

REVIEW

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Heterocorroles: corrole analogues containing heteroatom(s) in the core or at a *meso*-positionBooruga Umasekhar,^a Vijayendra S. Shetti^{ID}*^b and Mangalampalli Ravikanth^{ID}*^a

Corroles are 18 π aromatic macrocyclic systems having one direct pyrrole–pyrrole linkage leading to a contracted cavity compared to porphyrins. Corroles exhibit contrasting coordination chemistry and properties compared to porphyrins. Structural modification of corroles by introducing a heteroatom in their aromatic conjugation circuit *i.e.*, either in the core or at a *meso* position leads to a new class of corrinoids called heterocorroles. The core modification strategy includes replacing one or two core nitrogen atom(s) with O, S or C atoms and *meso*-modification involves replacing the *meso*-carbon atom at the 10-position with NH, NR, O, S, Se or Si atoms. This review article presents an overview of the progress in heterocorrole chemistry including their syntheses, key structural aspects and properties.

1. Introduction

Hodgkin's work on the X-ray crystallography of β -substituted free-base corrole revealed for the first time that corroles resemble corrin, a cobalt coordinating ligand in vitamin B₁₂.¹ Corroles are tetrapyrrolic macrocycles which have a direct pyrrole–pyrrole linkage similar to corrin and preserve the aromaticity of porphyrins despite having one less *meso*-carbon atom (Fig. 1).² Corroles differ from their congener porphyrin in having a contracted cavity and three ionisable core NH atoms.

This leads to a contrasting coordination chemistry of corroles where they can stabilize metal ions in a higher oxidation state unlike porphyrins and also can exhibit unique reactivity for low valent metallocorroles.³ Corroles' ability to absorb light

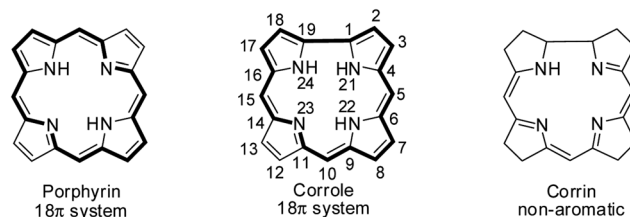


Fig. 1 Skeletal structure of porphyrin, corrole and corrin.

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assignments at Taiwan and South Korea, he joined BMS College of Engineering, Bengaluru as an Assistant Professor of Chemistry in 2014. His current research is focused on synthesis of novel π conjugated molecules for organic electronic applications.

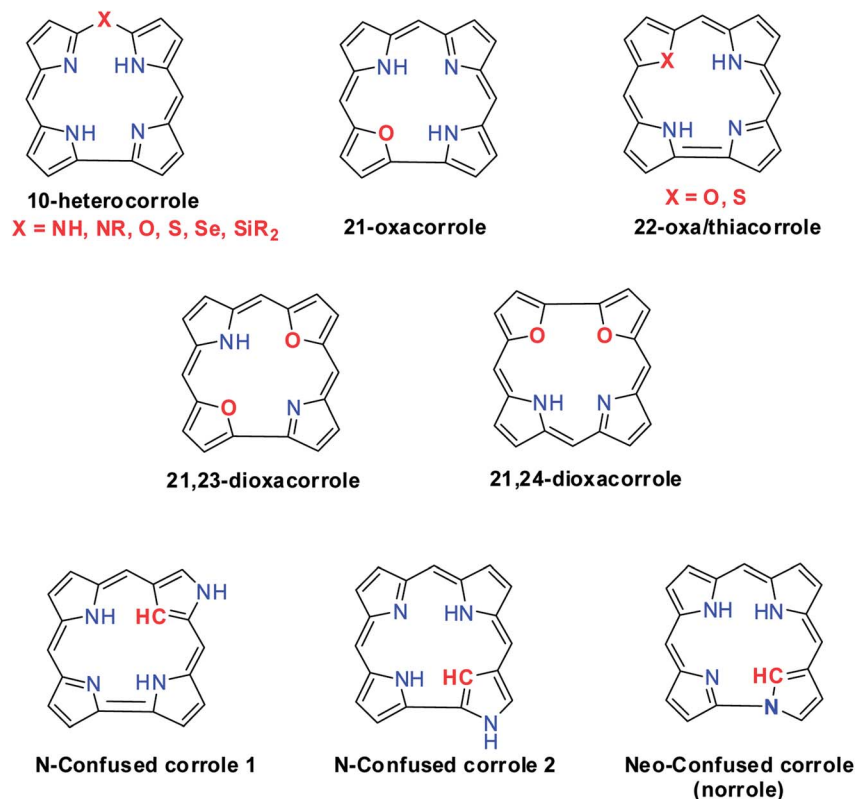
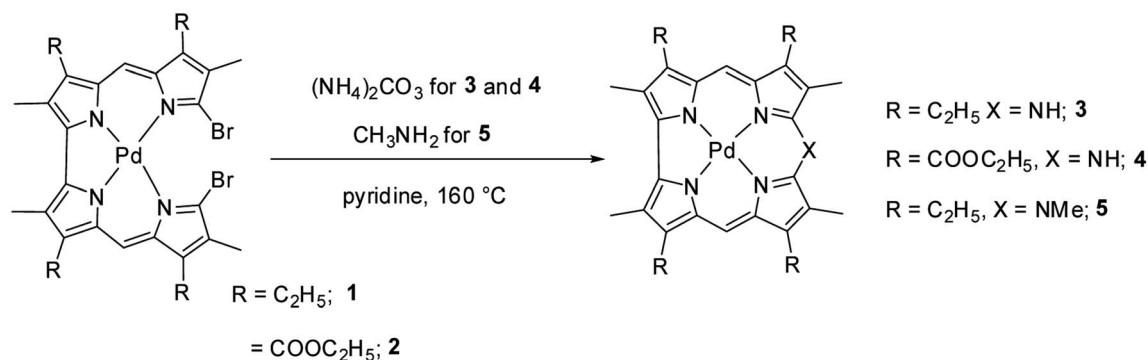


Fig. 2 Various examples of heterocorrole framework.



Scheme 1 Synthesis of β -substituted 10-azacorroles **3**–**5**.

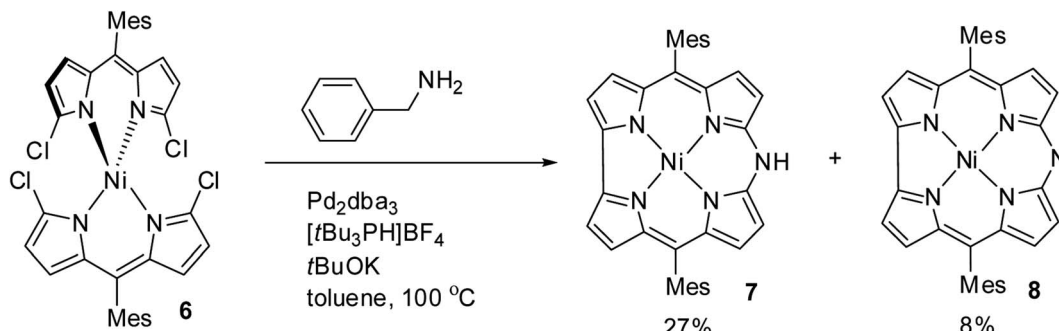


M. Ravikanth was born in India, in 1966. He received his BSc and MSc from Osmania University, Hyderabad and PhD from Indian Institute of Technology, Kanpur in 1994. After his postdoctoral stay in USA and Japan, he joined as a faculty at Indian Institute of Technology Bombay, where he is currently a full professor. His current research interest includes porphyrin and related macrocycles and boron dipyrromethenes.

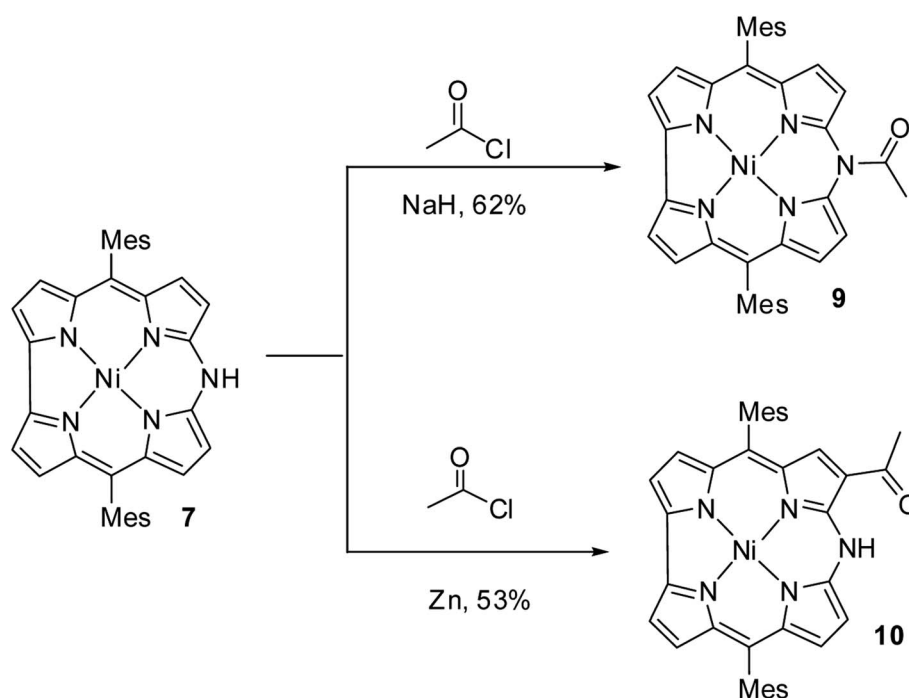
throughout the visible spectral range, their enhanced luminescence yield, their good photostability in most solvents and their relative ease of oxidation give them an edge over porphyrins in many applications.^{4–6}

Structural modification of corroles by introducing a heteroatom on their aromatic conjugation circuit *i.e.*, either at the core or at a *meso* position leads to a new class of corrinoids called heterocorroles.^{2,7,8} The core modification strategy includes replacing one or two core nitrogen atom(s) by oxygen, sulfur (21-, 22-, 21,23-heterocorroles) or carbon atoms (confused corrole) and *meso*-modification involves replacing *meso*-carbon atom at the 10-position by NH, NR, O, S, Se or even Si atoms (10-heterocorroles) (Fig. 2). Heterocorroles represent a hybrid class of molecules which have contracted cavity similar to corroles and dipyrromethenes.





Scheme 2 Synthesis of 10-azacorroles 7 and 8 by Buchwald–Hartwig reaction.



Scheme 3 N- and C-acylation reactions of 10-azacorrole 7.

are dianionic ligands like porphyrins. Because of this, they can serve as appropriate reference systems to study metallocorroles. In this review article, we wish to present an overview of the progress in the area of heterocorroles, especially in the area of core-modified corroles and mono *meso*-modified corroles. However, the discussion on corrolazines is not included in this article.

2. Corroles containing heteroatom at 10-position (*meso*-modified corroles)

2.1. 10-Azacorroles

Johnson, Kay and Rodrigo in their pioneering work⁹ synthesized β -substituted palladium corroles 3–5 containing nitrogen atom at one of the *meso* positions starting from Pd(II) chelates of dibromo 5,5'-bi(dipyrromethene) 1 and 2 (Scheme 1). Authors' attempts to obtain free base corroles by demetallating the

central metal atoms by acid treatment was not completely successful.

Shinokubo and co-workers¹⁰ reported the earliest synthesis of *meso*-aryl substituted azacorroles 7 & 8 by utilizing Buchwald–Hartwig coupling reaction between α,α' -dichlorodipyrin Ni(II) complex 6 and benzylamine (Scheme 2). Though, this reaction resulted in a mixture of azacorroles 7 and 8, column chromatographic separation enabled authors to isolate 7 in 27% yield. ¹H NMR spectral feature and nucleus independent chemical shift (NICS) studies revealed the aromatic character of azacorrole 7. To demonstrate regio-selective functionalization and the acidic nature of N–H bond at *meso*-position, authors performed acylation reaction of 7 with acetyl chloride (Scheme 3). The acylation reaction of 7 in the presence of a base resulted in the formation of N-acetyl product 9 in 62% yield. When the C-acylation was performed by using acetyl chloride and zinc, C8-acylated product 10 was obtained in 53% yield which is in



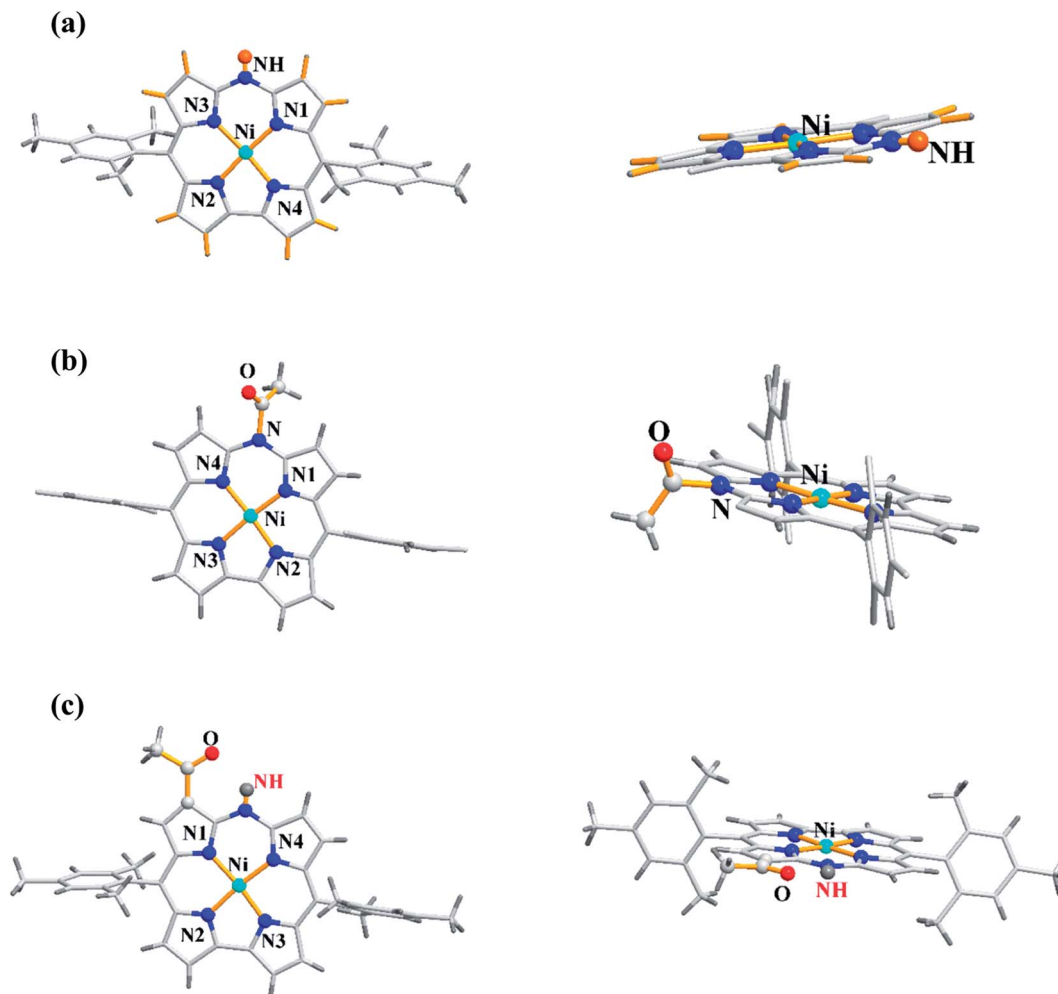


Fig. 3 Top and side view of X-ray crystal structures of (a) 7 (CCDC 861382) (b) 9 (CCDC 861384) and (c) 10 (CCDC 861385).

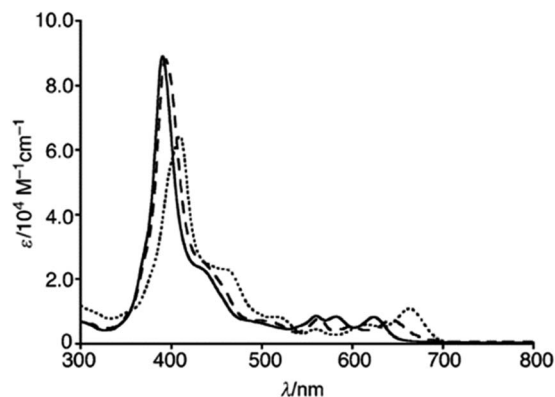


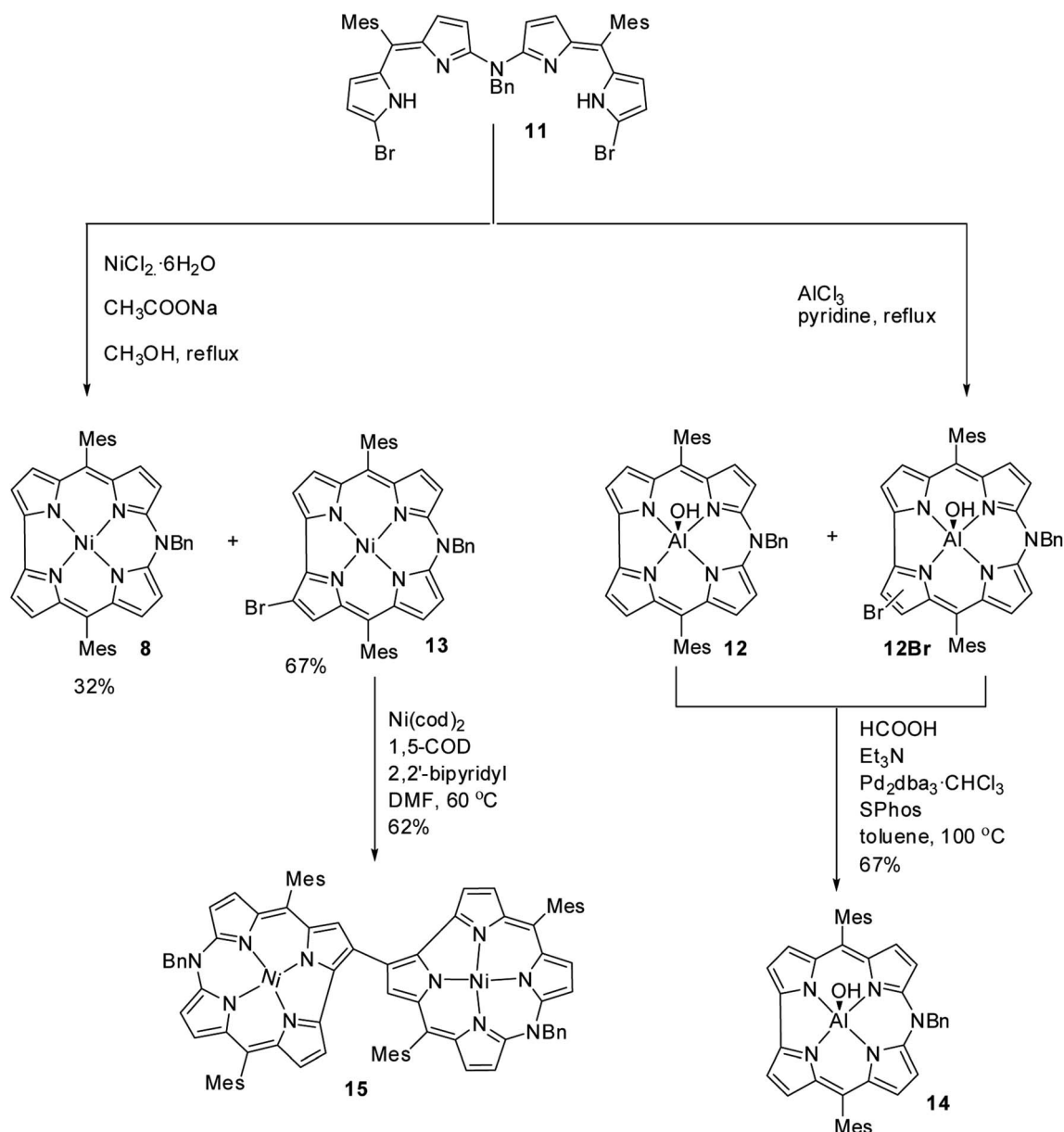
Fig. 4 UV-vis spectra of 7 (normal line), 9 (dashed line) and 10 (dotted line) recorded in dichloromethane. Reproduced from ref. 10 with permission. Copyright© 2012 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

contrast to the electrophilic substitution pattern noticed in normal corroles which generally yield C2 and C17 substituted products.

The single crystal X-ray analysis revealed highly planar structures of azacorroloides 7, 9 and 10 (Fig. 3). The orientation of acetyl moiety in 9 was found to be perpendicular to the corrole plane, whereas in 10, the acetyl group was coplanar with corrole plane due to its intramolecular hydrogen bonding interaction with *meso*-NH proton. This resulted in an expanded conjugation leading to a substantial bathochromic shift of Q-like bands of 10 compared to 7 and 9 (Fig. 4). The electrochemical studies showed that the first oxidation of corrole 7 was rather easy compared to acylated corroles 9 and 10.

Shinokubo, Hiroto and co-workers synthesized Ni(II) and Al(III) complexes of 10-azacorroloide by using coordination induced cyclization of nitrogen bridged bisdipyrrin.¹¹ The bisdipyrrin containing two bromo groups at α -position 11 when reacted with excess $\text{NiCl}_2 \cdot 6\text{H}_2\text{O}$ yielded Ni(II) azacorroloide 12 along with 2-bromo-10-azacorroloide 13. The reaction of same bisdipyrrin with excess AlCl_3 resulted in a mixture of Al(III) azacorroloides including regioisomeric bromoazacorroloides which upon subsequent





Scheme 4 Synthesis of 2-bromo-10-azacorrole **13**, Al(III) azacorrole **14** and corrole dyad **15**.

