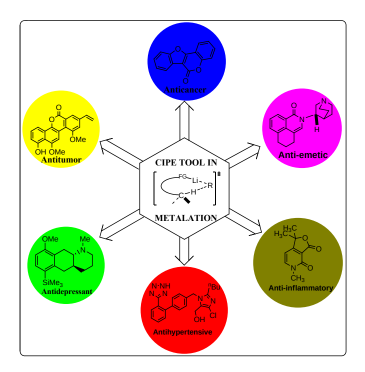
# Open Access

# Complex-Induced Proximity Effect in Lithiation: Unveiling Recent Potentials in Organic Synthesis of Biologically Relevant Heterocyclic **Compounds**

Olayinka O. Ajani\*, Michael O. Shomade<sup>1</sup>, Abiola Edobor-Osoh<sup>1</sup>, Christiana O. Ajanaku<sup>1</sup> and Alice O. Ajani<sup>2</sup>

Abstract: Reactions that convert carbon-hydrogen (C-H) bonds into carbon-carbon (C-C) or carbon-heteroatom (C-Y) bonds are attractive tools for organic chemists, potentially expediting the synthesis of target molecules via functional group interconversion. More explorative studies have shown Complex Induced Proximity Effect (CIPE) to be a solutionprovider for the synthesis of bioactive compounds. This might act as excellent pathfinders to new drugs for combating microorganisms' resistance challenges to old existing drug. So, a constant review into CIPE and lithiation chemistry is crucial because they offer excellent pathways to new heterocyclic compounds which are essential agents in drug design and discovery.



**Keywords:** Aziridine, bioactive heterocycles, cipe mechanism, enantioselective, lithiation, stereoselectivity.

### 1. INTRODUCTION

Over the years, there has been a considerable attention and interest in the aromatic metalation procedure, especially

<sup>&</sup>lt;sup>1</sup>Department of Chemistry, Covenant University, Canaanland, Km 10 Idiroko Road, P.M.B. 1023, Ota, Ogun State, Nigeria

<sup>&</sup>lt;sup>2</sup>Nigerian Stored Products Research Institute, Onireke, Ibadan, Oyo State, Nigeria

<sup>\*</sup>Address correspondence to this author at the Department of Chemistry, Covenant University, Canaanland, Km 10 Idiroko Road, P.M.B. 1023, Ota, Ogun State, Nigeria; Tel: +2348061670254; E-mail: ola.ajani@covenantuniversity.edu.ng

lithiation reaction which is an efficient synthetic method for heterocycle substitution [1]. A useful strategy for intra molecular cyclization and synthetic information transfer is by using proximity and shape of one part of a molecule to stereo selectively control reaction occurring nearby [2]. This idea in lithiation is known as complex induced proximity effect (CIPE). It originated in 1986, when novel carbanions formation by organolithium bases was described as a twostep process [3] in which the formation of a prelithiation complex brings reactive groups into proximity for directed deprotonation [4]. It is however, very interesting to note that relative ease of lithiation is attributed to a prelithiation phenomenon. This prelithiation phenomenon that occurred in the transition state [2] before deprotonation was achieved was termed the complex induced proximity effect, CIPE [5]. This factor provided alternative way to C-C bond formation which was facilitated by heteroatoms of the substituted organolithium reagents [6]. Among the most widely used are metalated sulfones [7], acyl anion [8], acetic acid dianion [9], cyanohydrin anion [10], homoenolate anion [11] and dithianes [12]. In this regard, reactions in which lithiated secondary amides provide CIPE control of regioselectivity and stereoselectivity have been a recurrent theme in this area [13].

Complex induced proximity effect provides the chemists with a well equipped synthetic toolbox for functionalization of the aromatic ring and intramolecular cyclization [14]. Organolithium synthesis generally involves an alkyllithium-promoted halogen-lithium exchange or the simple direct deprotonation of the most acidic and stabilizating position of the heterocyclic nucleus with alkylithiums or lithium amides [15]. This is a stabilization which can be produced by an adjacent atom or group in the so-called "directed ortho-metalation" (DoM) [16]. The potential of DoM, as amplified by the versatile lithium species, has been largely exploited in the total synthesis of natural products bearing bioactive heterocyclic cores [16]. Heterocycles are found in all kind of compounds of interest in medicinal chemistry research. They could be inserted in other class of compounds to boost therapeutic efficiency. Among all the possible synthetic methods of achieving this insertion into any structure, probably the use of lithiation chemistry is the most direct strategy [17]. Although, the principal enabling force in the development of organolithium chemistry is the commercial availability of inexpensive stable solutions of *n*-butyllithium [18] and the more potent and selective s-BuLi and t-BuLi, which on the contrary, are expensive and difficult to handle [19]. Nevertheless, the scope of the metalation reaction has been expanded by the use of complexing and chelating reagents such as hexamethylphosphoric triamide (HMPA), N,N'-dimethylpropyleneurea (DMPU) and tetramethylethylenediamine (TMEDA) which increase the rate of metalation and thus, extend the range of compounds which can be deprotonated [20].

The aim of this present work is to evaluate recent advance in the CIPE approaches to new compounds via lithiation chemistry covering from year 2000 to 2014. Hence, this review presents CIPE as a resourceful tool in metalation chemistry for the designing of valuable scaffolds which may serve as great opening to new drug discovery and development.

#### 2. MECHANISM IN CIPE

### 2.1. CIPE: The Underlying Mechanism Behind DOM Methodologies

The direct metalation group (DMG) is typically a Lewis basic moiety that interacts with the Lewis acidic lithium cation allowing for deprotonation by the alkyl-lithium species from the nearest ortho-position on the arene [21]. Applications of CIPE to synthetic goals have been particularly well-developed for directed ortho metalation (DOM) methodologies which involved the use of DMGs as the coordinating functional group necessary for CIPE [22]. Although, DMGs do not function alone in determining the site of metalation, but sterics and other functional groups on the arene also have a great deal of influence [23]. The general CIPE is illustrated in 4, a lithiation/substitution sequence as shown in the Scheme (1) below. The coordinative interaction of 1 with an appropriate organolithium reagent provides the complex 2 which upon subsequent directed lithiation via transition state 3 affords lithio species 4 [24]. The quenching of 4 by addition of an electrophile is highly favoured over traditional electrophilic substitution to achieve 5 because of the regioselective preference displayed. Cases that can be cited for CIPE include not only deprotonative mono- and dilithiations but hetero atom-lithium exchanges. inventive displacements' and additions. CIPE process appears to arise in a wide variety of reactions of organolithium compounds [17]. Thus, we have herein reviewed recently reported carbolithiations which are consistent with the general process outlined in Scheme (1).

## 2.2. Two Schools of Thought for One Mechanism: Lithiation

Selective ortho lithiation of aryl rings bearing heteroatom containing functional groups is a powerful synthetic strategy in organic and organometallic synthesis. Two major mechanisms theorized to drive ortho-lithiations: "Coordination only" substituent coordinates "complexes" with organolithium reagent to increase kinetic basicity, and directs deprotonation to ortho position. A typical example of which could be explained by the lithiation of dimethylbenzylamine 6 which afforded compound 7 and (ii) "Acid-base" - inductive and/or resonance effects from heteroatomic substituent make ortho proton more acidic [25]. A typical illustration of this mechanism could be explained by why the lithiation of pyrazole 8 gave the 5lithio derivative 9 and not the 3-lithio counterpart (Scheme 2). Some lithiations are driven entirely by one factor or the other, but the majority of lithiations occur by a combination of both. Organolithiums were thought to coordinate to heteroatoms in  $\alpha$ -lithiation of heterocycles [26]. One thing is sure, the mechanistic proposal must explain two main observations which depend on heteroatomic substituent; they are: (a) increase reactivity of substrate and (b) direct regioselectivity of deprotonation.

### 2.3. CIPE in $\alpha$ -Lithiation of Amine

The access to  $\alpha$ -lithiated amine *via* direct deprotonation has been established to occur by CIPE mechanism through a pathway of an intermediate complex in which the organolithium compound is pre-coordinated by the amine ligands [27]. Notwithstanding, the bottleneck associated with

DMG 
$$+ (RLi)_n$$
  $+ (RLi)_n$   $+ (RLi)_n$ 

Scheme (1). A lithiation/substitution strategy showing influence of CIPE.

direct α-lithiation of amine was earlier reported to be due to the destabilization of the carbanions by the interaction with the lone pair electron density of the adjacent nitrogen atom [28]. This had caused unmet demand in the formation of the templates as a desirable synthetic route to polar heteroorganometallics [29]. However, CIPE has been projected as the basic concept in the predicted and investigating basic metalation of 11 which was obtained from partial deprotonation of bis(3-methyl-1,3-diazacyclohex-1-yl)methane 10 (Scheme 3) [30]. This close proximity of the carbanion C(23) (tert-butyl group) and the hydrogen atom H(18a) of 10 is also represented as an atomic interaction line in the charge density topology [30]. This atomic interaction line has a bond critical point of low density  $(0.046 \text{ e Å}^{-3})$  and positive Laplacian (0.35 e Å<sup>-5</sup>) expressing its closed shell nature as shown in Fig. (1) [30]. Kamp and coworkers also demonstrated that the concept of a complex-induced proximity effect can be underlined and supported with charge density topology features [30].

