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Recent synthetic strategies for the functionalization of fused bicyclic heteroaromatics using organo-Li, -Mg and -Zn reagents†

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This review highlights the use of functionalized organo-Li, -Mg and -Zn reagents for the construction and selective functionalization of 5- and 6-membered fused bicyclic heteroaromatics. Special attention is given to the discussion of advanced syntheses for the preparation of highly functionalized heteroaromatic scaffolds, including quinolines, naphthyridines, indoles, benzofurans, benzothiophenes, benzoxazoles, benzothiazoles, benzopyrimidines, anthranils, thienothiophenes, purine coumarins, chromones, quinolones and phthalazines and their fused heterocyclic derivatives. The organometallic reagents used for the desired functionalizations of these scaffolds are generally prepared in situ using the following methods: (i) through directed selective metalation reactions (DoM), (ii) by means of halogen/metal exchange reactions, (iii) through oxidative metal insertions (Li, Mg, Zn), and (iv) by transmetalation reactions (organo-Li and Mg transmetalations with ZnCl₂ or ZnO(Piv)₂). The resulting reactive organometallic reagents allow a wide range of C-C, C-N and C-X cross-coupling reactions with different electrophiles, employing in particular Kumada or Negishi protocols among other transition metal (Pd, Ni, Co, Cu, Cr, Fe, etc.)-catalyzed processes. In addition, key developments concerning selective metalation techniques will be presented, which rely on the use of RLi, LDA and TMP metal bases. These methods are now widely employed in organic synthetic chemistry and have proven to be particularly valuable for drug development programs in the pharmaceutical industry. New and improved protocols have resulted in many Li, Mg and Zn organyls now being compatible with functionalized aryl, heteroaryl, alkenyl, alkynyl and alkyl compounds even in the presence of labile functional groups, making these reagents well-suited for $C(sp^2)-C(sp^2)$, $C(sp^2)-C(sp)$ and $C(sp^2)-C(sp^3)$ cross-coupling reactions with fused heteroaryl halides. In addition, the use of some transition metal-catalyzed processes occasionally allows a reversed role of the reactants in cross-coupling reactions, providing alternative synthetic routes for the preparation of fused heteroaromatic-based bioactive drugs and natural products. In line with this, this article points to novel methods for the functionalization of bicyclic heteroaromatic scaffolds by organometallic reagents that have been published in the period 2010-2023.

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1 Introduction

Organometallic reagents (R-M; M = Li, Mg, Zn; R = alkyl, alkenyl, alkynyl, aryl, heteroaryl) are widely used in synthetic organic chemistry and various areas of catalysis. 1-10

These reagents play an important role in drug discovery and development, in the synthesis of natural products and bioactive compounds, 11-17 as well as in programs dedicated to medicinal and materials chemistry. 18-23 A variety of synthetic methods have been developed for the preparation of organometallic reagents such as organolithium, organomagnesium, and organozinc reagents and their derived cognates. Their potential use

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[†] Dedicated to Prof. Bert Maes

in catalytic applications has been investigated by several leading research groups such as those of Snieckus, 24,25 Knochel, ^{26–30} Harutyunyan, ^{31–35} Aggarwal, ^{36–40} Feringa, ^{41–44} Marek, ^{45–50} Mongin, ^{51–53} Kürti, ^{54–56} Maes, ^{57–59} Buchwald, ⁶⁰ Smith, 61 Li, 62 and others. 63-65 All of these methods most likely utilize one of the following procedures depicted in Fig. 1: (i) directed oxidative metal insertions in the presence of LiCl salts or InCl₃-Lewis acid (Mg, Zn, Al, Mn, In), 66,67 (ii) halogen-metal exchange reactions (X/M, X = Br, I; M = Li, Mg, Zn, Sm, La, Mn) using Turbo Grignard reagents (i-PrMgCl·LiCl) or other organometallic reagents, 68-72 and (iii) chemo- and regioselective

direct metalations using Knochel-Hauser bases (TMPLi, -Mg and -Zn).73-75

In the last two decades, organolithium, magnesium and zinc reagents derived from TMP-H (2,2,6,6-tetramethylpiperidinyl) have played an important role in organic synthesis. These organometallic reagents have proven to be very effective as they facilitate the chemo- and regioselective metalation of various aromatic and non-aromatic heterocyclic scaffolds. 73-75 The various alkylamine bases, including TMP-M, R¹R²N-M (M = Li, Mg, Zn; R^1 & R^2 = alkyl) and LDA, are synthesized from commercially available secondary amine sources such as



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Vishal S. Dodke received an MSc degree in Organic Chemistry from Savitribai Phule Pune University, India in 2010. He was a Doctoral Student of Chemistry at Jacobs University Bremen, Germany where he focused on the Subsequently, organocatalysis. he worked with the private companies and earned experience in organic synthesis. He joined as a PhD student under the supervision of Dr Rambabu Dandela at Institute of

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Marappan Pradeep Kumar

Marappan Pradeep Kumar, originally from Tiruppur, Tamil Nadu, holds a BSc in Chemistry Bharathiar University (2018-2021) and an MSc in Chemical Sciences from Pondicherry University (2021-2023). He gained research experience through a summer research fellowship at IIT Madras under the guidance of Prof. G. Sekar. Currently, he serves as a Project Assistant in the OMC research group at the National Institute

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Hatice Seher Korkmaz

Hatice Seher Korkmaz was born 1996 in İzmir (Turkey). In 2021, she obtained her bachelor's degree at the Bilkent University (Turkey). In 2023, she completed her master's degree at the same university. Her master thesis focused on inverse electrondemand Diels-Alder reactions of 1,2-diazines. In 2024, she moved to the Munich (Germany) and she is currently PhD candidate at the Ludwig-Maximilians-University in Munich (Germany) by Profs. A.

Hoffmann-Röder and P. Knochel. Her research interest focuses on preparing organometallic reagents and methodologies for application in organic synthesis and the synthesis of natural products.

TMPH, DIPA and their derivatives employing readily available alkyl lithium reagents. These methods, developed by researchers such as Knochel, Snieckus, Mongin and others, have demonstrated the broad applicability of TMP-bases in numerous catalytic and synthetic transformations. 73-75 Recent reports have repeatedly emphasized the influence of ligands on the reactivity and stability of organometallic reagents. For example, organozinc pivalates (R-ZnOPiv) exhibit higher stability than halogenated organometallic

compounds (RMX, M = Mg, Zn; X = Cl, Br, I). In addition, recent kinetic and mechanistic reports have described that salt-stabilized organozinc pivalates show a significant counterion effect due to -OPiv coordination, making them easy to handle even under an oxygen-containing atmosphere. Since these reagents can be stored with no noticeable degradation or loss of yield for up to 48 h under air, they are ideally suited for various transition metal-catalyzed Negishi cross-coupling reactions.76-83



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Rambabu Dandela

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Ragupathy Dhanusuraman

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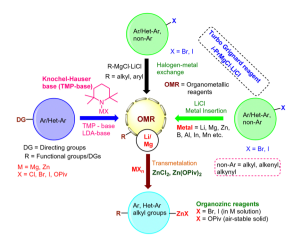


Fig. 1 Reported methods for the preparation of organometallic reagents (RMX).

Another major synthetic challenge is the regio- and enantioselective synthesis and functionalization of small organic molecules, as some of these N-heterocyclic systems can serve as chiral ligands or catalysts in synthetic transformations to produce chiral organic intermediates (Fig. 2).84,85

Highly selective and reactive aryl and alkyl organometallic reagents (Li, Mg, Zn) can be attached to electrophilic carbonyl or imine scaffolds to generate complex tetrasubstituted chiral alcohols and amines by catalysis. 86-90 The preferred organometallic compounds for these transformations are organo-Li, -Mg and -Zn reagents, due to their accessibility, low cost and nontoxicity, which favours their use in research laboratories and the pharmaceutical industry. Extensive work has shown that many functionalized fused bicyclic heteroaromatic molecules



Paul Knochel

Paul Knochel has been a full professor for organic chemistry at the Ludwig-Maximilians-Munich/Germany University (LMU) since 1999. He did his undergraduate studies at the University of Strasbourg (France) and his PhD at the ETH-Zürich with Prof. D. Seebach. He spent 4 years at the CNRS at the University Pierre and Marie Curie in Paris with Prof. J.-F. Normant and one year of postdoctoral studies at Princeton

University in the laboratory of Prof. M. F. Semmelhack. In 1987, he accepted a position as Assistant Professor at the University of Michigan at Ann Arbor, MI. In 1991, he became Full Professor at this university and in 1992, he moved to Philipps-University at Marburg (Germany) as C4-Professor in Organic Chemistry. His research interests include the development of novel organometallic reagents and methods for use in organic synthesis, asymmetric catalysis and natural product synthesis.

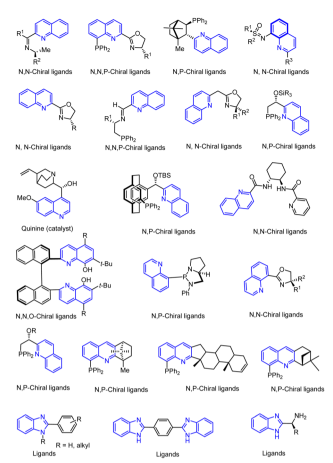


Fig. 2 N-Heterocyclic aromatic compounds for use as ligands and

are preferred structural motifs, due to their (potential) biological activity. Thus, such compounds are of particular interest for drug development (Fig. 3) but also for the production and modification of novel catalysts, as well as for material science applications.

Such applications require access to substituted scaffolds of diverse heteroaromatic core structures such as quinoline, quinoxaline, naphthyridine, indole, benzofuran, benzimidazole, benzothiophene, benzoxazole, benzothiazole, benzopyrimidine, anthranile, thienothiophene, triazole, purine, coumarin, but also chromones, quinolones and phthalazines. 91-95

Fused heterocyclic scaffolds have been decorated by various catalytic methods such as metal-free and metal-catalyzed C-H activation reactions, radical transformations, photocatalytic processes, etc. 96-98 However, only a few cases require stoichiometric amounts of transition metal catalysts or excess amounts of organometallic reagents to perform these transformations. Moreover, transition metal-catalyzed processes mediated by organometallic reagents can be associated with undesirable side effects, including β-hydride elimination and homocoupling reactions.⁹⁹ The functionalization of bicyclic fused heteroaromatic compounds using transition metal catalysts (Co, Fe, Ni, Cr) and organometallic reagents has therefore attracted

Fig. 3 Selective examples of bicyclic fused heteroaromatic natural products and bioactive compounds.

considerable attention in organic synthesis, particularly in the pharmaceutical industry and sustainable catalysis.

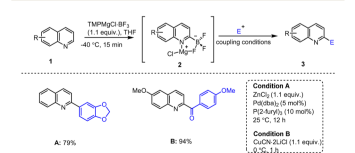
2 Functionalization of fused bicyclic hetero-aromatic compounds with TMPLi, -Mg, -Zn

In 2010, one of the best strategies for the functionalization of quinoline scaffolds 1 utilizing TMPMgCl·BF₃-mediated selective metalations of N-heterocycles was reported by Knochel's group, leading to the preparation of type 2 intermediates. ¹⁰⁰ Subsequent transmetalation with ZnCl₂ (1.1 equiv.) followed by Pd-catalyzed Negishi cross-coupling with aryl iodide afforded the 2-arylated quinoline 3a in 79% yield. Similarly, a rapid transmetalation of the magnesium species with CuCN·2LiCl and subsequent acylation reaction with acid chloride led to the corresponding heteroaryl ketone in 94% yield (Scheme 1). ¹⁰¹

Rohbogner *et al.* developed a method for the preparation of pharmaceutically active quinoline scaffolds such as Talnetant 7 and the *P*-selectin inhibitor 9 using TMP-M-mediated selective metalations of quinoline derivative 4. Cross-coupling reactions, deprotection, saponification and amination describe the reaction sequences that provide the expected target quinoline scaffolds 7 and 9 in a few steps under mild conditions and in excellent yields (Scheme 2).¹⁰²

Furthermore, to obtain the reactive magnesium intermediates 11, many N-heterocyclic phosphorodiamidate derivatives of type 10 including quinoline and quinoxaline molecules were subjected to a directed *ortho*-metalation (DoM) procedure with TMPMgCl-LiCl or TMP₂Mg·2LiCl. Later, the resulting organometallic reagents 11 were transmetalated with ZnCl₂ or CuCN-2LiCl and subsequently reacted with various electrophiles to give the corresponding arylation, acylation, thiolation and allylation products providing *e.g.*, the highly functionalized quinoline and quinoxaline scaffolds 12 in good yields of up to 87% (Scheme 3). ^{102,103}

A facile and efficient palladium-catalyzed direct benzylation of methylquinoline derivatives using TMPZnX-derived bases was accomplished in 2011 by Duez *et al.*¹⁰⁴ The desired zinc-containing quinoline derivatives were prepared by direct TMPZnCl-mediated metalations of 2/4-methylquinoline 13 in THF at 25 °C for 1 h. Subsequently, palladium-catalyzed Negishi cross-couplings of the resulting benzylic zinc reagents with a variety of aryl bromides (0.8 equiv.) were performed. When using Pd(OAc)₂ (2 mol%) and SPhos (4 mol%) as the catalyst, functionalized quinoline scaffolds 15 were obtained in

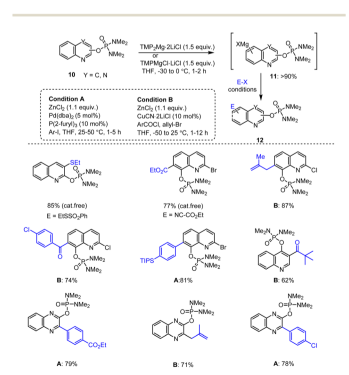


Scheme 1 TMPMgCl·BF $_3$ -mediated regioselective metalation of quinoline for subsequent cross-coupling reactions.

Scheme 2 TMP-Metal base-mediated synthesis of biologically active Talnetant and a P-selectin inhibitor scaffold

excellent yields of up to 97%, with this method also tolerating sensitive functional groups such as OH, NH2, CN and CF3 (Scheme 4).

In 2016, Mongin, Halauko and co-workers showed that a series of chloroquinolines 16 could be deprotometalated with a TMEDA-based mixed n-BuLi and TMPLi combination. Good regioselectivities were observed with a corresponding lithiumcopper bimetallic combination and were confirmed by consistent trapping with reactive acid chlorides in THF at r.t. The



Scheme 3 TMP-Base-mediated metalation of quinolines and quinoxalines for subsequent functionalization with different electrophiles.

