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Jonathan Benny 💿 ; Toru Saito 🗢 💿 ; Jianbo Liu 🗢 💿

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# Nitrosation mechanisms, kinetics, and dynamics of the guanine and 9-methylguanine radical cations by nitric oxide—Radical-radical combination at different electron configurations



- <sup>2</sup>The Ph.D. Program in Chemistry, the Graduate Center of the City University of New York, 365 5th Ave., New York, New York 10016, USA
- <sup>3</sup>Department of Biomedical Information Science, Graduate School of Information Science, Hiroshima City University, 731-3194 Hiroshima, Japan
- <sup>a)</sup>Authors to whom correspondence should be addressed: tsaito@hiroshima-cu.ac.jp and jianbo.liu@qc.cuny.edu. Tel.: 1-718-997-3271

## ABSTRACT

As a precursor to various reactive nitrogen species formed in biological systems, nitric oxide ( $^{\circ}NO$ ) participates in numerous processes, including enhancing DNA radiosensitivity in ionizing radiation-based radiotherapy. Forming guanine radical cations is another common DNA lesion resulting from ionization and oxidation damage. As such, the interaction of  $^{\circ}NO$  with guanine radical cations ( $G^{\bullet+}$ ) may contribute to the radiosensitization of  $^{\circ}NO$ . An intriguing aspect of this process is the participation of multiple spin configurations in the reaction, including open-shell singlet  $^{1,CS}[G^{\bullet+}(\uparrow)\cdots(\downarrow)^{\bullet}NO]$ , closed-shell singlet  $^{1,CS}[G(\uparrow\downarrow)\cdots NO^+]$ , and triplet  $^{3}[G^{\bullet+}(\uparrow)\cdots(\uparrow)^{\bullet}NO]$ . In this study, the reactions of  $^{\circ}NO$  with both unsubstituted guanine radical cations (in the 9HG $^{\bullet+}$  conformation) and 9-methylguanine radical cations (9MG $^{\bullet+}$ , a guanosine-mimicking model compound) were investigated in the absence and presence of monohydration of radical cations. Kinetic-energy dependent reaction product ions and cross sections were measured using an electrospray ionization guided-ion beam tandem mass spectrometer. The reaction mechanisms, kinetics, and dynamics were comprehended by interpreting the reaction potential energy surface using spin-projected density functional theory, coupled cluster theory, and multiconfiguration complete active space second-order perturbation theory, followed by RRKM kinetics modeling. The combined experimental and computational findings revealed closed-shell singlet  $^{1,CS}[7-NO-9MG]^+$  as the major, exothermic product and triplet  $^{3}[8-NO-9MG]^+$  as the minor, endothermic product. Singlet biradical products were not detected due to high reaction endothermicities, activation barriers, and inherent instability.

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#### I. INTRODUCTION

Among the natural DNA nucleobases, guanine exhibits the lowest ionization<sup>1,2</sup> and oxidation potential,<sup>3,4</sup> which renders it a dominant target in DNA for photolysis,<sup>5,6</sup> ionizing<sup>7,8</sup> and ion-beam radiation,<sup>9</sup> transition metal-mediated oxidation,<sup>10</sup> electrocatalytic oxidation,<sup>11</sup> and photooxidation.<sup>12</sup> The formation of guanine radical cations ( $G^{\bullet+}$ ) as a result of these oxidative damages exacerbates

DNA lesions.<sup>13–18</sup> A chemically and biologically relevant question arises regarding whether and how the  $G^{\bullet+}$  lesion interacts with reactive oxygen species (ROS)<sup>19</sup> and reactive nitrogen species (RNS)<sup>20,21</sup> generated during pathologic events. In recent years, our laboratory has investigated the oxidative modifications of  $G^{\bullet+}$  by singlet O<sub>2</sub> (one of the major ROS in biological systems).<sup>22–26</sup> The present study expands our research to the reaction of  $G^{\bullet+}$  with nitric oxide ( $^{\bullet}$ NO) radical.<sup>21</sup> The latter serves as a precursor to various RNS (e.g., N<sub>2</sub>O<sub>3</sub>, ONOO<sup>-</sup>, transitional metal-nitrosyl) in biological systems.<sup>21</sup> The intriguing aspects of this radical-radical reaction are the formation, internal conversions (ICs), and inter-system crossing (ISC) of multiple electron configurations (i.e., open-shell singlet, closed-shell singlet, and triplet), which controls reaction mechanisms and dynamics as demonstrated in our recent study on the spin-orbit charge transfer between G<sup>++</sup> and <sup>•</sup>NO and the induced tripletsinglet ISC.<sup>27</sup> Charge transfer represents only one possible pathway in the radical-radical reaction. The complexity, variability, and versability of the reaction can manifest through other pathways that emerge as the radicals come into close proximity. In this work, we traced spin dynamics from singlet and triplet ion-molecule encounters through reaction precursors to covalent nitrosation adducts such as [X-NO-G]<sup>+</sup> (wherein a nitroso group -N=O is bonded to the X position at guanine), [X-ON-G]<sup>+</sup> (wherein the O terminal of nitric oxide is bonded to the X position at guanine), [X, Y-NO-G]<sup>+</sup> and [X, Y-ON-G]<sup>+</sup> [wherein the N and O termini (or vice versa) of NO are cyclo-added to guanine across the X-Y bridge]. We aimed to elucidate the roles that different spin configurations play in short-range interaction and combination of the two radicals. The biological relevance of this work is evident in the increased radiosensitivity of DNA in ionizing radiation-based radiotherapy in the presence of •NO.<sup>28-38</sup> Radiosensitization by •NO arises from its reaction with short-lived DNA radicals generated by radiation.<sup>31,34,37,39,40</sup> However, studies on this subject have been scarce, with the existing literature primarily focusing on the reactions between \*NO and the \*OH adducts of purine and pyrimidine.34

The present work started with the formation of the radical cations of guanine (in its O6-keto-N9H form, abbreviated as  $9HG^{\bullet+}$ ) and 9-methylguanine (abbreviated as 9MG<sup>•+</sup>, where the methyl group mimics the nucleoside sugar) in the gas phase, followed by the reactions of these radical cations with \*NO under single-collision conditions provided in a guided-ion beam scattering experiment. The electronic structures and cross sections of product ions were determined as a function of kinetic energy. One of the primary objectives of this research was to benchmark theoretical methods capable of handling multireferential characters and non-adiabatic transitions in the reactions of doublet-doublet electronic states. To this end, experimental results were compared with reaction potential energies calculated using various theoretical approaches, including density functional theory (DFT), coupled cluster theory (CCSD),<sup>41–43</sup> multireference complete active space self-consistent field method (CASSCF),44,45 and complete active space secondorder perturbation theory (CASMP2<sup>46-48</sup> and CASPT2<sup>48,49</sup>). The combined experimental and computational findings were able to unveil explicit, electron configuration-dependent nitrosation mechanisms.