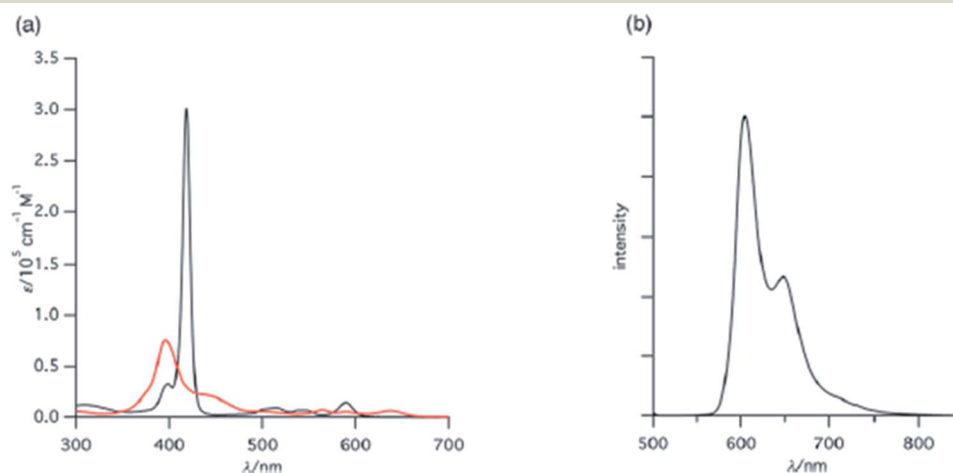
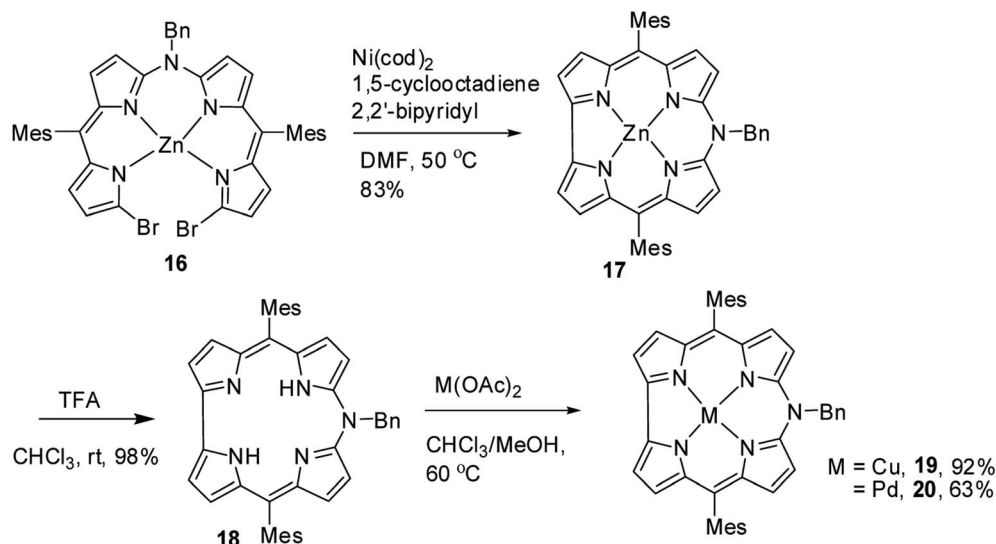


Fig. 5 (a) UV-vis spectra of **12** (red), **14** (black) (b) emission spectrum of **14** ($\lambda_{\text{ex}} = 420$ nm) recorded in dichloromethane. Reproduced from ref. 11 with permission. Copyright© 2016 Royal Society of Chemistry.





Scheme 5 Synthesis of free base 10-azacorrole **18** and its metal complexes **17**, **19** & **20**.

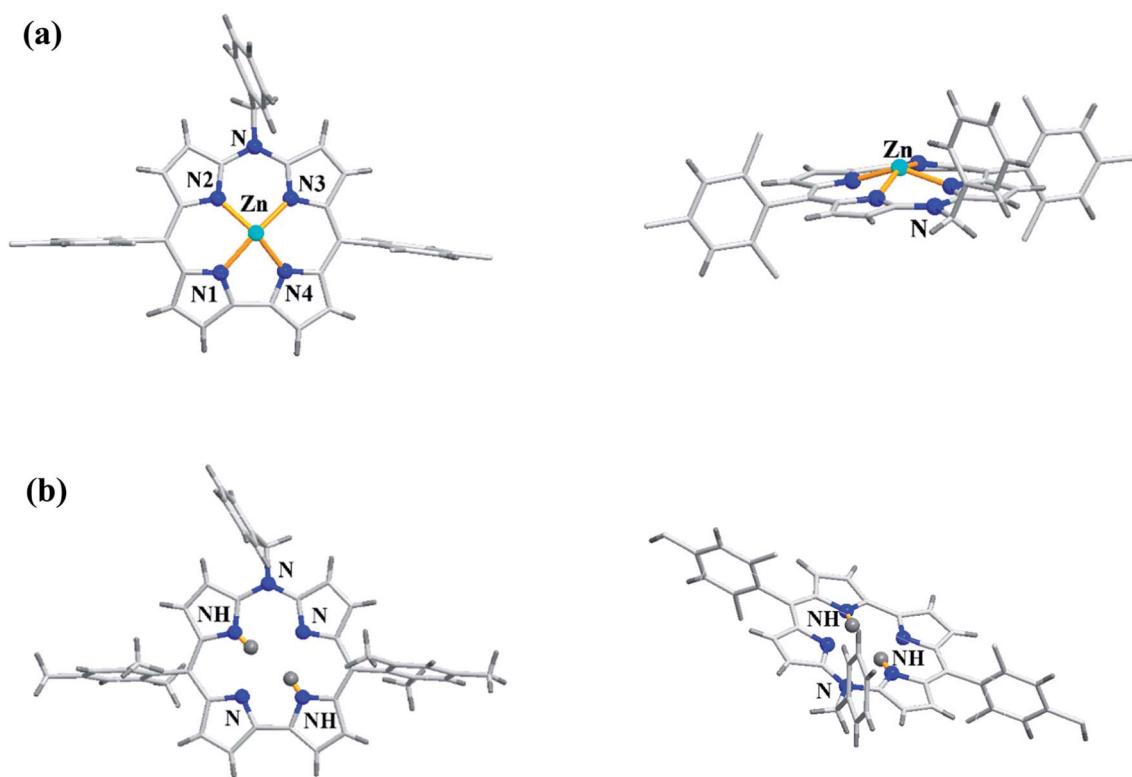
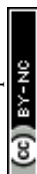


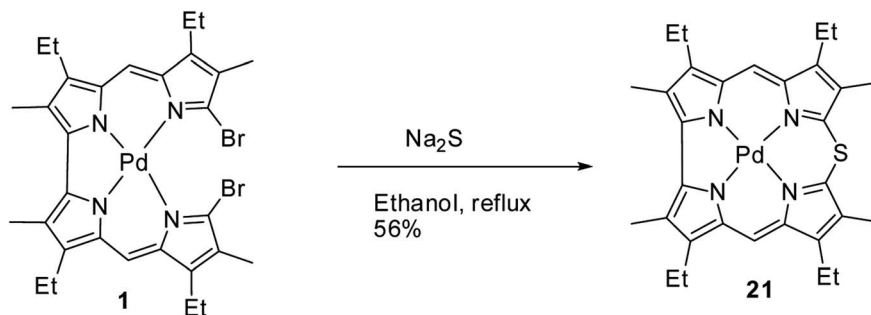
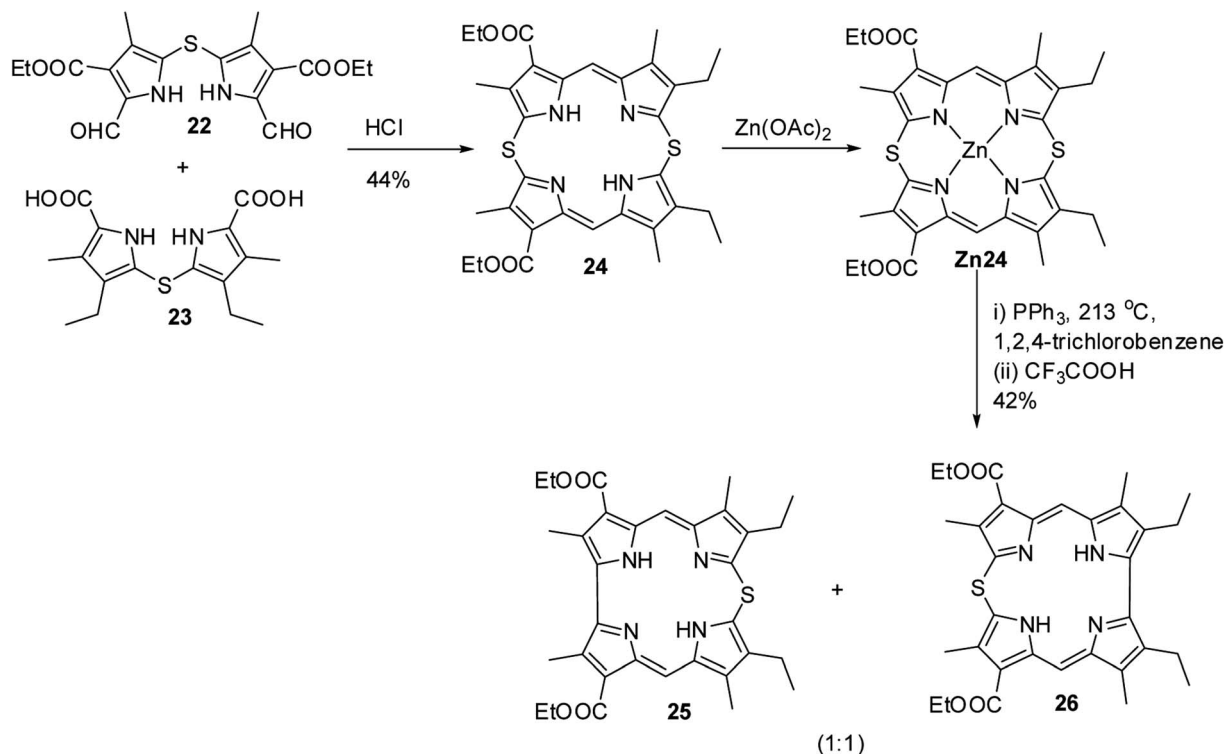
Fig. 6 Top and side view of X-ray crystal structures of (a) **17** (CCDC 1477606) (b) **18** (CCDC 1477608).

reductive dehalogenation afforded Al(III) 10-azacorrole **14** in 67% yield (Scheme 4). The 2-bromo-10-azacorrole **13** was further subjected to coupling reaction in presence of $\text{Ni(cod)}_2/2,2'$ -bipyridyl (Yamamoto conditions) yielding to a 2,2'-linked corrole dimer **15** (Scheme 4). The Al and Ni corroles displayed contrasting absorption spectral features compared to their porphyrin counterparts. The Al corrole **14** exhibited a sharp Soret band and hypsochromically shifted lowest energy

absorption band compared to the Ni corrole **12** (Fig. 5). The Al corrole **14** exhibited larger oxidation and reduction potential values compared to the Ni corrole due to electron withdrawing nature of Al(III). The Al(III) corrole **14** also represented the first examples of emissive 10-azacorroles with moderate quantum yield (Fig. 5).

As demetallation of Ni(II) and Al(III) from above corroles was challenging and unsuccessful, the same authors, in their



Scheme 6 Synthesis of β -substituted Pd(II) 10-thiacorrole **21**.Scheme 7 Synthesis of β -substituted free base 10-thiacorroles **25** and **26**.

pursuit to find a synthetic protocol to make free base *meso*-azacorroles, devised a method involving Zn(II) complex of bisdipyrrin **16**.¹² The Zn(II) bisdipyrrin **16** was subjected to Yamamoto reaction conditions to facilitate cyclization leading to Zn(II) 10-azacorrole (Scheme 5). The TFA induced demetallation yielded corresponding free base 10-azacorrole **18** in quantitative yield. The single crystal X-ray structures of compound **17** and **18** are shown in Fig. 6. The single crystal X-ray analysis of zinc corrole **17** showed a pentacoordinated Zn(II) in the core and the mean plane deviation of zinc (0.088 Å) was found to be greater than the corresponding Ni complex **8** (0.020 Å). The X-ray structure of free base corrole **18** showed a highly planar structure with two core-NH atoms. The chemical shift value of the core NH resonance in ¹H NMR spectrum (5.23 ppm) reflected on its lower aromatic character. The Cu(II) **19** and Pd(II) **20** azacorroles were synthesized by metallating free base azacorrole **18**

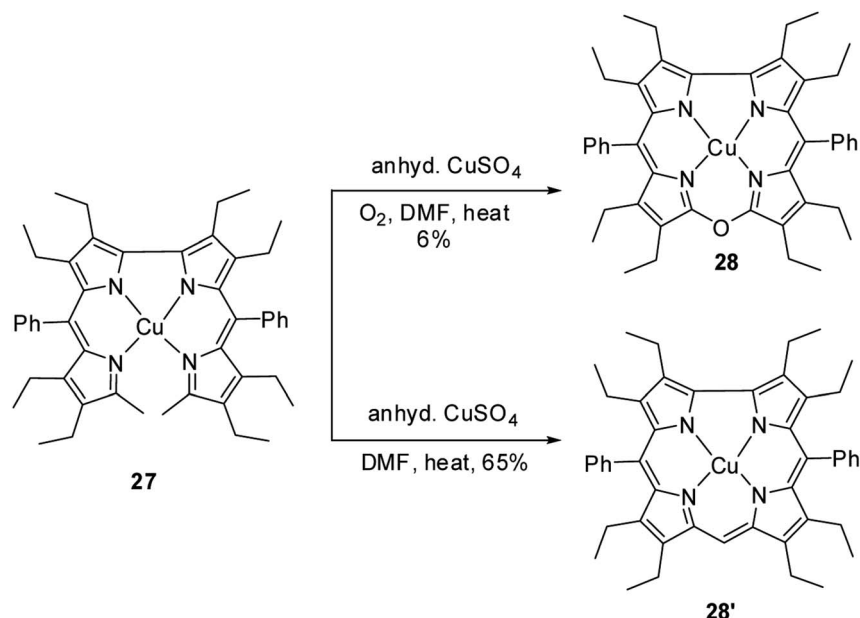
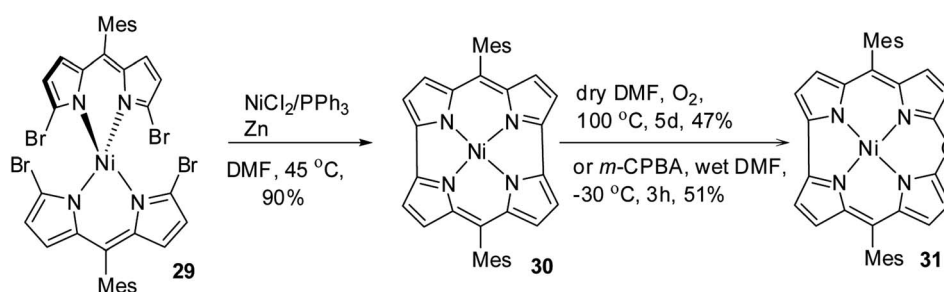
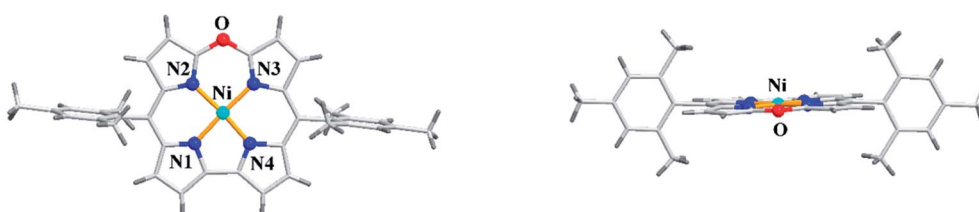
with acetates of copper and palladium respectively. The Zn(II) **17** and Cu(II) **19** azacorroles showed red shifted absorption bands compared to free base azacorrole **18**. The electrochemical studies revealed marginally reduced HOMO–LUMO gap for Zn(II) **17** and Pd(II) **20** corroles compared to their free base analogue. The fluorescence quantum yield of zinc corrole **17** was comparable to earlier reported Al(III) azacorrole **14**.

2.2. Corroles containing chalcogen atom at 10-position

Johnson, Kay and Rodrigo in their 1963 paper,⁹ also reported the synthesis of palladium derivative of β -substituted *meso*-thiacorrole **21** by treating palladium chelate of 5,5'-bi(dipyrromethene) **1** with sodium sulphide in ethanol at refluxing temperature (Scheme 6).

Further, in 1972, Grigg, Broadhurst and Johnson¹³ reported the synthesis of β -substituted *meso*-thiacorroles **25** & **26** by



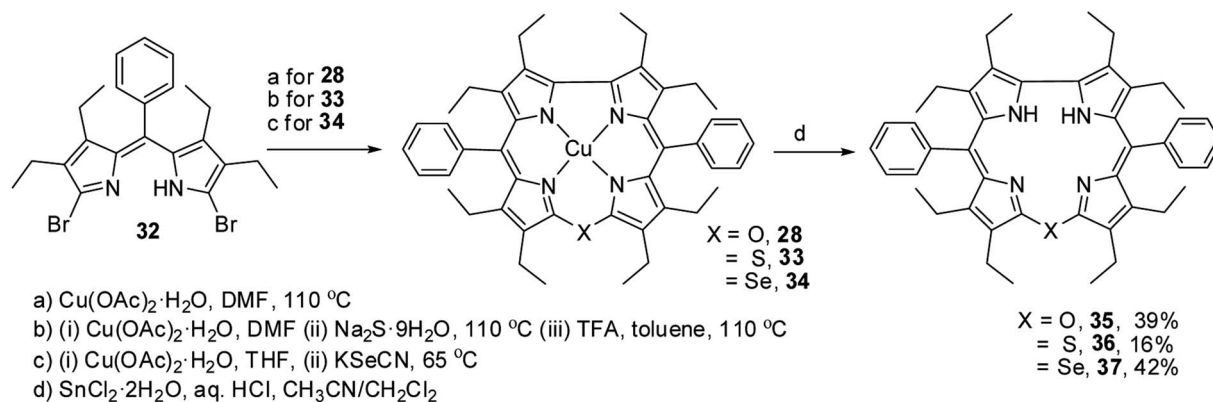
Scheme 8 Synthesis of copper derivatives of 10-oxacorrrole **28** and normal corrrole **28'**.Scheme 9 Synthesis of *meso*-substituted Ni(II) 10-oxacorrrole **31**.Fig. 7 Top and side view of X-ray crystal structure of **31** (CCDC 883625).

thermally driven sulfur extrusion reaction of Zn(II) complex of *meso*-dithiamacrocycle **Zn24** in presence of triphenylphosphine (PPh_3) followed by TFA assisted demetallation (Scheme 7). However, after these two initial reports on *meso*-thiacorrroles, the further developments in this area were halted for almost four decades presumably owing to the difficulties involved in the synthesis of such macrocycles.

Bröring, Holthausen and co-workers¹⁴ reported the synthesis of copper derivative of 10-oxacorrrole **28** by anhydrous copper(II) sulphate assisted oxidative macrocyclization of 2,2'-bidipyrins **27** in the presence of oxygen (Scheme 8). The oxacorrrole **28** was

used as an appropriate reference compound to study the electronic ground state of copper in normal corrrole **28'** as it has similarity in cavity size to **28'** and its resemblance to porphyrins in having double negative charge arising from its core. Absorption spectroscopy of both **28** and **28'** confirmed their typical porphyrinoid macrocyclic structure. The normal corrrole **28'** was EPR silent whereas the oxacorrrole **28** exhibited a EPR pattern typical of Cu(II) porphyrinoid. However, the comparison of X-ray structures of copper corrrole **28'** with isostructural **28** followed by a detailed computational study on **28'** highlighted the possible electronic interaction between central Cu ion and





Scheme 10 Synthesis of octaethyl Cu(II) 10-heterocorroles **28**, **33**, **34** and their free base analogues **35**–**37**.

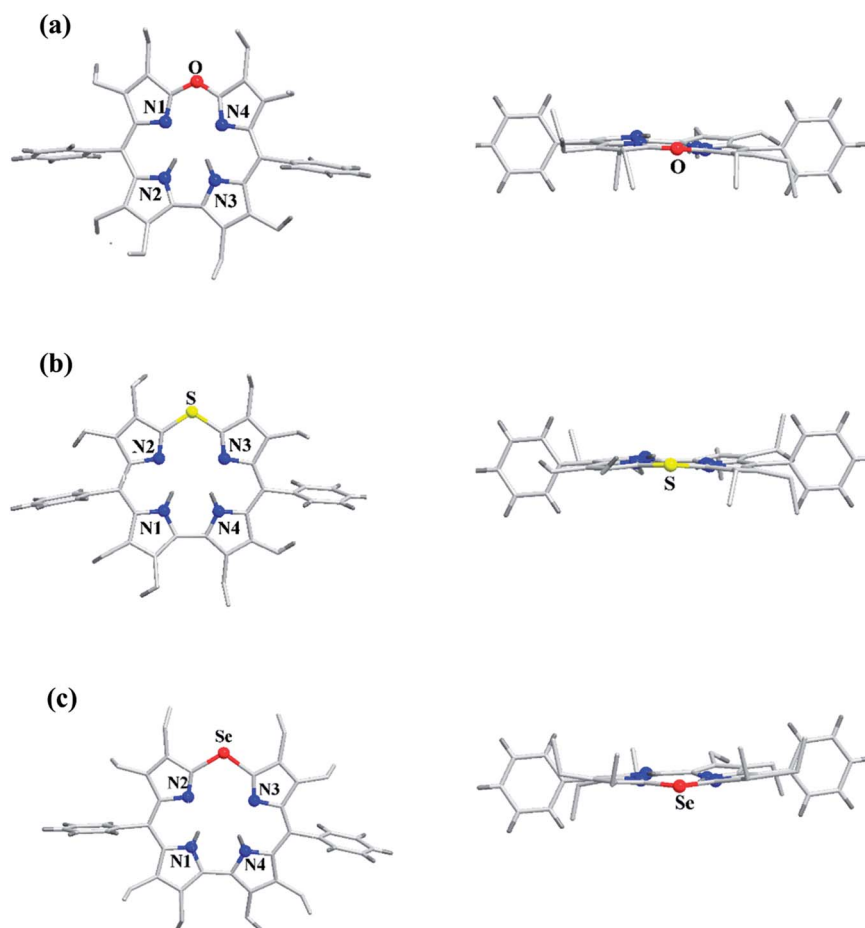


Fig. 8 Top and side view of X-ray crystal structures of (a) **35** (CCDC 921878) (b) **36** (CCDC 921879) and (c) **37** (CCDC 921880).

macrocyclic π -orbital and hence led authors to conclude that copper's oxidation state in corrole **28'** can be better addressed as Cu(II) rather than Cu(III).

Shinokubo, Kobayashi and co-workers¹⁵ reported a remarkable gram-scale synthesis of strained macrocycle called Ni(II) norcorrole **30**, which lacks two *meso*-carbon atoms compared to porphyrins.¹⁶ This Ni(II) norcorrole **30** showed a very good solid and solution state stability for an antiaromatic compound.

However, authors observed that norcorrole was slowly getting oxidized in the presence of air at elevated temperatures to form an aromatic Ni(II) 10-oxacorrole **31** (Scheme 9), where the lone pair of electron on the *meso*-oxygen contributes to aromaticity.

