Interstellar Formation of Glyceric Acid (HOCH2CH(OH)COOH) –

The Simplest Sugar Acid

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**Abstract:** Glyceric acid (HOCH2CH(OH)COOH) — the simplest sugar acid — represents a key molecule in biochemical processes vital for metabolism in living organisms such as glycolysis. Although critically linked to the *Origins of Life* and identified in carbonaceous meteorites with abundances comparable to amino acids, the underlying mechanisms of its formation have remained elusive. Here, we report the very first abiotic synthesis of racemic glyceric acid via the barrierless radical-radical reaction of the hydroxycarbonyl radical (HOĊO) with 1,2-dihydroxyethyl (HOĊHCH2OH) radical in low-temperature carbon dioxide (CO2) and ethylene glycol (HOCH2CH2OH) ices. Utilizing isomer-selective vacuum ultraviolet photoionization reflectron time-of-flight mass spectrometry, glyceric acid was identified in the gas phase based on the adiabatic ionization energies and isotopic substitution studies. This work reveals the key reaction pathways for glyceric acid synthesis through non-equilibrium reactions from ubiquitous precursor molecules advancing our fundamental knowledge of the formation pathways of key biorelevant organics – sugar acids – in deep space.

**Teaser:** Glyceric acid was prepared in interstellar ice analogs through barrierless reaction of hydroxycarbonyl with 1,2-dihydroxyethyl.

**Introduction**

Since the first identification of the simplest sugar acid — glyceric acid (HOCH2CH(OH)COOH, **1**) — in the Murchison and Murray meteorites by Cooper *et al*. more than twenty years ago (*1*), **1** has been at the center of attention from the astronomy, astrobiology, and laboratory astrochemistry communities (*2-7*). **1** plays a key role in contemporary biochemical processes vital to cellular metabolism such as glycolysis (Fig. 1). It serves as the starting point to synthesis of 2-phosphoglyceric acid (C3H7O7P, **2**) and 3-phosphoglyceric acid (C3H7O7P, **3**) via phosphorylation (*8*). **1** plays a central role in glycolysis, the tricarboxylic acid (TCA) cycle, and photosynthesis in which the Calvin cycle contributes to carbon metabolism (TCA cycle), carbon fixation, and sugar phosphate production (*9*). Through nucleophilic substitution, **1** may react with ammonia (NH3) to yield theproteinogenic amino acid serine (HOCH2CH(NH2)COOH, **4**), eventually producing complex amino acids and peptides. Through the oxidation with nicotinamide adenine dinucleotide (NAD+), **1** can be formed from glyceraldehyde (HOCH2CH(OH)CHO, **5**), which can then be reduced by nicotinamide adenine dinucleotide phosphate (NADP+) to glycerol (HOCH2CH(OH)CH2OH, **6**) — a key molecular precursor to phospholipids – the main component of cell membranes (*10, 11*).

In prebiotic chemistry, **1** may have served as a fundamental building block for crucial biomolecules (Fig. 1) (*8*). The methylation of **1** can form deoxy sugar acids such as 2-methyl glyceric acid (HOCH2C(CH3)(OH)COOH), **7**). Undergoing carbon-carbon bond cleavage, **1** can be converted into formic acid (HCOOH, **8**) and glycolic acid (HOCH2COOH, **9**). Recently, Marks *et al*. revealed the formation of **9** in irradiated interstellar model ices of carbon dioxide (CO2) and methanol (CH3OH, **10**); **9** was formedthrough barrierless radical-radical recombination of the hydroxycarbonyl radical (HOĊO, **11**) with the hydroxymethyl (ĊH2OH) radical (*12*). Oxidation of **1** results in the formation of biorelevant hydroxypyruvic acid (HOCH2C(O)COOH, **12**) and tartronic acid (HOOCCH(OH)COOH, **13**) (*13*), which can be decarboxylated into **9**. Further, the cleavage of the carbon-oxygen bond in **1** prepares 3-hydroxypropionic acid (HOCH2CH2COOH, **14**) and the biomolecule lactic acid (CH3CH(OH)COOH, **15**) (*14*).