# 2.4. Kinetically Enhanced Metalation by CIPE Mechanism

Directed lithiations are a topic of considerable deliberation. The selective deprotonation of an ortho or benzylic position assisted by an electron withdrawing group bearing electron lone pairs may be explained through the

complex-induced proximity effect (CIPE) model [31]. This mechanism considers the lithiation as a two-step process. First, the coordination of the lithium cation of the base with one Lewis basic heteroatom of the substrate results in the formation of a complex [24]. This complex brings the carbanionic center of the base close to the acidic proton, thus favoring the transfer of the proton in the second step. Orthodirected deprotonations have been interpreted by an alternative mechanism involving a one-step reaction [32]. In this model, the metalation was described as a kinetically controlled transformation for which the term "kinetically enhanced metalation" has been coined [32]. It was of investigated using the reaction N-alkvl-Nbenzyl(diphenyl)phosphinamides 12 with s-BuLi leading to de-aromatized products via the isotopic-labeling and NMR study of the mechanism. This was one of the crucial cases in which pre-lithiation complexes have been structurally identified in the directed deprotonation of a phosphorusbearing substrate (Scheme 4) [32].

# 2.5. Trans-Esterification Mechanistic Occurrence by CIPE

Rate enhancements of trans-esterification associated with a second Lewis base centre can be considered as a manifestation of CIPE, which have shown to play an important role in a number of reactions of organic

(i). Typical Coordination Only Mechanism

(ii). Typical Acid-Base Mechanism

**Scheme (2).** Two schools of thought in lithiation mechanism.

Scheme (3). Formation of 11 by partial deprotonation of bis(3-methyl-1,3-diazacyclo hex-1-yl)-methane 10.

Scheme (4). Dearomatization of diphenylphosphinamides 12 through anionic cyclization. The isotopic-labeling study was conducted using the general reaction conditions for the synthesis of the benzoazaphospholes 15 which involved the treatment of a THF solution of 12 and 6.0 equiv of DMPU (or HMPA) with 2.5 equiv of s-BuLi at -90 °C for t<sub>1</sub> min, followed by the addition of the appropriate electrophile and stirring for  $t_2$  min at the same temperature.

compounds of lithium [17]. The significantly increased reactivity in trans-esterification displayed by the esters containing second Lewis base centre proximal to the ester functionality has been attributed to the CIPE as reported by Jackman and co-workers, 1991. Based on this mechanistic assumption, the free energies of activation for the transesterification of six 3,5-dimethylphenyl esters 16a-f possessing a second Lewis base centre was investigated wherein the predicted and the observed values were compared (Table 1) [33]. It is therefore possible that the presence of other Lewis base centres in the acid moiety of the esters might provide more effective means of attachment [34], even though the electrophilic activation of the carbonyl group associated with coordination of its oxygen to lithium would be lost [33]. The essential role of the pre-equilibrium step is to attach the ester to the tetramer in order to effect electrophilic attack of the putative phenolate ion on the carbonyl carbon atom [33].

#### POLYFUNCTIONAL **ORGANOLITHIUM** 3. REAGENTS VIA CIPE

The use of Lewis bases to increase the reactivity of lithium organics is an important tool in synthetic chemistry [34]. N,N,N',N'-tetramethyl ethylenediamine (TMEDA), one of the most powerful and most often used Lewis bases, is known to undergo a direct R-lithiation, the regioselectivity of which depends on the used deprotonation agent [20]. The α-lithiation of TMEDA in the presence of t-BuLi was reported to give, after electrophilic quenching, a tridentate ligand 17 according to Scheme (5) [28]. The presence of heteroatoms in close proximity to the carbon-lithium bond facilitates the formation of an organolithium species as long as the various functional groups are tolerated [17]. The direct lithiation with lithium powder in the presence of a catalytic amount of 4,4'-di-tert-butybiphenyl (DBB), as popularized, proves to be a very convenient method for preparing a broad range of polyfunctional organolithium reagents [35]. Thus,

imidoyl 18, carbamoyl 19a, or thiocarbamoyl 19b lithium compounds which are formerly difficult to prepare, could now be generated using this facile approach which was made possible by CIPE. The direct preparation of acyllithium compounds such as 20, either by a direct low-temperature route from RLi/CO or a lithium-tellurium exchange reaction, has been successfully performed [36]. In these direct preparation methods, the acyllithium species are generated in the presence of an electrophile [37].

# 4. UNVEILING SYNTHETIC POTENTIALS OF METALATION CHEMISTRY

# 4.1. Diversity of Aziridine and its CIPE Accomplice

Aziridines are important compounds because of their widespread use in organic synthesis and their presence in many natural products and biologically active molecules [38]. They have been extensively investigated either from the synthetic or reaction points of view. Concerning the

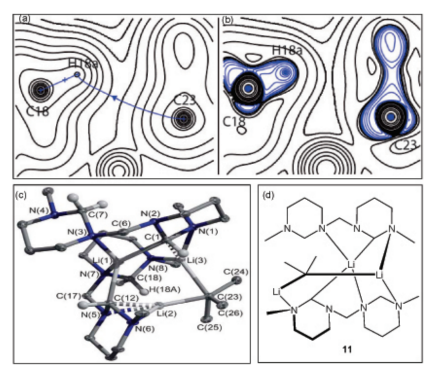


Fig. (1). (a) Electron density map (b) and Laplacian in the C918)-H(18a)-C(23) plane of 11, showing interaction between the tert-butyl carbanion and the proton next to be abstracted. (c) Molecular Structure of 11 in the solid state. Only hydrogen atoms bound to the endocyclic NCN units are shown. Topological links to the Li atoms drawn on the basis of distance criteria are dashed. (d). Chemical structure of 11 formed by partial deprotonation of bis(3-methyl-1,3-diazacyclohex-1-yl) methane 10.

Table 1. Observed and predicted free energies of activation for the transesterification, at 30 °C in DME, of 3,5-dimethylphenyl esters possessing a second lewis base center.

	OH + O KOH (109)  Me Me CI stirred at then at rt	0 °C for 1h	O O R Me					
$\Delta G^*$ (kcal mol <sup>-1</sup> )								
Comp. No	R	obsd	pred	diff				
16a	CH <sub>2</sub> OCH <sub>3</sub>	19.6	21.3	1.7				
16b	CH <sub>2</sub> CH <sub>2</sub> OCH <sub>3</sub>	21.6	22.7	1.1				
16c	CH <sub>2</sub> CH <sub>2</sub> CH <sub>2</sub> OCH <sub>3</sub>	22.5	22.8	0.3				
16d	2-tetrahydrofuryl	20.4	21.7	1.3				
16e	CH <sub>2</sub> N(CH <sub>3</sub> ) <sub>2</sub>	20.9	22.2	1.3				
16f	CH <sub>2</sub> (2-pyridyl)	19.9	22.1	2.2				

Comp. No = Compound Number; obsd = observed; pred = preserved; diff = difference

reactivity, the most common transformations of these springloaded three-membered ring systems are the ring-opening reactions that can be initiated by both electrophilic and nucleophilic reagents [39]. Numerous synthetic approaches have been developed for the preparation of aziridine due to their widespread applications in medicinal chemistry. Some of these methods include aza-Darzen approaches [40], transfer of nitrogen to olefins [41], addition across the carbon-nitrogen double bond of aziridines [42], and more recently, metalation approach via CIPE [43, 44].

### 4.1.1. Stereoselective lithiation of N-Alkyl-(o-tolyl)aziridine in Isochromans

Six membered-ring oxygen-bearing aromatic heterocycles with isochroman and related skeletons occur in nature and among bioactive compounds of interest, including drugs (medicines, agrochemicals, etc.) and drug candidates [45]. The lateral lithiation of ortho-tolylaziridine 21 followed by electrophile trapping gave the intermediate orthohydroxyalkylated aziridines 22 which has been recently reported as an excellent route towards a range of bioactive isochromans 23 via acid-catalyzed cyclization approach (Scheme 6) [42]. The results of the lithiation/trapping sequence above clearly demonstrated the directing group ability of the aziridine ring [42]. It is likely that the nitrogeninduced stabilization in 21-Li, and CIPE could act synergically making the lateral benzylic position the kinetically and thermodynamically favored one [46]. This work reported a new and convenient methodology for the preparation of ortho-functionalized aziridines based on the benzylic lithiation of simple and easily available otolylaziridines [47]. It is, indeed, worth pointing out that the lithiation of the related acyclic derivative, 2-N,Ndimethylaminomethyltoluene, is comparatively much slower requiring more than 6 h at room temperature for complete deprotonation [42].

