NJC

COMMUNICATION



Cite this: New J. Chem., 2024. **48**, 9424

Received 26th February 2024, Accepted 9th May 2024

DOI: 10.1039/d4nj00913d

rsc.li/njc



View Article Online View Journal | View Issue

Deoxygenation of heterocyclic N-oxides employing iodide and formic acid as a sustainable reductant*

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We present a novel deoxygenation method for heterocyclic Noxides utilizing iodide as a catalyst. Iodide acts as a reducing catalyst that is regenerated by formic acid, which also serves as a Brønsted activator and solvent. The method demonstrates high efficiency and excellent selectivity in the reduction of a variety of heterocyclic N-oxides and tertiary amines. Our computational DFT investigation revealed that the reduction mechanism entails a direct interaction between iodide and the oxygen of the N-oxide within a Mg²⁺/formic acid framework, resulting in the formation of the *N*-heterocycle and the release of a hypoiodite unit. Additionally, a molecular mechanism for the regeneration of iodide from hypoiodite, facilitated by formic acid, is suggested. This method provides an environmentally friendly approach for the deoxygenation of N-oxides and related species.

Pyridines and quinolines are fundamental heterocyclic structures, extensively found in natural products,¹ and playing pivotal roles across various fields. Their significance ranges from their inherent presence in biologically active compounds² to applications in medicinal chemistry (Fig. 1a),³ and materials science.⁴ In consequence, compounds containing N-pyridinic moieties are among the most commonly used in the manufacture of drug candidates.⁵ Unfortunately, the direct functionalization of pyridines and their derivatives through direct electrophilic substitutions imposes a challenge due to poor chemoselectivity and a lower π -system energy compared to benzene.⁶ A compelling strategy for effective pyridine functionalization involves the use of N-activated pyridines,⁷ among which *N*-oxide derivatives (Fig. 1b), with remarkable availability and reactivity, enable functionalization

through various strategies.8 Therefore, after the functionalization of pyridine/quinoline N-oxides, regeneration of the Nheterocycle through deoxygenation would be needed. Among the reductant systems used for heterocyclic N-oxide deoxygenation (NOD) many are based on transition metals (TM)⁹ including Mo, Pd, In, Ir, Ru, Sm, etc., while other TM-free based methods are found in lesser extent.¹⁰

Iodine-based deoxygenations of N-functionalities have emerged as alternatives,^{11,12} particularly since some methods employ green sacrificial reductants.¹² In the case of iodidebased methodologies for NODs, iodide is often used as a sacrificial reductant, in stoichiometric or superstoichiometric quantities, and often paired with metals, including TMs.¹³

The search for cost-effective, efficient, and environmentally friendly methods for the selective reduction of heterocyclic N-oxides is crucial. Formic acid (FA) has emerged as a prominent reducing agent.14 Furthermore, its production from biomass or CO₂ valorization adds a sustainable appeal.¹⁵ Recently, we introduced the I^{-/}FA as a green reagent for reducing methylsulfinates and sulfoxides (Fig. 1c).16 This method utilizes iodide as the catalytic reductant, which is regenerated by formic acid. Additionally, formic acid serves both as the medium/solvent and in the elaboration of an activating Brønsted template. Under the optimal conditions, we observed exceptional reducing efficiency, coupled with remarkable orthogonality to various reducible functional groups.

Following our research program, which focuses on the development of TM-free^{16,17} and iodine/iodide-based synthetic methods,^{16,18} we present a practical method that enables the deoxygenation of pyridine N-oxides (and related species such as N-oxides derived from quinolines, isoquinolines and tertiary amines) using the I⁻/FA reagent. Our experimentation began exposing quinoline N-oxide (1a) to KI (10% mol) in formic medium, under MW irradiation, delivering 40% yield of quinoline (2a) (entry 1, Table 1). As expected, iodide was essential for the reduction as the absence of KI or its replacement by potassium salts such as K₂SO₄ and K₂HPO₄ delivered no reduced product (entries 2-4). Other iodide sources such as

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[†] Electronic supplementary information (ESI) available: Cartesian coordinates of the calculated reaction mechanism. See DOI: https://doi.org/10.1039/d4nj00913d

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Fig. 1 (a) Examples of relevant *N*-oxide pyridine species in medicinal chemistry (MedChem). (b) Pyridine *N*-oxide features enhanced reactivity compared to pyridine. (c) I^-/FA reagent for the efficient reduction of sulfoxides (shown) and methyl sulfinates (not shown). (d) Mgl₂ as a convenient iodide source for a TM-free reduction based on the I^-/FA reagent.