Glyceric acid (**1)** represents a simple representative of sugar-related compounds and can form from the simplest sugar molecule glyceraldehyde (**5**) via oxidation (Fig. 1). Sugar acids and sugar alcohols are vital precursors to the molecular building blocks of biomolecules and could have seeded the evolution of life as we know it (*15*). Therefore, the elucidation of their formation routes can aid in understanding the molecular mass-growth processes of astrobiologically relevant molecules necessary for the origins of life. Among some 300 molecules identified in the interstellar medium (ISM) (*16*), sugar-related molecules such as glycolaldehyde (HCOCH2OH) have been identified toward the hot molecular core Sagittarius B2 (*17, 18*). Although **1** has not yet been identified in the ISM (*16*), it has been observed in laboratory simulation experiments. Ices composed of electron-irradiated methanol (**10**) ice (*4*) and ultraviolet-irradiated ice mixtures of **10**, ammonia (NH3) and water (H2O) (*19, 20*) reveal its formation through the hydrolysis of organic residues remaining after temperature-programmed desorption (TPD) to room temperature. Furthermore, **1** has been identified in the hydrolyzed organic matter of carbonaceous chondrites with an abundance (80 nmol g−1) comparable to amino acids (*1*), indicating that **1** is able to survive the entrance of the meteorite into the atmosphere of the early Earth when embedded in a matrix (*21*). However, unraveling the abiotic formation mechanisms of molecular **1** in interstellar analog ices is still elusive. Under interstellar conditions, the formation of biologically relevant molecules is feasible via abiotic (non-biological) reaction mechanisms, in which these molecules can be formed in interstellar ices composed of simple organic molecules when subjected to ionizing radiation. This understanding is of fundamental importance to the astronomy and astrochemistry communities to rationalize the origin and formation routes of biorelevant molecules linked to the *Origins of Life* and the simplest sugar acid **1** in particular.

Here, we demonstrate the very first abiotic synthesis of **1** in low-temperature (5 K) carbon dioxide and ethylene glycol (HOCH2CH2OH, **16**) ice mixtures via the barrierless radical-radical reaction of the hydroxycarbonyl (HOĊO, **11**) with the 1,2-dihydroxyethyl (HOĊHCH2OH, **17**) radicals (Figs. 1 and 2). These model ices were exposed to energetic electrons mimicking secondary electrons generated in the track of galactic cosmic rays (GCRs) penetrating ices in cold molecular clouds aged a few million years (*22*). By merging our experiments with calculations, **1** was identified in the *gas phase* through vacuum ultraviolet (VUV) photoionization reflectron time-of-flight mass spectrometry (PI-ReTOF-MS) during the temperature-programmed desorption (TPD) of irradiated ices based on computed adiabatic ionization energies (IEs) and isotopic substitution studies. This finding significantly advances the knowledge of the formation pathways of key biorelevant organics — sugar acids — in deep space. Carbon dioxide is one of the most ubiquitous molecules in interstellar ices and has been detected at fractions of up to 40% with respect to water toward the AFGL 989 source (*23*). Ethylene glycol (**16**) generated via radical-radical recombination of the hydroxymethyl (ĊH2OH) radicals (*4, 24*) is abundant in the ISM, Murchison meteorites, and comets (*1, 18, 25*); the abundance of **16** has been found to be 0.25% with respect to water in the Comet C/1995 O1 (Hale-Bopp) (*26*). Therefore, the formation of **1** in interstellar ices containing carbon dioxide and ethylene glycol represents a plausible mechanistic pathway as demonstrated here. Once formed, **1** can be incorporated into planetesimals and ultimately delivered to planets like early Earth via meteorites thus participating in a complex chain of chemical reactions leading to the molecular precursors important for the *Origins of Life*.

