c** with pinacolborane **25** catalysed by Ru-complex **117**.

hydrogen atom transfer from NHC-borane **148**. Alkenyl borane **154** has two remaining B–H bonds and can undergo a second hydroboration to give product **156** (Scheme 29).¹⁴⁵

3.3. Hydroboration of separated 1,*n*-diynes in the synthesis of macromolecular compounds

Non-catalytic hydroboration of internal and terminal diynes with alkyl or aryl spacers between ethynyl groups with dihydroboranes can lead to the formation of polymers, which possess a boryl group attached to C_{sp^2} determining their further reactivity. In most cases, sterically hindered thexylborane **157**, mesitylborane **158**, or tripylborane **159** have been used as hydroboration agents.

The foundation research carried out by Chujo *et al.*, used thexylborane **157** in the polyaddition process to terminal octa-1,7-diyne **160**, as well as internal 3,9-dodecadiyne **161a**, 3,8undecadiyne **161b**, 3,10-tridecadiyne **161c**, 3,9-dodecadiyne **161d**, which occurred in THF, at 0 °C. When terminal diyne **160** was used, the polymer **162**, possessing 20% of the branched structure was obtained, which was visible by the gelation of the reaction mixture. The cross-linking structure occurred because



Scheme 23 Mechanism of Z-selective hydroboration of alkynes and diynes using Ru-pincer complex 117.

of the easy access of the second borane molecule to the unshielded vinylborane bond in the subsequent hydroboration process. This branched structure caused the broadening of the molecular weight distribution of the polymer. When internal diynes **161** were used, the linear polymers **163** were mostly formed, which was determined by the observation of vinylic groups in ¹H NMR spectra (Scheme 30).¹⁴⁵

When mesitylborane **158** was used as a hydroborating agent, no gelation and crosslinking were observed due to its high steric hindrance. The linear polymers **165** were formed from terminal separated diynes **116b**, **c**, **160**, and **164a–i** with good or moderate yields (35–95%) and moderate molecular weights (Scheme 31 and Table 2). The solvent type was observed to have a big influence on the products yields (Table 2, entries 1–4). Moreover, the temperature and the diyne type are also important factors for the reaction course. The best results were obtained for polymerisation of **160** in deuterated CDCl₃, but for most of the examples, THF was used. The polymers were more stable to air-oxidation than the products obtained by hydroboration with thexylborane **157**. The application of diynes with chromophores permitted the synthesis of polymers with optoelectronic properties.¹⁴⁶ The organoboron polymer prepared from diethynylbenzene **116c** and mesitylborane **158** was subsequently subjected to reaction with iodine to form poly(phenylene–butadienylene) **166** (Scheme 32).^{146–150}

The same authors described the application of hydroboration with tripylborane **159** as a method for producing optoelectronic polymers **167a–d** using various 1,4-diethynylbenzenes as starting monomers (Scheme 33). The obtained polymers with chromophores **167a–d** emitted green or blue light, while their



photostability depended on the electron density of the substituents in the comonomers. Moreover, conjugated polymers containing boron atoms in their backbone are known to extend π -conjugation through the vacant p-orbital of the boron atom. The authors used Gaussian 03 and theoretical calculations using DFT methods at the B3LYP/6-31G(d,p)/B3LYP/6-31G(d,p) level to calculate the bandgap in the polymers. The results showed that the bandgap decreased significantly with increasing the number of repeating mers in the polymer, showing that conjugation length was extended in polymer *via* the vacant p-orbital of the boron atoms.^{147–150}

Using the same theoretical methods, it was possible to calculate the reactivity of C=C bonds depending on the R-groups attached to the aryl ring in the polymer by calculating the bond order. The bond order was found to be $OCH_3 < CH_3 < H$, while the stability was in the opposite order

 $OCH_3 > CH_3 > H.^{150}$ The polymerisation was also carried out for optically active diyne **168** with tripylborane **159**. A chiroptical activity was induced to the polymer **169** *via* the chiral side chain (Scheme 34).¹⁵¹

4. Hydroalumination of conjugated and separated diynes

The addition of an Al–H bond to the $C \equiv C$ bond of a diyne may proceed *via* mono- or bishydroalumination for the synthesis of metallated enynes or dienes, which can be further used in the chemical transformation towards the synthesis of natural products or *fine chemicals*. In comparison to boranes, there is a limited availability of organoaluminium hydrides which is



Scheme 25 Proposed mechanism for the Co-catalysed hydroboration/ cyclisation of 1,6-diynes **116a** and **127a–r**.

responsible for just a small number of papers focused on the hydroalumination of alkynes and diynes.^{152,153}

In 1977, Zewifel described the hydroalumination of 1,3divnes with lithium di(iso-butyl)methylaluminium hydride 172, which was formed in the reaction of di(iso-butyl)aluminium hydride 170 with methyllithium 171. The reaction occurred in diglyme at room temperature furnishing lithium enynylaluminate 173. The rate of hydroalumination was found to be dependent upon the solvent and, when 1,2-dimethoxyethane or THF were used, the yields were much lower. The hydroalumination of the second $C \equiv C$ bond in 13a-c, 65d was not observed even when a 50% excess of aluminum hydride 172 was used (Scheme 35).¹⁵⁴ The reaction was highly stereoselective, which was confirmed with the exclusive formation of (*E*)-enynes 174a-d after hydrolysis of obtained aluminate.¹⁵⁴ Deuterolisys of the aluminate in D₂O was used to prove the reaction regioselectivity. More than 98% deuterium was placed at the less shielded internal carbon bond 175. The products 173 were additionally transformed to envonic acids 176 in the reaction with CO2. The reaction was only selective for symmetrical diynes 13a-c, 65d (Scheme 35). In the case of 2,2-dimethyldeca-3,5-diyne 177, two regioisomers 178 and 179 were obtained with comparable yields (Scheme 36). The opposite (Z)-isomer 16 to 173 was obtained by the same authors using hydroboration reaction.¹²⁶ To apply this transformation to unsymmetrically substituted diynes, reagents with electronically different substituents attached to $C \equiv C$ bonds



Reaction conditions: (a) N-phenylmaleimide, THF, r.t., 15 h; (b) $H_2O_2/NaOH$, THF, r.t., 3 h; (c) $CuCl_2$ **143**, THF/ H_2O 70 °C, 10 h; (d) pnitroiodobenezne **76**, Pd(PPh₃)₄ **35**/K₂CO₃, toluene/ $H_2O/MeOH$, 80 °C, 12 h; (e) alkenyl bromides or iodides, Pd(PPh₃)₄ **35**/MeONa, toluene/MeOH, 80-100 °C, 15 h; (f) 2,3-dicyano-5,6-dichlorobenzoquinone, toluene, 100 °C, 3 h; (g) 1. glyoxilic, morpholine, hexafluoropropan-2-ol, r.t., 8 h; 2. TMSCH₂N₂, THF, 4 h.

Scheme 26 Application of product 128a as a building block in organic chemistry.





were used **180a–c**. The trimethylsilyl group was attached to one alkyne, while an alkyl or cycloalkyl group was included on the other alkyne. The presence of the silyl protecting group made the second triple bond more susceptible to nucleophilic attack by the aluminum hydride **181** (Scheme 37). This strong activating effect from the silyl group was proved by the reaction of equimolar amounts of two different diynes: silyl-substituted **180a** and alkyl-substituted deca-4,6-diyne **65b**. Within the process, the silyl-substituted diyne **180a** was converted to enyne **182a**, while diyne **65b** was unreactive.¹⁵² The obtained enynes

following deprotonation **182a–d** were subjected to a second hydroalumination reaction with *i*-Bu₂AlH **170**. Here, the aluminate was attached to the C₁ atom with silyl group **183a–d** and then hydrolysed to **184a–d**. This regiochemistry was analysed according to the deuterolysis reaction, indicating more than 95% of D atoms at the C₁ position (Scheme 37).

The hydroalumination and hydroboration reactions were applied in the synthesis of insect pheromone *Bombykol* **187** applying a desilylation procedure with KF × 2H₂O in DMF. The product **187** was obtained in high *trans*-selectivity in 81% yield (Scheme 38).¹⁵²

27% (2 h)

31% (4 h)

1) NCS **78** (2.0 equiv.) CH₂Cl₂, 0 °C, 15 min 2) 4-iodoanisole **58** (1.1 equiv.), PdCl₂(dppf) **149** \cdot CH₂Cl₂ (10 mol%) Na₂CO₃ (5.0 equiv.) THF, reflux, 2.5 h







44% (2 h)

42% (4 h)





PATH B: R = NO₂

Scheme 29 Proposed mechanism for the formation of borepin 156 based on radical hydroboration.



Scheme 30 Noncatalytic hydroboration of terminal octa-1,7-diyne 160 and internal separated diynes 161a-d with thexylborane 157.



Scheme 31 Non-catalytic hydroboration of terminal separated diynes (116b–c, 160, 164a–i) with mesitilborane 158. The results of the polymerisation are presented in Table 2.

Table 2Results of polymerisation of terminal separated diynes 116b-c,160, 164a-i with mesitilborane 158 based on the hydroboration reaction

Entry	Diyne	Diyne:158	Mn	$M_{ m w}$	$M_{\rm w}/M_{\rm n}$	Yield of 165 [%]
1	160	n.a. ^d	5600	9600	1.7	35
2^a	160	n.a. ^d	4200	5900	1.4	63
3^b	160	n.a. ^d	12600	28600	2.3	80
4^c	160	n.a. ^d	4700	8100	1.7	56
5	164a	n.a. ^d	7100	13200	1.9	39
6	116b	n.a. ^d	6300	12600	2.0	38
7	164b	n.a. ^d	6100	13400	2.2	48
8	164c	n.a. ^d	810	1270	1.6	n.a.
9	116c	n.a. ^d	10500	24400	2.3	47
10	116c	1.17	6500	16000	2.5	71
11	164d	1.27	3000	4800	1.6	58
12	164e	1.03	3000	5900	2.0	57
13	164f	1.21	2900	4500	1.6	36
14	164g	1.27	5100	10500	2.1	95
15	164ĥ	1.24	2800	4200	1.5	71
16	164i	1.22	1300	1700	1.5	67

Reaction conditions: THF, room temperature, isolated after precipitation in MeOH. ^{*a*} 50 °C. ^{*b*} CDCl₃. ^{*c*} CH₂Cl₂. ^{*d*} A small excess of diyne **116b–c**, **160** or **164a–i** was added to the THF solution of **158** (1.0 M).

Hydroalumination of 1,4-bis(trimethylsilyl)-1,3-butadiyne **180c** and 1,4-bis(trimethylsilylethynyl)benzene **188** with di(*tert*butyl)aluminium hydride **189** proceeded *via cis*-addition of the Al–H bond to both $C \equiv C$ bonds in the diyne. Due to the directing effect of the silyl group, both organoaluminium groups were

attached to the carbon atoms possessing the silicon atom. Within this reaction, the kinetic dienes 190 and 191 with (Z,Z)-stereoselectivity were formed. Increasing the temperature to 60 °C degrees caused the rearrangement of diene towards the thermodynamic product with (E,E)-configuration 193. The exclusive formation of this isomer occurred when 1,4-bis-(trimethylsilylethynyl)benzene 188 was used as an initial reagent. In the case of 1,4-bis(trimethylsilyl)-1,3-butadiyne 180c upon heating, a mixture of different products was obtained. The formation of both products: kinetic and thermodynamic were confirmed using NMR spectroscopy and X-ray analysis. The rearrangement of isomers from (Z,Z) 191 to (E,E) 192 took 7 days, while the total consumption of initial diynes in the first hydroalumination step was carried out for 15 or 3 hours respectively (Scheme 39). The formation of products 190, 191, 193 was confirmed by X-ray analyses (Fig. 2).155

5. Hydrosilylation

The subject focused on the hydrosilylation of diynes is the most documented of all hydroelementation processes discussed in this review. This is owing to the fact that the products are useful synthons in organic chemistry. The presence of the silyl group in the product structures, as well as other functional groups (hydrosilylation is a highly tolerant reaction), these compounds



Scheme 32 Synthesis of poly(phenylene-butadienylene) 166











Scheme 35 Synthesis of (E)-envnes 174a-d and envonic acids 176 in the hydroalumination/hydrolysis or oxidation reactions.



Scheme 36 Hydroalumination/hydrolysis of unsymmetrical 2,2-dimethyl-deca-3,5-diyne 177 with 172.

might be applied in various transformations leading to fine organic and organometallic compounds and materials.^{67,68,156–162} This broad applicability is a result of the ease of substitution of the silyl group with a broad range of functional groups, as well as the formation of different silylated products: 1,3-enynes, allenes, polymers, or cyclic compounds, depending on the type of diyne starting material (conjugated or separated with alkyl or aryl spacers). Moreover, silyl-substituted compounds are easy to handle, simple for isolation, stable in air, and active in many chemical transformations. Additionally, the hydrosilylation of diynes, when an appropriate catalyst is chosen, might be carried







Bombykol

i) Li[AlH(*i*-Bu)₂*n*-Bu] **181**/DME-hexane, 25 °C, 1 h; 3 M HCl; ii) KF x 2H₂O/DMF, 25 °C; iii) *n*-BuLi **85**/hexane-diglyme, -78 °C to 25 °C, *n*-C₃H₇Br/25 °C to 80 °C, 18 h; iv) Disiamylborane **14**/THF, 0 °C, 3 h; AcOH, 60 °C, 5 h, H₂O₂, NaOH, 30 °C to 50 °C; v) MeOH, 3 M HCl.

Scheme 38 Synthesis of insect pheromone Bombykol 187 with hydroalumination, desilylation, and hydroboration steps.



Scheme 39 Kinetic 190 and 191 and thermodynamic products 193 in the hydroalumination of diynes 180c and 188 with HAl(t-Bu)₂ 189

out in a 100% atom economic way yielding a single product. Such an approach is especially important owing to the simplification of separation steps. Therefore, considering the reaction methodology, conditions, and application of a specific type of catalyst (often tailored-made) is of prior importance, especially when such complex diyne molecules are used as reagents.^{68,73,74,156,163} The simplicity of the hydrosilylation process, its high tolerance towards various functional groups present in the reagent



Fig. 2 Molecular structures of kinetic **190**, **191** and thermodynamic **193** wich were obtained in the hydroalumination reaction of **180c** and **188** with HAI(t-Bu)₂ **189**.¹⁵⁵

structures, as well as the diversity of the selectivities, which can be tuned by the proper choice of the catalyst, has rendered this transformation the first choice for the synthesis of organosilicon compounds. The reactivity of the silyl group in coupling reactions or desilylation processes has allowed the application of the resulting compounds (*e.g.*, 1,3-enyne or 1,3-diene fragments), in the synthesis of natural or biologically active compounds.^{67,68,156–160}

To systemise the results in this section, the information is ordered according to the hydrosilylation of conjugated 1,3-diynes, 1,*n*-diynes, as well as the formation of various products, 1,3-enynes, allenes, polymers, or cyclic compounds.

5.1. Hydrosilylation of conjugated 1,3-diynes towards molecular and macromolecular unsaturated organosilicon compounds

The hydrosilylation of conjugated 1,3-diynes is a straightforward and 100% atom economic method, which occurs *via* the addition of the Si–H bond to the C \equiv C bond, but due to the presence of two such alkyne groups, the formation of a specific single product with high selectivity is a challenging task. Depending on the type of the catalyst, reagents, their concentration, ratio, and process conditions, silylated 1,3-enynes, 1,3dienes, or allenes can be formed, frequently as a complex mixture of products (up to nine different compounds can potentially be formed).^{62,73,74,156,164} Many papers describe the hydrosilylation of monoalkynes,^{14,41,48,49,51} but the addition of the Si–H bond to 1,3-diynes is much more demanding and limited only to a few papers. The hydrosilylation of 1,3-diynes occurs mainly in the presence of noble metal complexes (Rh, Pt, Pd, Ru). There are also some examples of the application of less expensive Ni or Co catalysts. However, in the majority of examples there was little discussion of the influence of the reagent structure, reaction conditions, or the nature of the catalyst on the reaction outcome. Our recent papers focused on the hydrosilylation of 1,3-diynes with silanes or silsesquioxanes in the presence of commercially available platinum complexes give the first detailed research which discusses the influence of several parameters on the hydrosilylation selectivity.^{73,74,165}

In a report by Perry et al. the synthesis of conjugated polymers from 1,3-diynes and bis(silylhydrides) were described. The products were obtained via hydrosilylation reactions in the presence of Karstedt's catalyst 194. To check whether the double Si-H addition to the 1,3-diyne had occurred, the authors carried out model reactions using monohydrosilanes 195a-f with methyl, phenyl, or trimethylsiloxy groups with 1,4-diphenylbuta-1,3-diyne 1a, dodeca-5,7-diyne 13a, and 2,2,7,7tetramethyl-octa-3,5-diyne 13c. Bissilyl adducts 197 were obtained under harsh reaction conditions (120-145 °C) in xylene. The silyl-substituted but-3-en-1-ynes 196 were obtained for less bulky silanes 195a and 195b under lower temperature (80 °C) in toluene. Not only were the steric properties of silanes important but also the 1,3-diyne used influenced the formation of monosilyl or bissilylated adducts. The functionalisation of bulky 13c gave exclusively the silvl-substituted envne 196 (75–99%) (Scheme 40). The hydrosilylation occurred according to the syn-addition with the silyl group attached to the most internal carbon atom.

Furthermore, the polymerisations were carried out with dihydrosilanes **198a-c** which furnished polymers **199** with a



Scheme 40 Hydrosilylation of symmetrical 1,3-diynes 1a, 13a, and 13c with silanes 195a-f in the presence of Karstedt's catalyst 194.



Scheme 41 Hydrosilylation of symmetrical 1,3-diynes **1a**, **13b** with dihydrosilanes **198a–c** catalysed by Karstedt's catalyst **194**.

Table 3 Parameters of polymers 199 obtained in the hydrosilylation reaction of conjugated diynes 1a and 13b with dihydrosilanes 198a-c

				199					
Entry	Diyne	198	M _n	$M_{ m w}$	$M_{\rm w}/M_{\rm n}$	State			
1	1a	198a	9540	4130	2.31	Solid			
2	1a	198b	n.a.	n.a.	n.a.	n.a.			
3	1a	198c	16190	6490	2.50	Solid			
4	13b	198a	19770	8020	2.47	Liquid			
5	13b	198c	17 610	7190	2.45	Liquid			

(Z,Z)-2,3-disubstituted-1,3-butadiene mers. The main products were linear polymers **199**, but a few percent of cyclic oligomers were formed as well. Most of the cyclic products and low molecular weight linear polymers were separated by precipitation in MeOH: acetone = 3:1 solution, which presence was confirmed by MALDI and SEC analysis. The authors reported also that the rate of polymerisation depended on the catalyst **194** concentration, but did not influence the molecular weight of the polymers and polydispersity, which varies from 2.0 to 2.61 for specific reagents (Scheme 41 and Table 3).⁶²

The hydrosilylation of poly[(dimethylsilylene, methylphenylsilylene, and diethylsilylene)but-1,3-diyne] **200a–c** with 1,4-bis-(methylphenylsilyl)benzene **201** was selectively carried out at 80 °C in the presence of 0.5 mol% of Rh₆(CO)₁₆ **202**, with a **200 : 201** ratio = 1 : 0.3–0.42 (Scheme 42). Applying the catalyst **202**, the addition to only one alkynyl group in the polymer occurred, while other Rh complexes, *e.g.*, Rh(acac)(CO)₂ **204** and RhCl(PPh₃)₃ **205** also yielded allenes. H₂PtCl₆ **206** additionally catalysed depolymerisation reaction. The polymers **203** were obtained with 49–79% yield, the $M_w = 112\ 000-424\ 000$, and $M_w/M_n = 2.1$. The higher the molecular weight of **203**, the longer reaction time was needed. The catalyst activity was checked in the model reaction of poly[(dimethyl-silylene)buta-1,3-diyne] **200a** with triethylsilane **207a**.¹⁶⁶

Escribano's group used a heterogeneous monometallic or bimetallic catalyst with active calcinated or non-calcinated platinum supported on titania in the hydrosilylation of symmetrical 1,4-diaryl **1a**, **1c**-**d**, **208a**, or 1,4-dialkyl-substituted-1,3diynes **60a**, **65f**, **208b**, and one unsymmetrical diyne **208c** with



Scheme 42 Rh-catalysed crosslinking of polymers with conjugated $C \equiv C$ bonds 200 with 1,4-bis(methylphenylsilyl)benzene 201.

silanes 195e and 207a-b. The best activity occurred using Pt/ TiO_2 catalyst 209 (Scheme 43). Under the optimised reaction conditions (0.25 mol% of Pt/TiO₂ 209, 70 °C, solvent-free conditions), depending on the reagent structures, monohydrosilylation or bishydrosilylation resulted. Electronically different diaryl-1,3-diynes underwent hydrosilylation using three silanes (Et₃SiH 207a, Ph₃SiH 195e, (MeO)₃SiH 207b) giving silylated 1,3-enynes 210a-g with high yields (85-98%). The hydrosilylation of electron-rich reagents was much faster than for electronpoor diynes and occurred with higher yields. Bishydrosilylation of divnes was possible only for the dialkyl-substituted reagents 60a, 65f, and 208b using 2.5 equiv. of silane 207a. Additionally, the hydrosilylation of unsymmetrical 5-phenylpenta-2,4-diyn-1ol 208c with Et₃SiH 207a gave silvlated 1,3-envne 210g as a product with 75% isolated yield. For the reaction of 1,4-diphenylbuta-1,3-divne 1a with triethylsilane 207a, catalyst 209 was filtered and three times recycled, giving the product 210a with 100, 70, and 15% in the following cycles. The significant decrease of the product yield was caused by the Pt leaching, which was confirmed by ICP-MS analysis (the catalyst 209 contained 22% of initial Pt loading after the third cycle). The products 210 and 211 were obtained with (E)-stereochemistry and with the silicon atom bonded to the internal carbon atom of the conjugated system.¹⁶⁷

Another heterogeneous catalyst, which was used in the hydrosilylation of alkynes and 1,3-diynes was based on Rh nanoparticles **214** synthesised by the reduction of RhCl₃ **213** with NaBH₄ **212** and their further stabilisation in a nitrogenrich poly(oxyethylate) derivative. The catalyst was used for hydrosilylation of dodeca-5,7-diyne **13a** and 2,2,7,7-tetramethylocta-3,5-diyne **13c** with triethylsilane **207a** used in a 4.0–6.0 fold excess. The more hindered diyne **13c** gave monosilylated enyne **215**, while the less shielded **13a** gave bissilylated diene **216** with excellent isolated yields (95% and 98% respectively) (Scheme 44).¹⁶⁸ Palladium nanoparticles were also tested for single alkyne examples.^{169,170}

The hydrosilylation of conjugated symmetrical 1,3-diynes 1a, 13c, and 180c with mono- or dihydrosilanes 195a, 207a, 217 was carried out in the presence of various Ni(0) 218–220 or



Scheme 43 Solvent-free mono- and bishydrosilylation of 1,3-diynes 1a, 1c-d, 60a, 65f, 208a-c with silanes 195e, 207a-b catalysed by Pt/TiO₂ 209.



poly(oxyethylenate) derivative 214

Rh(I) **221** complexes with the addition of different chiral or nonstereoselective ligands **222a–i**. In all experiments, the silane was used in a 3.5 to 4.0-fold excess with respect to the diyne. The bishydrosilylated allene **224** was obtained for 1,4bis(trimethylsilyl)buta-1,3-diyne **180c**, while for diynes substituted with *tert*-butyl **13c** and phenyl **1a** groups, a mixture of silyl-substituted enyne **223** and allene **224** was formed (Scheme 45). The synthetic procedure was quite enigmatic, and the silane was used in high excess with no equimolar reagent ratios tested (Table 4).^{164,171,172}

A recent publication from our group details the application of commercially available catalysts: $Pt_2(dvs)_3$ **194**, $Pt(PPh_3)_4$ **225**, or PtO_2 **226** in the hydrosilylation of various symmetrical 1,4disubstituted buta-1,3-diynes (**1a**, **13c**, **65a**, **227a-b**) with sterically and electronically different triethyl-**207a** and triphenylsilane **195e**. Comprehensive optimisation studies were carried out to find the most suitable conditions that permitted obtaining either the monosilylated enynes or bissilyated dienes with high stereo- and regioselectivity. The application of a Pt catalyst led to the syn-addition of silane to the $C \equiv C$ bond and the formation of the alkenyl silane with the silvl group attached to the internal carbon atom. This was confirmed by the crystal structures of the products 228h and 228i (Fig. 3), as well as with ¹H-¹³C HSQC and NOESY 2D NMR. Within the study, an equimolar ratio of reagents was reported for the first time, which is in agreement with the atom economy policy and simplifies the separation procedure, additionally reducing the process costs. Pt Karstedt's catalyst 194 was used for the synthesis of bisadducts 229, whereas the less active PtO₂ 226 and $Pt(PPh_3)_4$ 225 were capable of the synthesis of monosilylated enynes 228 (Scheme 46). Moreover, the influence of the reaction temperature on reaction selectivity was noticeable. For monohydrosilylation, 40 °C or lower temperature gave better selectivity. The structure of the reagents has also an important role in reaction selectivity. For sterically hindered diynes 13c and 227b and triphenylsilane 195e, only monoadducts 228h and 228e were obtained. The products 228a–j and 229a–d were isolated with 82-98% yield and were fully characterised.⁷³



Scheme 45 Catalytic hydrosilylation of symmetrical conjugated diynes 1a, 13c, 180c with silanes 195a, 207a, and 217 in the presence of L₂Ni(0)-butadiyne 218-220 and [Rh(cod)Cl]₂ 221 + L 222 complexes.

Table 4 Hydrosilylation of 1,3-diynes with L₂Ni(0)-butadiyne 218-220 and [Rh(cod)Cl]₂ 221 + L 222a-i complexes

Entry	Cat ^a	Diyne	Silane	[Diyne]:[silane]	T[°C]	<i>t</i> [h]	Yield [%]	Selectivity of (223/224) [%]	Isolation method	Isolated yield (223/224) [%]
1	218a	180c	217	1.0/3.5	80	2	100	2/91	Distillation	—/49
2	218a	180c	217	1.0/2.5	80	2	92	20/61	Distillation	
3	218a	13c	217	1.0/3.5	80	1	100	3/78	Chromatography	
4	218a	13c	217	1.0/3.5	80	6	100	1/80	Chromatography	—/78
5	218a	13c	195a	1.0/3.5	80	2	100	51/31	Solvent evaporation	
6	218a	13c	195a	1.0/3.5	80	6	100	53/33	Chromatography	
7	218a	13c	207a	1.0/3.5	80	6	66	59/—	Chromatography	50/—
8	218a	13c	207a	1.0/3.5	80	30	100	89/—	Precipitation/crystallization	74/—
9	218b	180c	217	1.0/3.5	80	6	100	—/92	Chromatography	78(0/78)
10	218c	180c	217	1.0/3.5	80	6	100	—/92	Chromatography	93(6/93)
11	219	180c	217	1.0/3.5	80	6	100	—/90	Chromatography	77(0/77)
12	220	180c	217	1.0/3.5	80	12	87	20/59	Chromatography	n.a.
13	222a	13c	195a	1.0/4.0	70	24	91	90/1	Chromatography	75(75/0)
14	222b	13c	195a	1.0/4.0	70	24	90	90/1	Column chromatography	n.a.
15	222c	13c	195a	1.0/4.0	70	24	99	92/7	Column chromatography	n.a.
16	222d	13c	195a	1.0/4.0	70	24	71	67/4	Column chromatography	n.a.
17	222e	13c	195a	1.0/4.0	70	24	78	53/25	Column chromatography	n.a.
18	222f	13c	195a	1.0/4.0	70	24	71	41/30	Column chromatography	n.a.
19	222g	13c	195a	1.0/4.0	70	24	77	56/21	Column chromatography	n.a.
20	222ĥ	13c	195a	1.0/4.0	70	24	91	56/35	Column chromatography	n.a.
21	222i	13c	195a	1.0/4.0	70	24	93	66/27	Column chromatography	n.a.

^a For 222a-i the catalyst [Rh(cod)Cl]₂ 221 was used. Only ligand is placed in the table.





The same catalytic systems **194**, **225**, **226**, and heterogeneous Pt/SDB **232**, and the equimolar ratio of reagents were used in the hydrosilylation of 1,4-symmetrically substituted 1,3-diynes

1a, **13c**, **65a**, **227a-b**, **230a-d** with 1-dimethylsiloxy-3,5,7,9, 11,13,15-hepta-*iso*-butylpentacyclo-[9.5.1.1.^{3,9}1.^{5,15}1^{7,13}] octasiloxane ((HSiMe₂O)(*i*-Bu)₇Si₈O₁₂) **231** yielding silsesquioxane



Scheme 46 Hydrosilylation of conjugated 1,3-diynes 1a, 13c, 65a, 227a-b with triphenylsilane 195e and triethylsilane 207a in the presence of commercially available platinum catalysts 194, 225, and 226.