TBAI, NaI, KIO₃, ZnI₂, I₂, and MgI₂ were also evaluated, with the last two (entries 9 and 10) exhibiting superior performance compared to KI. Although KI, I2 and MgI2 delivered comparable results in terms of yield, MgI₂ delivered better reproducibility and chemoselectivity, prompting us to proceed with it for subsequent experiments. We had prior evidence indicating that iodide demonstrated superior performance compared to other halides in similar reductions.¹⁶ This was reinforced for the N-O reduction through additional experiments employing magnesium chloride and bromide (entries 16 and 17), delivering low conversion. No contribution from Mg²⁺ was observed, further supporting the role of iodide as the reducing entity in MgI_2 (cf. entries 10 and 11). From tuning of reaction conditions, we identified 10% mol of MgI2 and 3 h of reaction time as optimal (entry 14). An additional experiment conducted with conventional heating underscored the necessity of microwave activation (cf. entries 13 and 18).^{19,20}

With the optimal conditions at hand, diverse *N*-oxides with varying substituents were submitted to the optimal deoxygenation conditions. Isoquinoline, benzo[h]quinoline, and phenazine *N*-oxides (**1b–1d**) were successfully deoxygenated with comparable yields to those obtained using TM-based technologies (Table 2).⁹

Table 1 Halide source screening for the deoxygenation of quinoline 1-oxide $\left(1a\right)$

$1a \stackrel{\text{[I^{-}]}(x \text{ mol}\%)}{0} \underbrace{HCO_2H}_{MW, 3 \text{ h } 140 \ ^{\circ}\text{C}} 2a$				
Entry	Halide source [I ⁻]	(% mol)	Time (h)	Yield (%) ^a
1	KI	10	1	40
2	_		1	
-	K ₂ SO ₄	10	1	
1	K ₂ HPO ₄	10	1	_
5	TBAI	10	1	39
5	NaI	10	1	37
7	KIO ₂	10	1	37
3	ZnI ₂	5	1	14
9	I ₂	5	1	42
10	 MgI2	5	1	46
11	MgSO ₄	5	1	_
12	MgI ₂	5	3	57
13	MgI ₂	10	1	47
14	MgI ₂	10	3	91/91 ^b
15	MgI_2	10	4	$95/85^{b}$
16	$MgCl_2$	10	3	2
17	MgBr ₂	10	3	2
18 ^c	MgI ₂	10	3	$65/49^{b}$

 a Determined by ¹H-NMR. b Isolated yields. c Reaction performed using a conventional heating reactor at 140 $^\circ C$ (Anton-Paar Monowave[®] 50).

Interestingly, sterically hindered 2,6-methylated pyridine *N*-oxides (**1e**, **1f**) underwent deoxygenation without any detrimental effects, yielding the corresponding pyridines with excellent yields. Furthermore, complete orthogonality was observed for carbonyl-containing *N*-oxides, such as 4-acetylpyridine *N*-oxide (**1h**) and 4-benzoylpyridine *N*-oxide (**1i**). These results significantly support a wider range of conditions under which the I Γ /FA reductive reagent is effective (90–140 °C), without affecting carbonyl groups.²¹ Additionally, a cyano-substituted substrate (**1j**) and 6-MeO-quinoline *N*-oxide (**1m**) underwent chemoselective *N*-deoxygenation, producing the corresponding heterocycles with excellent yields.

