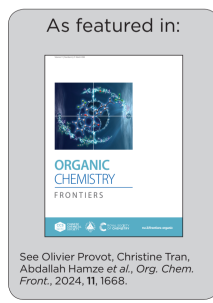


Showcasing research from Professor Abdallah Hamze's laboratory, Université Paris-Saclay, CNRS, BioCIS, 91400 Orsay, France.

Mo-catalyzed cyclization of *N*-vinylindoles and skatoles: synthesis of dihydroindolo[1,2-c]-quinazolines and dihydroindolo[3,2-b]-indoles, and evaluation of their anticancer activities

A novel synthesis method using *N*-vinylazoles produces unexpected compounds DINQ (R= Me) and DINI (R= H). Mocatalyst is crucial for DINQ formation from skatole and for accelerating the DINI cyclization. Select compounds exhibit potent cytotoxic effects against HCT-116 cells.









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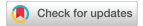




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Mo-catalyzed cyclization of *N*-vinylindoles and skatoles: synthesis of dihydroindolo[1,2-c]-quinazolines and dihydroindolo[3,2-b]-indoles, and evaluation of their anticancer activities†

Shannon Pecnard,^{†a} Xinya Liu,^{‡a} Olivier Provot,*^a Pascal Retailleau,^b Christine Tran*^a and Abdallah Hamze • **p***

A novel approach has been developed for synthesizing unexpected dihydroindolo[1,2-c]-quinazolines (DINQ) and dihydroindolo[3,2-b]indole (DINI) compounds using N-vinylazoles as starting materials. The presence of a Mo-catalyst was found to be essential for the formation of DINQs from skatole derivatives and for accelerating the cyclization of DINI precursors. Biological evaluation revealed that compound **30**, a 2-phenyl-1-(1-phenylvinyl)-1H-indole derivative and DINI analog **6g** exhibited nanomolar cytotoxic effects in vitro against a human colon cancer cell line (HCT-116).

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Introduction

Heterocyclic compounds play a pivotal role in medicinal chemistry due to their wide range of applications in drug design and discovery. 1 More than 85% of all biologically active chemical entities contain a heterocycle unit,2 with prevalent nitrogen-containing heterocycles usually found in the form of fiveor six-membered rings. Therefore, developing new synthetic methodologies that enable rapid access to diverse functionalized heterocyclic compounds holds great significance to medicinal chemists. Such advancements not only broaden the chemical space available for drug development but also streamline drug discovery programs. Due to their diverse biological activities, nitrogen-containing heterocycles have consistently captivated the interest of synthetic organic chemists.³⁻⁶ In cancer therapy, microtubules crucial for cell division serve as pivotal targets. Clinically used antimicrotubule agents include vinca alkaloids and taxanes (e.g., paclitaxel and docetaxel). Paclitaxel (Taxol), an antimicrotubule agent, is employed in the treatment of malignant solid tumors such as

Vinca alkaloids are approved for clinical use; vincristine, vinblastine, vinorelbine, vindesine, and vinflunine serve as agents in the treatment of hematological and lymphatic neoplasms. The principal toxicity symptom of vincristine is neurotoxicity, while neutropenia is the main toxicity sign of the other vinca alkaloids.

Compounds derived from combretastatin, acting as antitubulin agents, exhibit the potential to inhibit tubulin polymerization into microtubules, offering a promising avenue in cancer treatment.¹¹ Over the past decade, our research group has dedicated efforts to designing and synthesizing anti-tubulin agents.^{12,13} Despite the notable anticancer activity of natural combretastatin A-4 (CA-4), its efficacy is hampered by the presence of a Z double bond, resulting in the formation of an inactive E isomer (Scheme 1A).

