Chemical Science



EDGE ARTICLE

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2020, 11, 5037

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cobalt-nitrosyl $\{Co(NO)\}^8$ to Co^{II} -nitrito $\{Co^{II}(NO_2^-)\}$: base induced hydrogen gas (H_2) evolution†

Nitric oxide monooxygenation (NOM) reaction of

Sandip Das,^a Kulbir,^a Somnath Ghosh,^a Subash Chandra Sahoo^b and Pankaj Kumar (10 **a)

Here, we report the nitric oxide monooxygenation (NOM) reactions of a Co^{III} -nitrosyl complex (1, {Co-NO}^8) in the presence of mono-oxygen reactive species, *i.e.*, a base (OH⁻, tetrabutylammonium hydroxide (TBAOH) or NaOH/15-crown-5), an oxide (O²⁻ or Na₂O/15-crown-5) and water (H₂O). The reaction of 1 with OH⁻ produces a Co^{III} -nitrito complex {3, (Co^{III} -NO₂⁻)} and hydrogen gas (H₂), *via* the formation of a putative N-bound Co-nitrous acid intermediate (2, {Co-NOOH})⁺). The homolytic cleavage of the O-H bond of proposed [Co-NOOH]⁺ releases H₂ *via* a presumed Co^{III} -H intermediate. In another reaction, 1 generates Co^{III} -NO₂⁻ when reacted with Co^{III} - via an expected Co^{III} -nitro (4) intermediate. However, complex 1 is found to be unreactive towards H₂O. Mechanistic investigations using Co^{III} -NO₂ and Co^{III} -labeled-NaO²H (NaOD) evidently revealed that the N-atom in Co^{III} -NO₂ and the H-atom in H₂ gas are derived from the nitrosyl ligand and OH⁻ moiety, respectively.

Received 17th March 2020 Accepted 24th April 2020

DOI: 10.1039/d0sc01572e

rsc.li/chemical-science

As a radical species, nitric oxide (NO) has attracted great interest from the scientific community due to its major role in various physiological processes such as neurotransmission, vascular regulation, platelet disaggregation and immune responses to multiple infections.1 Nitric oxide synthase (NOS),2 and nitrite reductase (NiR)3 enzymes are involved in the biosynthesis of NO. NOSs produce NO by the oxidation of the guanidine nitrogen in L-arginine.4 However, in mammals and bacteria, NO₂⁻ is reduced to NO by NiRs in the presence of protons, *i.e.*, $NO_2^- + e^- + 2H^+ \rightarrow NO + H_2O.5$ Biological dysfunctions may cause overproduction of NO, and being radical it leads to the generation of reactive nitrogen species (RNS), i.e., peroxynitrite (PN, OONO⁻)⁶ and nitrogen dioxide ('NO₂),⁷ upon reaction with reactive oxygen species (ROS) such as superoxide (O2. -),8 peroxide (H₂O₂), and dioxygen (O₂). Hence, it is essential to maintain an optimal level of NO. In this regard, nitric oxide dioxygenases (NODs)11 are available in bio-systems to convert excess NO to biologically benign nitrate (NO₃⁻).¹²

$$NO_2^- + Fe^{II} + H^+ \leftrightarrow NO + Fe^{III} + OH^-$$
 (1)

$$[M-NO]^n + 2OH^- \rightarrow [M-NO_2]^{(n-2)} + H_2O$$
 (2)

