Photo-Induced Reactions between Glyoxal and Hydroxylamine in Cryogenic Matrices

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Abstract: In this paper, the photochemistry of glyoxal–hydroxylamine (Gly–HA) complexes is studied using FTIR matrix isolation spectroscopy and ab initio calculations. The irradiation of the Gly–HA complexes with the filtered output of a mercury lamp (λ > 370 nm) leads to their photoconversion to hydroxyketene–hydroxylamine complexes and the formation of hydroxy(hydroxyamino)acetaldehyde with a hemiaminal structure. The first product is the result of a double hydrogen exchange reaction between the aldehyde group of Gly and the amino or hydroxyl group of HA. The second product is formed as a result of the addition of the nitrogen atom of HA to the carbon atom of one aldehyde group of Gly, followed by the migration of the hydrogen atom from the amino group of hydroxylamine to the oxygen atom of the carbonyl group of glyoxal. The identification of the products is confirmed by deuterium substitution and by MP2 calculations of the structures and vibrational spectra of the identified species.

Keywords: matrix isolation; infrared spectroscopy; photochemistry; carbonyl; hydroxyketene; hemiaminal

1. Introduction

Glyoxal, HCOHCO (Gly), the simplest α-dicarbonyl, plays a significant role in atmospheric photochemistry. It is formed in the atmosphere as a product of the oxidation of volatile organic compounds (VOCs) [1], and the direct photolysis of glyoxal is its main removal route from the atmosphere [2,3]. The gas-phase photodissociation of glyoxal involves several fragmentation channels, whose relative contributions depend on the excitation wavelength and other factors such as the occurrence of intermolecular collisions [4–10]. Moreover, glyoxal is also absorbed by cloud droplets, where it participates in complex photochemical transformations [11–15]. Our group has reported systematic study of the photochemical behavior of molecular complexes of glyoxal with atmospherically relevant molecules such as water, methanol and hydrogen peroxide isolated in cryogenic matrices [16–18]. For the glyoxal–methanol complex, a very interesting photochemical behavior was observed, namely, under irradiation in the visible range, glyoxal complexed with methanol isomerized to the rather exotic hydroxyketene molecule, H(OH)C≡C=O (Equation (1)) [18].

\[ \text{CHOCHO} \cdots \text{MeOH} \rightarrow \text{H(OH)C≡C=O} \cdots \text{MeOH} \] (1)

This reaction only occurred for the complex with methanol. The complex of glyoxal with H₂O₂ shows a different photochemical behavior, in which the hydrogen peroxide molecule undergoes addition to glyoxal [17]. In contrast, the complex of glyoxal with water does not undergo a photochemical isomerization reaction [16]. The complex of methanol with glyoxal derivative, methylglyoxal, shows similar behavior to the glyoxal–methanol complex [19]. Irradiation of this complex leads to the formation of methylhydroxyketene.
An experimental study in argon matrices shed some light on the isomerization reaction of glyoxal [18], and the mechanism of this reaction has also been extensively studied using theoretical methods [20,21]. Irradiation of the complex with deuterium-enriched methanol, CH$_3$OD, produced deuterium-enriched ketene, H(OD)C=C=O. This experimental result indicates that the isomerization occurs via a double hydrogen exchange reaction between glyoxal and methanol (or hydrogen and deuterium in the case of CH$_3$OD). Recent advanced theoretical studies on the photoisomerization reaction in the CHOCHO···MeOH complex have revealed that it follows a complicated mechanism triggered by the excitation of the complex to the lowest single excited electronic state. The mechanism involves two complementary pathways. One of them takes place entirely in the singlet manifold, and the other also involves the lowest triplet state (T$_1$) [20,21].

Here, we present the results of a study of the photochemical behavior, under visible irradiation, of the glyoxal complex with hydroxylamine, CHOCHO···NH$_2$OH (Gly–HA). The structures and energetics of CHOCHO···NH$_2$OH complexes embedded in solid argon and nitrogen have recently been reported [22]. Studies of the CHOCHO···NH$_2$OH complex provide new, interesting data on hydrogen exchange reactions in glyoxal complexes. They also indicate the existence of an additional reaction channel by which hydroxylamine undergoes addition to glyoxal, forming a product of hemiaminal structure (hydroxy(hydroxyamino)acetaldehyde). In this way, this unique product could be detected and characterized spectroscopically.

Hemiaminals are important intermediate species in the formation of oximes, which are created in a reaction between a carbonyl compound (aldehyde or ketone) and a nucleophile (alkoxyamine or hydroxylamine). This reaction is stepwise: The first step involves the formation of an unstable intermediate hemiaminal, and the next step, usually catalyzed by acids or bases, proceeds via hydrolysis of the hemiaminal and oxime formation. Oxime formation is employed in numerous scientific fields, e.g., chemical biology, organic synthesis and polymer chemistry [23–25], as a versatile conjugation strategy. Studies on hemiaminals are relatively rare due to their low stability, but they can be observed under special conditions. For example, hemiaminals have been observed in solutions at low temperatures (−78, −30 °C in THF) [26] or detected with the help of host-guest chemistry as molecules trapped in deep molecular cavities [27] or in a porous coordination network [28]. They have also been identified in a solid-state reaction as an unstable compound [29]. The first stable open-chain hemiaminal, stabilized by an intramolecular hydrogen bond, was obtained in a crystalline form by condensation of di-2-pyridyl ketone with 4-cyclohexyl-3-thiosemicarbazide [30]. A recent study on a hemiaminal presents a synthesis method for aminomethanol, the simplest of hemiaminals, in low-temperature binary ices of methylamine and oxygen. This study demonstrated that aminomethanol can be formed under astrophysical-like conditions [31]. In the case of our work, the elusive hemiaminal formed in the addition reaction was stabilized by the inert environment of the cryogenic matrix.

The products that were formed under irradiation of the Gly–HA complexes embedded in cryogenic matrices were identified by FTIR spectroscopy and ab initio calculations. The results of our study are described below.