Furthermore, the isomerization of CA-4's active *Z*-double bond to the less active *E*-CA-4 is readily observed during administration and metabolism, presenting a challenge to its clinical application.¹⁴ To address this limitation, our group identified isoCA-4¹⁵ and isoFCA-4,¹⁶ 1,1'-diarylethylenes compounds with antiproliferative activity equivalent to CA-4 but with enhanced stability. Additionally, we demonstrated that derivatives such as isoCbzCA-4,¹⁷⁻¹⁹ isoChromCA-4,²⁰ and isoDBzxCA-4,²¹ featuring carbazole, arylchromene, and dihydrobenzoxepine scaffolds, respectively, exhibit outstanding antiproliferative activities against colon cancer cell lines HCT-116 in the low nanomolar range (Scheme 1A). Recently, we synthesized *N*-vinylazoles by coupling *N*-tosylhydrazones

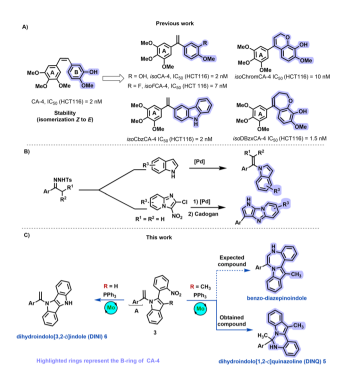
breast, lung, head and neck, and ovarian cancer. Common side effects encompass cardiotoxicity, myelosuppression, and peripheral neuropathy. 8

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[†]Electronic supplementary information (ESI) available: Detailed experimental procedures, characterization data, and copies of ¹H, ¹³C, and ¹⁹F NMR spectra. CCDC 2310238–2310243. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d3q001973j

[‡]These authors contributed equally.



Scheme 1 (A) Structure of CA-4, isoCA-4 analogs, (B) Our previous work, and (C) Targeted structures.

derivatives and N-H azoles,22 enabling the construction of a compound library with a pyrrolo-imidazo[1,2-a]pyridine backbone (Scheme 1B).²³

Building on our comprehension of the structure-activity relationships of isoCA-4 and our advancements in the chemistry of N-vinylazoles, we aimed to create compound collections based on privileged scaffolds (Scheme 1C). Consequently, our investigation focused on synthesizing novel isoCA-4 analogs featuring seven-membered ring heterocycles, specifically benzo-diazepinoindole, using N-vinylazoles as starting materials.

In this study, we will investigate the synthesis of various terminal N-vinylazoles 3, and their cyclization under Mo-Cadogan-reductive cyclization conditions. This cyclization could afford benzo-diazepinoindole (Scheme 1).

Results and discussion

Our investigation commenced with the preparation of N-vinylazole substrates (3) (Table 1). In accordance with our prior work,22 we performed the coupling reaction between indole derivative (1a) and N-tosylhydrazone (2a) using Pd2dba3·CHCl3 as the catalyst and iodobenzene as the oxidant agent. Unfortunately, under these conditions, the desired product (3a) was obtained but in a low 25% yield (entry 1, Table 1). The presence of a hindered *ortho*-nitrophenyl substituent on the C-2 position of (1a) resulted in decreased reactivity of the indole. Changing the oxidant from iodobenzene to oxygen and gradually introducing compound (2a) over 1 h using a syringe, led to a slight increase in yield (entry 2). Increasing the catalytic load to 20 mol% resulted in a yield of

Table 1 Optimization of *N*-vinylazole-2-(2-nitrophenyl) formation^a

	[Cat.]	2	${f L}$	Base	Solvent	Time (h) temp °C	$Yield^{b}$ (%)
1	Pd₂dba₃·CHCl₃	2a	_	NaO <i>t</i> Bu	CPME	5/115	25 ^c
2	Pd(OAc) ₂	2a	L1	K_3PO_4	DMF	5/100	36^d
3	Pd(OAc) ₂	2a	L1	K_3PO_4	DMF	5/100	$45^{d,e}$
4	PdG ₃ (XPhos)	2a	L1	K_3PO_4	DMF	5/100	55^d
5	Cu powder	2b	L2	K_2CO_3	Toluene	20/135	80
6	Cu powder	2c	L2	K_2CO_3	Toluene	20/135	85
7	Cu powder	2c	L2	K_2CO_3	Toluene	20/110	25
8	Cu powder	2c	L3	K_2CO_3	Toluene	20/135	55
9	Cu powder	2c	L2	K_2CO_3	Dioxane	20/135	45

^a Reaction conditions: indole 1a (0.2 mmol), reagent 2 (0.4 mmol), catalyst (10 mol%), ligand (20 mol%), base (0.3 mmol) and 0.8 mL of solvent in a sealed tube under argon for 20 h. ^b Isolated yield. ^c Reaction was performed in the presence of iodobenzene (1.2 equiv.) as oxidant. ^d Reaction was performed under O₂ balloon, and the N-tosylhydrazone was added with a push syringe in 1 h. ^e Catalyst 20 mol%, ligand 40 mol%.