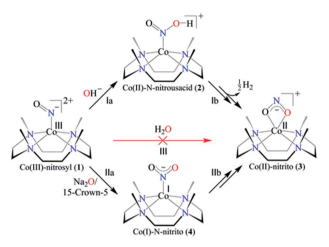
NOD enzymes generate NO₃⁻ from NO;^{11b,12-13} however, the formation of NO₂ from NO is still under investigation. Clarkson and Bosolo reported NO2 formation in the reaction of Co^{III}-NO and O₂. 14 Nam and co-workers showed the generation of Co^{II}-NO₂ from Co^{III}-NO upon reaction with O₂. Recently, Mondal and co-workers reported NO₂ formation in the reaction of Co^{II}-NO with O₂. ¹⁶ Apart from cobalt, the formation of Cu^{II}-NO₂ was also observed in the reaction of Cu^I-NO and O₂. ¹⁷ For metal-dioxygen adducts, i.e., Cr^{III}-O₂. and Mn^{IV}-O₂. NOD reactions led to the generation of Cr^{III}-NO₂⁻ (ref. 18) and Mn^V= O + NO₂⁻, respectively. However, the NOD reaction of Fe^{III}-O₂. and Fe^{III}-O₂²⁻ with NO and NO⁺, respectively, generated Fe^{III}-NO₃ via Fe^{IV}=O and 'NO₂.20 Ford suggested that the reaction of ferric-heme nitrosyl with hydroxide leads to the formation of NO2 and H+.12 Lehnert and co-workers reported heme-based Fe-nitrosyl complexes21 showing different chemistries due to the Fe^{II}-NO⁺ type electronic structures. On the other hand, Bryan proposed that the one-electron reduction of NO₂ to NO in ferrous heme protein is reversible (eqn (1)).²² Also, it is proposed that excess NO in biological systems is converted to NO₂ and produces one equivalent of H⁺ upon reaction with 'OH.23 Previously reported reactivity of M-NOs of Fe24 with OHsuggested the formation of NO₂⁻ and one equivalent of H⁺, where H⁺ further reacts with one equivalent of OH⁻ and produces H₂O (eqn (2)).25

Here in this report, we explore the mechanistic aspects of nitric oxide monooxygenation (NOM) reactions of the Co^{III}-nitrosyl complex, [(12TMC)Co^{III}(NO⁻)]²⁺/{Co(NO)}⁸ (1), ^{15,26}

[&]quot;Department of Chemistry, Indian Institute of Science Education and Research (IISER), Tirupati 517507, India. E-mail: pankaj@iisertirupati.ac.in

^bDepartment of Chemistry, Punjab University, Chandigarh, Punjab, India

[†] Electronic supplementary information (ESI) available. CCDC 1974616. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/d0sc01572e



Scheme 1 Nitric oxide monooxygenation (NOM) reactions of cobalt-nitrosyl complex (1) in the presence of a base (OH $^-$), sodium oxide (Na₂O) and water (H₂O).

bearing the 12TMC ligand (12TMC = 1,4,7,10-tetramethyl-1,4,7,10-tetraazacyclododecane) with mono-oxygen reactive species (O²⁻, OH⁻ and H₂O) (Scheme 1). Complex 1 reacts with the base (OH-, tetrabutylammonium hydroxide (TBAOH)/or NaOH in the presence of 15-crown-5 as the OH source) and generates the corresponding Co^{II}-nitrito complex, [(12TMC) $Co^{II}(NO_2^{-1})^+$ (3), with the evolution of hydrogen gas (H_2) via the formation of a plausible N-bound Co-nitrous acid intermediate ([Co-NOOH]⁺, 2) in CH₃CN at 273 K (Scheme 1, reaction (I)). Also, when 1 reacts with the oxide (O²⁻ or Na₂O in the presence of 15-crown-5), it generates the Co^{II}-nitrito complex (3) via a probable Co^I-nitro, [(12TMC)Co^I(NO₂⁻)] (4), intermediate (Scheme 1, reaction (II)); however, 1 does not react with water (Scheme 1, reaction (III)). Mechanistic investigations using 15Nlabeled-15NO, D-labeled-NaOD and 18O-labelled-18OH demonstrated, unambiguously, that the N and O-atoms in the NO2 ligand of 3 resulted from NO and OH moieties; however, the Hatoms of H2 are derived from OH-. To the extent of our knowledge, the present work reports the very first systematic study of Co^{III}-nitrosyl complex reactions with H₂O, OH⁻ and O²⁻. This new finding presents an alternative route for NO₂⁻ generation in biosystems, and also illustrates a new pathway of H₂ evolution, in addition to the reported literature. 12,27