Scheme 47 Hydrosilylation of 1,3-diynes 1a, 13c, 65a, 227a-b, and 230a-d with silsesquioxane 231 using equimolar amounts of reagents and commercially available Pt-complexes.

products with several functionalities attached to the enyne **233a–m**, **234a–m**, or diene moieties **235a–m**, **236a–m**, *e.g.*, 4-boronic acid pinacol ester, 4-bromophenyl, hydroxyl groups, making them potentially useful nanobuilding blocks in polymerisation or Suzuki–Miyaura, Sonogashira, Heck, and Hiyama coupling reactions.⁷⁴ The process selectivity depended on the catalyst type and concentration, as well as the structure of the reagent. For hindered 1,3-diynes as **13c** or **227b**, only silsesquioxyl-substituted enynes **235** were formed (Scheme 47

and Table 5). Alkenylsilsesquioxanes have already been used in materials chemistry, in the synthesis of OLEDs, liquid crystals, or porous biocompatible materials. The attachment of silsesquioxanes as pendant groups to the conjugated molecular or macromolecular compounds is known to increase material brightness, color stability, and their solubility in organic solvents, or to improve the mechanical or thermal properties of the final products.^{173–176} Similar systems were obtained by the use of incompletely condensed silsesquioxanes **238a–b**. In these cases,

Table 5The optimised reaction conditions for the hydrosilylation of1,3-diynes 1a, 13c, 65a, 227a-b, and 230a-d with silsesquioxane 231

Entry	Diyne	[Pt]	[231]:[diyne]:[Pt]	$T [^{\circ}C]$	<i>t</i> [h]	Selectivity of 233/234/235/236/237 ^a
1	1a	194	$1\!:\!1\!:\!4 imes 10^{-4}$	100	24	a, 85/0/5/2/8
2	1a	225	$1\!:\!1\!:\!4 imes 10^{-2}$	40	24	a, 85/0/2/2/11
3	1a	232	$1\!:\!1\!:\!4 imes 10^{-2}$	100	24	a, 83/0/9/0/8
4	13c	194	$1\!:\!1\!:\!2 imes 10^{-4}$	100	$24\ 2$	b , 93/7/0/0/0
			$1\!:\!1\!:\!2 imes 10^{-3}$			b , 93/7/0/0/0
5	65a	194	$1\!:\!1\!:\!4 imes 10^{-2}$	100	0.5	c, 76/24/0/0/0
6	65a	194	$1\!:\!1\!:\!4\times10^{-2}$	100	2	c, 86/5/9/0/0
7	65a	194	$1\!:\!1\!:\!4\times10^{-2}$	40	24	c, 83/17/0/0/0
8	65a	194	$2.3\!:\!1\!:\!2 imes 10^{-4}$	100	24	d , 0/0/91/9/0
9	227a	194	$2\!:\!1\!:\!2 imes 10^{-4}$	100	6	e, 17/0/83/0/0
10	227b	194	$1\!:\!1\!:\!4\times10^{-4}$	100	48	f, 100/0/0/0/0
11	230a	194	$1\!:\!1\!:\!4\times10^{-4}$	100	48	g, 95/0/5/0/0
12	230a	194	$2\!:\!1\!:\!4 imes 10^{-3}$	100	48	h , 0/0/69/0/31
13	230b	194	$1\!:\!1\!:\!4 imes 10^{-4}$	100	24	i, 91/9/0/0/0
14	230b	194	$2\!:\!1\!:\!2 imes 10^{-4}$	100	24	j, 0/12/88/0/0
15	230c	194	$2\!:\!1\!:\!2 imes 10^{-3}$	100	72	k, 0/0/87/13/0
16	230d	194	$1\!:\!1\!:\!4 imes 10^{-2}$	40	48	l, 73/0/5/3/19
17	230d	194	$2\!:\!1\!:\!2 imes10^{-4}$	100	96	m , 6/0/80/0/14

^{*a*} Conversion of diynes in all experiments was complete. Toluene was used as a solvent: $m_{S1}/V_{tol.} = 50 \text{ mg mL}^{-1}$.

a stoichiometric amount of diyne was used for unsymmetrical diynes with Si(i-Pr)₃ groups **239a–c**. An excess of diyne (6–12 mol) compared to silsesquioxanes was necessary when symmetrically substituted diynes **1a–b**, **13c**, **27c**, and **227b** were tested. All the target products **240a–i** were formed with a very high selectivity of 99% (Scheme 48).¹⁶⁵

The hydrosilylation of 1,4-bis(trimethylsilyl)buta-1,3-diyne **180c** with triethylsilane **207a** using RhCl(PPh₃)₃ **205**, H₂PtCl₆ **206**, Pt(PPh₃)₄ **225**, and Pd(PPh₃)₄ **35** was also described in 1984 by Hiyama *et al.*, but complex mixtures of bissilylated allenes and monosilyl-substituted enynes were obtained, regardless of the catalyst used.¹⁷⁷ Better selectivity was observed when silyl-substituted butenynes **241** in analogous reactions were used.¹⁷⁸ Chem Soc Rev

Recent work published by Ge et al. focused on the hydrosilvlation of symmetrically or unsymmetrically substituted 1,3-diynes 1a-b, 1d-e, 13a, 27c, 37a, 37c-n, 37s, and 242a-c catalysed by inexpensive $Co(acac)_2$ complex 38 with xantphos 39a, dppf 39b, or dppp 243 ligands with dihydrosilanes (Scheme 49). The authors previously reported the effectiveness of this system in the hydrosilylation of alkynes.^{179,180} Moreover, other Co-catalysed systems for terminal and internal alkynes hydrosilylation have been recently published.¹⁸¹⁻¹⁸⁵ Under the optimised conditions of 2 mol% of Co(acac)₂ 38, 2 mol% of dppp 243, 50 °C, toluene, after 24 h, several silvlated envnes 246a-z were obtained with high yields and selectivity as confirmed by GC-MS and NMR analyses. The electronic effects of substituents attached to the aryl ring were not noticeable, and the catalyst was tolerant towards many functional groups. Mainly diphenylsilane 217, but also diethylsilane 244 and methylphenylsilane 245 were used as silvlating agents.¹⁵⁶

The authors proposed the mechanism of this transformation, which started from the generation of Co-hydride complex 247 in the reaction with H₂SiPh₂ 217, in the presence of dppp 243. The insertion of 1,3-diyne 37 or 242 into the Co–H bond generated the vinylcobalt intermediate 248, which directly reacted with dihydrosilane 217 with the elimination of the desired product – (*E*)-1-en-3-yn-2-ylsilane 246 and regeneration of initial catalyst 247 (Scheme 50).¹⁵⁶ The utility of silylsubstituted 1,3-enynes 246e and 246p as building blocks in organic synthesis was presented in the desilylation reaction by oxidation to ketone 250 and silanols (251, 253), protodesilylation to enynes 249, and Hiyama and Sonogashira coupling reactions (Scheme 51) furnishing products 252 and 255 respectively.¹⁵⁶

Another example of Co-catalysed hydrosilylation of 1,3diynes (symmetrical and one nonsymmetrical reagent) was recently reported by Chen *et al.*¹⁶³ Cobalt tridentate complexes $N^{C}NN$ -CoX₂ were previously reported as effective systems for hydrosilylation of alkynes.¹⁸⁶ The catalyst obtained from CoBr₂



Scheme 48 Hydrosilylation of 1,3-diynes 1a-b, 13c, 27c, 227b, and 239a-c with silsesquioxanes 238a-b catalysed by Karstedt's catalyst 194.



Scheme 49 Co-Catalysed selective hydrosilylation of conjugated symmetrical and nonsymmetrical diynes **1a–b**, **1d–e**, **13a**, **27c**, **37a**, **37c–n**, **37s**, and **242a–c** with dihydrosilanes **217**, **244**, and **245**.

256 and tridentate ligand 257 transpired to be highly active in the hydrosilylation of various 1,3-diynes 1a–e, 27b–c, 37t, 208a, 208c, 230d, 242b–c, and 258a–q with electron-donating and electron-withdrawing groups (Scheme 52). Several ligands were tested, but the best results were obtained when 257 was used. The high conversion of diynes was obtained within 5 minutes at room temperature. A longer reaction time was required for

fluoro, chloro, bromo, trifluoromethyl, and cyano electronwithdrawing groups to obtain satisfying yields of **260a-ad**. The alkenyl-substituted diyne reacted in the hydrosilylation process under the applied conditions without addition to the C==C bond. Excellent regioselectivity was observed in the hydrosilylation of an unsymmetrical diyne. The mechanism of this transformation started from the formation of active



Scheme 50 Proposed catalytic cycle for the hydrosilylation of 1,3-diynes with dihydrosilanes in the presence of Co-catalyst formed *in situ* from Co(acac)₂ **38** and dppp **243**.

complex 261 in the reaction with NaHBEt₃ 259 and Ph₂SiH₂ 217 followed by the coordination of 1,3-diyne 1a–e, 27b–c, 37t, 208a, 208c, 230d, 242b–c, 258a–q to form intermediate 262. The 1,3-diyne inserts to the Co–Si bond yielding the vinylcobalt species 263. The reaction with the second molecule of Ph₂SiH₂ 217 causes the catalyst 261 regeneration and evolution of the silylated enyne product 260a–ad (Scheme 53).

The enyne **260a** was desilylated according to the procedure described by Ge,¹⁵⁶ and then hydrosilylated again with diphenylsilane **217** to silylated 1,3-diene **264**. Two regioisomers with silyl groups attached to the internal and external C bond were formed in the ratio 15:85 with a high yield of 95%. The double bond in enyne was unreactive under the applied reaction conditions.¹⁶³

Chen et al. tested several cobalt complexes in hydrosilvlation of 1,3-diynes 1a-e, 27b-c, 37t, 208a, 208c, 230d, 242b-c, 258a-i, 258k-q, 265a-c (Scheme 54). Among tested catalytic systems synthesised from the commercially available materials, CoCl₂dppp (1 mol%, 266) exhibited the best regio- and stereoselectivity (in the presence of 3 mol% NaHBEt₃ 259). A variety of (E)-2-silyl-1,3-enynes 268a-ah were obtained in high yields through monohydrosilvlation at the internal carbon of the 1,3diyne unit via syn-addition. Good functional (alkoxy, amine, halides, esters, heterocyclics) tolerance was achieved by testing more than thirty substrates. A mechanism of 1,3-divne hydrosilvlation was proposed (Scheme 55) in which CoCl₂-dppp 266 initially reacts with NaHBEt₃ 259 to afford the low-valent cobalt(1) hydride intermediate 269. Subsequently, the coordination of 1,3-diyne with 269 is followed by the migratory insertion of one of the alkynyl groups into the Co-H bond and forms the intermediate 270. In the end, Ph2SiH2 217 reacts with 270 and as a result, the alkenylsilanes (268a-ah) are obtained, accompanied by the regeneration of 269.187

Zhan *et al.* studied the hydrosilylation of 1,3-diynes catalysed by $Ni(acac)_2$ 273 with a series of organophosphine ligands screened in THF at room temperature. First, the use of xantphos 39a as the ligand showed moderate regioselectivity and



Scheme 51 Applications of silyl-functionalised enynes 246e and 246p as building blocks.



Reaction time: ^a 15 min; ^b 1 h; ^c 3 h.

Scheme 52 Monohydrosilylation of conjugated 1,3-diynes **1a–e**, **27b–c**, **37t**, **208a**, **208c**, **230d**, **242b–c**, **258a–q** with diphenylsilane **217** catalysed by a CoBr₂ **256**/N^cNN-tridentate ligand **257** system.



Scheme 53 Proposed mechanism of the hydrosilylation of 1,3-diynes with Co-catalyst with tridentate N^CNN-ligand **256–257**.

yield. Several other commercially available phosphorus ligands were examined, however, the results were less than satisfactory. Therefore, from vinyl-functionalised xantphos monomer, through solvothermal polymerisation, POL-xantphos 274 was obtained and employed as a heterogeneous ligand for nickel catalysed 1,3-diyne hydrosilylation. Unsymmetrical and symmetrical 1,3-diynes 1a, 1e, 37f, 37h, 180a, 239a, and 271a-b were reacted with silanes 217, 245, 272 yielding the corresponding silyl-functionalised 1,3-enynes 275a-x (Scheme 56). The authors claimed that, due to the microporous structure of immobilised system Ni(acac)₂/POL-xantphos 273/274, the selectivity of the process increased compared to the system based on the monomeric xantphos ligand. Based on the experimental results a hydrometalation pathway with a Ni(0) intermediate for this Ni-catalysed hydrosilylation of 1,3-divnes was proposed (Scheme 57). In the mechanism, the nickel precursor is reduced in situ by phenylsilane 272 to form Ni(0) 276, and then oxidative addition of the silane generates 277. Reaction with the 1,3diyne generates 278 which then leads to the alkenyl nickel intermediate 279 after the insertion of the alkyne into the Ni-Si bond. The final product 275a-x is obtained by C-H reductive elimination with the return of the Ni(0) active species 276 into the catalytic cycle. The recyclability of the catalytic system based on Ni(acac)₂/POL-xantphos 273/274 was examined for the hydrosilylation of 1,4-diphenylbuta-1,3-diyne 1a and PhSiH₃ 272. After five runs, the Ni/POL-xantphos 273/274 reacted with nearly no loss of activity and selectivity demonstrating the good reusability of this catalytic system.188

When the hydrosilylation occurs as a *trans*-addition, cyclic siloles **283** and **284** are formed. The reaction proceeded in the

presence of 20 mol% of $[Cp*Ru(MeCN)_3]PF_6$ **281**, which was described as an effective *trans*-hydrosilylation catalyst of alkynes.¹⁸⁹ The symmetrical and nonsymmetrical 1,4-disubs-tituted-buta-1,3-diynes **1b–d**, **27c**, **60e**, **258i**, **258o**, and **280a–f** reacted with 9-silafluorene **282** or diphenylsilane **217** to the corresponding 2,5-diarylsiloles **283** and **284** with moderate to good yields. It was found that electron-donating groups attached to the aryl ring facilitated the process, while electron-withdrawing functions, *e.g.*, acetyl group **280b**, rendered the process more sluggish (Scheme 58).

The process occurred stepwise. After the first *trans*-hydrosilylation, the intramolecular second *trans*-addition proceeds. The obtained siloles were characterised by high fluorescence maxima.¹⁹⁰ The same catalytic system ($[Cp*Ru(MeCN)_3]PF_6$ **281**) was used by Trost *et al.* in the hydrosilylation of diynols which was one of the stages in the total synthesis of biologically important natural products.¹⁹¹ In this study, diynols **285** and **289** were reacted with dimethylethoxysilane **286** and benzyldimethylsilane **290**, respectively. It was observed that the partial reduction of C \equiv C to *trans* C \equiv C led to the enynols **287** and **291**, and is directed by the propargyl alcohol fragments of the diynols (Scheme 59).¹⁹¹

5.2. Hydrosilylation of separated diynes – synthesis of molecular unsaturated linear compounds

Selective hydrosilylation of α, ω -divides with CH₂OCH₂ 127p, C₄H₈ 160, and CH₂NHCH₂ 293 spacers were successfully carried out using the [Pt(IPr*OMe)(dvs)] 294 catalyst with bulky NHC ligand (where IPr*OMe = 1,3-bisimidazol-2-ylidene) and dimethylphenylsilane 195a under the typical anti-Markownikow manner. The hydrosilylation of the $C \equiv C$ bonds leads to (E)-products, while the formation of monosilylated enyne 295 or bissilylated diene 296 can be distinguished with different reagents stoichiometry: diyne : silane = 1:1 or 1:2. The exclusive formation of silylated enyne was furnished for diyne 160, while in the case of diynes with heteroatoms 127p and 293 a small amount (up to 4%) of bissilylated diene 296 was observed when equimolar reagents ratio were used. Bishydrosilylation was carried out quantitatively for 127p and 293, while for octa-1,7-divne 160 a complex reaction mixture consisting of $(\beta - E)/(\beta - Z)/\alpha$ isomers in 77:20:3 ratio were formed (Scheme 60).¹⁹² The single example of hydrosilylation of deca-1,9-diyne 116b with diphenylsilane 217 was also carried out by Leitner et al., who used ruthenium pincer complex $[Ru(t-BuPNP)(H_2)(H)_2]$ 117 [t-BuPNP = 2,6-bis(di(tertbutyl)phosphinomethyl)pyridine], which was also active in the addition of B-H bonds to alkynes and diynes.¹³⁶ The bissilylate diene 297 (E): (Z) = 91:9 was obtained using an equimolar ratio of the neat reagents and 0.2 mol% of Ru 117 within 16 h at 50 °C. No dehydrogenative coupling reaction occurred, which was visible in the case of sterically hindered terminal alkynes (Scheme 61).¹⁹³

The thermal hydrosilylation of α,ω -diynes was also used for the modification of silica surface Si(100) **298** used in the preparation of monolayers for electrodes in the reaction of diynes (*e.g.*, nona-1,8-diyne **164a**) with silica enriched with the Si-H bonds **299**. The products **300** were next modified in



Scheme 54 Monohydrosilylation of conjugated 1,3-diynes 1a-e, 27b-c, 37t, 208a, 208c, 230d, 242b-c, 258a-i, 258k-q, and 265a-c with diphenylsilane 217 catalysed by CoCl₂-dppp 266 or CoCl₂-dppp 267.



Scheme 55 Proposed mechanism of the hydrosilylation of 1,3-diynes with Co-dppp 266.

"click" chemistry with azides **301a–d** by Husigen-type cycloaddition (Scheme 62).^{194–197}

A broad range of diyne and triyne π -electron bridges arenes **116c**, **304a–f** (aryl cores = phenyl, azulene, fluorene, carbazole, 9,9'-spirobi[fluorene], 1,1'-binaphtalene) were hydrosilylated with chlorodimethylsilane **305** in the presence of Karstedt's catalyst **194** in a *syn*-addition manner towards (*E*)-silylated products **306** with high yields and selectivities. The reactions occurred with the best yield in Et₂O, at room temperature for 6–12 h with 3 mol% of a Pt-catalyst **194** (Scheme 63). The influence of the solvent on reagents conversion was visible and when THF or toluene was used a much lower conversion was observed. Good results were also obtained for the hydrosilylation of 1,3,5-triethynyllbenezene **304a** with H₂PtCl₆ **206** and Pt/ C **307** in toluene. The rhodium catalysts on the other hand were less active.

The product **306a** possessing halogen attached to silicon atom was further functionalised by a substitution reaction with lithiated chromophores **312**, **314**, or LiAlH₄ **308** in stoichiometric



Reaction conditions: $R^3R^4SiH_2$ (0.3 mmol), 1,3-diyne (0.25 mmol), Ni(acac)₂ (2 mol%), POL-xantphos (10 mg), THF (2 mL), r.t., N₂ atmosphere, 3 h. ^a Et₂O instead of THF. ^b At 70 °C. ^c r.r = 6:1

Scheme 56 Monohydrosilylation of conjugated 1,3-diynes 1a, 1e, 37f, 37h, 180a, 239a and 271a-b with Ni(acac)₂ 273 immobilised on POL-xantphos 274.



Scheme 57 Proposed mechanism of the hydrosilylation of 1,3-diynes with Ni $(acac)_2$ /POL-xantphos 273/274.

reactions, followed by the subsequent hydrosilylation reaction with 4-ethynylbenzonitrile **310**.¹⁹⁸ The presence of the SiMe₂ bridge between chromophores facilitates intramolecular photoinduced charge transfer process and interrupts the π -conjugated chains (Scheme 64).^{199–201}

5.3. Hydrosilylation of separated diynes – synthesis of conjugated polymers

In 2008, Trogler and Sanchez published a review regarding the synthesis and application of functionalised polymers delocalised through silicon, which can be obtained by a hydrosilylation reactions of diynes. The authors discussed the properties and applications of such polymers in detail, therefore within this review we focus only on the synthetic aspects in the formation of such macromolecular compounds.⁶⁸ Luh *et al.* reported the RhCl(PPh₃)₃ **205** catalysed hydrosilylation of 1,4-diethynylarenes **116c**, **164k**, and **316a–d** with disilanes **317a–b** obtained in the NiCl₂(PPh₃)₂ **319** catalysed reaction of dithiolano-substituted arenes **320** with Me₂(*i*-PrO)SiCH₂MgCl **321**, followed by the reduction of alkoxy group with LiAlH₄ **308**. The process was carried out with 0.5 mol% RhCl(PPh₃)₃ **205** with an equimolar ratio of reagents. The molecular weight of the obtained polymers was a function of reagents concentration and reaction time. Increasing both parameters led to a higher molecular weight of **318** being obtained (Scheme 65).^{202,203}

The synthesis of poly(silyl-vinylenes) was also carried out in the presence of Pt-catalysts, in particular Speier's catalyst H_2PtCl_6 **206**, with the predominant formation of polymer **322** with (*E*)- β -regioselectivity, sometimes in addition to α -silylated mers in residual amounts. The formation of both isomers was due to the steric freedom of the monomers, which did not possess any bulky substituents responsible for preventing α -silylation. The products were obtained with low M_w (Schemes 66, 67 and Table 6).^{204–206}

Platinum-based Karstedt's catalyst 194 was used in the polymerisation of aromatic diynes 327a-d with two different types of silsesquioxanes: octakis(hydrodosilsesqioxane) T^H₈ 328²⁰⁷ and double-decker-shaped silsesquioxane (DDSQ) with two hydrido functions **329**.²⁰⁸ Hydrosilylation with T^H₈ **328** was carried out with the use of 2.1-2.4 mol% of 194. The total consumption of reagents was observed in 2 h, but the reaction mixture was homogeneous even after 24 h, with the M_w ranging from 10 000 to 34 000, depending on the divne 327a-d used. The polymerisation was carried out in an equimolar ratio of the reagents, at room temperature. When the divne $327:T_8^{H_8}$ 328 ratio was increased to 1:1.55, the $M_{\rm w}$ of polymer 330a-e increased to 87 000, but crosslinking with TH8 328 was observed. ¹H NMR spectra did not detect CH-sp³ bonds in the post-reaction mixture, which indicated that the $C \equiv C$ bonds are much more reactive than the C=C bonds (Scheme 68 and Table 7).²⁰⁷

The same conclusions were obtained for the polymerisation with DDSQ **329**. The linear polymers **331a–d** were obtained for



Scheme 58 Double trans-hydrosilylation of 1,4-diarylbuta-1,3-diynes leading to 2,5-diarylsiloles 283-284 catalysed by Ru complex 281.
Review Article



internal diynes 327 (yield 90–97%, $M_n = 11\,900-29\,100$, $M_w/M_n = 2.9-4.9$) after 24 h at 100 °C with 0.2 mol% of Karstedt's catalyst **194**. When 1,4-diethynylbenzene **116c** was used as a monomer, the insoluble polymer was achieved within 30 minutes, due to the subsequent crosslinking reaction of the less shielded vinyl bonds (Scheme 69 and Table 8).²⁰⁸

The same group synthesised polymer **333** by the hydrosilylation of (1,4-bis(4-(tetrahydroxypyranyloxy)phenyl)ethenyl)benzene**332**with dihydrido-DDSQ**329**used in equal amountswith Pt₂(dvs)₃**194**as the catalyst. The reaction was carried out in toluene, at 100 °C for 7 h. The THP groups in **333** were further hydrolysed to give **334**, and the polymer with 10 wt% MBHP **335** and **1.5** wt% PTMA **336** was applied as a chemically amplified negative-working photoresist system **337** (Scheme 70).²⁰⁹ Incorporation of silsesquioxane into the polymer chain or as a pendant group enhances many properties of the final materials: mechanical, thermal, fire or oxygen resistance.^{173,176}

 $Pd_2(dba)_3$ **338**/PCy₃ was also used as an effective catalyst in the polymerisation of diethynylarenes with dihydrosilanes.^{210–212} Yamashita *et al.* used 0.1 mol% $Pd_2(dba)_3$ **338**/0.2 mol% PCy₃ and







Scheme 63 Hydrosilylation of 1,*n*-diynes **116c**, **304b**–**f** and 1,3,5-triethynylbenezene **304a**, with chlorodimethylsilane **305** in the presence of Karstedt's catalyst **194**.

an equimolar ratio of silane (Ph₂SiH₂ **217**, MePhSiH₂ **245** and Ph(H₂C=CH)SiH₂ **339**) in the hydrosilylation of *p*-diethylnybenzene **116c** and *m*-diethynylbenzene **324** at 70–110 $^{\circ}$ C for 0.5–9 h yielding polymers **340a–f** with 84–91% yield and $M_{\rm w} = 12\,000$ to 49000. The reaction occurred mostly through the *syn*-addition with the silyl group attached to β -carbon atom, but some other possible isomers (β , α) and (α , α) were also presented in the reaction mixture what was distinguished



Scheme 65 Polyaddition of disilanes 317a-b to diynes 116c, 164k, and 316a-d towards the synthesis of conjugated polymers 318.



Scheme 66 Hydrosilylation of dignes 116c, 164g with 217 and 245 catalysed by H₂PtCl₆ 206.



Scheme 67 Hydrosilylation of 1,4-diethynylbenzene 116c and 1,3-diethynylbenzene 324 with 1,4-dimethylsilylbuta-1,3-diyne 323

Table 6 Hydrosilylation of diynes $116c,\ 164g$ with 217 and 245 (Scheme 66)

Entry	Silane	Diyne	Polymer	Yield [%]	M _n	$M_{\rm w}$	$M_{\rm w}/M_{\rm n}^{\ a}$	n ^{bc}	x^d
1	217	116c	322a	92	2800	7200	1.7	$9^{b}(19)^{c}$	0.92
2	217	164g	322b	77	1550	3000	3.1	4(4)	0.96
3	245	116c	322c	77	3000	5200	2.6	12(11)	0.82
4	245	164g	322d	79	3000	9200	1.9	9(5)	0.85

 a Poly(styrene) standard. b Number of repeat units calculated from GPC data. c Number of repeat units calculated from $^1{\rm H}$ NMR data. d x-value was calculated from the relative intensity of vinyl peak to vinylidene peak in $^1{\rm H}$ NMR spectrum.

with ¹H and ²⁹Si NMR. The signals at $\delta = -15.7$ and -15.6 ppm for **340** seemed to arise from the -CH=CH-Si-CH=CH-(β , β) linkages, while the signal at $\delta = -13.4$ ppm in the ²⁹Si NMR spectrum could be signed to the -CH=CH-Si-C(=CH₂)-(β , α) linkages. The coupling constants J_{H-H} in the ¹H NMR spectrum also confirmed the predominant formation of the (*E*)product. Since the Pd-complex **338** only catalysed the addition of the Si-H bond to the C=C bond it was possible to prepare polymers with vinyl function attached to Si-atom **340e-f**, which can be then transformed in further reactions (Scheme 71 and Table 9). 210

Rao *et al.* used the same strategy for the formation of macromolecular compounds, but to build a crosslinked matrix. They used 1,3,5-triethynylbenzene **304a** or 1,3,5-triethynyl-2,4,6-trimethylborazine **342** in different ratios **245**:(**116c** or **324**):(**304a** or **342**) = 100:95:5, 100:90:10, and 100:80:20. The reaction time was 4.5 h and the polymers **343a-i** were precipitated in propan-1-ol before gelation. Depending on the ratio of momers and crosslinking agents, different molecular weights of **343a-i** were obtained, with the highest $M_w = 110\ 000-130\ 000$ for a 100:80:20 ratio. The degree of crosslinking influenced the thermal stability of polymers (Scheme 72 and Table 10).²¹¹

The same group also used phenylsilane **272** as a monomer possessing three Si–H bonds in an equimolar hydrosilylation reaction with 1,4-diethynyl-**116c** or 1,3-diethynylbenzene **324** with the Pd₂(dba)₃ **338**/PCy₃ catalytic system to furnish polymers **344** with different ratios of regioisomers $(\beta,\beta):(\beta,\alpha):(\alpha,\alpha) = 60:35:<5$. The presence of a free Si–H bond in the polymer structure **344** allowed its further modification with different



Scheme 68 Hydrosilylation of diynes 327a-d with silsesquioxane 328 in the presence of Karstedt's catalyst 194.

Table 7 Hydrosilylation of diynes 327a-d with silsesquioxane 328 (Scheme 68)

Entry	Diyne	Polymer	Diyne/ 330 in polymer ^b	$M_{ m w}^{\ \ c}$	$M_{\rm w}/M_{\rm n}^{\ c}$	$T_{d1}^{d} [^{\circ}C]$	$T_{d5}^{\ \ d} \left[{}^{\circ} \mathbf{C} \right]$	Residue at 984 $^{\circ}C^{d}$ [%]
1	327a	330a	1.23	21 000	2.03	534	1000	95.4
2^a	327a	330b	1.55	87 000	2.29	501	748	94.1
3	327b	330c	1.22	34 000	1.79	477	788	93.8
4	327c	330d	1.17	26 000	1.65	454	649	93.4
5	327d	330e	1.59	10000	1.79	486	841	92.9

Reaction conditions: 327:328 = 1:1, Karstedt's catalyst Pt₂(dvs)₃ **194** 5 µL, toluene, r.t., 2 h. ^{*a*} **327a:328** = 1.5:1. ^{*b*} Based on elemental analysis. ^{*c*} Determined by GPC with poly(styrene) standard. ^{*d*} Based on TGA in N₂.