### 4.1.2. Regioselective lithiation of Aziridine

Aziridines are widely used as versatile building blocks for the synthesis of a variety of biologically and pharmaceutically important molecules [48]. Data from the literature indicate that N-alkyl-2-phenylaziridines undergo smooth ortho-lithiation [49]. In contrast, trans-N-alkyl-2,3diphenylaziridines undergo exclusive R-lithiation with a stereochemistry strongly depending on the coordinating ability of the solvents [43]. The aziridino-borane complexes 25a,b were prepared by treating 2-phenylaziridines 24a,b with 1M THF solution of BH<sub>3</sub>·THF complex. When 25a was reacted with s-BuLi (1.2 equiv) in THF at -50 °C for 2 h, the corresponding aziridinyllithium was generated as proved by its trapping with D<sub>2</sub>O to furnish complex **26a** (Scheme **7**). The BH<sub>3</sub> removal was easily achieved by adding a small amount of H<sub>2</sub>O at room temperature and the corresponding 2-deuterated aziridine 27a was recovered almost

**Scheme (5).** Synthesis of tridentate ligand **17** and structures of other selected ligands.

Scheme (6). Preparation of isochromans 23 via lithiation and acid-catalyzed cyclization.

Scheme (7). Synthesis of Aziridino-borane complex 26a,b and its double functionalization.

quantitatively and as a single stereoisomer after the work-up [43]. This showed the ability of the aziridino group to act as a directing metalation group (DMG) [49].

### 4.1.3. Stereospecific Lithiation of Arylaziridine

Normally, the presence of an electron withdrawing group (EWG) on the nitrogen or the carbon atoms of the heterocyclic ring is crucial for successful metalation [50]. Recently, lithiation/electrophile trapping of unsubstituted and 2-alkylsubstituted N-Bus-aziridines has been reported [51]. However, no efficient methods for the  $\alpha$ -lithiation of N-Bus-substituted monoarylaziridines have been disclosed. In view of this, Musio and coworkers developed stereospecific lithiation route for the synthesis of optically active trisubstituted arylaziridines and further assessed the role of N-Bus in the lithiation reaction [44]. The enantiomerically enriched N-Bus-2-phenylaziridine (R)-28 was prepared from (-)-phenylglycinol by a high-yielding sequence that involved N-sulfinylation, oxidation, o-tosylation, and cyclization (Scheme 8). Upon lithiation / trapping sequencing of (R)-28, a stereospecific route was provided for obtaining α,αdisubstituted aziridines 30a-e as single enantiomers (er > 98:2). This indicates that the intermediate organolithium (R)-**29** is configurationally stable [44].

# **4.2.** Enantioselective Carbolithiation in Heterocyclic Construction

The carbolithiation reaction has attracted considerable interest among synthetic organic chemists, as it offers an attractive pathway for the efficient construction of heterocyclic compounds of medicinal interest [52]. These reactions can be carried out either in inter- or intramolecular fashion [50].

- (a). Taking advantage of CIPE, Woltering and coworkers reported that the deprotonation of (carbamoyloxy)methyl-N-cinnamyl piperidine 31 with sbutyllithium/(-)-sparteine (L1), and subsequent anionic 5exo-trig cyclization, leads to the formation of (2R,8aR)-2benzyloctahydroindolizin-1-yl 2,2,4,4-tetramethyloxazolidine-3-carboxylate 32 with high diastereomeric and enantiomeric ratios, in moderate yield. (Scheme 9) [53]. It is important to note that the resulting benzyllithium can also be trapped with electrophiles in order to achieve other synthetic manipulations [18].
- (b). Barluenga and coworkers reported the enantioselective synthesis of benzo fused furan derivative *via* intramolecular carbolithiation with special regard to CIPE [54]. Enantiomerically enriched 2,3-dihydrobenzofurans

Scheme (8). Synthesis of optically active trisubstituted arylaziridine 30a-e.

**Scheme (9).** Carbolithiation of cinnamyl piperidine to indolizidine derivative **32**.

R<sup>2</sup>

$$R^2$$
 $R^2$ 
 $R^2$ 

Scheme (10). Intramolecular cabolithiated synthesis of 2,3-dihydrobenzofurans 34.

34a-d were obtained in moderate to good yields and high enantiomeric purity from intramolecular carbolithiation of 3,5-disubstituted 2-(allyloxy)-1-bromobenzene **33a-d** by using (-)-sparteine (L1) as chiral ligand, and diisopropyl ether as solvent (Scheme 10). The resulting organolithium can be trapped with several electrophiles [54]. However, the presence of a substituent at the 3-position of the aromatic ring  $(R^1 \neq H)$  is very crucial otherwise, the aryllithium intermediate undergoes a tandem carbolithiation-γ-elimination leading to enantio-enriched 2-cyclopropylphenols [55].

(c). The presence of a substituent in the position ortho to the lithium atom of lithio species generated from 35 leads to lower yields and the opposite enantiomer (S)-3-methyl indoline (S)-36 with low enantiomeric excess (22% ee) when (-)-sparteine (L1) is used (Scheme 11) [56]. However, Mealy and coworkers has proved that the (1R,2R)-N,N,N',N'-tetramethylcyclohexane-1,2-diamine (**L4**) to be a more efficient ligand for lithium, leading to (R)-1-allyl-3,4dimethyl indoline (R)-36 in 70% ee [56].

(d) The lithiation sequences from 38a-c to 37 and 39 were quite illustrative as shown in Scheme (12). Treatment of 38a with LDA afforded 37, the result of a directed lithiation and an anionic ortho-fries rearrangement [22]. In the case of 38b or 38c, the normal site of metalation was

Scheme (11). Comparative study of efficiency of ligands in synthesis of 36.

Scheme (12). Remote carbolithiation toward synthesis of dibenzopyranones 40.

Scheme (13). Synthesis of bioactive natural product Coumestan, 45.

blocked and deprotonation of the remote ring became thermodynamically favoured by the initial formation of a complex between the directing group and the organolithium reagent [50]. Rearrangement and acid catalyzed cyclization followed to provide the dibenzopyranones **40** in good yields [22].

(e) Aliyenne and coworkers has developed a more convenient method for the preparation of chiral saccharins than the earlier one by Soubh and coworkers [57]. Aliyenne and coworkers reported that their process was driven by CIPE and constitutes a mild method for the LDA-HMPA mediated regiospecific conversion of *N*-arylsulfonyl oxazolidin-2-ones **41a**–**f** readily available from optically pure amino acids into novel chiral analogues of saccharins **42a**–**f** (Table **2**) [58]. The resulting optically active benzisothiazolinone 1,1-dioxides **42a**–**c** and naphthaiso thiazolinone 1,1-dioxides **42d**–**f** were obtained in good yields [58].

# 4.3. Natural Product Synthesis

(a) Recently, Tricotet and coworkers successfully applied vinyl-lithiation/electrophile trapping/ring closure reaction sequence for the synthesis of the medicinally important natural product Coumestan **45** [59] which is a potential

anticancer agent [60]. Vinyl lithiation of bis-*ortho*-methoxy *cis*-stilbene **43** followed by  $CO_2$  quench provided routine access to intermediate **44** upon which demethylation with BBr<sub>3</sub>, treatment with base, and oxidative cyclization completed the synthesis of **45** as shown in Scheme (**13**) [59]. The vinyl lithiation which resulted into the formation of key intermediate **44** was made possible by kinetically favoured CIPE mechanism.

- (b) The combination of amide and *o*-carbamate DreM strategies was illustrated in the synthesis of dengibsin **49** [24], a member of the rare class of naturally occurring fluorenones [61]. Thus, the differentially protected biaryl *o*-carbamate **46** (available by Suzuki–Miyaura cross-coupling followed by DoM-mediated silylation) was treated with excess LDA under vigorous conditions to afford the amide **47**. Protection and desilylation leads to **48**, which upon the second LDA-mediated reaction (under milder conditions) and subsequent de-isopropylation afforded the natural product **49** as shown in Scheme (**14**) [22, 24].
- (c) Compound **54** which is coded as RS-42358 and its analogs are a class of 5-HT3 receptor antagonists that show promise as anti-emetic agents. The total synthesis of this biologically active compound **54** was achieved by Kowalczyk as shown in Scheme (**15**) [62]. This involved

Table 2. Cyclization of 3-N-Arylsulfonyloxazolidin-2-ones 41a-f to 42a-f.

		O R S-N O O 41a-f	i. LDA, HMPA ii. NH₄Cl	./THF/-78 °C	Ar N—	R —ОН	
D 1 4	R	Ar	[α] <sub>D</sub>	Typical Conditions		37.00	T71 11/0/
Product				Temp. °C	Additive	Mp °C	Yield(%)
42a	Me	C <sub>6</sub> H <sub>5</sub>	+45	-78	НМРА	132–134	69
				-78	TMEDA	_	n.r.
42b	s-Bu	C <sub>6</sub> H <sub>5</sub>	-53.57	-78	HMPA	73–75	65
42c	Bn	C <sub>6</sub> H <sub>5</sub>	-55.0	-78	HMPA	107–109	71
42d	Me	2-Naphthyl	+40	-78	НМРА	156–158	65
42e	s-Bu	2-Naphthyl	+38	-78	HMPA	141–143	65
42f	i-Pr	2-Naphthyl	+55.55	0	HMPA	99–101	62

n.r. = no reaction

Scheme (14). Synthetic route to naturally occurring dengibs in 49.