To further explore the chemistry of $[(12TMC)Co^{III}(NO^-)]^{2+}$ (1), ^{15,26} and the mechanistic insights of NOM reactions, we have reacted it with a base (OH^-) , an oxide (O^{2-}) , and water (H_2O) . When complex 1 was reacted with TBAOH in CH_3CN , the color of complex 1 changed to light pink from dark pink. In this reaction, the characteristic absorption band of 1 (370 nm) disappears within 2 minutes (Fig. 1a; ESI, Experimental section (ES) and Fig. S1a†), producing a Co^{II} -nitrito complex, $[(12TMC)Co^{II}(NO_2^-)]^+$ (3), with H_2 (Scheme 1, reaction (Ib)), in contrast to the previous reports on base induced NOM reactions (eqn (2)). ^{12,25,28} The spectral titration data confirmed that the ratio-metric equivalent of OH^- to 1 was 1:1 (ESI, Fig. S1b†). 3 was determined to be $[(12TMC)Co^{II}(NO_2^-)](BF_4)$ based on various spectroscopic and structural characterization experiments (*vide infra*). ^{15,26b}

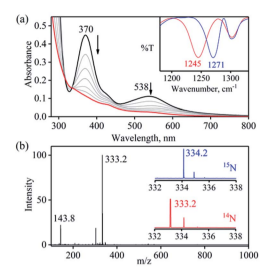


Fig. 1 (a) UV-vis spectral changes of 1 (0.50 mM, black line) upon addition of OH $^-$ (1 equiv.) in CH $_3$ CN under Ar at 273 K. Black line (1) changed to red line (3) upon addition of OH $^-$. Inset: IR spectra of 3^{-14} NO $_2$ $^-$ (blue line) and 3^{-15} NO $_2$ $^-$ (red line) in KBr. (b) ESI-MS spectra of 3. The peak at 333.2 is assigned to [(12TMC)Co II (NO $_2$)] $^+$ (calcd m/z 333.1). Inset: isotopic distribution pattern for 3^{-14} NO $_2$ $^-$ (red line) and 3^{-15} NO $_2$ $^-$ (blue line).

The FT-IR spectrum of 3 showed a characteristic peak for nitrite stretching at 1271 cm $^{-1}$ (Co $^{\text{II}}_{-}$ ¹⁴NO $_{2}$) and shifted to 1245 ${\rm cm}^{-1}({\rm Co}^{\rm II_15}{\rm NO}_2^{\rm I})$ when 3 was prepared by reacting $^{15}{\rm N}$ -labeled NO (Co^{III}₋¹⁵NO) with OH⁻ (Inset, Fig. 1a and Fig. S2†). The shifting of NO_2^- stretching ($\Delta = 30 \text{ cm}^{-1}$) indicates that the Natom in the NO₂ ligand is derived from Co^{III}₂ NO. The ESI-MS spectrum of 3 showed a prominent peak at m/z 333.2, [(12TMC) $Co^{II}(^{14}NO_2^{-})]^+$ (calcd m/z 333.2), which shifted to 334.2, $[(12 \mathrm{TMC})\mathrm{Co^{II}}(^{15}\mathrm{NO_2}^-)]^+$ (calcd m/z 334.2), when the reaction was performed with CoIII_15NO (Inset, Fig. 1b; ESI, Fig. S3a†); indicating clearly that NO₂ in 3 was derived from the NO moiety of 1. In addition, we have reacted 1 with Na¹⁸OH (ES and ESI†), in order to follow the source of the second O-atom in 3-NO₂. The ESI-MS spectrum of the reaction mixture, obtained by reacting 1 with Na¹⁸OH, showed a prominent peak at m/z 335.2, [(12TMC) $Co^{II}(^{18}ONO^{-})]^{+}$ (calcd m/z 335.2), (SI, Fig. S3b†) indicating clearly that NO₂⁻ in 3 was derived from ¹⁸OH⁻. The ¹H NMR spectrum of 3 did not show any signal for aliphatic protons of the 12TMC ligand, suggesting a bivalent cobalt center (Fig.-S4†).266 Furthermore, we have determined the magnetic moment of 3, using Evans' method, and it was found to be 4.62 BM, suggesting a high spin Co(II) metal center with three unpaired electrons (ESI† and ES).29 The exact conformation of 3 was provided by single-crystal X-ray crystallographic analysis (Fig. 2b, ESI, ES, Fig. S5, and Tables T1 and T2†) and similar to that of previously reported Co^{II}-NO₂⁻/M^{II}-NO₂⁻.^{15,26b} Also, we have quantified the amount of nitrite (90 \pm 5%), formed in the above reaction, using the Griess reagent (ESI, ES, and Fig. S6†).