Scheme 4 Negishi cross-coupling of 2/4-methylpyridine after zincation using TMPZnCl.

resulting carbonyl compounds 17 were produced in moderate yields (Scheme 5).105

Jaric et al. developed a successful method for the functionalization of bioactive quinine cores 18 to be used as organocatalysts by employing MeLi and TMPMgCl in the presence of BF₃·OEt₂. The subsequent trapping reactions such as Pdcatalyzed Negishi couplings, Cu-catalyzed allylations, and direct quenching with bromo- and iodo-electrophiles furnished the corresponding C-3 functionalized quinoline heterocycles **19–21** in good yields of 41–66% (Scheme 6). 106

Knochel described an efficient TMP-base-mediated protocol for the synthesis of functionalized aminoquinolines 24 via transition metal-free secondary amination of quinoline 2/8sulfonamides and 8-napthylsulfonyl chlorides. This was accomplished using R₂NMgCl·LiCl in THF at 25 °C for 1-8 h. The selective magnesiation of quinoline-2/8-sulfonamides 22 was also described using TMPMgCl·LiCl. A variety of quinoline derivatives 24 functionalized at the 2/8 position were prepared by successive C-N couplings with various amine-based organometallic reagents (R2NMgCl·LiCl) under mild conditions to afford the expected amino quinolines in good yields (Scheme 7).107 In addition, the possible mechanism of this metal-free direct amination was described based on two possible scenarios (Scheme 7). According to mechanistic pathway A,

Scheme 5 Deproto-metalation of chloroquinolines with amido-based bimetallic species and subsequent quenching with acid chlorides.

C-3- or C-2-Functionalization of quinine using TMPMgCl-LiCI

the first step is the selective addition of the magnesium reagents R'NMgCl·LiCl to the C=N bond of the quinoline skeleton 23, which leads to the formation of the key intermediate A, ultimately providing aminated product 24 after elimination of R2NSO2MgCl. Alternatively, addition of R2NMgCl·LiCl to the 2-quinolinylsulfonamide group was proposed first, resulting in species B, which can undergo an intramolecular transfer reaction of the amino group to magnesiated intermediate A. After elimination, the latter then also affords the desired aminated quinoline 24 (route B, Scheme 7).

Manolikakes et al. demonstrated a simple method for the preparation of pyridylmethyl alcohols using TMPMgCl·BF₃, a

Scheme 7 Functionalization of quinoline 2- and 8-sulfonamides by TMPMgCl·LiCl followed by desulfonylation and addition of R₂NMgCl.

Metalation of quinoline followed by 1,2-addition to aromatic Scheme 8 aldehydes and ketones.

frustrated Lewis pair that mediates selective metalations to organotrifluoroborates. The latter readily adds to a variety of aromatic aldehydes and ketones in the absence of a transition metal catalyst to afford diarylmethyl alcohols 26 in good yields of 65% (Scheme 8).108

Knochel, Zipse and co-workers reported a protocol for the full functionalization of pyrazolo[1,5-a]pyrimidine scaffolds 27, 30, 33 at positions 2, 3 and 7 using TMP-M bases. Reaction of 5chloro-pyrazolo[1,5-a]pyrimidine with TMPZn under mild conditions afforded heterocyclic zinc intermediates 28, 31, 34. Subsequent reactions with various readily accessible electrophiles in the presence of palladium or copper catalysts afforded the corresponding fused pyrazolo[1,5-a]pyrimidine derivatives 29, 32, 35 in good to excellent yields (Scheme 9-11). These functionalized heterocyclic compounds are often used in pharmaceutical applications, and the described catalytically active system tolerates both electron-rich and electron-poor functional groups such as Me, SPh, OMe, Cl, I, CN, NO2, and CO₂Et. 110,111

Unsinn et al. investigated a simple, mild, and efficient protocol for the regioselective C-3-metallation of 1H-indazoles 36 with TMP₂Zn. The resulting indazolylzinc reagents of type 37 could be smoothly arylated by Pd-catalyzed Negishi crosscoupling reactions in THF at 50 °C with various aryl iodides. The process took 8-24 h to afford the indazolyl analogs 38 in good yields. In addition, copper-mediated acylations and

Scheme 9 Selective zincation at position 7 of the pyrazolo[1,5-a]pyrimidine core for subsequent cross-coupling and acylation reactions.

Scheme 10 Selective C2-metalation of the pyrazolo[1,5-a]pyrimidine scaffold followed by electrophilic trapping reactions.

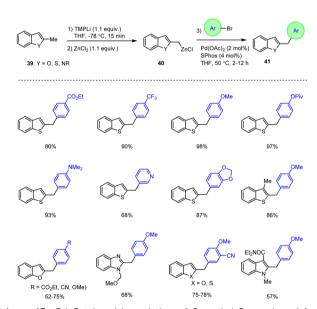
Scheme 11 Selective metalation of pyrazolo[1,5-a]pyrimidine a 2-position using TMPMgCl-LiCl and quenching with electrophiles.

allylations are also applicable under these mild conditions (Scheme 12). This synthetic method is suitable for the preparation of various biologically active N-heterocycles.

In 2012, Duez et al. developed an efficient method for the Pdcatalyzed arylation of different 2-methyl-5-membered fused heterocycles of type 39 with TMPLi bases. This innovative synthetic process involves TMPLi-mediated selective metalation at the benzylic position in THF at -78 °C, followed by transmetalation with ZnCl2 to form the corresponding organozinc intermediate 40. Subsequent Pd-catalyzed Negishi crosscoupling reaction under mild conditions provided the desired arylated fused bicycles 41 with indole, benzothiophene and benzofuran heterocyclic cores in good to excellent yields of up to 98% (Scheme 13).113 Under the optimized reaction conditions, fused bicyclic heteroaromatic compounds containing different functional groups with electron-poor and electronrich substituents were obtained, making this approach widely applicable for the synthesis of pharmaceutically active molecules (API).

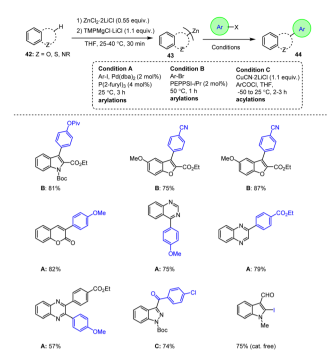
Unsinn *et al.* presented an improved strategy for the synthesis of bis-heteroaryl zinc reagents 43 and reported on their subsequent reaction with various electrophiles. The refined procedure using a TMP-Mg base in the presence of $ZnCl_2$ is

Scheme 12 Pd-Catalyzed Negishi cross-coupling reactions of indazolyl zinc reagents.



 $\begin{tabular}{lll} Scheme 13 & Pd-Catalyzed benzylation of 2-methyl-5 membered fused heterocycles using TMPLi base. \end{tabular}$

preferable to the methods previously developed by Knochel *et al.* in which zinc bases are prepared from commercially available 2,2,6,6-tetramethylpiperidinyl. Most importantly, this novel protocol enables the isolation of heterocyclic products in high yields under mild conditions and with shorter reaction times. This is particularly important for the synthesis of organozinc reagents on an industrial scale and for the subsequent arylation and acylation reactions. Remarkably, the organozinc

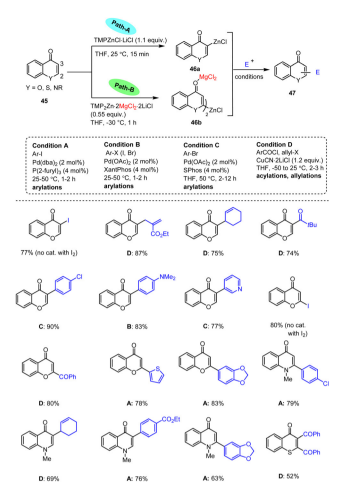


Functionalization of fused heteroaromatic indole, benzofuran, coumarin, guinoxaline and benzothiophene derivatives.

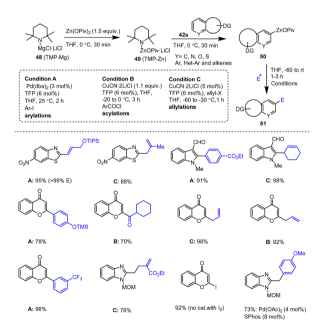
reagents not only exhibited excellent reactivity, but also tolerated a wide range of sensitive functional groups. Consequently, a direct conversion to the corresponding highly desirable organometallic zinc reagents 43 was possible, which could then be readily reacted with various electrophiles to generate functionalized heterocyclic compounds 44 in high yields (Scheme 14).114

Klier et al. presented a Lewis acid that triggered the regioselective metalation of several chromones and quinolones 45. In the absence of the Lewis acid MgCl₂, zincation is observed at the C-3 position, whereas in the presence of MgCl₂, zincation is observed at the C-2 position. Subsequent Pd-catalyzed Negishi C(sp²)-C(sp²)-coupling under mild conditions afforded the corresponding desired functionalized fused 6-membered heterocycles 47 in good to excellent yields in the range of 63-90% (Scheme 15).115

Stathakis et al. used a wide range of interesting organozinc pivalates 50 prepared by TMPZnOPiv-mediated selective metalation methods. These organozinc reagents showed high stabilities and good reactivities in C-C cross-coupling reactions with various readily available electrophiles. In addition to CuCNmediated acylation and allylation reactions, Pd-catalyzed Negishi couplings were performed in the presence of catalytic amounts of Pd(dba)2 and the ligand TFP in THF. The resulting solid organozinc pivalates of type 50 are easy to handle in industrial experiments because their stability is maintained (>90%) for 4-6 h. As shown in Scheme 16, a broad spectrum of aryl and alkenyl zinc pivalates efficiently reacted with different electrophiles, such as aryl halides, acid chlorides, allyl bromides and iodine, leading to the production of functionalized heterocyclic compounds 51 in good to excellent yields. 116



Scheme 15 TMP-Base mediated functionalization of chromones, guinolones and thiochromones using Pd-catalysis.



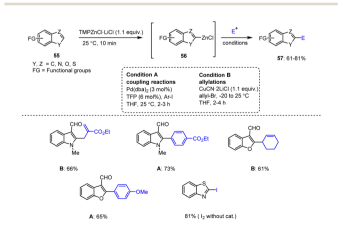
Scheme 16 Pd- and Cu-Catalyzed cross-couplings of organozinc pivalates.

Scheme 17 Cobalt-catalyzed cross-coupling reactions of heteroaryl zinc reagents with alkyl halides.

A simple and efficient cobalt-catalyzed Negishi-type cross-coupling reaction of heteroaromatic zinc reagents 53 with secondary and primary alkyl iodides or bromides using a THF-soluble homogeneous catalytic system of $CoCl_2$ -2LiCl (20 mol%) and TMEDA (30 mol%) allowed production of the desired alkylated heterocycles 54 in yields of up to 88%. As shown in Scheme 17, the required organozinc reagents were prepared from readily available heteroaromatic compounds 52 via a TMPZn-mediated selective metalation reaction. 117

Bresser *et al.* reported a wide range of functionalized heteroaryl-zinc reagents synthesized by directed zincation of the sensitive and weakly deactivated heteroaromatic compounds 55 with TMPZnCl under distinct optimized conditions. The resulting heteroaryl-zinc organometallics 56 further exhibited excellent reactivity in various electrophilic addition reactions, and afforded the corresponding heteroaromatic compounds 57 in moderate to high yields (Scheme 18). 118

Crestey *et al.* described a strategy for the functionalization of phthalazine scaffolds *via* regioselective zincation of chlorophthalazines using TMPZnCl·LiCl under microwave irradiation. This approach led to novel polysubstituted phthalazine derivatives of type **60** after trapping the resulting organozinc reagents with various electrophiles. In addition, Negishi crosscoupling reactions between chlorophthalazines and organometallic reagents were conducted using a Pd catalyst, providing



Scheme 18 Regio- and chemo-selective zincation of fused heteroaromatics using TMPZnCl.

Scheme 19 Synthesis of polyfunctionalized phthalazines using TMPZnCl·LiCl.

highly functionalized phthalazine scaffolds **59** in excellent yields of up to 94% (Scheme 19). 119

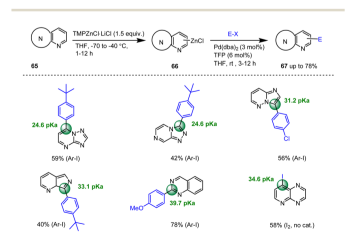
Zimdars *et al.* demonstrated the selective metalation of positions 4 and 7 of the benzo[c][1,2,5]thiadiazole scaffold **61** using TMP-bases in THF at $-40\,^{\circ}$ C for 14 h. The corresponding reactive Mg-intermediate **62** was readily transmetalated with ZnCl₂ followed by electrophilic quenching. In this way, Pd catalysis enabled the preparation of functionalized asymmetric disubstituted benzothiadiazole derivatives **64** in high yields, as shown in Scheme 20. ¹²⁰

In 2020, Balkenhohl *et al.* reported a powerful model for predicting site-selective metalation approaches with TMPZnCl-LiCl. The pKa values of the functionalized condensed N-heterocycles were calculated and compared with experimental results of deprotonations. Thus, the fused heteroaromatic bicycles **65** such as pyrido[2,3-b]pyrazine, imidazo[1,2-b]pyridazine, [1,2,4]triazolo[4,3-a]pyrazine, [1,2,4]triazolo[1,5-a]pyrimidine, and imidazo[1,5-a]pyridine as well as quinazoline were smoothly deprotonated at the predicted positions (p K_a = 24.6–39.7), leading to the corresponding aryl zinc reagents **66** in 40–70% yield. After iodolysis or palladium-catalyzed Negishi

Double functionalization of benzo[c][1,2,5]thiadiazole via TMPMg-mediated coupling reactions.

cross-coupling reactions, the expected highly functionalized Nheterocycles of type 67 were obtained in 42-59% yields (Scheme 21).121

Regioselective magnesiation of benzothiophene and benzofuran scaffolds 68 with TMPMgCl·LiCl in THF at 0 °C for 2 h, followed by a 12 h trapping reaction using the thio-electrophile (Me2NC(S)S)2 afforded the desired benzothienyl dithiocarbamates 70 in excellent yields of 76-90%. Kienle et al. also prepared substituted 2-aryl and 3-aryl benzothiophene derivatives under similar reaction conditions (Scheme 22).122 This method allows polycyclic heteroaromatic compounds such as dibenzothiophenes and dibenzothieneothiophenes scaffolds to be prepared in good yields. Derivatives of benzothiophenes, dibenzothiophenes, dibenzothieno-thiophenes and their S-heterocyclic congeners are broadly applied in various fields

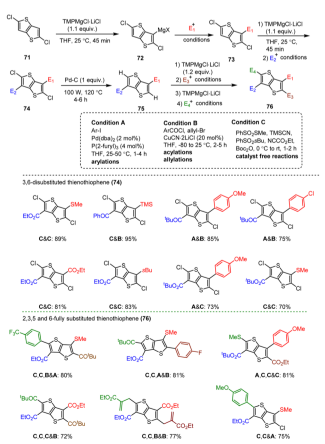


Scheme 21 TMPZnCl·LiCl-promoted regioselective deprotonations and subsequent functionalizations of N-heterocycles

Scheme 22 Preparation of heteroaryl dithiocarbamates using TMPMgCl-LiCL for metalation

such as in agriculture, for pharmaceuticals and dyes, as well as in building blocks for conductive polymers.