Table 8 Hvdro	silvlation of di	vnes 116c .	327b and 327	e with silsesc	uioxane [DDSQ 329	(Scheme 6	69)
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489	156
518	153
301	_
	518 301

ethynylarenes **345a–b**, **346** in the next step towards functional polymers **347a–d**. Therefore, some additional functional groups or chromophores were included in the polymer **347a–d** as pendant groups (Scheme 73 and Table 11).²¹²

RhI(PPh₃)₃ **351** was reported as the selective catalyst for hydrosilylation of alkynes, in which the stereoselectivity can be tuned by altering the reaction temperature. When the reaction was carried out at 80 °C, the formation of (*E*)-alkenylsilane occurred, while at 0 °C the (*Z*)-isomer was predominantly formed.²¹³ The same observations were visible for hydrosilylative polyaddition of dihydrosilanes with diethynylarenes. The products were obtained with high yields and stereoselectivity depended on the reaction conditions. The coupling constants for ethenyl hydrogens were typical for (*Z*)- or (*E*)-isomers. At elevated temperatures (80 °C), polymers **353** with (*E*) regioselectivity were mainly formed ((*E*) > 93%), while at 0 °C (*Z*)-isomers **354** were synthesised ((*Z*) > 91%) With more sterically hindered silanes **1**,4-bis[methyl(3,3,3-trifluoropropylsilyl)]benzene **349** and 1,3-bis[methyl(3,3,3-trifluoroprorylsilyl)]benzene **350**, the polymerisation was carried out in the presence of $[RhI(cod)]_2$ **352**. All polymers **353–354** were obtained with high yields 54–96% and $M_n = 5000-22\ 000$ (Scheme 74).^{213–215}

The same catalyst was used in the synthesis of hyperbranched polymers by the homopolymerisation of bis(4ethynylphenyl)methylsilane **355**. The process resulted in polymer **356** with the (*E*)-regularity in 95% yield.²¹⁶ Dendrimeric structures were also obtained in the reaction of bis(1ethynylphenyl)dimethylsilane **327c** with dichloromethylsilane **357** in the presence of a Pt/C catalyst **307** (10% of Pt).

Bishydrosilylation leading to saturated products and α hydrosilylation was not visible. The further substitution of halogen in **358** with 1,4-bislithium-1,2,3,4-tetraphenylbuta-1,3diene **359** led to siloles **360**, while the reaction with lithiumethynylbenzene **361** led to the product **362** that can be further hydrosilylated with dichloromethylsilane **357** to build a more branched product (Scheme 75).²¹⁷



Scheme 71 Pd-catalysed hydrosilylation polyaddition of dihydrosilanes (217, 245, 339) to diynes (116c, 324) catalysed by Pd₂(dba)₃ 338/PCy₃.

Table 9Results of Pd 338-catalysed hydrosilylation of diynes (116c, 324)with dihydrosilanes (217, 245, 339) with diynes (Scheme 71)

						340					
Entry	Arene	Silane	$T\left[^{\circ}\mathbf{C}\right]$	<i>t</i> [h]		Yield [%]	$M_{\rm w}^{\ a}$	$M_{\rm w}/M_{\rm n}^{\ a}$	α : β^b		
1	116c	217	70	1	a	90	53 000	6.8	82:18		
2	324	217	70	8	b	94	12000	3.6	79:21		
3	116c	217	70	0.5	с	86	49000	6.6	78:22		
4	324	217	70	2	d	85	20000	4.2	78:22		
5 ^c	116c	339	110	6	e	91	20000	4.6	70:30		
6 ^{<i>c</i>}	324	339	110	9	f	88	12000	3.8	75:25		

^{*a*} Estimated by GPC using poly(styrene) standards. ^{*b*} Estimated from the ¹H and/or ²⁹Si NMR spectra. ^{*c*} PdCl₂(PCy₃)₂ **341** was used in place of Pd₂(dba)₃ **338**/PCy₃.

Sanchez *et al.* reported a special type of hydrosilylation of diynes using dihydrosiloles **282** and **368**. The reaction was carried out with different transition metal catalysts based upon Rh, Pt, and Pd catalysts (*e.g.*, $Pt_2(dvs)_2$ **194**, $RhCl(PPh_3)_3$ **205**, $Pd(PPh_3)_4$ **35**). The best results according to polymer molecular

weight, yield, and selectivity were obtained when heterogeneous H₂PtCl₆ 206 was used in boiling toluene. The reactions were carried out for 10 min-12 h. Very bulky 2,3,4,5-tetraphenylsilole 368, as well as silafluorene 282, were used in these polyaddition reactions. The process occurred by cis-addition of the Si-H bond to the $C \equiv C$ bond of divide forming exclusively (E)-products. In the case of other complexes, α -hydrosilylation or desilylative coupling was also observed. During the reaction, the selectivity was controlled sterically and kinetically. Less bulky groups such as silafluorene 282 required more accurate temperature control. At lower temperatures, (E)-products were obtained, while at higher temperature complex mixture of β - and α -hydrosilylation was observed. Bulky reagents such siliptycene (1,1-dihydrido-4,5,8,9-bis(triptycene)silafluorene) as 369d remained completely unreactive towards polyaddition. The structure of divne also influenced the polydispersity and molecular weight of the polymers 370 and 371 (Scheme 76). Obtained polymers were used as luminescence chemosensors for explosives. Cyclic siloles increase the efficiency of application of



Scheme 72 Pd-catalysed 338 polyaddition of methylphenylsilane 245 to diynes 116c and 324 with different ratios of crosslinking agent 304a and 342.

Table 10 Results of polyaddition of silane 245 to diynes 116c, 324 with different ratios of crosslinking agent 304a, 342 in the presence of Pd-catalyst 338 (Scheme 72)

Entry	Diyne	Crosslin. agent	Diyne:crosslin. agent	$T [^{\circ}C]$	<i>t</i> [h]	Polymer, yield [%] ^a	$M_{ m w}^{\ \ b}$	$M_{\rm w}/M_{\rm n}^{\ b}$
1	116c	_	100:0	60	4	343a , 75	24 000	3.8
2	116c	304a	95:5	60	4	343b, 70	26 000	4.1
3	116c	304a	90:10	60	4.5	343c, 68	30 000	5.0
4	116c	304a	80:20	60	3.5	343d, 80	130 000	16
5	116c	342	80:20	80	4.5	343e , 70	59 000	7.2
6	324	_	100:0	70	5	343f , 65	15000	3.1
7	324	304a	95:5	70	4	343g , 70	21000	4.0
8	324	304a	90:10	70	5	343h, 64	59 000	9.7
9	324	304a	80:20	70	3	343i , 75	110000	14

^{*a*} Reaction conditions: 245 (0.5 mmol), diyne + crosslinking agent (0.5 mmol), 338 (0.005–0.01 mmol, P/Pd = 2), toluene. ^{*b*} Estimated by GPC using poly(styrene) standard.



 Table 11
 Results of the synthesis of polymers
 347a-d
 via
 hydrosilylation

 reactions (Scheme 73)
 Figure 10
 <t

1 116c 345a 347a 80 112 000 7.5 2 324 345a 347b 79 56 000 9.4 3 116c 345b 347c 85 61 000 5.7 4 324 346 347d 85 42 000 4.1	Entry	Diyne	Alkyne	Polymer ^a	Yield ^b [%]	$M_{\rm w}^{\ c}$	$M_{\rm w}/M_{\rm n}^{\ c}$
	1 2 3 4	116c 324 116c 324	345a 345a 345b 346	347a 347b 347c 347d	80 79 85 85	$112\ 000\\56\ 000\\61\ 000\\42\ 000$	7.5 9.4 5.7 4.1

^{*a*} 272 (0.3 mmol), diyne **116c** or **324** (0.3 mmol), alkyne **345a–b**, **346** (0.315 mmol), $Pd_2(dba)_3$ **338**/PCy₃ (0.042 mmol in total, P/Pd = 2), benzene. ^{*b*} Purified by precipitation in benzene/2-propanol. ^{*c*} Estimated by GPC with poly(styrene) standards.

these vinylene-silole polymers as light-emitting diodes (LEDs), luminescent sensors, or organic charge carrier materials.^{68,218–220}

5.4. Cyclisation of 1,n-diynes by hydrosilylation reactions

Silanes and 1,*n*-diynes were also used in the hydrosilylation/ cyclisation to silylated (*Z*)-1,2-dialkylidenecyclohexanes, useful synthons in the synthesis of fine chemicals. The reactions occurred in the presence of metal complexes including Ni, Rh, Ru, Pd, Pt, but there are several limitations of each catalytic system. Described by Tamao et al., Ni(0) complexes catalysed the cyclisation of 1,7-diynes 160, 372-375, while 1,6-diynes were not active in this transformation.^{157,221} Different types of trialkyl, trialkoxysilanes, and silazanes 376a-e were applied as reagents. The reactions were carried out in the presence of Ni(acac)₂ (1 mol%) 273/DIBAH (2 mol%) 170, at 50-100 °C, for 6-24 h. The higher the temperature and the longer the reaction time, the lower yield of exocyclic diene was obtained, due to the subsequent polymerisation process. The cyclisation of terminal diynes occurred with moderate or good yield (47-73%) with the exclusive or predominant formation of (Z)-product. The process was effective also for optically active diyne 372. The asymmetric diyne 373 containing nitrogen led to the silyl-substituted tricyclic alkaloid-type dienes 379 with lower selectivity (Z)/(E)(379a/379b) = 79:21, suggesting a directing effect of the



Scheme 74 Temperature tunable stereoselective hydrosilylation of diynes 116c, 324 with disilanes 198a, 348–350 in the presence of RhI(PPh₃)₃ 351 and [RhI(cod)]₂ 352 catalysts.

nitrogen atom, which can easily coordinate to the metal centre. The internal diyne **374** was less reactive and reacted only under higher reaction temperatures and using accelerating triphenyl-phosphine as an additive. Unsymmetrical diyne **375** containing one terminal and one internal $C \equiv C$ bond reacted with the silane from the less shielded terminal acetylene furnishing a single regioisomer **381** in the post-reaction mixture (Scheme 77).^{157,221}

The mechanism of this transformation (Scheme 78) started from the insertion of one of the $C \equiv C$ bonds (less shielded) to the Ni-Si bond, generated by the oxidative addition of silane 376a-e to metal centre 382. The insertion of the second acetylene group to the Ni-vinyl bond 384 and reductive elimination of the exocyclic diene 381 closed the catalytic cycle. The insertion of acetylene to the Ni-H bond can be eliminated, because of the lack of other isomers in the post-reaction mixture.^{157,221} This Ni-catalysed reaction was possible only for 1,7-diynes. 1,6- or 1,8-diynes in the hydrosilylation process gave only polymeric products. To cyclise 1,6-diynes with hydrodisilanes, the reaction was catalysed with 5 mol% of Ni(acac)₂ 273/DIBAH 170/PEt₃ and the mechanism proceeded with the formation of Ni-silylene intermediate.¹⁵⁷ The obtained exocyclic dienes with (Z)-selectivity were used as reagents in Diels-Alder reactions, or the silyl groups were reacted in C-C bond forming reactions with aryl halides in Hiyama coupling reactions (Scheme 79).157,221

Widenhoefer *et al.* developed cationic Pt-complex, formed *in situ* from (phen)PtMe₂ **399** (phen = phenanthroline) and B(C₆F₅)₃ **401** that was highly active and selective in the cyclisation/hydrosilylation reactions of 1,6- and 1,7-diynes **127a**, **127k**, **127p**, and **394a-l** leading to silylated 1,2-dialkylidenecyclopentanes **402–410** and 1,2-dialkylidenecyclohexane **411**, with high (*Z*)-selectivity ((*Z*)/(*E*) > 8 : 1). The catalyst was found to be inactive in the cyclisation/hydrosilylation of separated dienes (for which palladium analogs were active), making this process highly selective.^{222–225} The reactions were carried out for 10 min–3 h at 110 °C in toluene for different silanes **207a**, **395–398** (Scheme 80).¹⁵⁸

The same group developed a diimine cationic Pt complex, $[PhN=C(Me)C(Me)=NPh]PtMe_2$ 400, which was much more active and selective than the complex with phenanthroline **399**. The products **402–411** were obtained in 15 min at 110 °C or 85 min at 70 °C with higher selectivity towards (*Z*)-isomer (*Z*):(*E*) > 30:1. The electronic and steric properties of the diimine ligands were found to have an important influence on the cyclisation/hydrosilylation reaction. The rate of the process decreased with the increase of the electron density and steric bulk of the ligand. The structure of silane and diyne also influenced the reaction rate. When $HSi(i-Pr)_3$ **398** was used instead of $HSiEt_3$ **207a**, the reaction was 10 times slower. The catalytic system was tolerant towards many functional groups including *inter alia* sulfones, amides, ketones (Scheme 80).¹⁵⁹

The authors proposed the mechanism of this transformation (Scheme 81). Initially CO or $B(C_6F_5)_3$ **401** abstracts the methyl group from the pre-catalyst **400** forming the Pt-cationic complex **412** upon coordination to the diyne substrate. Oxidative addition of the silane, which occurred readily, even at -30 °C leads to complex **413**. Loss of CH₄ leads to **414**. Next, the insertion of the alkyne into the Pt–Si bond occurs leading to **415**, followed by the β -migratory insertion of the coordinated second alkyne group and formation of the platinum dienyl intermediate **416**. The oxidative addition of silane **207a** or **395–398** formed **417**. Elimination of the product **402–411** and the coordination of diyne regenerates the initial catalyst **414**. The obtained cyclic products were used in protodesilylation and Diels–Alder transformations. Examples of these processes are presented in Scheme 82 using **402a** as a reagent.^{158,159}

Several papers discussed the application of Rh complexes in the synthesis of 1,2-dialkylidenecyclopentanes. The use of the popular Wilkinson's complex **205** in this transformation was reported by Matsuda *et al.*^{226,227} The exact catalyst, which facilitated the formation of cyclic compounds was the complex $Rh(H)(SiR_3)Cl(PPh_3)_2$ **438**, which was obtained by oxidative addition of silane to the metal centre. The order and time of addition of silane and diyne were important for the reaction course. When reagents **127a**, **127k**, **127p**, **195a** and **394a–l** were added 1,2-dialkilidenecyclopentane **434** was formed immediately. In other cases, indane **435** was formed as the main product (Scheme 83). Scheme 84 shows various dialkilidenecyclopentanes



Scheme 75 Preparation of hyperbranched organosilicon compounds using hydrosilylation of diynes.

434 obtained within this transformation in the presence of **438**. Depending on the catalyst structure (*E*)- or (*Z*)-cyclic isomers were obtained. When mono- or bidentate electron-donating phosphine ligands are coordinated to the metal centre RhCl(PPh₃)₃ **205** or

 $[Rh(cod)(dppb)][PF_6]$ **439**, the insertion of the second alkyne group is slowed down giving time to convert the (*Z*)-isomer **434** into (*E*)-product **434**. For $Rh_4(CO)_{12}$ **440** with electron-withdrawing CO ligands, the insertion process is much faster, and there is no time





Scheme 77 Synthesis of 1,2-dialkylidenecyclohexanes 377–381 via catalytic cyclisation of 1,7-diynes 160, 372–375 with silanes 376a–e catalysed by Ni(0) complex 273.



Scheme 78 Mechanism of cyclisation *via* hydrosilylation of 1,7-diynes with silanes catalysed by Ni(0) complex generated *in situ* from **273/170**.

for the formation of (*E*)-isomer (Scheme 85).^{226,227} The obtained products were used in Diels–Alder transformations with different dienophiles, as well as in the hydrogenation process catalysed by the Pd/C **446** system, followed by the homologation reaction (Scheme 86).

Ojima *et al.* reported several papers based on the cyclisation/ hydrosilylation of 1,6-diynes **1270–p**, **450a–f** in the presence of rhodium complexes **204**, **451–452** and different pressures of CO.^{66,67,228–230} The course of the reaction strictly depended on the reagents (silane, diyne) structure, the type of the catalyst, as well as the pressure of CO. When $Rh_2Co_2(CO)_{12}$ **451**, $Rh(acac)(CO)_2$ **204** or Rh(t-BuNC)₄CO(CO)₄ **452** were used, the corresponding bicyclo[3.3.0]octenones **453–455** were obtained in 82–93% yield *via* carbobicyclisation with the incorporation of CO (15–50 bar) (Scheme 87 and Table 12). Under lower CO pressures (1–2 bar), no reaction with CO was observed and typical dialkilidenecycloalkanes were formed.²³⁰ Moreover, the steric hindrance of silane or diyne influenced the formation of a specific product. Additionally, the C₄ position in 1,6-diynes **450a–f** exerts marked influence on the product distribution. When the heteroatom is at the C₄ position 1,2-hydrosilylation is the main process, while 1,4-hydrosilylation is favoured with 4,4-*gem*-disubstitution with ester groups.²²⁸ Products **453** can easily isomerise quantitatively to **454** in the presence of RhCl₃·3H₂O **213** as a catalyst in ethanol under 50 °C.

A detailed mechanism of this transformation was also presented which explained the formation of various cyclic products. The product outcome was found to be dependent on the further transformations of complex **459** formed in the carbocylisation of **457** to **458**, followed by CO insertion and subsequent carbocyclization to bicyclic **460**. When a 1,3-[Rh]shift occurred from **460**, the complex **461** is formed, which then after reductive elimination furnishes product **453**. When β -hydride elimination occurred from **460**, the dienone–M–H complex **462** is formed or/and bicyclic diene **464**. The addition of the M–H species leads to intermediates **463** or **465**, which next (*via* addition of the next silane and reductive elimination of R₃Si–[Rh]) accomplishes the products **454** or **455**. The formation of **455** was observed only for the product which was able to form a stable aromatic pyrrole product (Scheme 88).²²⁹

Ojima discussed also that endiynes **466** (dodec-11-ene-1,6diyne or their heteroatom analogs) reacted with silanes (PhMe₂-SiH **195a**, Et₃SiH **207a**, or (*t*-Bu)Me₂SiH **395**) in the presence of Rh(acac)(CO)₂ **204** in unique silylative cascade carbonylative carbotricyclisation process, at room temperature and under ambient pressure of CO. The reaction yielding fused 5-7-5 tricyclic products 5-oxo-1,3*a*,4,5,7,9-hexahydro-3*H*-cyclopenta[*e*]azulenes **467** or their heteroatom congeners. Within this process, functionalised polycyclic compounds were obtained that are useful synthons in the synthesis of natural products (Scheme 89).^{66,67}

Using the same $[Rh(acac)(CO)_2]$ complex 204, it was possible to carry out hydrosilylative cyclisation with carbonylation of



Scheme 79 Transformations of 1,2-dialkylidenecyclohexanes with vinylsilyl group 377 in Diels-Alder and Hiyama coupling reactions



Scheme 80 Cyclisation/hydrosilylation of diynes 127a, 127k, 127p, and 394a–l catalysed by a 1:1 mixture of Pt catalyst (399 or 400) and $B(C_6F_5)_3$ 401 in toluene at 110 °C towards exocyclic dienes 402–411.

various 1,5-diynes with aromatic, olefinic, and ethylene tethered spacers **468** under the ambient pressure of CO (1 atm.). The reaction furnished various 2,5-dialkylidenecyclopentanones **469** in good yields. In this example, the insertion of CO was favoured to build a five-membered ring and avoid high strains. The products **469a–m** were obtained with moderate or high yields 30–92%, which varies with both reagent structures (Scheme 90).²³¹ The mechanism of carbonylative cyclisation of 1,5-diynes **468** using [Rh(acac)(CO)₂] **204** started from silylrhodation, followed by the insertion of CO to **471** to form acylrhodium species **472**, then acylrhodation to the second alkynyl group forms the 5-membered ring **473** (Scheme 91).²³¹

Cyclisation/hydrosilylation of 1,6-, 1,7- and 1,8-diynes 127a, 127o-p, 160, 164a, 474a-d was carried out in the presence of

ionic Pd complex $[(\eta^3 \cdot C_3H_5)Pd(cod)][PF_6]$ **476** with chlorodimethyl-**305**, dichloromethyl-**357**, and trichlorosilane **475**. The reaction occurred at room temperature in CH₂Cl₂ and products (*Z*)-1-methylene-2-silylmethylenecycloalkanes **477** were obtained in good yields, which were further transformed to their ethoxy analogs **478** (Scheme 92). For unsymmetrical diyne 2-butynyl propargyl ether **474b**, it was found by NOE analysis that the silyl group is attached to the internal C \equiv C bond, suggesting that the formation of the regioisomer **478f** was due to the fact, that the reaction started from the hydropalladation at the terminal alkyne site to **481**, instead of the insertion of the alkyne into the metal–Si bond. The further steps in the plausible mechanism are: intramolecular carbopalladation, the formation of cyclised (*Z*)-alkenylpalladium intermediate **482**, and finally σ -metathesis with a hydrosilane,



Scheme 81 Mechanism of the hydrosilylation/cyclisation reaction of separated diynes in the presence of Pt-diimine complex 400 and B(C₆F₅)₃ 401.

followed by the product release **477** and regeneration of the initial catalyst **479** (Scheme 93).^{232–234}

Liu and Wiedenhoefer reported that the cationic rhodium complex [Rh(BINAP)(cod)][BF₄] **485** (BINAP = ((\pm)-2,2bis(diphenylphosphino)binaphthyl)) is as an effective catalyst (10 mol%) in the cyclisation/hydrosilylation reaction of terminal and internal 1,*n*-diynes **127a**, **483a–m** with silanes **195d**, **207a**, and **484** leading to 1,2-dialkilidynecycloalkanes **486a–p** with high yield and high diastereoselectivity (Scheme 94). The higher the concertation of silane the lower the selectivity. The best results were obtained when triethylsilane **207a** was used (1.0–1.7 M) and was added slowly to the reaction mixture. The mechanism of this transformation is similar to that previously described for Pt-complexes (Scheme 81).¹⁶⁰ The obtained silylated 1,2-dialkilidynecycloalkanes **486** were further used in the Diels–Alder reaction with 4-phenyl[1,2,4]triazole-3,5-dione **430** at 0 °C for 30 min (Scheme 95).

Another example that involved the reaction of diynes **116a**, **127c**, **127o-p**, **488a-c**, *tert*-butyldimethylsilane **395**, and CO, which furnished two different catechol derivatives **490** and **491**, was carried out in the presence of $[Ru_3(CO)_{12}]$ **489**/PCy₃. Product **490** was the primary product, which can be readily hydrosilylated to **491** when 6 equiv. of silane **395** was used.

In the case of a lower excess of **395**, product **490** was visible in the reaction mixture. The best results were obtained using acetonitrile as a solvent. Various terminal and internal diynes were used as reagents giving products with moderate yields (Scheme 96).²³⁵ The mechanism of the process involves the formation of an oxycarbyne complex **493** as an intermediate and the process requires the introduction of two CO molecules into the diyne structure. A carbyne/CO coupling yields intermediate **493** was previously tested for tungsten.²³⁵ Analog complexes to **495** for alkynes were determined for other metals, *e.g.*, Nb, Ta, V, Ta. Katz *et al.* reported the formation of a similar product in the reaction of (CO)₄BrM \equiv CCH₃ (M = Cr, W) with diyne.²³⁶ This proved that the proposed mechanism is plausible (Scheme 97).

Lewis acids such as AlCl₃ **496** and EtAlCl₂ **497** were successfully applied for the hydrosilylation of alkynes with trialklylsilanes, which occurred as a *syn*-addition of the Si–H bond to the C \equiv C bond with the formation of the *trans*-product.^{57,237} The mechanism of this transformation assumes the formation of a zwitterionic intermediate by the coordination of **496** or **497** to the acetylenic bond. Next, the hydride of silane attacks the electron-deficient carbon atom from the opposite site to AlX₃ with the formation of ate-complex. The coupling between the





Scheme 83 Possible silylative cyclisation reactions of diynes **394a–l** with dimethylphenylsilane **195a**. Different products were formed depending on the order of reagent addition.



Scheme 84 Dialkilidenecyclopentanes 434a-l obtained from the silylative cyclisation of diynes with silanes in the presence of Rh(H)(SiR₃)Cl(PPh₃)₂ 438.



Scheme 85 Proposed mechanism for the formation of (*Z*)- and (*E*)isomers of **434** in the silylative cyclisation reaction in the presence of Rh-catalysts **205**, **439**, or **440**.

silyl cation and vinyl group furnishes the silylated olefin with retention of configuration. The same catalysts **496** and **497** were also used in the hydrosilylation of hepta-1,6-diyne **116a** and octa-1,7-diyne **160**, using 4 equiv. of triethylsilane **207a**. For a shorter chain of terminal diyne **116a**, the cyclic product **498** was obtained in 60% yield, while for octa-1,7-diyne **160**, 1,8-bistriethylsilyl-octa-1,7-diene **499** was formed predominantly

(Scheme 98).²³⁷ Formation of bissilylated diene using this Lewis catalyst contrasts with the cyclization process *via* hydrosilylation, which occurred in the presence of Ni or Rh catalysts.^{157,221}

6. Hydrogermylation of conjugated and separated diynes

Hydrogermylation of diynes is limited only to two examples, which describe the formation of 2,5-disubstituted germoles²³⁸ or germylene–divinylene polymers.²³⁹

Murakami *et al.* developed a *trans*-hydrogermylation of conjugated symmetrical and nonsymmetrical 1,3-diynes **1a–b**, **1d**, **27c**, **60e**, **258i**, **258o**, **500a–d** with diphenylgermane **501** in the presence of [Cp*Ru(MeCN)₃][PF₆] **281** wich yielded cyclic germoles **502a–o** with good or moderate yields (Scheme 99).

The same complex **281** was previously used by Trost *et al.* in the *trans*-hydrosilylation of alkynes, 56,189,240,241 but its activity in the reaction with conjugated diynes was much lower than for hydrogermylation (Table 13). The double addition of diphenyl-germane **501** to 1,4-diphenyl-buta-1,3-diyne **1a** occurred with a much higher yield in comparison to the hydrosilylation reaction (93% *vs.* 29%) (Scheme 58).¹⁵⁶ The hydrogermylation reaction was carried out with 3 equiv. of germane **501** and 10 mol% of Ru catalyst **281**, but a lower excess of reagent **501** was also possible (1.2 equiv.).

The reaction was efficient for diaryl-substituted diynes with different functional groups (silyl, boryl, methoxy, bromo, fluoro) or compounds with heteroaryl substituents **27c**, **500b**. The presence of strongly electron-withdrawing nitro groups in the para position **500a** was responsible for the lower product yield (40%, **502i**). No reactions were observed for diynes with alkyl groups (hexa-2,4-diyne **65a**) and silyl functionalities (1,4-bis-(trimethylsilyl)buta-1,3-diyne **180c**). When dibutylgermane **503** was used as a reagent, the reaction was less effective, even with 20 mol% of **281**. Applying conjugated 1,8-diphenylocta-1,3,5,7-tetrayne **504**, diphenylgermane **501** (6 equiv.), and 20 mol% of



Scheme 86 Hydrogenation of dialkilidenecyclopentane 434d followed by a Ti-catalysed homologation reaction.



Table 12 Results of silacarbocyclisation of 1,6-diynes1270-pandR450a-f catalysed by various Rh complexes204, and 451-45256

Entry	Diyne	Silane	Rh	CO (bar)	$T\left[^{\circ}\mathrm{C}\right]$	Yield ^a [%] 453/454/455
1	450a	395	451	15	50	93/0/0
2			204	15	50	93/0/0
3			452	15	50	82/0/0
4	450b	395	204	50	50	70/0/0
5	450c	395	204	50	50	47/16/0
6	450d	395	204	50	120	73/0/0
7	450e	395	204	50	65	0/18/70
8	450e	395	452	50	65	0/10/57
9	450f	207a	204	50	65	0/0/58
10	450f	207a	452	50	65	0/62/0
11	1270	395	204	50	66	0/22/56
12	127p	395	204	50	65	27/22/0

^{*a*} Isolated yield, reaction time 10–20 h.

Ru complex **281** it was also possible to obtain 2,2'-bigermole **505** with 56% yield (Scheme 100).²³⁸

Diphenylgermane **501** was also used as a reagent in the hydrogermylation of various diynes, with aryl or alkyl spacers between the $C \equiv C$ bonds leading to germylene-divinylene polymers. The polymerisation was effectively catalysed in the presence of 0.9 mol% of Pd catalysts (Pd₂(dba)₃ **338**/2PCy₃, PdCl₂(PCy₃)₂ **341**). The reactions were carried out at 50–90 °C and the polymers **506** were obtained with $M_w = 12\,000-83\,000$ and $M_w/M_n = 3.3-12.0$ (Scheme 101). They were isolated by precipitation in benzene/propan-2-ol solution. Due to the high conjugation, the germylene-divinylene polymers **506a-d** indicated intense light emission depending on the structures of the monomers. The best results were obtained for anthrylene



Scheme 88 Various catalytic pathways in silacarbocyclisation of 1,6-diynes 1270-p and 450a-f furnishing products 453-455.



Scheme 89 Carobocyclisation of enediynes 466 catalysed by Rh catalyst204. Construction of functionalised fused 5-7-5 ring systems 467.

polymer which gave intense and broad UV-Vis spectra from 420 to ≥ 600 nm with λ_{max} peaks at 440, 464, and 534 nm.²³⁹

7. Hydrostannation

Alkenylstannanes are useful building blocks in the synthesis of various organic compounds (also complex molecules as pharmaceuticals or natural compounds) due to their ability to the formation of the new C–C bonds in Stille coupling reactions.^{242–247} The hydrostannation of alkynes, which can occur under a free radical manner, in the presence of a transition–metal catalyst or *via* a hydrogen atom transfer reaction (with trialkyltin hydride used as a nucleophilic species), is the most convenient and popular method for the synthesis of alkenylstannanes.^{18,54–56} Despite several papers focused on the hydrostannation of alkynes, the literature concerning the addition of the Sn–H bond to the C \equiv C bonds in diynes is limited to a few examples based on radical or transition metal-catalyzed transformations.