As is known from the literature, a metal-nitrous acid intermediate may form either by the reaction of a metal-nitrosyl with a base²⁷ or by the metal-nitrite reaction with an acid (nitrite reduction chemistry);²⁶ however, the products of both the

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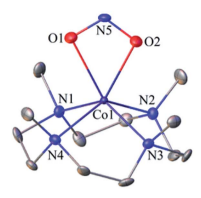
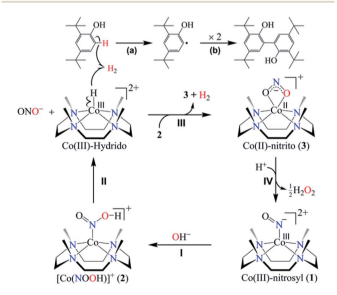


Fig. 2 Displacement ellipsoid plot (20% probability) of **3** at 100 K. Disordered C-atoms of the TMC ring, anion and H-atoms have been removed for clarity.

reactions are different. Here, for the first time, we have explored the reaction of Co^{III}-nitrosyl (1) with a base. In this reaction, it is clear that the formation of Co^{II}-nitrito would be accomplished by the release of H2 gas via the generation of a transient Nbound [Co-(NOOH)] intermediate (Scheme 2, reaction (II)). The formation of Co^{II}-NO₂⁻ (3) from the [Co-(NOOH)]⁺ intermediate is likely to proceed by either (i) homolytic cleavage of the O-H bond and release of H2 via the proposed CoIII-H transient species (Co^{III}-H = Co^{II} + 1/2H₂)³⁰ (Scheme 2, reaction (III)), as reported in previous literature where the reduced cobalt, in a number of different ligand environments, is a good H⁺ reduction catalyst and generates H2 gas via a CoIII-H intermediate31 or (ii) heterolytic cleavage of the O-H bond and the formation of Co^I-NO₂⁻ + H⁺.²⁷ In the present study, we observed the formation of 3 and H₂ via the plausible homolytic cleavage of the NOO-H moiety of 2 as shown in Scheme 2, in contrast to the previous reports on base-induced reactions on metalnitrosyls (eqn (3)).²⁷ Taking together both possibilities, (i) is the most reasonable pathway for the NOM reaction of complex 1 in the presence of a base (as shown in Scheme 2, reaction (III)). And the reaction is believed to go through a Co^{III}-H intermediate as reported previously in Co^I-induced H⁺ reduction in different ligand frameworks and based on literature precedence, we believe that complex 1 acts in a similar manner.³¹

In contrast to an O-bound Co^{II} -ONOH intermediate, where N–O bond homolysis of the ON-OH moiety generates H_2O_2 (Scheme 2, reaction (IV)), set the N-bound [Co-(NOOH)] intermediate decomposes to form NO_2^- and a Co(III)-H transient species, arising from β -hydrogen transfer from the NOO-H moiety to the cobalt-center (Scheme 2, reaction (II)). soa, c, 32 The Co(III)-hydrido species may generate H_2 gas either (a) by its transformation to the Co(II)-nitrito complex (2) and H_2 gas as observed in the case of Co^{III} -H intermediate chemistry soa, c, e-g as proposed in the chemistry of the Co^I complex with H^+ reduction and other metal-hydrido intermediates and also explained in O_2 formation in PN chemistry $I^{17,33}$ or (b) by the reacting with another $[Co-(NOOH)]^+$ intermediate (Scheme 2, reaction (III)).

Furthermore, we have confirmed the H_2 formation in the NOM reaction of $\mathbf{1}$ with OH^- by headspace gas mass spectrometry (Fig. 3a). Also, carrying out the reaction of $\mathbf{1}$ with NaOD leads to the formation of the [Co-(NOOD)]⁺ intermediate, which then transforms to a Co^{III} -D transient species. Further, as described above, the Co^{III} -D species releases D_2 gas, detected by headspace gas mass spectrometry (Fig. 3b), which evidently established that H_2 gas formed in the reaction of $\mathbf{1}$ with OH^- . In this regard, we have proposed that in the first step of this reaction, the nucleophilic addition of OH^- to $\{Co\text{-NO}\}^8$ generates a transient N-bound $[Co\text{-(NOOH)}]^+$ intermediate that is generated by an internal electron transfer to Co^{III} (Scheme 2, reaction (I)). By following the mechanism proposed in the case



Scheme 2 NOM reaction of complex 1 in the presence of OH $^-$, showing the generation of Co II -nitrito (3) and H $_2$ via a Co(III)-hydrido intermediate.