Kunz et al. showed that direct magnesiation with TMPMgCl-LiCl enables highly regioselective and complete functionalization of the thieno[3,2-b]thiophene core 71 under mild conditions. A wide variety of sensitive functional groups can be successfully introduced as substituents, yielding various polyfunctionalized fused thienothiophenes 74 and 76 (Scheme 23), 123 which are otherwise difficult to process into



Scheme 23 Synthesis of 2,3,5 and 6-substituted thienothiophenes.

Scheme 24 Synthesis of GSK-3 and Lck protein kinase inhibitor *via* chemoselective magnesiation, followed by Pd-catalyzed coupling reaction and deprotection.

promising heterocyclic scaffolds. This protocol could allow the fine-tuning of material properties of such S-heterocycles (*e.g.* absorption bands, overlap of frontier orbitals) by introducing specific side chains into monomeric building blocks. The conjugated heterocyclic aromatic compounds represent a new class of S-containing condensed bicyclic heterocycles and their polymers, which may also be of interest as materials for OLEDs or solar cells.

Knochel and co-workers developed a practical magnesiation protocol for trifluoromethylated pyrazinamide 77, which is carried out at 0 $^{\circ}$ C and is compatible with carbonyl functionalities. Subsequent quenching with heteroaryl halides led to satisfactory yields of arylated indole 79. After deprotection of the pyrazinamide with K_2CO_3 under green conditions, the targeted free heterocyclic amine 80 was obtained in good yield (Scheme 24).¹²⁴

Starting from alkynyl(aryl)thioethers **81**, Kunz *et al.* developed a novel intramolecular carbomagnesiation protocol for the synthesis of magnesiated benzothiophene intermediate **82**. Other heteroaromatic Mg species also reacted with readily accessible electrophiles to give the highly functionalized benzo[*b*]thieno[2,3-*d*]thiophenes of type **83** in excellent yields (Scheme 25). The method tolerates a wide range of functional groups, and the authors further elaborated the cyclization process to produce highly diverse condensed benzothiophene scaffolds as well as new complex heterocyclic analogs under mild conditions.

Frischmuth *et al.* prepared a wide range of polyfunctional SF_5 -substituted indole analogs **86** using TMPMgCl·LiCl. This was accomplished *via* the formation of organomagnesium intermediate **85** through the reaction between indole **84** and

Scheme 25 Functionalized benzo[*b*]thiophenes obtained by magnesiation and subsequent quenching with electrophiles.

Scheme 26 Functionalization at the position C-2 and C-3 of protected SF_s-substituted indole derivative using TMPMqCl-LiCl.

the TMP base over a period of 0.5–2 h at moderate conditions (Scheme 26). 126 A library of SF₅-substituted heteroaromatic compounds could be accessed with the help of this organometallic strategy to enable the discovery of new biologically active indole compounds.

Groll *et al.* showed that the combination of BF₃·OEt₂ with TMPZn bases enables regioselective functionalization of fused heterocycles and thieno-pyrimidines. Remarkably, the use of BF₃·OEt₂ together with TMPZn bases enabled efficient zincation of the C-2 position, whereas metalation at the C-6 position was observed in the absence of the Lewis acid. The pre-formed organozinc species subsequently reacted with different electrophiles smoothly to produce the desired functionalized heterocycles **88** and **89** in good yields (Scheme 27).¹²⁷

The regioselective functionalization of a broad spectrum of N-protected purine scaffolds was successfully performed by Crestey *et al.* For this purpose, they applied a TMP base to substituted purine derivatives **90** for forming a zinc or magnesium intermediate **91**. This is furnished after trapping with various electrophiles such as I₂, Br₂, Ar–I, and allyl bromide the desired highly substituted purine analogs **92** (Scheme 28). Subsequent arylation reaction was achieved with Pd(dba)₂ (2 mol%) in combination with the ligand TFP (4 mol%) at 45 °C for 14 h, while for the corresponding allylation reaction CuCN-2LiCl (10 mol%) in THF at -60 to 25 °C for 2 h was used. ¹²⁸

Klatt *et al.* reported a new and efficient protocol for the regioselective metalation of the condensed heteroaromatic cinnoline backbone using two complementary methods. While

Scheme 27 Switchable, regioselective metalation of thieno-pyrimidines with TMPZn-base.

A: 70%

TMPMgCI·LiCI (1.1 equiv.) 91 Condition A ArCOCI, allyl-Br CuCN-2LiCI (10 mol%) THF, -60 to 25 °C, 2-5 h I₂, PhSO₂SPh, (CCI₂Br)₂, THF, -60 °C to rt, 2-4 h Pd(dba)₂ (2 mol%) P(2-furyl)₃ (4 mol%) THF, 25-45 °C, 2-14 h

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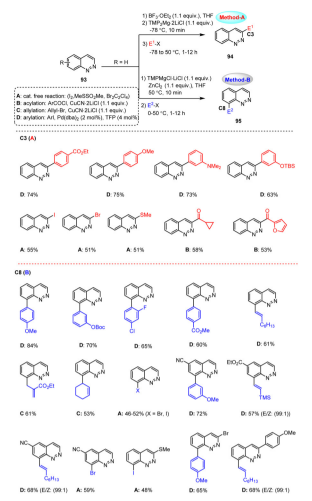
A: 63% Regioselective functionalization of purine scaffolds with TMP metal bases

A: 53%

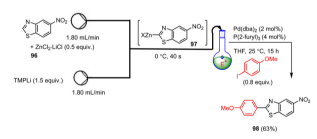
the use of BF3·OEt2 and TMP2Mg·2LiCl (method A) allowed magnesiation at the C-3 position, application of TMPMgCl·LiCl (method B) enabled selective zincation at the C-8 position of the cinnoline skeleton. By using the TMP-Mg base, this reaction allowed the formation of reactive Mg intermediates of both simple and substituted cinnoline derivatives. Subsequent coupling reactions with various electrophiles such as X2, aryl halides, allyl bromide and acid chlorides in the presence of Pd(dba)₂ (2 mol%) with TFP (4 mol%) or CuCN-2LiCl (1.1 equiv.) furnished the desired polyfunctionalized heterocycles 94 and 95 in good to excellent yields (Scheme 29). 129

By using a flow process, highly sensitive, electron-poor benzothiazoles 96 can be efficiently transmetalated upon treatment with TMPLi in the presence of MgCl2 or ZnCl2·2LiCl to yield the corresponding organomagnesium or organozinc reagents such as 97 (Scheme 30). According to Becker et al., 130 these flow reactions take place under significantly milder conditions than in the batch process within 40 s (0 °C instead of -78 °C). The resulting heteroaromatic metalation intermediates can then be reacted with various electrophiles, whereby the reaction scope of the flow metalations is also significantly larger than in the corresponding batch processes. In addition, these flow reactions can be easily scaled up by extending the reaction time without further optimization steps. As a result, easily modified benzothiazole scaffolds such as 98 can be produced in high yields.

An efficient method for the functionalization of substituted quinoxalines by metalation at the C-6 and C-8 positions was



Regioselective metalations for subsequent functionalization of the cinnoline scaffold



Scheme 30 Continuous-flow zincation of benzothiazole followed by Pd-catalyzed trapping with Ar-I.

developed by Nafe et al. using TMPLi and 2,3dichloroquinoxalines. This protocol enables the synthesis of interesting, highly functionalized quinoxaline scaffolds via subsequent Pd-catalyzed cross-couplings (Scheme 31). 131

In addition, the resulting functionalized products can be used further to construct expanded O- or S-heterocyclic compounds via anellation reactions. Such expanded quinoxaline scaffolds are characterized by robust photoluminescence with high molecular extinction coefficients within the blue and

Scheme 31 Synthesis of substituted quinoxalines using TMPLi, ZnCl₂, and electrophiles

green spectral range and thus represent interesting potential fluorescent imaging tools with fine-tuned optical properties (Scheme 32).132

Balkenhohl et al. developed regioselective metalation and functionalization reactions of the condensed 1,5-naphthyridine scaffold 103 mediated by TMP-metal bases. A clever combination of TMPMg and TMPZn bases allowed regioselective bisfunctionalization of the 1,5-naphthyridine core, which is an important heteroaromatic scaffold. Furthermore, the C-8 substituted 1,5-naphthyridine 105 allows additional regioselective functionalization at the C-4 position by using TMPMg or TMPZn bases under mild conditions (Scheme 33). 133 Thus, this reaction method is not only a key technique for the design and synthesis of OLED materials, but can also be used for pharmaceutical applications, such as the production of potentially antibacterial and antiviral agents.

A recent discovery of a highly regioselective functionalization of pyrazolo[1,5-a]pyridine by TMPMgCl·LiCl at the C-2 and

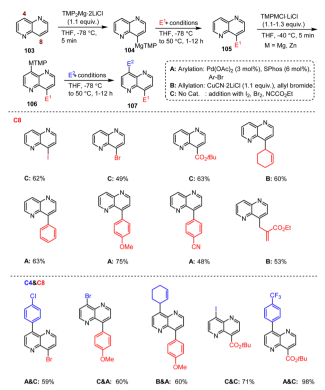
$$\begin{array}{c} R^{1} \\ \downarrow \\ N \\ Cl \\ R^{2} \\ \hline 101 \\ Y = OH, SH \\ \end{array}$$

$$\begin{array}{c} K_{2}CO_{3} (5 \text{ equiv.}) \\ DMF, 25 °C, 22 \cdot 48 \text{ h} \\ R^{2} \\ \hline \end{array}$$

$$\begin{array}{c} R^{1} \\ N \\ N \\ \end{array}$$

$$\begin{array}{c} R^{3} \\ R^{3} \\ \hline \end{array}$$

Functionalization of quinoxalines at C-2 and C-3 for anella-Scheme 32



Scheme 33 Regioselective metalation and functionalization of 1,5naphthyridines.

C-7 positions, guided solely by the presence or absence of BF₃. OEt2, was also described by Balkenhohl et al. A wide range of functionalized pyrazolo[1,5-a]pyridine derivatives 109 and 110 thus obtained under moderate metalation conditions with suitable regioselectivities. The organomagnesium reagents prepared in situ reacted smoothly with various electrophiles, such as I2, C2Cl4Br2, NCCO2Et, MeSSO2Me, and Tietze's reagent. Moreover, CuCN-mediated acylation and allylation reactions were also successfully carried out (Scheme 34). 134

Knochel, Wagschal et al. reported a highly regioselective metalation of various aryl-substituted azoles by using a sterically hindered TMPMgBu base. In this reaction, arylazole 111 for example, was allowed to react with TMPMgBu in a toluene/ hexane combination at room temperature for 1-6 h before it was subjected to Negishi cross-coupling reactions with various (hetero)aryl halides in the presence of organozinc reagents and a suitable palladium catalyst. The resulting polyfunctionalized arylazole scaffolds 112 were isolated in excellent yields of up to 91% (Scheme 35). 135,136 This protocol could be useful for preparing key intermediates of active pharmaceutical ingredients (API), as well as for several late-stage modifications of drug-related aromatic compounds. Mechanistic studies emphasize the key role of the TMPMg base for the observed selectivity, which could be exploited for different cross-coupling reactions and synthetic applications in organic chemistry.

Furthermore, Knochel, Bein and co-workers reported a selective functionalization sequence of readily available 7substituted SEM-protected 1H-imidazo[1,2-b]pyrazole scaffold.

Scheme 34 Metalation of pyrazolo[1,5-a]pyridine for functionalization at C-2 and C-7.

Successive functional group installations in the 3- and 2positions were achieved through consecutive metalations using metal amides followed by quenching reactions with suitable electrophiles. For example, the cyano-substituted 1Himidazo[1,2-b]pyrazole 116 was selectively metalated at C-3 with

Scheme 35 Regioselective metalation and functionalization of various aryl substituted azole systems using a sterically hindered TMPMgBu base.

Scheme 36 Selective metalation and guenching reactions of 7-CN substituted 1H-imidazo[1,2-b]pyrazole derivatives.

TMPMgCl·LiCl to generate the magnesiated intermediate 117, which was then successfully reacted with different electrophiles to provide e.g., allylated, acylated, thiolated as well as Negishitype arylation products 118 in 65-84% yields (Scheme 36). 137

Further functionalization in position 2 of the 3-ester substituted N-heterocycle 119 was achieved via a bis-organozinc species 120 generated upon treatment with bis-base TMP₂Zn-MgCl₂·2LiCl at 0 °C in THF. Subsequent Cu-catalyzed acylations with various types of acylchlorides and Negishi-type crosscouplings proceeded smoothly to afford the desired trisubstituted heterocycles 121 in good yields (Scheme 37). 137 The high chemoselectivity of the reactive intermediate zinc species even allowed the use of electrophiles containing sensitive functional groups such as an ester or a nitro group.

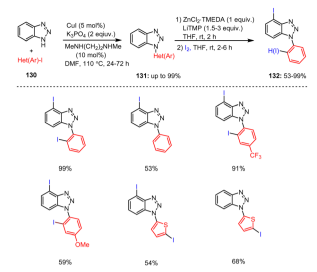
Scheme 37 Selective metalation and functionalization of the 7-CN and 3-CO₂Et disubstituted 1*H*-imidazo[1,2-*b*]pyrazoles.

Consecutive functionalization of 1H-imidazo[1,2-b]pyrazole within the synthesis of the pruvanserin isostere 126.

Moreover, this consecutive functionalization sequence was applied to the synthesis of a non-classical isostere 126 of the indolyl drug pruvanserin (Scheme 38). The latter is a selective 5-HT2A serotonin receptor antagonist suffering from low solubility under physiological conditions. Comparative assays between the original drug and the isostere revealed a significantly improved solubility in aqueous media due to the substitution of the indole ring with the 1*H*-imidazo[1,2-*b*]pyrazole core. Fused five-membered N-heterocyclic scaffolds such as 1Himidazo[1,2-b]pyrazoles recently attracted much attention as key structural elements for many pharmaceutical, agrochemical and material science applications. 138

Melzig et al. established an efficient two-step protocol for the 2,3-difunctionalization of benzofuran scaffolds 127 (Scheme 39). Here in the first step, a sulfoxide group acts as a metalation-directing group (DoM) in the presence of TMPMgCl-LiCl to allow smooth ortho-magnesiation and electrophile trapping. In the second step, the sulfoxide group of 128 works instead as a leaving group enabling a sulfoxide-magnesium exchange in the presence of commercially available Turbo-Grignard (i-PrMgCl·LiCl). Upon further reaction of the in situ prepared novel organomagnesium species with electrophiles, highly functionalized heterocyclic compounds 129 were obtained in good yields. The chemoselective TMPMgCl·LiCl and i-PrMgCl·LiCl reagents are compatible with a wide range of functional groups (e.g. F, Cl, CF₃, CN, CO₂^tBu, alkynyl, ether, thioether), so that this method is particularly suitable for gramscale syntheses in standard laboratories. 139

Scheme 39 TMPMgCl-Mediated metalation of benzofuran followed by a sulfoxide-Mg exchange reaction and trapping with various electrophiles.