Alternatively, the treatment of norcorrole **30** with *m*-chloroperbenzoic acid (*m*-CPBA) led to formation of Ni(II) 10-oxacorrole **31** in 51% yield (Scheme 9). The single crystal X-ray analysis showed a planar conformation for oxacorrole **31** (Fig. 7)



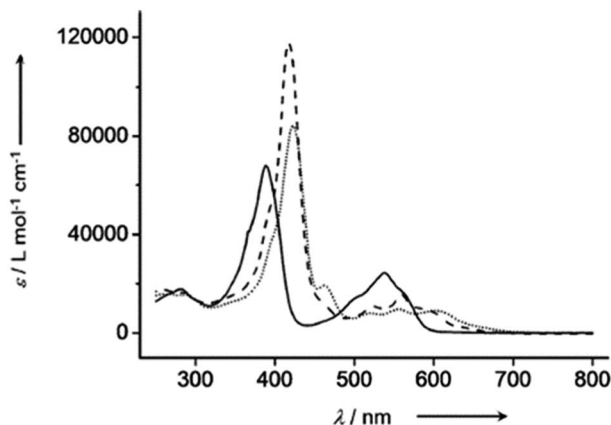
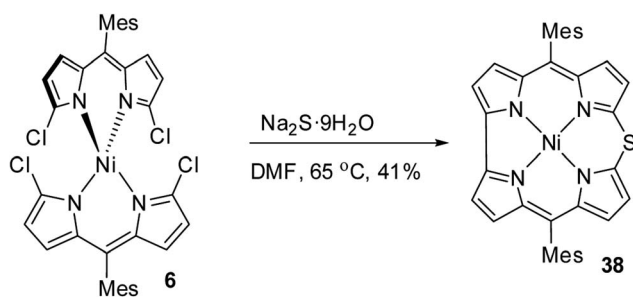


Fig. 9 UV-vis spectra of **35** (normal line), **36** (dashed line) and **37** (dotted line) recorded in dichloromethane. Reproduced from ref. 17 with permission. Copyright© 2013 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.



Scheme 11 Synthesis of *meso*-substituted Ni(II) 10-thiacorrole **38**.

and the absorption and Magnetic Circular Dichroism (MCD) studies revealed features typical of aromatic porphyrinoid with lower molecular symmetry.

Bröring and co-workers¹⁷ synthesised a series of octaethyl derivatives of 10-heterocorroles **35–37** containing O, S, Se atoms at the 10-position by a metal-promoted two-step macrocyclization of dipyrin **32** as depicted in Scheme 10.

The chemical shift position of core NH resonance for free base corroles **35–37** and *meso*-H resonance for *meso*-

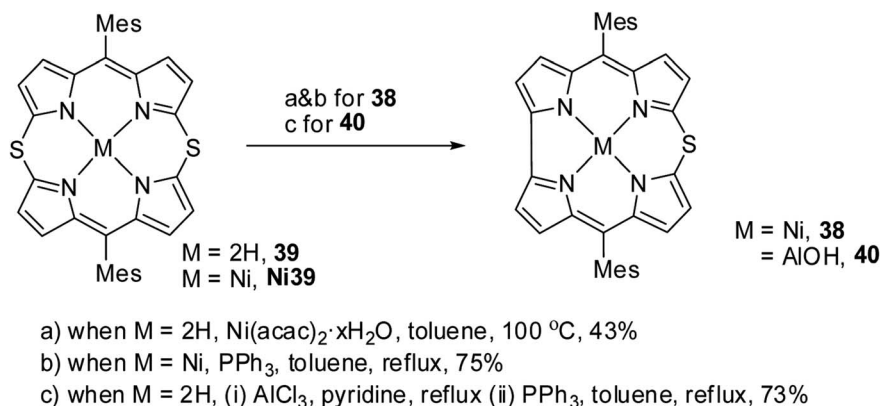
unsubstituted analogues of **35–37** (not shown in the scheme) revealed that the aromaticity of these macrocycles follow the order: 10-thiacorrole **36** > 10-selenacorrole **37** > 10-oxacorrole **35**. The X-ray structure of 10-heterocorroles **35–37** showed that they are almost planar with little saddle-shaped distortions and both the core-NH protons were located on the N-atoms of bipyrrolic unit (Fig. 8). The macrocyclic cavity size was found to increase from oxa **35** (7.05 Å²) to thia **36** (7.57 Å²) to selenacorroles **37** (7.77 Å²). The thia **36** and selenacorroles displayed absorption spectral bands typical of porphyrinoids whereas the oxacorrole **35** showed broadened and hypsochromically shifted absorption bands expected for a macrocycle with diminished aromatic nature (Fig. 9).

Shinokubo and co-workers¹⁸ achieved the synthesis of *meso*-substituted Ni(II) 10-thiacorrole **38** by sulfidation of α,α' -dichlorodipyrin Ni(II) complex **6** as shown in Scheme 11.

Alternatively, the insertion reaction on 5,15-dithiaporphyrin **39** in refluxing toluene and sulfur extrusion reaction of nickel dithiaporphyrin **Ni39** upon treatment with triphenylphosphine resulted in the formation of Ni thiacorrole **38**. Also, Al(III) thiacorrole **40** was synthesized by refluxing dithiaporphyrin **39** with AlCl₃ in pyridine (Scheme 12). The same research group also observed similar sulfur extrusion reaction when Cu(II) 5,15-dithiaporphyrin was treated with PPh₃ to yield Cu(II) 10-thiacorrole (not shown in the scheme).¹⁹ The ¹H NMR spectral studies and absorption spectral patterns revealed that Al corrole **40** possessed strong aromaticity compared to Ni corrole **38**.

The X-ray structure of Ni(II) 10-thiacorrole **38** showed a planar conformation and the bond length of C–S (1.698(8) Å and 1.680(8) Å) was decreased compared to typical C–S bond length (1.82 Å) (Fig. 10) highlighting the partial double bond character of C–S bond and hence an indication of effective delocalization of sulfur's lone pair into the macrocyclic π system.

Kong and co-workers²⁰ reported the synthesis of a *meso-meso* linked porphyrin–corrole hybrid (triad) in which 10-thiacorrole was flanked by two porphyrin units (Scheme 13). Porphyrinyl dipyrromethane **41** upon consecutive bromination, 2,3-dichloro-5,6-dicyano-benzoquinone (DDQ) oxidation and treatment with Ni(II) salt resulted in an unstable intermediate **42**, which on treatment with Na₂S·9H₂O yielded triad **43** in 41%



Scheme 12 Synthesis of Ni(II) 10-thiacorrole **38** and Al(III) thiacorrole **40** from 5,15-dithiaporphyrin.

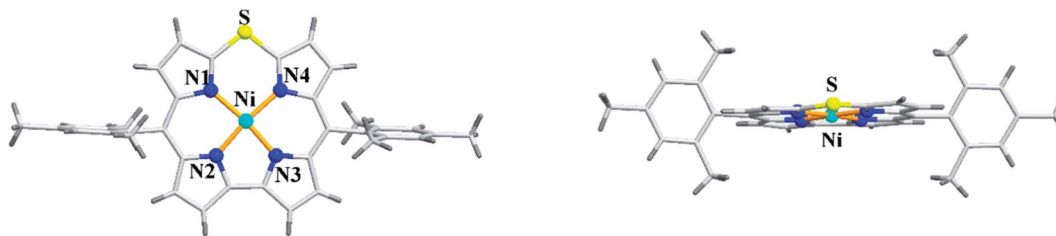
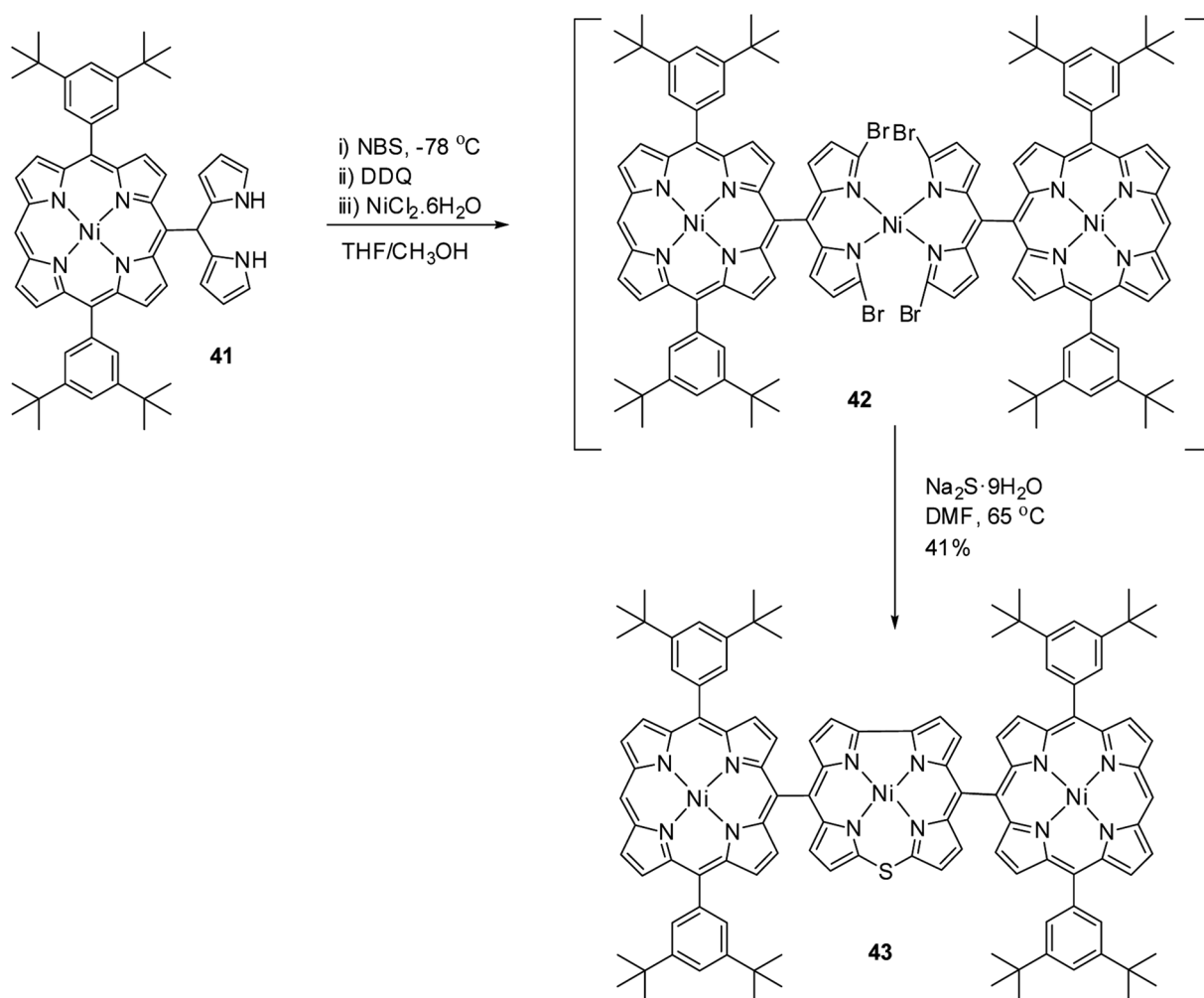


Fig. 10 Top and side view of X-ray crystal structure of **38** (CCDC 893774).



Scheme 13 Synthesis of *meso-meso* linked porphyrin-corrole hybrid **43**.

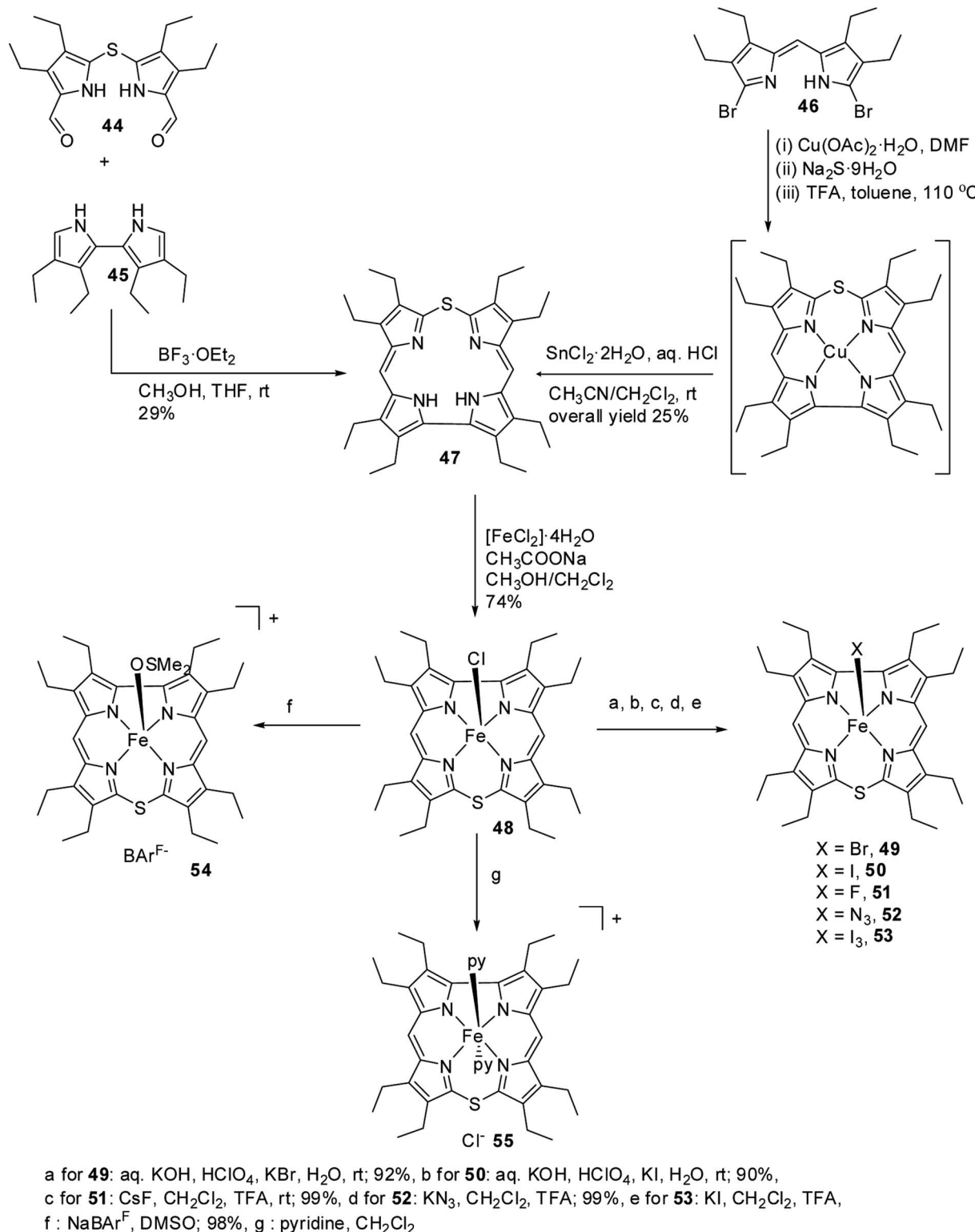
yield. In triad **43**, the β -CH resonances of porphyrin subunits were slightly downfield shifted compared to thiacorrole unit and **43** displayed a broad absorption spectral band.

Bröring and co-workers^{21,22} carried out a systematic study on a series of iron 10-thiakorroles **48–55** containing halogenido, pseudo-halogenido and solvent-derived axial ligands to study the ground state spin state of Fe(III) ion in such complexes. The free base octaethyl-10-thiakorrole **47** was synthesized by either MacDonald-type [2 + 2] condensation of bis(3,4-diethyl-5-formylpyrrole)sulphide **44** and tetraethyl-2,2'-bipyrrole **45** or

Cu(II) templated macrocyclization of dibromodipyrin **46** (Scheme 14).

The synthesis of Fe(III) complexes of **47** with halogenido **48–51**, pseudo-halogenido **52, 53** and solvent-derived axial ligands **54, 55** are also shown in the Scheme 14. The X-ray structures of compounds **48, 52**, and **54** are shown in Fig. 11. The iron corroles containing halogen axial ligand **48–51** showed porphyrinoid like absorption features such as intense Soret band and several minor Q-bands. Most of these iron corroles (**48–50, 52–54**) displayed elongated Fe-axial ligand and short equatorial Fe–

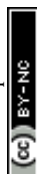




Scheme 14 Synthesis of octaethyl-10-thiacorrole **47** and its Fe(III) complexes **48–55** with various axial ligands.

N bonds compared to iron porphyrins. The chemical shift values for *meso*-CH protons were in the down field region ($\delta = 35\text{--}50$ ppm). They also displayed rhombic EPR signals; g_z strongly split by superhyperfine coupling with axial donor's

nuclear spin. The Mossbauer spectra revealed their large quadrupole splitting values (ΔE_Q of >2.5 mm s⁻¹) (Fig. 12). The redox potential values of these corroles exhibited cathodic shift in comparison to iron porphyrins. All these observations served



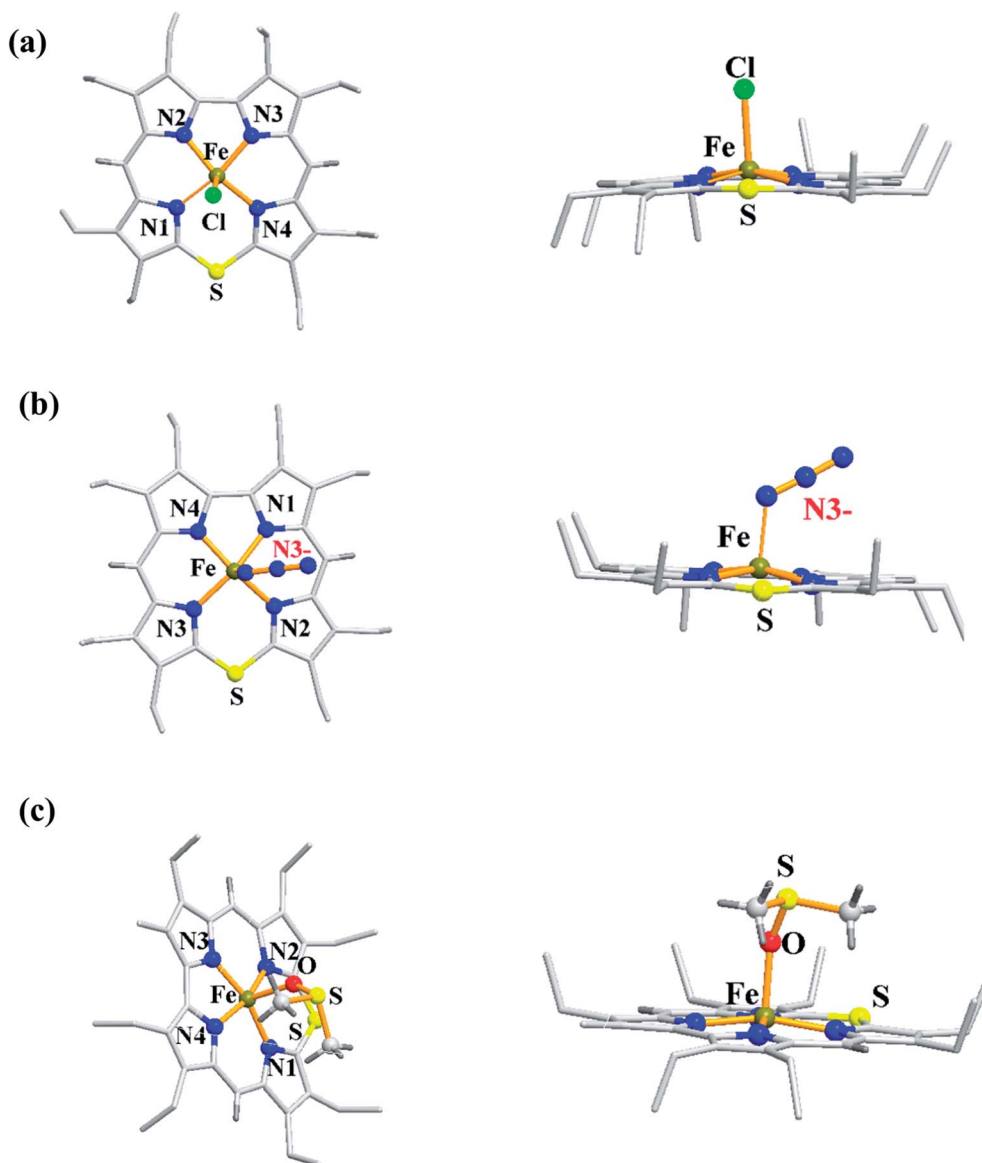


Fig. 11 Top and side view of X-ray crystal structures of (a) **48** (CCDC 962051) (b) **52** (CCDC 962050) and (c) **54** (CCDC 962046).

as a corroborative evidence to support the fact that smaller cavity size of these iron *meso*-thiacorroles resulted in an intermediate spin (i.e., $S = 3/2$) state for ground state of Fe(III) ion. This intermediate spin state was found to be unperturbed and was temperature independent. This observation is in contrast to Fe(III) porphyrins which typically display high spin ($S = 5/2$) nature.

Bröring and co-workers²³ further studied Ni(II), Cu(II) and Pd(II) complexes of 10-heterocorrole ligands containing O, S, Se at 10-position (35–37) to understand macrocycle's adaptability towards metal ions of different sizes while forming complexes. The synthesis of Ni, Cu, and Pd derivatives of these corroles is shown in Scheme 15. Their findings, mainly based on the X-ray structural data, revealed that ruffling dominates as out-of-plane distortion mode in complexes where a mismatch between the

ligand cavity size and the radius of metal ion exists. The maximum ruffling was observed in the case of nickel 10-selenacorrole **58** (Fig. 13). This paper, for the first time, revealed that corroles can indeed exhibit ruffling conformations while forming complexes despite the presence of a direct pyrrole-pyrrole connection.

2.3 10-Silacorroles

Recently, Shinokubo, Hiroto and co-workers²⁴ reported the synthesis of Ni(II) (**62**, **63**) free base (**64**, **65**), and Zn(II) (**66**, **67**) complexes of 10-silacorroles as shown in the Scheme 16. The Pd catalyzed silylation reaction of bis(α, α' -dibromodipyrin) Ni(II) complex **29** resulted Ni(II) silacorrole **62** and **63**, which upon demetallation and subsequent zinc insertion reaction yielded free base (**64**, **65**) and Zn(II) silacorrole (**66**, **67**) respectively.