7.1. Radical hydrostannation of conjugated and separated diynes

Radical hydrostannation was successfully applied in the reaction with 1,3-diynes,²⁴⁸ as well as diynes possessing an aryl spacer between the $C \equiv C$ bonds.²⁴⁹



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Scheme 91 Proposed mechanism of carbonylative silylcarbocyclisation of 1,5-diynes 468 with silanes catalysed by [Rh(acac)(CO)₂] 204.

Konno *et al.* reported selective radical hydrostannation of 5-benzyloxy-1-trifluoromethyl-5-methyl-hexa-1,3-diyne **507** with tributyltin hydride **508**. The radical is generated from $HSn(n-Bu)_3$ **508** in the presence of Et₃B **509** and oxygen (Scheme 102).

Despite the fact that even eight different products might be obtained in hydrostannation due to the presence of double $C \equiv C$ bonds and different substituents in terminal positions, some of the products might be eliminated. The attack of the radical on the carbon in position β - or γ -can be excluded



Scheme 90 Carbonylative hydrosilylation of 1,5-diynes 468 with silanes 195a, 195e, 207a, 376a in the presence of [Rh(acac)(CO)₂] 204 and 1 bar of CO.



Scheme 92 Cyclisation-hydrosilylation functionalisation of 1,6-, 1,7- and 1,8-diynes 127a, 127o-p, 160, 164a, 474a-d catalysed by $[(\eta^3-C_3H_5)-Pd(cod)][PF_6]$ 476.



Scheme 93 Mechanism of the cyclisation-hydrosilylation reaction of separated diynes catalysed $[(\eta^3-C_3H_5)Pd(cod)][PF_6]$ **476**.

because of the lack of resonance of the vinyltin radical. The radical, which has a nucleophilic character attacks the more electrophilic α -carbon atom in **507** with a strong electron-withdrawing CF₃ group. Moreover, the bulky group in δ -position limits the access of the organotin group, therefore product **510** is formed with high regio- and stereoselectivity (Scheme 103). The obtained enyne **510** was generated in 75% yield and was further used in the synthesis of CF₃-substituted (*Z*)-enediyne **517** compounds in iododestannylation/Sonogashira coupling reactions (Scheme 104).²⁴⁸

The radical hydrostannation of various diynes and triynes **102c** and **264a–e** was carried out stereoselectively with tributyltin hydride **508**. In the two cases, the (*E*)-products **521a** and **521c** were exclusively formed. The hydrostannation of other diynes **264a–c**, and **264e** occurred with lower selectivity, but still with an excess of the (*E*)-products **521** (Scheme 105). To obtain high selectivity, an elevated temperature (80 °C) has to be maintained. Under lower temperatures, the conversion was not complete and other isomers were also formed. The authors proved the (*E*)-selectivity of the products through ¹H NMR spectroscopy by the large coupling constant of the vinyl group ($J_{H-H} = 18-19$ Hz) and the characteristic values for tin hydrogen coupling ($J_{Sn-H} = 124-138$ Hz). For 1,3,5-tris[(*E*)-2-(tributylstannyl)vinyl]benzene **521f**, the authors carried out Stille coupling with various bromo-substituted chromophores **522** in the





phenyl[1,2,4]triazole-3,5-dione at 0 °C **430** via Diels–Alder reactions.

presence of PdCl₂(PPh₃)₂ **94** catalyst and CsF **523** or CuI **519** as additives. The products were obtained with moderate or good yield with the retention of the configuration, showing the utility of organotin compounds (Scheme 106).²⁴⁹ Previously published papers described that radical hydrostannation of diynes led to the mixture of various isomers, which is in opposition to the above-reported results.²⁵⁰

The non-catalysed addition of organotin compounds to the $C \equiv C$ bond in elevated temperatures occurs relatively easily due to the weak Sn–H bond. The application of dihydrides in the hydrostannation of diynes may lead to cyclic or polymeric products which can be controlled by appropriate selection of the substrates and reaction conditions.²⁵¹ The addition of Bu₂SnH₂ **525** to penta-1,4-diyne **526a** in refluxing heptane followed by heating the reaction mixture to 200 °C gave a sixmembered heterocycle **528a** with 43% yield. The product was distilled from a viscous polymeric residue together with the

small amount of five-membered by-product 530a. Generally, the terminal addition (path A) of the Sn–H bond to the $C \equiv C$ bonds yielded a six-membered heterocycles 528, whereas the non-terminal addition (path B) led to five-membered adducts 530 (Scheme 107). The regioselectivity could be controlled by the proper selection of the substituents attached to C_{sp} carbon. When the hepta-2,5-diyne 526b or 1-phenyl-1,4-pendadiyne 526f were used the five-membered heterocycles 530b and 530f were formed as the major regioisomers. The application of monoalkyl-substituted 1,4-diynes 526c-e on the other hand gave in an excess stannabenzene derivatives 528c-e. The authors suggested that radical-stabilising substituents in 1,4-divnes mainly led to stannoles 530, whereas 6-substituted hexa-1,4-divnes 526g-i gave the six-membered adducts 528g-i (Scheme 107).^{252,253} The substitution of CH₂ spacer between alkynyl groups, in the case of 3-organyl-substituted 1,5-diynes, did not influence process regioselectivity leading mainly to the six-membered products.²⁵⁴⁻²⁵⁶

The hydrostannation of diynes possessing the p-block element as a linker between alkynyl groups gave in major an attractive sixmembered rings with two heteroatoms, which are useful synthons in organic synthesis. For instance, the hydrostannation of (dialkylamino)dialkynylboranes **531a–c** with dimethylstannane **532** yielded 1,1-dimethyl-1-stanna-4-bora-2,5-cyclohexadies **533**, which could be further converted *via trans*-amination to 4-amino derivatives **534** or *via* solvolysis of **533** to alkoxy derivatives **535**. These latter were precursors for 4-alkyl-1,1dimethyl-1-stanna-4-boracyclohexadienes **537** or lithium-1,1,2,4,4,6-hexamethyl-1-stanna-4-borata-2,5-cyclohexadiene which



Scheme 96 Ru₃(CO)₁₂ 489 catalysed carbocyclisation of 1,6-diynes 116a, 127c, 127o-p, 488a-c with HSi(t-Bu)Me₂ 395 and CO.



Scheme 97 Mechanism of the Ru-catalyzed 489 reaction of 1,6-diynes with silanes and CO

were obtained in high yield 536 (Scheme 108).²⁵⁷ Analogous ring systems with different heteroatoms could also be obtained for the hydrostannation of diynes containing Si, Sn, or P atoms as spacers between alkynyl groups.258-260

The diyne structure, as well as reaction conditions, have a crucial influence on the product formed. The hydrostannation of α, ω -divnes, such as 1,4-diethynylbenzene 116c, nona-1,8diyne 164a, and hexa-1,5-diyne 541 with diorganotin dihydrides 525, 532, 538-540 at high temperatures gave rubber-like polymers. However, for hexa-1,5-diyne 541 small amounts of 1-stanna-2,6-cycloheptadiene 544 derivatives were isolated as well. The polymer formation occurred via intermolecular



Scheme 98 Lewis acid (496 or 497) catalysed hydrosilylation of hepta-1,6-diyne 116a or octa-1,7-diyne 160

poly-addition of alkenyldiorganoltin hydride 542 whereas, the cyclic product is obtained through its intramolecular cyclisation (Scheme 109). The molecular weight of the polymers depended on both the α,ω -divnes, and organotin compounds (Table 14).²⁶¹ Similar observations were made when p-phenylene-bis(dimethyltin hydride) 545 was used in the poly-addition to α,ω -diynes.²⁶²

The appropriate selection of the reaction condition was also crucial for the synthesis of tin-containing seven-membered heterocycles (stannepines) by the hydrostannation of (Z)-endiynes. Mild reaction conditions and the presence of base led to the desired heterocycles instead of polymeric material.²⁶³

The hydrostannation of o-diethynylbenzene 545 with diorganotin hydrides 532, 538, 540, and 546 yielded, in addition to polymers 549, the seven and fourteen-membered tin-containing heterocycles (547 and 548) with low or moderate yields.^{264,265} The highest yield of the fourteen-membered ring system was observed for 548d when ethylphenyltin dihydride 546 was used, whereas the benzostannepin 547b was formed in 22% yield when diethyltin dihydride 538 was applied. Nevertheless, in all cases, the polymers were the main products (Scheme 110 and Table 15).



Scheme 99 Double *trans*-hydrogermylation of 1,3-diynes **1a–b**, **1d**, **27c**, **60e**, **258i**, **258o**, and **500a–d** with diphenylgermane **501** in the presence of [Cp*Ru(MeCN)₃][PF₆] **281**.

 Table 13
 Synthesis of 2,5-disubstituted germoles
 502a-o
 via
 double

 trans-hydrogermylation of 1,3-diynes with diphenylgermane
 501

Entry	\mathbb{R}^1	R^2	Diyne	Product	Isolated yield [%]						
1	Ph	Ph	1a	502a	93 $(90)^b$						
2	$4-FC_6H_4$	$4-FC_6H_4$	1b	502b	66						
3	4-MeOC ₆ H ₄	4-MeOC ₆ H ₄	1d	502c	80						
4	3-Thienyl	3-Thienyl	27c	502d	94						
5	$3-BrC_6H_4$	$3-BrC_6H_4$	258i	502e	93						
6	Cyclohexen-1-	Cyclohexen-1-	2580	502f	70						
	yľ	yl									
7	2-Naphtyl	2-Naphtyl	280a	502g	87						
8	$3-(pin)BC_6H_4$	3-(pin)BC ₆ H ₄	280d	502ĥ	91						
9	$4 - O_2 NC_6 H_4$	$4 - O_2 NC_6 H_4$	500a	502i	40						
10	5-Pyrimidyl	5-Pyrimidyl	500b	502j	69						
11	Ph	4-MeOC ₆ H ₄	60e	502k	95						
12	Ph	4-CNC ₆ H ₄	500c	502l	71						
13	Ph	4-ViC ₆ H ₄	280e	502m	87						
14	4-MeC ₆ H ₄	$n-C_6H_{13}$	500d	502n	44						
15	4-Me ₃ SiC ₆ H ₄	4-Me ₃ SiC ₆ H ₄	500e	502o	75						
^a Diyı MeCH	^a Diyne: 501 = 1:3, 1,2-dicholoroethane, r.t., 10 mol% of $[Cp*Ru(-MeCH)_{a}][PE_{a}]$ 281 ^b 1.2 equiv of 501										

The obtained heterocyclic compounds and polymeric materials could be readily transformed with the retention of configuration into alkenyl iodides by the reaction with I₂ **418**. Polymer degradation with iodine **418** revealed that the polymer product contained *Z*,*Z*-, *Z*,*E*- and *E*,*E*-units.²⁶⁵

The tin-containing six-membered heterocycles are attractive precursors in the preparation of various 15 group heterobenzenes.^{266,267} In 1971, Ashe reported the synthesis of arsabenzene 555 based on the arsenic/tin exchange. The l,4dihydro-l,l-dibutylstannobenzene 528a was converted in a one-step procedure to desired product 555 through the reaction with the arsenic trichloride 553. Similarly, 528a reacted with phosphorus tribromide 550 to give phosphabenzene 552.268 The same research group extended the scope of 15 group heterobenzenes to stibabenzenes 558²⁶⁹ and bismabenzene 561²⁷⁰ in an analogous manner. However, treatment of l,4-dihydro-l,l-dibutylstannobenzene 528a with SbCl₃ 556 or BiCl₃ 559 gave 1-chloro-1stibacyclohexa-2,5-diene 557 or 1-chloro-1-bismacyclohexa-2,5diene 560, respectively. The group 15 heterobenzenes underwent Diels-Alder reactions with hexafluorobutyne 562 to give 563. The reactivity of heterobenzenes increased with the higher atomic number of heteroatom. For instance, stilabenzene 558 reacted rapidly with hexafluorobutyne 562 at 0 °C, arsabenzene 555 at





Scheme 101 Polymerisation of 1,*n*-diynes 116c, 164a,164i, and 324 with diphenylgermane 501 *via* hydrogermylation reaction catalysed by Pd-complexes 338 and 341.



Scheme 102 Radical hydrostannation of conjugated diyne 507 with tributyltin 508 followed by the assignment of the stereoselectivity in the destannation reaction.



room temperature, whereas phospabenzene **552** was converted to Diels–Alder adduct at 100 °C (Scheme 111).²⁷⁰ The 2- and 4-subtituted heterobenzenes could be also synthesised through hydrostannation of appropriate 1,4-diynes. Further transformation of the stannabeznes to the phospha- or arsabeznene derivatives could also be achieved.^{271–273} The same procedure was adopted to the formation of borabenzenes.²⁵³

The hydrostannation of penta-1-4-diyne 526a with Bu_2SnH_2 525 was used in the synthesis of 13-thiaarachidonic acid 573. Compounds 573 is a time- and O_2 -dependent irreversible inhibitor of soybean lipoxygenase and was prepared in reaction sequence presented in Scheme 112.²⁷⁴ The process was characterised by excellent stereoselectivity and satisfactory yields of each of the individual reaction steps.

7.2. Transition metal-catalysed hydrostannation of diynes

The phosphine-free palladium Pearlman's catalyst $Pd(OH)_2/C$ 575 was found to be effective in the hydrostannation of 1,6-diynes **127a**, **127j**, **127m**, **127p**, **394d** and **574a–c** with $HSn(n-Bu)_3$ **508**, which generated 1,2-dialkylidenecyclopentenes with the



Scheme 106 Cross-coupling reaction of electrophiles RX 522 with 1,3,5-tris[(E)-2-(tributylstannyl)vinyl]benzene 521f catalysed by PdCl₂(PPh₃)₂ 94.

tributylstannyl group **576**. This stannylative coupling was effective for various 1,6-diynes, including those possessing hydroxyl groups or protected alcohols, as well as reagents with heteroatoms in the propargylic position. The reactions occurred with high yields of the products **576** (58–95% yield) (Scheme 113). Several other complexes such as Pd₂(dba)₃ **338**, Pd/C **446**, Pd(acac)₂ **577** gave the desired cyclised product **576a** with the yield >75%. Adding 1 or 2 equiv. of PPh₃ or dppb to Pd₂(dba)₃ **338**, gave a complex postreaction mixture with less than 15% of **576a**. The authors suggested that the phosphine coordinates to the metal centre, blocking the possibility chelate formation with the 1,6-diynes **127a**, **127j**, **127m**, **127p**, **394d**, and **574a–c**. The mechanism of the process begins with the oxidative addition of HSn(*n*-Bu)₃ **508** and chelation of the 1,6-diyne to give **579**. The formation of the product might occur within two possible pathways based on stannylpalladation (path A) or hydropalladation/carbopalladation (path B) (Scheme 114). There was no information on which cycle is more probable. Terminally substituted 1,6-diynes **584a–c** were also reactive in this cyclisation





Scheme 108 Hydrostannation of (dialkylamino)dialkynylboranes 531a-c with dimethylthin dihydride 532, and further transformations.



Scheme 109 Synthesis of linear polymers 543 through the hydrostannation of α, ω -diynes 116c, 164a, 541 with diorganotin dihydrides 525, 532, 538-540.

Table 14Results of hydrostannation of α, ω -diynes116c, 164a, 541 withdiorganotin dihydrides525, 532, 538–540 (Scheme 109)

Review Article

Entry	α,ω-Diyne	R_2SnH_2	Molecular weight, $M_{\rm w}$	Degree of polym., <i>n</i>
1	541	540	75 000 ^a	170
2	541	532	b	_
3	541	538	b	_
4	541	539	50 000	180
5	541	525	50 000	160
6	164a	540	100 000	250
7	164a	525	45 000	130
8	116c	540	65 000	160
a			h h h	

^{*a*} Molecular weight of benzene-soluble fraction. ^{*b*} Polymer insoluble in benzene after heating under vacuum.

reaction, but the electronic properties of the diyne substituents strongly influenced the selectivity, and a mixture of cyclised **585** and **586** and linear **587** vinylstannanes were generated. The linear product was predominantly formed (**587c**, 59%) in the case of hydrostannylation of the silyl-substituted reagent **584c** (Scheme 115).^{275,276} The obtained dialkylidenecyclopentenes functionalised with stannyl group **576a–c** were used in several destannylation reactions: Diels–Alder with *N*-phenyl maleimide **387** to **588** followed by protodestannylation to **589**, Stille coupling with *p*-iodoanisole **58**, and homocoupling of **576a**, showing the high utility of this reagent in organic synthesis (Scheme 116).

Furstner et al. reported that conjugated 1,3-diynes 594, as well as non-conjugated 1,n-diynes 595 (with an unprotected hydroxyl group in the propargyl position), underwent double or site-selective trans-monohydrostannation depending on the reaction conditions in the presence of catalytic [Cp*RuCl]₄ 596. The process was found to be temperature-dependent. When the reaction was carried out in boiling 1,2-dichloroethane (at 80 °C), the site-selective reaction is favoured, while at a lower temperature (especially at -40 °C) bishydrostannylation occurred in a large amount. Irrespective of the alcohol type (primary, secondary, or tertiary) the *trans*-hydrostannylation occurred with high selectivity (Scheme 117). Additionally, the type of substituent attached to the second alkyne influence the process selectivity with bulkier groups giving better selectivity towards trans-hydrostannation. The selectivity of the stannylation of divne 594 from the propargylic side resulted from the hydrogen bonding of OH with the polarised [Ru-Cl] bond of 596. The propargylic alcohol readily forms an adduct with the Ru-complex under room or higher temperature, while binding the alkyne

 Table 15
 Results for hydrostannation of o-diethynylbenzene
 545 with

 diorganotin hydrides
 532, 538, 540, and 546 (Scheme 110)
 532, 538, 540, and 546 (Scheme 110)

	$R^1 R^2$	SnH_2			Yield [%]				
Entry					547	548	549		
1	532	Ме	Ме	a	$10(6)^{a}$	_	80		
2	538	Et	Et	b	$22(17)^{a}$	_	50		
3	540	Ph	Ph	с		$17(12)^{a}$	70		
1	546	Ph	Et	d	5	$41(25)^{a}$	50		
7 • • •	(2) 6								

^a Yield (%) after extensive purification.

occurs only at a lower temperature. The reaction was also effective for *trans*-monohydrostannation of 1,*n*-diynes **595** to give products **599a-e** (Scheme 118). The strong directing effect of the hydroxyl group in the propargylic position was responsible for the high process selectivity. The stannyl-substituted products might be directly transformed to (*E*)-conjugated enynes by the protodestannation reaction with copper diphenylphosphinate CuOP(O)Ph₂ **601** in DMF. The site-selective *trans*-hydrostannation was applied in the total synthesis of typhonoside series of glycolipids **608** and **614**, which have neuroprotective properties (Scheme 119). Moreover, the application of this transformation permitted for late-stage modification of the bioactive compound, which was illustrated by the synthesis of the fluoroalkene sphingosine analog. The replacement of tin with fluorine was carried out with F-TEDA-PF₆ **615** in the presence of silver phosphinate AgOP(O)Ph₂ **616**.⁶³

In 1990 Zhang et al. reported the palladium- and molybdenum-catalysed addition of Sn-H to C≡C bonds leading to vinylstannanes in high regio- and stereoselectivity. Although the authors described in detail the hydrostannation of mono alkynes, a few examples of diyne reactivity was also reported. The readily available and air-stable catalyst PdCl₂(PPh₃)₂ 94 was applied in the hydrostannation of symmetrical and unsymmetrical 1,3-diynes. The addition of HSn(n-Bu)₃ 508 to symmetrically substituted dodeca-3,5-diyne 13a under mild reaction conditions and a short reaction time (10 min) gave (E)-enyne 618a in 78% yield. The n-Bu₃Sn moiety was attached to the carbon atom contiguous to the $C \equiv C$ unit. The further addition of Sn-H bond to the unreacted triple $C \equiv C$ bond was not possible and led to the decomposition of 618a. Similar regioand stereoselectivity was observed when unsymmetrically substituted diyne 617 terminated with ethynyl group was used. In turn, the hydrostannation of deca-1,3-diyn-1-yltrimethylsilane



Scheme 110 Hydrostannation of o-diethynylbenzene 545 with diorganotin hydrides 532, 538, 540, and 546.





i) heptan, reflux, AIBN **67**, ii) *n*-BuLi **85**, -40 °C, 1.5 h, iii) BF₃•Et₂O **565**, iv) a)TsCl, pyridine, 0 °C–r.t., 5 h, b) Nal, acetone, reflux, 4 h, c) PPh₃, MeCN, 14 h, d) LDA, -78 °C, 1 h, v) HMPA, -78 °C, 40 min. vi) l₂**418**, DCM, pyridine, -45 °C, 1.5 h, vii) DMF, 105 °C, 6 h, viii) LiOH/DME = 1/1, r.t., 4 h.

Scheme 112 Synthetic path to 13-thiaarachidonic acid 573 via hydrostannation of penta-1-4-diyne 526a.



Scheme 113 The stannylative coupling of 1,6-diynes 127a, 127j, 127m, 127p, 394d, and 574a-c with tributyltin hydride 508 catalysed by Pd(OH)₂/C 575.



Scheme 114 Proposed mechanism of stannylative coupling of 1,6-diynes with tributyltin hydride **508**. Both pathways A and B are possible.

180a gave monohydrostannation product **618b** in 86% yield. The presence of trimethylsilyl moiety caused the addition of Sn-H to C \equiv C bond adjacent to the alkyl substituents. Intriguingly, the hydrostannation of 1,2-bis(trimethylsilyl)ethyne did not occur at all, thus the SiMe₃ moiety in **180a** could be considered as a directing group (Scheme 120).²⁷⁷

The same palladium catalyst $PdCl_2(PPh_3)_2$ **94** was applied for the tin-functionalised dienynes by the hydrostannation of (*Z*)- or (*E*)-endiynes with $HSn(n-Bu_3)$ **508** in just 20 minutes at room temperature. The protocol was suitable for the symmetrical and unsymmetrical (*Z*)-endiynes **619a–j** with various (aryl, alkyl, alkoxy, silyl) substituents. Among many possible isomers only α -products with the tin atom located on the carbon atom adjacent to C=C bond, were formed. Nevertheless, very high selectivity was noticed only for symmetrical (*Z*)-endiynes. In the case of unsymmetrical (*Z*)-trideca-5-en-3,7-diyn-1-ol **619e** an equimolar mixture of α -isomers and α' -isomers was observed since, the HSn(*n*-Bu₃) **508** did not distinguish in its addition between the two triple bonds. Intriguingly, the hydrostannation of SiMe₃ substituted unsymmetrical diynes with HSn(*n*-Bu)₃ **508** gave exclusively α -isomers, thus the silyl moiety acted as a directing group. The addition of Sn–H bond occurred on the silylunsubstituted C \equiv C bond with tin moiety attached at C_{α} (Scheme 121).²⁷⁸

Notably, the geometry of endiynes double bond had a crucial influence on reaction regioselectivity. The hydrostannation of unsymmetrical (*E*)-endiynes **621a–d** in the same reaction conditions gave a mixture of α - and β -isomers even in the presence of the directing SiMe₃ group. However, the silyl-substituted C = C bond, similar to hydrostannation of (*Z*)-endiynes, remained unreactive. The ratio of α - and β -isomers was dependent on the second substituent and ranging from 64:36 to 94:6 (Scheme 122).²⁷⁸

(*Z*,*E*)-Stannylated dienynes **620** were also found to be attractive building blocks in organic synthesis. Bujard *et al.* reported the synthesis of (*Z*,*E*)-dienediynes through the iododestannylation of **620d** and **620i** with NIS **624** and subsequent Pd/Cu catalysed coupling of vinyl iodide **625** with terminal alkyne **626a–c.** The process was highly stereoselective and gave desired products **627a–c** in good isolated yields (47–51%) (Scheme 123). The authors suggested that the obtained acyclic dienediynes are promising substrates for the synthesis of more complex molecules such as neocarzinostatin chromophore which was found to be an antitumor antibiotic.^{279,280}

Kazmaier *et al.* described the Mo(CO)₃(NC-*t*-Bu)₃ **629** catalysed hydrostannation of C \equiv C bonds. Although in the report a detailed research on hydrostannation of alkynes was presented, a single example of hydrostannation of a diyne was presented. The hydrostannation of diynoic ester **628**, possessing internal and terminal triple C \equiv C bonds, with HSn(*n*-Bu)₃ **508** occurred preferentially at the internal C \equiv C bond bearing electron-withdrawing group. The reaction was relatively selective leading to a mixture of α - and β -isomers (**630/631** = 82/18) in 74% isolation yield (Scheme 124).²⁸¹

8. Hydroamination

Compounds (acyclic and heterocyclic) possessing carbon-nitrogen bonds are omnipresent in an array of chemicals, especially



Scheme 115 Hydrostannation of terminally substituted diynes 584 with tributyltin hydride 508 catalysed by Pd(OH)₂/C 575.







Scheme 117 Temperature tunable *trans*-hydrostannation of 1,3-diynes 594 with tributyltin hydride 508 catalysed by [Cp*RuCl]₄ 596.

in natural compounds, agrochemicals, pharmaceuticals, or cosmetics.^{22,282–288} They are produced on a gram scale as *fine*

chemicals, as well as in feedstock in tonnage scale in the industry. A limited number of chemical transformations leading







Scheme 119 Synthesis of Typhonoside F 608 and Thphonoside E 614 with trans-stannylation step of diyne 602.

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to the formation of the C–N bonds in stoichiometric reactions led to the intensive development of hydroamination reactions, which simply introduce the N-atom to the compound structure, and occurs by the addition of the N–H bond to the unsaturated C–C bonds in olefins and alkynes.^{19–22,29,289–293} This 100% atom economic method mostly requires the application of a catalyst to (i) overcome the repulsion electrostatic effect between the high electron-dense unsaturated C=C bond and the strong Lewis base (electron-rich amine 1° or 2°, ammonia, or hydrazine), and (ii) to facilitate this addition reaction due to the high energy difference between both types of bonds.²⁹⁴ The hydroamination of (non)conjugated diynes leads to various products, but intramolecular cyclisation is of utmost importance to produce N-heterocyclic compounds, *e.g.*, indoles, pyrroles, pyrazoles, pyrimidines.

8.1. Noncatalytic hydroamination of conjugated 1,3-diynes

The origins of noncatalytic hydroamination of 1,3-diynes date back to the 1960s and 1970s, which was briefly described in the review published in 2002, which focused on the heterocyclisation of diynes.²⁹⁵ Different hydroamination agents (*e.g.*, ammonia **632**, hydrazine **633**, substituted hydrazines **634**, amines **635**, diamines **636–637**, hydroxylamine **638**, 2-aminoethan-1-ol **639**, guanidine **640**) were used in this transformation. Depending on the type of



Scheme 122 Synthesis of dienynes 622 and 623 via hydrostannation of (E)-dienynes 621a-d with $HSn(n-Bu)_3$ 508 catalysed by $PdCl_2(PPh_3)_2$ 94 (5 mol%).



Scheme 123 Synthesis of (*Z*,*E*)-dienediynes *via* iododestannylation of **620d**, **620i**, and vinyl iodide **625** coupling with terminal alkynes **626a**–c catalysed by Cul/Pd(PPh₃)₄ **518/35**.



reagents and reaction conditions various heterocyclic products (*e.g.*, pyrazoles **645–647**, pyridines **649**, diazepines **651**, pyrymidines **650**, isoxazole **648**) were obtained (Scheme 125).^{295–298}

The Cope-type hydroamination of conjugated 1,3-diynes occurs under noncatalytic and relatively mild conditions, while

the reactivity of the substrates depends on the electronic structure of the 1,3-diyne. A reduction of the electronic density on the $C \equiv C$ bond has a positive influence on the reaction yield and formation of the hydroaminated product. Therefore, the electron-withdrawing groups attached to the benzene ring in



655a–g Scheme 126 Synthesis of 3,5-disubstituted isoxazoles (656, 657) and pyrazoles (658, 659) in the Cope-type hydroamination reactions of 1,3-diynes.