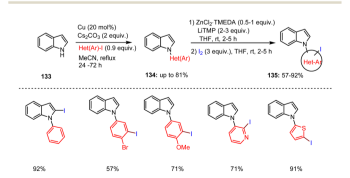
Scheme 40 Deproto-metalation of benzotriazole by TMP-base for subsequent iodination

Mongin, Halauko, Chevallier and co-workers developed a deproto-metalation method for several N-arylated 1Hbenzotriazoles 130 with TMP-bases. The in situ prepared organometallic reagents reacted readily with commercially available iodine reagents and gave mono- or diiodinated heterocyclic compounds 132 in good to high yields. Varying amounts of the TMP-base (1.5-3 equiv.) were used with substrates bearing electron-rich and electron-poor substituents (Scheme 40). 140

The same group also succeeded in functionalizing indole derivatives 134 under very similar conditions, i.e., in the presence of TMPLi and ZnCl2·TMEDA. For example, selective deproto-metalation of the protected indoles followed by quenching with I2 led to the formation of the iodinated indole derivatives 135 in satisfactory yields (Scheme 41). 141

In addition, a strategy for the direct metalation of methylated quinoline and quinoxaline scaffolds 136 using LDA and TMPLi as bases was reported in 2023. Subsequent capture reactions with various electrophiles at low temperatures led to quinolinyl alcohol and amine derivatives 137 in moderate yields (Scheme 42).142

Clososki et al. described the straightforward C-2 and C-5 functionalization of indolizine motifs 138 with an ester group



Scheme 41 Deproto-metalation of protected indole by TMP-base for

Scheme 42 Deproto-metalation of methylated guinolines and guinoxalines using LDA or TMPLi bases for subsequent electrophilic trapping reactions

at C-1. The directed metalation process took place under mild conditions using the organometallic bases LDA or TMPMgCl-LiCl, whereby the reaction of the corresponding organometallic intermediates (A and B) with different electrophiles enabled the production of difunctionalized indolizines 139 and 140 in high yields. While LDA favoured C-5 functionalization, the TMPMg base yielded mostly C-2 functionalized derivatives via selective ortho-metalation. However, the regioselectivity of these reactions was not only dependent on the choice of the base. Rather, in the case of the TMPMgCl·LiCl-mediated reaction, electrophile-controlled regioselectivity was observed due to a dynamic equilibrium between the two reactive C-2/C-5organomagnesium species. The scope of applicable substrates for this process is summarized in Scheme 43143 affording a

Scheme 43 LDA- and TMPMgCl-mediated selective directed metalation of Indolizine derivatives followed by reaction with various electrophiles.

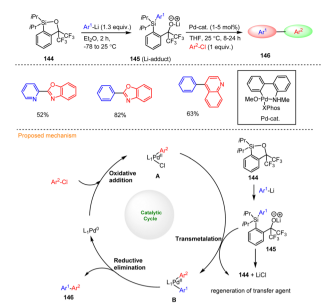
wide range of potentially biologically active heterocyclic compounds in good yields and with high functional group tolerance.

3 Functionalization of fused bicyclic heteroaromatics with organolithium reagents RLi

Snieckus and co-workers presented a general method for preparing air-stable ortho-boropinacolato aryl sulfonamides 142 via the directed ortho-metalation (DoM) method. Metalation of aryl sulfonamides of type 141 with LDA at −78 °C generates highly reactive aryl-lithium species. Subsequent trapping with i-PrOBPin boron reagents (4 equiv.) produced the corresponding ortho-boropinacolato aryl(heteroaryl) sulfonamides 142 in good yields. Moreover, 142 was subjected to a Pdcatalyzed C(sp²)-C(sp²) Suzuki-Miyaura cross-coupling reaction with readily accessible aryl and heteroaryl halides in DMF/H2O at 80 °C providing biaryl and heterobiaryl sulfonamides and in particular functionalized indole scaffolds 143 (Scheme 44). 144 This method overcomes previous failings in the preparation of aryl sulfonamide boronic acids and should be of value for medicinal chemistry programs centered on the sulfonamide functional group.

Recently, Houk, Smith and co-workers developed an efficient protocol for the palladium-catalyzed cross-coupling reaction of readily available aryl and pyridyllithium reagents 145 at room temperature using a reusable siloxane transfer reagent 144. The crystalline, bench-stable siloxane reagent is easily prepared in a one-step protocol, and its use eliminates the need for prefunctionalization, as well as isolation of organometallic crosscoupling partners. Importantly, this reagent can be recovered and reused without sacrificing reactivity. Both electron-rich and electron-poor substrates can be efficiently cross-coupled, and a variety of common functional groups on the electrophilic partner were well tolerated (i.e. esters, nitriles, azaheterocycles, fluorinated aromatics and quinolines) as well as sterically burdened aryl chlorides. Hence, functionalized quinoline and benzoxazole derivatives 146 can be obtained in acceptable yields by C(sp²)-C(sp²) cross-coupling reactions of aryl lithium and aryl chlorides in the presence of Pd pre-catalyst (1-5 mol%) and the XPhos ligand (Scheme 45). The

Scheme 44 LDA-Mediated one-pot directed ortho-metalation of aryl sulfonamides followed by Suzuki Miyaura cross-couplings.



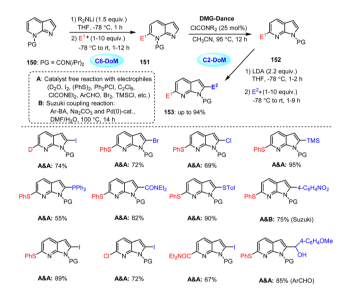
Scheme 45 Pd-Catalyzed cross-coupling reactions of heteroaryl chlorides with aryl lithium reagents in the presence of siloxane transfer reagent 144

Pd-catalyzed cross-coupling reaction of the heteroaryl chlorides with aryllithium reagents comprises the following typical steps: oxidative addition to the intermediate A, followed by transmetallation with ArLi to form intermediate B. Reductive elimination finally yields the desired heterocyclic product 146 in good yield and regenerates the active Pd catalyst for the next cycle run.

Efficient regioselective functionalization of fused 7azaindole heterocycles using LDA and amide bases was reported by Kitching, Snieckus and co-workers. Carbamoylprotected 7-azaindole 147 underwent a rapid LDA-mediated regioselective metalation followed by trapping with various electrophiles to generate C-2-substituted 7-azaindole derivatives 149 in excellent yields (Scheme 46).146

Under similar mild conditions, the regioselective functionalization of 7-azaindole by controlled ring isomerization using directed metalation group (DMG) migration was also achieved (Scheme 47). 147 By using a TMPLi base, azaindole 150 was first regioselectively metalated at the C-6 position and then quenched with an electrophile to obtain a C-6-substituted derivative 151. Subsequently, a carbamoyl group shift (a dance from N7 to N1) was performed in the presence of a catalytic amount of ClCONR₂, leading to the formation of product 152. A second directed metalation and electrophile quench sequence then provided 2,6-substituted azaindoles 153. Overall, the controlled migration of the carbamoyl group enables multiple functionalization events of the bioactive azaindole scaffold, bypassing the removal and introduction of another DMG and instead allowing the same DMG group to direct functionalization at a new, distant site. Furthermore, the use of the directed metalation group dance strategy could be applied to a latestage deuteration of an antipsychotic compound (L-745870)

Scheme 46 Regioselective functionalization of Carbamoyl-protected 7-azaindoles using LDA.



Scheme 47 Regioselective functionalization of 7-azaindole at C-6 and C-2 positions via controlled DMG migration.

demonstrating a new strategy for the functionalization of the bioactive azaindole scaffold and related N-heterocycles.

The same research group also reported a method for a rather unusual C-4 peri-metalation of NH-free azaindoles 154, in which CONEt2 and SO2NEt2 groups as DMG on C-3 allowed the highly regioselective synthesis of 4-substituted derivatives via anionic shielding of C-2. The reaction is robust and scalable. If such anionically shielding DMG is introduced at C-4 (156), successive ortho-metalations (DoM) of C-2 (157) and C-5 (158) are possible. The multiple, sequential DoM reactions (i.e., ring-walk metalation sequences) provide a rational and generally regioselective route to polysubstituted 7-azaindoles 159 and other heterocycles of potential pharmaceutical significance (Scheme 48).148

Scheme 48 Organolithium-mediated C-4-peri-metalation of substituted 7-azaindole derivatives followed by electrophilic additions.

4 Functionalization of fused bicyclic heteroaromatics by organo-Mg reagents (RMgX)

In 2017, Provot and co-workers developed an efficient Kumada coupling reaction between chlorovinyl benzothiophene 160 and Grignard reagent 161 in the presence of Fe(acac)₃ (10 mol%) in THF at -30 °C leading to the highly substituted benzothiophene 162 in good yields (Scheme 49). 149

Knochel and co-workers have shown that similar to Fe(III) catalysis, chemoselective cross-coupling reactions of the Kumada type can also be carried out using catalytic amounts of chromium(II) chloride (Scheme 50). Thus, substrates considered challenging such as isoquinoline and quinoline derivatives 163 were successfully coupled with a wide range of functionalized aryl(heteroaryl)-Grignard reagents 164 in the presence of CrCl₂ (3 mol%). These reactions, which led exclusively to the formation of the α-arylated heterocycles 165 without significant amounts of homocoupling products, proceed rapidly within minutes at room temperature in cyclopentyl methyl ether (CPME). Several functional groups, including esters and acetals, are tolerated by this method and the

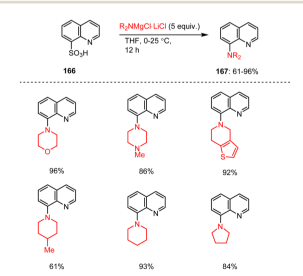
Scheme 49 Fe-Catalyzed Kumada coupling for the synthesis of 2,3disubsituted benzothiophene derivatives

Scheme 50 CrCl₂-Catalyzed cross-couplings of 2-quinolinyl halides with ArMgX.

required chromium salts can be successfully separated after the reaction using solid supports.

Later, the research group also developed a strategy for the transition metal-free amination of 8-quinoline sulfonic acids 166 using magnesium amides of the type R2NMgCl·LiCl. Thus, numerous cyclic secondary amines were first successfully converted by i-PrMgCl·LiCl into the corresponding magnesium amides, which in turn reacted readily with the quinoline sulfonic acids under mild conditions and provided the desired aminoquinolines 167 in excellent yields (Scheme 51).151 Aminoquinoline scaffolds derived from heteroaryl sulfonic acids are of major interest in pharmaceutical chemistry and drug development.

Steib et al. reported on a ligand-free Cr-catalyzed amination reaction of various N-heterocyclic quinoline and quinoxaline chlorides. The catalytic regioselective amination of 2-chloroquinolines, 1-chloroisoquinolines and 2,3-dichloroquinoxalines 168 with a



Scheme 51 Catalyst-free amination reaction of 8-quinoline sulfonic acid using magnesium amides

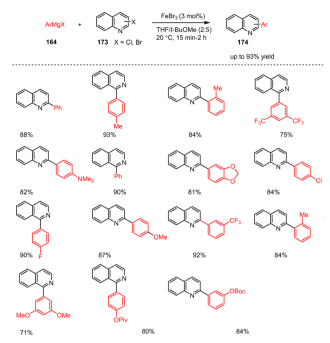
Scheme 52 Cr(II)-Catalyzed amination of 2-quinonyl chlorides with magnesium amides.

wide range of aliphatic, benzylic, and saturated (hetero-)cyclic magnesium amides were described. The C-N coupling reactions were carried out in THF at 50 °C for 2-24 h and led to the desired aminated bicyclic heteroaromatic compounds 169 in 54-95% yields (Scheme 52).152

The highly regioselective synthesis of functionalized Sheterocycles 172 under mild conditions and with the use of five-membered fused 2-thienyl-magnesium intermediates 171 was presented by Sämann et al. For this purpose, an efficient Br/Mg exchange reaction between unsymmetrically substituted dibromothienothiophenes 170 and i-PrMgCl·LiCl in THF at 0 °C was carried out in good yields. Ring substituents, such as thioether or trimethylsilyl groups, as well as pyridyl and thienyl groups or ortho-substituted aryl groups of the thienothiophenes directed the Br/Mg exchange at position C5 with excellent regioselectivity of up to >99:1 (Scheme 53). Subsequently, these heterocyclic magnesium derivatives were selectively functionalized with electrophiles such as aldehydes, aryl iodides, acyl chlorides or aryl sulfinyl chlorides, providing the targeted thienothiophene derivatives. Finally, the resulting bromo heterocycles can be readily subjected to a second Br/Mg exchange, followed by further electrophilic functionalizations.

A simple and practical method was developed by Kuzmina et al. using a non-toxic iron catalyst. This new approach for 2-substituted quinoline and isoquinoline scaffolds allowed smooth C(sp²)-C(sp²) cross-couplings of N-heterocyclic halides 173 with various electron-poor and electron-rich aryl

Scheme 53 Preparation of functionalized thienothiophenes by reaction of heteroaryl magnesium reagents with electrophiles.



Scheme 54 FeBr₃-Catalyzed cross-coupling reactions of quinoline- and isoquinoline-based halides with functionalized organomagnesium reagents.

magnesium reagents (ArMgX, 164) in the presence of FeBr₃ (3 mol%) (Scheme 54). 154 The inexpensive and efficient iron cross-coupling reaction was carried out in a mixture of THF and tBuOMe at 20 °C and turned out to be a key procedure for the preparation of highly functionalized N-heterocycles 174 in excellent yields with no formation of undesired homocoupling by-products. Thus, organomagnesium reagents with a variety of sensitive functional groups such as F, Cl, CF₃, OPiv, OBoc, OMe, Me, and NMe₂ were successfully employed in this method. The resulting quinolines and heterocyclic derivatives thereof have shown major promise for the treatment of several diseases, including inflammation, cancer, diabetes, and malaria, as well as for various viral infections.