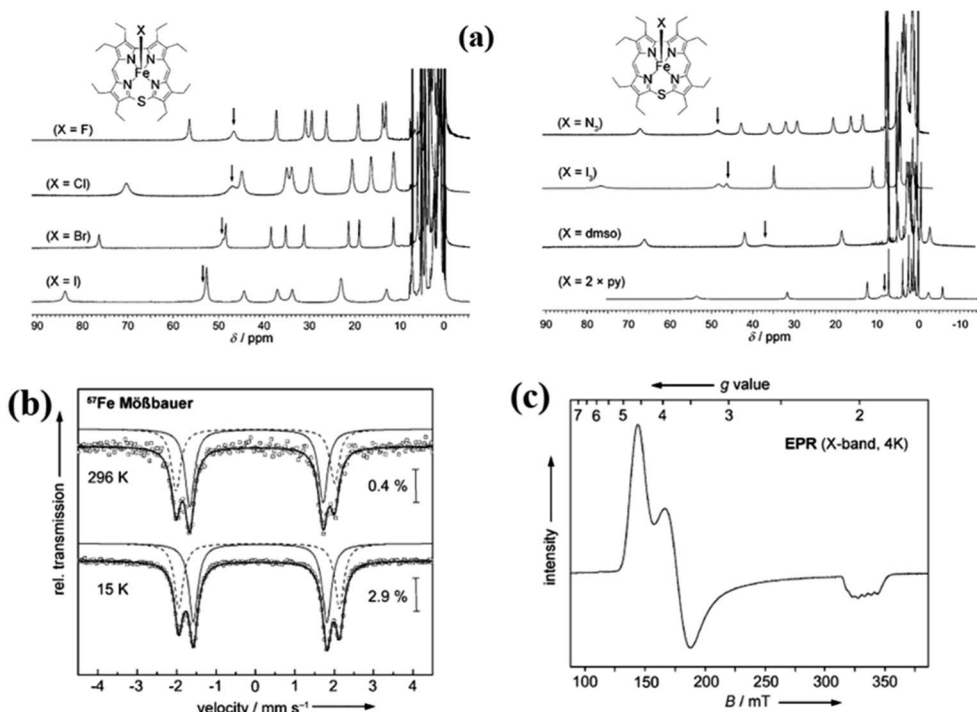
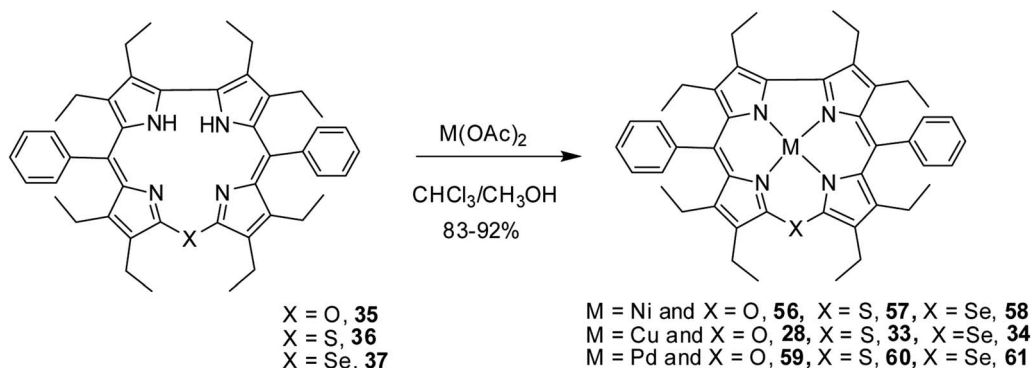


Fig. 12 (a) ^1H NMR spectra of iron(III) thiacorroles **48–55** in CDCl_3 (b) ^{57}Fe Mössbauer spectra of **50** (powder) (c) EPR spectrum of **50** (solution, $\text{CHCl}_3/\text{CH}_2\text{Cl}_2$ 1 : 1, 4 K). Reproduced from ref. 21 with permission. Copyright© 2014 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.



Scheme 15 Synthesis of Cu(II) (**28**, **33**, **34**) Ni(II) (**56–58**), and Pd(II) (**59–61**) complexes of 10-heterocorrole **35–37**.

Although the Ni(II) and free base silacorroles adopt highly planar conformation as per their X-ray structures (Fig. 14), the ^1H NMR spectral features such as β -pyrrolic proton resonances ($\delta = 5.99$ to 6.31 for Ni silacorrole **62**) and core NH resonance ($\delta = 15.1$ ppm for free base silacorrole **64**) were indicative of their non-aromatic nature. The absorption spectral band of Ni corrole was far red-shifted than zinc and free base corroles and reaches into NIR region (Fig. 15). The zinc silacorrole **66** also exhibited emission in NIR region (Fig. 15) with fluorescence quantum yield = 0.014 and emission lifetime = 1.78 ns. The electrochemical and theoretical studies carried out on these silacorroles showed the presence of $\sigma^*-\pi^*$ conjugation between the silylene group and the tetrapyrrole system. Also, their electronic structures were susceptible to the central metal ions and the substituents present on silicon at *meso*-position.

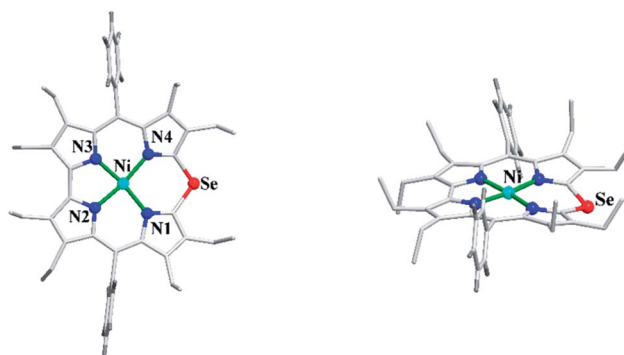
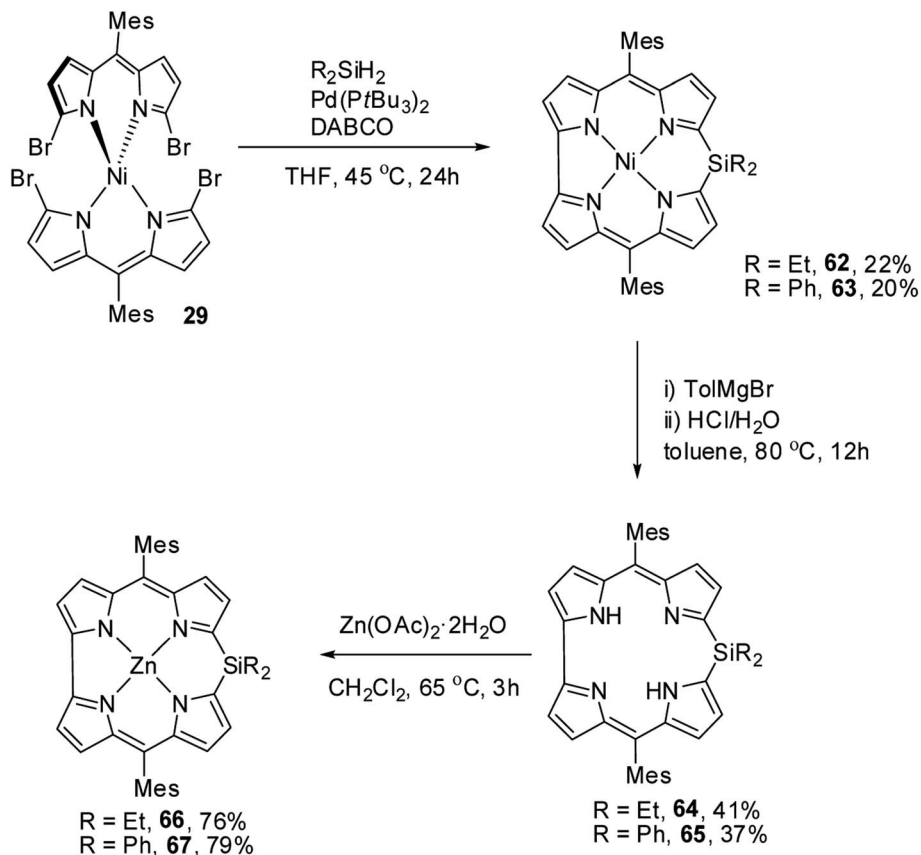


Fig. 13 Top and side view of X-ray crystal structure of **58** (CCDC 1544170).



Scheme 16 Synthesis of Ni(II) (**62**, **63**) free base (**64**, **65**), and Zn(II) (**66**, **67**) complexes of *meso*-substituted 10-silacorrroles.

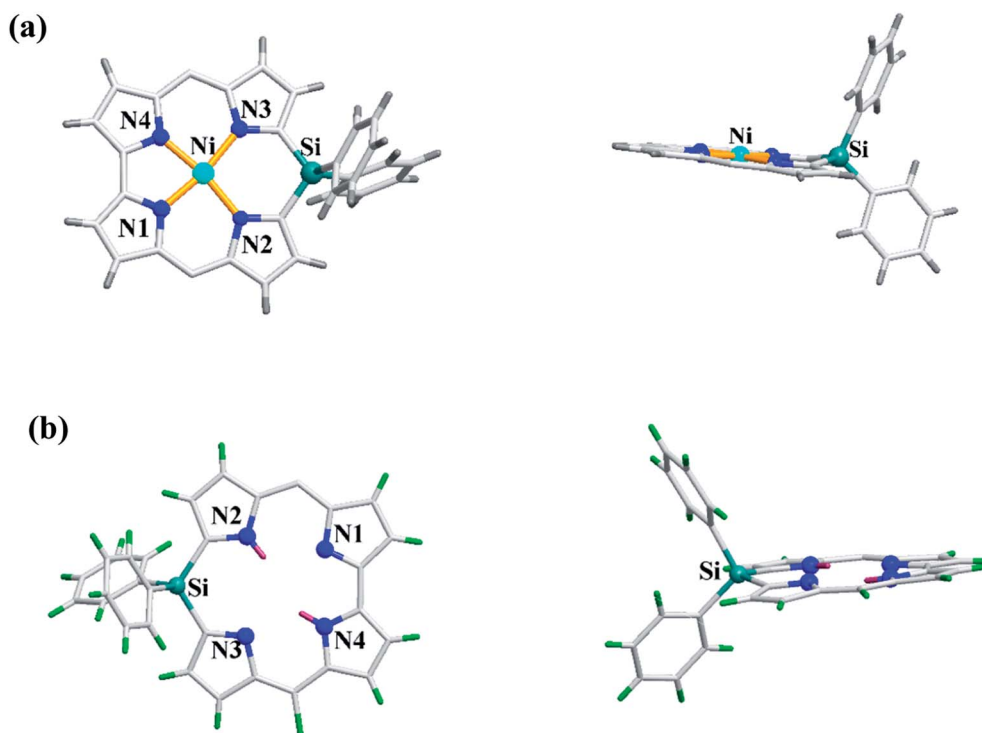


Fig. 14 Top and side view of X-ray crystal structures of (a) **63** (CCDC 1520052) (b) **65** (CCDC 1520055) (*meso*-mesityl groups are omitted for clarity).



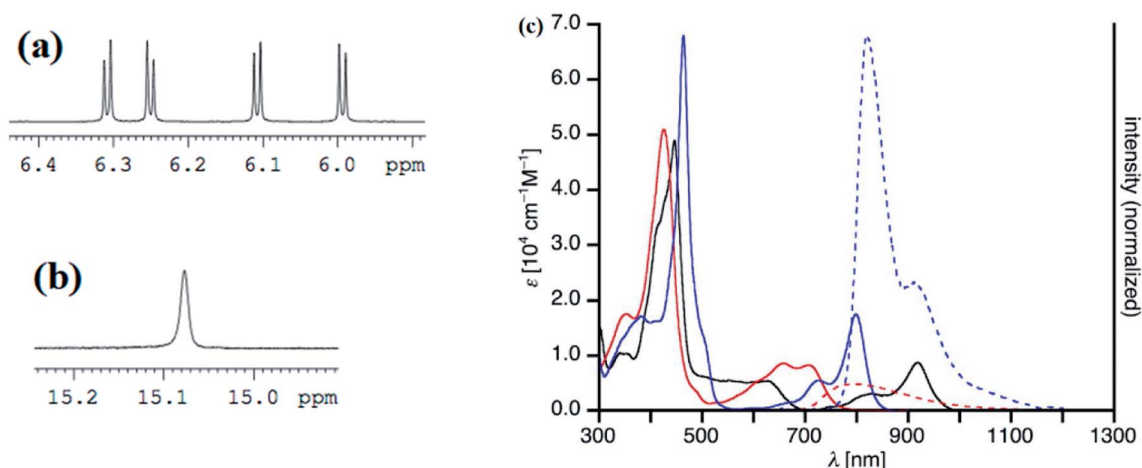
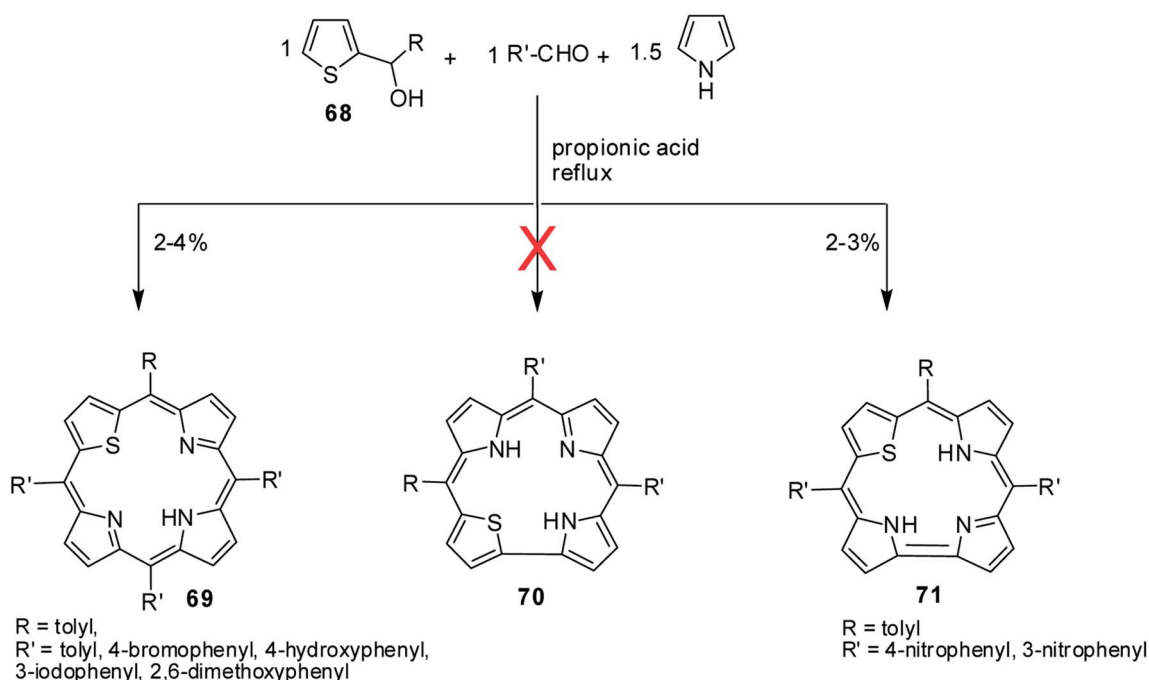


Fig. 15 Selected regions of ^1H NMR spectra of (a) **62** (b) **64** and (c) UV-vis/NIR absorption spectra (normal line) of **62** (black), **64** (red), **66** (blue) and emission spectra (dashed line) of **64** (red, $\lambda_{\text{max}} = 426$ nm), **66** (blue, $\lambda_{\text{max}} = 463$ nm) recorded in dichloromethane. Reproduced from ref. 24 with permission. Copyright© 2017 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.



Scheme 17 Synthesis of *meso*-substituted 22-thiacorrole **71**.

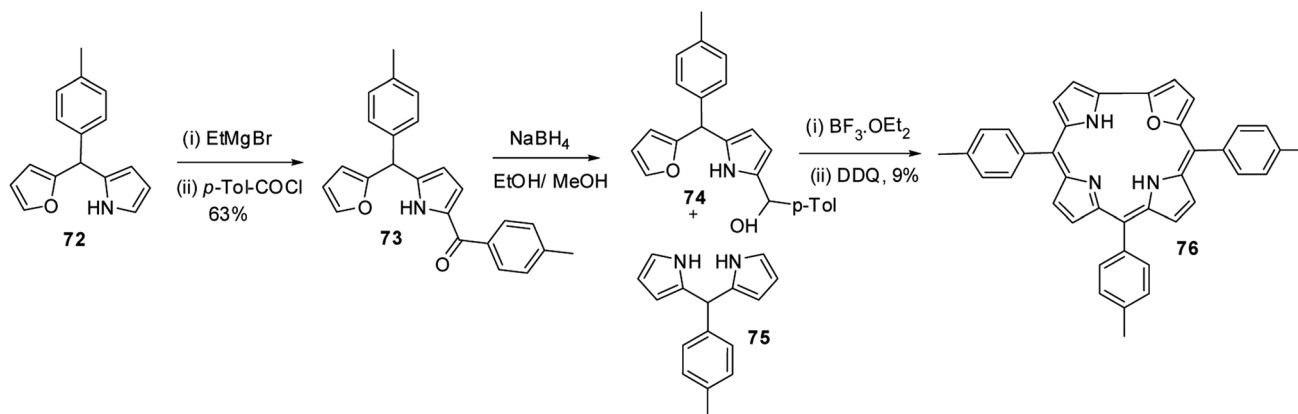
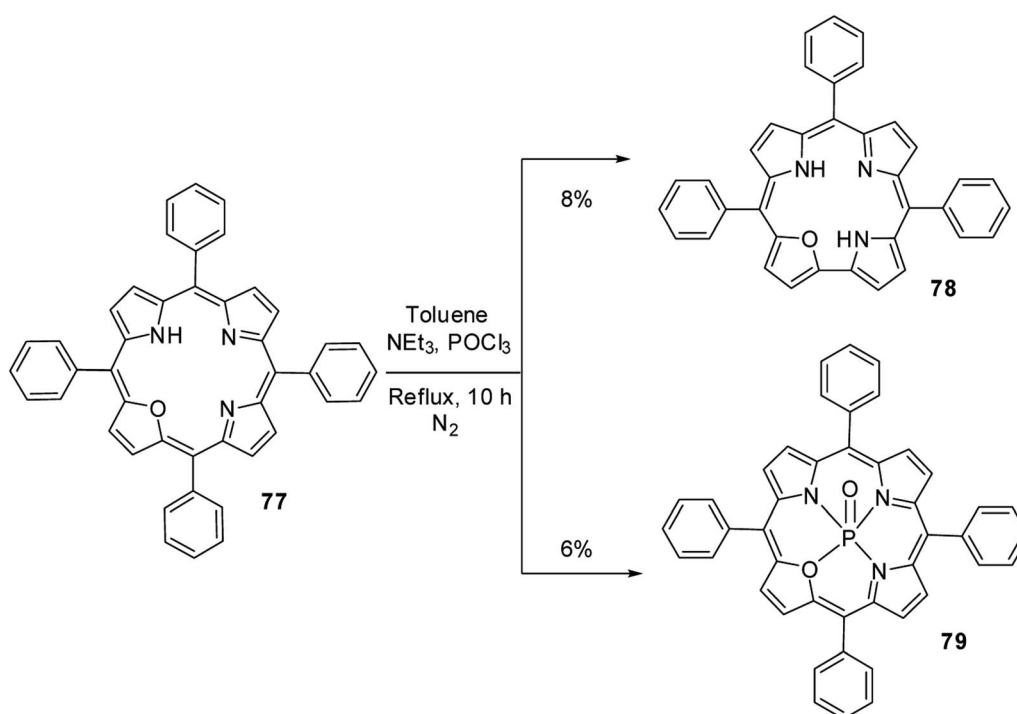
3. Corroles containing heteroatom(s) at their core (core modified corroles)

3.1. 22-Thiacorroles

A perusal of literature reveals that there is only one report available on 22-thiacorroles that has direct pyrrole-pyrrole bond.²⁵ The other possible isomer, 21-thiacorrole having direct pyrrole-thiophene link and dithiacorroles containing two core sulfur atoms in place of two pyrrole nitrogens are yet to be synthesized. Ravikanth and co-workers' attempts to make thiacorroles by condensing thiophene mono-carbinol **68** (1 eq.),

aromatic aldehyde (1 eq.) and pyrrole (1.5 eq.) in refluxing propionic acid predominantly resulted in the formation of 21-thiaporphyrin **69** (ref. 26) (Scheme 17). Although authors observed the formation of thiacorrole in few cases, their isolation was not possible due to the decomposition of corrole during column chromatography. However, authors were successful in isolating stable thiacorroles **71** in ~3% yield when 4-/3-nitrobenzaldehyde was used in the above condensation in otherwise similar conditions. Interestingly, no formation of 21-thiaporphyrin was noticed in this reaction. Authors found that changing the reaction conditions to mild Lindsey's protocol²⁷ did not result in formation of thiacorroles. The thiacorrole



Scheme 18 Synthesis of *meso*-substituted 21-oxacorrole **76**.Scheme 19 Synthesis of *meso*-substituted 21-oxacorrole **78** from 21-oxaporphyrin **77**.

formed in this condensation was expected to have direct α - α thiophene-pyrrole link **70** (21-thiacorroles) but the structural elucidation by detailed NMR studies revealed that the corrole was having direct α - α pyrrole-pyrrole link **71** (22-thiacorroles). Unfortunately, no crystal structure was available for 22-thiacorrole, but the combined NMR and Density Functional Theory (DFT) studies [B3LYP/6-31G(d) level] revealed their diminished aromatic character and non-planar conformation of 22-thiacorrole macrocycle.

The 22-thiacorroles **71** exhibited one strong Soret-like band at ~ 435 nm and one broad Q-band like transition at ~ 620 nm due to their less symmetric nature and were also weakly fluorescent with one broad ill-defined fluorescence band at ~ 660 nm. The electrochemical studies revealed that 22-

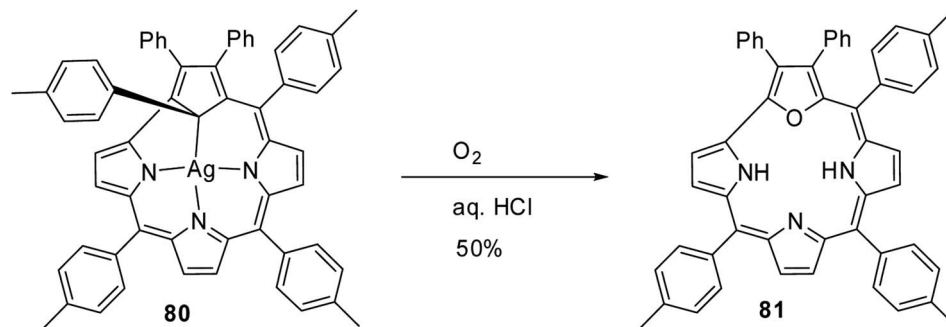
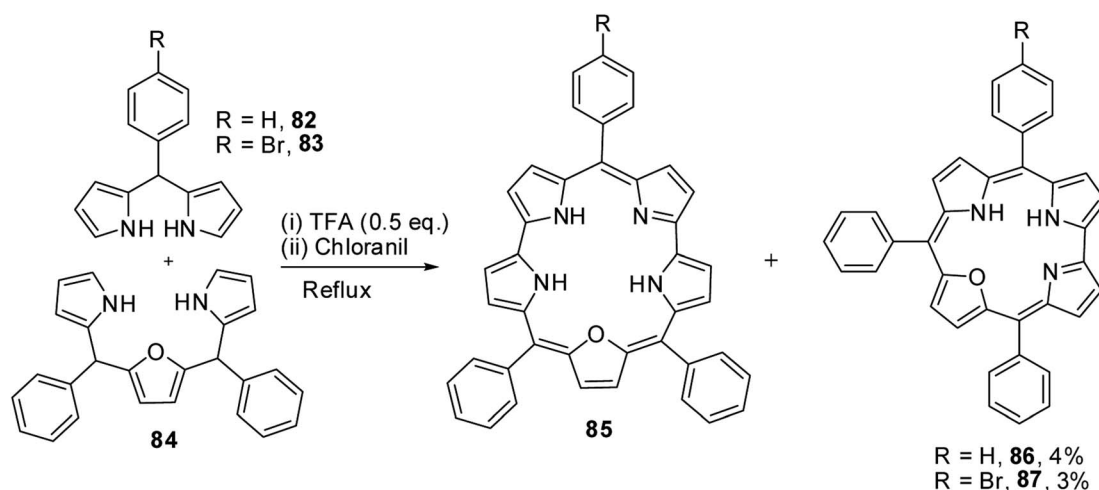
thiacorroles **71** were easier to reduce compared to 21-thiaporphyrins supporting their electron deficient nature.