Scheme (15). Total synthesis of anti-emetic agent 54 via lateral lithiation.

synthetic conversion of acid 50 to the intermediate N,Ndiethyl substituted amide 51 which was lithiated, followed by formylation to afford 52. Lateral lithiation was the key step towards closure of the intermediate 53 which upon condensation with amine generated the targeted compound **54** [62].

(d) As a foray into the synthetic potential of metalation, the versatile CIPE concepts of Beak & Meyers led, as a direct consequence, to the establishment of Directed remote Metalation (DreM)-induced reaction to afford naphthobenzopyrone 55 which is an essential building block to the antitumor natural product defucogilvocarcin V, 57 via the intermediate Stille product 56 (Scheme 16) [63]. Defucogilvocarcin V 57 is also reported as a new antibiotic from Streptomyces arene with potential activity against lung cancer cell line [64]. In these processes, the effective use of the carbamate moiety as a carbonyl dictation equivalent was demonstrated [65]. These examples demonstrated that the DoM reaction has not only a recognized potential in the modification of a DMG's ortho-environment but, through its privileged connections with rapidly growing methods (metalcatalyzed cross coupling, RCM, DreM, direct arylation) imposes the choice and exploration of new synthetic routes [66]. In a similar manner, enantioselective total synthesis of (-)-hyperforin was reported in 18 steps starting from ortholithiation of 1,3-dimethoxybenzene [19].

(e) An iodination reaction for the synthesis of 6-aza-Ltryptophan [67] and 4-alkoxy carbonylations in the preparation of pyridopyrimidinones [68] are recent examples of synthetic application of lithiation technique. In the case of 3-bromopyridine, LDA has been used as metalating agent for the generation of the corresponding brominated 4pyridyllithium reagent, which has been employed in an addition reaction to acrolein in the synthesis of restricted

Scheme (16). DreM-induced synthetic approach to defucogilvocarcin V, 57.

nicotine analogues [69]. An illustrative example of a DoM reaction for preparing a synthetically useful 4-pyridyllithium species is the one-pot synthesis shown in Scheme (17). Thus, 3-pyridyl carboxylic acid 58 was treated with *n*-butyllithium to give the corresponding carboxylate anion 59, and further lithiation with LiTMP afforded the organolithium species 60. Subsequent reaction with acetone gave the dilithium salt 61, which was transformed into a lactopyridone after acid treatment. Final *N*-methylation under basic conditions gave the pyridinone alkaloid cerpegin 62 (Scheme 17) [70] which is an anti-inflammatory and selectively inhibits the post-acid activity of mammalian 20S proteasomes [71].

(f) 5-Stannylated *N*-Boc-protected 2,3-dihydro-1*H*-pyrrole **63** has been recently obtained by direct lithiation-stannylation of the corresponding *N*-Boc-pyrroline, and has been used in a Stille cross-coupling reaction with the vinyl

triflate **64** to give the trienecarbarbamate **65**. This compound has been heated to effect an electrocyclic ring closure and oxidized in situ with manganese(IV) oxide to give the marine sponge metabolite  $(\pm)$ -cis-trikentrin A (**66**) after Bocdeprotection and aromatization (Scheme **18**) [72]. Starting from a related stannylated pyrroline,  $(\pm)$ -cis-trikentrin B has been obtained [72].

# **4.4.** Unprecedented Approach of CIPE to 3,4-fused Pyridine-2-one

The synthesis of the 3,4-fused pyrimidine skeleton is limited to three general methods, which include (i) intermolecular annulation and intramolecular cyclization of 3- or 4-substituted pyrimidine derivatives (route i) [73] (ii) [3+3] annulation of a C–C–N fragment, such as an  $\alpha$ -acidic imine derivative, with a C–N–C fragment, such as an acyl

Scheme (17). Total synthesis of anti-inflammatory agent, cerpegin.

Scheme (18). Synthetic application in preparation of marine metabolite, (±)-cis-trikentrin A, 66.

Scheme (19). Merit of one-pot, three-component synthesis of 3,4-fused pyrimidin-2-one, 70.

heterocumulene derivative (route ii) [74]; and (iii) [5+1] annulations of an N-C-C-N fragment with a C-1 unit, such as a carbonyl compound or a heterocumulene (route ii) [75]. However, these methods are limited by the fact that they often require the isolation of intermediates, the synthesis of starting materials, high reaction temperatures, and a prolonged reaction time, which decreases product yield [76]. Hence, Sasada and coworkers recently identified a practical three-component coupling reaction via CIPE using a picoline derivative 67, a nitrile 68, and triphosgene 69 as a C-1 unit that produced 3,4-fused pyrimidin-2-one 70 in a direct, one-pot synthesis as shown in route iv of Scheme (19) [76].

### 4.5. De-protonative Metalation of Nicotine

In earlier study, Gros and coworkers reported a new base composed of *n*-BuLi and  $Me_2N(CH_2)_2OLi$  [77]. This unimetal superbase called *n*-BuLi-LiDMAE induced a regioselective lithiation of pyridine derivatives even when an ortho-directing group was present on the heterocyclic ring [78]. Further investigation by Gros and coworkers, revealed that bidentate tertiary diamines such as TMEDA led also to addition products as well as sterically hindered aminoalkoxides such as 2-diisopropylaminoethoxide. The aggregates were also found to be highly sensitive to solvents [77]. Later, a variety of novel, as well as known, C-2- and C-6-substituted nicotines have been synthesized directly from (S)-nicotine 71 in moderate to high yield with the help of conceptual information from CIPE in unimetal superbase [79]. The complete inhibition of the DoM effect of the C-2 chlorine of 72 with n-BuLi-LiDMAE was explained by the formation of aggregates between n-BuLi-LiDMAE and the substrate via lithium complexation by the pyridine nitrogen atom which upon quenching with electrophile afforded 73 (Scheme **20**) [79]. It has also been shown that *t*-BuLi in Et<sub>2</sub>O promotes an exclusive regioselective metalation of 2-aryl-6chloropyridine compounds at the aromatic ortho position [80].

**Scheme (20).** Evidence for regiospecificity at C-6 of 2-heterosubstituted pyridine.

## 4.6. Ortho-lithiation of N-Benzoyl Iminophosphoranes

Ortho-lithiation is most commonly achieved through deprotonation reactions with organolithium bases [81]. According to the CIPE model, the polar group linked to the aromatic ring directs the approach of the base to the deprotonation site by coordination to the lithium atom and contributes to the stabilization of the ortholithiated species through intramolecular coordination [82]. Although, regioselective ortho-deprotonation of iminophosphoranes 74 and 75 at either side of the PNCO moiety is feasible, the synthetic usefulness of these anions is rather limited due to the intramolecular quench observed for the ortho-PN anion of the parent compound 74 to afford the benzophenone 76 (Scheme 21a) [83] and the poor performance in the case of the ortho-CO anion arising from the methoxy derivative 75 to give the ortho-methylated product 77 (Scheme 21b) [84].

Moreover, owing to these drawbacks encountered in **74** and **75** above, Aguilar and coworkers pressed further and attempted halogen/lithium exchange reactions on **78** as a method for accessing ortholithiated iminophosphoranes **79**. This attempt was not only successful in producing **79** but

also served as opening to various new compounds **80a-h** because of tolerable electrophile quenching attributable to intermediate **79** as shown in Scheme **(22)** [84]. It is interesting to note that the trapping reactions with a representative series of electrophiles allowed the transformation of the C-Li bond of **79** into a wide variety of C-X (X = Hg, I, P, Sn, Si) and C-C bonds, providing access to new stabilized iminophosphoranes **80a-h** not easily accessible through other synthetic pathways [84].

## 4.7. Accessibility of New Polyphosphazenes

The degree of accessibility of the reactive centers to the incoming reagents largely dictated by the behavior of the polymers in solution is one of the great determining factors toward new functionalized polymeric material [85]. In view of this, Carriedo and Valenzuela reported that the halogenations of their earlier synthesized precursor phosphazene  $\{[NP(O_2C_{12}H_8)]\}_n$  81 [86] led to a new type of well-defined chemically regular chlorinated- 82a or iodinated polyphosphazenes 82b which upon silylation afforded 83a and 83b respectively (Scheme 23) [87]. There is an evidence of prelithiation disparity as 82a could not be substitutionally silylated on the chlorine position, but have to

Scheme (21). (a). Ortho-PN deprotonation of 74 and intramolecular quench (b). Ortho-CO deprotonation of 75 and subsequent methylation.

Scheme (22). Trapping reaction of lithio species 79 for convenient synthesis of new stabilized iminophosphoranes 80a-h.