45% (entry 3). The use of the precatalyst PdG3(XPhos) did not provide conclusive results (entry 4), and (3a) was obtained in a moderate 55% yield.

Copper demonstrates remarkable versatility, efficiently catalyzing reactions involving both radical and polar mechanisms. In the context of an extended reaction time (20 hours), copper's unique reactivity profile, cost-effectiveness, and sustainability. Accordingly, our focus shifted to the evaluation of copper catalysts (entries 5-9).²⁴ Employing Cu(0) and N,N-dimethylethylenediamine (L2) as a ligand (entry 5) in toluene for the coupling reaction between (1a) and 1-bromovinyl benzene (2b) resulted in an 80% yield of (3a). Encouraged by this favorable outcome, we explored additional reaction conditions employing copper as the catalyst. Substituting 1-bromovinyl benzene with 1-iodovinyl benzene led to a slight increase in yield to 85% (entry 6). However, reducing the temperature to 110 °C resulted in a significant drop in yield to 25% (entry 7). Switching the ligand to ethylenediamine (L3) yielded a lower yield of (3a) compared to L2 ligand (entry 8 vs. entry 6). Furthermore, replacing toluene with dioxane as the solvent led to a substantial decrease in the yield of (3a) (entry 9 vs. entry

Using the optimized conditions outlined in entries 5 and 6 of Table 1, we proceeded with the synthesis of N-vinylazole-2-(2-nitrophenyl) substrates (3) (Scheme 2). Depending on the availability of starting materials, specifically vinylhalides, and in some cases, due to the instability of (1-iodovinyl)phenyl derivatives, we mainly employed (1-bromovinyl)phenyl derivatives for the coupling reaction. The cross-coupling reaction exhibited good compatibility with electron-rich vinyl halides when paired with skatole or indole derivatives.

These conditions proved efficient even with sterically hindered vinyl halides, resulting in the successful synthesis of compounds (3c, 3e, 3j, and 3l) with good yields. While the coupling reaction was successful in synthesizing unsubstituted indoles on the C3 position (3i-o), slightly lower yields were obtained for (3i, 3j, and 3l) due to partial vinylation at the C3 position of $(1, R^1 = H)$.

Notably, the structure of (3i) was confirmed by 2D NMR analysis, including COSY, HSQC, HMBC, and NOESY experiments. Except for compounds (3g) and (3h) containing a fluorine atom, the standard conditions were less effective in synthesizing terminal N-vinylazoles bearing an electron-withdrawing group (e.g., $R^2 = CN$, 3p or $R^2 = 2'$, 3', 4'-Cl, 3r). Also, no reaction was observed with free phenol derivative (3s). However, to our delight, we successfully attempted the reaction with a C3-arylated indole, affording compound (3q) with a good 83% yield.

Applying our optimal conditions to alkyl substrates such as 2-bromopropene and 2-bromo-3,3,3-trifluoro-1-propene, does not provide the corresponding coupling products 3t and 3u.

After obtaining N-vinylazoles (3), our investigation shifted towards exploring the cyclization process. Our initial focus was on converting N-vinylazole derivatives derived from skatole, such as 3a. However, when the reaction was conducted using the standard Cadogan reaction conditions (4 equiv. of PPh₃ in

Scheme 2 Substrate scope for the synthesis of N-vinylazole-2-(2nitrophenyl) substrates 3a. Reaction conditions: all reactions were conducted with 0.2 mmol of indole 1, vinylhalide 2 (0.4 mmol), Cu(0) (10 mol%), N,N'-dimethyl-ethylenediamine (L2) (20 mol%), K2CO3 (1.5 equiv.), 0.8 mL of dry toluene in a sealed tube under argon for 20 hours at 135 °C. ^a The reactions were conducted with 1.9 mmol of indole 1, vinylhalide 2 (2.4 mmol), Cu(0) (20 mol%), L2 (40 mol%), K_2CO_3 (1.5 equiv.), 7.2 mL of dry toluene in a sealed tube under argon for 20 hours at 135 °C.