### II. EXPERIMENTAL AND COMPUTATIONAL SECTION

### A. Experimental procedures

All chemicals used, including 2'-deoxyguanosine (dGuo, Sigma-Aldrich, >99%), 9MG (Chemodex, >98%), Cu(NO<sub>3</sub>)<sub>2</sub> (Alfa Aesar, 99.999%), nitric oxide gas (Linde, >99.5%), and HPLC-grade methanol and water, were purchased from commercial vendors and used without further purification. Ion-molecule reactions

were carried out on a home-built guided-ion beam tandem mass spectrometer equipped with an electrospray ionization (ESI) source, a radiofrequency (rf) hexapole ion guide, a reactant quadrupole mass filter, an rf octopole ion guide surrounded by a scattering cell, a product quadrupole mass filter, and a pulse-counting electron multiplier detector. The apparatus and operational procedures were previously published.<sup>27,50</sup>

Following literature methods for producing radical cations peptides and nucleosides,<sup>51-53</sup> 9HG<sup>++</sup> was produced by of collision-induced intramolecular electron-transfer dissociation of a Cu<sup>II</sup>-nucleoside complex, i.e.,  $[Cu^{II}(dGuo)_3]^{\bullet 2+} \rightarrow [Cu^{I}(dGuo)_2]^+$ + dGuo<sup>++</sup> followed by cleavage of the N-glycosidic bond via dGuo<sup>++</sup>  $\rightarrow$  9HG<sup>•+</sup> + C<sub>5</sub>H<sub>8</sub>O<sub>3</sub>. In the experiment, a mixture of 0.25 mM dGuo and 0.25 mM Cu(NO<sub>3</sub>)<sub>2</sub> in methanol/water (v/v = 3:1) was electrosprayed into a heated (190 °C) desolvation capillary, which leads into the mass spectrometer source chamber. A 1.0-mm-diameterorifice skimmer, positioned 3 mm from the capillary end, separates the source chamber and the hexapole ion guide. An electric field between the capillary (biased at 117 V with respect to ground) and the skimmer (biased at 18 V with respect to ground) prompted collision-induced dissociation of [Cu<sup>II</sup>(dGuo)<sub>3</sub>]<sup>•2+</sup> formed in the electrospray to dGuo<sup>•+</sup> and 9HG<sup>•+</sup>, with the latter adopting the spe-cific O6-keto-N9H conformer.<sup>54,55</sup> Similarly, 9MG<sup>•+</sup> was produced by the dissociation of [Cu<sup>II</sup>(9MG)<sub>3</sub>]<sup>•2+</sup> formed from equimolar 0.25 mM 9MG and Cu(NO<sub>3</sub>)<sub>2</sub> in methanol/water. The primary ion beam of 9HG<sup> $\bullet^+$ </sup> had an intensity of 1 × 10<sup>5</sup> counts/s, while that of 9MG<sup>•+</sup> had an intensity of  $5 \times 10^5$  counts/s. When the desolvation capillary was mildly heated (160-180 °C), monohydrated  $9MG^{\bullet+}$ ·H<sub>2</sub>O (4 × 10<sup>4</sup> counts/s) was produced alongside anhydrous radical cations.

The skimmed radical cation beam was directed into the hexapole ion guide for energy dampening and collisional <sup>6,57</sup> with the background gas at a pressure of ~20 mTorr. focusing After passing the hexapole, ions were thermalized to a kinetic energy determined by the hexapole DC bias potential (1.0 V). The ions also exhibited 310 K Maxwell-Boltzmann vibrational (Evib) and rotational (Erot) energy distributions.<sup>50</sup> Radical ions of interest were then mass-selected by the reactant quadrupole mass filter and injected into the octopole ion guide that passes through an 11-cm-long scattering cell containing the nitric oxide target gas at a pressure of 0.02 mTorr. Similar to the hexapole, the octopole ion guide was operated with a combination of DC bias and rf potential. The rf potential trapped ions radially, while the DC potential accelerated or decelerated reactant ions at the octopole entrance to achieve the desired ion kinetic energy in the laboratory frame ( $E_{lab}$ ). The absolute zero and full width at half-maximum (FWHM) of  $E_{\text{lab}}$  were calibrated by retarding potential measurements.<sup>50,58</sup> The center-of-mass collision energy  $(E_{CM})$  for the reaction of  $\bullet$ NO with radical cations was determined by  $E_{\rm CM} = E_{\rm lab} \times m_{\rm neutral}/(m_{\rm neutral})$ +  $m_{\rm ion}$ ), where  $m_{\rm neutral}$  and  $m_{\rm ion}$  are the masses of the neutral and ionic reactants, respectively. The uncertainty for the absolute zero of  $E_{lab}$  was <0.1 eV, and the FWHM of  $E_{lab}$  was 0.55 eV. These corresponded to an  $E_{\rm CM}$  uncertainty of  $\leq 0.015$  eV and a FWHM of  $\leq 0.09$  eV. The nitrosation reaction was studied over an  $E_{\rm CM}$ range of 0.05-1.0 eV and under single ion-molecule collision conditions rendered by sufficiently low <sup>•</sup>NO gas pressure within the cell. Product ions and the remaining reactant ions were collected by the octopole and refocused into the second quadrupole mass

filter for analysis and counting. The thin-target collision condition allowed the calculation of reaction cross sections based on the ratio of product to reactant ion intensities, the <sup>•</sup>NO pressure in the scattering cell, and the effective cell length for collisions, following the Beer–Lambert law.<sup>59,60</sup> Each experiment was repeated three times to determine the relative measurement uncertainty.

#### **B.** Calculations

#### 1. DFT geometry optimization

Geometries of reactants, precursor complexes, transition states (TSs), and products for the reaction of 9MG<sup>•+</sup> with <sup>•</sup>NO were optimized using the DFT  $\omega$ B97XD functional coupled with the 6-31+G(d,p) basis set. The range-separated  $\omega$ B97XD functional mitigated self-interaction errors and improved the orbital description of radical cations.<sup>61</sup> To distinguish between open- and closed-shell singlet structures, singlet biradicals were computed using the broken symmetry (BS) unrestricted DFT with the guess = mix option (i.e., mixing HOMO and LUMO to break  $\alpha$ - $\beta$  and spatial symmetries in an initial guess),<sup>62</sup> whereas closed-shell singlet structures were computed using restricted DFT. All TSs exhibited only one imaginary frequency, consistent with expected reaction pathways. Intrinsic reaction coordinate (IRC) calculation was carried out for each TS to verify that it connects to the correct reactant and product structures.

DFT calculations were conducted using Gaussian 16.<sup>63</sup> Reaction enthalpies ( $\Delta$ H) were computed at 298 K, incorporating zero-point energies (ZPEs, scaled by 0.975<sup>64</sup>) and thermal corrections. Atomic charges, spin densities, and  $\alpha$ - and  $\beta$ -spins were analyzed using the Natural Bond Orbital Analysis Program, NBO 6.0.<sup>65</sup> A computational difficulty arose due to the open-shell, multi-reference character of °NO, which is akin to triplet instability in a doublet system.<sup>66,67</sup> Consequently, the  $\omega$ B97XD/6-31+G(d,p) method overestimated the ionization energy (IE) of °NO by 0.5 eV compared to the experimental adiabatic IE (9.2643 ± 0.0005 eV, measured by pulsed-field ionization photoelectron spectroscopy),<sup>68</sup> as shown in Table S1 of the supplementary material. For this reason, the DFT energy of the °NO reactant was obtained by subtracting the experimental IE(°NO) from the DFT energy of NO<sup>+</sup>.