### 2. Results and Discussion

A study on the interaction of glyoxal with hydroxylamine in solid argon and nitrogen was published recently [22]. The study showed that the non-planar Gly–HA complex I$_{GH}$ was formed in the argon matrix. In the nitrogen matrix, two types of complexes were observed: a non-planar type, I$_{GH}$, and a cyclic planar type, II$_{GH}$ (see Figure 1). The cyclic planar structure of Gly–HA complex was more favorable than the non-planar one. The infrared spectra and experimental wavenumbers in comparison with the calculated ones for the CHOCHO···NH$_2$OH and CHOCHO···ND$_2$OD complexes are given in the Supplementary Materials (Figure S1 and Table S1).
The UV photochemistry of hydroxylamine has not yet been studied in matrices. Gericke et al. report that the end product of the photolysis of hydroxylamine vapor at 193 nm is $\text{H}_2 + \text{HNO}$ [32]. The main dissociation channel leads to $\text{H} + \text{H} + \text{HNO}$ with a quantum efficiency of 1.7 for hydrogen atoms. According to Luckhaus et al., the single-photon irradiation of hydroxylamine at 240 nm produces $\text{NH}_2$ and $\text{OH}$ radicals (mostly in their vibrational ground state) [33]. The photolysis of glyoxal monomers and “cage” dimers in an argon matrix leads to the formation of formaldehyde and carbon monoxide when they are excited in the $S_2 \leftarrow S_0$ absorption region ($320 \text{ nm} > \lambda > 260 \text{ nm}$) [34]. No photoproducts were observed under excitation in the $S_1 \leftarrow S_0$ absorption region ($\lambda \sim 445 \text{ nm}$). Our present study concerning the photolysis of glyoxal in argon and nitrogen matrices showed that only trace amounts of formaldehyde and carbon monoxide were formed when glyoxal was irradiated with a wavelength of $\lambda \geq 370 \text{ nm}$. Irradiation with the full output of a medium-pressure mercury lamp led to the appearance of a small concentration of carbon dioxide in addition to $\text{HCHO}$ and $\text{CO}$. The exposure of the $\text{NH}_2\text{OH}/\text{Ar}(N_2)$ matrices to the $\lambda \geq 370 \text{ nm}$ radiation led to the appearance of trace amounts of nitrogen oxide dimers [35] as a result of hydroxylamine degradation, but no HNO was detected [36].

As can be seen in Figure 2, the exposure of $\text{Gly}/\text{HA}/\text{Ar}(N_2)$ to irradiation at $\lambda \geq 370 \text{ nm}$ led to a decrease in bands due to the glyoxal–hydroxylamine complexes, and simultaneously a set of new bands appeared. The new bands diminished when the matrices were additionally irradiated with the full output of a medium-pressure mercury lamp. The bands that appeared after irradiation were separated into three groups, 1a, 1b and 2, based on their behavior in all the performed experiments. The wavenumbers of the bands assigned to groups 1a, 1b and 2 are listed in Tables 1 and 2.

### 2.1. Formation of Hydroxyketene–Hydroxylamine Complexes

The sets of bands belonging to groups 1a and 1b (see Figure 2) involve a very intense band in the 2140–2090 cm$^{-1}$ region that is characteristic of a ketene group, which suggests that the $\text{HCOHCO} \cdot \cdot \cdot \text{NH}_2\text{OH}$ complex undergoes a similar photoconversion reaction to that of $\text{HCOHCO} \cdot \cdot \cdot \text{CH}_2\text{OH}$, during which double hydrogen transfer occurs and then a hydroxyketene–hydroxylamine complex is formed (Equation (2)).

$$\text{HCOHCO} \cdot \cdot \cdot \text{NH}_2\text{OH} \rightarrow \text{H(OH)C}=C=O \cdot \cdot \cdot \text{NH}_2\text{OH} \quad (2)$$

For the $\text{H(OH)C}=C=O \cdot \cdot \cdot \text{CH}_2\text{OH}$ complex, the $\nu_{as}(C=O)$ band was observed at ca. 2105 cm$^{-1}$.

Three new bands assigned to groups 1a and 1b appear in the $\nu(\text{OH})$ region, one of which has a close wavenumber to the $\nu(\text{OH})$ of $\text{HK}$ in the $\text{H(OH)C}=C=O \cdot \cdot \cdot \text{CH}_2\text{OH}$ complex (ca. 3400 cm$^{-1}$) [18]. The other absorptions occur in the vicinity of $\delta\text{NOH}$, $\omega\text{NH}_2$ of $\text{HA}$ and $\delta\text{CH} + \nu_3\text{CCO}(\nu\text{CO})$ vibrations of $\text{HK}$ (see Table 1). These spectroscopic data support
the conclusion that during the irradiation of the HCOHCO···NH₂OH complexes, their photoconversion to H(OH)C=C=O···NH₂OH complexes (HK–HA) takes place.

![Figure 2](image_url)  
Figure 2. Selected regions of the difference spectrum of the CHOCHO/NH₂OH/Ar(N₂) matrix after 90 min of irradiation minus freshly deposited CHOCHO/NH₂OH/Ar(N₂) matrix at 11 K.

| H(OH)CCO–NH₂OH | Approximate Description | Exp. | Calc. | Exp. | Exp. | Calc. | Approximate Description |
|----------------|------------------------|------|------|------|------|------|-------------------------|
| 3608.0         | ν(OH) HA, IVHKH        | 3573.2 | 3607.1 | 3667 (70) | 3607.1 | 3667 (70) | ν(OH) HA, IVHKH          |
| 3537.2         | ν(OH) HK, IVHKH        | 3600.0 | 3394.5 | 3488 (548) | 3488 (548) | 3488 (548) | ν(OH) HK, IVHKH          |
| 3260 (697)     | ν(OH) HK, I HKH        | 2116.4 | 2119.7 | 2115 (391) | 2115 (391) | 2115 (391) | ν₃COCO HK, IHKHK         |
| 799.8 sh       | τ(OH) HK, I HKH        | 1393.6 | 1358.5 | 1349 (34)  | 1349 (34)  | 1349 (34)  | δNOH HA, IVHKHK          |
| 792.2          |                         | 1153.7 | 1158.8 | 1167 (144) | 1167 (144) | 1167 (144) | δCH +  ν₃COCO HK, IHKHK, IVHKHK |

| 792.2          |                         | 799.8 sh | 752 (119) | 623 (102) | 623 (102) | 623 (102) | τ(OH) HK, I HKH          |