3.2. 21-Oxacorroles

Lee and co-workers^{28,29} developed a rational method to synthesize the first examples of 21-oxacorroles containing direct furan-pyrrole bond **76** as shown in Scheme 18.

The *p*-tolyl-(furan-2-yl)-(pyrrol-2-yl)methane **72** was treated with EtMgBr in THF followed by *p*-toluoyl chloride resulted in an exclusive formation of pyrrole acylated product **73** in 63% yield. The pyrrole acylated compound **73** was then reduced to the corresponding alcohol **74** with excess of NaBH₄. The 21-oxacorrole **76** was synthesized in 9% yield by acid-catalyzed [2 +



Scheme 20 Synthesis of 21-oxacorrole **81** from Ag(III) carbacorrole **80**.Scheme 21 Synthesis of *meso*-substituted 22-oxacorroles **86** and **87**.

2] condensation of furyl-pyrrolyl mono-ol **74** with *meso*-aryl dipyrromethane **75** followed by the column chromatographic purification.

Ravikanth and co-workers³⁰ serendipitously noticed the formation of 21-oxacorrole **78** while preparing the phosphoryl complex of 21-oxaporphyrin **77**. The *meso*-tetraaryl 21-oxaporphyrin **77** was treated with excess $POCl_3$ in toluene in the presence of triethylamine at reflux followed by column chromatographic purification afforded *meso*-triaryl-21-oxacorrole **78** in 8% yield along with phosphoryl complex of *meso*-tetraaryl 21-oxaporphyrin **79** in 6% yield (Scheme 19). The structure of 21-oxacorrole **78** was also confirmed by X-ray crystallography. As per the mechanism proposed by authors, the *meso*-tetraaryl 21-oxaporphyrin **77** underwent ring contraction upon treatment with $POCl_3$ via dearylation of *meso*-aryl group present between furan and pyrrole moieties to form *meso*-triaryl-21-oxacorrole **78**.

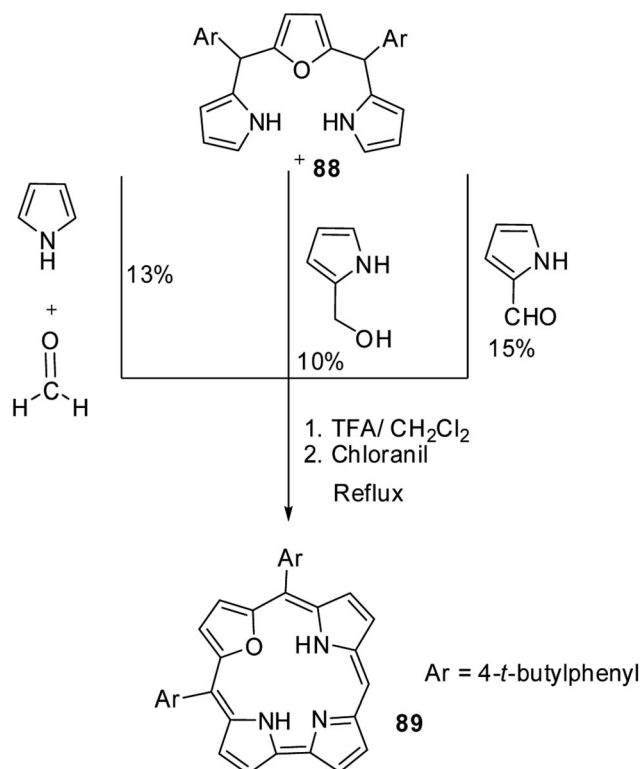
In an interesting report, Grażyński and co-workers noticed the formation of 21-oxacorrole **81**, when the silver(III) carbacorrole **80** was reacted with dioxygen in presence of aqueous HCl (Scheme 20).³¹

3.3. 22-Oxacorroles

Chandrashekar and co-workers³² observed the formation of 22-oxacorrole containing direct pyrrole–pyrrole bond **86**, **87** as a byproduct during their synthesis of expanded porphyrin, 25-oxasmaragdyrin **85**. The TFA-catalyzed oxidative coupling between 16-oxatripyrrane **84** and *meso*-aryl dipyrromethanes **82**, **83** followed by oxidation with chloranil resulted in the formation of 25-oxasmaragdyrin **85** as a major product (~50% yield) and 22-oxacorroles **86**, **87** in 3–4% yield (Scheme 21). Although this method yielded 21-oxacorroles in low yields, it was later proved to be an easy and versatile method to make *meso*-substituted 22-oxacorroles.

Chandrashekar and co-workers³³ subsequently developed three different [3 + 1] synthetic routes to prepare 22-oxacorroles containing one *meso*-free carbon **89** as shown in Scheme 22. The acid catalyzed condensation of *meso*-substituted 16-oxatripyrrane **88** was done in three ways: in the first method, the 16-oxatripyrrane **88** was condensed with pyrrole and para-formaldehyde; in second method, **88** was condensed with 2-(hydroxymethyl)pyrrole and in the third method, **88** was condensed with pyrrole-2-carboxaldehyde. In all three methods, the condensation was followed by a chloranil oxidation resulting in mono *meso*-free 22-oxacorrole **89** in 10–15% yields.



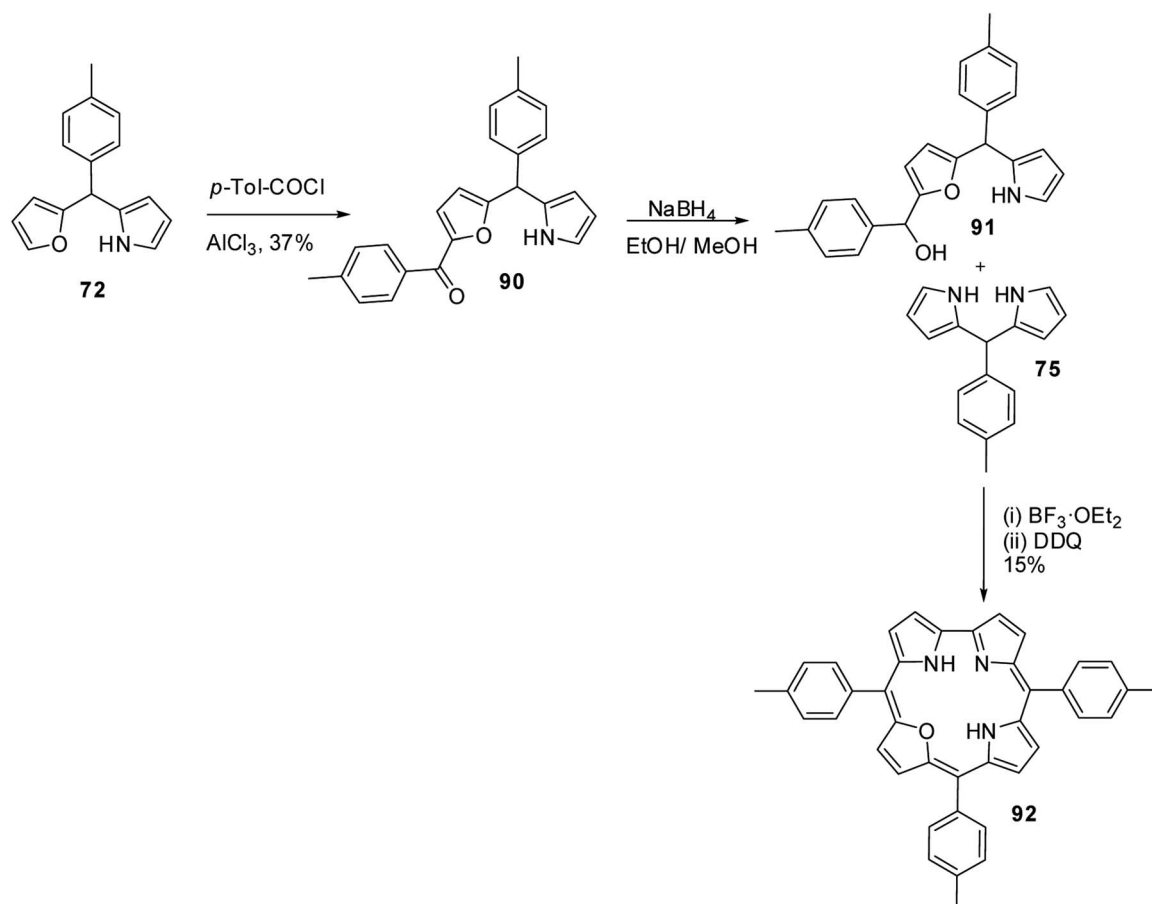


Scheme 22 Three different synthetic routes to prepare *meso*-substituted 22-oxacorrole **89**.

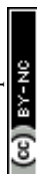
Lee and co-workers also successfully synthesized 22-oxacorroles by [2 + 2] condensation strategy as shown in Scheme 23.^{28,29} The AlCl_3 catalysed acylation of *p*-tolyl-(furan-2-yl)-(pyrrol-2-yl)methane **72** followed by NaBH_4 reduction yielded corresponding alcohol **91**, which upon Lewis acid catalyzed condensation with *meso*-aryl dipyrromethane **75** followed by column chromatographic purification afforded 22-oxacorrole **92** in 15% yield.

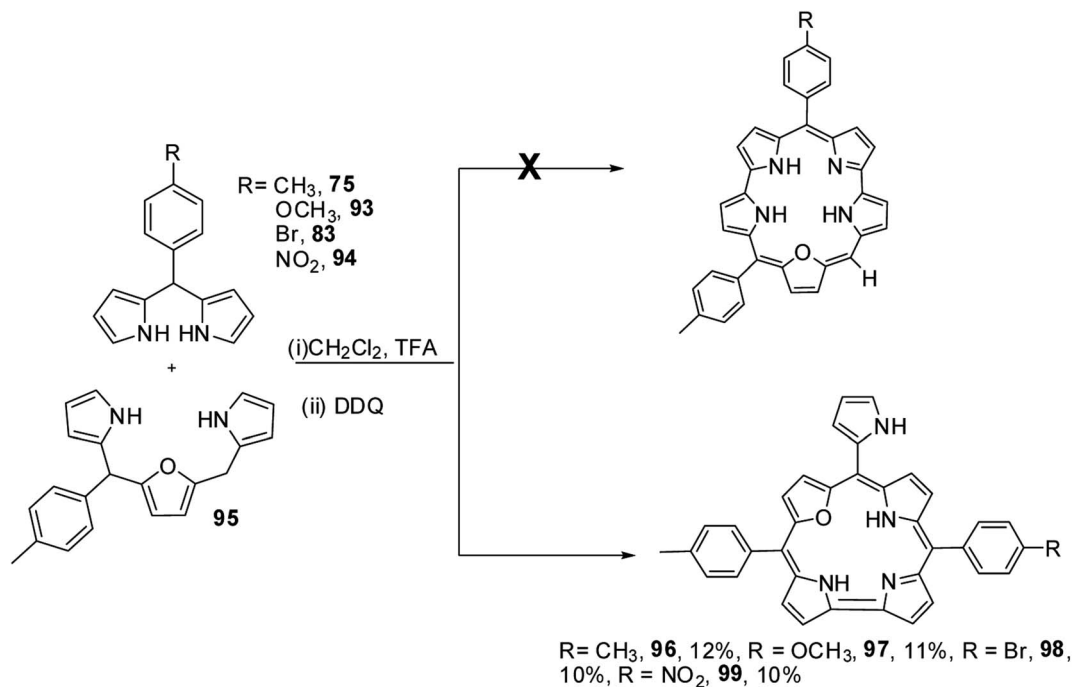
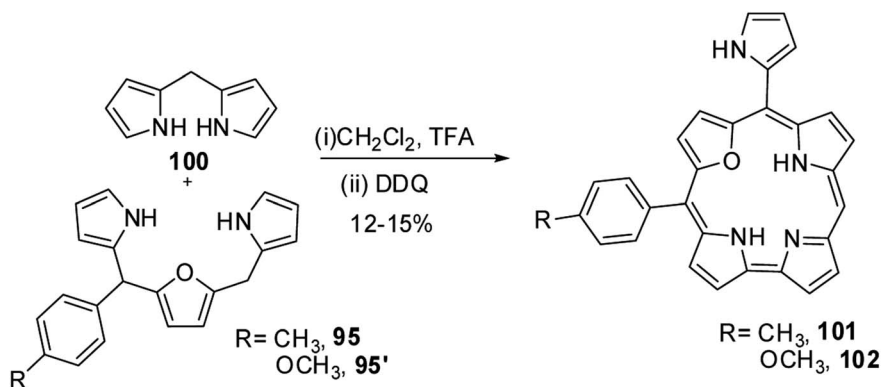
Ravikanth and co-workers adopted the same [3 + 2] synthetic strategy to prepare mono *meso*-pyrrolyl 22-oxacorroles **96–99**.³⁴ The 22-oxacorroles containing two *meso*-aryl and one *meso*-pyrrolyl groups **96–99** were synthesized by condensing 16-oxatripyrrane **95** with *meso*-aryl substituted dipyrromethanes **75**, **83**, **93** and **94** under mild acid catalyzed conditions followed by DDQ oxidation (Scheme 24). Kalita and Ravikanth further extended this synthetic strategy to prepare *meso*-free mono-*meso*-pyrrolyl 22-oxacorroles **101**, **102** as shown in Scheme 25.³⁵

Chandrashekar and co-workers³⁶ obtained *meso*-ferrocenyl 22-oxacorroles **113–117** as minor products (~3% yield) along with *meso*-ferrocenyl 25-oxasmaragdyrins **108–112** (~20% yield) by condensing appropriate 16-oxatripyrranes **88**, **104–107** with *meso*-ferrocenyl dipyrromethane **103** under TFA-catalyzed conditions (Scheme 26).



Scheme 23 Synthesis of *meso*-substituted 22-oxacorroles **92**.



Scheme 24 Synthesis of *meso*-pyrrolyl 22-oxacorrole **96–99**.Scheme 25 Synthesis of mono *meso*-free 22-oxacorrole **101** and **102**.

3.4. X-ray structures of 21-oxacorrole and 22-oxacorrole

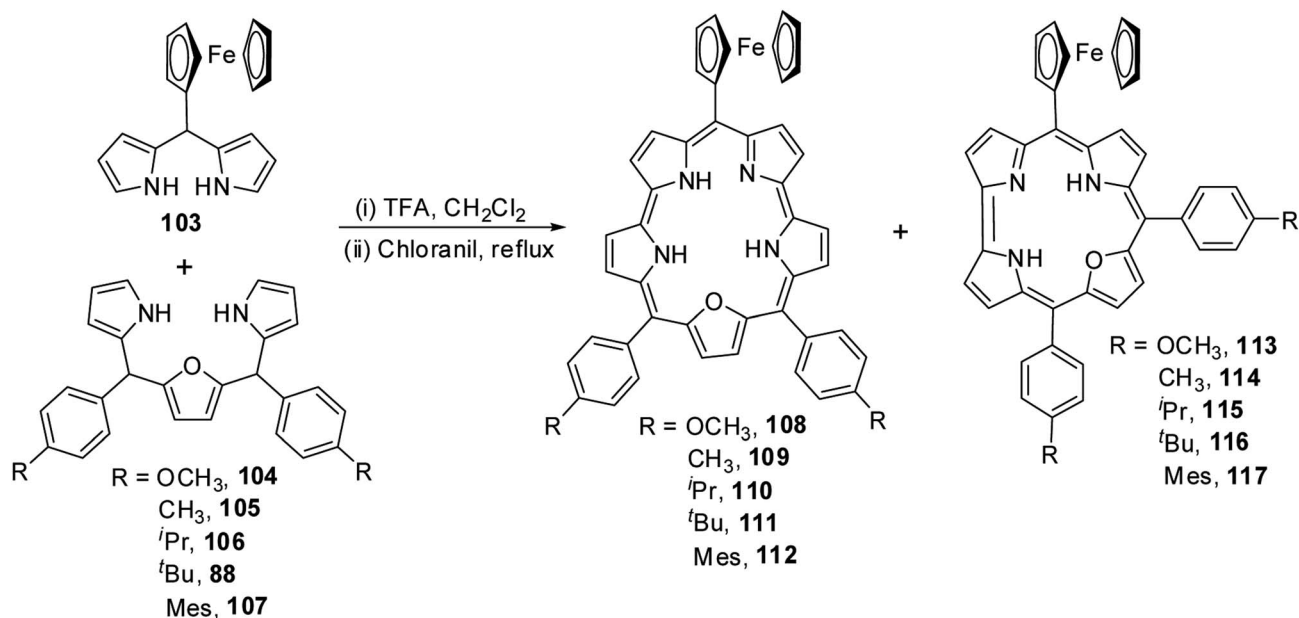
Gratifyingly, the crystal structures for both *meso*-triaryl 21-oxacorrole **78** and *meso*-triaryl 22-oxacorrole **86** were available by two different research groups and are presented in Fig. 16.^{30,32} Compared to *meso*-tetraaryl 21-oxaporphyrin, the 21-oxacorrole and 22-oxacorroles **78** and **86** adopted a near-planar conformation owing to the presence of direct furan-pyrrole bond and direct pyrrole-pyrrole bond respectively. The deviation of the furan ring from the mean plane of the macrocycle was much less in 21-oxacorrole **78** (5.69°) and 22-oxacorrole **86** (5.50°) compared to 21-oxaporphyrin (11.35°). Similarly, the deviation of three pyrrole rings from the mean plane of the macrocycle was more pronounced in 21-oxaporphyrin compared to 21-oxacorrole and 22-oxacorrole. Furthermore, the direct distance between the pyrrole and furan ring in 21-oxacorrole (1.40 Å) was little shorter

than the direct pyrrole-pyrrole distance in 22-oxacorrole (1.43 Å). The *meso*-phenyl groups were more coplanar with the macrocycle in 21-oxacorrole and 22-oxacorrole compared to 21-oxaporphyrin.

3.5. Spectral and electrochemical properties of 21-oxacorroles and 22-oxacorroles

The comparison of Soret and Q-band absorption spectra, fluorescence spectra and cyclic voltammograms of 21-oxacorrole **78** and 22-oxacorrole **86** are presented in Fig. 17. The 21-oxacorrole displayed Q-type bands in the region 480–610 nm and a split Soret band in 400–420 nm, whereas the 22-oxacorroles showed four well defined Q-type bands in the region 490–650 nm and a strong Soret band at 411 nm. The blue shifted absorption bands of 21-oxacorrole compared to 22-oxacorrole indicate that the 21-oxacorrole is less flexible and more resonance stabilized





Scheme 26 Synthesis of mono *meso*-ferrocenyl 22-oxacorroles 113–117.

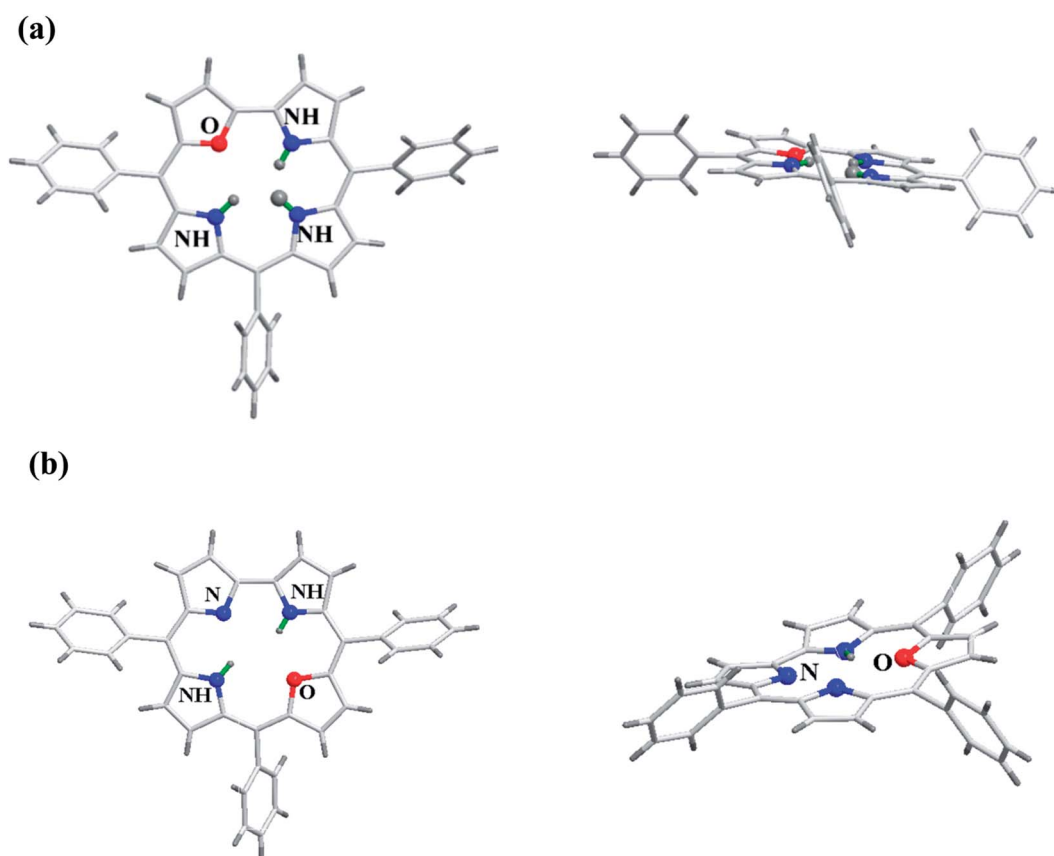


Fig. 16 Top and side view of X-ray crystal structures of (a) *meso*-substituted 21-oxacorrole **78** (CCDC 913385) (b) *meso*-substituted 22-oxacorrole **86** (CCDC 112074).

than 22-oxacorrole. The nature and presence of *meso*-substituent on the absorption spectral pattern was also evident as *meso*-pyrrolyl 22-oxacorroles and *meso*-ferrocenyl 22-oxacorroles

showed slight bathochromic shifts in their peak maxima whereas the *meso*-free 22-oxacorroles showed slight hypsochromic shifts compared to *meso*-triaryl 22-oxacorroles.