X = I; 3r 0%

dioxane),²⁵ no desired product 4a was observed, and the starting material (3a) was fully recovered (Table 2, entry 1). It appears that the presence of steric hindrance in the structure

Table 2 Preliminary results of intramolecular cyclization with 3a. and an optimization study for the formation of dihydroindolo[1,2-c]quinazoline **5a**^a

Entry	[Mo] Cat.	PR ₃ (equiv.)	Solvent	Time (h)	Yield ^b (%)
1		PPh ₃ (4)	Dioxane	16	0
2	10 mol%	$PPh_3(4)$	Dioxane	16	77
3	5 mol%	$PPh_3(4)$	Dioxane	16	78
4	5 mol%	$PPh_3(4)$	DMF	16	75
5	5 mol%	$PPh_3(4)$	Toluene	16	94
6	5 mol%	$PPh_3(4)$	Toluene	16	72^{c}
7	5 mol%	$PPh_3(4)$	Toluene	6	95
8	5 mol%	PPh ₃ (2.4)	Toluene	6	95
9	5 mol%	$PCy_3(2.4)$	Toluene	6	n.r. ^d
10	5 mol%	$P(o-tol)_3$ (2.4)	Toluene	6	n.r. ^d
11	5 mol%	TFP (2.4)	Toluene	6	35

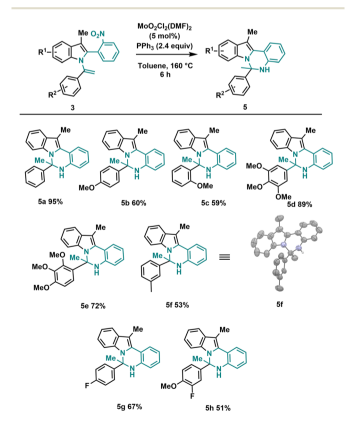
^aGeneral conditions: reactions were conducted with 0.1 mmol of N-vinylazole (3a), PPh3, MoO2Cl2(DMF)2, solvent (1 mL) in a sealed tube, 160 °C. b Isolated yield. Reaction was conducted at 135 °C. No reaction.

of skatole hampers the reaction compared to the standard deoxygenation of ortho-nitrostyrenes. Arnáiz et al., developed a Mo(vI)/PPh3 catalyst system that is compatible with labile substrates.26

In our recent work, we successfully applied molybdenumcatalyzed cyclization conditions to synthesis pyrrolo-imidazo [1,2-a]pyridines.²³ Adapting these conditions to N-vinylazole (3a) involved utilizing MoO₂Cl₂(DMF)₂ (10 mol%) in the presence of PPh₃ (4 equiv.) in dioxane at 160 °C in a sealed tube, resulting in complete conversion of the starting material (entry 2). However, a careful analysis of the crude product's ¹H NMR spectrum did not reveal the characteristic proton signal of the anticipated 7-membered ring (proton H3). Instead, a singlet was observed for 3 protons at 2.17 ppm in deuterated acetone. Additionally, mass spectrometry (ESI+) analysis displayed a major peak of (M + H)⁺ 325.1705, deviating from the expected mass of 323.1548 for (4a). To unequivocally confirm the structure of the newly obtained compound, we successfully obtained a crystalline solid suitable for X-ray analysis.²⁷ This analysis confirmed the formation of the six-membered ring dihydroindolo-quinazoline (5a), isolated with a yield of 77% (Table 2). Encouraged by this outcome, we continued optimizing the reaction conditions to enhance the formation of this novel heterocycle (5a). Initially, reducing the catalytic amount of the Mo catalyst to 5 mol% did not impact the yield, and (5a) was obtained with a good yield of 78% (entry 3). Subsequently, exploring of different solvents revealed that employing toluene increased the yield (entries 4-6). Altering the temperature to 135 °C resulted in a slight decrease in yield (entry 6). Ultimately, fine tuning of the reaction time and the amount of PPh₃ allowed us to identify the optimal conditions for the formation of (5a), which involved the use of 2.4 equivalents of PPh₃ for 6 h (entry 8). Using other reductant phosphine sources gave no reaction with tricyclohexylphosphine or tri-(otolyl)phosphine (entries 9 and 10), and only 35% yield was obtained with tri-(2-furyl)phosphine (entry 11).