*MP2-calculated harmonic intensities (km mol⁻¹) are given in parentheses; b broad band maximum (see in the text).
Table 2. Comparison of the observed and calculated anharmonic wavenumbers for CHOCH(OH)NHOH.

| CHOCH(OH)NHOH | Exp. a | Calc. b |
|---------------|-------|---------|
| 2 (Ar)        |       |         |
|                | 2 (N2)|         |
| 3623.8        | 3676 (63) | 3655 (73) | 3677 (73) |
| 3558.0        | 3656 (71) | 3578 (71) | 3656 (66) |
| 1769.2, 1758.7, 1750.0, 1745.8, 1733.4 | 1754.3 | 1718 (85) | 1720 (87) | 1721 (92) |
| 1390.9        | 1393 (7) | 1416 (46) | 1385 (16) |
| 1373.5        | 1358 (32) | 1394 (12) | 1363 (12) |
| 1302.5        | 1293 (29) | 1299 (37) | 1303 (19) |
| 1228.7, 1226.5, 1204.6 | 1203.7 | 1169 (55) | 1179.4, 1176.5 |
| 1185.1, 1179.4, 1176.5 | 1169 (55) | 1162 (85) | 1153 (23) |
| 1089.3, 1080.4, 1075.7, 1052.5 | 1081.7, 1060.0 | 1060 (72) | 1036 (29) | 1073 (68) |
| 1038.5, 1021.2, 1014.7 | 1013.6 | 1036 (10) | 998 (59) | 1042 (17) |
| 996.4, 990.2 | 984 (94) | 953 (14) | 990 (59) |
| 920.8, 893.1, 887.3, 884.0 | 916 (34) | 871 (68) | 951 (3) |
| 835.7, 827.5, 824.0 | 806 (36) | 806 (31) | 884 (66) |

a the wavenumber of the most intense line is in bold; b the MP2-calculated harmonic intensities (km mol\(^{-1}\)) are given in parentheses.

The bands assigned to groups 1a and 1b are attributed to two different types of HK–HA complexes as discussed below. Bands of group 1a are only identified in an argon matrix, while bands of group 1b are observed in both solid argon and nitrogen.

The structures of HK–HA complexes of 1:1 stoichiometry were optimized by the MP2/6-311++G(2d,2p) method. The calculations resulted in twelve stationary points (IHKH–XIIHKH), whose structures and ∆ECP(ZPE) binding energies are shown in Figure S2 in the Supplementary Materials. The six most stable structures, IHKH–VIHKH, are also presented in Figure 3. The geometrical parameters for these structures are given in Table S2 in the Supplementary Materials. The three most stable structures, IHKH–IIIHKH, are stabilized by the OH . . . N hydrogen bond between the hydroxyl group of HK and the N atom of the amino group of HA. Structures IIHKH and IIIHKH are additionally stabilized by OH . . . O interactions, where the hydroxylamine OH group serves as a proton donor and the O atom of the hydroxyl group of hydroxyketene serves as a proton acceptor. The next three structures, IVHKH–VIHKH, are stabilized by the OH . . . O hydrogen bond that occurs between the hydroxyl group of HK and the oxygen atom of the hydroxyl group of HA. Complex VHKH is additionally stabilized by the NH . . . O interaction of the amino group of HA and the hydroxyl group of HK. Structures IVHKH–VIHKH are predicted to be about 6–7 kJ mol\(^{-1}\) less stable than structures IHKH–IIIHKH.

In Tables S3–S5 in the Supplementary Materials, the harmonic and anharmonic wavenumbers predicted at the MP2/6-311++G(2d,2p) level for HA and HK monomers and HA–HK complexes (IHKH–VIHKH) are listed. In Table 1, the experimental wavenumbers observed for the photoproducts 1a and 1b are compared with their corresponding calculated wavenumbers of structures IHKH and IVHKH, respectively. Because the wavenumbers and intensities of the vibrational bands of complexes IHKH–IIIHKH are quite similar (see Table S5 in the Supplementary Materials), in Table 1 we only present the data calculated for the most stable structure in the group, IHKH. The same applies for complexes IVHKH–VIHKH, as in Table 1 we only present the data for the structure IVHKH. Comparison of the experimental and calculated wavenumbers suggests that the bands of group 1a are due to HA–HK complexes stabilized mainly by an OH . . . N bond and represented by structure IHKH, while the bands assigned to group 1b correspond to complexes stabilized by an OH . . . O bond and represented by structure IVHKH. The bands of the ν(OH) and
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was not identified. This is most probably due to the fact that the absorption is broad and
(calculated intensity $I = 365 \text{ km mol}^{-1}$) originating from disturbed hydroxylamine as well
as $\nu$(OH) at 3260 cm$^{-1}$ ($I = 697 \text{ km mol}^{-1}$) and $\tau$(OH) at 752 cm$^{-1}$ ($I = 119 \text{ km mol}^{-1}$) due
to perturbed hydroxyketene. In the spectra of the argon matrices, we observed two bands
of group I at 3537.2 cm$^{-1}$ and 792.2 cm$^{-1}$ corresponding to the disturbed $\nu$(OH) vibration
of HA and $\tau$(OH) vibration of HK, respectively. The $\nu$(OH) absorption of disturbed HK
was not identified. This is most probably due to the fact that the absorption is broad and
diffuse, which is a commonly observed feature for the $\nu$(OH) stretch of the relatively strong
O–H· · · O or O–H· · · N bonds [37]. Moreover, possible overlapping of this band with the
$\nu$(OH) absorption of the HA dimer that occurs in this region [38] makes identification
of the band even more difficult. Two additional bands are identified for $I_{\text{HKH}}$ at 1339.6 and
1153.7 cm$^{-1}$. The first one is attributed to the $\delta$CH + $\gamma$,CCO mode of HK and the second
one to the perturbed $\omega$NH$_2$ mode of HA, in accordance with the calculations (see Table 1).
The calculations predict that for complex $\text{IV}_{\text{HKH}}$, the $\nu$(OH) bands due to disturbed HA
and HK are located at 3667 cm$^{-1}$ ($I = 70 \text{ km mol}^{-1}$) and 3488 cm$^{-1}$ ($I = 548 \text{ km mol}^{-1}$),
respectively. These bands are identified at 3608.0, 3607.1 cm$^{-1}$ (HA) and 3400.0, 3394.5 cm$^{-1}$
(HK) in the spectra of the argon and nitrogen matrices, respectively. The $\nu$(OH) band of
HK in $\text{IV}_{\text{HKH}}$ is observed at a very similar wavenumber to the corresponding band of the
HK–Me complex (3395.7 cm$^{-1}$) [18], which confirms that group 1b belongs to a complex in
which HK plays the role of proton donor toward the oxygen atom of the OH group of HA,
forming an O–H· · · O bond (which is the case for structures $\text{IV}_{\text{HKH}}, \text{V}_{\text{HKH}}$ and $\text{VI}_{\text{HKH}}$).