Reagents and conditions: (i) Cl<sub>2</sub> /H<sub>2</sub>SO<sub>4</sub>; (ii) [lpy<sub>2</sub>]BF<sub>4</sub>/HSO<sub>3</sub>CF<sub>3</sub> in Cl<sub>2</sub>CH<sub>2</sub>; (iii) 1, LiBu<sup>t</sup> THF,-78 °C; 2, CISiMe<sub>3</sub>.

Scheme (23). Synthesis of functionalized silylated polyphosphazenes.

occur on the trimeric group as seen in 83a. However, because of large atomic size of iodine and the lithium charge effect, the silvlation occurred on the dimeric position to replace the iodine to afford 83b [87]. Thus, Carriedo and Valenzuela stated clearly that the chemical reactivity of polyphosphazenes with 2,2'-dioxybiphenyl phosphorus rings in the repeating units is limited by conformational changes induced by the new groups incorporated to the ring carbons and by the proximity of the reaction centers to the main chain [87].

### 4.8. Variation of Directing-Group Orientation

For deprotonative lithiation reactions, the geometrical constraints within a complex in the transition state for transfer of the proton to the lithiating reagent have been shown to be important for efficient reaction [23]. For reactions that provide  $\alpha$ -lithioamine derivatives of amides an orthogonal relationship between the lithio carbanion and the pi system of the amide has been established to be favorable. These results along with semi-empirical calculations suggested that a small dihedral angle and a calculated distance of 2.78 Å between the carbamate carbonyl oxygen and the proton to be removed were favorable for a carbamate-directed lithiation. Based on the careful study of the effect of directing-group orientation, new series of selected bicyclic carbamates were obtained. The direct lithiation of N-Boc pyrrolidine 84 and reaction with diisopropyl ketone or di-tert-butyl ketone afforded intermediate alkoxide which underwent cyclization upon warming up to room temperature to afford the bicyclic carbamates 85a and 85b respectively [88]. The pyrrolidinederived oxazolidinones 85a and 85b, upon treatment with sec-butyllithium (s-BuLi)/TMEDA at -78 °C followed by electrophiles provided the substituted products  $86a_1 - 86b_4$  in good yields as shown in Table 3 [88]. Recently, the reactions of Hoppe's lithiated carbamates with appropriately

Table 3. Stereoselective lithiation-substitutions of bicyclic carbamate.

substituted vinylboranes or boronic ester have been reported [89].

#### 4.9. Essentiality of DoM in Scale-up

In the past decade, the DoM reaction has enjoyed increasing application in large-scale process chemistry for the preparation of required amounts for advanced drug discovery studies and commercial drugs. By way of illustration, the synthesis of 2-bromo-6-chlorobenzoic acid 88 on a 60 kg scale and in excellent yields (89–90%) was achieved by Merck chemists [90] via painstaking optimization study of the experimental conditions in metalation of 3-chloro bromobenzene 87 followed by quenching with CO<sub>2</sub> as the suitable electrophile (Scheme 24a). Similarly, at Novartis, a pilot plant synthesis of the lead tricyclic compound 91 was devised in large scale [91]. This involved the metalation of the 1,7-dimethoxynaphthalene 89 and electrophilic quenching to give the intermediate 90 in 83% vield which upon further transformation afforded the targeted 6-trimethylsilyl-9-methoxy-1-methyl-3,4,4a,5,10,10 a-hexahydro-2*H*-benzo[g]quinoline, **91** [91] (Scheme **24b**) which was a dopamine D2/D3 agonist useful as antidepressant and anti-psychotic agent [92]. In one pot, chemists at BMS were able to use DoM to synthesize a tetrazole boronic acid 93 from the tetrazolo-starting material, 92 in the first step of the preparation of Losartan™ 94, an antihypertensive drug which is produced 1000 kg/Year [93] (Scheme **24c**).

### 4.10. Hydroxymethylation via Deprotonation

Since the availability of  $\alpha$ -silyl carbanion played a significant role in the first stage of the reactions, many synthetic strategies have been developed in order to access

this synthon [94]. Among these  $\alpha$ -silyl carbanion-generating methods, deprotonation by butyllithium is by far the most convenient way and easily accessible technique, since αhalosilanes. α-heteroatom substituted silanes. vinylsilanes are not always readily available. deprotonation of 2-pyridyltrimethylsilane using t-BuLi in diethyl ether was reported to proceed through CIPE strategy of the 2-pyridyl group on silicon 95 [95]. This is because, in most cases, together with the stabilization of the carbanion by the α-silyl group, additional stabilization effects by neighboring heteroatoms or electron withdrawing groups have been exploited for their generation [95]. In the second stage, the lithiated species of 2-pyridyldimethylsilyl (2-PyMe<sub>2</sub>-Si) **96** formed was quenched with appropriate electrophiles to afford substrates 97a-e which were considered to be excellent hydroxymethylating agents because they possessed versatile silyl group that could be converted to the hydroxyl group with much milder conditions compared to the well-known PhMe<sub>2</sub>Si group [95]. The oxidative cleavage of carbon-silicon bonds were performed using 30 % H<sub>2</sub>O<sub>2</sub> (30 equiv), KF (2.0 equiv), and KHCO<sub>3</sub> (2.0 equiv) in MeOH/THF (1/1) at 50 °C to afford the corresponding alcohols **98a-e** in excellent yields (Table 4). Hence, this two-step transformation provided an efficient method for the nucleophilic hydroxymethylation [95].

#### 4.11. Fluorenone: Carbocyclic Ketone from CIPE

Other reported cipe-based techniques have been reported to include lithiation of a silyl ether in the preparation of ortho-Fries hydroxyketone [96], directed remote aromatic metalation to access carbocyclic compounds [97], regioselective ring lithiation of BF<sub>3</sub>-complexed 3-picoline

<sup>&</sup>lt;sup>a</sup> A careful search for diastereomers in each case did not reveal their presence. It was estimated that at least 2% of any diastereomers would have been detected by GC analyses. <sup>b</sup> A compound tentatively identified as the *trans* diastereomer was isolated in 8% yield. <sup>c</sup> Additional product was present but was not separated from benzophenone. <sup>d</sup>Two diastereomers were formed in a 1.5:1 ratio.

**Scheme (24).** Large scale production and industrial application of DoM in metalation.

Table 4. New Alcohol by Hydroxymethylation of Substrate via H<sub>2</sub>O<sub>2</sub> Oxidation.

95  1.1 t-BuLi, Et <sub>2</sub> O  N SiMe <sub>3</sub> -78 °C, 30 min  SiMe <sub>2</sub> 96  P SiMe <sub>2</sub> N Si E KF, KHCO  MeOH/THF, 50 °C  98a-e							
Entry	Substrate	Electrophile	E	Product 98a-e	Time (h)	Yield <sup>a</sup> (%)	
1	97a	Ph(CH <sub>2</sub> ) <sub>3</sub> Br	Ph(CH <sub>2</sub> ) <sub>3</sub>	HO Ph	6	98	
2	97b	PhCHO	PhC-OH	HO Ph OH 98b	6	96	
3	97c	Ph(CH <sub>2</sub> ) <sub>2</sub> CHO	Ph(CH <sub>2</sub> ) <sub>2</sub> C-OH	HO Ph	6	90	
4	97d	PhC(O)Me	PhC(OH)Me	HO Me OH 98d	6	93	
5	97e	HO-C <sub>6</sub> H <sub>10</sub> -Br	HO-C <sub>6</sub> H <sub>10</sub>	HO OH	6	95	

[98], benzyne intermediate product formation via media effect on n-BuLi reactivity [99], enolate formation for one step synthesis of an optically active β-substituted ketone [100] and LDA-mediated ortho metalation of N,N-dialkyl-2biphenyl carboxamides for the synthesis of cyclic ketone [101]. A typical cyclic ketone produce via this route is called fluorenone which is reported to be achieved through the pathway presented in Scheme (25) below according to the investigation by Tilly and coworkers [101]. In the absence of an electrophile, 100 undergoes equilibration via 99° with 102, whose fate is instantaneous cyclization to a stable tetrahedral carbinolamine oxide 103 which, only hydrolysis, affords fluorenone 104 [101].

#### **CONCLUSION**

In summary, heteroatom-facilitated lithiation reactions have assumed an increasingly important role in the elaboration of carbocyclic aromatic and heteroaromatic systems. The reactivity profile of the lithio species in a variety of C-C-bond-forming reactions is quite broad and useful. The complex-induced proximity effect (CIPE) in deprotonation may serve as a heuristic to discover new

Scheme (25). Synthetic pathway to accessing fluorenone as a valuable cyclic ketone.

modes for C–H activation, which could be extended to carbanion chemistry. CIPE is an area that demands more careful examination in order to gain insight into the design of highly active and synthetically useful heterocyclic compounds *via* metalation chemistry. This review underlined the importance of initial lithiation site knowledge to understand the course of a metalation reaction as well as the crucial role of selective site complexation in directed lithiations. It therefore, provides a vista of opportunity towards constructing new biologically active heterocyclic compounds for present and future drug design and development.