Using the optimized reaction conditions outlined in Table 2, we investigated the substrate diversity of this protocol. As depicted in Scheme 3, various substrates with different electronic nature and steric properties underwent the reaction under the standardized conditions, affording the dihydroindolo-quinazoline derivatives (5a-h) in good yields. Notably, the reaction displayed tolerance towards steric hindrance substrates, as evidenced by the successful formation of compounds (5c) and (5e). Similarly, electronic factors did not impede the reaction, as demonstrated by the effective synthesis of compounds (5b) and (5g). We also successfully obtained a crystalline solid suitable for X-ray analysis for 5f.

Having explored the scope of the Mo/PPh3 cyclization process with skatole derivatives, next, we proceeded to investigate the standard cyclization conditions using unsubstituted C3-indole substrates (3i-o). This cyclization process can occur either on the terminal double bond of N-vinylazole (3) as observed with skatole derivatives, or on the C3 position of the indole moiety. By conducting the reaction under the standard conditions (Table 1), we achieved complete selectivity for cyclization at the C3 position of the indole, resulting in the formation of dihydroindolo[3,2-b]indole (DINI) derivatives



Scheme 3 General conditions: 3 (0.2 mmol), MoO₂Cl₂(DMF)₂ (5 mol%), PPh₃ (2.4 equiv.), toluene (3 mL), sealed tube 160 °C, 6 h.

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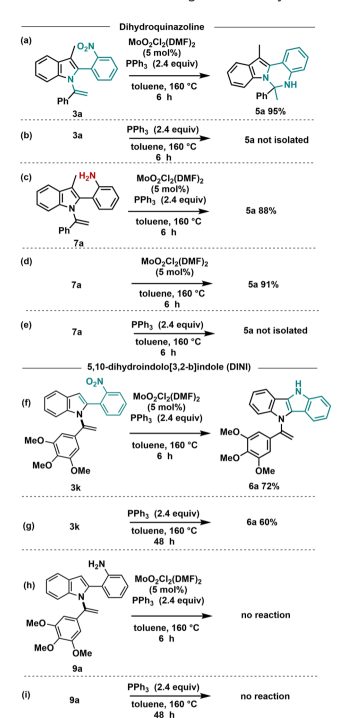
Scheme 4 General conditions: 3 (0.2 mmol), MoO₂Cl₂(DMF)₂ (5 mol%), PPh₃ (2.4 equiv.), toluene (3 mL), sealed tube 160 °C, 6 h.

(Scheme 4). The absence of the methyl group on the skatole allows C-H insertion on the carbon of the C3 position of the indole, preferentially over the enamine terminal carbon (see infra for the mechanism). Owing to the electron-rich and ease of oxidation of the DINI scaffold, materials incorporating DINI structures²⁸ have found extensive applications in organic electronics, including organic photovoltaics²⁹ and organic lightemitting diodes (LEDs).30 Also, compounds 6a-g were successfully synthesized using our standard conditions in good yields. It is worth noting that the aryl groups of these compounds do not contain electron-withdrawing groups (EWGs). However, we have not yet been able to obtain the corresponding N-vinylazole derivatives for this type of compounds. Interestingly, when utilizing 3-(2-nitrophenyl)-indole (3q), we obtained 5,10-dihydroindolo[3,2-b]indole 8a with a yield of 79%. The structure of this novel heterocycle was confirmed through X-ray analysis.

To gain insights into the mechanism of these reactions, we conducted several controlled experiments (Scheme 5). Initially, we investigated the formation of dihydroindolo[1,2-c]quinazoline compounds 5. In the absence of the Mo-catalyst, no conversion of N-vinylazole 3a to 5a was observed even after 6 h (Scheme 5b), indicating the crucial role of the Mo-catalyst in this transformation.

To determine whether the cyclization reaction proceeds through an amine or a nitrene intermediate, we subsequently reduced the nitro group of 3a to the corresponding amine 7a using an iron catalyst (Fe/HCl cat.).