![Figure 3](image-url)

**Figure 3.** The MP2/6-311++G(2d,2p) optimized structures of the six most stable hydroxyketene–
hydroxylamine complexes. The $\Delta E^{\text{CP}}_{\text{(ZPE)}}$ binding energies in kJ mol$^{-1}$ are given in parentheses. The
intermolecular distances are given in Å.

The shape of the very broad band of the $\nu_{\text{as}}$,C=O vibration of HK with few sub-
peaks (or folds) on the band (see Figure 4) suggests that different structures of the HA–HK
complex with similar wavenumbers of the $\nu_{\text{as}}$,C=O mode are probably isolated in the
argon and nitrogen matrices. This band is broader in the Ar than in the N$_2$ matrix, which is
in accordance with the fact that in solid argon all six structures $I_{\text{HKH}}$–$\text{VI}_{\text{HKH}}$ of HA–HK
may contribute to its broadness whereas in solid nitrogen it is only the three $\text{IV}_{\text{HKH}}$–$\text{VI}_{\text{HKH}}$.
The MP2 calculations predict that the position of this band in each particular structure
differs by a few wavenumbers (from 2114 cm$^{-1}$ to 2123 cm$^{-1}$, see Table S5 in the
Supplementary Materials).
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Figure 4. The 2150–2080 cm⁻¹ region of the difference spectra of: (A) the CHOCHO/NH₂OH/Ar(N₂) matrix after 90 min of irradiation minus the freshly deposited CHOCHO/NH₂OH/Ar(N₂) matrix; (B) the CHOCHO/ND₂OD/Ar(N₂) matrix after 90 min of irradiation minus the freshly deposited CHOCHO/ND₂OD/Ar(N₂) matrix.

To obtain some information about the mechanism of photoconversion of Gly–HA to HK–HA, we performed experiments with deuterated hydroxylamine, ND₂OD (HA_d). The isotopic enrichment was ca. 90% as estimated from the measured and theoretically predicted intensities of the bands due to the ν(OH) and ν(OD) vibrations. The spectra of the matrices doped with Gly and HA_d indicate that the complexes Gly–HA_d formed in solid argon and nitrogen have the same structures as those observed for non-deuterated hydroxylamine. The spectra are presented in Figure S1, and the wavenumbers of the bands of Gly–HA_d complexes are listed in Table S1 in the Supplementary Materials. The irradiation (λ > 370 nm) of the Gly/HA_d/Ar(N₂) matrices leads to a decrease in bands due to the Gly–HA_d complexes and due to the formation of the new bands of the photoproducts. Like in the experiments with non-deuterated Gly–HA complexes, the bands of photoproducts can be separated into three groups: 1a, 1b and 2. The difference spectra presented in Figure 5 show the bands attributed to all three groups. The identified 1a and 1b bands evidence the formation of the HK_d–HA_d complexes of hydroxyketene with hydroxylamine, with structures analogical to those observed in the experiment with the non-deuterated HA (IHKH and IVHKH). The wavenumbers of the identified bands assigned for the HK_d–HA_d complexes are listed in Table 3.

The spectra recorded after irradiation of the matrices with deuterated hydroxylamine involve numerous bands due to a number of species. The HK_d–HA_d complexes are formed with relatively low yield, and the identification of their absorptions is quite a difficult task. However, we were able to unequivocally identify in the photolyzed CHOCHO/ND₂OD/Ar matrices four bands corresponding to group 1a (3536.1, 2614.2, ~2113 and 1000.0 cm⁻¹) and eight bands belonging to group 1b (2661.4, ~2113, 1360.4, 1153.6 (subpeaks 1161.2, 1157.2 cm⁻¹), 1136.8 and 1017.6 cm⁻¹). In the spectra of photolyzed CHOCHO/ND₂OD/N₂ matrices, only bands due to group 1b occurred (3349.6, 2664.5, 2495.7, 2116.7, 1360.3, 1160.7, 1155.6, 1137.2 and 1021.4 cm⁻¹). The wavenumbers of all bands belonging to groups 1a and 1b are listed in Table 3.
Among the four identified bands for group 1a in solid argon, the most informative are
whether in the exchange process the OD or ND part participates, leading to the formation of H(OD)CCO complexes
is attributed to the δ band for both complex I and II. The broad, intense ν of this band indicates that one of the deuterium atoms of the ND
the complex formed involves ND···NHDOD characterized by similar structures to the non-deuterated complexes I and IV. The additional subscript d or d2 is applied to mark whether the complex formed involves ND2OH, I, IV, or ND2OH, I, IV (i.e., whether in the exchange process the OD or ND group of hydroxylamine participates). Among the four identified bands for group 1a in solid argon, the most informative are those observed at 3536.1 and 2614.2 cm−1. The first one appears in the OH stretching region of hydroxylamine and shows very good agreement with the wavenumber calculated for the OD stretch of NHDOD in complex I, which provides strong evidence that in the exchange reaction with the hydrogen atom of CH, the deuterium of the OD group of hydroxylamine participates, leading to the formation of H(OD)CCO···ND2OH, complex I. In turn, the wavenumber of the identified 2614.2 cm−1 band has practically the same value as that calculated for the OD stretch of NH2 in complex I (2615 cm−1). The appearance of this band indicates that one of the deuterium atoms of the ND2 group participates in the exchange reaction and the H(OD)CCO···NH2 complex I is formed. The broad, intense νC=O absorption at ca. 2113 cm−1 probably involves the corresponding band for both complex I and complex I. The band observed at 1000 cm−1 is attributed to the δCOD vibration of I in accordance with calculations, and the corresponding band is also observed for the 1b complex.