**Results**

**Infrared spectroscopy**. Fourier-transform infrared spectroscopy (FTIR) was utilized to monitor the chemical evolution of the ices before (black line) and after (red line) the irradiation (figs. S1 to S4). Detailed assignments of the FTIR spectra are summarized in tables S1 to S4. The absorptions of the deposited ices can be attributed to the fundamentals and combination modes of carbon dioxide and ethylene glycol (**16**). After irradiation at 5 K, several new absorption features were detected. The region between 2200 cm−1 and 1400 cm−1 can be deconvoluted into several Gaussian peaks (fig. S1B). The infrared absorption band at 2140 cm−1 is assigned to the stretch of carbon monoxide (CO); this is confirmed from the observation of C18O at 2090 cm−1 and 13CO at 2092 cm−1 (figs. S2B and 4B) (*27*). The absorptions at 1767 cm−1 and 1657 cm−1 are linked to the formation of one or more carbonyl (C=O) containing species and the bending mode (ν2) of water, respectively (*12*). After accounting for vibrational anharmonicity at the B3LYP/aug-cc-pVTZ level of theory, the strongest vibration mode of **1** is predicted at 1803 cm−1 for **1g**, i.e.,the lowest energy conformer of **1** (table S5). Since vibrational frequencies of matrix-isolated species are often shifted from their gas-phase values (*28*)and FTIR absorptions can be broadened due to the ice matrix, the anharmonic frequency is close to the observed absorptions at 1767 cm−1, indicating that this absorption may be linked to **1** in a polar environment. The absorptions at 1722 cm−1 and 1500 cm−1 are associated with the CO stretching mode (ν2) and CH2 bending mode (ν3) of formaldehyde (H2CO) (*27, 29*); these features can also be identified in deuterium (fig. S3) and 13C (fig. S4) labeled ices through absorptions at 1703 cm−1 (D2CO, ν2), 1681 cm−1 (H213CO, ν2) and 1500 cm−1 (H213CO, ν3) (*12, 27, 29, 30*). The absorptions at 1849 cm−1 in processed CO2−HOCH2CH2OH ice and 1811 cm−1 in irradiated 13CO2−HO13CH213CH2OH ice (figs. S1 and S4) can be assigned to the *trans*-hydroxycarbonyl radical (HOĊO, ν2) and/or formyl radical (HĊO, ν3). Because of the matrix effect, the absorptions of fundamental (CO stretch) of *trans*-hydroxycarbonyl radical (ν2) and formyl radical (ν3) are close to one another (*31, 32*) and cannot be clearly distinguished here. Due to the overlapping absorption features resulting from a wide variety of complex organic molecules produced during radiation processing, infrared spectra cannot uniquely identify complex organic molecules such as **1**, but only their functional groups, highlighting the need for an alternative analytical technique to detect *individual* reaction products (*33, 34*).

**PI-ReTOF-MS**. Photoionization reflectron time-of-flight mass spectrometry (PI-ReTOF-MS) represents an ideal technique to identify individual C3H6O4 isomers in the gas phase based on their distinct adiabatic ionization energies and mass-to-charge ratios (*m/z*) (*33, 35*). Separate experiments exploiting two photon energies of 11.10 eV and 8.77 eV were selected to distinguish the first-generation products **1** and 2-hydroxyethyl hydrogen carbonate **(**HOCH2CH2OCOOH, **18**) formed through radical-radical reactionsas well asthesecond-generation product propene-1,1,2,3-tetrol (HOCH2CH(OH)C(OH)2, **19**)accessedvia tautomerization of **1** (Fig. 2). 11.10 eV photons aid in the photoionization of all three isomers if formed, while 8.77 eV photons only ionize **19** (IE = 7.24 – 7.53 eV), but neither **1** (IE = 9.69 – 10.17 eV) nor **18** (IE = 9.79 – 11.09 eV). Fig. 3 compiles the PI-ReTOF mass spectra of the subliming molecules from the exposed carbon dioxide–ethylene glycol ices during the TPD phase. Tentative assignments of C2H4O3 and C3H6O3 isomers are presented in the Supplementary Materials (figs. S7-S8). Concentrating on the C3H6O4 isomers, the TPD profile of the ion signal at *m/z* = 106 for the irradiated CO2–HOCH2CH2OH ice at 11.10 eV (black line, Fig. 4A) shows sublimation events at 240 K (peak 1) and 273 K (peak 2). Since the signal at *m/z* = 106 can be associated with the molecular formulae C2H2O5, C3H6O4, C4H10O3, C6H2O2, C7H6O, and C8H10, isotopically labeled reactants were exploited to assign the molecular formula(e). The replacement of the CO2–HOCH2CH2OH ice by 13CO2–HO13CH213CH2OH ice shifts the *m/z* by 3 amu from *m/z* = 106 (C3H6O4+) to *m/z* = 109 (13C3H6O4+); this finding confirms the presence of three carbon atoms (Fig. 4A). The substitution of CO2 by C18O2 results in products with two 18O atoms that can be observed at *m/z* = 110 (C3H6O218O2+) in the C18O2–HOCH2CH2OH ice. Finally, deuterated methylene (CD2) moieties (CO2–HOCD2CD2OH ice) result in products with five deuterium atoms; the observed signals shift to *m/z* = 111 (C3HD5O4+) (Fig. 4B). Hence, the sublimation events (peak 1 and peak 2) at *m/z* = 106 can be clearly assigned to a molecule of the formula C3H6O4.