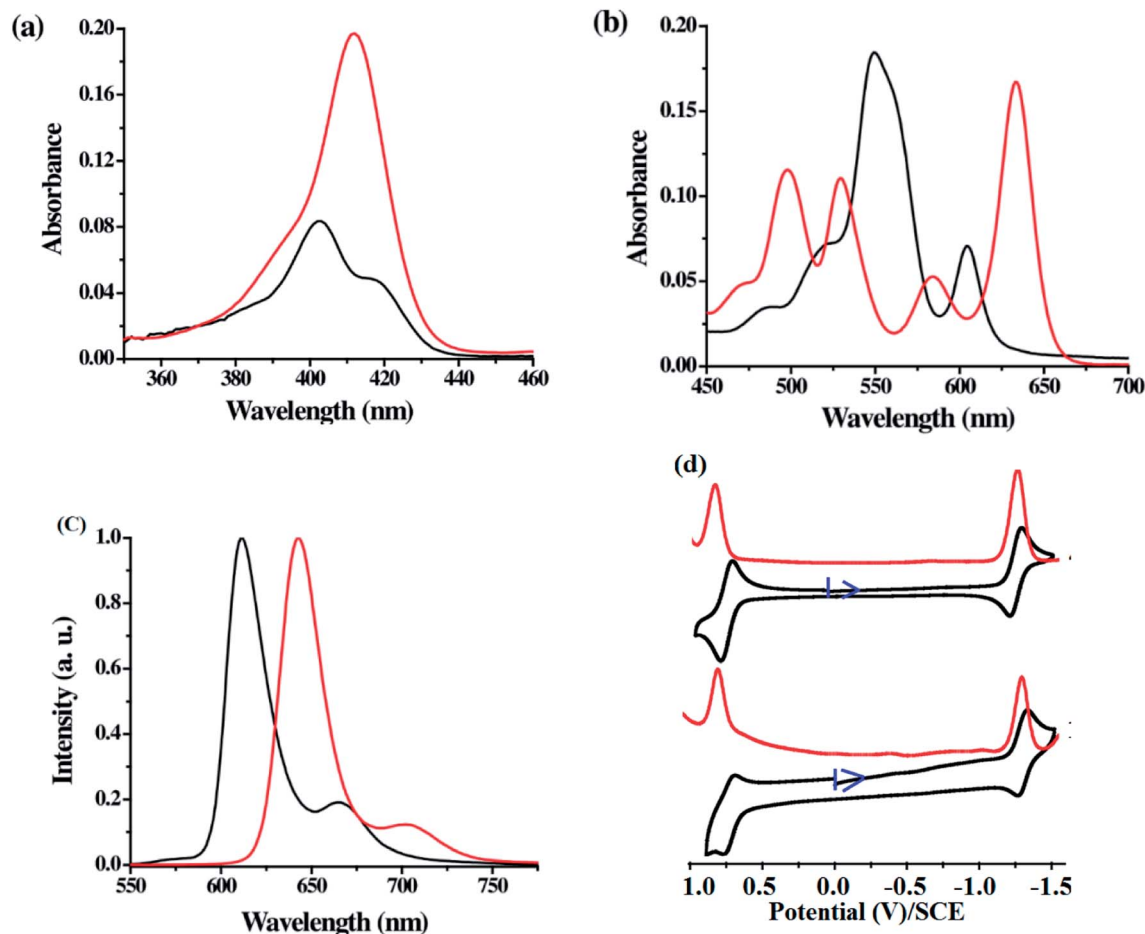


Fig. 17 (a) Soret band (b) Q-bands of *meso*-substituted 21-oxacorrole **78** (black), 22-oxacorrole **86** (red), (c) emission profiles of 21-oxacorrole **78** (black, $\lambda_{\text{ex}} = 402$ nm), 22-oxacorrole **86** (red, $\lambda_{\text{ex}} = 412$ nm) recorded in dichloromethane (d) cyclic voltammogram (black) and DPV plots (red) of 22-oxacorrole **78** (top) and 21-oxacorrole **86** (bottom) recorded in dichloromethane with TBAP as supporting electrolyte. Reproduced from ref. 30 with permission. Copyright© 2013 American Chemical Society.

Both 21-oxacorrole **78** and 22-oxacorrole **86** were brightly fluorescent with emission maxima in 610–650 nm range and the quantum yields were in the range of 0.3 to 0.5. In general, the oxacorroles exhibit one quasi-reversible and one irreversible peaks for both oxidation and reduction processes. Although both 21-oxacorrole **78** and 22-oxacorrole **86** were stable under redox conditions and exhibited similar redox behaviour, their oxidation/reduction was found to be difficult compared to 21-oxaporphyrins. The presence of different *meso*-substituents such as pyrrole and ferrocenyl groups slightly altered their redox behaviour. For example, the *meso*-pyrrolyl 22-oxacorroles were easier to reduce compared to *meso*-triaryl 22-oxacorroles.

3.6. Metal complexes of 22-oxacorroles

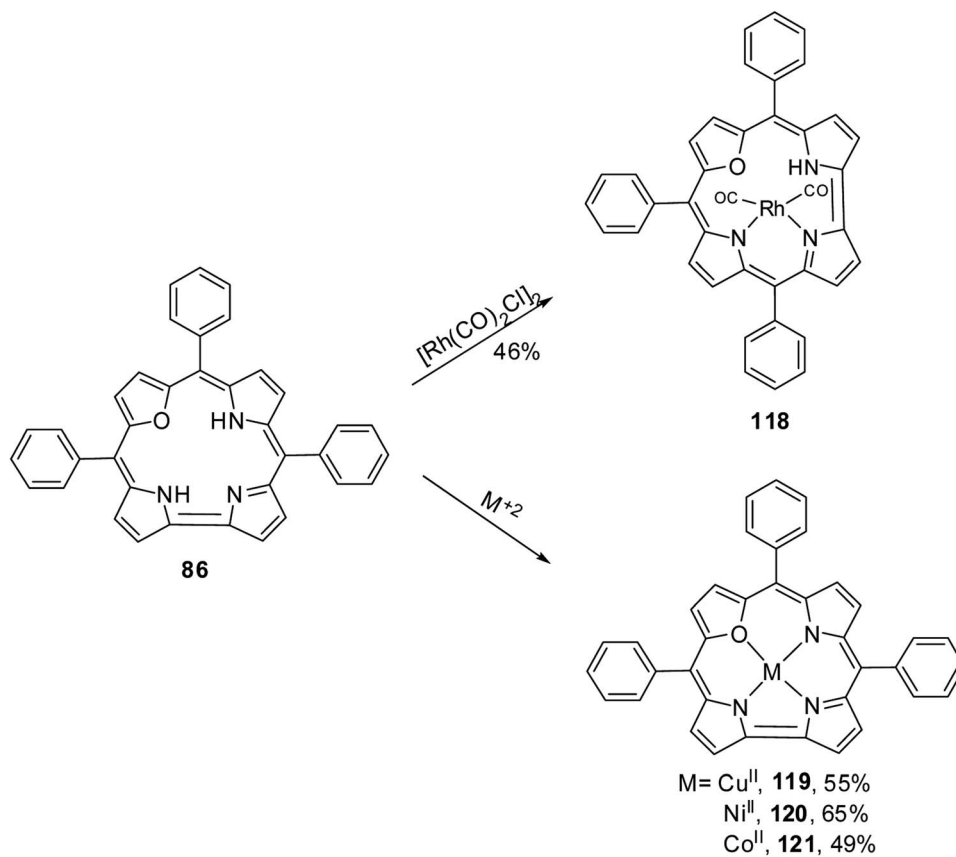
The 22-oxacorroles with two ionizable protons are expected to form complexes readily with divalent metals unlike normal corroles which are known to stabilise metals in higher oxidation states due to three ionizable protons. Chandrashekar and co-workers³⁷ reported one monovalent Rh(I) complex **118** and three divalent metal complexes such as Cu(II) **119**, Ni(II) **120** and Co(II) **121** by treating the 22-oxacorrole **86** with appropriate metal

precursors under standard metallation reaction conditions (Scheme 27). The X-ray structures of Rh(I) **118** and Ni(II) **120** complexes are presented in Fig. 18. The crystal structure of Rh(I) complex **118** revealed that Rh(I) was located above the macrocyclic plane in a near square planar geometry coordinated by one imino and one amino nitrogen of the 22-oxacorrole macrocycle and remaining two coordination sites were occupied by carbonyl ligand (Fig. 18). Since Rh(I) was situated out of plane, the pyrrole rings of dipyrromethene moiety twisted above the mean corrole plane by an angle of 23.51° and 19.69° respectively. The angle between the plane containing the Rh(I) with its coordinated atoms and the mean corrole plane was 62.64°.

The Ni(II) insertion into the 22-oxacorrole core flattens the macrocycle to almost planar structure and Ni(II) was coordinated in distorted square planar conformation. The Ni(II) ion lies above the mean plane of the macrocycle by only 0.008 Å. The Ni–N and Ni–O distances were shorter compared to the corresponding distances in Ni(II) derivative of 21-oxaporphyrin due to the reduced core size of 22-oxacorrole.

The Cu(II) **119** and Co(II) **121** complexes of 22-oxacorrole were paramagnetic and EPR spectra of these complexes displayed





Scheme 27 Synthesis of metal derivatives of 22-oxacorrole **118**–**121**.

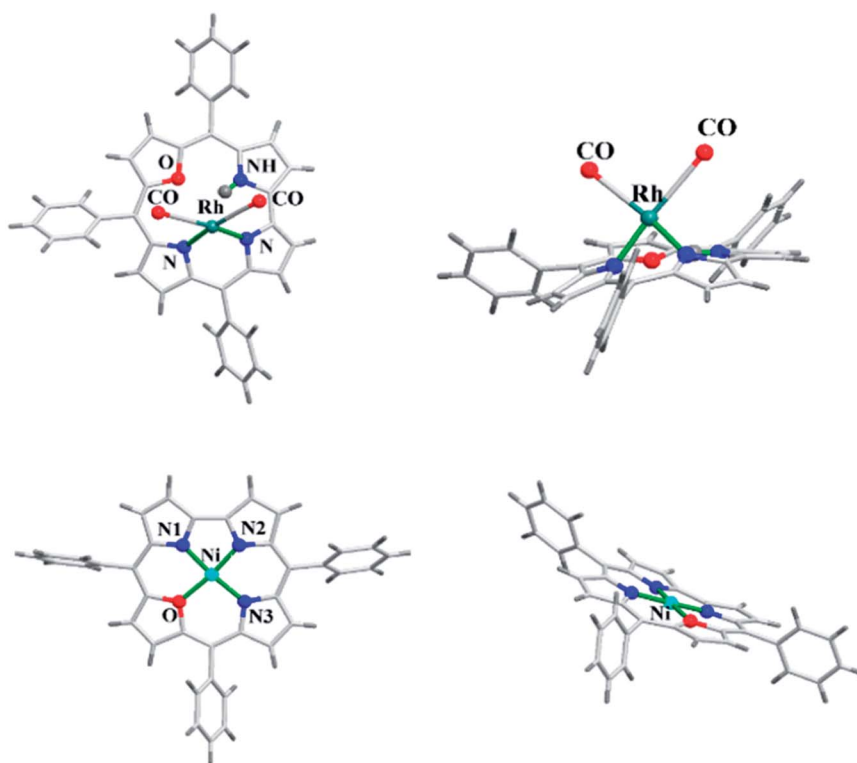
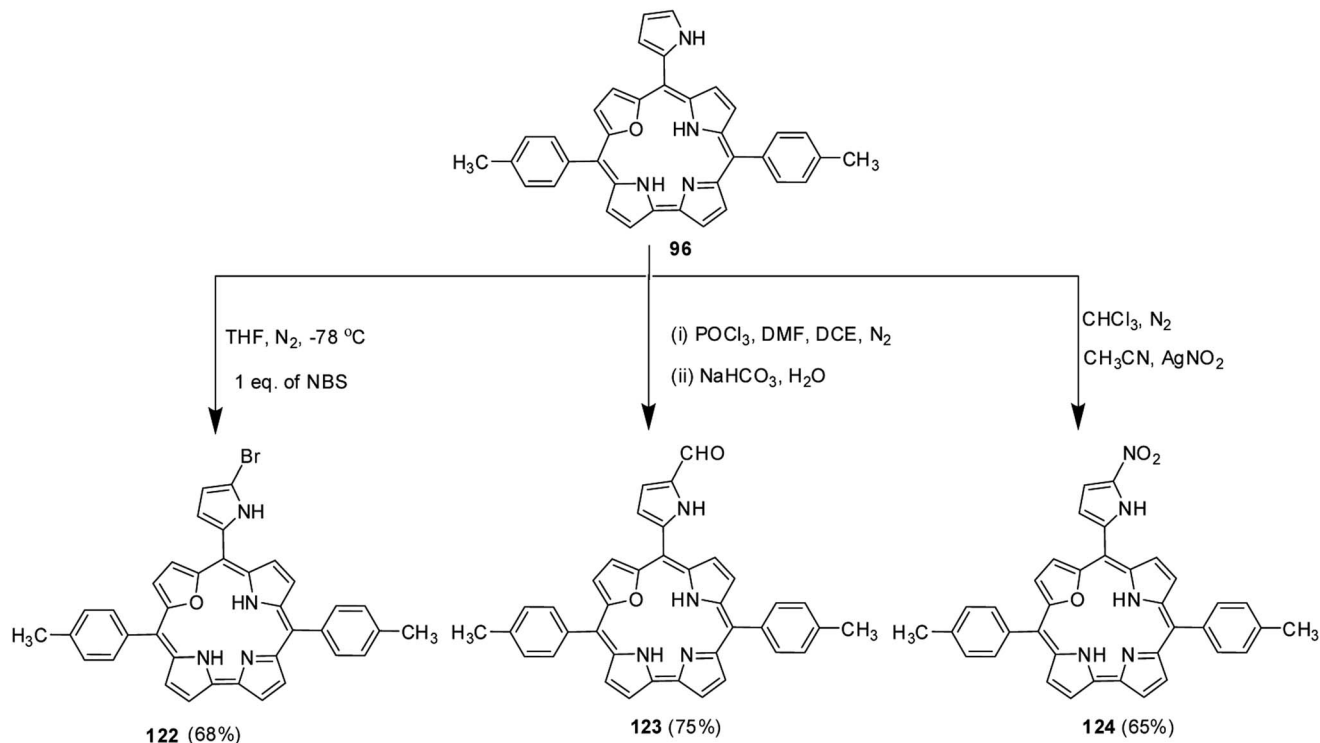


Fig. 18 Top and side view of X-ray crystal structures of Rh(I) **118** (CCDC 135916) (top) and Ni(II) **120** (CCDC 135917) (bottom) complexes of *meso*-substituted 22-oxacorrole.





Scheme 28 Functionalization reactions of *meso*-pyrrolyl 22-oxacorrole **96**.

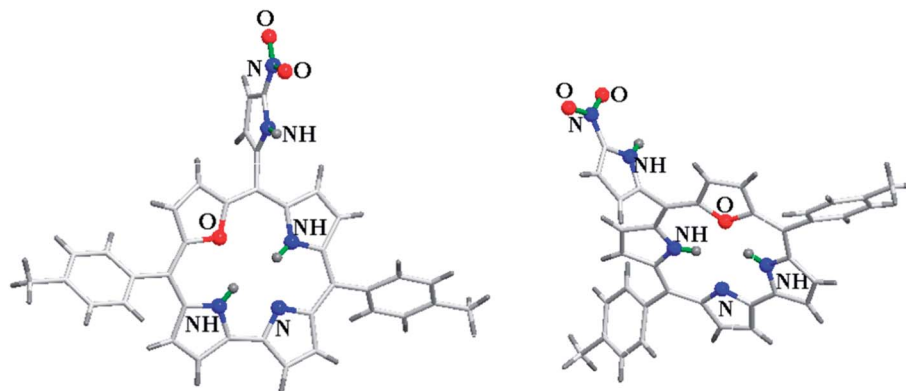


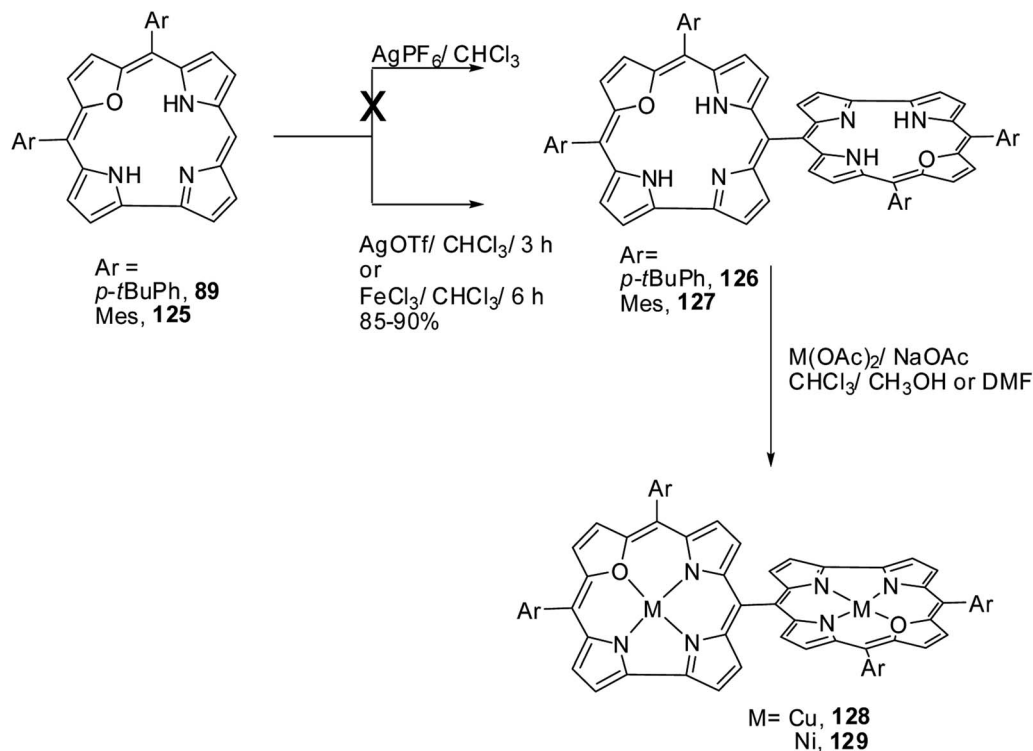
Fig. 19 Top and side view of X-ray crystal structures of **124** (CCDC 973515).

significant decrease in the metal hyperfine couplings compared to the corresponding porphyrin complexes. The absorption spectra of the metallated derivatives of 22-oxacorrole showed characteristic Soret-type and Q-type bands in the region 400–700 nm like their free base counterparts. The Cu(II) and Co(II) complexes showed broad Soret type band whereas the Rh(I) and Ni(II) showed split Soret band due to their lower symmetry in solution. The extinction coefficient values of all the metallated derivatives were smaller than the free base 22-oxacorrole due to decreased π -electron conjugation in metal derivatives. The electrochemical studies of these metal complexes revealed that the redox process was only macrocycle centered.

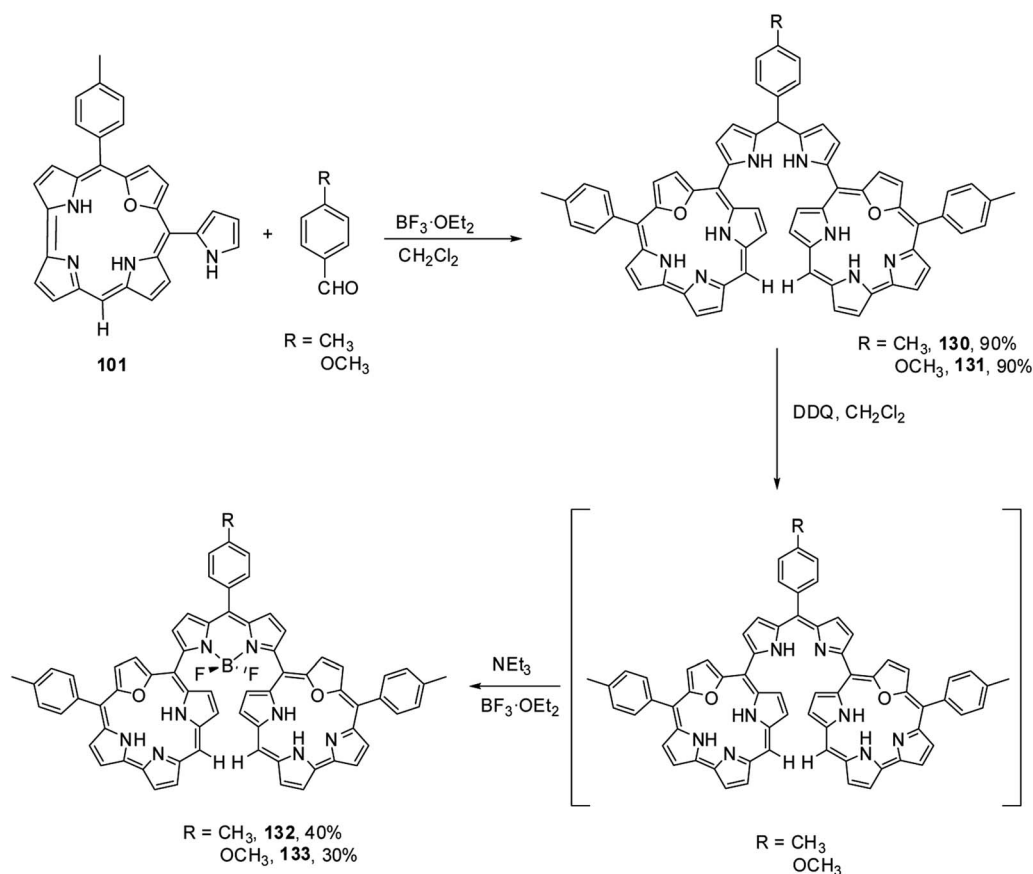
3.7. Functionalized 22-oxacorroles and covalently linked 22-oxacorrole dyads

Ravikanth and co-workers explored the functionalization reactions of *meso*-pyrrole substituted 22-oxacorroles by taking advantage of the reactivity of the α -position of pyrrole ring present at the *meso*-position.³⁴ Bromination, formylation and nitration reactions were carried out on *meso*-pyrrolyl substituted 22-oxacorrole **96** by using N-bromosuccinimide (NBS), Vilsmeier reagent and AgNO₂ respectively, as shown in Scheme 28. The crystal structure of *meso*-pyrrole nitrated 22-oxacorrole **124** showed that (Fig. 19) the corrole macrocycle was almost planar with slight deviations ranging from 2.98 to 11.52° with respect to the mean plane defined by three *meso*-carbon atoms, three





Scheme 29 Synthesis of meso-meso linked oxacorrrole dimer 126, 127 and their bis metal derivatives 128, 129.



Scheme 30 Synthesis of BODIPY bridged 22-oxacorrrole dyad 132, 133.