Figure 5. Selected regions of the difference spectrum of the CHOCHO/ND2OD/Ar(N2) matrix after 90 min of irradiation minus the freshly deposited CHOCHO/ND2OD/Ar(N2) matrix.
Numerous bands were identified for the 1b complexes in both solid argon and nitrogen, and they have close wavenumbers in the spectra of the two matrices. Comparison of the identified wavenumbers with the calculated ones for various types of deuterated complexes shows good agreement with the wavenumbers calculated for complex \( \text{IV}_{\text{HKH-d2}} \) (Table 3). Strong evidence for \( \text{IV}_{\text{HKH-d2}} \) formation is provided by the band observed at 2661.4, 2664.5 cm\(^{-1}\) in the spectra of solid argon and nitrogen, respectively, which is due to the \( \nu \text{OD} \) vibration of perturbed hydroxylamine. Its wavenumber is ca. 45 cm\(^{-1}\) higher than the \( \nu \text{OD} \) of complex \( \text{I}_{\text{HKH-d2}} \) (2614.2 cm\(^{-1}\)), as expected when the OD group of \( \text{HA}_d \) serves as a proton acceptor. The appearance of this band also shows that in the process of exchange with the H atom of glyoxal, one of the deuterium atoms of the ND\(_2\) group participates. The bands identified at 3349.6, 1155.6 cm\(^{-1}\) (solid nitrogen) and at 1153.6 cm\(^{-1}\) (solid argon) are attributed to the \( \nu \text{NHD} \) vibrations according to calculations and are direct evidence for the formation of NHDOD. The other four bands identified for complex \( \text{IV}_{\text{HKH-d2}} \) are attributed to perturbed \( \text{HK}_d \) vibrations (see Table 3).

As discussed above, the experiments with ND\(_2\)OD prove that the irradiation of CHOCHO\(\cdots\)ND\(_2\)OD leads to the formation of H(OD)CCO\(\cdots\)NHDOD and H(OD)CCO\(\cdots\)ND\(_2\)OH complexes as bands due to perturbed NHDO and ND\(_2\)OH molecules can be clearly identified in the spectra. This fact demonstrates that in the hydrogen and deuterium exchange reaction between glyoxal and hydroxylamine, the deuterium atom of the OD group or one of the deuterium atoms of the ND\(_2\) group participates: (i) CHOCHO\(\cdots\)ND\(_2\)OD \(\rightarrow\) H(OD)CCO\(\cdots\)ND\(_2\)OH or (ii) CHOCHO\(\cdots\)ND\(_2\)OD \(\rightarrow\) H(OD)CCO\(\cdots\)NHDOD. In the argon matrix, three complexes are probably formed, namely \( \text{I}_{\text{HKH-d2}}, \text{I}_{\text{HKH-d4}}, \) and \( \text{IV}_{\text{HKH-d2}} \), but in the nitrogen matrix only the latter. Whereas the formation of \( \text{IV}_{\text{HKH-d2}} \) is very well documented by a number of bands identified for this complex, the presence of \( \text{I}_{\text{HKH-d2}} \) and \( \text{I}_{\text{HKH-d4}} \) is evidenced by one band characteristic of the particular complex \( \nu \text{OD} \text{HA}_d \) at 2614.2 cm\(^{-1}\) for \( \text{I}_{\text{HKH-d2}} \) and \( \nu \text{OH} \text{HA}_d \) at 3536.1 cm\(^{-1}\) for \( \text{I}_{\text{HKH-d4}} \). The other two identified bands of group 1a \( (2113, 1000 \text{ cm}^{-1}) \) attributed to \( \nu \text{NHD} \text{C}=\text{O} \) and \( \delta \text{COD} \) vibrations may be due to both complex \( \text{I}_{\text{HKH-d2}} \) and complex \( \text{I}_{\text{HKH-d4}} \).

The obtained data show that the non-planar complexes of glyoxal with hydroxylamine, \( \text{Gly} \text{HA}_d, \text{IGH} \) (stabilized by the OH\(\cdots\)O bond), that are formed in the argon matrices may photoconvert to the \( \text{I}_{\text{HKH-d2}}, \text{I}_{\text{HKH-d4}} \) and \( \text{IV}_{\text{HKH-d2}} \) \( \text{HK}_d \text{HA}_d \) complexes of hydroxyketene with hydroxylamine. In the nitrogen matrices, both the non-planar, \( \text{I}_{\text{GH}} \) and

### Table 3. Comparison of the observed and calculated anharmonic wavenumbers (cm\(^{-1}\)) for the H(OD)CCO–ND\(_2\)OH, H(OD)CCO–NHDOD complexes.