The TPD profile of *m/z* = 106 (C3H6O4+) at 11.10 eV reveals sublimation events at 240 K (peak 1) and 273 K (peak 2) (Fig. 4A). A blank experiment conducted without exposing the ices to ionizing radiation under otherwise identical conditions does not reveal any ion signal at *m/z* = 106 between 210 K and 300 K (Fig. 4A). This finding confirms that peaks 1 and 2 are the result of the irradiation of the ices. At 11.10 eV, all three isomers **1** (IE = 9.69 – 10.17 eV), **18** (IE = 9.79 – 11.09 eV) and **19** (IE = 7.24 – 7.53 eV) can be ionized (Fig. 2; table S5). Thereafter, the photon energy was reduced to 8.77 eV, at this energy only the enol tautomer **19** can be ionized if present. However, upon reducing the photon energy to 8.77 eV, peaks 1 and 2 are absent (Fig. 4C). An additional experiment was performed with a higher dose of irradiation (100 nA, 120 minutes) to probe the formation of **19**. Even at this increased dose, which corresponds to 12 ± 2 eV molecule−1 of carbon dioxide and 21 ± 3 eV molecule−1 of **16**, no ion signal was detectable at *m/z* = 106 confirming the absence of **19**. Therefore, peaks 1 and 2 can only be associated with isomers **1** and/or **18**. Due to the overlap of their IEs,it is imperative to confirm their formation using isotopically labeled ices. Here, the identification of isomers **1** and **18** was achieved by exploiting partially deuterated reactants (CO2–HOCD2CD2OH ice) during photoionizing at 11.10 eV. This strategy is highlighted in fig. S5 and reveals that their ion signals can be separated through the detection of *m/z* = 109 (HOCD2CD(OH)COOH+) for **1** and *m/z* = 111 (HOCD2CD2OCOOD+) for **18**. In the processed CO2–HOCD2CD2OH ice (Fig. 4B), the TPD profile at *m/z* = 109 only shows one sublimation event (peak 2) indicating that peak 2 is linked to **1**.For ion signals of *m/z* = 111, the TPD profile shows both peak 1 and peak 2. This is counterintuitive since only one peak connected to *m/z* = 111 (HOCD2CD2OCOOD+) for **18** should be present. In the processed CO2–HOCD2CD2OH ice, it is likely that 2-hydroxyethoxy-d4 (ȮCD2CD2OH, **20**) can isomerize to **17**-d4 (DOĊDCD2OH), which then reacts with hydroxycarbonyl-d1(DOĊO, **11**) radical to form **1**-d5(HOCD2CD(OD)COOD+; *m/z* = 111) leading to the presence of peak 2. Therefore, only peak 1 can be clearly associated with **18**. This isomerization was also tackled computationally. Using the AE-CCSD(T)/CBS//AE-MP2/aug-cc-pVTZ level of theory, Wang et al. (*36*) calculated the isomerization reaction from 2-hydroxyethoxy (ȮCH2CH2OH, **20**) radicalto1,2-dihydroxyethyl (HOĊHCH2OH, **17**)radicalto be exoergic (44 kJ mol−1) with a reaction barrier of 117 kJ mol−1; this barrier can be overcome by the kinetic energy of the incident electrons. Moreover, the sublimation sequences of **18** at 240 K and **1** at 273 K correlate with an enhanced polarity and increased number of hydroxyl groups (OH) of the subliming isomers (*24, 37-39*). In particular, **1** carries an extra hydroxyl group in comparison to **18**, leading to an increase of the sublimation temperature by 33 K. The detected counts of **1** and **18** in the CO2 – HOCH2CH2OH system were 230 ± 20 and 270 ± 15 counts, respectively. In addition, the TPD profile of the ion signal at *m/z* = 106 for the irradiated CO2–HOCH2CH2OH ice matches the TPD profile at *m/z* = 89 (fig. S6), which may be caused by the photodissociation of **1** and **18** by losing a hydroxyl group (OH) upon ionization at 11.10 eV.