#### CONFLICT OF INTEREST

The authors hereby declare that there is no conflict of interest as regard this present work.

# ACKNOWLEDGEMENT

Declared none.

### REFERENCES

- [1] Parra, S.; Vitse, O.; Bénézech, V.; Deleuze-Masquéfa, C.; Subra, G.; Bompart, J.; Escale, R.; Chapat, J.P.; Bonnet, P.A. Metalation and halogen-metal exchange in the imidazo [1,2-a]quinoxaline series. J. Heterocycl. Chem., 2001, 38, 41-44.
- [2] Clayden, J.P. In patai series: The chemistry of functional groups. the chemistry of organolithium compounds; Rappoport, Z.; Marek, I., Eds.; Wiley: Chichester, 2004; Part 1, pp. 495-646.
- [3] Beak, P.; Meyers, A. Stereo- and regiocontrol by complex induced proximity effects: reactions of organolithium compounds. Acc. Chem. Res., 1986, 6, 356-363.
- [4] Breit, B. Controlling stereoselectivity with the aid of a reagent-directing group: hydroformylation, cuprate addition, and domino reaction sequences. *Chem. Eur. J.*, 2000, 6, 1519-1524.
- [5] O'Brien, P.; Wilberg, K.B.; Bailey, W.F.; Hermet, J.-P.; McGrath, M.J. An experimental and computational study of the enantioselective lithiation of *N*-Boc pyrrolidine using sparteine-like chiral diamines. *J. Am. Chem. Soc.*, 2004, 126, 15480-15489.
- [6] Engel, K.M.; Mei, T.S.; Wasa, M.; Yu, J.Q. Weak coordination as a powerful means of developing broadly useful C-H functionalization reactions. Acc. Chem. Res., 2012, 45, 788-802.
- [7] Hassner, A.; Ghera, E.; Yechezkel, T.; Kleiman, V.; Balasubramanian, T.; Ostercamp, D. Stereoselective and enantioselective synthesis of five-membered rings *via* conjugate

- additions of allylsulfone carbanions. *Pure Appl. Chem.*, **2000**, *72*, 1671-1683.
- [8] Chinchilla, R.; Najera, C. Acylvinyl and vinylogous synthons. Chem. Rev., 2000, 100, 1891-1928.
- [9] Parra, M.; Sotoca, E.; Gil, S. A convenient generation of acetic acid dianion. Eur. J. Org. Chem., 2003, 1386-1388.
- [10] Nagendrappa G. Benzoin condensation The cyanide connection with Tapioca and vitamin B1. Resonance, 2008, 13, 355-368.
- [11] Maki, B.E.; Chan A.; Scheidt K.A. Protonation of homoenolate equivalents generated by *N*-heterocyclic carbene. *Synthesis*, **2008**, *8*, 1306-1315.
- [12] Smith A.B.; Adams C.M. Evolution of dithiane-based strategies for the construction of architecturally complex natural products. Acc. Chem. Res., 2004, 37, 365-377.
- [13] Linnane, P.; Magnus, N.; Magnus, P. Induction of molecular asymmetry by a remote chiral group. *Nature*, 1997, 385, 799-801.
- [14] Donskaya, O.V.; Dolgushin, G.V.; Lopyrev, V.A. Vicarious nucleophilic substitution of hydrogen in nitro-substituted pyrroles, azoles and benzannelated systems based on them. *Chem. Heterocycl. Comp.*, 2002, 38, 371-384.
- [15] Hermet, J.P.R.; Porter, D.W.; Dearden, M.J.; Harrison, J.R.; Koplin, T., O'Brien, P.; Parmene, J.; Tyurin, V.; Whitwood, A.C.; Gilday, J.; Smith, N.M. Synthesis of sparteine-like chiral diamines and evaluation in the enantioselective lithiation-substitution of N-(tert-butoxycarbonyl)-pyrrolidine. Org. Biomol. Chem., 2003, 1, 3977-3988.
- [16] Hartung, C.G.; Snieckus, V. The directed ortho metalation reaction - a point of departure for new synthetic aromatic chemistry. *Modern Arene Chem.*, 2002, 330-367.
- [17] Chincilla, R.; Nájera, C.; Yus, M. Metalated heterocycles in organic synthesis: recent applications. Arkivoc, 2007, 2007, 152-231.
- [18] Raposo, M.L.; Fernández-Nieto, F.; Garcia-Rio, L.; Rodríguez-Dafonte, P.; Paleo, M.R.;Sardina F.J. Mechanism of the deprotonation reaction of alkyl benzyl ethers with n-butyl lithium. Chem., Eur. J., 2013, 19, 9677-9685.
- [19] Sparling, B.A.; Moebius, D.C.; Shair, M.D. Enantioselective total synthesis of hyperforin. J. Am. Chem. Soc., 2013, 135, 644-647.
- [20] Beng, T.K.; Tyree, W.S.; Parker, T.; Su, C.; Williard, P.G.; Gawley, R.E. Dynamic of catalytic resolution of 2-lithio-N-Bocpiperidine by ligand exchange. J. Am. Chem. Soc., 2012, 134, 16845-16855.
- [21] Anderson, D.R.; Faibish, N.C.; Beak, P. Complex-induced proximity effects in directed lithiations: Analysis of intra- and intermolecular kinetic isotope effects in directed aryl and benzylic lithiations. J. Am. Chem. Soc., 1999, 121, 7553-7558.
- [22] Wang, W.; Snieckus, V. Remote directed metalation of biaryl o-carbamates. Ring to ring carbamoyl transfer route to biaryls, dibenzo[b,d]pyranones and the natural fluorenone dengibsin. *J. Org. Chem.*, **1992**, *57*, 424-426.

- Resek, J.E.; Beak, P. Complex-induced proximity effects evidence for a complex on the reaction pathway of α-lithiation of a benzylic urea. J. Am. Chem. Soc., 1994, 116, 405-406.
- Whisler, M.C.; MacNeil, S.; Snieckus, V.; Beak, P. Beyond [24] thermodynamic acidity: a perspective on the complex-induced proximity effect (CIPE) in deprotonation reactions. Angew. Chem. Int. Ed., 2004, 43, 2206-2225.
- Clark, R.D.; Jahangir, A. Lateral lithiation reactions promoted by [25] heteroatomic substituents. Organic Reactions, 2004, 1-314.
- Quanch, T. Heteroatom-facilitated lateral lithiation: generation and application in organic synthesis. Department of Chemistry, University of Toronto, Canada, 2001.
- [27] Gessner, V.H. The complex-induced proximity effects in organolithium chemistry In: Ideas in Chemistry and Molecular Sciences Pignataro B. (Ed.), John Wiley and Sons, 2010, pp. 96-
- [28] Gessner, V.H.; Strohmann, C. Lithiation of TMEDA and its higher homologous TEEDA: understanding observed α- and βdeprotonation. J. Am. Chem. Soc., 2008, 130, 14412-14413.
- [29] Gessner, V.H.; Däschlein, C.; Strohmann, C. Structure formation principles and reactivity of organolithium compounds. Chem. Eur. *J.*, **2009**, *15*, 3320-3334.
- Kamps, I.; Mix, A.; Berger, R.J.F.; Neumann, B.; Stammler, H.G.; [30] Mitzel, N.W. Two diamino-substituted lithiocarbanions in one molecule. Chem. Commun., 2009, 5558-5560.
- [31] Saá, J.M. An HF and DFT ab initio study on the mechanism of ortho-directed lithiation of hydric and non-hydric aromatic compounds incorporating aggregation and discrete solvation: the role of N,N,N',N'-tetramethylethane-1,2-diamine (TMEDA) in lithiation reactions. Helv. Chim. Acta, 2002, 85, 814-840.
- [32] Fernández, I.; González, J.; López-Ortiz, F. Deuterium-labeling and NMR study of the dearomatization of N-alkyl-Nbenzyldiphenylphosphinamides through anionic cyclization: ortho and benzylic lithiation directed by complex-induced proximity effects. J. Am. Chem. Soc., 2004, 126, 12551-12564.
- Jackman, L.M.; Petrei, M.M.; Smith, B.D. [33] Degenerate transesterification of 3,5-dimethyl phenolate/3,5-dimethylphenyl esters in weakly polar, aprotic solvents. Reaction of aggregates and complex-induced proximity effects. J. Am. Chem. Soc., 1991, 113,
- [34] Denmark, S.E.; Beutner G.L. Lewis base catalysis in organic synthesis. Angew. Chem. Int. Ed., 2008, 47, 1560-1638.
- [35] Yus, M.; Torregrosa, R.; Pastor, I.M. Masked ω-lithio ester enolates: synthetic applications. *Molecules*, **2004**, *9*, 330-348.
- Petragnani, N.; Stefani, H.A. Advances in organic tellurium [36] chemistry. Tetrahedron, 2005, 61, 1613-1679.
- [37] Boudier, A.; Bromm, L.O.; Lotz, M.; Knochel, P. New applications of polyfunctional organometallic compounds in organic synthesis. Angew. Chem. Int. Ed., 2000, 39, 4414-4435.
- Hu, X.E. Nucleophilic ring opening of aziridines. Tetrahedron, [38] 2004, 60, 2701-2743.
- D'hooghe, M.; Van Speybroeck, V.; Waroquier, M.; De Kimpe, N. Regio- and stereo- specific ring opening of 1,1-dialkyl-2-(aryloxymethyl)aziridinium salts by bromide. Chem. Commun., 2006, 1554-1556.
- [40] Sweeney, J.B. Aziridine synthesis via nucleophilic attack of carbene equivalents of imines: The aza-Darzens reaction. Eur. J. Org. Chem., 2009, 4911-4919.
- Jeong, J.U.; Tao, B.; Sagasser, I.; Henniges, H.; Sharpless, K.B. Bromine-catalyzed aziridination of olefins. A rare example of atom-transfer redox catalysis by a main group element. J. Am. Chem. Soc., 1998, 120, 6844-6845.
- [42] Dammacco, M.; Degennaro, L.; Florio, S.; Luisi, R.; Musio, B.; Altomare, A. Lithiation of N-alkyl-(o-tolyl)aziridine: Stereoselective synthesis of isochromans. J. Org. Chem., 2009, 74, 6319-6322.
- Azzena, U.; Dettori, G.; Pisano, L.; Musio, B.; Luisi, R. BH<sub>3</sub>-Promoted stereoselective  $\beta$ -lithiation of N-alkyl-2-phenylaziridines. J. Org. Chem., 2011, 76, 2291-2295.
- [44] Musio, B.; Clarkson, G.J.; Shipman, M.; Florio, S.; Luisi, R. Synthesis of optically active arylaziridines by regio- and stereospecific lithiation of N-Bus-phenylaziridine. Org. Lett., 2009,
- Ralph, J.; Peng, J.; Lua, F. Isochroman structures in lignin: a new [45] β-1 pathway. *Tetrahedron Lett.*, **1998**, *39*, 4963-4964.
- [46] Clayden J., In Organolithiums: Selectivity for Synthesis; Pergamon: Oxford, UK, 2002; Chapter 2.