1,4-diphenyl-buta-1,3-diyne permitted the desired products to be isolated in higher yields, while electron-donating groups caused the opposite effect. Bao et al. have reported the synthesis of 3,5-disubstituted isoxazoles or pyrroles (Scheme 126) by the Cope-type intramolecular hydroamination of 1,3-diynes 1a, 1c-d, 27a, 60a, 60e, 208b, 230d, 500a, 500d, 655a-g with hydroxylamine 639 or hydrazine 633 respectively.^{299,300} Both reactions occurred at elevated temperatures (110 °C or 60 °C) in DMSO and using an excess of hydroaminating reagent 633 or 639 (1.5-4.0 equiv.) to provide the full conversion of 1,3-diynes. Triethylamine (Et₃N) was used as the most effective base in the synthesis of isoxazoles 656-657. The reactions yielded isoxazoles 656 and 657 in 61-92% or pyrazoles 658 and 659 in 60-93% isolated yields (Table 16). The high selectivity for unsymmetrical divnes was obtained when the reagent was substituted with groups with a distinct difference in electronic properties (e.g., hexyl- and 4-nitrophenyl).^{299,300} The mechanism of intermolecular Cope-type hydroamination of 1,3-diynes occurred via the formation of intermediate 661 in a protontransfer process, which further undergoes isomerisation to the allenyl oxime intermediate 662, followed by the electrophilic

cycloaddition towards 3,5-disubstituted isoxazoles **656** or pyrazoles **658** (Scheme 127).^{299,300} Moreover, the same group developed a one-pot procedure for the synthesis of heterocycles *via* Glasser coupling of alkynes followed by intramolecular hydroamination. The final products were obtained in comparable yields.^{299,300}

The hydroamination with hydroxylamine **638** or hydrazine **633** was carried out also for symmetrical **663a–f** (Scheme 128) and nonsymmetrical **1**,3-diyne indole derivatives **668** (Scheme 129). The reactions were conducted in eco-friendly PEG-400 as a solvent, which facilitates a proton transfer to the allenyl intermediate, which according to the DFT calculations is the rate-determining step of the process. Applying PEG-400 as a solvent, it was possible to shorten the reaction time from 20 h to 2–6 h, and to carry out the reactions under milder conditions.^{301,302} Additionally, *N*-substituted products were obtained by the application of arylhydrazines **665a–d** (Schemes 128 and 129).³⁰²

3,5-Disubstituted pyrazoles **674a–d** were synthesised using the Cope-type hydroamination in a sustainable manner by the application of a continuous flow process, starting from terminal alkynes **518a** and **672a–c** and hydrazine **633**. Two coil Table 16 Synthesis of 3,5-disubsituted isoxazoles (656, 657) pyrazoles (658, 659) in Cope-type hydroamination reactions with hydrazine 633 and hydroxylamine 638

	Yield [%]		Diyne		Yield [%]	Yield [%]	
Entry	656	657	R^1	R^2		658	659
	656 = 657					658 = 659	
1	656a, 86		Ph	Ph	1a	658a, 83	
2	656b, 81		<i>p</i> -MeOC ₆ H ₄	<i>p</i> -MeOC ₆ H ₄	1d	658b, 76	
3	656c, 84		$p-MeC_6H_4$	$p-MeC_6H_4$	1c	658c, 78	
4	656d, 81		<i>m</i> -MeC ₆ H ₄	<i>m</i> -MeC ₆ H ₄	27a	658d, 78	
5	656e, 91		p-BrC ₆ H ₄	p-BrC ₆ H ₄	230d	658e, 93	
6	656f, 98		$p-NO_2C_6H_4$	$p-NO_2C_6H_4$	500a	658f, 93	
7	656g, 66		<i>n</i> -Hexyl	<i>n</i> -Hexyl	208b	658g, 60	
8	656h, 89		<i>c</i> -Hexyl	<i>c</i> -Hexyl	60a	658h, 76	
	656 ≠ 657					658 ≠ 659	
9	656i, 41	657i, 32	Ph	p-MeOC ₆ H ₄	60e	658i, 47	659i, 35
10	656j, 72	657j, 9	p-FC ₆ H ₄	p-MeOC ₆ H ₄	655a	658j, 52	659j, 31
11	656k, 87	657k, 0	$p-NO_2C_6H_4$	p-MeOC ₆ H ₄	655b	658k, 58	659k, 30
12	656l, 78	657l, 0	Ph	<i>n</i> -Hexyl	655c	658l, 72	659l, 11
13	656m, 81	657m, 0	Ph	c-Hexyl	655d	658m, 70	659m, 18
14	656n, 89	657n, 0	$p-NO_2C_6H_4$	n-Hexyl	655e	658n, 88	659n, 0
15	6560, 94	6570, 0	$p-NO_2C_6H_4$	c-Hexyl	655f	6580, 70	6590, 0
16	656p, 64	657p, 12	$p-MeC_6H_4$	n-Hexyl	500d		Hexyl
17	_ • ′		p-NO ₂ C ₆ H ₄	Ph	655g	658q, 54	659q, 27



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reactors were combined, the temperature, the volume of the coils, and reagent flow rates, which influence the residence times, which were carefully chosen to obtain high product yields of 1a, 1c, 27c, and 258a in the Glasser coupling of alkynes and hydroamination process. For the homocoupling of alkynes 518a, 672a-c a 3.5 mL coil, alkyne concentration 0.75 M in DMSO, 120 $^{\circ}$ C, and 0.1 mL min⁻¹ flow (residence time: 35 minutes) were used. After the reactor outlet, the thiourea 673 scavenger column was applied to trap the copper (CuBr₂ 52) used as a catalyst for the Glaser reaction. The hydroamination was carried out in a 17.5 mL coil with hydrazine 633 in DMSO $(0.1-0.2 \text{ mL min}^{-1})$ at 140 °C. The 87.5 min residence time was sufficient for the total conversion of 1,3-diynes 1a, 1c, 27c, and 258a yielding the appropriate pyrazoles 674a-d in 90-98% (isolated yields: 84-90%). The system was active for 16 hours for subsequent continuous flow Glaser coupling/hydroamination of 3-ethynylthiophene 27c with hydrazine 633, leading to 0.52 g of pure 3-(thiophen-3yl)-5-(thiophen-3-ylmethyl)-1H-pyrazole 674d in 81% isolated

yield. The ICP analysis detected only residue amounts of Cu (3 ppm), showing a high efficiency of the in-line Cu-scavenger (Scheme 130).³⁰³

This noncatalytic hydroamination was also used in the synthesis of 2,4,6-pyrimidines **677a–o**, which possess biological activities (*e.g.*, antitumor, antifungal, anticancer, anticonvulsant), luminescence properties, or are the component of nucleic acids. They can be effectively synthesised from diaryl or monoaryl-substituted 1,3-diynes (**1a–d**, **27b**, **258a**, **258e**, **258h**, and **675**), and amidines **676** (acetamide hydrochloride **676a**, benzamidine hydrochloride **676b** or formamidine acetate **676c**), which are used as bidentate nucleophiles in the presence of Et₃N as a base. The reaction occurred effectively in DMSO under a high temperature (160 °C), with the reagent ratio [diyne]: [**676**]: [Et₃N] = 1:3:3. The products **677a–o** were obtained with 46–88% isolated yields, with the highest efficiency for electron-poor 1,3-diynes with electron-withdrawing groups (Scheme 131).³⁰⁴







Scheme 129 Synthesis of 3,5-disubstituted-1,2-isoxazoles 669 and 3,5-disubstituted 1*H*-pyrazoles 670 and 671 from nonsymmetrical diynes 668 in the hydroamination process.

8.2. Catalytic hydroamination of conjugated 1,3-diynes and separated diynes

The hydroamination of conjugated, as well as nonconjugated diynes, is often catalysed by homogeneous transition metal

catalysts (Au, Ag, Cu, Pd) as well as non-noble metal or main group element complexes (Ti, Ni, Co, Ca). Among them, Au complexes have found a prominent position in their application for catalytic hydroamination. The hydroamination reaction







Scheme 131 Synthesis of substituted pyrimidines 677a-o by the hydroamination reaction of diynes 1a-d, 27b, 258a, 258e, 258h, and 675 with aminides 676.

constitutes one of the steps in the synthesis of natural products as *e.g.*, indolizidine alkaloid (±)-Monomorine, pharmaceuticals or agrochemicals.^{64,305–307}

Skrydstrup *et al.* reported the synthesis of electron-rich 2,5-diamidopyrroles, 1,2,5-trisubstituted pyrroles, as well as pyrazoles using a hydroamination reaction in the presence of Au(1)-complexes. These products are difficult to synthesise according to other methods. Using (Ph₃P)AuNTf₂ **680** and only a slight excess of aniline **679a** (1.05 equiv.), appropriate 2,5-diamidopyrroles were obtained in 30 min, under low temperature

(30 °C) in CH₂Cl₂. The anilines with electron-withdrawing groups in the *para* position required a longer reaction time (60 min). The trisubstituted products were obtained after 24 h using different Au **680** and **682** catalysts in toluene and at elevated temperatures (80 °C), (Scheme 132 and Table 17).³⁰⁸

The cationic gold(1) catalyst supported by a cyclic(alkyl)-(amino)carbene (CAAC) generated *in situ* from an equimolar mixture of AuCl(CAAC) **684**/KB(C_6F_5)₄ was able to activate NH₃ **632** and NH₂NH₂ **633** in the hydroamination reactions of alkynes, and conjugated and non-conjugated diynes **1a**, **65a**,



Table 17 Synthesis of pyrroles **681** in Au-catalysed hydroamination reactions of diynes **1a**, **1d**, **208b**, and **678a–c** with anilines **679a–g** and phenylhydrazine **665a** (Scheme 132)

Entry	Diyne	Amine	Reaction conditions	Product	Yield [%]
1	1a	679a	(ii)	681a(b) ^a	49 $(56)^a$
2	1d	679a	(ii)	681c(d) ^{<i>a</i>}	48(63) ^a
3	678a	679a	(i)	681e	96
4	678a	679c	(i)	681f	95
5	678a	679d	(i)	681g	95
6	678a	679e	(i)	681ĥ	94
7	678b	679b	(i)	681i	95
8	678c	679f	(i)	681j	93
9	678c	679g	(i)	681k	94
10	208b	679a	(ii)	681l	24

^a Phenylhydrazine 665a was used as a reagent.

541, 690 (Scheme 133). These hydroaminating reagents are powerful reductive agents, which can form saturated products, as well as metal nanoparticles, therefore their use in the formation of the new C–N bonds is problematic. The gold centre is capable of NH₃ **632** or NH₂NH₂ **633** addition if it is coordinated by the CAAC ligand and rendered cationic by Cl⁻ abstraction. The same Ag complex: (CAAC)AgCl **692**/KB(C₆F₅)₄ **685** did not cause the activation of NH₃ **632**. The coordination of NH₃ **632** or NH₂NH₂ **633** led to a typical Werner complex immediately. The same happened when the alkyne was added to the initial catalyst, η-2 bounded to the gold atom. The reaction occurred according to the insertion mechanism. The addition of NH₃ **632** to 1,4-diphenyl-buta-1,3-diyne **1a** or hexa-1,5-diyne **541**, occurred according to the Markovnikow rule, followed by the ring-closing hydroamination to give pyroles



Scheme 133 Au-catalysed (683–686) hydroamination of diynes with NH₃ 632 and NH₂NH₂ 633. Synthesis of pyrroles and pyrazoles.
with high yields: 87% for **688** and 96% for **689a**. The same reaction with 3,3-dimethyl-1,5-diphenylpenta-1,4-diyne **690** formed two products: Markovnikov six-membered ring and anti-Markovnikov five-membered heterocycles **691** and **691**' in a 2:3 ratio. Similar activity was observed for the reaction with hydrazine **633** where pyrroles or pyrazoles were formed (Scheme 133).^{309,310}

Amphiphilic gold nanoparticles: Au-HS/SO₃H-PMO(Et) **693**, obtained with a narrow particle distribution 1–2 nm (which is important for their high catalytic activity) permitted the reactions to be carried out with organic reagents in an aqueous solution without using any organic solvents. The intramolecular hydroamination of hexa-2,4-diyne **65a** in water occurred with high yields of the product **676** (87%). Moreover, the addition of catalytic amount of H_2SO_4 to AuCl(PPh₃) **694** (used as a homogeneous catalyst) was also successful, but the yields were much lower than for the reaction catalysed by nanoparticles **693**.³¹¹

Nolan *et al.* described the application of 5 mol% of [Au(IPr)OH] **695** (IPr = 1,3-bis-(2,6-di-*iso*-propylphenyl)imidazol-2ylidene **696**) as a precatalyst for the hydroamination and hydration of conjugated 1,3-diynes **1a** and **2580** to pyrroles **697a–d** and furans respectively. The active cationic form of the catalyst is formed in the presence of 7.5 mol% of HNTf₂. Microwave irradiation was used as a heating source, and the reaction was carried out at 120 °C for 90 min (Scheme 134).³¹² The results were similar to those obtained by Skrydstrup (see Scheme 132).³⁰⁸

Ohno *et al.* developed a method for the formation of various fused indoles and indolines using gold catalysts. Depending on the catalyst and ligands type different products were selectively formed with very good yields.^{313–315} The first paper focused on the synthesis of aryl-annulated[α]carbazoles **700** *via* gold-catalysed 5-*endo-dig* hydroamination of diynes followed by 6-*endo-dig* hydroarylation. The type of phosphine ligand attached to the gold atom **699a–c** has a strong impact on the diyne **698a–s** conversion. Particularly, when bulky biarylphosphine ligands were used, the dissociation of the catalyst from a substrate is accelerated, improving the possibility for activation of the appropriate C \equiv C bond for hydroamination, even in the case of reagents with electron-rich aryl groups (*p*-MeC₆H₄, **698a**). The reaction was sluggish when *o*-CNC₆H₄

698e was used probably due to the interaction of CN group with the catalyst. The process was carried out in the presence of R₃PAuCl **699**/AgOTf systems yielding aryl-annulated[α]-carbazoles **700** with very good yields (Scheme 135).³¹³ Carbazoles **700c** and **700n** showed good antifungal activity against *T. metagrophytes* and modest activity against *T. rubrum*.³¹³

Applying this method it was also possible to synthesize dihydrobenzoindole **702** and **703** and azepino-**705a** oxepino-[3,4-*b*]indole **705b** and cyclohepta[*b*]indole **705c** derivatives with moderate to good yields (Scheme 136). The authors proposed the mechanism of this transformation, which started from the activation of diyne **698t** by gold catalysts **699** to **706**. Next the 5-*endo-dig* cyclisation furnishes the indolylgold intermediate **707**. After proto-deuaration the cyclised product **708** is formed. It is activated by the gold catalyst, which promotes 6-*endo-dig* cyclisation at the C-3 position of the indole, followed by the rearomatisation to arylgold species **709**. The cycle is finished with the proto-deuaration of **709** and production of fused carbazole **700t**, with the subsequent regeneration of the initial catalyst **699** (Scheme 137).

The same group developed a method for the synthesis of fused indolines **716** and indoles **712** from anilines functionalised with diyne group **710** catalysed by gold complexes. The formation of both products is controlled by the reagent, ligand, and solvent. When IPr **696** ligands and protic solvents were used the fused indoles **712** were predominantly formed. While Buchwald's type ligands (*e.g.*, JohnPhos **714** and BrettPhos **715**) and nonpolar solvents (*e.g.*, toluene) promoted the synthesis of indolines **716** as the main products. The most active catalyst for the preparation of indoles was IPrAuNTf₂ (5 mol%) **711**, while for the synthesis of indolines John-PhosAuNTf₂ **713** was applied (Scheme 138 and Table 18).

The catalytic systems **711** and **713** were tolerant to many electron-donating and electron-withdrawing functional groups in the diyne structures. For **710g** (with highly electron-withdrawing CN group R^1) the indole **712g** was formed with very low yield (17%), while propellane type indoline **716g** was not formed regardless of the method A or B. Moreover the influence of the position of substituents in phenyl ring of aniline influences the products yields. When the ring was



Scheme 134 Synthesis of pyrroles 697a-d in the hydroamination of 1,3-diynes in the presence of [Au(IPr)OH] 695



Reaction conditions: 5 mol% of 699c, AgOTf (5 mol%), 3–24h, ^a For 694, ^b For 699a, ^c For 699b, ^d 20 mol% of 699c and AgOTf.

Scheme 135 Synthesis of aryl-annulated[α]carbazoles via gold-catalysed 5-endo-dig hydroamination of diynes followed by 6-endo-dig hydroarylation.



substituted in the *ortho* position to alkyne, **710h** propellane type indoline **716h** was formed in moderate yield (44%), while oxocine-fused indole was not detected at all. The steric repulsion

between *o*-Br and phenyl groups interferes with the formation of indole **712h**. When the length of the chain between both $C \equiv C$ bonds was shorter as in **710u**, the propellane type indoline **716u** is



Scheme 137 Proposed mechanism of the synthesis of fused carbazole **700t** in the hydroamination/cyclisation reactions catalysed by gold complexes **699**.



Scheme 138 Synthesis of fused indolines 716 and indoles 712 catalysed by gold complexes 711 and 713.

Table 18	Reagents	scope in t	the gold	catalysed	synthesis	of fused	indolines	716a–w an	id indoles	712a-v	v
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Entry	710	\mathbb{R}^1	R^2	R ³	n	Z	Method	Yield 716 [%]	Yield 712 [%]
1	710a	Н	Ph	Ме	1	0	А	716a, 11	712a , 82
2							В	716a, 88	712a, 8
3	710b	4-F	Ph	Me	1	0	Α	716b, 3	712b, 68
4							В	716b, 67	712b, 22
5	710c	4-Cl	Ph	Me	1	0	Α	716c, trace	712c, 64
6							В	716c, 79	712c, 13
7	710d	4-Br	Ph	Me	1	0	Α	716d, 6	712d, 62
8							В	716d, 82	712d , 11
9	710e	4-Me	Ph	Me	1	0	Α	716e, 3	712e, 77
10							В	716e, 77	712e, 6
11	710f	4-MeO	Ph	Me	1	0	А	716f, 3	712f , 77
12							В	716f, 77	712f, 6
13	710g	4-CN	Ph	Me	1	0	A ^a	716g. 0	712g. 17
14	0						В	716g. 0	712f. 5
15	710h	3-Br	Ph	Me	1	0	А	716h . 0	712h . 0
16							В	716h , 44	712h . 0
17	710i	2-Br	Ph	Me	1	0	Ā	716i , trace	712i, 71
18	, 101	2 01		1110	-	0	B	716i, 80	712i, 9
19	710i	н	4-ClC _c H ₄	Me	1	0	A	716i < 10	712i 73
20	/10 j		1 0106114	1010	1	U	B	716j, <10	712j, 70
20	710k	н	4-BrC.H.	Me	1	0	Δ	716k < 15	712J, 0
21	/108	11	4 0106114	ivic	1	0	B	716k, < 10	712k, 50
22	710]	ч	4-CNC-H	Me	1	0	Δ	7161 < 11	712k, 4 712l 73
23	/101	11	4 0106114	wie	1	0	B	716 , 7 1	7121, 73
25	710m	н	4-MeC-H	Me	1	0	Δ^a	7101, 74 716m < 12	712n, 0
25	/1011	11	4 101006114	me	1	0	B	716m, <12	712m , 30
20	710n	ч	4-MeOC H	Me	1	0	Λ^a	716n < 14	712m, 10
20	71011	11	4 1010006114	wie	1	0	P	7101, < 14 716n 67	712n, 22
20	7100	ч	4 CIC H	Мо	1	0	B A	7160 < 7	7120, 71
29	/100	п	$4 - C_1 C_6 \Pi_4$	WIC	1	0	A D	7100, < 7	7120, 71
30	710m		a cle u	Ма	1	0	D A	7100, 74	7120, 8
31	710b	п	$2-CIC_6H_4$	Me	1	0	A	710p, < 12	712p, 70
32	710a		1 nonhthul	Ма	1	0	В	716p, 26	712p, 27
33	710q	п	1-naphtnyi	Me	1	0	A	716q, < 9	712q, 63
34	710-		Мо	Ма	1	0		710 4 , 52	712q, 0
30	/100	п	Me	Me	1	0	A	710F, 8	712F, 10
36							В	716r, 34	712 r , 5
3/	710-		pl		4	0	C A	716r, 82	712 r , trace
38	/105	н	Ph	н	1	0	A	/16S, /	7128, 32
39	-101					<u> </u>	В	7 16s , 18	712s, 0
40	710t	Н	Ph	Bn	1	0	A	7 16t , trace	712t, 56
41			-1		_		В	7 16t , //	7 12t , 8
42	710u	Н	Ph	Ме	0	0	A	7 16u , 0	7 12u , 63
43			-1				В	7 16u , 0	7 12u , 44
44	710v	Н	Ph	Me	2	0	A	716v, 0	712v, 0
45			-1				В	716v, 28	712v, 0
46	710w	Н	Ph	Me	1	NTS	Α	716w, 0	712w, 63
47							В	716w, 0	712w, 67

Reaction conditions: A: 5 mol% IPrAuNTf₂ 711;EtOH or *i*-PrOH, 40–60 °C, 5 min–6.5 h; B: 5 mol% JohnPhosAuNTf₂ 713, *i*-PrOH, 80 °C, 10 min–6.5 h. ^a Additional 5 mol% of 711 was added. ^b The reaction was carried out in *i*-PrOH with the addition of MS3 Å at 80 °C.

not obtained due to the higher ring strain, while for longer chains as in **710v** no nine-membered ring fused indole **712v** was produced, while **716v** was obtained in 28% (Table 18).

The mechanism of this transformation, in which hydroamination is a crucial step was proposed according to the experiments. Activation of alkyne with gold **717** is responsible for 5-*endo-dig* cyclisation followed by the protodeauratiom towards the indole. Next, the activation of the second alkyne group promotes the 8-*endo-dig* hydroarylation of **719** to intermediate **720**. The subsequent protodeuaration of **720** furnishes oxocine fused indole **712**. The intermediate **720** can be easily opened to cationic intermediate **721**. Elimination of the gold from **721** leads to allene **722**, which is essential for obtaining propellane-type indoline **716** (Scheme 139).³¹⁵ Protic solvents accelerate the protodeauration of vinyl-gold intermediate **720** yielding oxocine-fused indoles **712**. The same influence is observed for electron-donating IPr ligand **696**. When allene **722** is formed the mechanism is favoured to obtained propellane-type indolines **716**. The DFT calculations for this transformation was also used to help underpin the reaction mechanism.³¹⁶ A detailed discussion on the influence of substituents attached to the nitrogen atom in aniline, in the diyne, and the aryl ring on process selectivity and product yields and mechanism of the process were comprehensively discussed by the authors in several papers.^{313–315}

Wiest, Helquist *et al.* applied a hydroamination reaction for the desymmetrisation of diynes 727a-c, 730a-c, 732 in the

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presence of Ag(phen)OTf **728** yielding to 1-pyrrolines **729a-c**, **731a-c**, or **733** with two entirely different, orthogonal functional groups, which are capable of further functionalisation. The reaction occurred under mild reaction conditions (25–50 °C), with the low catalyst **728** loadings (0.5–2.0 mol%) (Scheme 140).⁶⁴ Additionally, this method was applied to the synthesis of natural indolizidine alkaloid (±)-monomorine **743**, which started from the hydroamination/cyclisation of diyne **734** synthesised from 4-bromo-1-butyne followed by the several steps illustrated in Scheme 141.⁶⁴

In the hydroaminative cyclisation of diynes was used also AgSbF₆ **746** as a catalyst. The process was developed for the synthesis of naphthol-indole derivatives **750a–l** from 1,3-diynes **745a–l** and sulfoxonium ylides **744** in a one-pot cascade reaction (i) intramolecular hydroamination/cyclisation of diynesubstituted anilines **745a–l** to **750a–l**, and (ii) [RhCp*Cl₂]₂ **748** catalysed arene *ortho*-C–H bond activation. Indoles functionalised in the C2 position **750** were obtained with good yields with high functional groups tolerance (Scheme 142).³¹⁷

The system generated *in situ* from TiCl₄ **448** by the addition of *t*-BuNH₂ **751** in toluene is an active catalyst for the hydroamination of alkynes and 1,3-diynes with hydrazine **633** leading to indole or pyrrole derivatives respectively. Using 20 mol% of TiCl₄ **448**, at 105 °C for 18 h, the reagents (anilines **679a–b** and dodeca-5,7-diyne **13a**) are quantitatively converted to the products mixture of mono- and bishydroamination of diyne **13a**. The pyrroles **752** were obtained as the main products in 30% yield (Scheme 143).³¹⁸

CpCo(C_2H_4)₂ 755 was applied in the hydroaminative coupling of α, ω -diynes 79b, 160, 753a–j with various amides 754a–f, which resulted in the formation of dienamides 757–758 with high regio- and stereoselectivity (Scheme 144 and Table 19). Such compounds can be used as reagents in Diels–Alder reaction, in the synthesis of polycyclic compounds as well as natural product derivatives. They can be also synthesised from alkynes by a co-oligomerisation reaction with *N*-vinyl amides or Ticatalysed coupling with ynamides.^{319,320} The mechanism of Co-mediated reaction started from the oxidative addition of









diyne **79b**, **160**, or **753a–j** to the metal centre of **755** with the formation of cobalt–cyclopentadiene **760**, followed by the formation of 18 electron *N*-coordinated complex **761**. Proton transfer from nitrogen to carbon then takes place to generate intermediate **762**, which subsequently rearranges to *N*-coordinated cobaltcyclopentene **763**, that tautomerises to product **764**. The regioselectivity is controlled by the proton transfer step to the least hindered carbon atom in cobaltcyclopentadiene (Scheme 145). The reaction occurring according to this mechanism permitted several amidated 1,2-dimethylenecycloalkanes to be obtained in moderate to good yields (24–81%) (Scheme 144 and Table 19).³²¹

Shimada and Yamamoto have developed a different approach applying hydroamination reaction in the C–C bond cleavage in diynes **1a**, **13a**, **617**, and **765a–f** with *o*-aminophenols **766a–h**. The transformation leads to 2-substituted

benzoxazoles **769** and **771** and ketones **770** and **772**. The reaction occurred according to two possible pathways in the presence of $\operatorname{Ru}_3(\operatorname{CO})_{12}$ **489** with $\operatorname{NH}_4\operatorname{PF}_6$ **767** by the $C \equiv C$ (path A) or C–C single (path B) bond cleavage. The formation of more sterically hindered benzoxazoles **769** is favourable. Additionally, the bulky groups in the diyne (*e.g.*, *t*-butyl **765c**) led to the almost exclusive formation of product (**769**, **769**:**771** = 30:1) with 81% yield. For substituted *o*-amimophenols **766d–f** in positions 4 and 5 with strong electron-donating or electron-withdrawing groups, the reactions were sluggish. Ru-catalyst **489** was found to be more effective in the reaction with terminal diynes **765a–f**, while internal diynes **1a** and **13a** proceeded better with $\operatorname{Pd}(\operatorname{NO}_3)_2$ **768** (Scheme 146). The key step in the bond cleavage is the hydroamination of one of the C \equiv C bonds of **765a–f** with **766**, followed by the



Scheme 142 Synthesis of naphtol-indole derivatives 750a-l from 1,3-diynes 745a-l and sulfoxonium ylides 744 catalysed by [RhCp*Cl₂]₂ 748/AgSbF₆ 746 system.





tautomerisation leading to α , β -unsaturated imines 774. The addition of the second molecule of *o*-aminophenols 766 to 765 yielded β -aminoimines 775 and their tautomers 777, which further undergoes intramolecular cyclisation to ketals 776 and 778. The final step leading to benzoxazoles 769 and 771

occurred by the C–C bond cleavage through a retro-Mannich-type reaction (Scheme 147).³²²

The copper-catalysed synthesis of pyrroles *via* hydroamination of diynes was first published in 1965.³²³ 0.1 mol% of CuCl 55 was used for the hydroamination/cyclisation of 1,3-diynes

Table 19 Hydroaminative coupling of substituted α, ω -diynes with dienamides **754**

Entry	Diyne	754	Yield 757:758:759 [%]
			757 = 758:759
1	$R^1 = R^2 = H, n = 2, 160$	754a	24:0
2		754b	55:0
3		754c	65:0
4		754d	32:0
5		754e	81:0
6		754f	69:0
7	$R^1 = R^2 = SiMe_3, n = 1, 753a$	754e	64:0
8	$R^1 = R^2 = Ph, n = 1, 753b$	754e	48:0
9	$R^1 = R^2 = CMe_2OH, n = 2, 753c$	754e	58:0
10	$R^1 = R^2 = CO_2Me, n = 2, 753d$	754e	0:70
11	$R^1 = R^2 = Bpin, n = 2, 79b$	754e	0:60
	1 / /		757:758:759
12	$R^1 = H, R^2 = SiMe_3, n = 1, 753e$	754e	72:0:0
13	$R^1 = H, R^2 = SiMe_3, n = 2, 753f$	754e	66:0:0
	, ,	754c	14:65:0
14	$R^1 = H, R^2 = Ph, n = 2, 753g$	754e	70:0:0
15	$R^1 = H, R^2 = Ph, n = 2, 753g$	754c	78:0:0
16	$R^1 = Ph, R^2 = SiMe_3, n = 2.753h$	754e	62:18:0
17	$R^1 = H, R^2 = CMe_2OH, n = 2, 753i$	754e	70:0:0
18	$R^1 = H, R^2 = Bpin, n = 2, 753i$	754e	22:13:0

 $\label{eq:rescaled} \begin{array}{l} \mbox{Reaction conditions: (1) } CpCo(C_2H_4)_2 \ 755 \ (1 \ equiv.), 754 \ (5 \ equiv.), THF, \\ r.t., \ 24 \ h, \ then: \ (2) \ Fe(NO_3)_2 \cdot 9H_2O \ 756 \ (1 \ equiv.), 0 \ ^{\circ}C, \ 5 \ min. \end{array}$

1a, **1d**, **65a**, **208b**, **258n**, **781a–c** with aromatic and aliphatic primary amines **679a**, **679c**, **679g**, **782a–i** and ammonia **632**. The reaction was carried out in MeOH, EtOH, 1,4-dioxane, or DMF for 1 h, at 150–180 °C, furnishing pyrroles **783a–w** in moderate yields. Increasing the catalyst **55** concentration to 10 mol%, under solvent-free conditions, and with 10 equiv. of amine **679** and **782** it was possible to obtain almost quantitative yields of pyrroles **783** in 24 h (Scheme 148).^{324,325} The same catalytic system was applied in the synthesis of 2,2'-bipyrolle derivatives possessing four aryl groups in 1, 1', 5, 5' positions. The reaction was carried out with CuCl **55** as a catalyst, in DMF at 90–150 °C.^{326,327}

Modified Ullmann conditions (CuI **519**/L/Cs₂CO₃, where L = 1,10-phenatrholine, L-proline, (*E*)-4-hydroxy-L-proline) were used for the synthesis of *N*-alkenynes in hydroamination/amidation reaction of 1,4-diaryl-1,3-diynes **1a**, **1c**-**d**, **27b** with heterocyclic indoles **784a**-**d**, azoles **784e**-**h**, pyrazole **784i** and cyclic or acyclic amides **754b**, **785a**-**c**. The reaction yielded a mixture of (*Z*)- and (*E*)-*N*-alkenynes **786**-**796** with an excess of the (*Z*)-isomer in the range of 60–95%, and 75–95% yields for cyclic reagents **784a**-**i**, **754b**, and exclusive formation of (*E*)-isomer for acyclic amides **785a**-**c**. In the latter case, the yield was reduced to low to moderate values 10–41% (Scheme 149).