| Exp. | Calc. \(^a\) | Calc. \(^a\) | Exp. \(^b\) | Exp. \(^b\) | Calc. \(^a\) | Approximate Description |
|------|-------------|-------------|-------------|-------------|-------------|------------------------|
| 1a (Ar) | \( \text{I}_{\text{HKH-d2}} \) | \( \text{I}_{\text{HKH-d4}} \) | 1b (Ar) | 1b (N\(_2\)) | \( \text{IV}_{\text{HKH-d2}} \) | \( \nu \text{OH} \text{HA}_d, \text{I}_{\text{HKH-d4}} \) |
| 3536.1 | 3533 (305) | 3349.6 | 3356 (43) | \( \nu \text{NH} \text{HA}_d, \text{IV}_{\text{HKH-d2}} \) |
| 2614.2 | 2615 (184) | 2661.4 | 2658.4 | 2710 (38) | \( \nu \text{OD} \text{HA}_d, \text{I}_{\text{HKH-d2}}, \text{IV}_{\text{HKH-d2}} \) |
| 2432 (382) | 2472 (261) | 2495.7 | 2579 (278) | \( \nu \text{OD} \text{HA}_d, \text{I}_{\text{HKH-d2}}, \text{IV}_{\text{HKH-d2}} \) |
| 2113 \(^c\) | 2114 (441) | 2119.6 | 2118 (394) | \( \nu \text{asCCO} \text{HK}_d, \text{I}_{\text{HKH-d2}}, \text{IV}_{\text{HKH-d2}} \) |
| 1000.0 | 1029 (45) | 1026.5 | 1021.4 | 1009 (53) | \( \delta \text{COD} \text{HK}_d, \text{I}_{\text{HKH-d2}}, \text{IV}_{\text{HKH-d2}} \) |

\(^{a}\) MP2-calculated harmonic intensities (km mol\(^{-1}\)) are given in parentheses; \(^{b}\) the most intense line of the 2 or 3 lines is in bold; \(^{c}\) broad band maximum (see in the text).
planar, \( \text{II}_{\text{GH}} \) (main product), \( \text{Gly–HA}_d \) complexes are present (the latter stabilized by OH . . . O and CH . . . N bonds), and after conversion the \( \text{IV}_{\text{HKH-d2}} \) \( \text{HK}_d–\text{HA}_d \) complexes are formed. These data suggest that irradiation of the non-planar \( \text{Gly–HA}, \text{HA}_d \) complexes stabilized by the OH . . . O hydrogen bond, \( \text{I}_{\text{GH}} \), leads to double hydrogen exchange (or hydrogen and deuterium exchange) between glyoxal and hydroxylamine in which the hydroxyl or the amino group of \( \text{HA}, \text{HA}_d \) may be involved. The irradiation of the planar complex, \( \text{II}_{\text{GH}} \), stabilized by OH . . . O and CH . . . N hydrogen bonds, proceeds by double hydrogen exchange between the CH group of glyoxal and the amino group of \( \text{HA} \) only.

In the case of the \( \text{CHOCHO–CH}_2\text{OD} \) complex, the photoconversion resulted in an exchange of proton and deuterium between the CH group of glyoxal and the OD group of methanol, and finally the \( \text{H(OOD)CCO–CH}_3\text{OH} \) complex was formed. According to the coupled-cluster calculations, the non-hydrogen-bonded (non-planar) glyoxal–methanol complex photoconverted initially to a planar hydrogen-bonded complex, which then converted to the final photoproduct (hydroxyketene–methanol complex) [18,20].

2.2. Formation of Hydroxy(hydroxyamino)acetaldehyde (Hemiaminal)

Irradiation of \( \text{Gly/HA/Ar(N}_2\text{)} \) matrices with a wavelength of \( \lambda \geq 370 \text{ nm} \) also leads to the appearance of the group of bands labeled as group 2 (see Figure 2 and Table 4). This group is assigned to different conformers of the product of the addition of hydroxylamine to one of the CHO groups of glyoxal, namely, hydroxy(hydroxyamino)acetaldehyde, \( \text{HHA} \). The formation of \( \text{HHA} \) proceeds via the addition of the nitrogen atom of the amino group of hydroxylamine to the carbon atom of the carbonyl group of glyoxal and the subsequent migration of the hydrogen atom (from the \( \text{NH}_2 \) group of \( \text{HA} \)) to the oxygen atom of the carbonyl group of glyoxal, as presented in Scheme 1. This molecule is formed readily in an argon matrix, but we only observe trace amounts of this photoproduct in a nitrogen matrix. The molecule has not been characterized so far, and its infrared spectrum is unknown. The formation of \( \text{CHOCHO–NH}_2\text{OH} \) has been reported [12], but we are not able to unequivocally assign the experimental wavenumbers to this form. The irradiation of the planar \( \text{GH}–\text{HA} \) complex, the photoconversion resulted in an exchange of proton and deuterium between the CH group of glyoxal and the OD group of methanol, and finally the \( \text{H(OOD)CCO–CH}_3\text{OH} \) complex was formed. According to the coupled-cluster calculations, the non-hydrogen-bonded (non-planar) glyoxal–methanol complex photoconverted initially to a planar hydrogen-bonded complex, which then converted to the final photoproduct (hydroxyketene–methanol complex) [18,20].

Table 4. Comparison of the observed and calculated anharmonic wavenumbers for CHOCH(OD)NDOD.

| Exp. \(^a\) | Calc. \(^b\) |
|-------------|-------------|
| \( 2 (\text{Ar}) \) | \( 2 (\text{N}_2) \) | \( \text{I}_{\text{HHA-d}} \) | \( \text{II}_{\text{HHA-d}} \) | \( \text{III}_{\text{HHA-d}} \) |
| 2674.2 | 2702 (39) | 2702 (40) | 2715 (41) |
| 2624.3 | 2702 (39) | 2642 (41) |
| 1764.2, 1749.6, 1745.4, 1744.0 | 1761.9, 1745.0 | 1719 (84) | 1711 (88) | 1721 (90) |
| 1339.3 | 1346.6 | 1347 (3) | 1367 (12) | 1341 (18) |
| 1254.9, 1245.4, 1233.0 | 1255.9, 1251.8, 1246.0 | 1143 (67) | 1176 (40) | 1169 (20) |
| 1094.0 | 1136 (29) | 1101 (29) | 1143 (9) |
| 986.2 | 1013.3, 1010.5 | 990 (8) | 974 (41) | 992 (2) |
| 788.5, 785.3, 782.9 | 800.3, 788.0, 783.4 | 750 (58) | 729 (35) | 745 (43) |

\(^a\) the wavenumber of the most intense line is in bold; \(^b\) the MP2-calculated harmonic intensities (km mol\(^{-1}\)) are given in parentheses.