- [47] Florio, S.; Luisi, R. Aziridinyl anions: generation, reactivity, and use in modern synthetic chemistry. Chem. Rev., 2010, 110, 5128-
- [48] Wu, Y.-C.; Zhu, J. Asymmetric total syntheses of (-)-renieramycin M and G and (-)-joru mycin using aziridine as a lynchpin. Org. Lett., 2009, 11, 5558-5561.
- [49] Capriati, V.; Florio, S.; Luisi, R.; Musio, B. Directed ortho lithiation of N-alkylphenyl aziridines. Org. Lett., 2005, 7, 3749-3752.
- Gómez-SanJuan, A.; Sotomayor, N.; Lete, E. Inter- and intramolecular enantioselective carbolithiation reactions. Beilstein J. Org. Chem., 2013, 9, 313-322.
- [51] Hodgson, D.M.; Hughes, S.P.; Thompson, A.L.; Heightman, T.D. Terminal aziridines by α-deprotonation/electrophile trapping of Nprotected aziridine. Org. Lett., 2008, 10, 3453-3456.
- [52] Martínez-Estíbalez, U.; Gómez-SanJuan, A.; García-Calvo, O.; Arrasate, S.; Sotomayor, N.; Lete, E. Intramolecular carbolithiation reactions of aryllithiums in the synthesis of carbocyclic and heterocyclic compounds. In: Targets in Heterocyclic Systems; Attanasi, O.; Spinelli, D., Eds.; Italian Society of Chemistry: Rome, **2010**; *14*, 124-149.
- Woltering, M.J.; Fröhlich, R.; Wibbeling, B.; Hoppe, D. Stereoselective synthesis of hydroxylated indolizidines via (-)-Sparteine-mediated kinetic resolution coupled with intramolecular carbolithiation. Synlett, 1998, 797-800.
- Barluenga, J.; Fañanás, F.J.; Sanz, R.; Marcos, C. Intramolecular [54] carbolithiation of allyl o-lithioaryl ethers: A new enantioselective synthesis of functionalized 2,3-dihydrobenzo furans. Chem., Eur. J., **2005**, 11, 5397-5407.
- [55] Barluenga, J.; Fañanás, F.J.; Sanz, R.; Marcos, C. Diastereo- and enantioselective carbo-lithiation of allyl o-lithioaryl ethers: new chiral cyclopropane derivatives. Org. Lett., 2002, 4, 2225-2228.
- Mealy, M.J.; Luderer, M.R.; Bailey, W.F.; Sommer, M.B. Effect of [56] ligand structure on the asymmetric cyclization of achiral olefinic organolithiums. J. Org. Chem., 2004, 69, 6042-6049.
- Soubh, L.; Besch, A.; Otto, H.H. Synthesis and properties of N-[57] substituted saccharin derivatives. Pharmazie, 2002, 57, 384-392.
- [58] Aliyenne, A.O.; Khiari, J.E.; Kraïem, J.; Kacem, Y.; Hassine, B.B. Efficient access to chiral N-substituted saccharin analogues via the directed ortho-lithiation of 3-N-aryl sulfonyl oxazolidin-2-ones. Tetrahedron Lett., 2006, 47, 6405-6408.
- [59] Tricotet, T.; Fleming, P.; Cotter, J.; Hogan, A.M.L.; Strohmann, C.; Gessner, V.H.; O'Shea, D.F. Selective vinyl C-H lithiation of cisstilbenes. J. Am. Chem. Soc., 2009, 131, 3142-3143.
- [60] Sarveswaran, S.; Gautam, S.; Ghosh, J. In: Wedelolactone, a medicinal plant-derived coumestan, induces caspase-dependent apoptosis in prostate cancer cells via down-regulation of PKCepsilon without inhibiting Akt. [abstract]. Proceedings of the 104th Annual Meeting of the American Association for Cancer Research; 2013 Apr 6-10; Washington, DC. Philadelphia (PA): AACR; Cancer Res., 2013, 73 (8 Suppl.) doi:10.1158/1538-7445.AM2013-
- Jones Jr, W.D.; Ciske, F.L. A convenient synthesis of dengibsin. J. [61] Org. Chem., 1996, 61, 3920-3922.
- Kowalczyk, B.A. Total synthesis of RS-42358 and analogs using [62] lateral lithiation. Synthesis, 2000, 8, 1113-1116.
- [63] James, C.A.; Snieckus, V. Combined directed metalation - cross coupling strategies. Total synthesis of the aglycones of gilvocarcin V, M and E. Tetrahedron Lett., 1997, 38, 8149-8152.
- [64] Liu, T.; Kharel, M.K.; Zhu, L.; Bright S.A.; Mattingly, C.; Adams, V.R.; Rohr, J. Inactivation of the ketoreductase gilU gene of the gilvocarcin biosynthetic gene cluster yields new analogues with partly improved biological activity. ChemBioChem, 2009, 10, 278-
- Chauder, B.; Green, L.; Snieckus, V. The directed ortho [65] metalation-transition metal-catalyzed reaction symbiosis in heteroaromatic synthesis. Pure Appl. Chem., 1999, 71, 1521-1529.
- [66] Kűrti, L.; Czakó, B. Strategic Applications of Named Reactions in Organic Synthesis. Elsevier Academic Press, Burlington, USA. 2005, pp 418-425.
- Li, X.; Yin, W.; Sarma, P.V.V.S.; Zhou, H.; Ma, J.; Cook, J.M. Synthesis of optically active ring-A substituted tryptophans as IDO inhibitors. Tetrahedron Lett., 2004, 45, 8569-8573.
- [68] Dishington, A.P.; Johnson, P.D.; Kettle, J.G. Synthesis of a 5alkoxypyrido[3,4-d] pyrimidin-4(3H)-one derivative via directed