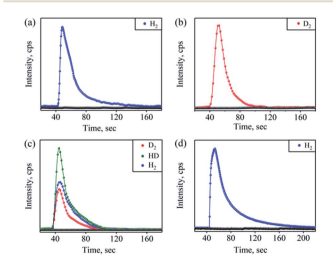
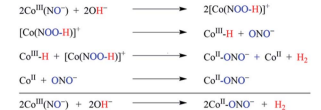


Fig. 3 Mass spectra of formation of (a) H_2 in the reaction of 1 (5.0 mM) with NaOH (5.0 mM), (b) D_2 in the reaction of 1 (5.0 mM) with NaOD (5.0 mM), (c) D_2 , HD, and H_2 in the reaction of 1 (5.0 mM) with NaOD/NaOH (1:1), and (d) H_2 in the reaction of 1 (5.0 mM) with NaOH in the presence of 2,4 DTBP (50 mM).



Scheme 3 NOM reaction of complex 1 in the presence of OH⁻, showing the different steps of the reaction.

of Co^{III} -H, $^{30a-c}$ O_2 , 15 and H_2O_2 (ref. 26b) formation, we have proposed the sequences of the NOM reaction of 1, which leads to the generation of Co^{II}-nitrito and H₂ (Scheme 2, reaction (I)-(III) and Scheme 3). In the second step, O-H bond homolytic cleavage generates a Co^{III}-H transient species + NO₂⁻ via a βhydrogen elimination reaction of the [Co-(NOOH)]+ intermediate.32 The CoIII-H intermediate may undergo the following reactions to generate H2 gas and CoII-nitrito either (a) by the natural decomposition of the Co^{III}-H transient species to generate H₂, 30a,c,e-g or (b) by the H-atom abstraction from another [Co-(NOOH)]⁺ intermediate (Scheme 3). Also, to validate our assumption that the reaction goes through a plausible N-bound [Co-(NOOH)]+ intermediate followed by its transformation to the CoIII-H species (vide supra), we have performed the reaction of 1 with NaOH/NaOD (in 1:1 ratio). In this reaction, we have observed the formation of a mixture of H2, D2, and HD gases, which indicates clearly that the reaction goes through the formation of Co^{III}-H and Co^{III}-D transient species via the aforementioned mechanism (Fig. 3c). This is the only example where tracking of the H atoms has confirmed the H2 generation from an N-bound NOO-H moiety as proposed for H₂ formation from Co^{III}-H.³⁰

While, we do not have direct spectral evidence to support the formation of the transient N-bound [Co-(NOOH)]⁺ intermediate and its decomposition to the CoIII-H transient species via βhydrogen transfer from the NOOH moiety to the cobalt center, support for its formation comes from our finding that the reactive hydrogen species can be trapped by using 2,4-di-tertbutyl-phenol (2,4-DTBP).34 In this reaction, we observed the formation of 2,4-DTBP-dimer (2,4-DTBP-D, ~67%) as a single product (ESI, ES, and Fig. S7†). This result can readily be explained by the H-atom abstraction reaction of 2,4-DTBP either by [Co-(NOOH)]⁺ or Co^{III}-H, hence generating a phenoxyl-radical and 3 with H2 (Fig. 3d and Scheme 2, reaction (a)). Also, we have detected H2 gas formation in this reaction (ESI,† ES, and Fig. 3d). In the next step, two phenoxyl radicals dimerized to give 2,4-DTBP-dimer (Scheme 2c, reaction (II)). Thus, the observation of 2,4-DTBP-dimer in good yield supports the proposed reaction mechanism (Scheme 2, reaction (a) and (b)). Further, the formation of 2,4 DTBP as a single product also rules out the formation of the hydroxyl radical as observed in the case of an O-bound nitrous acid intermediate.26b