# 2. Assessment of spin contamination using coupled-cluster theory

T1 diagnostic<sup>69,70</sup> for spin contamination in DFT-optimized reaction structures was carried out using the domain-based local pair-natural orbital coupled-cluster method with single, double, and perturbative triple excitations [DLPNO-CCSD(T)]<sup>71</sup> paired with the aug-cc-pVTZ basis set. The T1 value was calculated by  $T1 = ||t_1|| / \sqrt{n}$ , where  $t_1$  is the Frobenius norm of the single-excitation amplitude vector and n is the number of correlating electrons.<sup>69</sup> T1 of <0.02 for closed-shell systems and that of <0.03 for open-shell systems indicate minimal spin contamination, T1 ranging from 0.06 to 0.07 indicates the moderate multireference character, and T1 of  $\geq 0.16$  indicates the severe multireference character.<sup>73</sup> DLPNO-CCSD(T) has been validated against the gold standard method CCSD(T).<sup>41-43</sup> The comparisons have demonstrated an accuracy of 4 kJ/mol or better for DLPNO-CCSD(T) in estimating reaction enthalpies<sup>74,75</sup> and barriers.<sup>73,76</sup> The inclusion of perturbative correction for triple excitation helps mitigate

deficiencies arising from a single-determinant reference, enabling DLPNO-CCSD(T) to effectively manage moderate spin contamination.

DLPNO-CCSD(T) calculations were carried out using ORCA 4.2,<sup>77,78</sup> with the NormalPNO (pair natural orbital) parameters<sup>73,74</sup> of TCutPair =  $10^{-4}$ , TCutPNO =  $3.3 \times 10^{-7}$ , and TCutMKN =  $10^{-3}$ . The DLPNO-CCSD(T)-calculated IE(•NO) (9.20 eV, see Table S1 of the supplementary material) is only 0.06 eV less than the experimental value.

#### 3. Spin purification using approximate spin projection

Open-shell singlets are prone to spin contamination in unrestricted DFT calculations;<sup>79</sup> therefore, Yamaguchi's approximate spin projection  $(AP)^{27,80-82}$  was utilized for spin purification in energy calculations as  $E = \frac{\langle S^2 \rangle^{HS}}{\langle S^2 \rangle^{HS} - \langle S^2 \rangle^{BS}} E^{BS} - \frac{\langle S^2 \rangle^{BS}}{\langle S^2 \rangle^{HS} - \langle S^2 \rangle^{BS}} E^{HS}$ , where  $E^{BS}$  and  $\langle \hat{S}^2 \rangle^{BS}$  represent the total energy and the expectation value of the total spin angular momentum operator for a target broken-symmetry singlet, respectively, and  $E^{HS}$  and  $\langle \hat{S}^2 \rangle^{HS}$  are the counterparts for the triplet.

# 4. Complete active space second-order perturbation theory

Reaction energies were further explored using CASSCF<sup>44,45</sup> coupled with the 6-31+G(d,p) basis set. Active spaces were defined as (9,7) for 9MG<sup>•+</sup>, (11,8) for <sup>•</sup>NO, and (20,15) for adducts, including critical orbitals  $\sigma_{O(2s)-N(2s)}$ ,  $\sigma^*_{O(2s)-N(2s)}$ ,  $2 \times \pi_{O(2p)-N(2p)}$ ,  $\sigma_{O(2p)-N(2p)}$ ,  $2 \times \pi^*_{O(2p)-N(2p)}$ , and  $\sigma^*_{O(2p)-N(2p)}$  in  ${}^{\bullet}NO$ , as well as the  $\pi$  and  $\sigma_{N-H}$  in 9MG<sup>•+</sup>. However, CASSCF significantly underestimated IE(<sup>•</sup>NO) (see Table S1 of the supplementary material), likely because it included primarily nondynamic electronic correlation. To capture a significant part of dynamic correlation energy, second-order Møller-Plesset perturbation theory was incorporated by using CASMP2.  $^{46-48}$  Pople basis sets 6-31+G(d,p), 6-311+G(d,p), and 6-311++G(d,p) and the atomic natural orbital-relativistic corecorrelated valence triple-& plus polarization functions ANO-RCC-VTZP<sup>83,84</sup> were employed for the CASMP2 calculations of IE. Surprisingly, computational errors escalated notably with the expansion of the basis set, as listed in Table S1 of the supplementary material. The CASMP2 seems to have achieved the best result for  $IE(^{\circ}NO)$  (9.06 eV) with the 6-31+G(d,p) basis set, albeit potentially fortuitous.

To rectify these issues, an alternative version of the complete active space perturbation theory, CASPT2,<sup>48,49</sup> was adopted. A potential drawback of CASPT2 concerns an intruder state in which the zeroth-order energy associated with the CASSCF wavefunction is nearly equal to that of a perturber (correction function).<sup>85</sup> This problem was reduced by adopting an imaginary shift correction (0.1).<sup>86</sup> In addition, an ionization potential-electron affinity (IPEA) shift parameter (0.25 a.u.)<sup>87</sup> was applied to balance the descriptions of open- and closed-shell configurations in the zerothorder Hamiltonian.<sup>88</sup> The IE( $^{\circ}$ NO) of 9.12eV (see Table S1 of the supplementary material), calculated using a combination of CASPT2 and the ANO-L-VTZP basis set (it contains the same primitive basis functions as ANO-RCC but differs in contraction coefficient, which is determined for nonrelativistic calculations),<sup>89,90</sup> showed good agreement with the experimental value. CASSCF and CASMP2 calculations were carried out using Gaussian 16,<sup>63</sup> with the ANO-RCC-VTZP orbital obtained from the basis set exchange.<sup>91</sup> CASPT2 calculations were carried out using OpenMolcas 23.10.<sup>92,93</sup> Reaction enthalpies reported at CASPT2/ANO-L-VTZP included the CASPT2-calculated electronic energy and the 298 K thermal correction calculated at  $\omega$ B97XD/6-31+G(d,p) (including scaled ZPE).

#### 5. Unimolecular kinetics modeling

Rice–Ramsperger–Kassel–Marcus (RRKM) theory<sup>94</sup> densities of states (DOS) and rate constants were calculated using the direct count algorithm<sup>95</sup> available in the RRKM code of Zhu and Hase<sup>96</sup> and the scaled frequencies, energies, and moments of inertia of the complexes and TSs determined from the CASPT2/ANO-L-VTZP//  $\omega$ B97XD/6-31+G(d,p) calculations. For reactions involving no reverse barrier, orbit transition states<sup>97</sup> were assumed. The orbital angular momentum *L* for the reaction was determined from the collision cross section ( $\sigma_{collision}$ ), i.e.,  $L = \mu \cdot v_{rel} \cdot \sqrt{\sigma_{collision}/\pi}$ , where  $\mu$  and  $v_{rel}$  are the reduced mass and relative velocity for collision partners, respectively.