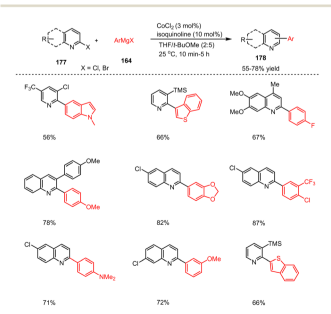
In 2013, Knochel and co-workers published a simple and efficient catalytic process for the preparation of 2- or 4substituted heterocyclic quinoline motifs 176. Using CrCl₂ (3 mol%) catalyst, N-heterocyclic chlorides 175 could be successfully used in a Kumada cross-coupling reaction with aryl(het)-magnesium reagents 164 under sustained conditions. When operated for 15 min up to 2 h in THF at 25 °C, this technique provided substituted quinolines 176 with good yields of up to 89% (Scheme 55). 155

In addition, the same group reported the serendipitous discovery that the addition of quinoline or isoquinoline dramatically increased the rate and yield of Fe- and Co-catalyzed cross-coupling reactions. This ligand acceleration allowed the general scope of the described cross-coupling reactions to be extended to complex functional groups and the formation of heteroaryl-heteroaryl bonds, where the desired products were previously obtained at very low yields. For example, by using

Scheme 55 CrCl₂-Catalyzed coupling reaction of quinoline-based chlorides with arylmagnesium reagents.

CoCl₂ (3 mol%) as a catalyst in the presence of isoquinoline (10 mol%), functionalized aryl and heteroaryl Grignard reagents **164** were successfully coupled with Br-, Cl-substituted quinolines **177** in reasonable yields (Scheme 56). 156

Starting from easily accessible bromoaniline derivatives 179, Frischmuth et al. investigated a mild and efficient intramolecular Cu-mediated carbomagnesiation method for the preparation of functionalized 4-azaindoles of type 181. Subsequent further functionalization of these 4-azaindoles with various electrophiles thus provided access to highly functionalized N-heterocycles of type 184 in excellent yields of 74-85%. The preparation of key magnesium intermediates 180 for intramolecular cyclization was carried out by halogen-metal exchange reactions using i-PrMgCl·LiCl, tolerating a broad spectrum of functional groups in the substrate. Starting from TMS-containing heterocyclic precursor 181, which is available in a few steps, the 3-iodoazaindole 182 was prepared by treatment with ICl, and then subjected to the halogen-metal exchange followed by an electrophilic scavenging reaction with acid chlorides. The corresponding, highly functionalized



Scheme 56 Co-Catalyzed Kumada coupling of 2-quinonyl halides

Scheme 57 Synthesis of 2,3-disubstituted 7-azaindoles using i-PrMgCl.

4-azaindole ketones **184** were obtained in reasonable amounts (Scheme 57). 157

Barl *et al.* reported the complete functionalization of the 7-azaindole scaffold **188** using the following reaction sequence: halogenation, cyclization, directed metalation, and halogen/Mg as well as sulfoxide/Mg exchange. By using this procedure, a complex and fully functionalized 7-azaindole **197** was obtained starting from the commercially available aniline derivative **185** via sequential transformation of the corresponding substituted key intermediate **188** (Scheme 58). Using n-BuLi, TMPLi and i-PrMgCl as key metalating reagents, this multistep protocol afforded the desired key azaindole structure **197** with moderate yields and high tolerance of functional groups.

In 2013, Knochel and co-workers developed a formal regioselective C(sp²)–C(sp³) cross-coupling of substituted pyridines and quinoline derivatives **198** *via* mild BF₃·OEt₂-mediated nucleophilic addition reaction of Grignard or organozinc reagents, followed by chloranil-mediated oxidative aromatization. The high regioselectivity and broad tolerance towards functional groups make this method very valuable for the preparation of polyfunctional pyridines and quinolines **199** (Scheme 59).¹⁵⁹ Moreover, these reactions can be easily carried out on a larger scale with no reduction in yield.

A general and mild intramolecular method for a coppermediated carbomagnesiation reaction was presented by Nickel et al., which can be used to synthesize functionalized Nheterocycles such as pyrrolo[2,3-d]pyrimidines **201** and azaindole derivatives **208**. In this work, pyrrolo[2,3-d]pyrimidines were prepared from metalated pyrrolo[2,3-d]pyrimidines, which in turn were accessible by treatment of *N*-alkynyl-5iodo-6-sulfamido-pyrimidines with i-PrMgCl·LiCl, followed by transmetalation with CuCN·2LiCl and intramolecular carbocupration. Finally, the desired polyfunctional pyrrolo[2,3-d]pyrimidines **200** were obtained after an electrophilic quenching

Scheme 58 Method for full functionalization of the 7-azaindole core by selective metalation and sulfoxide/Mg exchange reaction.

Scheme 59 Lewis acid-promoted direct alkylation of quinoline derivatives with alkylmagnesium reagents.

reaction with allyl halides or acid chlorides. In addition, subsequent reaction with ICl, followed by Negishi cross-coupling in the presence of PEPPSI-iPr as the catalyst, allows these compounds to be further functionalized and converted into fully substituted N-heterocycles **204** (Schemes 60 and 61). ^{160,161}

The methods described could offer short synthetic routes to biologically relevant molecules, as shown by the formal synthesis of the marine alkaloid rigidin-A 205 and of 7-azaserotonin

Scheme 60 Synthesis of rigidin-A using Grignard reagents.

Scheme 61 Synthesis of 7-azaserotonin analog 211 using RMgX.

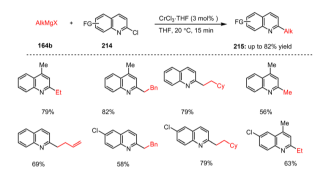
211, a closely related serotonin derivative (Schemes 60 and 61). 160,161

Greiner *et al.* reported a practical method for the modification of 1,8-disubstituted 2,7-naphthyridine derivatives by iron-catalyzed C(sp²)–C(sp³) cross-couplings with alkyl Grignard reagents. To incorporate alkyl substituents into the naphthyridine backbone, the 3,6-chloro substituents of 2,7-naphthyridine 212 have been replaced through cross-coupling reactions with various organomagnesium compounds 164a resulting in tetrasubstituted symmetric and unsymmetric naphthyridine motifs 213 (Scheme 62).¹⁶²

Similarly, iodoisoquinolines and chloroquinolines **214** can be alkylated with alkylmagnesium reagents **164b** using the tetrahydrofuran-soluble chromium(III) complex CrCl₃·3THF. This efficient chromium(III)-catalyzed C(sp²)-C(sp³) cross-coupling protocol proceeds within minutes at room temperature and yields the desired products **215** without the formation of homocoupling by-products that are commonly observed in the related iron-, cobalt- or manganese-catalyzed reactions (Scheme 63). ¹⁶³

In addition, Ziegler *et al.* developed a new halogen–Mg exchange reagent **217** using the alkylmagnesium alkoxide *s*-BuMgOR·LiOR (R = 2-ethylhexyl), which undergoes very fast Br/Mg and Cl/Mg exchange reactions in toluene. These processes are not only about 30 times faster than those of the previously

Scheme 62 Fe-Catalyzed cross-couplings 1,8-dimethylnaphthyridine with alkyl Grignard reagents



Scheme 63 Cr(III)-Catalyzed C(sp²)-C(sp³) Kumada coupling reactions of alkylmagnesium reagents with 2-chloroquinoline

used exchange reagent sBu₂Mg·2LiCl, but also about 110 times faster than those with Turbo-Grignard (i-PrMgCl·LiCl). In addition, the resulting Grignard reagents of the type ArMgOR·LiOR or HetArMgOR-LiOR can be easily added to ketones and acyl chlorides to form 218-type reaction products under mild conditions (Scheme 64).164 The synthesis of Grignard reagents in hydrocarbons or toluene is of great interest, as these weakly coordinated Grignard reagents can exhibit unusual reactivity and are also considered as industry-friendly reagents.

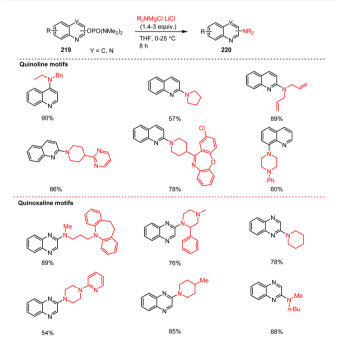
(1.2 equiv.) TMEDA (2 equiv.) Condition A Condition B Cul (10 mol%) ArCHO 25 °C,1 h A: 70% B: 62%

Scheme 64 Functionalized fused heteroarenes via Br/Mg-exchange reaction and quenching with electrophiles.

Aminated N-heterocycles play an important role in modern pharmaceutical chemistry, which makes the development of synthetic protocols for their efficient preparation very attractive. In 2018, Knochel and colleagues presented the amination of phosphorodiamidate-substituted quinolines and quinoxalines with the magnesium amides R2NMgCl·LiCl under catalyst-free conditions. Here, 2-, 4- and 8-hydroxyquinoline and 2-hydroxyquinoxaline derivatives were converted to the corresponding phosphorodiamidates 219 and subjected to an amination reaction using various pharmaceutically active amines to give the fused N-heterocycles of type 220 (Scheme 65).165,166

Since the phosphorodiamidate function is a strong direct metalating group (DMG), the amination can be combined with an ortho-functionalization step. Thus, several phosphorodiamidate-substituted N-heterocycles 221 were treated at 0 °C for 1 h with TMPMgCl·LiCl or TMP₂Mg·2LiCl in THF. The type 222 Mg species formed were then either quenched with electrophiles such as I₂ or (BrCl₂C)₂ or were successfully employed in Cu-catalyzed acylation reactions and Pd-catalyzed Negishi cross-couplings after Zn transmetalation steps. The resulting functionalized heterocycles 223 were finally subjected to the amination reaction that led to 2,3-difunctionalized quinoline derivatives 224 in 35-66% yields over two steps (Scheme 66). 165

Sulfur-containing organic molecules are often useful building blocks in organic synthesis, which is why, transition metalcatalyzed desulfinative cross-coupling reactions are used to produce heterobiaryl products, for example. As these processes usually require high temperatures due to catalyst deactivation, milder and transition metal-free desulfination protocols are desirable. Along this line, Wei et al. presented a cross-coupling reaction of heteroaryl sulfinates with Grignard reagents for the



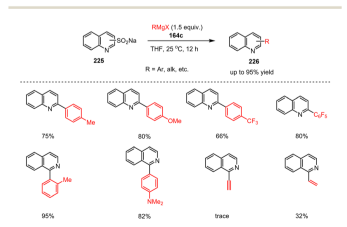
Scheme 65 Synthesis of 2-amino quinoline and quinoxalines with

TMP-base mediated directed ortho-metalation and functionalization of various aryl phosphorodiamidates, followed by amination with R2NMqCl-LiCl.

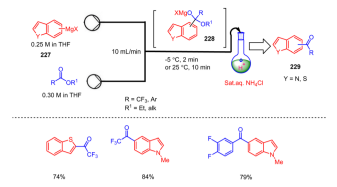
synthesis of heterobiaryls. Here, fused N-heteroaryl sulfinates 225 were reacted with functionalized aryl and alkyl Grignard reagents 164c to produce quinoline-based biaryl heterocycles 226 under mild conditions (Scheme 67). The transformation demonstrates great potential for the preparation of functionalized fused heteroaromatic compounds by utilizing quinoline and pyridine sulfinates as electrophilic starting materials for C-N cross-coupling reactions with no need for additional catalysts or bases.

A selective acylation protocol developed by the Knochel group for readily available heteroaryl magnesium reagents 227 and commercially available ester derivatives run at favourable temperatures (-5 to 25 °C) and short reaction times (2-10 min) under continuous flow conditions. The flow conditions prevent premature collapse of the hemiacetal intermediates despite non-cryogenic conditions, thereby providing satisfactory yields for heteroaryl ketones 229. The coordination ability of the ester and aryl(het)magnesium reagents was decisive for the outcome of the acylation reaction (Scheme 68). 168

Knochel, Bein and co-workers reported the selective functionalization of the 5-Br-substituted 1*H*-imidazo[1,2-*b*]pyrazole



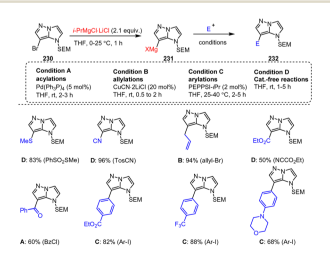
Scheme 67 Transition-metal-free functionalization of heterocycles via desulfinative cross-coupling of quinolinyl sulfinates with aryl and alkyl Grignard reagents.



Scheme 68 Synthesis of heteroaryl ketone using ester scaffolds and Grignard reagents

scaffold 230 using a Br/Mg exchange reaction with i-PrMgCl in THF at r.t. The resulting reactive Mg intermediate 231 reacted with various commercially available electrophiles, such as by direct quenching with highly reactive electrophiles or by Pd-catalyzed cross-coupling as well as Cu-mediated acylation or allylation reactions. Overall, these sequences led to the formation of functionalized condensed N-heterocycles 232 in good to excellent yields (Scheme 69). 137,169

Kürti, Ess and co-workers have described an intramolecular amination of arene C(sp²)-H bonds at low temperature, without the use of transition metals and with high regioselectivity. The reaction is operationally simple and scalable (1-10 mmol) and allows the formation of fused N-heterocycles 234 under mild conditions, using readily available 2-nitrobiaryls 233 and PhMgBr (Scheme 70). 170 Initially, the Grignard reagent attacks the nitro group of 233, whereby a reactive aryl nitroso intermediate B is formed after elimination of magnesium phenolate. This species then reacts with a second equivalent of PhMgBr to form an aryl nitrene intermediate D, which cyclizes intramolecularly to the desired carbazole derivative 234. This method also allowed the synthesis of the two bioactive carbazole alkaloids Clausin V and Glycoborin.



Scheme 69 Selective functionalization of the brominated fused aromatic 1H-imidazo [1,2-b]pyrazole via Br/Mg exchange reaction with i-PrMgCl.

Scheme 70 Highly regioselective synthesis of fused carbazoles using Grignard reagents.