Scheme 1. Schematic reaction path for the formation of CHOCH(OH)NHOH, \( \text{HHA} \), from the CHOCHO–NH\(_2\)OH complex.
Tables S8 and S9 in the Supplementary Materials show the harmonic and anharmonic wavenumbers and band intensities predicted at the MP2/6-311++G(2d,2p) level and the potential energy distribution (PED) for the two most stable forms of HHA. In Table S10, the calculated wavenumbers and band intensities of I\textsubscript{HHA}–VI\textsubscript{HHA} structures are given for comparison. In Table 2, the experimental wavenumbers observed for photoproduct 2 are compared with the calculated wavenumbers (and intensities) of the bands predicted for the three most stable conformers of HHA (I\textsubscript{HHA}–III\textsubscript{HHA}).

The appearance of several bands in the νC=O region and other distinct spectral regions indicates that more than one form of HHA is trapped in the argon matrix. The data suggest that one form probably dominates, and another few are isolated in smaller amounts. In the νOH region, we only observe two bands assigned to this group located at 3623.8 and 3558.0 cm\(^{-1}\), which, according to the calculations, most suit the II\textsubscript{HHA} form. Other identified bands in group 2 also mostly agree with the wavenumbers calculated for the conformer II\textsubscript{HHA}. However, we are not able to unequivocally assign the experimental bands to a specific structure because all the structures have similar vibrational characteristics and the amounts of the photoproducts in the matrices are small. Furthermore, the result of the experiment using deuterated hydroxylamine does not give clear evidence as to which conformers of HHA are formed in the matrices. The results of this experiment are presented in Figure 5 and Table 2 and in Tables S8, S9 and S11 in the Supplementary Materials.

A photoaddition reaction was also observed as a result of the irradiation (at λ > 345 nm) of the glyoxal–hydrogen peroxide complex in an argon matrix [17], in which 2-hydroperoxyethanal was formed. The formation of this photoproduct proceeds via the addition of one of the oxygen atoms of hydrogen peroxide to the carbon atom of the carbonyl group of glyoxal and the subsequent migration of the hydrogen atom (from H\textsubscript{2}O\textsubscript{2}) to the oxygen atom of the carbonyl group of glyoxal.

![Figure 6](image_url)  
**Figure 6.** The optimized structures of the three most stable forms of hydroxy(hydroxyamino)acetaldehyde, HHA. The computed ΔE\textsubscript{ZPE} energies of HHA conformers with respect to the energy of the most stable conformer, I\textsubscript{HHA}, in kJ mol\(^{-1}\) are given in parentheses.

3. Experimental and Computational Methods

Gaseous hydroxylamine (NH\textsubscript{2}OH, HA) was prepared from hydroxylamine phosphate salt (95%, Fluka) by heating the salt powder (at 50–65 °C) directly in the deposition line. Deuterated hydroxylamine (ND\textsubscript{2}OD, HA\textsubscript{d}) was prepared by heating hydroxylamine phosphate salt in deuterated water (D\textsubscript{2}O) solution and then evaporating off the water under a vacuum. This procedure was repeated several times until the deuteration degree was about 90%. Monomeric glyoxal (CHOCHO, Gly) was obtained from solid trimer dihydrate (98%, Sigma) topped with phosphorus pentoxide (P\textsubscript{2}O\textsubscript{5}) powder by heating the materials to 120 °C under a vacuum and then collecting the gaseous glyoxal in a liquid nitrogen trap.

The glyoxal/hydroxylamine/argon (or nitrogen) matrices were prepared by simultaneous deposition of CHOCHO/Ar(N\textsubscript{2}) and NH\textsubscript{2}OH vapor on a cold gold mirror kept at 11–17 K by a closed cycle helium refrigerator (Air Products, Displex 202A). The CHOCHO/Ar(N\textsubscript{2}) matrix concentration was varied in the range of 1/200–1/2000. The absolute concentration of hydroxylamine in the matrices could not be determined, but its concentration was varied by changing the rare gas flow rate and the heating temperature of the hydroxylamine salt. Infrared spectra (4000–600 cm\(^{-1}\)) with a resolution of 0.5 cm\(^{-1}\) were recorded in reflection mode using a Bruker 113v FTIR spectrometer and a liquid N\textsubscript{2}-cooled MCT detector.
The samples deposited in matrices were subjected to the radiation of a 200 W medium-pressure mercury lamp (Philips CS200W2). A 5 cm water filter (to reduce the infrared light) and a glass long-wavelength pass filter (WG370) were used.

The MP2 method with 6-311++G(2d,2p) basis set was used for geometry optimization of all the monomers and complex structures and calculation of harmonic and anharmonic vibrational spectra of the monomers, complexes and photoproducts [39,40]. Glyoxal under standard conditions exists in a trans form [34,41,42], so this conformer was considered exclusively in our study in terms of the nature of its interaction with hydroxylamine. Binding energies were corrected using the Boys–Bernardi full counterpoise procedure [43]. The calculations were performed using the Gaussian 03 program (version C.02) for geometry optimization and Gaussian 16 (version B.01) for frequency calculations [44,45]. The potential energy distribution (PED) of the normal modes was computed using the Gar2ped program [46].

4. Conclusions

The photochemistry of glyoxal–hydroxylamine complexes isolated in argon and nitrogen matrices has been investigated using FTIR and MP2/6-311++(2d,2p) calculations. The irradiation of the studied complexes in the visible range (λ > 370 nm) leads to the formation of two types of photoproducts: (i) hydroxyketene–hydroxylamine complexes, HK–HA, and (ii) hydroxy(hydroxyamino)acetdehyde, HHA.