Scheme 145 Mechanism of hydroaminative coupling of diyne 160 with amides 754 catalysed by CpCo(C₂H₄)₂ 755.



Scheme 146 Carbon-carbon bond cleavage of diynes 1a, 13a, 617, and 767a-f with 2-aminophenols 766a-h catalysed by $Ru_3(CO)_{12}$ 489 and $Pd(NO_3)_2$ 768.



Scheme 147 Mechanism of C-C bond cleavage in diynes 1a, 13a, 617, and 765a-f based on the hydroamination reaction with 2-aminophenol 766a catalysed by Ru₃(CO)₁₂ 489 or Pd(NO₃)₂ 768.





The authors assumed that the hydroamination reaction occurred *via* an oxidative addition/reductive elimination mechanism with

the addition of the N-H bond to Cu^I **519** as an initial step of the mechanism. The insertion of diyne **1a**, **1c-d**, **27b** to the Cu-N



bond, followed by the reductive elimination of *N*-alkenyne **786–796** is postulated as the next stage of the mechanism. The system was not efficient for alkyl-substituted **1**,3-diynes and unsymmetrical reagents.³²⁸

CuCl **55** was used also for the hydroamination of *meso,meso'*-1,3-butadiyne-bridged Zn(II) diporphyrin **797** with various amines **635**, **679a**, **782b**, **798a-c** to *meso,meso'*-pyrrole-bridged Zn(II) diporphyrins **799a-f**. The structure of diporphyrin **799** was confirmed by the single-crystal X-ray diffraction method. The bulky mesitylamine **798b** and octylamine **798c** were less active in the hydroamination reaction (Scheme 150). Moreover, it was possible to modify in Suzuki–Miyaura coupling reaction of diporphyrin with 4-bromophenyl substituted pyrrole **799c** with porphyrin possessing Bpin **800** groups to **801** with 15% yield. As a catalyst PdCl₂ **64**/dppf **39b** was used. (Scheme 151).³²⁹

3- or 4-Aminomethylpyrroles **806a–k** and aminomethylfurans **807a–i** bearing a sulfur group were obtained by the hydoamination/cyclisation reaction of *N*- or *O*-tethered 1,6-diynes **802** and **803a–d** with a sulfur substituent attached to one of the alkynyl group using two catalytic systems Ni(hfa)₂ hydrate **805** (10 mol%)/DBU (Method A) or Ni(hfa)₂ hydrate

805/PdCl₂(PPh₃)₂ 94/DBU (Method B) in DMSO at room temperature (Scheme 152). The products were obtained with good yields (50-92%) using cyclic and acyclic amines 804a-n between 2-72 h (Table 20). The possible mechanism of this transformation started from the isomerisation of diyne (802, 803) to alkyneallene 809 or allene-allene 810 intermediates via a carbanion 808, followed by its coordination to the Ni atom 805 with a sulfur ligand. This activates the alkyne moiety 812 towards intermolecular attack by the amine 804. This leads to the diamino metal intermediate 815 through intermediates 813 and 814. Next, the second intramolecular cyclisation towards 816 occurred, followed by the formation of 817. Its protonolysis and isomerisation yields the product 806 or 807 and regenerates the catalyst. Less nucleophilic amines might react with water according to path II with the formation of side product 811 (Scheme 153). The presence of the sulfur group in the products 806 permitted their further functionalisation such as the introduction of formyl or acetyl groups (819a-c, 820) which are then susceptible to subsequent modification in other chemical transformations, or the reaction with the strong base leading to 1H-pyrrole 821 (Scheme 154).³³⁰



Scheme 150 Synthesis of *meso,meso'*-pyrrole-bridged Zn(II) diporphyrins **799a-f** in Cu(II)-mediated annulation of *meso,meso'*-1,3-butadiyne-bridged Zn(III) diporphyrin **797** with various amines.



Scheme 151 Synthesis of Zn(II)-free base hybrid porphyrin pentamer 801 in Suzuki-Miyaura coupling reaction of 799c with 800.

Another approach to N-heterocyclic 1,2,5-trisubstituted pyrroles **826a–e** and **829a–c** was based on the hydroamination reaction of separated 1,4- or 1,5-diynes **822a–b**, **541**, or **827** with primary amines (aniline **679a**, benzylamine **635**, or 4-methoxy-benzylaniline **782b**) with the subsequent 5-*endo* dig or 5-*exo* dig cyclisation in the presence of $Ti(NMe_2)_2(dpma)$ **823** or $Ti(NMe_2)_2(dppm)$ **827** as a catalyst. The addition of amine **679a** or **635** occurred according to the Markovnikov rule. The hydroamination of unsymmetrical 1,4-diynes with aryl and alkyl substituents led exclusively to the product with amine attached to the β -carbon to aryl substituent **826a–e**, while nonsubstituted 1,4-diynes led to the dihydroamination product, since the second hydroamination of the terminal alkyne is faster

than the intermolecular cyclisation reaction. In the case of internal or terminal 1,5-diynes **541**, **827**, the cyclisation was faster than the hydroamination of the second $C \equiv C$ bond, and the cyclic pyrroles **829a-c** were formed exclusively (Scheme 155). This method is an alternative of Paal–Knorr synthesis to pyrroles, especially when unsymmetrical 1,4-diketones are used as reagents.³³¹

Wasterhausen *et al.* published several papers on the hydroamination reactions of alkynes and conjugated 1,4-diphenylbuta-1,3-diyne **1a** in the presence of heterobimetallic complexes constructed from Ca- and K-complexes **830** and **839**. The homometallic Ca- or K-catalysts were inactive in the hydroamination reactions. Depending on the structure of the complex



Table 20	Synthesis of 3- or 4-aminomethylpyrroles 806a-k and amino-
methylfura	ans 807a-i by hydroamination/cyclisation of 1,6-diynes 802 and
803a-d v	rith amines 804a—n catalysed by Ni(hfa) ₂ 805

Entry	Diyne	Amine	Method	Time [h]	Yield [%]
1	802	804a	А	4	806a , 79,
2			В	8	806a , 100
3		804b	Α	6	806b, 77
4			В	8	806b , 88
5		804c	Α	6	806c, 75
6			В	8	806c, 100
7		804d	Α	8	806d, 50
8			В	8	806d, 76
9		804e	В	8	806e, 43
10		804f	Α	8	806f, 49
11			В	8	806f, 82
12		804g	Α	8	806g, 71
13		-	В	72	806g, 84
14		804h	В	72	806h, 66
15		804i	Α	4	806i, 70
16			В	4	806i, 43
17		804j	Α	7	806j, 74
18		U U	В	1	806j, 92
19		804k	Α	4	806k, 48
20			В	2	806k, 53
21	803a	804a	Α	6	807a, 63
22		804c	Α	8	807b, 71
23		804j	В	8	807c, 71
24		804Î	В	8	807d, 70
25	803b	804m	В	8	807e, 61
26		804n	В	72	807f, 39
27	803c	804j	Α	4	807g, 55
28		804m	В	72	807h, 61
29	803d	804m	В	1	807i , 69

and amine, various products were formed such as cyclic cyclohepta-1,2,4,6-tetraenes **832a-b** and **833**, pyrroles **834a-d**, aminated enynes **836a-c**, **840–841**, or bisaminated dienes **837a-b**. The reactions under room temperature lead to thermo-dynamic products, *e.g.*, cyclohepta-1,2,4,6-tetraenes **832a-b** or **833**, while at higher temperatures, kinetic pyrrole products **834a-d** were formed. The structure of amines has a significant



Scheme 153 Mechanism of hydroamination/cyclisation of sulfur-substituted 1,6-diynes 802 and 803 with secondary amines 804 catalysed by Ni(hfa)₂ 805.



Scheme 154 Chemical transformations of aminomethylpyrroles 806 towards formyl or acetyl-functionalised products (819a-c, 820) or 1-H-pyrrole 821.



Scheme 155 Ti-Catalysed 823 and 824 hydroamination/cyclisation of 1,4-diynes 822a-b or 1,5-diynes 541, 827. Synthesis of 1,2,5-trisubstituted pyrroles 826a-e and 829a-c.

influence on the product type (Scheme 156.). The authors discussed in detail the mechanisms of these transformations, which differ according to the hydroamination reagent.²⁹⁴

9. Hydrophosphination

Unsaturated organophosphorus compounds have found several applications as building blocks in organic synthesis, (chiral) ligands for catalyst formation, biologically active compounds, or in the preparation of flame retardant materials.^{332–337} They are also used in medicinal- or agrochemistry, as components of drugs, which are used in *e.g.*, bone, calcium-metabolism or neurological diseases, antiviral and antibacterial systems, enzymes inhibitors.^{337–345} They are commonly applied as

monodentante as well as chelating ligands in various chemical transformations.^{332–334,339} The synthesis of vinylicphoshpines can be carried out using the hydrophosphination reaction.^{30,59,346} This type of addition reaction was also used in the reaction with conjugated and separated diynes, but, unlike hydroamination, the examples are limited only to a few papers.

9.1. Hydrophosphination of conjugated and non-conjugated diynes

Hydrophosphination of conjugated 1,3-diynes **1a**, **13a**, **13c**, **60a**, **208b**, **617**, **655c**, and **842a–c** with Ph₂PH **843** was carried out in the presence of ytterbium complexes $[Yb(\eta^2-Ph_2CNPh)(hmpa)_3]$ **844** or $[Yb[N(SiMe_3)_2]_3(hmpa)_2]$ **845**. The reaction occurred according to the double addition of two diphenylphosphine This article is licensed under a Creative Commons Attribution 3.0 Unported Licence.

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1a,13a,13c 60a, 208b, 617, 655c, 842a-c

R²

Scheme 157 Double hydrophosphination of conjugated diynes with diphenylphosphine 843 catalysed by Yb-complexes 844-845.

Ph₂(O)P

(Z,E)-847

P(O)Ph₂

843 molecules to the $C \equiv C$ bonds of diyne, even at low temperatures -35 to (-78) °C, and the formation of bis(diphenylphosphinyl)-dienes **846–849** with high yields but relatively low selectivities (Scheme 157 and Table 21). The stereo-chemistry of the process was kinetically and thermodynamically

controlled and the formation of the specific isomer depended on the structure of diyne. Hydrophosphination of disubstituted diynes predominantly formed (Z,Z)-846 isomers with a minor amount of (Z,E)-847 butadiene. Terminal diynes yielded (E,Z)butadiene 848 as the main product, while the sterically hindered

(E,E)-849

P(O)Ph₂

R²

850

 Table 21
 Double hydrophosphination of conjugated diynes with diphenylphosphine 843 catalysed by Yb-complexes 844–845

Entry	Diyne	Cat.	T[°C]	Total yield [%]	Selectivity [%] 846 : 847 : 848 : 849 : 850
1	$R^1 = R^2 = Ph, 1a$	845	-78	28	0:0:0:100:0
2	$R^1 = R^2 = n$ -Bu, 13a	844	-15	92	67:33:0:0:0
3	$R^1 = R^2 = t$ -Bu, 13c	844	r.t.	89	0:0:0:0:100
4		845	r.t.	80	0:0:0:0:100
5	$R^1 = R^2 = c - C_6 H_{11},$	844	-15	74	86:14:0:0:0
	60a				
6	$\mathbf{R}^1 = \mathbf{R}^2 = n\text{-}\mathbf{Hex},$	844	-15	82	74:26:0:0:0
	208b				
7		845	-15	82	61:39:0:0:0
8	$R^1 = Ph$,	844	-15	98	73:27:0:0:0
	$R^2 = n$ -Hex, 655c				
9		845	-15	95	72:28:0:0:0
10	$R^1 = H,$	844	-78^{a}	80	0:0:61:39:0
	$R^2 = n$ -Hex, 617				
11		845	-78^{a}	89	6:0:75:19:0
12	$R^1 = 4$ -MeOC ₆ H ₄ ,	844	-15	85	73:19:8:0:0
	$R^2 = n$ -Hex, 842a				
13	$R^1 = H, R^2 = n$ -Bu,	844	-78^{a}	89	16:0:64:20:0
	842b				
14	$R^1 = H, R^2 = Ph,$	844	-78	_	Polymerisation
	842c				-
^a -78	°C for 1 h then r.t.	for 2	h.		

1,4-ditertbutyl-buta-1,3-diyene **13c** was quantitatively converted to allenic product **850**. The reaction started from the formation of the [Yb]–PPh₂ complex **851**, which underwent anti-addition to diyne to form enynylyterrbium complex **852**. Protonation of **852** with Ph₂PH **843** yields diphenylphosphine-substituted enyne **853** and regenerates ytterbium–phosphide active complex **851**. Repetition of this process provided the bishydrophosphination product **846**. The products were easily oxidised with H₂O₂ to phosphine oxides, which were easier to isolate (Scheme 158).^{347,348} The formation of diphenylphosphine-substituted enyne **853** in the reaction was also possible using an equimolar ratio of reagents and a shorter the reaction time of up to 30 minutes. After oxidation with H₂O₂, the (*Z*)-products were predominantly formed.

Tanaka *et al.* developed the hydrophospinylation reaction of alkynes in the presence of Pd(PPh₃)₄ **35** or PdMe₂(PR₃)₂ **855** (PR₃==PPh₃ or PPhMe₂, PPh₂Me, PEt₃, PMe₃) catalysts. All these complexes catalysed the synthesis of alkenyldiphenylphosphine oxides from alkynes and Ph₂P(O)H **856**. Less basic phosphines (*e.g.*, PPh₃, PPh₂Me) accelerate the formation of anti-Markovnikov products, whereas application of more basic phosphines *e.g.*, PMe₃ or PEt₃ increases the amount of the geminal regioisomer. The best results were obtained when 5 mol% of Pd(PPh₃)₄ **35** was used at 35 °C. This method was used also for hydrophosphinylation of nona-1,8-dipne **164a** derivatives towards 1,9-diphosphinyl-nona-1,8-diene **857** when 2.2 equiv. of Ph₂P(O)H **856** and Pd(PPh₃)₄ **35** was used (Scheme 159).³⁴⁹

The same authors reported Pd-catalysed hydrophosphinylative carbocyclisation of α, ω -diynes. The reaction occurred in the presence of 5 mol% Pd(OAc)₂ **861** and chelating phosphine ligands such as ethylenebis(diphenylphosphine) (dppe) or 1,2-bis(diphenylphosphino)benzene (dppben) at 70 °C in chlorobenzene, toluene, ethylbenzene, or dioxane. The carbocyclisation



Scheme 158 The mechanism of hydrophosphination of 1,3-diynes catalysed by Yb-complexes 844 and 845.

was the most effective for 1,7-heptadiyne derivatives, while longer or shorter α, ω -diynes were less susceptible to cyclisation, and linear hydrophophinylatve products were mainly obtained. Scheme 160 presents the formation of various products, which depends on the reagents used (diynes **116a** or **1270**, and phosphorus compounds **856**, or **858–860**).³⁵⁰ For diyne **116a** product **862** is mainly formed, while for **1270** product **864** is predominantly obtained (Scheme 160).

Hydrophosphinylative carbocyclisation was also reported by Yamamoto *et al.* but, instead of Pd-complexes, ruthenium catalysts **281a–c** with cyclopentadiene ligands (responsible for the formation of the active ruthenacyclopentatriene intermediate) were used. The best results were obtained for [Cp*Ru(MeCN)₃]PF₆ **281a** for which exocyclic 1,3-dienylphosphine oxides **866a–j** were exclusively formed, under the optimised reaction conditions using HP(O)Ph₂ **856.** In the case of complexes **281b–c**, the hydrative cyclisation product **867** was formed as well. To suppress the formation of these by-products molecular sieves 4 Å were added to the reaction mixture. The [Cp*Ru(MeCN)₃]PF₆ **281a** was used in 5–10 mol%, depending on the reactivity of diynes **865a–j.** The substituents in the tether position have a significant influence on the product yields. The more hindered these groups, the lower the



yields of the desired products **866a–j** observed. To obtain the desired cyclic product, 1,6-heptadiyne derivatives need to be used with the aryl substituents in the terminal position (Scheme 161). When shorter chain dignes such as 1,5-hexadiyne, or reagents with alkyl substituents in terminal positions were used, the reaction did not occur or the products were formed in low yield. The aryl terminal groups accelerate the formation of active ruthenacyclopentatriene intermediate, which is essential for this transformation.

The mechanism of this transformation was proposed based on DFT calculations. The reaction started with the oxidative cyclisation of bis(alkyne) complex **868**. The rate-determining step was found to be H-atom transfer, which leads to the monocarbenoid with a phosphinate ligand **869**. The intramolecular attack of phosphorus on the remaining carbenoid carbon generates oxaphospharuthenatricycle **870** followed by the formation of (*Z*)-4-diene complex **871**. The addition of diyne **865** and HP(O)Ph₂ **856** generates product **866** and regenerates the active catalytic intermediate **868** (Scheme 162).³⁵¹

Hydrophosphinylation of symmetrical 1,4-diphenyl or 1,4*tert*-butyl-buta-1,3-diynes (**1a** or **13c**) was also carried out in the presence of a main group element catalyst [(thf)₄Ca(PPh₂)₂] **872**. The composition of the postreaction-mixture depended on the phosphorus reagent. When diphenylphosphane oxide HP(O)Ph₂ **856** was used as reagent, 1,4-diphenyl-2,3-bis(diphenylphosphoryl)-buta-1,3-diene **873** or 2,2,7,7-tetramethyl-3,6-bis-(diphenyl-phosphoryl)-4-octyne **874** were selectively formed in the reaction with 1,4-diphenylbuta-1,3-diyne **1a** or 1,4-di-*tert*butylbuta-1,3-diyne **13c** respectively in very good yields (80–82%). The reaction with Ph₂PH **843** yielded different products in 1,4- or 1,3 phoshponylation (**875–876**). These differences in process selectivity are due to the different base-acid interactions between calcium catalyst **872** and Ph₂PH **843** or HP(O)Ph₂ **856**. Rather, strong Ca–O interactions are responsible for the closeness of the alkali metal to reactive multiple C–C bonds (Scheme 163).³⁵²

10. Hydration of conjugated 1,3- and separated 1,*n*-diynes

Hydration of 1,3- and 1,*n*-diynes is limited to several examples, which are focused on the catalytic activation of the water molecule and diyne with various catalysts mostly based on transition metals. This transformation leads to many important



building blocks, especially in cyclisation reactions to furanes, 3-(2*H*)-furanones, or γ -pyrones. The obtained products are used in the synthesis of antitumor agents, antibiotics, natural and bioactive compounds.^{353–355} The addition of water to the C \equiv C triple bond may also yield carbonyl compounds *via* tautomerisation of the hydroxylated enyne.

Hydration or hydration/cyclisation reactions are simple and 100% atom economic transformations, which can provide the desired products in a straightforward procedure, without (or with a small number of) side-products. Therefore, they are a useful alternative to common methods, that require the application of complex reagents and multi-step procedures. Most of the catalytic systems for selective hydration reactions of diynes, which are based on predominately gold, ruthenium and palladium complexes, were developed within the last two decades.

The first papers on the hydration of diynes were published in the 1960s and apply mercuric salts. The addition of water to undeca-1,7-diyne **877** provided a mixture of two diketones, undecane-2,7-dione **878** and undecane-2,8-dione **879** with moderate yields and low selectivity.^{356,357} A modified procedure was used by Constantino *et al.* in the preparation of natural marine compound 1-(2,6,6-trimethyl-4-hydroxy-cyclohexenyl)-1,3-butanedione **880**, which possess antibiotic activity. They used HgSO₄ and formic acid (85%) in the hydration step. The compound was formed in 50% crude yield. The same system was applied for hydration of other cyclohexyl-substituted diynes. The terminal C \equiv C group was hydrated at first, followed by the reduction of the second alkynyl group.³⁵⁸

Ruthenium catalysed hydrative cyclisation of various diynes was studied in detail by Trost *et al.*^{359–364} They have found that simple cationic $[Cp*Ru(CH_3CN)_3]PF_6$ **281a** complex, which catalyses many different transformations such as alkyne–alkyne coupling reactions (*e.g.*, dimerisation, trimerisation)^{365–368} or cycloaddition reactions with dienes,³⁶⁹ isocyanates,³⁷⁰ nitriles,³⁷¹ can be effectively used in diyne hydrative cyclisation or cycloisomerisation reactions (Scheme 164).^{359–364} Depending on the structure of the diyne, different mechanisms for the reaction occur. Internal diynes can directly react with water in the presence of catalytic amounts of the Ru complex **281a** (3–10 mol%) producing five- or six-membered enones with moderate or excellent yields. The same catalyst was used for the dimerisation of propargylic alcohol and a further intramolecular cycloisomerisation reaction (Scheme 164). Tertiary or secondary propargylic



Scheme 162 Proposed catalytic cycle for hydrophosphinylative coupling reaction.

alcohols cycloisomerise to $\alpha, \beta, \gamma, \delta$ -unsaturated aldehydes and ketones, while primary propargylic alcohols also gave the hydrated cyclised product. The key step in both paths of

mechanism (Cycle A and Cycle B) starts from the reseonance invocation to ruthenacyclopentatriene 904. For primary propargylic alcohol divnes, the addition of water might occur to two carbene carbons yielding intermediates 905 and 910. The hydrative cyclisation process leads to the rearrangement of 905 to 907, followed by a hydride shift and protontion to the product 909. In the case of cyclodimerisation, compound 910 is rearranged to **911** which, after hydride shift and β -hydroxide elimination or protonation and water elimination, leads to product 913 (Scheme 165). The mechanism common for secondary and tertiary propargylic alcohols (which possess better-leaving groups) occurs mainly via Cycle B. The detailed mechanistic studies on the activation of a water molecule by ruthenacyclopentatriene 904 were studied by Yamamoto et al. Using DFT calculation, they postulated the formation of half-open oxaruthenocene as an initial step of the mechanism.³⁷² The methodology was used in the cyclisation of various divnes (Scheme 164). Moreover, the directing effect of carbonyl group attached to the $C \equiv C$ in the δ - or ϵ -position was observed, by the coordination of C=O to ruthenacyclopentadiene complex 914.³⁶⁰ Hydrative cyclisation was a step in the formation of natural compounds: tricyclic alkaloids Cylindricine C 920 (Scheme 166), while cycloisomerisation was used in the synthesis of (+)-a-kainic acid 933 (Scheme 167).^{362,364} Moreover, the cyclised products were applied in both intra- and intermolecular Diels-Alder reactions.^{360,361}

Another example of the application of hydration process is the formation of functionalised benzene derivatives **936a–e** in the aromatisation of enediynes **934a–e** catalysed by 10 mol% [TpRu(PPh₃)(CH₃CN)₂][PF₆] **935** (Tp = tris(1-pyrazolyl)borate) (Scheme 168).³⁷³ The process is also possible for the addition of other nucleophiles than H₂O (*e.g.*, aniline, acetylacetone, pyrroles, and dimethyl malonate) to non-functionalised enediynes



Scheme 163 Intermolecular hydrophosphanylation of butadiynes 1a and 13c with diphenylphosphane oxide 856 and diphenylphoshpine 843 catalysed by (thf)₄Ca(PPh₂)₂ 872.



Scheme 164 Hydrative cyclisation/cyclodimerisation of separated diynes catalysed by [Cp*Ru(CH₃CN)₃][PF₆] 281a.

934. The addition is highly selective and the attack occurs at the more electron-rich alkyne carbon yielding various functionalised aromatic compounds **936a–e** depending on the nucleophile. The mechanism was proposed according to the reactions with D₂O. These experiments proved that the catalytically active species is a ruthenium- π -alkyne complex instead of the ruthenium-vinylidene intermediate, which is a characteristic step in Saito-Myers cyclisation (Scheme 169).³⁷³

Gold complexes are another big class of catalysts, which have been used in the hydration of conjugated and separated diynes. The presence of water was essential for the hydrative cyclisation. In 2010, Skrydstrup *et al.* published that 1,3-diynes can be converted in a hydration reaction towards 2,5-disubstituted furans **946** (Scheme 170) or in a hydroamination process to 1,2,5-trisubstituted pyrroles **681** (Scheme 132). Au(1) complexes such as (Ph₃P)AuNTf₂ **680** and SPhosAuNTf₂ **945** were able to catalyse these two reactions under mild conditions. Complex **945** was more active in hydration reaction since H₂O is a better nucleophile when 1,4-diaryl or dialkylbuta-1,3-diynes were used. Within this methodology, it was possible to furnish a selection of 2,5-diamidofurans **946k–m** in 45 minutes with good to moderate yields (Table 22, entries 14–17). To obtain



Scheme 165 Hydrative cyclisation (Cycle A) and cyclodimerisation (Cycle B) mechanism based on the reaction of primary diynols with H₂O in the presence of [Cp*Ru(CH₃CN)₃][PF₆] **281a**.



high yields in the case of the hydration of symmetrical **1a**, **1c-d**, and non-symmetrical diraryl-**944b** or dialkyl-substitued **208b** diynes using complex **945**, 24 hour reaction times were necessary. Moreover, increasing the polarisation of the diyne by the introduction of electron-donating OMe groups led to a small amount of side products (Scheme 170). When D_2O was used instead of H_2O , furans **946e** with deuterium atom at 3,4-position were synthesied.³⁰⁸ The same products were formed, when [Au(IPr)OH] **695** was used as a precursor. The reaction proceeded only in the presence of Brønsted acid HX, which generated the active complex [Au(IPr)]X **947** *in situ*. HNTf₂ was the most effective in the model reaction with 1,4-diphenybuta-1,3-diyne **1a**. Poorer results were observed when HBF₄ **948** and HPF₆ **949** were used (77% *vs.* 37–39%). No catalyst activity was noticed for the complex with SbF₆⁻ or OTf⁻ groups. Elevated temperatures



(+)- α -Kainic acid

a) 10 mol% [Cp*Ru(CH₃CN)₃][PF₆] **281a**, 2% H₂O/acetone, 40 °C, 1 equiv. malonic acid; b) Li[SiMe₂Ph] **923**, CuCN, THF, -79–0 °C. 2.8:1 dr; c) DBU, beznene, reflux; d) 5% Pd/C **446**, HCOOH:MeOH (1:1); e) 20 mol% [Ir(cod)Py(PCy₃)][PF₆] **927**, 138 bar H₂, 1 equiv. B(O-*i*Pr)₃; f) Li(CH₂SiMe₃) **929**, THF, -78 °C; g) 1) HF, H₂O CH₃CN, 2) KH, *t*-BuOOH, TBAF **108**, DNF, 65 °C; h) 8 N Jones' reagent, acetone. r.t.; i) 1) Li, NH₃ **632**, THF, -78 °C, 2) ambertile CG-50.

Scheme 167 Multistep synthesis of $(+)-\alpha$ -Kainic Acid **933** with the cycloisomerisation step.



Scheme 169 The general mechanism of aromatisation reaction catalysed by [TpRu(PPh₃)(CH₃CN₂)][PF₆] 935 in the presence of various nucleophiles.