In 2010, the group of Knochel and co-workers developed a simple and extremely versatile one-pot strategy for the preparation of organo-fluorine compounds by converting functionalized aryl halides 235 into aryl and heteroaryl fluorides 237. This method enables efficient and direct, metal-free syntheses of fluorinated isoquinoline and benzothiophene derivatives as well as arylated fluorine compounds, which are otherwise difficult to prepare by the conventional catalytic methods (Scheme 71).171

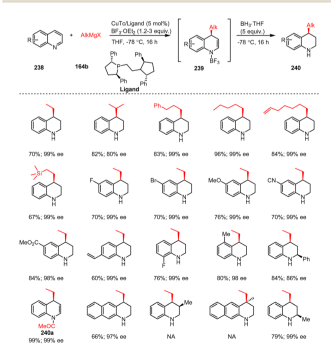
The dearomative functionalization of heteroaromatics is one of the most straightforward approaches for the synthesis of chiral heterocyclic systems, key building blocks for both

(PhSO₂)₂NF (1.2 equiv.) (1.1 equiv.) 4:1 mixture of DCM Y = N. S 237 PhO₂S PhO₂S PhO₂S

Electrophilic fluorination of heteroaryl magnesium reagents using NFSI.

synthetic chemistry and drug discovery. Yan et al. have recently presented a catalytic system that enables the nucleophilic dearomatization of quinolines 238 in combination with organometallic compounds very efficiently. A synergistic combination of Lewis acid, chiral copper(1) catalyst, and Grignard reagent allows to overcome the energy barrier of dearomatization and to obtain chiral C-4 addition products 240 with almost absolute regio- and stereo-control. Remarkably, in a preparative reaction, the amount of chiral copper catalyst can be reduced to 0.1 mol%, leading to the highest turnover number (TON) reported so far in any enantioselective reaction with Grignard reagents. Molecular modelling suggests that the role of the Lewis acid is not only to activate the substrate towards potential nucleophilic addition, but also to subtly direct the regiochemistry, by preventing C-2 addition (Scheme 72). 172

A convenient method for the synthesis of functionalized quinolines is based on site-selective modification by C-H functionalization of easily accessible quinoline scaffolds. An illustrative example of such an approach involving Ni-catalyzed C-H bond arylation of 8-aminoquinoline motifs at the distant C-4 position was presented by Qiu, Kambe and co-workers. The authors proposed the following reaction mechanism for this catalytic direct C-H arylation at C-4 of 8-aminoquinoline scaffolds: first, deprotonation of the quinolinylamide 241 with t-BuOK takes place, resulting after isomerization, in the formation of intermediate B. Its coordination to an in situ generated Ni(0) species leads to the key intermediate C, which then undergoes a nucleophilic addition of ArMgX to afford intermediate D. After HMgX elimination and subsequent protonation, the desired functionalized quinolines 242 are formed. The protocol shows a broad range of functional group tolerance to



Dearomative functionalization of quinoline position C-4 to access tetrahydroquinolines.

Scheme 73 Ni-Catalyzed remote C(sp²)-H bond arylation of 8-amino protected quinolines.

various functionalized aryl Grignard reagents 164 and aminoquinoline motifs 241 providing the desired arylated products 242 in good to excellent yields (Scheme 73). 173 As a result, rapid access to bioactive and multi-substituted aminoquinolines is made possible.

A practical and efficient solution for the direct amination of heteroaryl Li, Mg and Zn reagents, which has long been considered difficult, was developed in Kürti's group. Heteroaryl organometallics such as quinoline derivatives 243 could be converted to the desired amination products in the absence of directing groups by using an oxaziridine or hydroxylamine reagent 244 in the presence of Cu(I) salts. This new approach for direct electrophilic amination thus seems to represent a general yet simple method that could be used for the efficient production of many structurally complex active pharmaceutical compounds and natural products (Scheme 74). 174

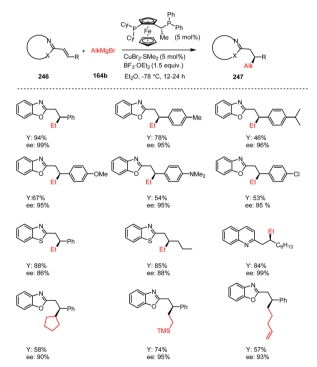
Recently, Harutyunyan's group developed a simple and chemoselective asymmetric addition protocol for the efficient construction of chiral heterocyclic molecules of type 247. Using readily available Grignard reagents 164b and an activating Lewis acid (BF₃·OEt₂), a wide range of β-substituted conjugated alkenyl-N-heteroaromatics 246 could be chemo- and enantioselectively alkylated by copper-catalyzed conjugate addition. This synthetic protocol is of particular interest for applications in medicinal chemistry as it allows the introduction of linear,

Scheme 74 Amination of structurally diverse heteroaryl-based organometallic reagents.

branched, and cyclic alkyl chains as well as a phenyl group at the β-carbon position of the alkenyl N-heteroaromatic compounds. The overall synthetic success depends largely on the interplay of the substrate-activating Lewis acid and the use of highly reactive Grignard reagents in the presence of a diphosphine-stabilized copper catalyst (Scheme 75). 175

The same group subsequently reported a related reaction in which alkyl Grignard reagents 161 are added enantioselectively under mild conditions to vinylogous imines synthesized in situ from sulfonylindoles 248. In the presence of a copper(1) salt and a chiral phosphoramidite ligand, high yields, and enantiomeric ratios of chiral 3-sec-alkyl-substituted indoles 249, which are important structural elements of several drugs and alkaloids, can be obtained (Scheme 76). 176 In addition, the reaction can also be conducted on a larger scale with only 1 mol% of the catalyst and with no loss of yield or enantiomeric purity of the product.

In addition to indole motifs, substituted benzofuran scaffolds, e.g. 251, are also important structural elements in



Scheme 75 Catalytic asymmetric conjugate addition of alkyl Grignard reagents to alkenyl-substituted benzothiazole and benzoxazole.

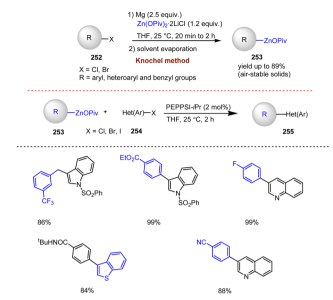
Scheme 76 Cu-Catalyzed enantioselective addition of ethyl Grignard reagents to indole-derived vinylogous imines

Scheme 77 Ni-Catalyzed Kumada cross-coupling of 2-thiomethyl benzofurans with alkyl Grignard reagents.

pharmacologically active compounds, for instance in protein tyrosine phosphatase inhibitors. Thus, Baralle et al. successfully demonstrated an efficient cross-coupling reaction between branched alkyl Grignard reagents 164b and 2-thiomethylbenzofurans 250 employing a Ni-NHC (Ni-N-heterocyclic carbene) complex as the catalyst. Even 3-(4-biphenylyl)-2-alkylbenzofurans can be assembled in one step employing reaction, rendering such compounds accessible in a diversity-oriented manner to be used as intermediates for the preparation of PTP 1B inhibitors (Scheme 77). 177

5 Organozinc-mediated functionalization of fused bicyclic heteroaromatics and their scaffolds

Organozinc reagents are well known for their transmetalation abilities and excellent tolerance towards functional groups.



Scheme 78 Pd-catalyzed C(sp²)-C(sp²)-cross-coupling of solid aryl, heteroaryl and benzylic organozinc pivalates with aryl electrophiles.

Organozinc pivalates (RZnOPiv) have been reported as saltstabilized solid aryl, heteroaryl and benzyl organozinc compounds, which not only exhibit improved resistance to air and moisture, but also show excellent reactivity in Negishi crosscoupling and carbonyl addition reactions. These organozinc compounds are prepared in a one-pot process under mild conditions from aryl and heteroaryl halides (Br, Cl) as well as benzyl chlorides 252 using Mg and Zn(OPiv)2·2LiCl (Scheme 78). 178 The solid materials 253 obtained after evaporation of the solvent can be stored at room temperature under argon for months and may even be exposed to air for short periods.

In a subsequent study, the pool of solid organozinc pivalates was further expanded and their stability in air was improved while their reactivity in Negishi cross-coupling reactions was maintained. The decisive factor here was the use of a directed metalation procedure of arenes and heteroarenes with TMPMgCl·LiCl followed by transmetalation with Zn(OPiv)2 (Scheme 79). 179 The Pd-catalyzed Negishi cross-coupling reaction of aryl halides 257 with heteroaryl-zinc reagents generates the depicted intermediates A-C by means of the well-known reaction sequence of initial oxidative addition to B, subsequent transmetalation with HetAryl-ZnOPiv 256 to form the key intermediate C and final reductive elimination to the desired cross-coupling product 258. This protocol also yielded fine powdered organozinc pivalates 256 that were easy to handle after evaporation of the solvent and that retained most of their activity (>85%), even when exposed to air for 4 h. Moreover, they could be readily used in Negishi cross-coupling reactions with a wide range of electrophiles 257 employing technical grade solvents, which is particularly important for industrial applications.

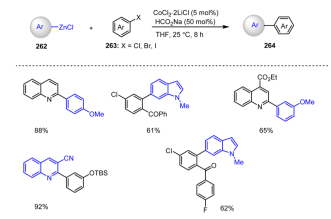
Barl et al. introduced a method for the functionalization of 7-azaindole via a cross-coupling reaction with alkylzinc

Proposed mechanism for Negishi cross-coupling reaction

Scheme 79 $C(sp^2)-C(sp^2)$ cross-coupling reaction of solid and air-stable organozinc reagents and addition of electrophiles enabled by palladium catalysts

reagents. Thus, by using a Pd-catalyzed Negishi reaction between the Reformatsky-type reagents 259 and silyl protected 3-bromo-7-azaindoles 260, targeted 7-azaindole carboxylic acid ester derivatives 261 were formed. Final TBAF-mediated removal of the TBDMS protecting group in THF at 0 °C afforded 2-(7-azaindolyl) carboxylic acid esters 261 in high yields of up to 94% (Scheme 80).180

Scheme 80 Pd-Catalyzed C(sp²)-C(sp³)-cross-coupling of 7-azaindoles and Reformatsky-type reagents.



Scheme 81 Co-Catalyzed cross-coupling reaction of aryl halides and organozinc reagents

A simple, cobalt-catalyzed procedure for the carbon-carbon coupling of halogenated ketones and N-heterocyclic chlorides/ bromides 262 with various (hetero)aryl zinc reagents 263 was developed by Haas et al. (Scheme 81). 181 Different electron-poor and electron-rich functionalized aryl zinc reagents were successfully coupled within a few hours at room temperature under the reported conditions. Moreover, the use of sodium formate HCO₂Na (50 mol%) proved to be decisive for the success of these cross-coupling reactions.

According to Hammann et al., the previously described airand moisture-stable aryl and heteroaryl zinc pivalates 265 can also be successfully subjected to Co-catalyzed cross-coupling reactions with various aryl and heteroaryl halides 266. The proposed method is again characterized by its broad applicability, as both electron-rich and electron-poor electrophiles are tolerated in the presence of CoCl2. Thus, the corresponding reactions of 5-indolylzinc pivalate with 2-bromopyrimidine or 2chloropyrazine, for example, proceeded in good yields (Scheme 82).182

Cobalt-catalyzed electrophilic amination reactions of organozinc pivalates with O-benzoylhydroxylamines under mild and sustained reaction conditions were first published by Chen

Scheme 82 Broadly applicable and robust Co-catalyzed cross-coupling reaction of functionalized organozinc pivalates with aryl halides.

Scheme 83 Co catalyzed amination of heteroaryl zinc pivalates with various O-benzoyl-hydroxylamines and anilines.

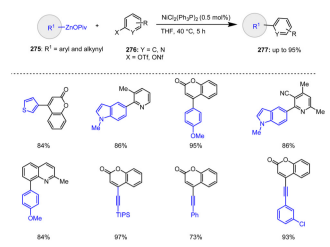
et al. Under the described reaction conditions, C-N crosscouplings between N-hydroxylamine benzoates and heteroaryl zinc pivalates 270 could be carried out at 25 °C within 2-4 h in the presence of CoCl₂·2LiCl (2.5-5.0 mol%), yielding the corresponding tertiary (hetero-) arylated amines in good amounts (Scheme 83). 183 In addition, this method provides access to pharmacologically relevant diarylamine and aryl(heteroaryl)amine units. For example, a clinical candidate for the treatment of tuberculosis was synthesized using this Cocatalyzed amination protocol as the key step.

In 2017, Hammann et al. delineated an improved method for the preparation of air-stable silyl group-protected alkynyl zinc reagents 273 that can be successfully employed in CoCl₂-catalyzed C(sp²)-C(sp) coupling reactions with various heteroaryl halides 272. The described process can be used to functionalize various biologically important heterocyclic scaffolds, such as quinoxalines and quinoline analogs 274 under sustainable conditions with acceptable yields, while incorporating synthetically beneficial alkyne residues (Scheme 84). 184

In 2019, Ni-catalyzed C(sp²)-C(sp²) and C(sp²)-C(sp) crosscoupling reactions between functionalized heteroaryl zinc pivalates 275 and various heteroaryl triflates as well as nonaflates 276 were reported with good to excellent yields. It is noteworthy that only 0.5 mol% of the Ni catalyst was required for these Negishi cross-couplings in THF at 40 °C (Scheme 85). 185

Organozinc pivalates are also useful reagents for the latestage functionalization of small peptides and cyclopeptides by means of Negishi cross-coupling reactions. For example, Leroux et al. were able to successfully couple peptides 278 containing tyrosine or phenylalanine scaffolds with highly functionalized

Scheme 84 CoCl₂-catalyzed C(sp²)-C(sp)-cross-coupling reaction of silyl-protected alkynylzinc pivalates with heteroaryl halides.



Scheme 85 Ni-Catalyzed Negishi cross-coupling of aryl and heteroaryl zinc pivalates with heteroaryl triflates.

aryl and heteroaryl organozinc reagents under Pd catalysis. For this purpose, the corresponding readily available iodotyrosineor iodophenylalanine-containing peptides 279 were sitespecifically reacted with the respective organozinc pivalates 280 in the presence of the Pd catalyst in THF at 25 °C for 4-8 h to provide the modified target peptides 281 (Scheme 86). 186

Cobalt-catalyzed electrophilic amination reactions of anthranilic derivatives 282 with functionalized aryl, heteroaryl, alkenyl and alkyl zinc pivalates 280 can be used to produce complex, condensed N-heterocyclic scaffolds 286 under mild conditions. Li et al. were able to show that the aniline derivatives 283 initially resulting from Co-catalyzed amination reactions can be cyclized under acidic conditions within a cascade reaction to form new complex, condensed quinolines (Scheme 87),187 some of which are of interest for materials

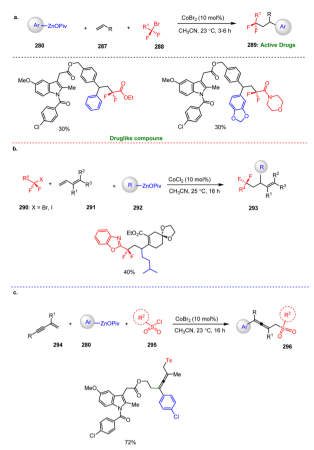
Scheme 86 Pd-Catalyzed late-stage functionalization of cyclopeptides and peptides with (hetero)aryl zinc reagents.

Scheme 87 Co-Catalyzed domino cascade amination reactions of aryl-zinc reagents with anthranil derivatives.

science applications such as organic light-emitting diodes or semiconductors in perovskite solar cells due to their promising properties.