Under standard conditions, 3-nitroquinoline N-oxide (1k) underwent reduction of both the N-oxide and nitro groups, accompanied by formylation to yield product 2ka. In contrast, 2-aminopyridine *N*-oxide (1g) was deoxygenated with excellent yield and without formylation, a result attributed to the significant electronic differences between positions 2 and 3 on the pyridine ring.²² After adjusting the reaction conditions, 2k was successfully obtained with the nitro group remaining unaffected (see ESI[†]). As for 4-nitroquinoline N-oxide (11), a Nef-type hydrolytic removal of the nitro group was impossible to avoid, delivering quinolin-4-(H)-one (21) as the sole product.²³ All attempts to adjust the conditions to obtain the nitro-containing deoxygenated product were unsuccessful (see ESI[†]). Finally, N-oxides of tertiary amines such as 1n and 10 were obtained with particularly good yields. Importantly, in all successful experimental trials, the reactor vial was pressurized during the reaction process and maintained a high pressure (>10 psi) after cooling to room temperature, which is consistent with the release of CO2 as byproduct.24 Upon completion of the reductions, the reaction mixtures typically



 a Reaction time 5 h. b Reaction time 6 h. c Reaction time 10 h @ 100 $^\circ C.$ d Determined by $^1 \rm H-NMR.$

developed a brownish hue, indicative of the presence of molecular iodine.

To get insight in the reaction mechanism for the N-O reduction, we performed DFT calculations (see ESI,† for further details) employing pyridine N-oxide as a model and considering the optimal conditions above discussed. First, we explored the interaction between MgI2 and solvent molecules (Fig. 2). In a recent previous work, we demonstrated that the coordination modes of the Mg metal center vary based on the σ -donor capability of the Lewis base.²⁵ Consequently, FA acts as a ligand and magnesium can coordinate up to four units, through the carbonyl O atom, resulting in Mg-O-C angles of approximately 120°. Contrary to what one might expect, FA-ligands are monodentate and not bidentate. The formation of the lower coordination modes A-1 and A-2 is spontaneous, while the penta- and hexacoordinated species, A-3 and A-4, corresponding to sp³d and sp³d² hybridizations at the Mg center, exist in equilibrium with the tetrahedral A-2 mode. The NOD mechanistic pathway and the subsequent iodide regeneration begins with a double substitution of iodide for two FA-ligands within the Mg coordination sphere $(A-4 \rightarrow I-1 \rightarrow I-2, Fig. 3)$. The formation of the $[Mg^{2+}(OCHOH)_6]2I^{-}$ complex (I-2) requires $\Delta G_{A-4 \rightarrow I-2} = +4.2$ kcal mol⁻¹. Subsequently, the spontaneous addition of a pyridine N-oxide (C5H5NO) unit forms I-3 ($\Delta G_{I-2} \rightarrow I-3 = -4.5 \text{ kcal mol}^{-1}$). This addition is deemed favorable due to the hydrogen bonding between the oxygen in pyridine N-oxide and protons from formic acid units. Despite the



Fig. 2 Relative Gibbs free energy (kcal mol⁻¹) profile for the progressive coordination of formic acid ligands to MgI₂ at the (SMD:HCO₂H) ω B97X-D/def2-TZVPP// ω B97X-D/def2-SVP level and 298.15 K.

pKa values of pyridine N-oxide $(0.79)^{26}$ and formic acid $(3.5)^{27}$ suggesting that proton transfer (PT) between these Brønsted species might not be favored, our findings show that this process is both barrierless and exergonic ($\Delta G_{I-3} \rightarrow I-4 = -3.5 \text{ kcal mol}^{-1}$). Thus, it seems that an effect of increased acidity of FA protons, through Mg-FA coordination, facilitates the PT process to access I-4 effortlessly. Next, a nucleophilic substitution (S_N2-type) takes place, centered on the oxygen atom via TS-1 ($\Delta G^{\ddagger}_{I-4} \rightarrow TS-1$ = +32.5 kcal mol⁻¹, in agreement with T = 140 °C), marking the rate-determining step and resulting in the formation of the pyridine-hypoiodous acid pair in I-5. Remarkably, formic acid acts as a Brønsted-activator enabling the electrophilicity of the pyridine N-oxide, so the nucleophilic attack of iodide is feasible. From I-5, a hydrogen transfer process was determined via TS-2A, with an energy barrier of $\Delta G^{\ddagger}_{1-5} \rightarrow TS-2A = +20.1$ kcal mol⁻¹, leading to the production of hydroiodic acid and carbon dioxide (P-A) through a net exothermic process. Then, hydroiodic acid spontaneously transfers its proton to the pyridine supported by a H₂O unit.