We then carried out the cyclization reaction using 7a as the starting material (Scheme 5c), and under the standard conditions, we successfully obtained 5a with an 88% yield. This experiment provided evidence that the reaction pro-



Scheme 5 Control experiments. (a) Reaction with 3a under standard conditions. (b) Reaction with 3a without Mo catalyst. (c) Reaction starting from amine derivative 7a under standard conditions. (d) Reactions starting from amine derivative 7a with Mo catalyst and without PPh3. (e) Reactions starting from amine derivative 7a with PPh3 without Mo catalyst. (f) Reaction with 3k under standard conditions. (g) Reaction with 3k with PPh3 without Mo catalyst. (h) Reaction with amine 9a under standard conditions. (i) Reaction with amine 9a with PPh3 without Mo catalyst.

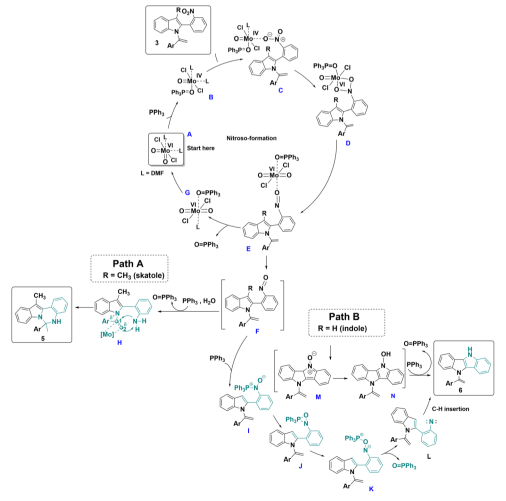
ceeds through the formation of an ortho-aniline intermediate. In the Cadogan reaction, PPh3 facilitates the deoxygenation of o-nitrostyrenes derivatives to nitrenes intermediates.

As this transformation proceeded with the amine 7a, we decided to carry out the reaction without the addition of PPh₃. As depicted in Scheme 5d, complete conversion of 7a to 5a was achieved in a 91% yield, suggesting that PPh3 does not play a significant role in this cyclization process starting from the amine. Finally, in the absence of Mo-catalyst, no conversion of the amine derivative to 5a was observed. These experiments led to the conclusion that the formation of dihydroindolo[1,2-c]quinazoline 5a from 3a necessitates the presence of both Mo and PPh₃. Additionally, it was observed that the formation of the amine intermediate occurred in the presence of a wet solvent, as previously reported in our work.31

Concerning the formation of DINI derivatives, the nitro compound 3k underwent conversion to the DINI derivative in a 72% yield within 6 h under the standard conditions (Scheme 5f). In the absence of Mo-catalyst, the reaction proceeded, albeit at a slower rate, requiring 48 hours for completion (Scheme 5g). To further validate the nature of the intermediate involved in this cyclization process, we conducted the reaction using the amine derivative 9a. No reaction was observed when starting from 9a in the presence of Mo-catalyst and PPh₃, nor in the absence of Mo-catalyst (Scheme 5h and i). These experimental results strongly support a Cadogan-type reaction mechanism.

Based on the control experiments and previous literature, 32 a mechanism is proposed in Scheme 6. Initially, in the presence of PPh₃, the Mo(v_I) catalyst A undergoes a reduction step, leading to Mo(IV) complex B. Subsequently, N-vinylazole 3 coordinates with complex B, forming complex C, which undergoes oxidative addition to provide the Mo(vi) metallacycle D. Then, D transforms into complex E, through oxygen atom transfer to the metal center leading to the loss of an oxygen atom from the initial nitro group. Complex E proceeds to generate the nitroso intermediate F and Mo-complex G. The departure of the O=PPh₃ ligand from complex G regenerates the initial Mocatalyst A, ready to initiate a new cycle.

In the case of skatole derivatives $(\mathbf{R} = \mathbf{Me})$, where the formation of the seven-membered ring is unfavorable, the nitroso intermediate F is reduced to the corresponding amine H in the presence of PPh3 and a wet solvent.33 The coordination of molybdenum to the ethylene moiety activates the double bond of the N-vinylazole and results in the formation of compound 5.