Recently, Lei, Li and co-workers presented a method for the regioselective difluoroalkylarylation of alkenes and 1,3-dienes using functionalized aryl zinc pivalates 280 and difluoroalkyl bromides 288 in the presence of a cobalt catalyst. This one-pot three-component cross-coupling reaction can be used to form difluoroalkylarylated products with predictable regioselectivity and high diastereoselectivity by cascade-like C(sp3)-C(sp³) and C(sp³)-C(sp²) bond formations. With activated and non-activated alkenes 287, the reaction proceeds under remarkably mild conditions with a broad range of substrates using user-friendly solid zinc reagents of excellent functional group tolerance. The process is therefore of particular interest for pharmaceutical applications and late-stage functionalization of drug candidates, as it is easily scalable and allows access to e.g., difluorinated indole scaffolds 289 (Scheme 88a). 188a

In addition, the preparation of solid, salt-stabilized branched alkyl–zinc reagents of the alkyl–ZnOPiv and R_3Si –ZnOPiv type was reported. Due to OPiv coordination, these reagents exhibited not only enhanced stability but also increased reactivity in Co-catalyzed difluoroalkylation reactions of dienoates, allowing modular, site-selective installation of CF_2 and $C(sp^3)$ groups to the double bonds. The twofold $C(sp^3)$ – $C(sp^3)$ cross-couplings of **291** proceeded under mild conditions and were characterized by a broad substrate spectrum and high compatibility with functional groups, thereby enabling an



Scheme 88 Fluoroalkylarylation of drugs scaffolds by cobalt catalysis and organozinc pivalates.

efficient late-stage functionalization of bioactive molecules and fluorinated products 293 (Scheme 88b). 188b

Furthermore, both salt-stabilized aryl and alkyl zinc pivalates **280** show superior reactivity for the cobalt-catalyzed 1,4-carbosulfonylation of 1,3-enynes **294** with sulfonyl chlorides **295** compared to conventional halide-supported organozinc reagents. In this way, highly functionalized α -allene sulfones **296** can be easily and selectively produced *via* the cascade formation of C-C/C-S bonds leading *inter alia* to functionalized bioactive indole scaffolds (Scheme 88c). ¹⁸⁸⁶

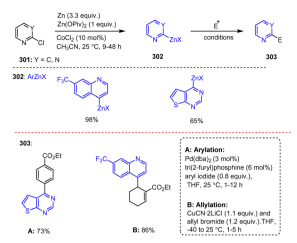
A new method for the preparation of solid, air-stable silyl zinc pivalates from the corresponding silyl lithium reagents by transmetalation with Zn(OPiv)₂ was recently presented by Li and co-workers. The resulting organosilyl zinc pivalates 298 could be successfully used for the silylative difunctionalization of alkenes. Using a convenient chelate-assisted Ni-catalyzed regioselective alkyl- and benzyl-silylation of 8-aminoquinoline-functionalized alkenes 297, the corresponding highly functionalized alkyl silanes 300 could be generated in very good yields. As before, OPiv coordination proved to be crucial for improving the reactivity of the silyl zinc pivalates and provided access to alkyl silanes with a broad substrate spectrum and a high tolerance towards functional groups. The synthetic utility of this alkene carbosilylation sequence was again demonstrated

Scheme 89 Catalytic applications of air-stable silylated zinc pivalates in nickel-catalyzed carbosilylation reactions of various olefins and alkyl halides

by late-stage functionalizations of natural products and druglike molecules as well as by simple, subsequent transformations of the resulting alkyl silanes (Scheme 89). 189

A protocol for converting various readily available Nheteroaryl chlorides 301 into the corresponding organozinc reagents by Co-catalyzed zinc dust insertion in the presence of zinc pivalates in benzonitrile was reported by Kremsmair et al. The resulting organozinc pivalates 302 were successfully reacted with a wide range of electrophiles, such as commercially available aryl halides or acid chlorides, employing Pd or Cu catalysts to give the targeted functionalized heteroarenes 303, as presented in Scheme 90. 190

Recently, Šebesta and co-workers presented a mechanochemical Negishi C(sp²)-C(sp²) cross-coupling reaction of aryl zinc pivalates 304 with aryl(heteroaryl) bromides 305 at room temperature under air. Using this simple Pd-catalyzed process,



Scheme 90 Synthesis of heteroaryl organozing reagents enabled by cobalt catalysis for cross-coupling reactions.

quinoline and benzothiophene derivatives 306 were prepared in good yields and within a short time by ball milling. The reaction tolerates various functional groups and can be carried out on a useful preparative scale (Scheme 91). 191

Melzig et al. reported the functionalization of various thiomethyl-substituted fused N-heterobicycles 307 (e.g., based on isoquinolines, quinazolines, benzothiazoles, or benzoxazole scaffolds) by nickel-catalyzed cross-coupling reactions with functionalized alkyl, aryl, heteroaryl, and benzylic zinc reagents 308. These reactions can be carried out using an inexpensive catalytic system of Ni(acac)2 and the DPE-Phos ligand at 25 °C, as shown in Scheme 92. 192

A mild Negishi cross-coupling of 2-pyridylorganozinc reagents 311 and heteroaryl chlorides 312 was described by Luzung, Patel, and Yin. Using catalytic amounts of Pd₂(dba)₃ (2 mol%) and XPhos as the ligand, many examples of the desired heterocycles 313 were accessible in high yields, complementing the existing coupling reactions for 2-heterocyclic organometallic reagents. The organozinc reagents 311 required for this purpose can be generated in situ under Knochel conditions from 2-pyridyl bromides 310 and i-PrMgCl at room temperature (Scheme 93). 193

Knochel's group has also developed a Ni-catalyzed crosscoupling of heteroaryl halides, including indole, quinoline and quinoxaline derivatives, with aminoalkyl zinc reagents 314 at room temperature. This method allows a convenient one-step

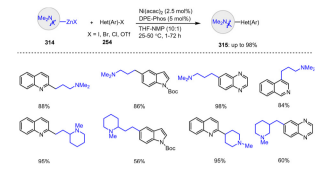
Scheme 91 Mechanochemical Pd-catalyzed Negishi coupling reaction of aryl halides and organozinc pivalates.

Scheme 92 Ni-Catalyzed $C(sp^2)-C(sp^2)$ and $C(sp^2)-C(sp^3)$ cross-coupling reactions of thiomethyl-substituted bicyclic heterocycles with organozinc reagents.

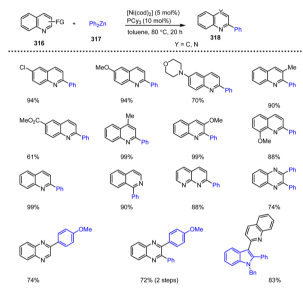
Scheme 93 Pd-Catalyzed cross-coupling of 2-pyridyl organometallic reagents and heteroaryl chlorides with X-Phos ligand.

preparation of various heterocyclic aminoalkyl products **315** (Scheme 94), which are structural components of numerous biologically active compounds. The required aminoalkyl zinc bromides were readily obtained from the corresponding aminoalkyl chlorides by Mg insertion and transmetalation with ZnCl₂. Suitable electrophilic coupling partners comprise aryl, heteroaryl and alkenyl iodides as well as the corresponding -bromides, -chlorides and -triflates. The viability of this method was impressively demonstrated by short total syntheses of the natural products (\pm) -galipinin and (\pm) -cusparin.

Hyodo *et al.* have developed an oxidative atom-economic alternative to the traditional cross-coupling reactions between organohalides and organometallic species enabling a one-step arylation of electron-deficient N-heteroarenes. In this process, functionalized N-heterocycles **316**, such as quinolines, indoles and quinoxalines, were arylated at the most electrophilic site in the presence of a nickel catalyst and PCy₃ as ligand using diaryl zinc reagents **317** as shown in Scheme **95**. ¹⁹⁵



Scheme 94 Direct amino alkylation of heteroarenes *via* Ni-catalyzed Negishi cross-coupling reactions.



Scheme 95 $\,$ Ni-Catalyzed arylation of quinoline scaffolds using Ar_2Zn reagents.

Buchwald and co-workers reported the Negishi crosscoupling of polyfunctionalized heteroaryl halides (Cl, Br, I) with 3,3-disubstituted allylzinc reagents using a previously established precatalytic palladacycle system with remarkable functional group tolerance. The method is characterized by an exceptionally low catalyst loading and mild reaction conditions. In addition, it enables the coupling of aryl/vinyl halides 266 with a wide range of complex heteroaromatic zinc compounds 319 that are often considered difficult. Many of the reported heterocyclic compounds 320 can be found as structural elements in pharmaceuticals (Scheme 96). 196 Moreover, Negishi couplings of 3,3-disubstituted allyl zinc reagents are of particular interest for the preparation of biologically active heteroaromatic compounds with prenyl-like side chains. A successful application of this method was demonstrated with the synthesis of the naturally occurring anti-HIV agent siamenol.

In 2013, Colombe *et al.* developed two protocols for the synthesis of solid and air-stable 2-pyridyl zinc reagents **322/323**, which were designed for bench-top handling and served as practical alternatives to organoboronates in cross-coupling

Scheme 96 Pd-Catalyzed Negishi couplings of hetero(aryl) zinc reagents with heteroaryl halides.

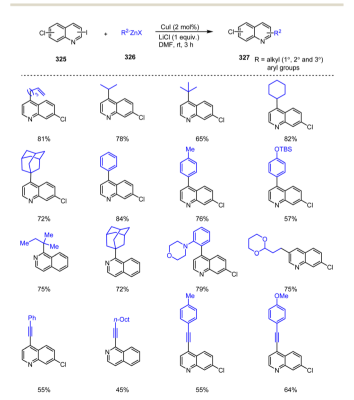
reactions. 197 The organozinc reagents described were subsequently employed in Negishi couplings with aryl halides under Pd catalysis using SPhos as the ligand (Scheme 97). Moreover, the influence of additional stabilizing ligands on the manageability of the organozinc reagents 323 was investigated, with dioxane proving to be particularly suitable for enhancing stability under atmospheric conditions.

Giri and co-workers reported a strategy for the bisfunctionalization of non-activated olefins with two carbon-based units in the 1,2-position using substituted organozinc reagents. For this purpose, suitable alkyl/arylzinc reagents 326 were first prepared and reacted in radical cyclization reactions in situ with LiCl in DMF in the presence of CuI (2 mol%) to give C(sp³)-Cu complexes. The latter were then captured in a second tandem step by a ligand-free Cu-catalyzed Negishi crosscoupling reaction with aryl and heteroaryl iodides 326. The resulting (arylmethyl-)carbocycles and heterocycles 327 were obtained in moderate to high yields and allow the rapid construction of complex biologically active molecular fragments with e.g., indanyl, dihydrofuranyl and indolinyl rings

Scheme 97 Preparation of solid 2-pyridyl zinc reagents for Negishi cross-couplings

from simple and readily available chemical starting materials (Scheme 98). 198

A report was published in 2017 by Greiner et al. on the preparation of polyfunctionalized naphthyridine derivatives 330 by Co-catalyzed Negishi cross-coupling reactions of halogenated naphthyridines 328 with alkyl/aryl magnesium halides, as well as functionalized aryl/heteroaryl zinc reagents 329 in the presence of CoCl₂·2LiCl (5 mol%) and sodium formate as an additive (50 mol%). Some of these naphthyridine derivatives



Scheme 98 Cu-Catalysed cross-coupling of aryl and alkyl zinc reagents with various heteroaryl iodides.

Scheme 99 Co-Catalyzed arylation of aryl(heteroaryl) zinc reagents with iodo-/chloro-naphthyridines.

330 are of interest for materials science applications due to their strong fluorescent properties (PLQE = 20-95%) with tuneable emissions from blue to yellow and long excited state lifetimes (3.8-12.0 ns, Scheme 99). ¹⁹⁹

Knochel and co-workers demonstrated that cobalt-catalyzed electrophilic aminations of heteroaryl zinc reagents 332 and aminating reagents (R¹R²N-OBZ) can proceed under moderate reaction conditions. This electrophilic process also enabled access to various C-N-coupled highly substituted amine products 334 in good to excellent yields using CoCl₂ (5 mol%) in THF at room temperature (Scheme 100).²⁰⁰ Furthermore, a candidate for a new anti-tuberculosis drug (Q203) was produced in a few steps and in very good yields with this method.¹⁸³

Knochel, Hevia and co-workers developed a method for the preparation of polyfunctional diaryl and diheteroaryl zinc species by I/Zn or Br/Zn exchange reactions with bimetallic reagents of the general formula $R_2'Zn \cdot 2LiOR$ (R' = s-Bu, t-Bu, p-tol). With the help of these atom economic I/Zn and Br/Zn exchange reactions, even highly sensitive functional groups such as triazines, keto and aldehyde groups, as well as nitro groups could be tolerated in the substrate. Thus, this protocol allows access to many functionalized (hetero)arenes after scavenging reactions with different electrophiles. Scheme 101 shows an example of such a reaction sequence, in which alcohol 335 is first converted by Et₂Zn into the organozinc species 336. Subsequent treatment with s-BuLi generates the reactive bimetallic reagent 337, which then allows functionalization of the iodoquinoline backbone via nucleophile 338, leading to the desired modified heteroaromatic compound 339 by Cu-catalyzed 1,4-addition to methyl vinyl ketone.

Scheme 100 Amination of solid heteroaryl zinc pivalates with *O*-benzo-ylhydroxylamines.

2ROH
$$Et_2$$
Zn (1 equiv.) [ROZnEt-ROH] $\frac{s\text{-BuLi}(2 \text{ equiv.})}{25 \, ^{\circ}\text{C}, 4 \text{ h}}$ $\frac{s\text{-Bu}_2\text{Zn} \cdot 2\text{LiOR}}{337 \cdot (0.6 - 1.0 \text{ M in Tol})}$ $\frac{s\text{-Bu}_2\text{Zn} \cdot 2\text{LiOR}}{337 \cdot (0.6 - 1.0 \text{ M in Tol})}$ $\frac{s\text{-Bu}_2\text{Zn} \cdot 2\text{LiOR}}{337 \cdot (0.6 - 1.0 \text{ M in Tol})}$ $\frac{s\text{-Bu}_2\text{Zn} \cdot 2\text{LiOR}}{337 \cdot (0.6 - 1.0 \text{ M in Tol})}$ $\frac{s\text{-Bu}_2\text{Zn} \cdot 2\text{LiOR}}{10 \text{ min}}$ $\frac{s\text{-Bu}_2\text{Zn} \cdot 2\text{Li$

Scheme 101 Example for the reparation of a mixed-metal reagent and its application.

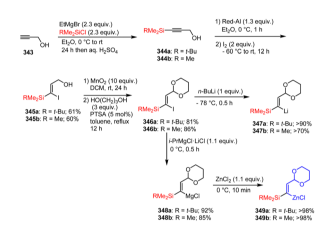
Structural and spectroscopic studies revealed the necessity of forming highly reactive bimetallic lithium bis(alkyl)bis(alkoxy) zincates to generate the polyfunctional aryl and heteroaryl zinc reagents from the corresponding organic iodides or bromides.²⁰¹

Graßl *et al.* were able to show that a broad spectrum of alkyl, aryl and heteroaryl zinc halides can be successfully aminated with highly functionalized alkyl, aryl, and heterocyclic azides in the presence of FeCl₃ (0.5 equiv.) and without further ligand addition. The reaction proceeded at 50 °C within 1 h in good yields and afforded highly functionalized secondary amines such as diarylamine 342 (Scheme 102). This non-toxic Femediated electrophilic amination protocol is particularly suitable for the synthesis of pharmaceutically active amine scaffolds and permits the successful use of peptidic azides.