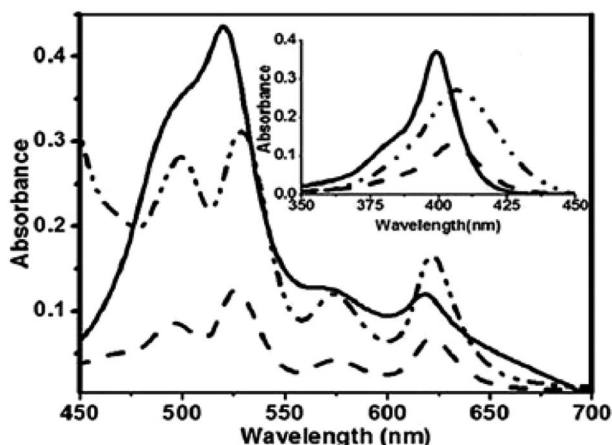
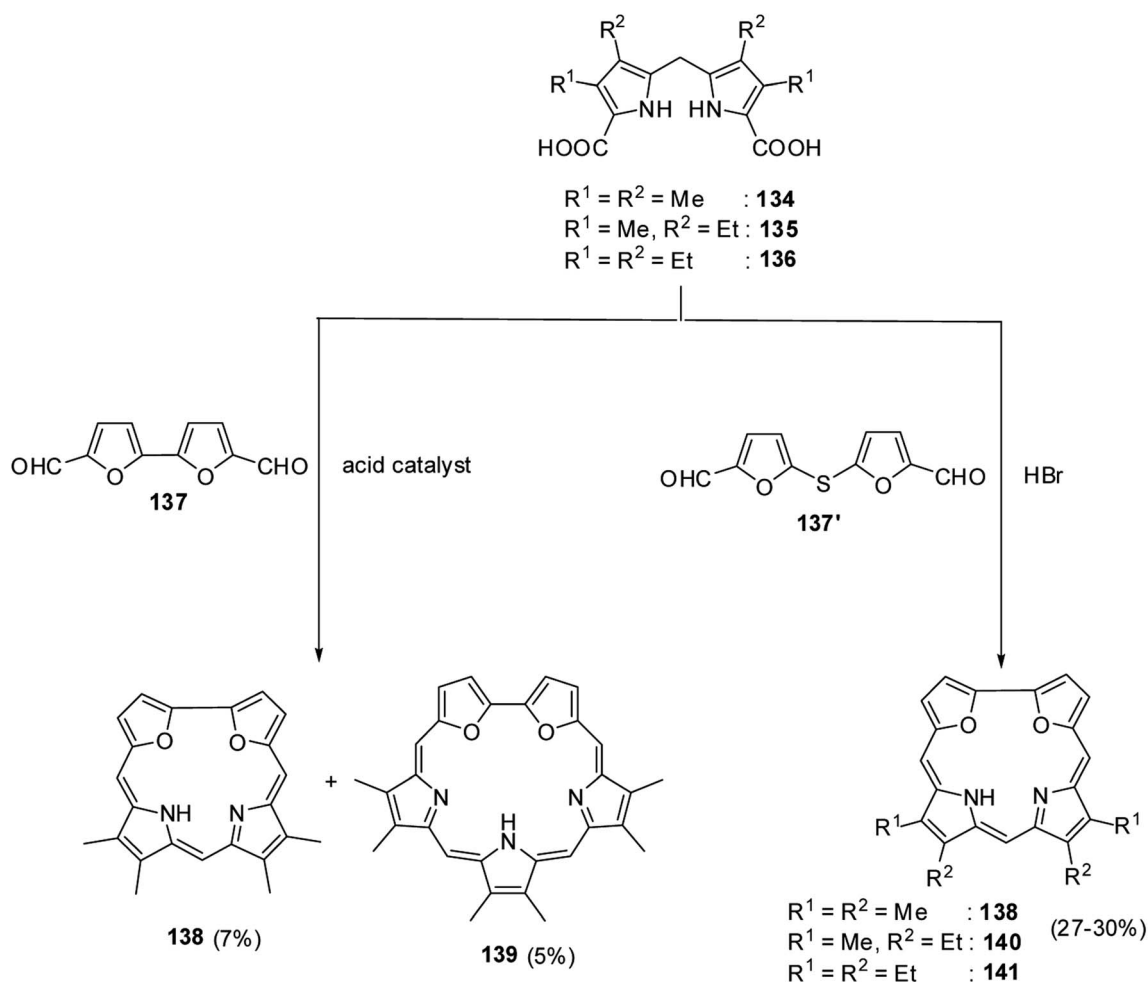


Fig. 20 Absorption spectra of BODIPY bridged corrole dyad **132** (normal line), corrole dyad (dotted line) **130** and monomer oxacorrole **101** (dashed line) recorded in dichloromethane. Reproduced from ref. 35 with permission. Copyright© 2015 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

pyrrole rings and one furan ring. The nitro group and *meso*-pyrrole groups were perpendicular to the plane of the macrocycle.

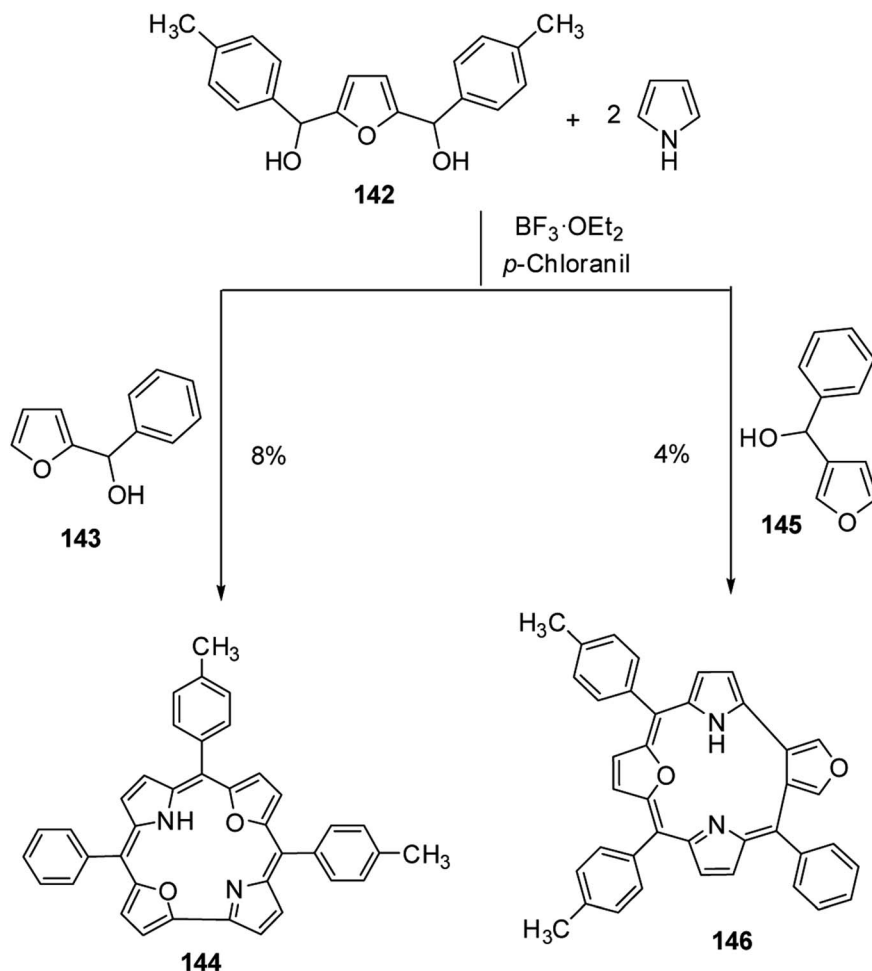
Chandrashekar and co-workers³⁸ successfully prepared the first examples of *meso-meso* linked 22-oxacorrole dimers **126**, **127** in near quantitative yield by AgOTf or FeCl₃ catalyzed coupling reaction of *meso*-free 22-oxacorroles **89** and **125** as shown in Scheme 29. The absence of *meso*-CH resonance signal in ¹H NMR spectrum provided a strong evidence for the formation of corrole dimer. The copper and nickel derivatives of *meso-meso* linked 22-oxacorrole dimers **128**, **129** respectively, were obtained by metallating dimers with appropriate metal salts (Scheme 29) However, authors did not succeed in obtaining hetero bis-metal derivatives of 22-oxacorrole dimer.

The absorption spectral studies revealed weak interactions between the two sub-units in dimers **126**, **127** and the exciton coupling observed for the free base 22-oxacorrole dimer was lower than the corresponding protonated derivatives. The NMR and DFT (B3LYP-631G-level) studies indicated a noncoplanar arrangement of two corrole units in dimers. The electron paramagnetic resonance and magnetic studies on bis-Cu(II) complex of 22-oxacorrole dimer **128** indicated that both copper ions behave as independent spins without any noticeable interaction. The fluorescence maxima of 22-oxacorrole dimers experienced substantial bathochromic shift (60 nm) compared to monomeric 22-oxacorrole. The authors also measured



Scheme 31 Synthesis of *meso*-free 21,24-dioxacorroles **138**–**141**.





Scheme 32 Synthesis of *meso*-substituted 21,23-dioxacorrole **144** and its isomer **146**.

hyperpolarizability (β) by hyper Rayleigh scattering (HRS) method which revealed that the β -values were doubled in 22-oxacorrole dimers compared to the corresponding 22-oxacorrole monomers due to enhanced π -conjugation in the dimers. The bis-copper complex was explored for the photocleavage of DNA and the studies revealed that the bis-copper complex selectively cleaves the nucleic acids without affecting the proteins. This observation suggests possible application of this bis copper complex in the removal of nucleic acid contamination from protein extracts through a simple photolytic pathway.

Kalita and Ravikanth³⁵ reported the synthesis of novel boron-dipyrromethene (BODIPY)-bridged 22-oxacorrole dyads **132**, **133** by using *meso*-pyrrolyl 22-oxacorrole **101** as a key synthon as shown in Scheme 30. The BODIPY-bridged corrole dyads **132**, **133** showed an absorption band at 520 nm due to the BODIPY moiety and three bands at 399, 572 and 618 nm due to corrole units (Fig. 20). The absorption bands of BODIPY-bridged 22-oxacorrole dyads were slightly blueshifted with an increase in their extinction coefficients compared to monomeric *meso*-pyrrolyl 22-oxacorrole. The BODIPY-bridged 22-oxacorrole dyads **132**, **133** were weakly fluorescent and showed one broad emission band at 640 nm with a quantum yield of 0.03–0.05.

3.8. Dioxacorroles

Broadhurst, Grigg and Johnson³⁹ reported the synthesis of *meso*-free 21,24-dioxacorroles **138**, **140** and **141** by utilizing the acid catalyzed, MacDonald type condensation of β -substituted dipyrrolylmethane diacids **134**–**136** with either bifuran dialdehyde **137** or diformyl-difuryl sulphide **137'** (Scheme 31). The condensation of β -substituted dipyrrolylmethane diacids **134**–**136** with bifuran dialdehyde **137** yielded dioxacorrole **138** along with an expanded porphyrin, heterosapphyrin **139** which were eventually separated by column chromatography. Alternatively, the acid-catalyzed condensation of diformyl-difuryl sulphide **137'** with dipyrrolylmethane diacids **134**–**136** yielded dioxacorroles **138**, **140** and **141** in 27–30% yields. Authors' attempts to make metal derivatives of 21,24-dioxacorroles were unsuccessful.

Latos-Grażyński and co-workers⁴⁰ reported the synthesis of 5-phenyl-10,15-bis(*p*-tolyl)-21,23-dioxacorrole **144** by condensing furan diol **142** with 2-phenylhydroxymethylfuran **143** in a synthetic route depicted in Scheme 32. The aromatic character of dioxacorrole **144** was evident from its NMR spectroscopic features such as downfield and upfield shifted resonance signals for β -CH of furan and core NH protons respectively.



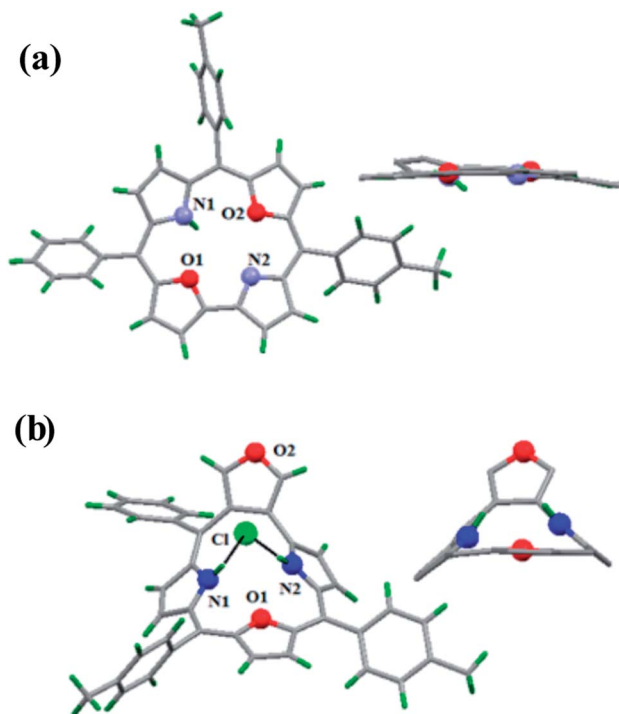
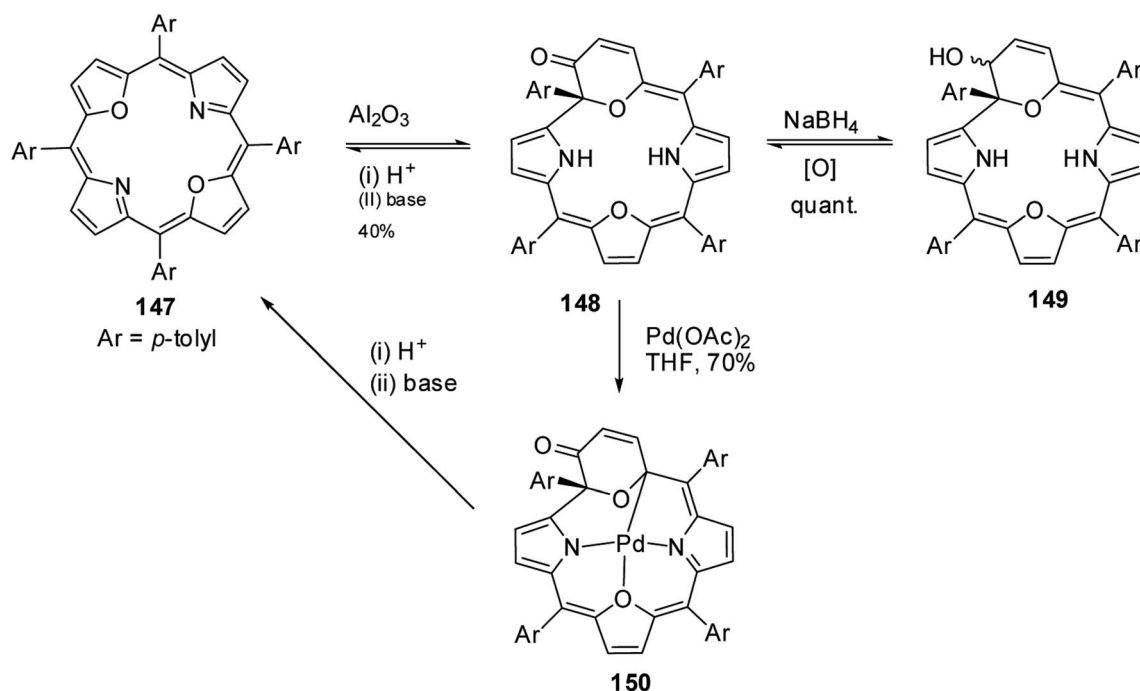


Fig. 21 Top and side view of X-ray crystal structures of (a) **144** (b) **146H⁺**.

The core NH protons exhibited rapid tautomerism by exchanging sites between two structurally inequivalent nitrogen atoms even at 188 K. Protonation by TFA arrested the tautomerism and resonance signals for inner core NH protons were observed at -1.46 and -1.98 ppm. Also, dioxacorrole **144** exhibited acid dependent chemical shift of NH resonance

emphasizing the role of counter-anion's interaction with monocation of dioxacorrole. The single crystal X-ray structure of dioxacorrole **144** is shown in Fig. 21. The furan moieties displayed longer and shorter bond lengths for $C_{\alpha}-C_{\beta}$ and $C_{\beta}-C_{\beta}$ respectively when compared to free furan indicating an alteration in π delocalization of furan rings in dioxacorrole. The dioxacorrole **144** exhibited markedly split Soret bands in 395–425 nm region reflecting reduced symmetry due to presence of two oxygen atoms in *trans* position along with series of Q-bands in 480–640 nm range.

Further, when 3-phenylhydroxymethylfuran **145** was condensed with **142** and pyrrole under identical reaction conditions, the formation of an unusual isomer of dioxacorrole containing protruding furan ring **146** was formed (Scheme 32). Relatively upfield shifted β -H resonances with respect to **142** and markedly down field shifted resonance of NH (17.71 ppm) convincingly demonstrated non aromatic character of **146**. The X-ray crystal structure of protonated corrole **146H⁺** showed a puckered structure arising due to the contraction of internal ring by one carbon atom compared to regular corrole core (Fig. 21). The protonated species **146H⁺** behaved as an anion receptor by exhibiting $N-H\cdots Cl$ hydrogen bond by two NH groups. The crystallographic data also revealed that the protruding furan ring preserved all the features typical of isolated furan whereas the second furan moiety in the macrocycle underwent perturbation in π -delocalization. The absorption spectroscopic features of **146** and its protonated form **146H⁺** were different from **144** and **144H⁺**. The electronic absorption spectrum of **146** exhibited a split Soret band at 375 and 409 nm along with a broad band at 753 nm which was remarkably red shifted to 962 nm in compound **146H⁺**.



Scheme 33 Synthesis of 3-pyranone dioxacorrole **148** and its Pd(II) derivative **150**.



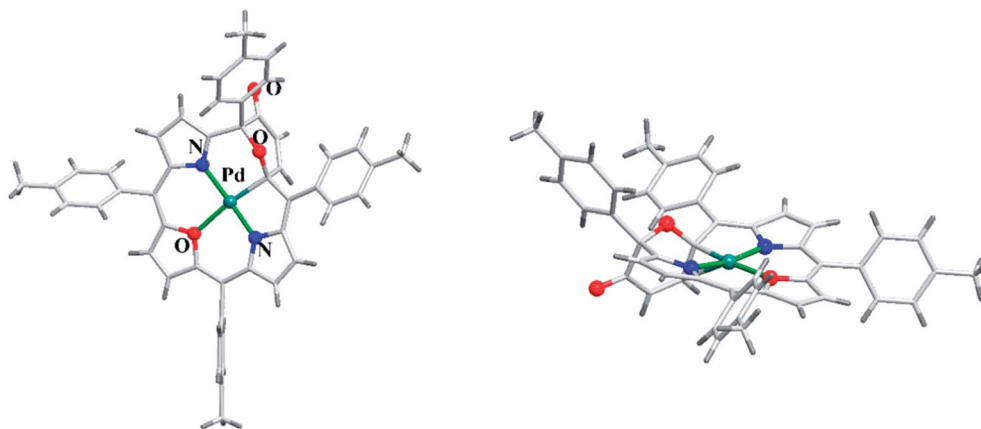
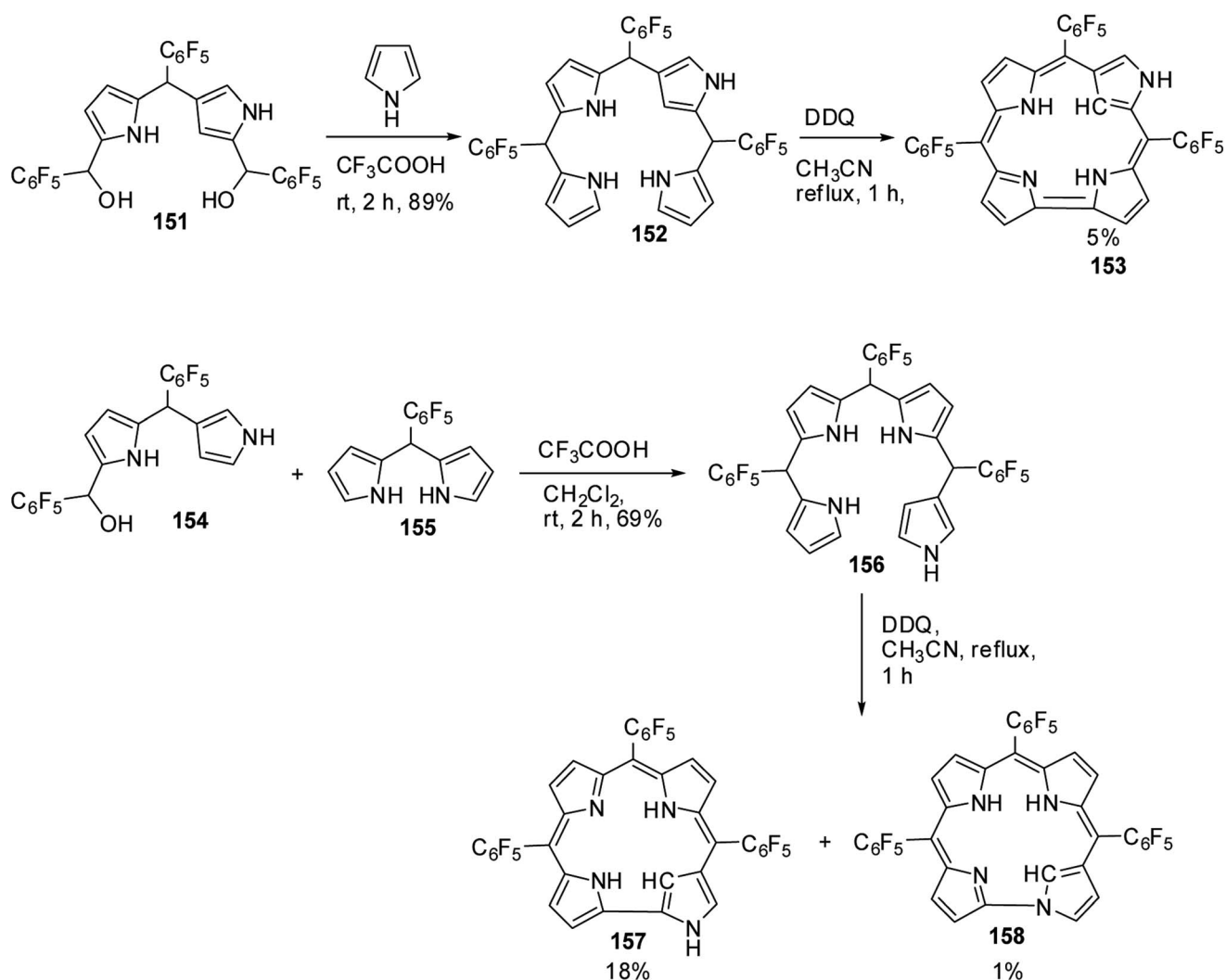


Fig. 22 Top and side view of X-ray crystal structures of **150** (CCDC 859245).

The five-coordinate high-spin (21,23-O₂Cor)Ni^{II}Cl complex **Ni144** was prepared by treating 21,23-dioxacorrolo with nickel(II) chloride in boiling DMF. The ¹H NMR spectral pattern

displayed considerably downfield shifted resonances for β-H of pyrrole and furan ring resonances of **Ni144** indicating its paramagnetic behavior. **Ni144** displayed an absorption pattern



Scheme 34 Synthesis of N-confused corroles **153**, **157** and norrole **158**.