### **III. EXPERIMENTAL RESULTS**

As previously reported,<sup>22,98</sup> the lowest-energy structures of 9HG<sup>•+</sup> and 9MG<sup>•+</sup> adopt an O6-keto-conformation, and each had an overwhelmingly dominant population in the ion beam. Note that the O6-keto-N7H isomer of the guanine radical cation (abbreviated as 7HG<sup> $\bullet$ +</sup>) lies at an energy of 0.11 eV higher than 9HG<sup> $\bullet$ +</sup>. We also note that the radical cations were generated during ion transfer within the source chamber under relatively high background gas pressure (1.6 Torr, predominantly air with a small fraction of solvent vapor). There was a possibility that radical cations underwent solvent-assisted isomerization.99 Despite this, the majority of 9HG<sup>•+</sup> and 9MG<sup>•+</sup> were expected to retain an O6-keto-N9H or O6-keto-N9-CH3 structure due to thermalization within the hexapole, where any higher-energy isomers would relax to equilibrium populations at 310 K. In addition, our CID experiments of various base pairs containing 9MG<sup>•+</sup> confirmed that only conformers containing an O6-keto-N9-CH3 structure match the measured dissociation threshold energies.<sup>100,</sup>



**FIG. 1.** Product ion cross section for the reaction of  $9HG^{\bullet+} + {}^{\bullet}NO$  as a function of kinetic energy in the center-of-mass frame ( $E_{CM}$ , bottom axis) and laboratory frame ( $E_{lab}$ , top axis). The insets show the ChemDraw for  $9HG^{\bullet+}$  and product ion mass spectrum recorded at  $E_{CM} = 0.05 \text{ eV}$ .

Scheme 1 illustrates the structures of reactants along with their NBO charge and spin density distributions. Both  $9HG^{\bullet+}$  and  $9MG^{\bullet+}$  delocalize the unpaired electron spin mainly among the N3, C5, and C8 positions and the positive charge among C2, C4, and C8. Consequently, it seems reasonable to infer that the N9-methyl substitution would not have a significant effect on the structures for guanine nitrosation.

#### A. Reaction of <sup>•</sup>NO with 9HG<sup>•+</sup>

We reported previously that the collisions of  $9HG^{\bullet+}$  (and  $9MG^{\bullet+}$ ) with  ${}^{\bullet}NO$  produced a charge-transfer product pair of 9HG (9MG) +  $NO^+$  at  $E_{CM} \ge 1.5 \text{ eV.}^{27}$  The present experiment thus specifically focused on measuring the formation of nitrosation complexes at low energies. Figure 1 depicts the reaction product ion cross section for  $9HG^{\bullet+}$  (m/z 151) +  ${}^{\bullet}NO$  as a function of kinetic energy. Error bars were derived from three sets of measurements. Product ions were detected at m/z 181, which corresponds to a covalent [NO-9HG]<sup>+</sup> adduct. The product cross section reaches a maximum at the lowest experimental kinetic energy. No complex ions



**SCHEME 1.** Structures of 9HG<sup> $\bullet+$ </sup>, 9MG<sup> $\bullet+$ </sup>, and  $\bullet$ NO with atomic numbering schemes. Spin (contour plots) and atomic charge (numbers) were calculated using NBO 6.0 at the  $\omega$ B97XD/6-31+G(d,p) level of theory.

were detected above  $E_{\rm CM}$  of 0.5 eV. This kinetic energy dependence indicates an exothermic reaction without an activation barrier above the reactants.

#### B. Reaction of \*NO with 9MG\*+

 $9MG^{\bullet+}$  was chosen as a model compound for dGuo<sup> $\bullet+$ </sup> with the bulky ribose sugar substituted by a methyl group. Figure 2(a) presents the nitrosation product cross section for  $9MG^{\bullet+}$  (*m*/z 165), measured as a function of  $E_{\rm CM}$  from 0.05 to 1.0 eV. The reaction efficiency, estimated by  $\sigma_{\rm reaction}/\sigma_{\rm collision}$  (where  $\sigma_{\rm collision}$  represents the ion-induced dipole capture cross section<sup>102</sup> or hard-sphere collision cross section calculated using the orientation-averaged projected area method of the IMoS program,<sup>103,104</sup> whichever is larger), is ~0.01% at  $E_{\rm CM} = 0.05$  eV, quickly decreasing to 0.003% at  $E_{\rm CM} = 0.1$  eV. For comparison, the reaction efficiency for  $9HG^{\bullet+}$  is 0.1% at  $E_{\rm CM} = 0.05$  eV, decreasing to 0.05% at 0.1 eV.

An intriguing finding in the nitrosation of  $9MG^{\bullet+}$  is the emergence of a distinct, endothermic product channel. The cross section for this product channel starts to rise at  $E_{CM} = 0.2$  eV, reaches its maximum at 0.22 eV, and then gradually decreases, becoming negligible at energies above 0.8 eV. Notably, this specific product channel is not prominently observed in the reaction of  $9HG^{\bullet+} + {}^{\bullet}NO$  in Fig. 1—there is a possibility that this channel has been overshadowed by the relatively large exothermic product channel for  $9HG^{\bullet+}$ .

Despite the similar kinetic energy dependence, the reaction efficiency of 9MG<sup>•+</sup> with <sup>•</sup>NO appears to be an order of magnitude lower than that of 9HG<sup>•+</sup>. It is plausible that, due to insufficient energy relaxation in the rarefied gas-phase environment, a fraction of the adducts formed through an exothermic reaction might have decomposed into the starting reactants before reaching the mass spectrometer detector (the ion time-of-flight time was 100–500  $\mu$ s within the mass spectrometer). A similar decomposition scenario was observed in the <sup>1</sup>O<sub>2</sub> addition reactions of 9MG<sup>•+</sup> and 8-bromo-9HG<sup>•+</sup>.<sup>22,24</sup>

No proton-transfer (PT) or hydrogen-transfer (HT) was observed between  $9HG^{\bullet+}/9MG^{\bullet+}$  and  $^{\bullet}NO$  within the experimental energy range. According to  $\omega B97XD/6-31+G(d,p)$  calculations, the PT  $\Delta H$  (298 K) leading to  $^{2}[9MG-H_{N1}]^{\bullet}$  + HNO<sup>+</sup> is 4.27 eV and that to  $^{2}[9MG-H_{N1}]^{\bullet}$  + HON<sup>+</sup> is 4.88 eV. The HT  $\Delta H$  (298 K)

leading to  $[9MG-H_{N2}]^+$  + HNO is 2.08 eV and that to  $[9MG-H_{N2}]^+$  + <sup>3</sup>HON is 2.92 eV.