Scheme 170 Gold(I) catalysed synthesis of 2,5-disubstituted furans 946a-r.

are needed to perform the reaction with the [Au(IPr)OH] **695**/HX system. Additionally, the type of substituents attached to buta-1,3-diyne skeleton is important for the reaction. Diynes with aryl groups in the terminal positions were the most active in the formation of 2,5-disubstituted furanes **946**. For the diyne with cyclohexene groups (**2580**) the reaction was less effective, while reagents with dialkyl sidechains in positions 1,4 did not

 Table 22
 Results of gold(I) catalysed synthesis of 2,5-disubstituted furans
 946a-r

Entry	Diyne	\mathbb{R}^1	\mathbb{R}^2	Method ^{<i>a,b,c,d</i>}	Yield	of 946a-r [%]
1	1a	Ph	Ph	b	a	73
2	1b	$4-FC_6H_4$	4-FC ₆ H ₄ l	c	b	82
3	1c	$4-MeC_6H_4$	$4-MeC_6H_4$	b	с	80
4				c	с	82
5	1d	$4-MeOC_6H_4$	$4-MeOC_6H_4$	b	d	84
6				b, d	e	79
7				c	d	82
8	27b	4-t-BuC ₆ H ₄	4-t-Bu-C ₆ H ₄	c	f	84
9	60e	4-MeOC ₆ H ₄	Ph	c	g	65
10	208b	<i>n</i> -C ₆ H ₁₃	$n - C_6 H_{13}$	b	ĥ	68
11	242b	$2-MeOC_6H_4$	$2-MeOC_6H_4$	c	i	71
12	2580	$c-C_6H_9$	$c-C_6H_9$	c	j	75
13	271a	Ph	t-Bu	c	k	62
14	678a	NTs(Bn)	NTs(Bn)	a	1	85
15	678b	NTs(Ph)	NTs(Ph)	a	m	51
16	944a	NTs(<i>i</i> -Pr)	NTs(<i>i</i> -Pr)	a	m	82
17				c	n	77
18	944b	4-MeOC ₆ H ₄	4-BrC ₆ H ₄	c	0	82
20	944c	4-MeOC ₆ H ₄	c-C ₆ H ₉	c	р	72
21	944d	4-MeOC ₆ H ₄	<i>n</i> -Bu	c	q	64
22	944e	4-MeOC ₆ H ₄	2-MeOC ₆ H ₄	с	r	71

 a 1.4 equiv. H₂O, 2 mol% (PPh₃)AuNTf₂ **680**, THF, 60 °C, 45 min. b 10.0 equiv. H₂O, 2–5 mol% SPhosAuNTf₂ **945**, THF, 60 °C, 24 h. c H₂O, 1 mol% [Au(IPr)OH] **695**, 1.5 mol% HNTf₂ dioxane, 80 °C, 4 h. d D₂O used.

lead to the desired products. When one of these group was substituted with an aryl ring, the reaction occurred with good yield (Scheme 170 and Table 22). According to stoichiometric experiments and DFT calculations, it was proved that the reaction proceeded *via* hydration of the one $C \equiv C$ bond. Two pathways are possible through the keto or enolate form. It was determined that the keto-pathway is favoured by 9.6 kcal mol⁻¹ (Scheme 171).³¹²

Hydration of conjugated diynes was used for the synthesis of 6,5,6-trioxabispiroacetal moieties, the spacers between the steroid cores. Steroid diynediols were used as reagents, while JohnPhos-Au(MeCN)SbF₆ **682** was applied as a catalyst.^{374,375}

Sanz et al. reported the Au-catalysed hydration-oxacyclisation reactions of 1,4-diyn-3-ones 959a-s, which were obtained from ethyl lactate as carbonyl source, a feedstock derived from biomass. Depending on the catalytic system composition it was possible to carry out the selective synthesis of 4-pyrones 960a-s or 3(2H)-furanones 961-962. Such compounds possess many biological activities, e.g., phenoxans, funicones and rapicones indicate anti-HIV activity.^{376,377} The reaction can be tuned by the ligand attached to the gold complex, the presence or absence of silver salts, and the counteranion. When 5 mol% of IPrAuNTf₂ 711 was used 4-pyrones 960a-s were predominantly formed (5:1-20:1), while applying 5 mol% of AuCl(PPh₃) 694/AgSbF₆ 746 3(2H)-furanones 961-962 were obtained (1:11-1:20). Both products were formed in moderate yields of 65-86%. The lowest yield of 3(2H)-furanones 961 was obtained for alkylsubstituted diynones. This pathway was much more effective for aryl- or heteroaryl-functionalised diynones, while 4-pyrones 960a-s were furnished with similar yields regardless of the type of substituents (Scheme 172 and Table 23). This is an alternative method towards 4-pyrones and furanones, which are typically made by multistep condensation cyclisation reactions of carbonyl compounds.378 The mechanism of this transformation was demonstrated from the reaction with D₂O. The key step in the formation of 4-pyrones 960 or 3(2H)-furanones 961 is the hydration of diynone 959, which might proceed according to Michael or anti-Michael addition. Both pathways are possible and depending on the catalytic system. Next, the intramolecular oxacyclisation occurred leading to Au-intermediates 965 or 966. Finally, protodeauration affords the final products with the



Scheme 171 Possible catalytic pathways for Au promoted hydrative cyclisation of 1,4-diphenylbuta-1,3-diyne 1a.



a: 5 mol% IPrAuNTf₂ 711, 1 h, 0.5. mmol of 959a-s, H₂O (1 mL), dioxane (2 mL)

b: 5 mol% AuCl(PPh₃) 694 / AgSbF₆ 746, 5 h, 0.5 mmol of 959, H₂O (1 mL), dioxane (2 mL)

Scheme 172 Synthesis of 4-pyrones 960a-s and 3(2*H*)-furanones 961 and 962 in the hydration-oxacyclisation of symmetrically and non-symmetrically substituted diynones 959 catalysed by Au(i) complexes.

Table 23 Results of the hydration-oxacyclisation reactions of symmetrically and non-symmetrically substituted diynones catalysed by Au(I) complexes

Entry	Diynones 959	R ¹	R ²	Method ^{ab}	Selectivity 960 /(961 + 962) ^{<i>c</i>}	961/962 ^c	Product, yield ^{d} [%]
					960/(961 = 962)	_	
1	a	Ph	Ph	а	9/1	_	960a , 73
2				b	1/18	_	961a, 80
3	b	<i>p</i> -Tol	<i>p</i> -Tol	а	10/1	_	960b, 81
4				b	1/11	_	961b, 77
5	с	$4-MeOC_6H_4$	4-MeOC ₆ H ₄	а	12/1	_	960c, 83
6				b	1/>20	_	961c, 81
7	d	3-MeOC ₆ H ₄	3-MeOC ₆ H ₄	а	> 20/1	_	960d , 78
8				b	1/>20	_	961d , 79
9	e	$4-FC_6H_4$	$4-FC_6H_4$	a ^e	10/1	_	960e , 79
10				\mathbf{b}^{e}	1/10	_	961e, 70
11	f	3-Th ^f	3-Th ^f	а	5/1	_	961f, 70
12				b	1/18	_	961f, 79
13	g	2-Th^{f}	2-Th^f	а	1/18	_	961g, 74
14	ĥ	<i>n</i> -Bu	<i>n</i> -Bu	а	> 20/1	_	960h, 81
15				\mathbf{b}^{g}	1.5/1	_	961h, 35
16	i	<i>c</i> -C ₃ H ₅	<i>c</i> -C ₃ H ₅	а	> 20/1	_	960i, 80
17				\mathbf{b}^{g}	2.5/1	_	961i, 26
18	j	$(CH_2)_2Ph$	$(CH_2)_2Ph$	а	> 20/1	_	960j , 86
19	k	c-C ₆ H ₉	c-C ₆ H ₉	а	> 20/1	_	960k, 67
20	1	$C(CH_3) = CH_2$	$C(CH_3) = CH_2$	а	> 20/1	_	960l, 74
21	m	$CH_2O(4-MeOC_6H_4)$	$CH_2O(4-MeOC_6H_4)$	а	> 20/1	_	960m, 65
22	n	$CH_2O[3,5-(MeO)_2C_6H_3]$	$CH_2O[3,5-(MeO)_2C_6H_3]$	а	> 20/1	_	960n, 70
					(960/(961 + 962))		
23	0	Ph	<i>n</i> -Bu	а	>20/1	_	9600, 82
24				b	1/1.25	1/10	9600, 35, 9610, 42
25	р	Ph	<i>c</i> -C ₃ H ₅	b	1/1.1	14/1	960p , 37, 961p, 35
26	q	Ph	4-MeOC ₆ H ₄	b	1/20	3/1	961q , 71
27	r	$4-FC_6H_4$	$4-MeOC_6H_4$	b	1/10	4/1	960r, 8, 961r, 72
28	S	Ph	Н	а	> 20/1	_	960s, 74
				b	>20/1	_	960s, 76

^{*a*} Method a: 5 mol% IPrAuNTf₂ 711, 1 h, 0.5. mmol of 959, H₂O (1 mL), dioxane (2 mL). ^{*b*} Method b: 5 mol% AuCl(PPh₃)₃ 694/AgSbF₆ 746, 5 h, 0.5 mmol of 959, H₂O (1 mL), dioxane (2 mL). ^{*c*} Determined by ¹H NMR analysis. ^{*d*} Isolated yields after column chromatography. ^{*e*} 10 mol% of catalyst was used. ^{*f*} Th = thienyl. ^{*g*} 8 hours.

elimination of the catalytic species. The regioselectivity is controlled by hydration step, not by a 6-endo vs. 5-exo oxacyclisation reaction (Scheme 173). The Michael or anti-Michael addition of water also had an influence on the synthesis of furanones and pyrones when unsymmetrically substituted diynones **9590-s** were used. Anti-Michael addition was favoured with the more electronpoor alkyne group causing the synthesis of furanones in a higher amount.³⁷⁸ This methodology was used in the preparation of *Polyporapyranone B* **969**, which is naturally occurring γ -pyrone in sea-grass derived fungi *Polyporales* (Scheme 174). The hydrationoxacyclisation reaction is the final step in the synthesis of this bioactive compound, proceeded by Sonogashira coupling of 2,4dimethoxyiodobenzene **967** with propargylic alcohol, oxidation, the addition of ethynylmagnesium bromide, and the next oxidation step. Finally, both products, which could be prepared on a gram scale, were utilised in further transformations leading to pyrylium salts, that can be used as photoredox catalysts or in the reaction with *N*-nucleophilic reagents to functionalised N-heterocycles (Scheme 175).³⁷⁸

Diynones **959** were also converted to 4-pyranones **960** in the presence of TfOH **974**, which promotes the hydration reaction, followed by cyclisation. The reaction occurred under metal-free conditions, making the process more legitimate in the case of the process economy (no expensive gold catalysts) and sustainability. Under the optimised conditions (1 equiv. TfOH **974**, 100 °C, 36 h) various symmetrically and non-symmetrically 2,6-substituted 4-pyranonens **960** were obtained with good yields (57–82%). Other acids as *e.g.*, *p*-TSA or PhCOOH were much less



Scheme 173 Michael and anti-Michael hydration as a key step in hydration-oxacyclisation reactions of diynones **959** in the presence of Au-complexes IPrAuNTf₂ **711** and AuCl(PPh₃)₃ **694**/AgSbF₆ **746**.





active than TfOH **974**. Diynones **959** substituted in the terminal position with aryl groups bearing electron-donating groups (*e.g.*, Me, *t*-Bu, OMe) gave products with slightly better yields,

than those with electron-withdrawing groups (e.g., F, Cl). The mechanism of the reaction starts from the activation of carbonyl group in diynone 959 by TfOH 974 and nucleophilic



Scheme 176Proposed mechanism of hydration/cyclisation of diynones959catalysed by TfOH974.

addition of water to the $C \equiv C$ triple bond followed by a ketoenol tautomerisation towards intermediate **963**. Subsequently the protonation and C–C bond rotation, which occurred under elevated temperature leads to species **975**. The cycle is subsequently closed by the intramolecular nucleophilic attack of the oxohydryl group to the second $C \equiv C$ to give cyclic intermediate **976**, which furnished 4-pyrorone **960** after deprotonation (Scheme 176).³⁷⁹

MeAuPPh₃ 978 or (PPh₃)AuNO₃ 979 with trifluoromethanesulfonic acid (TfOH) 974 are active catalysts in the hydrative cyclisation of 1,6-heptadiynes 116a, 127a, 127e, 127j, 394b, 394g, 450a, 450c, and 977a-o functionalised with various different groups, e.g., alkoxy, esters, carboxyl, carbonyl, phenyl, or nitrile (Scheme 177).^{380,381} Other acids as mineral H₂SO₄ or heteropolyacids H₃PW₁₂O₄₀ 982a, H₃PMo₁₂O₄₀ 982b, and H₄SiW₁₂O₄₀ 982c were also used as co-catalysts and permitted the isolation of the corresponding 3-methyl-hex-2-enone but with lower yields. The proposed mechanism of this transformation assumes the formation of an active Au⁺ species in the first step. The coordination of divne, followed by the H₂O attack then leads to intermediate 983, which further isomerises to gold cyclohexanone complex 984 by the intramolecular attack of enolic ion to the gold cation binding through the $C \equiv C$ bond. Product 981 is then released through a tandem double bond isomerisation process and gold catalyst elimination (Scheme 178).³⁸⁰

Moreover, ILs were used as solvents and immobilisation media for (PPh₃)AuNO₃ **979**. The best results were obtained for [BMIM][BF₄], allowing to obtain stable products with yields of 72–78% in six cycles of hydrative cyclisation of **127a**. Such strategy permitted the recycling of the expensive gold catalyst. No information about catalyst leaching was presented.³⁸¹

The same authors discovered that the $Pt(cod)Cl_2$ **980** catalyst with TfOH **974** as a co-catalyst is active in the hydrative cyclisation of the same reagents (1,6-heptadiynes): **116a**, **127a**,



Scheme 177 Au(I)- or Pt(II)-catalysed 978-980 hydrative cyclisation of terminal 1,6-diynes 116a, 127a, 127e, 127j, 394b, 394g, 450a, 450c, 977a-o.



Scheme 178 Plausible mechanism of hydrative cyclisation of 1,6-diynes catalysed by Au(i) complexes **978–979**.

127e, 127j, 394b, 394g, 450a, 450c, 977a-o functionalised in position 4. This catalytic system furnishes 3-methyl-hex-2-enone **981** with good yields (Scheme 177). The mechanism of the reaction was similar to that presented for Au-catalyst in Scheme 178. Interestingly Ru- or Pd-complexes were not active in this transformation.³⁸²

Liu *et al.* reported platinum and gold-catalysed hydrative cyclisation or carbocyclisation of oxo diynes or triynes, which led to benzopyrones and bicyclic spiro ketones.^{383–385} As a model reagent, diynone **986a** was used which gave products **987a–991a** depending on the catalyst used (Scheme 179 and

 Table 24
 Influence of the catalyst (Pt or Au) on the products and yields in the hydration/cyclisation reactions of **986a**

Entry	Cat ^a	Process conditions	Yield of products ^c [%]
1	992	25 °C, 12 h	987a , 65
2	992	100 °C, 12 h	988a , 63
3	993	100 °C, 14 h	988a, 74
4	694	25 °C, 14 h	989a , 78, $dr = 2:1$
5	994	100 °C, 5 h	990a, 61
6	995 ^b	100 °C, 12 h	991a , 47, 990a, 12
<i>a</i> =	(f		

^a 5 mol% for 694, 992-993 catalyst, 8 mol% for 994-995, 1,4 dioxane,
 [986a] = [0.15 M]. ^b 10 mol% of 2,6 lutidine. ^c Isolated yields.

Table 24). Simple AuCl 992 led to diketone product 987a, when the hydration step was carried out at room temperature. Increasing the temperature to 100 °C provided 1-H-inden-1one 988a as the main product, while spiroketone 989a was obtained using PPh₃AuCl 694/AgOTf as a catalyst. Product 987a is an intermediate in the synthesis of spiro ketone 989a. Switching from gold to a platinum catalyst, by application of PtCl₂/CO 994, the chemoselectivity was directed to benzoisochromene 990a. Triketone 991a was formed when 10 mol% of lutidine was added to the catalytic system. The yield towards 990a was improved by the application of 1 atm of CO, which role is to increase the nucleophilcity of Pt(II) by the formation of PtCl₂(CO)_n. Moreover, CO was essential to increase the process selectivity to 990a. PPh₃AuCl 694/AgOTf produced spiro ketones 989b-j with a very good yields (63-88%), depending on the substrate 986b-i (Scheme 180). The authors postulated that the ketone group accelerates the hydration of proximate C(1)-carbon of the neighboring alkyne (according to intermediate 996). The obtained diketone 987a undergoes a Conia-ene transformation³⁸⁶ based on the attack of its enol form 997 at the π -alkyne group to form indenyl ketone 998. A subsequent gold-or proto-catalysed aldol reaction formed spiro ketone 989a (Scheme 181). Whereas PtCl₂/CO 994 catalyses the transformation of various diynones 986 to isochromenes 990, hydrative cyclisation of diynones 986 and 999 or diynals 1002 catalysed by PtCl₂ 1000 furnishes benzoisochromenes 1001 or



Scheme 179 Chemoselectivity of the hydration/cyclisation process depending on the catalyst type: Pt- or Au-based.



Scheme 180 PPh₃AuCl 694/AgOTf catalysed synthesis of spiro ketones via hydration/cyclisation reactions.



Scheme 181 Mechanism of the hydration/cyclisation reaction of diynone 986a to spiro ketone 989a catalysed by PPh_3AuCl 694/AgOTf.

primary lactol derivatives **1003** (Schemes 182 and 183). The mechanism of this transformation was proposed on the basis of the reaction with D_2O . The diynone **986a** leads to the formation of benzopyriliums **1004**, which is transformed to triketone **1005**.

Next, the aldol condensation of **1005** catalysed by a Brønsted acid or $PtCl_2$ **1000** produces 1-naphtol **1007** *via* enol intermediate **1006**. Finally, the tetracyclic ketal **990**' is formed, which is reduced by D₂O. Oxonium intermediate **1010** then undergoes hydride



Scheme 182 Hydration/cyclisation of diynones 986b, 986g-h, 999a-b to benzoisochromenes 1001 catalysed by PtCl₂ 1000.





Scheme 184 Plausible mechanism including the secondary hydrogenation of primary ketal product **990**' to beznoisochromenes catalysed by PtCl₂ 1000.

addition by $DPtCl_2.$ Its formation from CO and $HOPtCl_2^-$ was reported in the literature (Scheme 184). $^{\rm 387}$

In the case of triynes **1012a–g** and **1015a–m**, applying PtCl₂/ CO **994** (or more active PtI₂/CO **1016**) as a catalyst led to nucleophilic hydration of the alkyne moiety, followed by the cyclisation led to tetracyclic **1013a–g** and **1014a–g**, or bicyclic spiro ketones **1017a–m** with excellent selectivity (Scheme 185). The authors postulated that the formation of products occurred according to two hydration processes, further alkyne insertion, and aldol condensation. The type of product which is formed depends on the order of the hydration process. Spiro ketones **1017a–m** are synthesised when the initial hydration occurred at the central diphenyl alkynes. When the outer alkyne is hydrated at first, tetracyclic ketones **1013a–g** and **1014a–g** were effectively synthesised. Both types of products were formed in good yields (Scheme 185).^{383,384}

Conjugated 1,3-diynes can be converted to 2,5-disubstituted furans using a simple and cheap copper(I) catalyst **519**, which constitutes an alternative to reactions catalysed by much more expensive Au(I) complexes.^{308,312,388} The formation of furans can occur directly from haloalkynes **1018a–n** and **1019a–n** *via* preliminary Glaser coupling to 1,3-diynes **1a–d**, **27c**, **37t**, **230d**, **258a**, **258g–h**, **265a**, **500b**, **1020a–b** or direct hydration of diynes **60e**, **655a**, **1020c–e**, followed by cyclisation. As a base, KOH was Synthesis of bicyclic spiro ketones



Scheme 185 PtCl₂/CO 994 or Ptl₂/CO 1016 catalysed synthesis of tetracyclic ketones 1013a–g and 1014a–g or bicyclo spiro ketones 1017a–m *via* hydration/cyclisation of triynes 1012a–g and 1015a–m.



Scheme 186 Synthesis of 2,5-disubstituted furans 1021a-s via Glaser coupling and hydration process catalysed by Cul 519/1,10-phen system.

used, and CuI **519**/1,10-phen was much more active than CuCl **55** or CuBr **1022** (Scheme 186). The mechanism of both subsequent processes: Glasser coupling and hydration is presented in Scheme 187.

The hydration of 1,3-diyne alcohols was also catalysed by non-metal systems, based on base-functionalised ionic liquids under an atmosphere of CO_2 . The best results were obtained using [HDBU][BenIm] which possess moderate basicity. DFT calculations proved that the process started from the reaction of 2-methyl-6-phenylhexa-3,5-diyn-2-ol **1028** with CO_2 , followed by intramolecular cyclisation, which was estimated to be the rate-determining step in this reaction. Then the cyclic



Scheme 187 Mechanism for the synthesis of 2,5-disubstituted furans 1021a-s from haloalkynes 1018a-n and 1019a-n.

carbonate is hydrolysed and CO_2 is released by the base [BenIm]. Finally 3(2*H*)-furanone is formed through isomerisation with the base catalyst and the intramolecular cyclisation. Much better results were obtained when protic ILs were used.³⁸⁹

Performing the hydration/cyclisation process with InI₃ 1030 as a catalyst and para-toluene sulfonic acid (p-TSA) 1031 as a co-catalyst, it was possible to obtain 2-disubstituted tetrahydrofurans 1032a-h and 1034a-b from 1,7- or 1,8-diynyl ethers 1029a-h and 1033a-b with moderate yields. The products contain an exocyclic enone part. The best yields were obtained for reagents bearing with nucleophilic aryl groups in the terminal positions (Scheme 188).³⁹⁰ The authors also postulated the mechanism of the reaction, which started from the activation of homopropargylic alkyne by chelation of InI_3 **1030**, with the ether oxygen atom. This accelerates the initial 7-endo-dig cyclisation with the nucleophilic aryl alkyne. The presence of such an aryl ring is necessary for the desired reaction course. Next, hydration occurred, which furnishes enol 1037, which via elimination process leads to acyclic cross-conjugated dienone 1038. The mechanism is concluded by the protonation of 1038 to tertiary carbocation 1039 and its cyclisation to the desired 2-disubstituted tetrahydrofurans (Scheme 189).390

In addition to the application of metal based catalysts in the hydration reaction of diynes, there are also some examples

focused on photocatalytic processes. Photohydration of nonsymmetricaly substituted conjugated 1,3-diynes 781c, 1040a-g with an aryl (naphthyl, phenyl, 4-MeOC₆H₄, 4-CH₃COOC₆H₄ 4or 3-CF₃C₆H₄, 4- or 3-NO₂C₆H₄) or alkyl groups (tert-butyl, methyl) occurred in an aqueous sulphuric acid solution. The acidity influences the ration of products 1041 and 1042, which differs in the hydration of a specific $C \equiv C$ bond. A mediumacidity gives quantitative yields of hydration, and azulenequenching postulating that the singlet excited state furnishes both 1041 and 1042 photoadducts. The triplet excited state yields only 1041 photoadducts when R is an alkyl group. Moreover, the type of the substituent attached to the aryl group has an influence on the photohydration process is in the order of $3-NO_2 > 4-NO_2 > 3-CF_3 > 4-CF_3 > 4-CO_2CH_3$. Depending on the reagent, various products with a carbonyl group attached to the $C \equiv C$ or allenic structures were obtained, which are presented in the mechanism shown in Scheme 190.391,392

11. Hydrothiolation of conjugated 1,3-diynes

Hydrothiolation of conjugated diynes is carried out mostly according to two pathways: (i) nucleophilic addition of thiols





Scheme 188 Synthesis of 2-disubstituted tetrahydrofurans 1032a-h and 1034a-b via hydrative cyclisations of 1,7-and 1,8-diynyl ethers 1029a-h and 1033a-b catalysed by InI_3 1030.

Scheme 189 Proposed mechanism for hydrative cyclisation of 1,7-diynyl ethers 1029 catalysed by InI_3 1030.

to unsaturated carbon–carbon bonds in the presence of various alkaline metal bases or, (ii) according to radical processes. Vinyl sulfides obtained in the hydrothiolation reactions are the components of several drugs used in the Alzheimer's, Parkinson's, cancer, or AIDS diseases.^{393,394} They are also important building blocks in organic synthesis, which might be easily converted to carboxylic acids, ketones, or aldehydes in a thio-Claisen rearrangement. They can also be used in Michael, Peterson, or Diels–Alder transformations as well being easily reductively cleaved.^{395–401} Vinylsulfides were isolated in the biologically active compounds *e.g.*, Griseoviridin from *Streptomyces graminofaciens* or benzylthiocrellidone from *Crella spinulata.*^{402–404}

Nucleophilic addition of thiolate anions to $C \equiv C$ bonds in alkynes and diynes occurs mainly according to a *trans*-addition reaction with the generation of (*Z*)-vinylic isomers. The formation of these nucleophilic species occurred predominantly in the reactions with strong bases (*e.g.*, hydroxides: KOH, NaOH, or alkoxides: NaOR or KOR). In most cases, the other possible isomers are accomplished by the post-reaction mixture. The addition of thiols **1046a–d** to conjugated 1,3-diynes **1a**, **180c**, **1045** led to the formation of 1,4-dithiol-1,4-disubstituted dienes **1047**. The reaction occurred in a stepwise process. First, the monothiolate 1,3-enyne is formed, which then is hydrothiolated to 1,4-dithiol-1,4-disubstituted dienes **1047**. The obtained products can be cyclised to dithiins **1048** by deprotection of the thiol group with Li in liquid NH₃ **632**,

followed by the oxidation of thiolate anions with I_2 **418** in KF. These cyclic compounds **1048** can be potentially used as antiviral compounds or antibiotics. During the hydrothiolation of 1,4-TMS-substituted buta-1,3-diyne **180c**, the desilylation reaction occurred (Scheme 191).^{405–408} Changing the reaction conditions, by applying a different solvent (DMSO), led to the formation of biologically active thiophenes **1051a–f** with moderate isolated yields (51–66%), instead of thio-substituted buta-1,3-dienes **1047** when EtOH or DMF were used.^{406,409} The 1,2-addition product of arylmethanethiol **1046a** or **1049a–c** led to the corresponding enyne **1050**. The thiophene was formed by the cyclisation of enenyne thiol **1050**, which occurs from the nucleophilic attack of benzyl anion on C_{sp} bond in the second alkynyl group of **1050** (Scheme 192).^{410,411}

The synthesis of thiophenes and other cyclic compounds from diynes was briefly reviewed by Maretina and Trofimov.²⁹⁵ The paper presented the procedures that were published mostly in 1960–1980 and are focused on the addition of sulfide ions to conjugated diynes yielding thiols. Very good results were obtained in the case of the formation of thiols when Na₂S **1052** was used as a reagent in KOH/DMSO. When a quantitative amount of Na₂S **1052** and KOH was applied thiophene **1053** was formed from buta-1,3-diyne with excellent yield up to 99%. The process occurred *via* hydrothiolation of diyne followed by cyclisation. Nonhydroxylic polar solvents *e.g.*, DMSO or *N*-methylpyrrolidone should be used in this transformation, because these solvents did not decrease the activity of anions by their solvation.²⁹⁵ By replacing KOH with TBAOH **1056** and benzylthiol







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1046a with butyl analog **1055** it was possible to shorten the reaction time to just 5 minutes and to increase the product yields and selectivity. (*Z*)-Thiobutenynes **1057a-k** were obtained exclusively for symmetrical and unsymmetrical diynes **1a**, **1d**, **13a**, **208b**, **258g**, **617**, **c**, **1028**, **1054a-b** with sterically different substituents in positions C_1 and C_4 . Hydrothiolation using the reductive system *n*- C_4H_9SH **1055**/TBAOH **1056** is more efficient because it is a stronger base which, due to its phase-transfer ability, increases the solubility of reagents in the organic phase and accelerates the formation of the butylthiolate anion (Scheme 193).⁴¹² In the presence of iodine **418**, the obtained (*Z*)-organylthioenynes underwent electrophilic cyclisation towards 3-iodothiophenes **1058**. The reaction was tested using **1.0** equiv. of **1057c** and **1.1** equiv. of **I_2 418** (Scheme 194).

Thiophenes **1061a–f** were obtained also from haloalkynes **1018** and **1059** from a Glaser reaction to conjugated 1,3-diynes **1a–c**, **258g**, **258n**, and **1060** followed by the hydrothiolation to sulfanyl substituted enynes, which further cyclises to thiophenes possessing different aryl or heteroaryl groups in positions 2- and 5- **1061a–f**. The reactions were catalysed by a CuI **519**/1,10-phen system and proceeded with high products yields. As a hydrothiolation agent, Na₂S·9H₂O **1052** was used (Scheme 195). The mechanism of this transformation was previously described for the analgous hydration process (Scheme 187).³⁸⁸

The synthesis of 3-halosubstituted thiophenes 1066 from simple aryl or alkyl-functionalised conjugated buta-1,3-diynes 1a, 1d, 13a, 27c, 65b, 208b, 258g, 258o, 617, 842b-c, 1028, and 1054a-b was described by Kesharwani et al. They proposed a two-step procedure yielding 3-chloro, 3-bromo and 3-iodothiophenes 1066 based on hydrothiolation reaction of 1,4-diaryl or 1,4-dialkyl-substituted divnes with methyl disulfide 1062 in the presence of NaBH₄ 212 as a hydrogen source and, electrophilic cyclisation of the obtained sulfanyl modified enynes 1063 with natrium halides 1064a-c (NaCl, NaBr, NaI) in the presence of CuSO₄·5H₂O 1065 (Scheme 196).⁴¹³ This methodology has a positive impact on the environment, because it uses the green solvent ethanol and simple inorganic salts. In many cases it also gave better results than typical methods used for the preparation of halothiophenes (Scheme 196).^{413–415} The mechanism of the cyclisation proposed the formation of CuCl₂ in the first step from CuSO₄ 1065 and NaCl 1064a, which can easily coordinate to the $C \equiv C$ triple bond in the enyne to 1067. Nucleophilic attack of sulfur provided intermediate 1068 that eliminates the methyl group attached to the sulfur atom via an S_N2 substitution reaction by the chloride anion yielding intermediate 1069. Reductive elimination furnished the desired



Scheme 195 Synthesis of 2,5-disubstituted thiophenes 1061 from haloalkynes in Glaser coupling/hydrothiolation/cyclisation reactions catalysed by Cul 519/1,10-phen.