We identified an equilibrium between I-5 and a formyl hypoiodite species (I-6, $\Delta G_{\text{I-5}} \rightarrow \text{I-6} = -1.9 \text{ kcal mol}^{-1}$). Finding a transition state connecting these minima proved to be difficult. Starting from I-6, I-5 can also lead to the formation of molecular iodine (I₂, **P-B**) through iodide addition, with an energy barrier of $\Delta G^{\ddagger}_{\text{I-6}} \rightarrow \text{TS-2B} = +9.6 \text{ kcal mol}^{-1}$. These equilibria towards non-iodide productive species determines the molar percentage required for the iodide reactant, as amounts of MgI₂ lower than standard significantly affect the conversion rate (*cf.* entries 12,14, Table 1). However, our thermochemical data suggest that iodine (**P-B**) can be converted into **P-A** by accessing **I-5** and then proceeding through **TS-2A**, requiring an energy investment of $\Delta G^{\ddagger}_{\text{P-B}} \rightarrow \text{TS-2A} = +33.1 \text{ kcal mol}^{-1}$.

Conclusions

We developed a methodology with a sustainable basis for the deoxygenation of pyridine *N*-oxides, as formic acid is employed as activator, solvent, and stoichiometric reductant, using MgI₂ as a convenient iodide source, which serves as a catalytic reductant. This method proved to be compatible with various potentially reducible functional groups, including carbonyl, cyano, benzyl, and to some extent, the nitro group. The same



Fig. 3 Relative Gibbs free energy (kcal mol⁻¹) profile for the pyridine N-oxide deoxygenation using FA-coordinated MgI₂ (A-4) at the (SMD:HCO₂-H) ω B97X-D/def2-TZVPP// ω B97X-D/def2-SVP level and 298.15 K.

system also allowed for the deoxygenation of tertiary amines. The mechanistic study through DFT suggests that the *N*-oxide deoxygenation occurs *via* an S_N2 -type mechanism. Formic acid performs Brønsted-activation, increasing its acidity through Mg^{2+} coordination (Lewis to Brønsted acid relay). Also, the molar concentration of MgI₂ must be such to overcome non-iodide productive oxidized iodine species, such as I₂. The regeneration of iodide mainly occurs through the reduction of hypoiodous acid by hydrogen transfer delivering a net exothermic process. To regenerate iodide from molecular iodine, the pathway involves the reversion towards hypoiodite followed by hydrogen transfer, requiring +33.1 kcal mol⁻¹ to complete the reduction.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work received support from CONACYT/CONAHCYT (Mexico) through a *Ciencia de Frontera* grant (CF-2019-51493). A. C.-J. would like to express gratitude to CONACYT/CONAHCYT for the PhD fellowship 708711 and V.A.L.-R. also acknowledges CONAHCYT for financial support through a MSc fellowship (1312084). P. A. A.-C. and J. B. M.-R. thank CONAHCYT for undergraduate and post-doctoral fellowships, respectively, within grant CF-2019-51493. The authors would like to acknowledge the computing time supported by CONACYT (Grant CB-2015-252356), and additional resources provided by LANCAD and CONACYT/CONAHCYT at the super-

computer hybrid cluster Xiuhcoatl at the General Coordination of Information and Communication Technologies (CGSTIC) of CIN-VESTAV (https://clusterhibrido.cinvestav.mx). Special thanks are extended to Teresa Cortéz, Géiser Cuellar, Ma. Luisa Rodríguez and Víctor González for assistance in spectral acquisition.

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