#### C. Reaction of \*NO with 9MG\*++H<sub>2</sub>O

Since the extremely low reaction efficiency observed for  $9MG^{\bullet+}$  with  $\bullet NO$  might be possibly attributed to the decomposition of nascent nitrosation adducts carrying the reaction's exothermicity, we adopted a workaround by using the monohydrated  $9MG^{\bullet+} \cdot H_2O$  as reactant ions for collisions with  $\bullet NO$ . In this modified reaction system, the heat release from nitrosation, which could otherwise lead to the decomposition of  $[NO-9MG]^+$  in dry conditions, instead facilitated the elimination of the water ligand along with the release of product kinetic energy.<sup>22,24,105</sup> The anticipated outcome was verified in the experiment.

As shown in Fig. 2(b), product ions for  $9MG^{\bullet+} H_2O$  (*m/z* 183) + •NO were detected at *m/z* = 195, which corresponds to the liberation of a water ligand from the •NO-adduct of  $9MG^{\bullet+} H_2O$ . The reaction efficiency for  $9MG^{\bullet+} H_2O$  is 0.54% at  $E_{CM} = 0.05$  eV, decreasing to 0.5% at 0.1 eV. Thus, the complex formation of  $9MG^{\bullet+} H_2O +$ •NO increases by two orders of magnitude than that of dry  $9MG^{\bullet+} +$ •NO.

The detection of the exothermic reaction of 9MG<sup>•+</sup>·H<sub>2</sub>O + •NO → [NO-9MG]<sup>+</sup> + H<sub>2</sub>O has imposed a constraint on the exothermicity of the reaction—it must exceed the water-binding energy of 9MG<sup>•+</sup>·H<sub>2</sub>O. The water-binding energy depends on the hydration structure (see Fig. S1 of the supplementary material). It is 0.48 eV when water is H-bonded to a N2–H in 9MG (i.e., referred to as W2), 0.58 eV when water is double H-bonded to N1–H and O6 (i.e., W16), and 0.68 eV when water is double H-bonded to N1–H and N2–H (i.e., W12).<sup>98,106</sup> According to direct dynamics trajectory simulations for 9MG<sup>•+</sup> hydration, <sup>98,106</sup> the W12 binding motif represents the most probable structure for 9MG<sup>•+</sup>·H<sub>2</sub>O. The implication is that the cross section in Fig. 2(b) should be mostly attributed to the reaction of the W12 structure of 9MG<sup>•+</sup>·H<sub>2</sub>O. It follows that the nitrosation enthalpy of 9MG<sup>•+</sup> should exceed 0.68 eV.

Since the exothermic nitrosation product for  $9MG^{\bullet+}H_2O$  +  $^{\bullet}NO$  remains dominant up to  $E_{CM}$  = 0.5 eV, the endothermic product, which would be otherwise observable between 0.2 and 0.5 eV, becomes obscured as reactant hydration could not enhance an endothermic reaction.



FIG. 2. Product ion cross section for the reactions of  $^{\bullet}$ NO with (a) 9MG $^{\bullet+}$  and (b) 9MG $^{\bullet+}$ ·H<sub>2</sub>O as a function of kinetic energy in the center-of-mass frame ( $E_{CM}$ , bottom axis) and laboratory frame ( $E_{lab}$ , top axis). The insets show the product ion mass spectra recorded at  $E_{CM} = 0.05 \text{ eV}$ .

#### **IV. THEORETICAL RESULTS**

### A. Spin dynamics

As 9MG<sup>•+</sup> more closely resembles dGuo<sup>•+</sup> in terms of structure and distributions of spin and charge than 9HG<sup>•+</sup>,<sup>22,98</sup> 9MG<sup>•+</sup> was chosen as a prototypical system for computational modeling. Figure 3 presents a relaxed potential energy surface (PES) for 9MG<sup>•+</sup> + •NO. Since the N-terminal of •NO holds the radical electron and thus has greater reactivity toward 9MG<sup>•+</sup> than the O-terminal, PES was calculated along the distance (r[9MG-N]) between the center-of-mass of 9MG and the N-terminal of •NO. The PES scans from r[9MG-N] = 15 to 1.8 Å (at a step size of 0.01 Å from 1.8 to 5–6.5 and 0.1 Å afterward) using the  $\omega$ B97XD/6-31+G(d,p) method. The PES includes three electron configurations: an open-shell singlet <sup>1,OS</sup>[9MG<sup>•+</sup>(†)···(↓)•NO] (black color PES in Fig. 3), a closed-shell singlet <sup>1,CS</sup>[9MG(†↓)···NO<sup>+</sup>] (red), and a triplet <sup>3</sup>[9MG<sup>•+</sup>(†)···(†)•NO] (blue). The surface for <sup>1,OS</sup>[9MG<sup>•+</sup>(†)···(↓)•NO] was corrected for spin contamination using approximate spin projection. Spin dynamics for the radical-radical combination varies at different stages of the reaction as described below:

(1) At the beginning of collisions, the radicals assemble into either <sup>1,OS</sup>[9MG<sup>•+</sup>(↑)···(↓)<sup>•</sup>NO] or <sup>3</sup>[9MG<sup>•+</sup>(↑)···(↑)<sup>•</sup>NO]. The two electronic states are energetically degenerate at long separation until r[9MG-N] reaches 5 Å from which the open-shell singlet becomes lower in energy than the triplet. Interaction in the ion-molecule encounter (mostly ion-dipole interaction) starts to become significant at the shallow PES minimum located at [9MG-N] = 5.8 Å. The structure of this encounter is shown in the supplementary material, where the <sup>•</sup>NO is approaching the 9MG<sup>•+</sup> in a parallel direction. The electrostatic binding energy of this encounter is 0.11 eV. The encounter rearranges the structure at a TS-like point with a forward barrier of 0.03 eV. The TS-like structure is also shown in the supplementary material, where the  $^{\circ}NO$  moiety moves over  $9MG^{^{\circ+}}$  to prepare for the formation of a stable [NO-9MG]<sup>+</sup> complex.