The Knochel group has developed an efficient approach to various polyfunctionalized 5-, 6- and 7-membered heterocycles, such as furans, pyrroles, quinolines, benzo[b]thieno-[2,3-b]pyridines, naphthyridines, fused pyrazoles and 2,3-dihydrobenzo[c]azepines, using conjugated β -silylated organometallic reagents. The required conjugated alkenyl Li, Mg and Zn reagents 347–349 combine an electrophilic acetal function with two 1,1-bimetallic nucleophilic moieties of well-differentiated reactivity and are readily accessible from propargylic alcohols 343 (Scheme 103). In addition, the silyl group can be converted into various carbon–carbon bonds during the construction of the heterocycles. 203

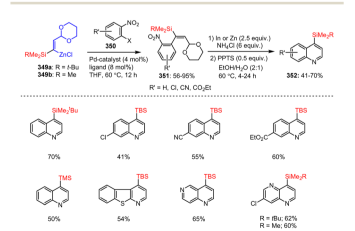
Examples of the preparation of fused 6-membered heterocycles such as quinolines, benzo[b]thieno-[2,3-b]pyridine and

Scheme 102 Fe-Mediated electrophilic amination of functionalized aryl zinc reagents with heteroaromatic azides.



Scheme 103 Preparation of acetal-protected organometallic Li-, Mg-, and Zn-reagents from propargylic alcohols.

naphthyridines relevant for pharmaceutical applications are depicted in Scheme 104. For this purpose, the alkenyl zinc compounds **349** were subjected to Pd-catalyzed Negishi crosscoupling reactions with various 1-halo-2-nitroarenes **350**,



Scheme 104 Synthesis of fused hetero aromatic compounds *via* sequences of Pd-catalyzed cross-coupling, reduction, acidic hydrolysis, and cyclization.

Scheme 105 Functionalization of the quinolines scaffolds *via* I/Si exchange, Negishi-coupling and subsequent re-aromatization.

yielding the corresponding alkenylated nitroarenes of type **351**. Subsequent In- or Zn-mediated reduction and acidic acetal cleavage afforded the desired condensed pyridines and quinoline **352** in 41–70% yields. The use of 3-bromo-2-nitrobenzo[b]thiophene or bromo-nitro-pyridines as coupling reagents thus enabled the rapid preparation of the valuable benzo[b]thieno[2,3-b]pyridines and the corresponding 1,5- and 1,6-naphthyridines.²⁰³

Further functionalization of the quinoline scaffolds through I/Si exchange proved to be extremely difficult for both the 4-TMS and the 4-TBS substituted derivatives 352a/b due to the electron deficiency in the pyridyl ring. However, the 1,2-addition of an organolithium reagent (nBuLi or PhLi) to this pyridyl ring, followed by a scavenging reaction with ClCO₂Et, led to its de-aromatization to form the alkenyl silane 353. Subsequent iodination with NIS and Ag₂CO₃ afforded the 4-iodo derivatives 354 in 95–96% yields, which were then successfully subjected to Negishi cross-coupling reactions with phenylzinc chloride or 1-methyl-1*H*-indol-5-yl-zinc bromide, leading to the coupling products 355. Finally, re-aromatization with KOH/N₂H₄ in ethylene glycol (190 °C, 2 h) afforded the desired 2,4-difunctionalized quinolines 356 (Scheme 105).²⁰³

Since the development of organozinc pivalates as versatile organometallic reagents with high reactivity and excellent air and moisture stability, these compounds have experienced an enormous surge in popularity for organic synthesis applications in recent years. The ease of handling and the absence of sophisticated inert gas techniques make these reagents widely applicable even in traditional (school) synthesis laboratories. In addition, novel protocols that guarantee increased tolerance even to highly sensitive functional groups facilitate the use of these reagents as tools for the construction of complex functional molecules in almost all areas of synthetic chemistry.

6 Selective examples of biologically active bicyclic fused heteroaromatic compounds

Many of the fused heterocyclic aromatic compounds presented so far in this review are fundamental scaffolds of biologically

Table 1 Application of biologically active fused bicyclic heteroaromatic compounds and their scaffolds

S. no.	Compound	Biological activity	Ref. no.
1	F_3CO S CH_3 CH_3	Neuroprotective agent	204
2	H ₂ N N HCI	Inhibitor of MRSA biofilms	205
3		Anti-RSV	206
4	H ₃ C H HN S CH ₃	Antibacterial	207
5	HO OH OH	Antimicrobial	208
6	HO S NH CI CH ₃	Inhibitor of bacterial enzyme	209
7	MeO Me OMe	Inhibitor of RSK2	210
8	HN-S N=O N=O N=O N=O N=O N=O N=O N=O N=O N=O	Inhibitor of 12-LOX	211
9	NH ₂ N · xHCl	Antitumor activity	212
10	CI N H Bn	Inhibitor of SIRT3	213
11	HO HN CF3	Disruptor of biofilm formation in <i>V. cholerae</i> bacteria	214

Table 1 (continued)

S. no.	Compound	Biological activity	Ref. no.
12	N N N N N N N N N N N N N N N N N N N	Inhibitor of pan-phosphodiesterase family	215
13	H ₂ N N N N N N N N N N N N N N N N N N N	Anti-castration resistant prostate cancer and CYP17 lyase-selective inhibitor	216
14	ON SCO2H	Antiviral	217
15	ONH2 HN-N CH3	Inhibitor of Topo and PARP-1	218
16	CI N N	Anti-HIV	219
17	R = N.N.N.N.N.N.N.N.N.N.N.N.N.N.N.N.N.N.N	Autofluorescent antimalarial	220
18	H ₃ C N N N N N N N N N N N N N N N N N N N	Anticancer	221
19	HO O O O	Anticancer	222
20	F OCH ₃	Anticancer	223
21	CI THE NAME OF THE PART OF THE	Anticancer	224

Table 1 (continued)

S. no.	Compound	Biological activity	Ref. no.
22	H ₃ C	Anticancer	225
23	N N CH ₃	Anticancer	226
24		Antiproliferative	227
25	CI-CIS-N-N	Anti-breast cancer	228
26		Anti-influenza virus activity	229
27		Potent inhibitor of ALK mutants	230
28	H ₃ CO NN N N CI	Anticancer	231
29		Anticancer	232
30	H ₂ N CN O	Anticancer	233
31	CY, CI	Anticancer	234
32	H ₂ N COMe	Telomerase inhibitor	235

Table 1 (continued)

S. no.	Compound	Biological activity	Ref. no.
33		Antiproliferative	236
34	CH ₃ CH ₃ CH ₃	Antifungal	237
35	HO	Antiproliferative	238
36	H ₂ N O O	Antitumor	239
37	H ₃ CO N	Inhibitor of microtubule formation	240
38	HO CH ₃ O O CH ₃	Antimitotic activity	241
39		c-Met inhibitor	242
40	HO OH	Leishmanicidal activity	243
41	Br N Me	Antioxidant	244
42		Anti-inflammatory	245
43	R = N N CI	Antidepressant	246

Table 1 (continued)

S. no.	Compound	Biological activity	Ref. no.
44	HO ₂ C OH OH OH	Anti-HIV	247
45	NH ₂ N	Human TLR7 agonist	248
46	NH OH	Human TLR8 agonist	249
47	NH ₂ C ₅ H ₁₁ NH ₂	Human TLR8 agonist	250
48	CH ₃ CF ₃ O CH ₃ CH ₃ CCH ₃ CCH ₃	Antifungal	251
49	N-V-F	Antibacterial	252
50	O N-N CI	Antibacterial	253
51	O N CF3	Antifungal	254
52		Anti-tubercular	255
53	MeO OEt	Anti-protozoal	256
54	OH N-N CF ₃	Antimalarial	257

S. no. Compound Biological activity Ref. no. 55 Anticancer 258 56 Anticancer 259 Antimicrobial agent 260 Potent antagonist 261 262 59 Antidepressant Bactericidal 60 263 Dual CDK2/CDK9 inhibitor and antitumor 61 264 62 CDK9 inhibitor 265 Antiangiogenic and antitumor 63 266 PI3K HDAC inhibitor and antiproliferative 64 267 Fungicidal 268 65 Anti-enteroviral 269 66

Table 1 (continued)

S. no.	Compound	Biological activity	Ref. no.
67	Nan Hand	Antibacterial	270

Table 2 Selective examples of fused bicyclic heteroaromatic system and their scaffolds for material applications

S. no.	Compound structure	Activity/properties	Ref. no.
1	Me F F F F F F F F F F F F F F F F F F F	Photochromo-phore	271
2	OH N N N C 12H25	OLED	272
3	R = N N N N N N N N N N N N N N N N N N	OLED	273
4	$Ph = \begin{bmatrix} & & & & \\ & & & & \\ & & & & \\ & & & &$	PLED	274
5	N NH S N n	Electrochromic devices	275
6	N N N OC6H17	LED	276
7	$C_{a}H_{17}$	Optoelectronics	277

Table 2 (continued)

S. no.	Compound structure	Activity/properties	Ref. no.
8	El ₂ N O	Optoelectronics	278
9	HN—On-C6H13	White light emission	279
10	$C_{12}H_{25}O$	Fluorescent material	280
11	$C_{10}H_{21}$	Organic electronics	281
12	$\bigcap_{S} \bigcap_{R} \bigcap_{R = C_{12}H_{25}} \bigcap_{R = C_{12}H_{$	Optoelectronics	282
13	HO N	Fluorescent sensor	283
14	CI—(S)—CI	Optoelectronics	284
15	C ₁₆ H ₃₃ ——N	Photovoltaics	285
16	Me F F F F F F F F F F F F F F F F F F F	Optoelectronics	286
17		OLEDS	287
18	$H_{2n+1}C_nO$ C_nH_{2n+1}	Liquid crystalline semiconductors	288
19	(CH ₂) ₁₂	Organic semiconductor	289
20	C ₆ H ₁₃ N, N, N	Electrochromic and optical properties	290

S. no.	Compound structure	Activity/properties	Ref. no.
21	R = 2-hexyldecyl	Field effect transistors	291
22	C ₁₂ H ₂₅ - S S S S S-C ₁₂ H ₂₅ - S N-N C ₀ H ₁₃	Optoelectronics	292
23	N C _n H _{2n+1}	Solid state fluorescence	293
24		Chromophores	294
25	NC CN N, N	Chromophores	295
26		OLEDS	296
27		OLEDS	297
28		OLEDS	298
29		OLEDS	299
30	CF ₃	Fluorophores	300
31		Fluorescent material	301

Table 2 (continued)				
S. no.	Compound structure	Activity/properties	Ref. no.	
32	C ₁₁ H ₂₅ O — R OC ₁₂ H ₂₅	Optoelectronics	302	
33		OLEDS	303	
34	O HO Br	Dye for bioimaging	304	
35		Light emitting diodes	305	
36	N.NH	Photo-switches in photonics	306	
37		OLEDS	307	
38	Ph—SSPh	Organic field effect transistors	308	
39	OHC—S———————————————————————————————————	Optical chromophores	309	
40	$C_{12}H_{25}$ $C_{12}H_{25}$ $C_{12}H_{25}$ $C_{10}H_{21}$ $C_{10}H_{21}$ $C_{10}H_{21}$	Field effect transistors	310	
41	NC COOH	Solar cells	311	
42	$C_{2}H_{5} \sim C_{6}H_{13}$	P-type organic semi-conductors	312	
43	C ₂ H ₁₇ O OC ₂ H ₁₇ (S) S S S S S S S S S S S S S S S S S S	Photovoltaics	313	

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S. no.	Compound structure	Activity/properties	Ref. no.
44	Bu-N-S-S-S-S-N-Bu	N-type molecular semi-conductors	314
45	$OC_8H_{17}O$	Electric device applications	315
46	$C_{10}H_{21}$ S N N S $C_{10}H_{21}$	Organic field transistors	316
47	Ar ————————————————————————————————————	Photo-luminescence	317
48	Ph N-Ar R = 9,9-diethylfluoren-2-yl; Ar=1-napthyl	Electro-luminescent materials	318
49	C ₉ H ₁₉	OLEDS	319

active natural products and drugs. While several key heterocyclic structures can be isolated from natural sources or built up stepwise using classical synthetic routes, over the last three decades research groups have been increasingly engaged in the development of synthetic methods that allow rapid, modular access to these scaffolds. Highly functionalized condensed heteroaromatic compounds are characterized by diverse biological activities and are therefore of central importance as fundamental lead structures in various fields of medicinal chemistry and the pharmaceutical industry. For example, active substances for the treatment of malaria, tuberculosis, HIV, cancer, and depression, to name just a few applications, are based on different suitably functionalized heteroaromatic scaffolds. Synthetic methods that facilitate the construction of heteroaromatics and their modification through late-stage functionalization are therefore of great importance for the further development of drugs and for medical advances. The following section provides an overview of various condensed heteroaromatic motifs found in active ingredients, along with their pharmaceutical and biological applications. 204-270 (Table 1)

7 Selective examples for bicyclic heteroaromatic system-based material applications

Functionalized condensed heteroaromatic compounds are not only of interest due to their prevalence in biologically active compounds. Rather, highly π -conjugated sulfur-, oxygen- and

nitrogen-based fused heterocycles show great application potential for materials sciences and can be used for instance in the development of organic light-emitting diodes (OLEDs), polymer LEDs, organic semiconductors, field-effect transistors, photoelectrically luminescent materials and solar cells. Further potential areas of application can be derived from the table compiled in this section. 271-319 (Table 2)

8 Conclusions

This review highlights important synthetic advances in the functionalization of 5- or 6-membered ring-fused bicyclic heteroaromatic frameworks using organo-Li, Mg and Zn reagents. The syntheses presented include complex sequences based on direct selective metalations, halogen/metal exchange reactions, oxidative metal insertions and transmetalation processes. Protocols for efficient preparations of different functionalized organometallic reagents with aryl, heteroaryl, alkenyl, alkynyl and alkyl groups allow their use in high-performance transition metal-catalyzed cross-coupling reactions with various electrophiles, which makes such reaction sequences widely applicable in synthesis programs and for the development of active pharmaceutical ingredients. Thus, selective functionalization of important heterocyclic scaffolds such as quinoline, indole, benzofuran, benzothiophene, benzoxazole, benzothiazole, benzopyrimidine, anthranil, thienothiophene, coumarin, chromones, quinolones, phthalazines and their condensed heterocyclic scaffolds can now be carried out in almost any

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desired position. This overview thus underlines the importance of modern synthetic technology with regard to functional group tolerance, cost efficiency and sustainable reaction conditions in the production of heterocyclic bioactive drugs and natural products.

Data availability

The authors confirm that the data supporting the findings of this study are available in the article.

Conflicts of interest

There are no conflicts to declare.

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