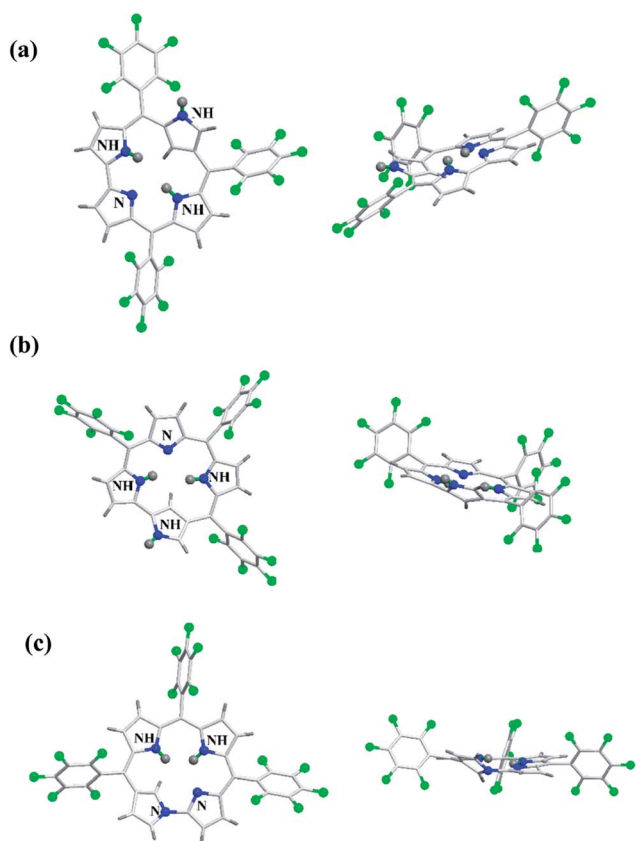


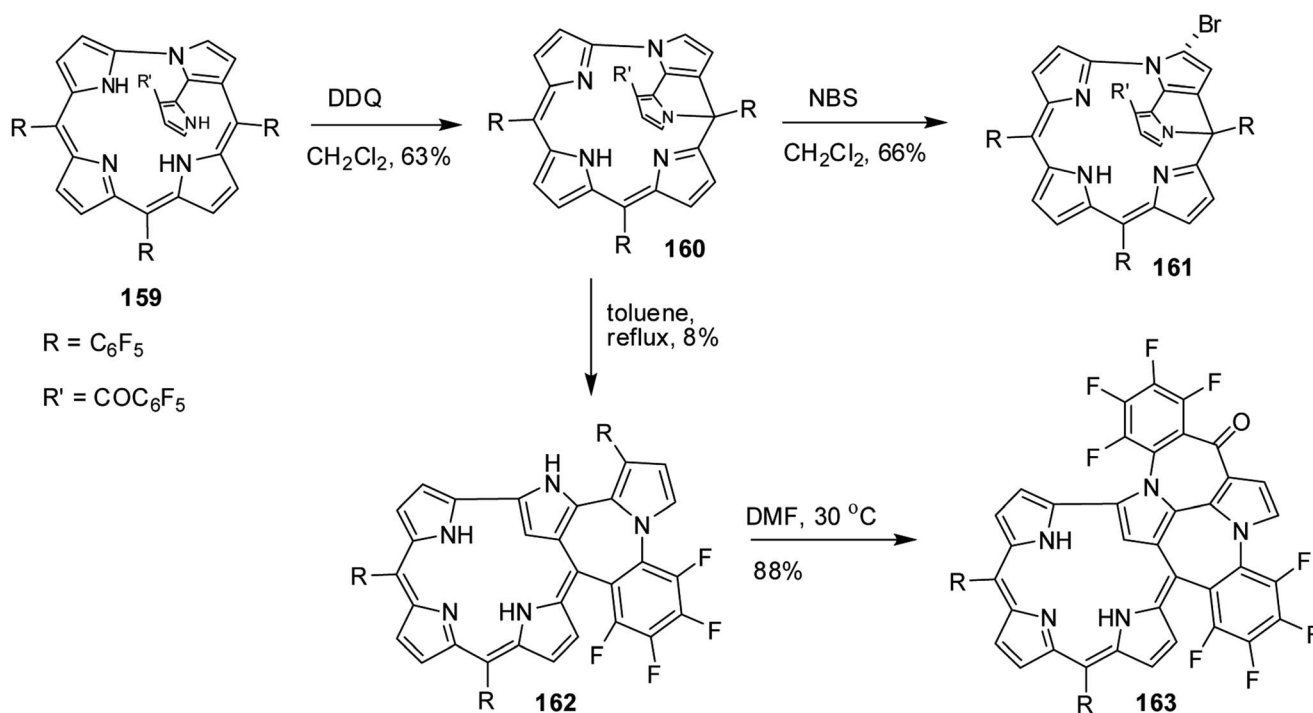
Fig. 23 Top and side view of X-ray crystal structures of (a) **153** (CCDC 719319) (b) **157** (CCDC 719320) (c) **158** (CCDC 719321).

typical of corroles. However, synthesis of Ni complex of dioxacorrole **146** under similar reaction conditions was unsuccessful.

Latos-Grażyński and co-workers⁴¹ reported a unique example of 3-pyranone dioxacorrole **148** by adopting an unprecedented route of Achmatowicz rearrangement in the tetraaryl-21,23-dioxaporphyrin **147** (Scheme 33). The aromatic, symmetrical tetraaryl-21,23-dioxaporphyrin **147** when placed on the basic alumina G II converts to a green compound which was eluted with dichloromethane and identified as 3-pyranone dioxacorrole **148** by detailed 1D & 2D NMR spectroscopy and mass spectral analysis. The formation of **148** was reversible and the acid treatment of **148**, regenerated tetraaryl-21,23-dioxacorrole **147** without the need of any reducing agent. The 3-pyranone dioxacorrole **148** readily forms Pd(II) complex upon treatment with Pd(OAc)₂. The X-ray structure obtained for Pd(II) complex of 3-pyranone dioxacorrole **150** showed that Pd(II) ion was in [CNON] coordination environment and forms an organometallic complex (Fig. 22). The coordination of Pd(II) ion forces the 3-pyranone dioxacorrole ligand to acquire an extraordinary conformation in which the pyranone ring was almost perpendicular to the plane of the rest of the macrocycle.

3.9. N-confused and neo-confused corroles

Furuta and co-workers⁴² reported the synthesis of corrole isomers **153**, **157** and **158** where one of the pyrrole rings adopted a confused conformation (Scheme 34). The N-confused corroles **153** and **157** displayed a relatively sharp singlet corresponding to the interior CH at -0.91 and 1.84 ppm respectively in their ¹H NMR spectra. The inner and peripheral NH protons of **153** and **157** displayed resonance signals in upfield and downfield



Scheme 35 Synthesis of singly and doubly N-C_{Ar}-fused N-Confused corroles **162** and **163**.

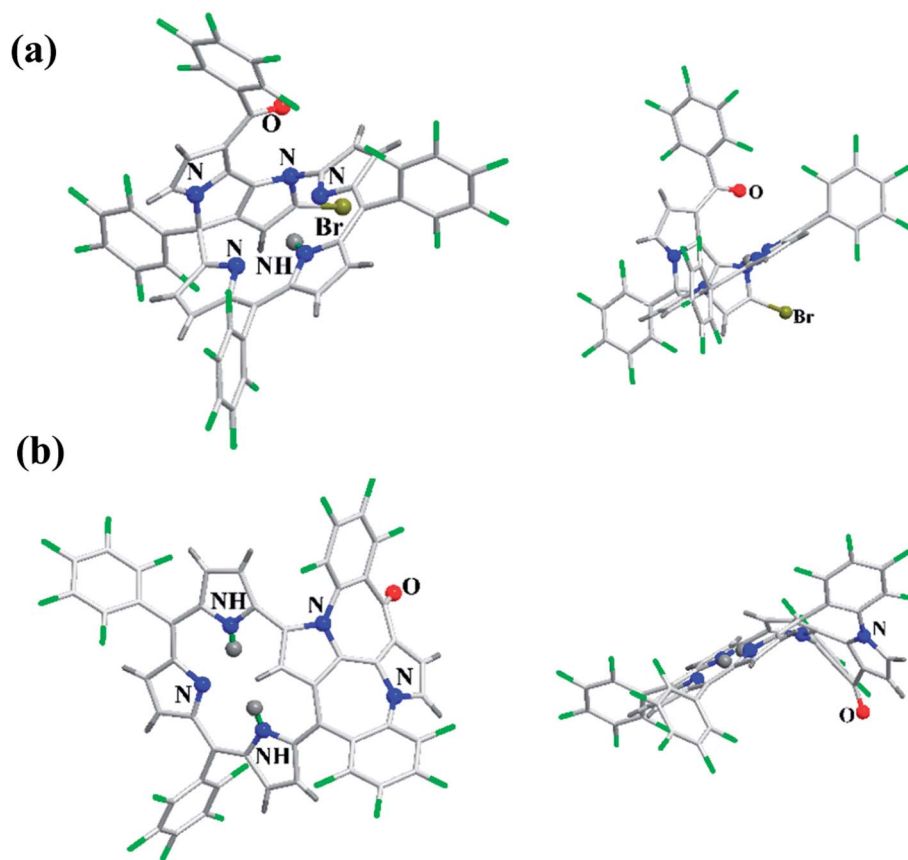


Fig. 24 Top and side view of X-ray crystal structures of (a) **161** (CCDC 1434096) (b) **163** (CCDC 1434097).

regions respectively throwing light on their aromatic nature. However, the corrole **157** was relatively less aromatic than **153**.

The neo-confused corrole (norrole) **158** displayed a unique resonance signal at $\delta = 1.21$ ppm for β -pyrrole CH in its ^1H NMR spectrum. The X-ray structures for N-confused corroles **153**, **157** and norrole **158** is shown in Fig. 23. The confused pyrrole moiety in **153** and **157** was found to be of amino type and was significantly tilted from the corrole mean plane. The confused corroles **153**, **157** and **158** displayed dramatic bathochromic shifts in their absorption bands compared to parent corrole. The neo-confused corrole **158** showed highest quantum yield compared to N-confused corroles **153** and **157** in dichloromethane (Fig. 25).

The other striking feature of these confused corroles was their large Stokes shift values ($774\text{--}1445\text{ cm}^{-1}$) compared to the corresponding regular corrole (221 cm^{-1}). The N-confused corroles **153**, **157** and **158**, by virtue of presence of NH group on confused pyrrole, displayed anion binding properties through hydrogen bonding.

Furuta, Xie and co-workers⁴³ recently disclosed a skeletal transformation reactions of norrole **159** (ref. 44) to its non-aromatic isomer isonorrole **160** and subsequently to N-confused corroles containing one and two N-C_{Ar}-fused ring **162** and **163** respectively (Scheme 35). The authors explored the reactivity of neo-confused corrole **159** containing an appended pyrrole moiety to generate these interesting molecules. The ^1H

NMR spectrum of **160** revealed decrease in number of resonance signals for NH protons confirming the oxidative dehydrogenation. The pyrrolic β -H signals in **160** were upfield shifted compared to **159** as an indication of its nonaromatic character. The bromination of **160** was regioselective and yielded brominated macrocycle **161** (Scheme 35). The compounds **162** and **163** were obtained respectively by refluxing **160** in toluene for ten hours and by treating **162** in DMF at 30°C . The compounds **162** and **163** displayed ^1H NMR spectral patterns typical for aromatic N-confused corroles. The X-ray structures of **161** and **163** are shown in Fig. 24. The absorption spectra of **160**, **162** and **163** revealed two sets of broad bands in the range $340\text{--}380\text{ nm}$ and $460\text{--}750\text{ nm}$, respectively (Fig. 25). The red shifted absorption bands of **162** and **163** compared to **160** is consistent with their respective aromatic nature. Surprisingly, corrole **160**, despite being non-aromatic, showed smallest HOMO–LUMO gap compared to **162** and **163**.

4. Conclusions and outlook

In this article, the synthesis, structure and properties of heterocorroles containing heteroatom(s) at the core and also at a *meso*-position are described. It is evident from the discussion that, although the first examples of heterocorroles were reported long ago, the measurable growth in this area has happened only since 1999. The incorporation of heteroatom on



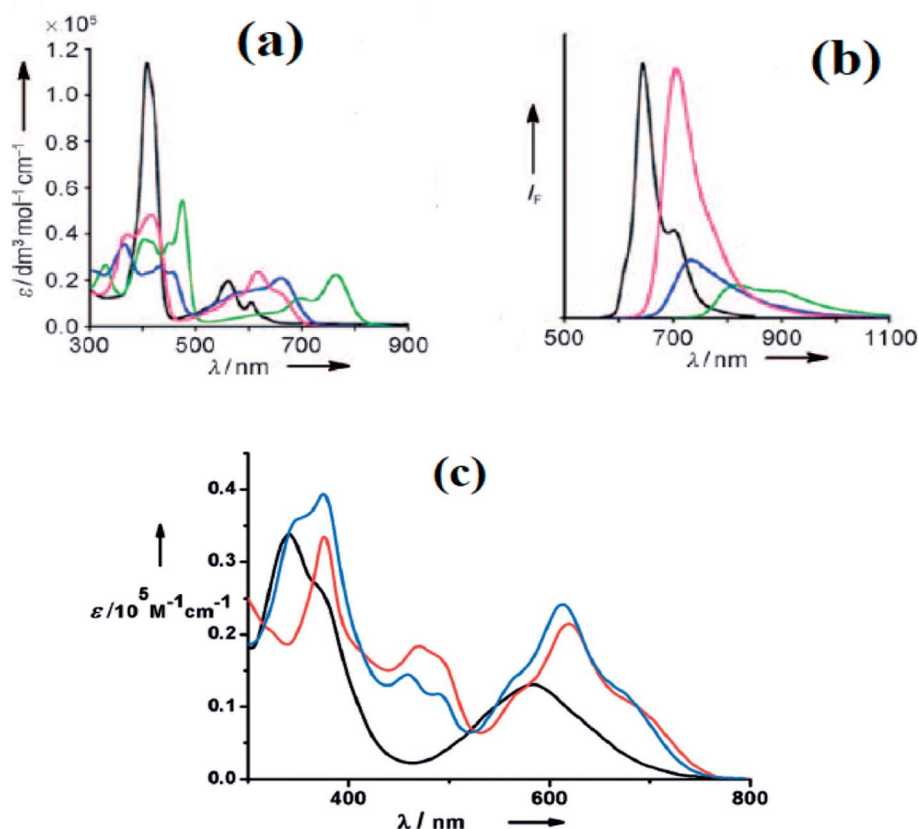


Fig. 25 (a) Absorption spectra of normal corrole (black), 153 (green), 157 (blue) and 158 (red) (b) their emission spectra (λ_{ex} = 634, 764, 662, 659 nm for normal corrole, 153, 157 and 158 respectively) (c) absorption spectra of ring fused norrole 160 (black), 162 (red) and 163 (blue) recorded in dichloromethane. Reproduced from ref. 42 and 43 with permission. Copyright© 2011 and 2016 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

macrocycle's aromatic conjugation pathway has imparted several new properties to heterocorroles. Owing to their contracted cavity size and presence of two inner ionisable NH atoms, heterocorroles have behaved as appropriate ligands to study coordination chemistry of porphyrins. The N-confused corroles and their isomer such as norrole have potential to exhibit novel coordination chemistry owing to the presence of carbon atom in the macrocycle's core. The near-infrared absorption and emission features of 10-silacorroles make them potential candidates for light harvesting applications. However, compared to heteroporphyrins, the progress in heterocorrole chemistry is rather slow presumably due to the difficulty involved in their synthesis and their inherent instability. The design of newer chemical synthesis of heterocorrole derivatives, their conjugates and multichromophoric systems containing heterocorroles would be an interesting proposition. The synthesis of heterocorroles containing heteroatom at a *meso*-position and also at the core without disrupting the aromaticity of the macrocycle will be interesting from their physico-chemical properties and coordination chemistry perspective. We are optimistic that such efforts will be undertaken in near future and the true potential of heterocorroles for various applications will be unveiled.

Conflicts of interest

There are no conflicts of interest to declare.

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References

- 1 H. R. Harrison, O. J. R. Hodder and D. C. Hodgkin, *J. Chem. Soc. B*, 1971, 640–645.
- 2 R. Orłowski, D. Gryko and D. T. Gryko, *Chem. Rev.*, 2017, **117**, 3102–3137, and references cited therein.
- 3 I. Aviv-Harel and Z. Gross, *Coord. Chem. Rev.*, 2011, **255**, 717–736, and references cited therein.



- 4 R. D. Teo, J. Y. Hwang, J. Termini, Z. Gross and H. B. Gray, *Chem. Rev.*, 2017, **117**, 2711–2729, and references cited therein.
- 5 L. Flamigni and D. T. Gryko, *Chem. Soc. Rev.*, 2009, **38**, 1635–1646, and references cited therein.
- 6 I. Aviv-Harel and Z. Gross, *Chem.–Eur. J.*, 2009, **15**, 8382–8394, and references cited therein.
- 7 T. Chatterjee, V. S. Shetti, R. Sharma and M. Ravikanth, *Chem. Rev.*, 2017, **117**, 3254–3328, and references cited therein.
- 8 D. T. Gryko, J. P. Fox and D. P. Goldberg, *J. Porphyrins Phthalocyanines*, 2004, **08**, 1091–1105, and references cited therein.
- 9 A. W. Johnson, I. T. Kay and R. Rodrigo, *J. Chem. Soc.*, 1963, 2336–2342.
- 10 M. Horie, Y. Hayashi, S. Yamaguchi and H. Shinokubo, *Chem.–Eur. J.*, 2012, **18**, 5919–5923.
- 11 H. Omori, S. Hiroto and H. Shinokubo, *Chem. Commun.*, 2016, **52**, 3540–3543.
- 12 H. Omori, S. Hiroto and H. Shinokubo, *Org. Lett.*, 2016, **18**, 2978–2981.
- 13 M. J. Broadhurst, R. Grigg and A. W. Johnson, *J. Chem. Soc., Perkin Trans. 1*, 1972, 1124–1135.
- 14 M. Bröring, F. Brégier, E. C. Tejero, C. Hell and M. C. Holthausen, *Angew. Chem., Int. Ed.*, 2007, **46**, 445–448.
- 15 T. Ito, Y. Hayashi, S. Shimizu, J.-Y. Shin, N. Kobayashi and H. Shinokubo, *Angew. Chem., Int. Ed.*, 2012, **51**, 8542–8545.
- 16 M. Bröring, S. Köhler and C. Kleeberg, *Angew. Chem., Int. Ed.*, 2008, **47**, 5658–5660.
- 17 D. Sakow, B. Böker, K. Brandhorst, O. Burghaus and M. Bröring, *Angew. Chem., Int. Ed.*, 2013, **52**, 4912–4915.
- 18 H. Kamiya, T. Kondo, T. Sakida, S. Yamaguchi and H. Shinokubo, *Chem.–Eur. J.*, 2012, **18**, 16129–16135.
- 19 N. Wachi, T. Kondo, S. Ito, S. Hiroto, J.-Y. Shin and H. Shinokubo, *J. Porphyrins Phthalocyanines*, 2014, **18**, 675–678.
- 20 R. He, H. Yue and J. Kong, *Molecules*, 2017, **22**, 1400.
- 21 D. Sakow, D. Baabe, B. Böker, O. Burghaus, M. Funk, C. Kleeberg, D. Menzel, C. Pietzonka and M. Bröring, *Chem.–Eur. J.*, 2014, **20**, 2913–2924.
- 22 M. Bröring, S. Köhler and C. Pietzonka, *J. Porphyrins Phthalocyanines*, 2012, **16**, 641–650.
- 23 J. Rösner, B. Cordes, S. Bahn Müller, G. Homolya, D. Sakow, P. Schweyen, R. Wicht and M. Bröring, *Angew. Chem., Int. Ed.*, 2017, **56**, 9967–9970.
- 24 H. Omori, S. Hiroto and H. Shinokubo, *Chem.–Eur. J.*, 2017, **23**, 7866–7870.
- 25 V. S. Shetti, U. R. Prabhu and M. Ravikanth, *J. Org. Chem.*, 2010, **75**, 4172–4182.
- 26 I. Gupta and M. Ravikanth, *J. Org. Chem.*, 2004, **69**, 6796–6811.
- 27 J. S. Lindsey, I. C. Schreiman, H. C. Hsu, P. C. Kearney and A. M. Marguerettaz, *J. Org. Chem.*, 1987, **52**, 827–836.
- 28 C.-H. C. Lee, W.-S. Cho, J.-W. Ka, H.-J. Kim and P. H. Lee, *Bull. Korean Chem. Soc.*, 2000, **21**, 429–433.
- 29 W.-S. Cho and C.-H. Lee, *Tetrahedron Lett.*, 2000, **41**, 697–701.
- 30 A. Ghosh, T. Chatterjee, W.-Z. Lee and M. Ravikanth, *Org. Lett.*, 2013, **15**, 1040–1043.
- 31 S. Janusz, L. Latos-Grażyński and L. Szterenber, *Chem.–Eur. J.*, 2008, **14**, 4861–4874.
- 32 S. J. Narayanan, B. Sridevi, T. K. Chandrashekar, U. Englisch and K. Ruhlandt-Senge, *Org. Lett.*, 1999, **1**, 587–590.
- 33 J. Sankar, H. Rath, V. PrabhuRaja, T. K. Chandrashekar and J. J. Vittal, *J. Org. Chem.*, 2004, **69**, 5135–5138.
- 34 H. Kalita, D. Kalita, W.-Z. Lee, J. Bellare and M. Ravikanth, *Chem.–Eur. J.*, 2014, **20**, 10404–10413.
- 35 H. Kalita and M. Ravikanth, *Chem.–Eur. J.*, 2015, **21**, 7399–7402.
- 36 S. Venkatraman, R. Kumar, J. Sankar, T. K. Chandrashekar, K. Sendhil, C. Vijayan, A. Kelling and M. O. Senge, *Chem.–Eur. J.*, 2004, **10**, 1423–1432.
- 37 B. Sridevi, S. Jeyaprakash Narayanan, T. K. Chandrashekar, U. Englisch and K. Ruhlandt-Senge, *Chem.–Eur. J.*, 2000, **6**, 2554–2563.
- 38 J. Sankar, H. Rath, V. Prabhuraja, S. Gokulnath, T. K. Chandrashekar, C. S. Purohit and S. Verma, *Chem.–Eur. J.*, 2007, **13**, 105–114.
- 39 M. J. Broadhurst, R. Grigg and A. W. Johnson, *J. Chem. Soc. D*, 1969, 23–24.
- 40 M. Pawlicki, L. Latos-Grażyński and L. Szterenber, *J. Org. Chem.*, 2002, **67**, 5644–5653.
- 41 M. Pawlicki, D. Bykowski, L. Szterenber and L. Latos-Grażyński, *Angew. Chem., Int. Ed.*, 2012, **51**, 2500–2504.
- 42 K. Fujino, Y. Hirata, Y. Kawabe, T. Morimoto, A. Srinivasan, M. Toganoh, Y. Miseki, A. Kudo and H. Furuta, *Angew. Chem., Int. Ed.*, 2011, **50**, 6855–6859.
- 43 M. Li, P. Wei, M. Ishida, X. Li, M. Savage, R. Guo, Z. Ou, S. Yang, H. Furuta and Y. Xie, *Angew. Chem., Int. Ed.*, 2016, **55**, 3063–3067.
- 44 Y. Xie, P. Wei, X. Li, T. Hong, K. Zhang and H. Furuta, *J. Am. Chem. Soc.*, 2013, **135**, 19119–19122.