- (2) Singlet reactions: The TS-like point marks the starting point where the paths of <sup>1,OS</sup>[9MG<sup>•+</sup>(↑)···(↓)•NO] and <sup>3</sup>[9MG<sup>•+</sup>(↑)···(↑)•NO] diverge. Depending on reaction orientation, <sup>1,OS</sup>[9MG<sup>•+</sup>(↑)···(↓)•NO] may form an electrostatically bonded complex at r[9MG-N] = 3.5 Å, referred to as <sup>1,OS</sup>precursor (shown in the bottom inset of Fig. 3). The <sup>1,OS</sup>precursor may converge to a singlet closed-shell <sup>1,CS</sup>[8-NO-9MG]<sup>+</sup>, wherein the positive charge is distributed over the entire complex, as manifested by the electrostatic potential (ESP)-fitted charge distribution in the middle inset of Fig. 3. Alternatively, <sup>1,OS</sup>[9MG<sup>•+</sup>(↑)···(↓)•NO] may converge to a singlet closed-shell <sup>1,CS</sup>[7-NO-9MG]<sup>+</sup> at [9MG-N] = 3.7 Å, and the latter may interconvert to <sup>1,CS</sup>[5-NO-9MG]<sup>+</sup>. The <sup>1,CS</sup>[5-NO-9MG]<sup>+</sup>, <sup>1,CS</sup>[7-NO-9MG]<sup>+</sup>, and <sup>1,CS</sup>[8-NO-9MG]<sup>+</sup> act as hubs for interconversion to other singlet nitrosation products to be described below.
- (3) Triplet reactions: The nitrosation reaction may also initiate from <sup>3</sup>[9MG<sup>•+</sup>(↑)···(↑)<sup>•</sup>NO]. The triplet first forms an electrostatically bonded complex at r[9MG-N] = 3 Å, referred to as <sup>3</sup>precursor. The spin sites of <sup>3</sup>precursor are characterized in the upper left inset of Fig. 3. The charge of <sup>3</sup>precursor, however, remains at 9MG. Similar to <sup>1,OS</sup>precursor, the <sup>3</sup>precursor allows for repeated encounters between reactants; consequently, the system proceeds to form various triplet nitrosation structures to be described below. As reported in our previous study,<sup>27</sup> <sup>3</sup>[9MG<sup>•+</sup>(↑)··· (↑)<sup>•</sup>NO] may undergo spin-orbit charge transfer at [9MG-N] = 5 Å and become correlated with the charge transfer (CT) asymptote 9MG + NO<sup>+</sup>.







**SCHEME 2.** Probable nitrosation pathways and products of 9MG<sup>++</sup> in the closed-shell singlet state.

#### B. Nitrosation pathways and energetics

Guided by spin dynamics, we mapped out possible nitrosation pathways using  $\omega$ B97XD/6-31+G(d,p). The results are illustrated using flow charts in Schemes 2–4. Cartesian coordinates for reaction structures are provided in the supplementary material. Scheme 2 presents three distinctive types of NO-additions leading from <sup>1,OS</sup> precursor to a closed-shell singlet product. The N-terminal addition of  $^{\circ}$ NO (illustrated in black) at the N7 and C8 positions produces  $^{1,CS}$ [7-NO-9MG]<sup>+</sup> and  $^{1,CS}$ [8-NO-9MG]<sup>+</sup>, respectively.  $^{1,CS}$ [7-NO-9MG]<sup>+</sup> may convert to  $^{1,CS}$ [5-NO-9MG]<sup>+</sup> and subsequently to  $^{1,CS}$ [3-NO-9MG]<sup>+</sup>, and  $^{1,CS}$ [8-NO-9MG]<sup>+</sup> may convert to  $^{1,CS}$ [7H-8-NO-9MG]<sup>+</sup> via intra-molecular proton transfer. The N-terminal addition also produces  $^{1}$ [2-NHNO-3H-9MG]<sup>+</sup> accompanied by a proton transfer from N2 to N3. The O-terminal addition of  $^{\circ}$ NO (illustrated in blue) occurs only at C8, leading to



SCHEME 3. Probable nitrosation pathways and products of 9MG<sup>++</sup> in the open-shell singlet state, wherein unpaired electrons are indicated in the ChemDraw structures and formulas.

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SCHEME 4. Probable nitrosation pathways and products of  $9MG^{\bullet+}$  in the triplet state.

the formation of  ${}^{1}$ [7H-8-ON-9MG]<sup>+</sup> with a simultaneous hydrogen transfer from C8–H to N7. Cycloadditions (illustrated in red) were identified for  ${}^{1,CS}$ [4,5-NO-9MG]<sup>+</sup> and  ${}^{1,CS}$ [7,8-NO-9MG]<sup>+</sup>. While both cycloadditions were mediated by a single TS, the reactions may be characterized as asynchronous concerted: although the cycloaddition involves no intermediate (i.e., concerted), it comprises two "events" that do not occur simultaneously (i.e., asynchronous). For example, the addition of the N-terminal to C4 precedes the addition of the O-terminal to C5 in the formation of  ${}^{1,CS}$ [4,5-NO-9MG]<sup>+</sup>.

Compared to the closed-shell singlet, the open-shell singlet presents fewer pathways, as depicted in Scheme 3. An O-terminal addition pathway leads to a  $^{1,OS}[8-ON-9MG]^+$  biradical, and the latter may interconvert to a  $^{1,OS}[7H-8-ON-9MG]^+$  biradical via a

hydrogen transfer. In addition, four cycloadditions were identified. Only  ${}^{1,OS}[7,8-O\dot{N}-9M\dot{G}]^+$  forms via an asynchronous concerted route starting with the 8-NO addition in  ${}^{1,OS}$ precursor, whereas  ${}^{1,OS}[2,4-\dot{N}O-9M\dot{G}]^+$ ,  ${}^{1,OS}[5,8-\dot{N}O-9M\dot{G}]^+$ , and  ${}^{1,OS}[5,8-O\dot{N}-9M\dot{G}]^+$ are all synchronous cycloadditions.

Of the three electronic states, the triplet presents the most versatile pathways. The top frame of Scheme 4 presents various N- vs O-terminal additions of  $^{\circ}$ NO to 9MG $^{\circ+}$ . The bottom frame presents concerted and sequential cycloadditions: the formation of  $^{3}$ [4,5-ON-9MG]<sup>+</sup>,  $^{3}$ [5,8-ON-9MG]<sup>+</sup>, and  $^{3}$ [7,8-ON-9MG]<sup>+</sup> is asynchronous, the formation of  $^{3}$ [2,4-NO-9MG]<sup>+</sup> and  $^{3}$ [7,8-NO-9MG]<sup>+</sup> is synchronous, and the formation of  $^{3}$ [5,8-NO-9MG]<sup>+</sup> can be either.

It is interesting to compare spin locations in the open-shell singlet vs triplet products. In the singlet O-terminal adducts, two unpaired electrons are each located at N and O of NO, whereas in the singlet biradical cycloadducts, two unpaired electrons each remain at  $9MG^{\bullet+}$  and  $\bullet$ NO, respectively. In the triplet N-terminal adducts, the unpaired electron of  $9MG^{\bullet+}$  shifts to the O of NO. In the triplet O-terminal adducts, the unpaired electrons being located at the N-terminal. Finally, in the triplet cycloadducts, two unpaired electrons are each retained at  $9MG^{\bullet+}$  and  $\bullet$ NO. Regardless of product electronic states, the charge largely remains at 9MG.

Tables S2 and S3 of the supplementary material compare 298 K reaction enthalpies calculated at the DFT, DLPNO-CCSD(T)/ aug-cc-pVTZ and CASPT2/ANO-L-VTZP levels, and  $\langle \hat{S}^2 \rangle$  and T1 diagnostic for reaction structures. All calculations used the  $\omega$ B97XD/6-31+G(d,p)-optimized reaction structures. All closed-shell singlet structures have a  $\langle \hat{S}^2 \rangle$  value of 0.00 and T1  $\leq$  0.019, and all triplet structures have a  $\langle \hat{S}^2 \rangle$  of 2.00 and T1  $\leq$  0.023, which indicates at most minor spin contamination for these species. Mean-while, biradical structures have a  $\langle \hat{S}^2 \rangle$  of 0.83–1.01, which represents a diradical character in a range of 59%–100% [using the diradical index<sup>62</sup>  $n_{DC} = \left(1 - \sqrt{\left(1 - \langle \hat{S}^2 \rangle\right)} \times 100\%\right]$ .