Scheme 196 Synthesis of 3-halothiophenes **1066** in hydrothiolation/electrophilic cyclisation reactions.

halothiophene **1066a**, while the Cu(0) is oxidised to CuCl **55** by CuCl₂ (Scheme 197). In the case of the application of NaBr **1064b** and NaI **1064c**, CuBr₂ **52** and CuI₂ can easily release I₂ **418** and Br₂, and by applying these as electrophiles forms of bromo- and iodothiophenes.⁴¹³ 3,4-Dichloro-substituted thiophenes **1071** can be formed in the reaction of 1,4-diarylsubstituted buta-1,3-diynes with sulfur chloride **1070**. Products **1071** were obtained with 18–80% yields.^{295,416} Excellent yields and selectivities of (*Z*)-thioenynes **1074a-l** were obtained when disulfides **1073** (PhSSPh **1073a**, BuSSBu **1073b**) were used as reagents. Oganylthiolate anions were generated *in situ* with NaBH₄ **212**. The application of disulfide **1073a-b** may constitute an

alternative towards the use of toxic and bad-smelling thiols (Scheme 198).⁴¹⁷

A sustainable and clean method for obtaining thiobutenynes **1074/1075** was carried out in the presence of KF/Al₂O₃ as a catalyst, using glycerol or poly(ethylene glycol) ($M_w = 400$, PEG400) as a green solvent. Applying this system, it was possible to decrease the amount of the catalyst employed, and the generation of inorganic products is reduced to a minimum. The products were extracted in hexane/ethyl acetate and KF/Al₂O₃ was directly used in following cycles. The reaction was effective for various diynes substituted in the terminal position with electron-withdrawing or electron-donating



 $\label{eq:scheme 197} Scheme 197 \mbox{ Proposed mechanism for chlorocyclisation of 1-methyl-sulphanyl-1,4-diphenyl-buta-1-en-3-yne 1063a catalysed by Cu(1).$

groups. The reaction was optimised for the use of an equimolar ratio of reagents. The best results were accomplished when 90 °C was used and the reaction was carried out for 6 hours. A lower temperature led to lower yields, while higher temperatures reduced the selectivity. Under the optimal conditions' product **1074** with (*Z*)-geometry was obtained in excess, in the ratio 90:10 to 100:0 depending on the reagent structure. The yields of the (*Z*)-1,4-diphenyl-2-(phenylthio)but-1-en-3-yne **1074a** in the three repetitive batches reached 93%, 89%, 80% respectively (64%, 55%, 48% isolated yield).⁴¹⁸

Perin *et al.* proved that the addition of phenyldisulfide **1073a** to **1,4-**diphenylbuta-**1,3-**diyne **1a** using the same conditions (NaBH₄ **212**, PEG400, 30 $^{\circ}$ C) may be accelerated applying

microwave irradiation as a heating source. It was possible to reduce the reaction time to 85 minutes from 24 hours under traditional conditions with a slightly better yield of **1074a** (96% *vs.* 82%). Moreover, by increasing the temperature to 90 °C, 1,4-diphenyl-1,4-di(phenylthio)buta-1,3-diene **1076** was selectively formed in good yield (65% *vs.* 69%).⁴¹⁹

The hydrothiolation of 1,3-butadiynes 1a, 208b, 655c, 1028, 1045. 1072a, and 1072c were carried also using various diarvl disulfides 1073a or 1077a-e, sodium hydroxymethanesulfinate 1078 (rongalite 1078, HOCH₂SO₂Na), and potassium carbonate. Rongalite 1078 was applied as a reducing agent cleaving the bond of disulfide. When disulfide was used in 0.5 equiv. to the divne, (Z)-1-sulfanyl-but-1-en-3-ynes 1079a-o were obtained with isolated yields in the range 45-86%. Increasing the temperature to 70 °C and an equimolar ratio of divne and disulfide, it was possible to obtain a mixture monothiolation 1079a-o and bisthiolation 1080 products with moderate yields. Moreover, the introduction of two different arylthiol groups to the product was possible by subsequent hydrothiolation of divne with two others disulfides. The reaction did not occur for benzyl and alkyl-substituted disulfides (Scheme 199). The mechanism of this transformation started by the decomposition of rongalite 1078 to formaldehyde 1081 and HSO₂⁻ 1082 in the presence of the base. Next, the single-electron transfer to disulfide 1073a leads to anionic 1084 and radical species 1085. The radical thiolate 1085 is then reduced to its ionic form 1084 by another single electron transfer from radical HSO2[•]. 1083. Addition of thiolate 1084 to diyne 1a followed by the protonation of the intermediate 1086 yields the desired product 1079a (Scheme 200).

Moreover, bishydrothiolation was also carried out in a sequence of one-pot reactions. Sonogashira coupling of



Scheme 198 Synthesis of (Z)-1-organothiobut-1-en-3-ynes 1074a–l by the hydrothiolation of buta-1,3-diynes using disulfides 1073a–b

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1,4-bis(trimethylsilyl)buta-1,3-diyne **180c** with aryl halides **1087** followed by bishydrothilation of the obtained 1,4-bisarylbuta-1,3-diyne with various thiols **1046a** and **1089a** to (*Z*,*Z*)-1,4-diarylbuta-1,3-dienes **1090**. The formation of the new C–C bonds was catalysed by Pd(OAc)₂ (1 mol%) **861** and Cu(xantphos)I (1 mol%) **1088**, while double hydrothiolation was promoted by basic Cs₂CO₃. The whole process occurred with moderate or good yields, with high regio- and stereoselectivity (Scheme 201).⁴²⁰ The mechanism of this transformation started with the

generation of the sulfur anion from thiol **1046a** or **1089** in the presence of base a (BH). The hydrothiolation step then occurs with the formation of intermediate **1091**. The (*Z*)-isomer **1093** is formed in the presence of thiol **1089a** or base according to the protonation step. Subsequent hydrothiolation of obtained enyne **1093** furnished (1*Z*,4*Z*)-bissulfanylbuta-1,3-diene **1090a** (Scheme 202).⁴²⁰

The addition of aminothiols **1094a–e** to buta-1,3-diyne **641** was carried out in ammonia **632**, which was used as a solvent








and base at the same time. HS⁻ anions are 280 times more reactive than NH_2^- , therefore the addition of aminothiols **1094a–e** to the C=C bonds occurred from the S-side. The enyne sulfides **1095a–e** were obtained in 78–98% yield (Scheme 203).⁴²¹

12. Hydroselenation of 1,3-diynes

Alkenyl selenides are an important class of organoselenium compounds, which have broad applications in organic chemistry

leading to a vast spectrum of valuable products.^{422–426} These molecules can be prepared by various synthetic pathways however, the most frequently employed method for their preparation is the hydroselenation of the $C \equiv C$ bond by nucleophilic organoselenolate anions. The synthesis and application of organoselenium compounds, especially selenophenes, has been summarised in many books and reviews,^{422,423,425,427–434} nevertheless the hydroselenation of diynes have not been comprehensively reviewed.



Scheme 202 Mechanism of one-pot Sonogashira/hydrothiolation reactions of 1,4-bis(trimethylsilyl)buta-1,3-diyne **180c** catalysed by Pd(OAc)₂ **861**/Cu(xanthphos)l **1088**/Cs₂CO₃ system.



The first example addition of a Se–H bond to diynes was reported by Taylor *et al.* in 1968.⁴³⁵ The addition of H₂Se **1097** to symmetrical and unsymmetrical **1**,3-diynes **1a**, **208c**, **1045**, **1072a**, **1096a–b** was catalysed by the Ag⁺ cations and led to 2,5-disubstituted selenophenes **1098a–f** in good yields (Scheme 204).

Dabdoub *et al.* developed an alternative and efficient synthetic protocol employing the phenylselenolate anion which was generated *in situ* by the reaction of Ph₂Se₂ **1099** with sodium borohydride **212** in ethanol, instead of using toxic hydrogen selenide. Hydroselenation of 1,4-substituted-1,3-butadiynes **1a**, **613a**, **842b–c**, **1072a–b**, **1072d** occurred with excellent regio-, stereo- and chemoselectivity smoothly leading

to (*Z*)-1-phenylseleno-1,4-diorganyl-1-buten-3-ynes **1100a-h** in high yields. However, reacting 2-hydroxy-2-methyl-3,5-dode-cadiyne **1072a** with Ph_2Se_2 **1099** and $NaBH_4$ **212** in ethanol under reflux, gave a mixture of regioisomers (**1100e**/1**100f** = 58/42) which was confirmed by ¹H NOESY experiments (Scheme 205).⁴³⁶

A similar strategy was applied by Zeni *et al.* who used various diorganodiselenides **1112a–e** in the preparation of (*Z*)-selenoenynes **1113a–o** through the hydroselenation of symmetrical and unsymmetrical 1,3-diynes **1a**, **1c**, **13a**, **37s**, **842b–c**, **1028**, **1045**, **1072a**, and **1111**. The obtained products were further cyclised with different electrophiles such as I₂ **418**, ICl **1114a**, PhSeBr **1114b**, or PhSeCl **1114c** to 3-substituted selenophenes



1115a–i in good yields. The electrophilic cyclisation did not occur for (*Z*)-1-(phenylseleno)-1,4-diphenyl-but-1-en-3-yne **1113f** even when using harsh reaction conditions. The authors also presented the versatility of these compounds for many transformations such as Sonogashira coupling, halogen-metal exchange reaction, or Ullmann-type C–O bond forming reactions (Scheme 206).⁴³⁷

An analogous protocol was applied by Kesharwan et al. who used sodium halides 1064a-c as a source of electrophilic halogens for the synthesis of halogenated selenophenes 1115a, 1116. The hydroselenation of 1,4-diphenyl-1,3-diyne 1a by Me₂Se₂ 1112a in the presence of NaBH₄ 212 in ethanol gave for (Z)-1-(benzylseleno)-1,4-diphenyl-but-1-en-3-yne 1113a in 60% yield which was transformed to halogenated selenophenes 1115a and 1116 (Scheme 207a). The protocol was also suitable for the preparation of halogenated thiophenes (Scheme 196).413 The replacement of electrophiles to a base such as t-BuOK in the cyclisation step is also efficient for the synthesis of selenophenes 1119 through selenoenynes 1113e and 1118 intermediates. The hydroselenation of 1,3-diynes 1a, 1c-d, 27a, 258a, or 280a with Bn₂Se₂ 1112e in the presence of NaBH₄ 212 in ethanol was limited to its symmetrical derivatives, and the reaction yields were significantly lower for bulky divnes (Scheme 207b). The authors suggested that 3-benzyl-substituted selenophenes 1119 could be highly promising building blocks for the preparation of polysubstituted selenophenes.438

The hydroselenation of diynes was also adopted to synthesise more complex structures such as tetrapyrrolic macrocycles. Chauhan *et al.* presented the preparation of selenium core-modified porphyrinogens based on the hydroselenation of diynediols **1072b**, **1121a–b** with *in situ* generated sodium selenide **1122** in the presence of MeOH and CH₃COOAg in the first step. The obtained selephones **1123** were further transformed to selenophene dipyrranes and used in 3 + 1 condensation reactions with the corresponding diols in the presence of boron trifluoride **565**. It was proved by UV-Vis, fluorescence and ¹H NMR spectroscopy that the selenium core-modified porphyrinogens have a coordination ability to detect of Hg²⁺

cations.⁴³⁹ The same group reported the preparation of porphomethenes **1128**, porphodimethenes **1127**, and porphotrimethenes **1129** using the same methodology including diynediol hydroselenation followed by 3 + 1 condensation of the selenatripyrranes **1125** with selenophene-2,5-diols **1123** and subsequent oxidation with 2,3-dichloro-5,6-dicyanobenzo-quinone **1126** (DDQ). Similar to porphyrinogens, the obtained macrocycles showed the binding affinity with Hg²⁺ anions (Scheme 208).⁴⁴⁰

Lopes *et al.* described the application of deep eutectic solvents (DES), commonly considered as the third generation of ionic liquids, in the preparation of vinyl selenides. Although the report concerns mainly the synthesis of (E)-1,2-bisorganylseleno alkenes through the hydroselenation of alkynes, the utilisation of symmetrical buta-1,3-divnes was also presented. Diphenyl selenide 1099 reacted with 1,4-diphenyl-1,3-diyne 1a or hexa-2,4-diyne-1,6-diol 1045 in the presence of NaBH₄ 212 in a choline chloride/urea (1/2) mixture at 90 °C to give corresponding (Z)-selenoenynes 1100g and 1130 with excellent selectivity and high yields (Scheme 209a).⁴⁴¹ A similar strategy based on the application of green solvents was used by Lara et al. who applied a poly(ethylene glycol) (PEG 400) as an alternative for DES. Depending on the reaction temperature, the (Z)-selenoenynes 1100g-h and 1132a-b or (Z,Z)-1,4-bisselenobuta-1,3dienes 1133a-c were obtained in the reaction of diorganodiselenides 1099, 1112c, and 1131 with the symmetrical 1,3-diynes 1a, 1045, 1072a in the presence of NaBH₄ 212. The process is highly stereoselective exclusively leading to the corresponding products in high yields. The reaction time was reduced from 24 to 1.25 h by the application of microwave radiation (MW) as a heating method (Scheme 209b).419

An interesting protocol for the synthesis of (*Z*)-1-(organoselanyl or sulfanyl)enynes was developed by Venkateswarlu *et al.* who utilised sodium hydroxymethanesulfinate (rongalite) **1078** as a reducing agent instead of commonly used NaBH₄ **212**. The hydroselenation of **1,3-**diynes **1a, 208b, 655c, 1028, 1045, 1072a-b** was carried out in the presence of potassium carbonate in a DMF-H₂O (20:1) mixture and under mild reaction





Scheme 207 Synthesis of 3-substituted selenophenes 1115a, 1116, and 1119 through (a) electrophilic cyclisation or (b) carbocyclisation of selenoenynes.



Scheme 208 Selected examples for the synthesis of porphomethenes 1128 and porphodimethenes 1127.



conditions leading exclusively to (*Z*)-isomers **1100g–h**, **1130** and **1134a–d**. It is worth noting that the discussed procedure was suitable for symmetrical and unsymmetrical 1,3-diynes, however, it failed when an aliphatic diacetylene such as hexadeca-7,9-diyne **208b** was applied. The low reactivity of **208b** in this

transformation can be explained in the same manner as hydrothiolation,⁴⁴² by weak stabilisation of transition state which is dependent on steric and electronic factors. Depending on reaction conditions it was possible to obtain mono- or bis-hydroselenation products. The hydroselenation of 1,3-diynes



Scheme 210 Hydroselenation of symmetrical and unsymmetrical substituted 1,3-diynes **1a**, **208b**, **655c**, **1028**, **1045**, **1072a**–**b** with diphenyl diselenide **1099** in a presence of rongalite **1078** as a reducing agent.



performed at a lower temperature (50 °C) and with 0.5 eq. of Ph_2Se_2 **1099** furnished (*Z*)-1-(organoselanyl)enynes **1100g–h**, **1130** and **1134a–d**, whereas application of 1.0 equiv. of Ph_2Se_2 **1099** and a higher temperature (70 °C) yielded a mixture of mono- and bishydroselenation products (Scheme 210). The presented protocol is also adequate for sulfanyl derivatives (Scheme 199).⁴⁴³ The mechanistic studies for the reaction were performed based on hydrothiolation of diynes (Scheme 200), however, it also can be extended to hydroselenation.⁴⁴⁴ It involves the reduction of Ph_2Se_2

1099 with the generation of the PhSe⁻ anion **1137** followed by hydroselenation of the 1,3-diyne. In the initial step, the rongalite **1078** is decomposed in the presence of K_2CO_3 to formaldehyde **1081** and $HSO_2^-Na^+$ **1082**. Single electron transfer (SET) gives anion **1137** and radical **1136**. Another SET reduces **1136** to **1137**. The *trans*-addition of the benzeneselenolate anion **1137** to the 1,3-diyne gives intermediate **1138** which is protonated to yields (*Z*)-1-(organoselanyl)enynes **1100g-h**, **1130** and **1134a-d** (Scheme **211**).⁴⁴³ The hydroselenation of 1,3-diynes was utilised by Męcik *et al.* for the synthesis of di(selenophen-3-yl)diselenides **1143a–e** and 3-methylene-3*H*-1,2-diselenoles **1144a–b**. These uncommon selenium heterocycles were only formed when 1-amino-4-aryl-buta-1,3-diynes or 1-amino-4-ester-buta-1,3-diynes **1141** (synthesised from 1-bromobutadiynes **1139** and secondary amine **1140**) were used. The reaction of 1-aminobutadiynes **1141** with generated *in situ* sodium selenide led to desired products **1143a–e** and **1144a–b**, instead of excepted selenophenes, in moderate to high yields.

Simple diaryl or dialkyl-1,3-butadiynes led to classical selenophenes or did not react at all thus, the presence of amine group in diyne structure was crucial for the synthesis of di(selenophen-3-yl)diselenides **1143a–e** and 3-methylene-3*H*-1,2-diselenoles **1144a–b**. It is worth noting that this protocol could be also adopted in a one-step strategy without the isolation of 1-aminobutadiynes **1141**, by the addition of sodium selenide solution to the reaction mixture directly after the amination step (Scheme 212). The authors proposed the mechanism of this transformation, which started from the hydroselenation of 1,3-butadiyne by the nucleophilic attack of generated *in situ* SeH⁻ anion **1145** to C=C bond.

Subsequently, the bisselenide **1147** is formed and transformed to selenirenium ion **1148**, which undergos nucleophilic attack to a carbon atom (for aryl-substituted diynes) **1149** and its dimerisation with the generation of di(selenophen-3-yl)diselenides **1143a–e**. For ester-substituted diynes occurs an internal nucleophilic attack of Se⁻ to Se⁺ **1151** and further rearrangement to 3-methylene-3*H*-1,2-diselenoles **1144a–b** (Scheme 213).⁴⁴⁵

13. Hydrotelluration of 1,3-diynes

Unsaturated organotellurium compounds, especially vinylic tellurides have found numerous applications in organic synthesis due to their high reactivity, tolerance towards many functional groups and the possibility for carbon–carbon bond formation. This has been covered in previous reviews.^{431,433,434,446–448} This versatile class of tellurium compounds is also an important intermediate in the synthesis of tellurophenes which have applications in material chemistry^{430,447,449–455} and biological chemistry.^{433,456} Among many synthetic strategies towards unsaturated organotellurium compounds, the hydrotelluration



Scheme 212 Synthesis of di(selenophen-3-yl)diselenides 1143 and diselenoles 1144 from 1-bromobutadiynes 1139





Scheme 214 Synthesis of π -conjugated 2,5-substituted tellurophene compounds 1154 and 1155.

of diynes is a highly efficient and stereoselective method for enynyl tellurides which are useful building blocks in modern chemistry. Since the first synthesis of tellurophene by the interaction of 1,3-butadiyne **1a** and **1152** with Na₂Te **1153** in methanol developed by Mack's in 1966,⁴⁵⁷ the several papers describing hydrotelluration of diynes appeared^{458–468} which was also covered by Detty⁴³¹ and Zeni.^{446,469} In this chapter the recent development of functionalisation of diynes by tellurium compounds will be presented. Seferos *et al.* reported π -conjugated 2,5-substituted tellurophene **1154** compounds which were synthesised *via* ringclosing reactions of 1,4-substituted butadiynes **1a** and **1152** in the presence of Na₂Te **1153** and a protic solvent (Scheme 214). This synthetic procedure avoids harsh reaction conditions and degradation of the tellurophene ring. The oxidation of tellurophene **1154** through Br₂ addition to **1155** changed the measured optical absorption spectrum and oxidation potential which was confirmed by absorption spectroscopy and DFT calculations. The authors suggested that this class of compounds might have potential applications as semiconducting materials or as transition metal-free catalysts for energy storage reactions.^{453,470,471}

A similar approach was applied by Chauhan *et al.* who synthesised a series of calixpyrroles and calixphyrins by the interaction of diynediols **1045**, **1072a**, **1121a–b**, and AgOAc in MeOH with an aqueous solution of Na₂Te **1153**. Subsequent 3 + 1 condensation of telluratdipyrranes **1157** or telluratripyrranes **1158** with corresponding tellurophene-2,5-diols **1156** in the presence of BF₃–etherate **565** gave desired products (Scheme 215). The obtained compounds had a binding affinity with Hg²⁺ cations which was confirmed by spectroscopy studies. A described



Scheme 215 Synthesis of core-modified porphyrinogens 1159.



synthetic protocol could also be applied for selenophenoediols (see Section 11.2). $^{\rm 439,440}$

Application of environmentally-friendly poly(ethylene glycol, $M_w = 400$) in the selective synthesis of (*Z*)-telluroenynes **1161** and (*Z*,*Z*)-1,4-bis-tellurobuta-1,3-dienes **1162** in the reaction of symmetrical diynes with diphenyl telluride **1160** and NaBH₄ **212** as a reducing agent was reported by Perin *et al.* The process was found to be temperature-dependent. When the reaction was carried out at 30 °C, the (*Z*)-telluroenynes **1162** were obtained with excellent selectivity while higher temperatures (90 °C) led to (*Z*,*Z*)-1,4-bis-tellurobuta-1,3-diene **1162** (Scheme 216). The use of microwave radiation as an alternative heating source furnished desired products in a few minutes instead of several hours. The protocol is also suitable for selenium and sulfur derivatives.⁴¹⁹

Deep eutectic solvents (DES) composed with the choline chloride (ChCl) and urea mixture (1:2) could be applied as

another green solvent in the synthesis of organoseleno alkenes and mono-chalcogenated (*Z*)-alkenynes. The hydrotelluration of **1a** with diphenyl telluride **1160** in the presence of NaBH₄ **212** led to the (*Z*)-1-phenyltelluro-1,4-diphenyl-but-1-en-3-yne **1161a** with excellent regio- and stereoselectivity and moderate isolated yield (42%) (Scheme 217).⁴⁴¹

Męcik *et al.* reported the synthesis of tellurophenes **1164a–d** by the reaction of 1-aminobutadiynes **1141** with sodium telluride **1153** which was generated *in situ* from Te **1163** and NaBH₄ **212**. The 1-aminobutadiynes **1141** were prepared from 1-bromobutadiynes **1139** and used without purification step leading to 2-aminotellurophenes **1164a–d** in good yields. Intriguingly, the application of sodium selenide **1122** gave di(selenophen-3-yl)-diselenides **1143** and methylene-3*H*-1,2-diselenoles **1144** instead of simple selenophenes (Scheme 212). The authors suggested that it might be caused by the lower stability of Te–Te bond compared with the Se–Se bond (Scheme 218).⁴⁴⁵

14. Conclusions and outlook

Conjugated and separated divnes constitute a special class of compounds, which due to their structural and electronic versatility that can be tuned by the presence of various functional groups attached to the $C \equiv C$ bonds, as well as by the spacer between both $C \equiv C$ bonds, create a "chemical mine" for developing fine organometallic and organic chemicals. The combination of these compounds with hydroelementation reagents, chosen from main group elements, permits an incredibly diverse array of products (enynes, dienes, allenes, cyclic compounds, heterocycles, polymers) to be obtained. Due to the presence of the unsaturated C-C bonds, as well as main group elements in their structures, these are extremely useful building blocks in the synthesis of tailor-made materials, pharmaceuticals, natural compounds analogs, and structurally advanced organic molecules. The review presents the library of the reactions, reagents, products, and catalysts for the hydroelementation of conjugated and separated divnes and can be used as a guidebook for planning the synthesis of advanced compounds via hydroelementation processes. The problems with the selective activation of the one or two $C \equiv C$ bonds, possible overreduction, and the stereo- and regioselectivity of hydroelementation processes are the biggest challenges that need to be overcome during the reduction of diynes. This can be achieved by the proper choice of reagents (steric and electronic properties have a significant influence on the process selectivity) catalyst, and reaction conditions. The number of examples that produce only one product and one isomer is however limited. Therefore, there is still a lot of scope for developing catalytic systems that might be highly selective, active, and stable as well as being straightforward from a synthetic perspective. Bearing in mind that the hydroelementation reaction is a 100% atom economic process, searching for highly effective and selective methods, which might be applied using an equimolar ratio of reagents is of prior importance. The products obtained in the hydroelementation reactions of conjugated and separated diynes can be used as important synthons in organic synthesis. Several demetallation, coupling, and addition reactions were presented in this review to demonstrate the power of the products obtained in the hydrometallation of diynes. In the future, improvements in the catalyst effectiveness and availability, selectivity, and productivity need to be undertaken to make the hydroelementation process straightforward for the formation of different products. Control of the process regio- and stereoselectivity is the biggest task for all chemists, which are focused on the synthesis of fine chemicals.

Abbreviations

acac	Acetylacetone
ACCN	((1,1-Azobis(cyclohexane-1-carbonitrile)))
BINAP	2,2'-Bis(diphenylphosphino)-1,1'-
	binaphthalene
bmin	1-Butyl-3-methylimidazolium

Bn
BPPI

Bn	Benzyl
BPPM	4-(Diphenylphosphino)-2-
	[di(nhenylphosphino)methyl]pyrrolidine
CAAC	[ulphenyiphospinio]neuryipyironume
CAN	Ammonium cerium(w) nitrate
CAN	Annonium certum(iv) intrate
CDZ	a f C alegate ling
cod	1,5-Cyclooctadiene
т-СРВА	3-Chiorobenzene-1-carboperoxoic acid
Cy	Cyclohexyl
<i>t</i> -BuPNP	2,6-Bis(di(<i>tert</i> -butyl)phosphinomethyl)pyridine
dba	Dibenzylideneacetone
DBN	1,5-Diazabicyclo[4.3.0]non-5-ene
DBU	1,8-Diazabicyclo[5.4.0]undec-7-ene
DCC	Dicyclohexylcarbodiimide
DDSQ	Double-decker-shaped silsesquioxane
DDQ	2,3-Dichloro-5,6-dicyanobenzoquinone
DES	Deep eutectic solvent
DIBAH	Di(iso-butyl)aluminum hydride
Dipp	2,6-Di(iso-propyl)phenyl
DMAP	4-(Dimethylamino)pyridine
dmpm	5,5-Dimethyldipyrrolylmethane
dpma	<i>N,N</i> -Di(pyrrolyl-alpha-methyl)- <i>N</i> -methylamine
DP	Polymerisation degree
dppb	1,4-Bis(diphenylphosphino)butane
dppben	1,2-Bis(diphenylphosphino)benzene
dppe	Ethylenebis(diphenylphosphine)
dppf	1,1'-Ferrocenediyl-bis(diphenylphosphine)
dppp	1,3-Bis(diphenylphosphino)propane
dvs	1.1.3.3-Tetramethyl-1.3-divinyldisiloxane
DIEA	<i>N.N</i> -Di-iso-propylethylamine
DIOP	2. 3-O-Iso-propylidene-2. 3-dihydroxy-1. 4-
2101	his(diphenylphosphino)butane
F-TEDA	1-Chloromethyl-4-fluoro-1 4-
I ILDII	diazoniabicyclo[2,2,2]octane
	his(tetrafluoroborate)
hfa	Hevafluoroacetone
hmpa	Hevamethylphosphoramide
шпра	Ionia liquid
	Di-ico-pinocampheulhorane
IPr [*] OMo	1.2 Picimidazol 2 vlideno
	Lithium diisannomdamida
	1. 4/ Motherlanghis[2. C
MBPH	4,4 -Methylenebis[2,6-
	bis(hydroxymentyl)jphenol
MS	Methanesulfonyl
MS	Molecular sieves
$M_{ m w}$	Molecular weight
MW	Microwave radiation
NCS	<i>N</i> -Chlorosuccinimide
NHC	N-Heterocyclic carbene
NIS	<i>N</i> -Iodosuccinimide
NMDPP	Neomenthyldiphenylphosphine
Norphos	2,3-Bis(diphenylphosphino)bicyclo[2.2.1]hept-
	5-ene
NTf_2	Bis(trifluoromethane)sulfonimide
OTf	Trifluoromethanesulfonate
PEG	Poly(ethylene glycol)

Review Article

nin

Dinacol

Pm	Timueon
PINBOP	2-Iso-propoxy-4,4,5,5-tetramethyl-1,3,2-
	dioxaborolan
pivOH	Pivalic acid
PNP	1,3-Bis(di- <i>tert</i> -butyl-phosphinomethyl)pyridine
POP-BZ	1-Benzyl-3,4-
	bis((diphenylphosphaneyl)oxy)pyrrolidine
PPM	4-(Diphenylphosphaneyl)-2-
	((diphenylphosphaneyl)methyl)pyrrolidine
РТМА	(5-Propylsulfonyloxyimino-5H-thiophen-2-
	ylidene)-2(methylphenyl)acetonitrile
pv	Pivaldehyde
PyrPhos	3,4-Bis-diphenylphosphino-pyrrolidine
QUINAP	1-(2-Diphenylphosphino-1-
	naphthyl)isoquinoline
$scCO_2$	Supercritical CO ₂
SET	Single electron transfer
TBAF	Tetrabutylammonium fluoride
TBHN	Di-tert-butyl hyponitrite
TBDMS	<i>Tert</i> -butyldimethylsilyl
TC	Thiophene 2-carboxylate
TDT	Tert-dodecanethiol
THP	Tetrahydropyranyl
TIPS	Tri(iso-propyl)silyl
TMEDA	<i>N,N,N',N'</i> -Tetramethylethylenediamine
TMS	Trimethylsilyl
Tos	Toluenesulfonyl
TP	Tris(1-pyrazolyl)borate
p-TSA	para-Toluenesulfonic acid
xantphos	4,5-Bis(diphenylphosphino)-9,9-
	dimethylxanthene

Conflicts of interest

There are no conflicts of interest to declare.

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