Furthermore, we have explored the NOM reactivity of 1 with $Na_2O/15$ -crown-5 (as the O^{2-} source) and observed the formation of the Co^{I-} nitrito complex (3) *via* a plausible Co^{I-} nitro (4)

intermediate (Scheme 1, reaction (IIa); also see the ESI† and ES); however, 1 was found to be inert towards H2O (Scheme 1, reaction (III); also see the ESI, ES and Fig. S8†). The product obtained in the reaction of 1 with O2- was characterized by various spectroscopic measurements. 15,26b The UV-vis absorption band of 1 ($\lambda_{max} = 370$ nm) disappears upon the addition of 1 equiv. of Na₂O and a new band ($\lambda_{max} = 535$ nm) forms, which corresponds to 3 (ESI, Fig. S9†). The FT-IR spectrum of the isolated product of the above reaction shows a characteristic peak for Co^{II}-bound nitrite at 1271 cm⁻¹, which shifts to 1245 cm⁻¹ when exchanged with ¹⁵N-labeled-NO (¹⁵N¹⁶O) (ESI, ES, and Fig. S10†), clearly indicating the generation of nitrite from the NO ligand of complex 1.26b The ESI-MS spectrum recorded for the isolated product (vide supra) shows a prominent ion peak at m/z 333.1, and its mass and isotope distribution pattern matches with $[(12\text{-TMC})\text{Co}^{\text{II}}(\text{NO}_2)]^+$ (calc. m/z 333.1) (ESI, Fig. S11†). Also, we quantified the amount of 3 (85 \pm 5%) by quantifying the amount of nitrite (85 \pm 5%) using the Griess reagent test (ESI, ES, and Fig. S6†).

In summary, we have demonstrated the reaction of Co^{III}nitrosyl, [(12-TMC)Co^{III}(NO⁻)]²⁺/{CoNO}⁸ (1), with mono-oxygen reactive species (O²⁻, OH⁻ and H₂O) (Scheme 1). For the first time, we have established the clear formation of a Co^{II}-nitrito complex, $[(12TMC)Co^{II}(NO_2^-)]^+$ (3), and H₂ in the reaction of 1 with one equivalent of OH via a transient N-bound [Co-(NOOH)]⁺ (2) intermediate. This [Co-(NOOH)]⁺ intermediate undergoes the O-H bond homolytic cleavage and generates a Co^{III}-H transient species with NO₂⁻, via a β-hydrogen elimination reaction of the [Co-(NOOH)]⁺ intermediate, which upon decomposition produces H2 gas. This is in contrast to our previous report, where acid-induced nitrite reduction of 3 generated 1 and H₂O₂ via an O-bound Co^{II}-ONOH intermediate.26b Complex 1 was found to be inert towards H2O; however, we have observed the formation of 3 when reacted with O^{2-} . It is important to note that H_2 formation involves a distinctive pathway of O-H bond homolytic cleavage in the [Co-(NOOH)]⁺ intermediate, followed by the generation of the proposed Co^{III}-H transient species (Co^{II} + 1/2H₂)³⁰ prior to H₂ evolution as described in CoI chemistry with H⁺ in many different ligand frameworks.31 The present study is the first-ever report where the base induced NOM reaction of Co^{III}-nitrosyl (1) leads to Co^{II}-nitrito (3) with H₂ evolution via an N-bound [Co-(NOOH)] intermediate, in contrast to the chemistry of O-bound Co^{II}-ONOH^{26b}, hence adding an entirely new mechanistic insight of base induced H2 gas evolution and an additional pathway for NOM reactions.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by Grants-in-Aid (Grant No. EEQ/2016/000466) from SERB-DST. We acknowledge Prof. Wonwoo Nam and co-workers for the stabilization of initial complexes and providing a ground to develop new chemistry. We acknowledge

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Prof. K. N. Ganesh (Director, IISER Tirupati) for continuous guidance and support, special thanks to Dr Sayam Sen Gupta (IISER Kolkata), Dr E. Balaraman (IISER Tirupati) and Dr R. O. Ramabhadran (IISER Tirupati) for fruitful discussion and support. SCS thanks DST-FIST for the single crystal facility at

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