Scheme 6 Proposed mechanism for the formation of 5 and 6

When $\mathbf{R} = \mathbf{H}$ (indoles derivatives), a second equivalent of PPh₃ reacted with complex F, resulting in the formation of intermediate I. Intermediate I can then undergo further transformation to yield oxazaphosphiridine intermediate I. The nitrogenphosphore bond of J is subsequently cleaved, generating intermediate K. This intermediate undergoes deoxygenation, leading to the formation of nitrene L and triphenylphosphine oxide. Finally, nitrene L undergoes C-H insertion, 34 resulting in cyclization and the formation of the dihydroindolo[3,2-b]indole derivative 6. Alternatively, nitroso intermediates F could undergo direct 6π electrocyclization, ³⁵ resulting in the formation of nitronene M and then the N-oxide derivative N. Finally, N would be further deoxygenated to 6 in the presence of PPh₃.

Biological results

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To assess the potential of the newly synthesized compounds to inhibit cell growth, an in vitro cytotoxicity assay was conducted using the human colon cancer cell line HCT116. The CellTiter-Glo® cell viability assay was employed, utilizing luminescence to measure the number of viable cells based on the quantification of ATP, an indicator of metabolically active cells (the transformation of luciferin to oxyluciferin in the presence of ATP). This assay enables the determination of the number of living cells in culture based on the amount of ATP, which is directly proportional to cell viability.

In this study, all the synthesized compounds were tested at a concentration of 10⁻⁶ M in triplicate on the HCT-116 cell

line. The results are expressed in Fig. 1 as cytotoxicity %. A higher percentage of cytotoxicity indicates a higher level of compound activity against the cancer cells. Among the tested compounds, two heterocycles (30 and 6g) showed significant cytotoxicity against the HCT116 cell line. (30) compound exhibited a cytotoxicity percentage of 66%, while (6g), also possesses a methoxy group on the indole ring and demonstrated a notably higher cytotoxicity percentage of 93%.

These results highlight the importance of the 3,4,5-trimethoxy phenyl moiety and the presence of the methoxy group on either the indole or the dihydroindolo[3,2-b]indole scaffold for achieving significant cytotoxic activity against the HCT-116 cell line. It is worth noting that compounds belonging to the DINQ family did not exhibit any significant cytotoxic activity against the HCT-116 cell line in this study. Next, we determined the required concentrations of (30) and (6g) for the reduction of the respective cell viability by 50% (IC_{50}) using a luminescent assay. After 24 h of culture, the cells were treated with the tested compounds at 10 different final concentrations. After a 72 h treatment of the HCT-116 cell line with the respective compound, CellTiter Glo Reagent containing luceferin was added and luminescence was recorded with a spectrophotometric plate reader.

The results revealed that compounds (30) and (6g) exhibited potent antiproliferative activity, with IC₅₀ values of 298 nM and 62 nM, respectively (Fig. 2).

As isoCA-4 derivatives act as antitubulin, 13,16,18,21,36,37 to elucidate the potential binding mode of compound (6g) in

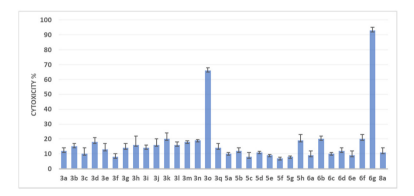
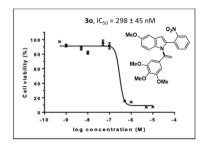


Fig. 1 Screening of generated analogs 3, 5, 6 and 8 in a human colon cancer cell line (HCT116). The cell proliferation inhibition was plotted for each of the analogs at 10^{-6} M concentration in HCT-116 cells. Three sets of experiments were carried out. Error bars represent standard deviation.



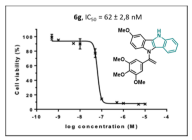


Fig. 2 Effect of compounds 30 and 6g on the HCT-116 cells viability

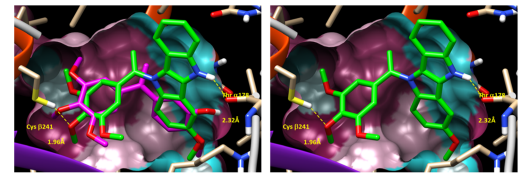


Fig. 3 (Left) Putative binding mode of DINI derivative 6 g (green color) within colchicine binding site of tubulin X-ray structure (accession code 6H9B) (left) with previously reported binding mode of reference compound isoCA-4 in overlay (magenta color). (Right) Same figure (without isoCA-4 reference binding mode).

tubulin, a docking study was conducted using the X-ray structure of tubulin (PDB code: 6H9B). 18 Compound (6g), which exhibited the highest cytotoxicity in this series, was docked and compared to isoCA-4. The docking results and superimposition of (6g) and isoCA-4 are shown in Fig. 3.