Compared to the CASPT2 results, the  $\omega$ B97XD level has underestimated the reaction energies by 0.2–0.4 eV, whereas the DLPNO-CCSD(T) level has overestimated the energies by 0.4–0.6 eV. The transition state TS-<sup>1,CS</sup>[4,5-NO-9MG]<sup>+</sup> for 4,5-cycloaddition shifts to an energy below the product at the CASPT2 level, indicating that this cycloaddition is barrierless. All theories have predicted that exothermic nitrosation pathways occur only in the closed-shell singlet, as the latter represents the ground electronic state; and all open-shell singlet and triplet pathways are either endothermic or have activation barriers above reactants, indicating that these electronic states are not thermodynamically favorable as expected from the Pauli exclusion principle.

#### V. DISCUSSION

# A. <sup>1,CS</sup>[7-NO-9MC]<sup>+</sup> represents the only exothermic nitrosation product

Figure 4 compiles the potential energies for the formation and interconversions of all nitrosation reactions mediated by the open- and closed-shell singlets. As aforementioned, the H-binding energy of  $9MG^{\bullet+}H_2O$  ranges from 0.48 to 0.68 eV, of which 0.68 eV



**FIG. 4.** Potential energy diagram for the reaction of  $9MG^{\bullet+} + {}^{\bullet}NO$  in the open- and closed-shell singlet states. Enthalpies (298 K) were calculated at CASPT2/ANO-L-VTZP// $\omega$ B97XD/6-31+G(d,p) (numbers in parentheses), except for open-shell singlet species, which were calculated at AP-DFT (indicated with the superscript AP). The shaded areas indicate the minimum and the most probable exothermicity range for the experimentally measured product.

is the most probable. This has imposed a constraint on the minimum and the most probable exothermicity range for experimentally measured products, as marked in the Y-axis of Fig. 4. Accordingly, the most probable product ions correspond to <sup>1,CS</sup>[7-NO-9MG]<sup>+</sup>. This agrees with the relaxed PES in Fig. 3, where <sup>1,CS</sup>[7-NO-9MG]<sup>+</sup> represents the first covalent structure as reactants approach each other.

other. <sup>1,CS</sup>[7-NO-9MG]<sup>+</sup> may continue conversion to <sup>1,CS</sup>[5-NO-9MG]<sup>+</sup> and <sup>1,CS</sup>[3-NO-9MG]<sup>+</sup> with barriers below the reactants. To determine relative contributions of different adduct structures, RRKM kinetics modeling was utilized. The time scale for the interconversion between <sup>1,CS</sup>[7-NO-9MG]<sup>+</sup> and <sup>1,CS</sup>[5-NO-9MG]<sup>+</sup> is less than 0.1 ns at the  $E_{\rm CM}$  range of 0.05–0.2 eV, and that for the interconversion between <sup>1,CS</sup>[5-NO-9MG]<sup>+</sup> and <sup>1,CS</sup>[3-NO-9MG]<sup>+</sup> is less than 1 ns at the same energy range. As a consequence, an equilibrium between <sup>1,CS</sup>[3-NO-9MG]<sup>+</sup>, <sup>1,CS</sup>[5-NO-9MG]<sup>+</sup>, and <sup>1,CS</sup>[7-NO-9MG]<sup>+</sup> could be anticipated within the product ion time-of-flight. We could then use the ratio of DOS in these complexes to estimate their relative populations and determine which complexes are important. The DOS is energy-dependent. It turns out that, at  $E_{\text{CM}} \leq 0.5 \text{ eV}$ , <sup>1,CS</sup>[7-NO-9MG]<sup>+</sup> is predominant (>99%); meanwhile, neither <sup>1,CS</sup>[3-NO-9MG]<sup>+</sup> (0%) nor <sup>1,CS</sup>[5-NO-9MG]<sup>+</sup> (<0.1%) makes a meaningful contribution. Furthermore, the RRKM kinetics modeling predicted that <sup>1,CS</sup>[7-NO-9MG]<sup>+</sup> has a lifetime of  $10^2 - 10^3 \mu s$  at  $E_{\rm CM}$  below 0.2 eV but decreased to less than  $\mu s$  at  $E_{\rm CM}$ above 0.5 eV. Therefore, while <sup>1,CS</sup>[7-NO-9MG]<sup>+</sup> accounted for the product ions at low energies, at high energies, it decomposed into

starting reactants before reaching the mass spectrometer detector. Such kinetics rationalizes the quick decline of exothermic product ions with increasing kinetic energy in Figs. 1 and 2.

Folkes and O'Neill<sup>40</sup> proposed the formation of <sup>1,CS</sup>[2-NHNO-G] during the modification of DNA by <sup>•</sup>NO under ionizing radiation, which corresponds to <sup>1,CS</sup>[2-NHNO-3H-9MG]<sup>+</sup> in the present reaction system. Despite this product being exothermic by 0.64 eV, it requires an activation barrier of 1.25 eV, making it less likely to occur in reality. Similarly, the product ion <sup>1,CS</sup>[7H-8-NO-9MG]<sup>+</sup>, although the most stable, requires a 1.15 eV activation energy above the reactants and is therefore insignificant.

Figure 4 also presents the two open-shell singlet products:  $^{1,OS}[8-\dot{ON}-9MG]^+$  and  $^{1,OS}[7H-8-\dot{ON}-9MG]^+$ , of which neither is thermodynamically favorable. Note that none of the biradical cyclo-adducts were included in this figure for two reasons: biradical cyclo-adducts demand high energetics for formation, and in fact, all these structures converged to different (non-cyclic) structures in restricted closed-shell calculations—implying their inherent instability.<sup>79</sup>

# B. Origin of the minor, endothermic nitrosation product

We now turn to the identification of the endothermic nitrosation product detected at  $E_{\rm CM} = 0.18-0.22$  eV for dry 9MG<sup>•+</sup> + •NO [Fig. 2(a)]. It is evident in Fig. 4 that the singlet product channels are either exothermic with no activation barrier above the

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FIG. 5. Potential energetic diagram for the reaction of 9MG<sup>++</sup> + •NO in the triplet state. Enthalpies (298 K) were calculated at CASPT2/ANO-L-VTZP//<sub>w</sub>B97XD/6-31+G(d,p) and DLPNO-CCSD(T)/aug-cc-pVTZ//<sub>w</sub>B97XD/6-31+G(d,p) (numbers in parentheses).

reactants (such as  ${}^{1,CS}$ [3-NO-9MG]<sup>+</sup>,  ${}^{1,CS}$ [5-NO-9MG]<sup>+</sup>,  ${}^{1,CS}$ [7-NO-9MG]<sup>+</sup>, and  ${}^{1,CS}$ [8-NO-9MG]<sup>+</sup>) or have product endothermicity or activation barrier of at least 1.15 eV (such as the closed-shell  ${}^{1,CS}$ [2-NHNO-3H-9MG]<sup>+</sup>,  ${}^{1,CS}$ [7H-8-NO-9MG]<sup>+</sup>,  ${}^{1,CS}$ [7H-8-ON-9MG]<sup>+</sup>,  ${}^{1,CS}$ [7H-8-NO-9MG]<sup>+</sup>,  ${}^{1,CS}$ [4,5-NO-9MG]<sup>+</sup>, and  ${}^{1,CS}$ [7,8-NO-9MG]<sup>+</sup> and the open-shell  ${}^{1,OS}$ [8- $\dot{ON}$ -9MG]<sup>+</sup> and  ${}^{1,OS}$ [7H-8 $-\dot{ON}$ -9MG]<sup>+</sup>). Therefore, none of these would be relevant to a moderate endothermic reaction.