- ortho-metallation of a pyridine analogue. Tetrahedron Lett., 2004, 45, 3733-3735.
- [69] Yang, X.; Luo, S.; Fang, F.; Liu, P.; Lu, Y.; He, M.; Zhai, H. Synthesis of conformationally restricted nicotine analogues by intramolecular [3+2] cycloaddition. *Tetrahedron*, 2006, 62, 2240-2246
- [70] Lazaar, J.; Hoarau, C.; Mongin, F.; Trécourt, F.; Godard, A.; Quéguiner, G.; Marsais, F. One-pot four-step synthesis of cerpegin. *Tetrahedron Lett.*, 2005, 46, 3811-3813.
- [71] Pham, H.; Hovhannisyan, A.; Bouvier, D.; Tian, L.; Reboud-Ravaux, M.; Melikyan, G.; Bouvier-Durand, M. A new series of N<sup>5</sup> derivatives of the 1,1,5-trimethyl furo[3,4-c] pyridine-3,4-dione (cerpegin) selectively inhibits the post-acid activity of mammalian 20S proteasomes *Bioorg. Med. Chem. Lett.* **2012**, *22*, 3822-3827.
- [72] Huntley, R.J.; Funk, R.L. Total syntheses (±)-cis-trikentrin A and (±)-cis-trikentrin B via electrocyclic ring closure of 2,3divinylpyrrolines. Org. Lett. 2006, 8, 3403-3406.
- [73] Qu, G.-R.; Ren, B.; Niu, H.-Y.; Mao, Z.-J.; Guo, H.-M. A novel one-step method for the synthesis of C-5-substituted O<sup>6</sup>,5'cyclopyrimidine nucleoside analogues in ammonia water. *J. Org. Chem.*, 2008, 73, 2450-2453.
- [74] Zhou, A.; Pittman Jr., C.U. Cyclizations of 2-alkylthiazolines and 2-alkyloxazolines with α,α-disubstituted diacid chlorides or N-(chlorocarbonyl) isocyanate. Tetrahedron Lett., 2005, 46, 2045-2048.
- [75] Fustero, S.; Piera, J.; Sanz-Cervera, J.F.; Román, R.; Brodsky, B.H.; Sánchez-Roselló, M.; Acea, J.L.; Ramirez de Arellano, C. New fluorinated 1,3-vinylogous amidines as versatile intermediates: synthesis of fluorinated pyrimidin-2(1*H*)-ones. *Tetrahedron*, 2006, 62, 1444-1451.
- [76] Sasada, T.; Moriuchi, M.; Sakai, N.; Konakahara, T. An unprecedented approach to the single-step synthesis of 3,4-fused pyrimidin-2-one and pyrimidin-2-thione derivatives by a [3+2+1] annulations. Eur. J. Org. Chem., 2009, 5738-5743.
- [77] Gros, P.; Choppin, S.; Mathieu, J.; Fort, Y. Lithiation of 2-heterosubstituted pyridines with BuLi-LiDMAE: Evidence for regiospecificity at C-6. J. Org. Chem., 2002, 67, 234-237.
- [78] Gros, P.; Fort, Y. nBuLi/lithium aminoalkoxide aggregates: New and promising lithiating agents for pyridine derivatives. Eur. J. Org. Chem., 2002, 3375-3383.
- [79] Février, F.C.; Smith, E.D.; Comins, D.L. Regioselective C-2 and C-6 substitution of (S)- nicotine and nicotine derivatives. Org. Lett., 2005, 7, 5457-5460.
- [80] Fort, Y.; Rodriguez, A.L. First regioselective ortho-lithiation induced by a 2-chloro pyridyl group complexation. *J. Org. Chem.*, 2003, 68, 4918-4922.
- [81] Clayden, J.; Kenworthy, M.N. Dearomatising disrotatory electrocyclic ring closure of lithiated *N*-benzoyloxazolidines. *Org. Lett.*, 2002, 4, 787-790.
- [82] Petrov, A.R.; Rufanov, K.A.; Harms, K.; Sundermeyer, J. Reinvestigation of ortho-metalated *N,N*-dialkylbenzylamine complexes of rare-earth metals. First structurally characterized arylates of neodymium and gadolinium Li[LnAr<sub>4</sub>]. *J. Organomet. Chem.*, **2009**, *694*, 1212-1218.
- [83] García-López, J.; Fernández, I.; Ruiz, M.S.; López-Ortiz, F. C<sub>u</sub>.C<sub>ortho</sub>-Dimetalated phosphazene complexes. *Chem. Commun.*, 2007, 4674-4676.
- [84] Aguilar, D.; Fernández, I.; Cuesta, L.; Yañez-Rodríguez, V.; Soler, T.; Navarro, R.; Urriolabeitia, E.P.; López-Ortiz, F. Synthesis, structure, and reactivity of *N*-benzoylimino phosphoranes ortho lithiated at the benzoyl group. *J. Org. Chem.*, 2010, 75, 6452-6462.
- [85] Andrianov, A.K.; Marin, A.; Chen, J.; Sargent, J.; Corbett, N. Novel route to sulfonated polyphosphazenes: single-step synthesis using "noncovalent protection" of sulfonic acid functionality. *Macromolecules*, 2004, 37, 4075-4080.

- [86] Carriedo, G.A.; García-Alonso, F.J.; Gómez E.P.; González, P.A.; Marco, C.; Gómez, M.P.; Ellis, G.J. Appl. Polym. Sci., 2000, 77, 568-576
- [87] Carriedo, G.A.; Valenzuela, M.L. Chlorination, iodination, and silylation of poly(2,2'-dioxy-1,1'-biphenylphosphazene). New halogenated polyphosphazenes with sterically hindered reactivity. *Macromolecules*, 2010, 43, 126-130.
- [88] Gross, K.M.; Beak P. Complex-induced proximity effects: the effect of varying directing-group orientation on carbamate-directed lithiation reactions. J. Am. Chem. Soc., 2001, 123, 315-321.
- [89] Althaus, M.; Mahmood, A.; Suárez, J.R.; Thomas, S.P.; Aggarwal, V.K. Application of the lithiation-borylation reaction to the preparation of enantioenriched allylic boron reagents and subsequent in situ conversion into 1,2,4-trisubstituted homoallylic alcohols with complete control over all elements of stereochemistry. J. Am. Chem. Soc., 2010, 132, 4025-4028.
- [90] Gohier, F.; Castanet, A.S.; Mortier, J. Ortholithiation of unprotected benzoic acids: application for 2-chloro-6-substituted benzoic acid syntheses. Synth. Commun., 2005, 35, 799-806.
- [91] Bänziger, M.; Küsters, E.; La Vecchia, L.; Marterer, W.; Nozulak, J. A new practical route for the manufacture of (4aR, 10aR)-9-methoxy-1-methyl-6-trimethyl silanyl-1,2,3,4,4a,5,10,10a-octahydrobenzo [g]quinoline. Org. Process Res. Dev., 2003, 7, 904-912.
- [92] Brown, D.A.; Kharkar, P.S.; Parrington, I.; Reith, M.E.A.; Dutta, A.K. Structurally constrained hybrid derivatives containing octahydrobenzo[g or f]quinoline moieties for dopamine D2 and D3 receptors: Binding characterization at D2/D3 receptors and elucidation of a pharmacophore model. J. Med. Chem., 2008, 51, 7806-7819.
- [93] Larsen, R.D.; King, A.O.; Chen, C.Y.; Corley, E.G.; Foster, B.S.; Roberts, F.E.; Yang, C.; Lieberman, D.R.; Reamer, R.A.; Tschaen, D.M.; Verhoeven, T.R.; Reider, P.J. Efficient synthesis of losartan, a nonpeptide angiotensin II receptor antagonist. *J. Org. Chem.*, 1994, 59, 6391-6394.
- [94] Bates, T.F.; Dandekar, S.A.; Longlet, J.J.; Wood, K.A.; Thomas, R.D. Regiospecific synthesis of alpha-lithiated alkoxysilanes. J. Organomet. Chem., 2000, 595, 87-92.
- [95] Itami, K.; Kamei, T.; Mitsudo, K.; Nokami, T.; Yoshida J. Pyridyl group assisted deprotonation of a methyl group on silicon: Complex induced proximity effect and novel hydroxymethylation. *J. Org. Chem.*, 2001, 66, 3970-3976.
- [96] Lo, H.-J.; Lin, C.-Y.; Tseng, M.-C.; Chein, R.-J. Lithiation of a silyl ether: formation of an ortho-Fries hydroxyketone. Angew. Chem. Int. Ed., 2014, 53, 9026-9029.
- [97] Tilly, D.; Magolan, J.; Mortier, J. Directed remote aromatic metalations: mechanisms and driving forces. *Chem., Eur. J.*, 2012, 18, 3804-3820
- [98] Dhau, J.S.; Singh, A.; Kasetti, Y.; Bharatam, P.V. Complexinduced proximity effect in the regioselective lithiation of pyridine derivatives. *Eur. J. Org. Chem.* 2012, 1746-1752.
- [99] Slocum, D.W.; Carroll, A.; Dietzel, P.; Eilerman, S.; Culver, J.P.; McClure, B.; Browna S.; Holman, R.W. Metalations in hydrocarbon solvents; media effects on n-BuLi reactivity. Tetrahedron Lett., 2006, 47, 865-868.
- [100] Tanaka, F.; Node, M.; Tanaka, K.; Mizuchi, M.; Hosoi, S.; Nakayama M.; Taga, T.; Fuji K. 1,13-Binaphthalene-2,23-diol as a chiral auxiliary diastereoselective alkylation of binaphthyl esters, complex-induced proximity effects in enolate formation, and onestep synthesis of an optically active β-substituted ketone. J. Am. Chem. Soc., 1995, 117, 12159-12171.
- [101] Tilly D.; Fu, J-M.; Zhao, B-P.; Alessi, M.; Castanet, A-S.; Snieckus, V.; Mortier, J. On the mechanism of the directed *ortho* and remote metalation reactions of *N*,*N*-dialkyl biphenyl 2carboxamides. *Org. Lett.*, 2010, 12, 68-71.

Received: August 13, 2014 Revised: December 20, 2014 Accepted: January 16, 2015