In the docking study, compound (6g) demonstrated a binding pose similar to that observed with isoCA-4. Similar to CA-4 and isoCA-4, the trimethoxyphenyl ring of (6g) was positioned in close proximity to Cysβ241, while the 3-methoxy-5,10-dihydroindolo[3,2-*b*]indole ring was situated Thrα178, forming a hydrogen bond between its NH group and the oxygen atom of Thra178. This binding arrangement suggests that compound (6g) interacts with tubulin in a similar manner to isoCA-4, which is known to act as an antitubulin agent.

Conclusions

In conclusion, this study presents a copper-catalyzed approach for the synthesis of various terminal N-vinylazoles. Through optimization of the Cadogan-reductive cyclization conditions, unexpected but welcome dihydroindolo[1,2-c]quinazoline compounds 5 were obtained from skatole derivatives. The mechanism of the formation of these compounds involves the formation of an aniline intermediate, which undergoes intramolecular cyclization in the presence of Mo-catalyst. This cyclization occurs through a nucleophilic aniline attack on the terminal double bond of the N-vinylazole substrate, resulting in the formation of compound 5. However, when N-vinylazoles bearing a hydrogen atom at the C3 position of the indole were used, the double bond of the indole reacted preferentially over the terminal double bond of the N-vinylindole. This preference led to the trapping of the nitrene intermediate and the formation of dihydroindolo[3,2-b]indole derivatives 6. Biological evaluation of the synthesized derivatives revealed for compounds 30 and 6g their potential as cytotoxic agents against the HCT-116 human colon cancer cell line. These compounds exhibited significant cytotoxicity, with IC50 values of 298 nM and 62 nM, respectively. Furthermore, docking studies

suggested that DINI derivative 6g interacts with tubulin in a similar manner to the antitubulin agent isoCA-4, supporting its potential as an antitubulin agent. These findings highlight the promising nature of the synthesized analogs as potential candidates for further development as anticancer agents.

Experimental

General

Full experimental procedures for the synthesis and spectroscopic characterization of all compounds and intermediates can be found in the ESI.†

General procedure for preparation of N-vinylazole compounds 3a-o

In a sealed tube under argon atmosphere, 2-(2-nitrophenyl)-1H-indole derivative (1) (0.2 mmol, 1.0 equiv.), copper powder (10 mol%), and K₂CO₃ (1.5 equiv.) were solubilized in dry toluene (0.8 mL). Then, N,N'-dimethylethylenediamine (DMEDA) (20 mol%), and the corresponding styrene (2) (2.0 equiv.) were added. The sealed tube was stirred at 135 °C for 20 hours.

After completion, the mixture reaction was filtered through a Celite[®] pad and concentrated under reduced pressure. Finally, the crude product was purified by silica gel chromatography with cyclohexane/ethyl acetate (0 to 15% of ethyl acetate) as eluent to give the corresponding *N*-vinylazole.

General procedure for Cadogan-like cyclization: synthesis of compounds 5a-h, 6a-g and 8a

In a sealed tube, 2-(2-nitrophenyl)-1-(1-aryllvinyl)-1H-indole (3) (0.1 mmol, 1.0 equiv.), MoO₂Cl₂(DMF)₂ (5 mol%) and PPh₃ (2.4 equiv.) were dissolved in toluene (3.0 mL). Next, the solution was degassed with an argon balloon from -196 °C in liquid nitrogen to room temperature before being warmed at 160 °C for 6 hours. After completion, the mixture reaction was filtered through a Celite[®] pad, and concentrated under reduced pressure. Finally, the crude product was purified by silica gel chromatography with cyclohexane/ethyl

acetate as eluent (0 to 20% of ethyl acetate) to give the corresponding compounds 5,6-dihydroindolo[1,2-c]quinazoline derivatives (5).

Conflicts of interest

Research Article

There are no conflicts to declare.

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