The HK–HA complexes are formed in the exchange reaction of two hydrogens between two complex subunits, namely, the CH group of Gly and the NH or OH group of HA. The experiments with deuterated hydroxylamine, ND\textsubscript{2}OD, provide evidence that hydrogen atoms from both the OH and NH\textsubscript{2} groups of hydroxylamine may participate in the exchange process. The H(OH)CCO . . . NH\textsubscript{2}OH complexes exist in two forms in the matrices. The first type is stabilized by the OH . . . N interaction between the hydroxyl group of HK and the amino nitrogen of HA. In the second type, the OH . . . O hydrogen bond is formed between the hydroxyl group of HK and the oxygen atom of the hydroxyl group of HA. These two types of HK–HA complex are formed in an argon matrix in noticeable amounts, but in nitrogen only HK–HA complexes bonded by the OH . . . O interaction are created.

Hydroxy(hydroxyamino)acetdehyde, HHA, belonging to the hemiaminals family, is formed by the addition of HA to the carbon atom of one aldehyde group of Gly and the subsequent migration of the hydrogen atom from the NH\textsubscript{2} group of HA to the oxygen atom of the carbonyl group of Gly. In an argon matrix, the HHA hemiaminal conformers are observed in noticeable amounts, whereas in a nitrogen matrix the yield of HHA is low. The difference in the yield of the products, both HK–HA complexes and the HHA adduct, in the argon and nitrogen matrices may relate to the fact that different types of glyoxal–hydroxylamine (Gly–HA) complexes exist in argon and nitrogen before irradiation.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/molecules27154797/s1, Table S1: Comparison of the observed wavenumbers (cm\textsuperscript{-1}) and wavenumber shifts (Δν = ν\textsubscript{GH} – ν\textsubscript{M}) for the Gly–HA (GH) complexes present in the Ar and N\textsubscript{2} matrices with the corresponding calculated values for the complexes I\textsubscript{GH} and II\textsubscript{GH}. The calculated band intensities are given (in km mol\textsuperscript{-1}) in parentheses; Figure S1: Spectra of the CHOCHO/Ar(N\textsubscript{2}) (a), NH\textsubscript{2}OH/Ar(N\textsubscript{2}) (b) and CHOCHO/NH\textsubscript{2}OH(ND\textsubscript{2}OH)/Ar(N\textsubscript{2}) (c) matrices recorded after matrix deposition at 11 K. The bands of Gly–HA complexes are indicated by arrows. Dashed and solid arrows correspond to the I\textsubscript{GH} and II\textsubscript{GH} structures, respectively; Figure S2: The MP2/6-311++G(2d,2p) optimized structures of the hydroxyketene–hydroxylamine (I\textsubscript{HKH}–XI\textsubscript{HKH}) complexes. The ΔE\textsubscript{CP} (ZPE) binding energies (in kJ mol\textsuperscript{-1}) are given in parentheses. The intermolecular distances are given in Å; Table S2: Selected geometrical parameters of the hydroxylamine and hydroxyketene subunits in their binary complexes (I\textsubscript{HKH}–VI\textsubscript{HKH}); Table S3: Calculated harmonic (ν) and anharmonic (γ\textsubscript{anh}) wavenumbers (cm\textsuperscript{-1}) and intensities (I, km mol\textsuperscript{-1}) for HA and H\textsubscript{AD}; Table S4: Calculated harmonic (ν) and anharmonic (γ\textsubscript{anh}) wavenumbers (cm\textsuperscript{-1}), intensities
(I, km mol\(^{-1}\)) and potential energy distribution (PED) for HK and HK\(_d\): Table S5: Calculated harmonic (\(\nu\)) and anharmonic (\(\nu_{\text{anh}}\)) wavenumbers (cm\(^{-1}\)) and intensities (I, km mol\(^{-1}\)) for HK–HA complexes (\(\text{HKKH}−\text{V}_{\text{HHK}}\)); Table S6: Calculated harmonic (\(\nu\)) and anharmonic (\(\nu_{\text{anh}}\)) wavenumbers (cm\(^{-1}\)) and intensities (I, km mol\(^{-1}\)) for deuterated hydroxyamine–hydroxyketene complexes (\(\text{HKKH}−\text{V}_{\text{HHK}}\); Table S7: Optimized structures of hydroxy(hydroxyamino)acetaldehyde (HHA). The computed \(\Delta E_{\text{ZPP}}\) energies of HHA conformers with respect to the energy of the most stable conformer. \(I_{\text{HHA}}\) are given in parentheses (in kJ mol\(^{-1}\)); Table S8: Calculated harmonic (\(\nu\)) and anharmonic (\(\nu_{\text{anh}}\)) wavenumbers (cm\(^{-1}\)), intensities (I, km mol\(^{-1}\)) and PED for \(I_{\text{HHA}}\) and \(I_{\text{HHA}-\text{d}}\); Table S9: Calculated harmonic (\(\nu\)) and anharmonic (\(\nu_{\text{anh}}\)) wavenumbers (cm\(^{-1}\)), intensities (I, km mol\(^{-1}\)) and PED for \(I_{\text{HHA}}\) and \(I_{\text{HHA}-\text{d}}\); Table S10: Calculated harmonic (\(\nu\)) and anharmonic (\(\nu_{\text{anh}}\)) wavenumbers (cm\(^{-1}\)) and intensities (I, km mol\(^{-1}\)) for conformers of HHA (\(\text{HHA}−\text{V}_{\text{HHA}}\)); Table S11: Calculated harmonic (\(\nu\)) and anharmonic (\(\nu_{\text{anh}}\)) wavenumbers (cm\(^{-1}\)) and intensities (I, km mol\(^{-1}\)) for the deuterated conformers of HHA (\(\text{HHA}-\text{d}−\text{V}_{\text{HHA}-\text{d}}\)).

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