It follows that the detected endothermic product may only arise from a triplet state. Figure 5 outlines all triplet addition pathways initiated at <sup>3</sup>precursor. All reactions present an activation barrier in the range of 0.73–3.38 eV, except for <sup>3</sup>[8-NO-9MG]<sup>+</sup>, which has a reaction enthalpy of 0.45 eV and an activation barrier of 0.62 eV. Therefore, <sup>3</sup>[8-NO-9MG]<sup>+</sup> represents the most likely endothermic product structure in the experiment. The fact that this channel was observed at the range of  $E_{\rm CM}$  (0.18–0.22 eV) below its calculated barrier was due to reactant internal energy and translational kinetic energy spread and Doppler broadening in ion–molecule collisions. The falloff of the <sup>3</sup>[8-NO-9MG]<sup>+</sup> cross section at high energies is due to the combination of the decreasing complex intermediacy and the emerging of the spin–orbit charge transfer reaction.<sup>27</sup>

#### C. Low yield for radical-radical combination

The low nitrosation product yield for 9HG<sup>•+</sup> and 9MG<sup>•+</sup>, even in the presence of hydration, warrants rationalization. The nitrosative reaction, or, in a general sense, combination of radicals, is spin selective.<sup>107</sup> The pair of guanine radical cation and <sup>•</sup>NO can form one anti-symmetric, singlet spin eigenfunction  $[\alpha(1)\beta(2) - \beta(1)\alpha(2)]/\sqrt{2}$ , and three symmetric, triplet spin eigenfunctions  $\alpha(1)\alpha(2)$ ,  $\beta(1)\beta(2)$ , and  $[\alpha(1)\beta(2) + \beta(1)\alpha(2)]/\sqrt{2}$ . Therefore, the partition between singlet and triplet is 0.25:0.75.

Nitrosation products may form from radical-radical combination to the lowest-energy triplet state. However, all triplet pathways are endothermic and face competition from CT as well as nonreactive collisions (i.e., escaping combination and decaying back to reactants), with these factors being cumulative. Of course, the triplet may interconvert to the singlet but must be driven by spin-orbit charge transfer-induced ISC;<sup>27</sup> consequently, the final state must correlate with CT products 9MG + NO<sup>+</sup>. Meanwhile, the charge in all nitrosation products remains predominantly localized at the 9MG moiety, thereby mitigating the driving force for the envisioned triplet–singlet transition (unless a spin flip is induced additionally by a heavy-atom substitute<sup>108</sup> and/or paramagnetic relaxation<sup>109</sup>). That implies that, of all ion–molecule collisions, 75%, which were born in the triplet, could be largely discounted for nitrosation.

The combination of the radicals into a singlet state may occur without any activation barrier (Fig. 4). It follows that a reaction yield of up to 25% could be expected at the lowest collision energy. This statement, however, was true only if all starting open-shell singlet ion-molecule encounters converged to closed-shell singlet and all singlet products were captured by the mass spectrometer. Unfortunately, the less-than-100% interconversion between openand closed-shell singlets (due to reaction orientation dependence) as well as the fast decay of nascent, singlet nitrosation products has in turn invalidated this condition. The monohydration of reactant ions had helped a fraction of the product ions survive, but the effect was not substantial enough to reverse the prevailing trend toward decay.

#### **VI. CONCLUSIONS**

Following our previous study on the spin-orbit charge transfer of •NO with 9HG•+ and 9MG•+, we carried out a combined guided-ion beam mass spectrometry and electronic structure investigation of the nitrosation reactions for these two reaction systems. 9HG<sup>•+</sup> and 9MG<sup>•+</sup> present similar reaction outcomes, and they are referred to collectively as  $G^{\bullet+}$ . Both open-shell singlet  $^{1,OS}[G^{\bullet+}(\uparrow)\cdots(\downarrow)^{\bullet}NO]$  and triplet  $^{3}[G^{\bullet+}(\uparrow)\cdots(\uparrow)^{\bullet}NO]$  formed reaction precursors in a ratio of 1:3. The  ${}^{3}[G^{\bullet+}(\uparrow)\cdots(\uparrow)^{\bullet}NO]$  precursor is responsible for the formation of a minor amount of endothermic product ions <sup>3</sup>[8-NO-G]<sup>+</sup> at relatively low collision energies and for the spin-orbit charge transfer at high energies. Meanwhile, the  ${}^{1,OS}[G^{\bullet+}(\uparrow)\cdots(\downarrow)^{\bullet}NO]$  precursor leads to the ultimate formation of the major, exothermic product ions <sup>1,CS</sup>[7-NO-G]<sup>+</sup> without any activation barriers above the reactants. Combined with our previous work, we have delineated the roles and evolution of the two charge states and three spin configurations and caught a glimpse of fertile radical-radical reaction dynamics. The findings have demonstrated synergistic oxidative nucleobase damage in the presence of ionization, one-electron oxidation, and nitrosation.

#### SUPPLEMENTARY MATERIAL

The supplementary material encompasses Cartesian coordinates, energetics,  $(\hat{S}^2)$ , and T1 diagnostic for all reaction structures.

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#### AUTHOR DECLARATIONS

#### **Conflict of Interest**

The authors have no conflicts to disclose.

#### **Author Contributions**

Jonathan Benny: Data curation (equal); Formal analysis (equal); Funding acquisition (supporting); Investigation (equal); Methodology (equal); Software (equal); Validation (equal); Writing – original draft (equal); Writing – review & editing (equal). Toru Saito: Data curation (equal); Formal analysis (equal); Funding acquisition (supporting); Investigation (supporting); Methodology (equal); Software (equal); Validation (equal); Writing – original draft (supporting); Writing – review & editing (supporting). Jianbo Liu: Conceptualization (lead); Data curation (lead); Formal analysis (lead); Funding acquisition (lead); Investigation (lead); Methodology (lead); Project administration (lead); Resources (lead); Software (lead); Supervision (lead); Validation (lead); Writing – original draft (lead); Writing – review & editing (lead).

#### DATA AVAILABILITY

The data that support the findings of this study are available within the article and its supplementary material.

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