Having provided compelling evidence on the preparation and detection of **1** and **18**, we shift our attention now to their formation mechanisms. *First*, reactions (1)–(3) lead to unimolecular decomposition of **16** to **17** (HOĊHCH2OH;reaction (1)) and to **20** (ȮCH2CH2OH; reaction (2)); the suprathermal hydrogen atoms generated add to the C=O bond in carbon monoxide forming the hydroxycarbonyl radical **11**(HOĊO) (reaction (3)). The decomposition of **16** is endoergic by 398 ± 4 kJ mol−1 for reaction (1) and 443 ± 4 kJ mol−1 for reaction (2) (*40, 41*); this endoergicity can be compensated for by an energy transfer from the impinging electrons to **16** (*42*). Reactions (1) and (2) closely resemble the decomposition of methanol (**10**) that produces hydroxymethyl (ĊH2OH) and methoxy (CH3Ȯ) radicals (*24, 43*). Previous work by Song et al. revealed that reaction (3) is exoergic by 22 kJ mol−1 with an entrance barrier of 106 kJ mol−1 (1.1 eV) calculated at the MRCISD+Q/cc-pVTZ//CASSSCF/cc-pVTZ level of theory (*44*). Note that gas-phase calculations may overestimate the real barrier of the reactions because nearby molecules in the ice phase could be involved in the reaction mechanism (*45, 46*). Furthermore, the suprathermal hydrogen atoms have excess kinetic energies of a few eV (*47*); this energy can be utilized to overcome the entrance barrier of reaction (3) leading eventually to **11** (*35*) as identified through FTIR spectroscopy at 1849 cm−1 for **11** (ν2;HOĊO) in irradiated CO2−HOCH2CH2OH ice and at 1811 cm−1 for **11-**13C (ν2;HO13ĊO) in irradiated 13CO2−HO13CH213CH2OH ice.

HOCH2CH2OH (**16**) → HOĊHCH2OH (**17**) + Ḣ (1)

HOCH2CH2OH (**16**) → ȮCH2CH2OH (**20**) + Ḣ (2)

CO2 + Ḣ → HOĊO (**11**) (3)

*Second*, the preparation of **1** and **18** relies on barrierless radical-radical recombination of **11** with **17** (reaction (4)) and **20** (reaction (5)) radicals, respectively. Recall that the TPD profile at *m/z* = 106 (C3H6O4+) in CO2–HOCH2CH2OH ice shifts 4 amu to *m/z* = 110 (C3H6O218O2+) in C18O2–HOCH2CH2OH ice and 5 amu to *m/z* = 111 (C3HD5O4+) in CO2–HOCD2CD2OH ice (Fig. 4) indicating the formation of **1** or **18** involves one carbon dioxide and one **16** molecule.

HOĊO (**11**) + HOĊHCH2OH (**17**) → HOCH2CH(OH)COOH (**1**) (4)

HOĊO (**11**) + ȮCH2CH2OH (**20**) → HOCH2CH2OCOOH (**18**) (5)

**Discussion**

In conclusion, this study presents an abiotic route to a key biomolecule glyceric acid (**1**) — the simplest sugar acid — via barrierless radical-radical recombination of **11** with **17**, providing crucial steps toward a systematic understanding of how sugar acids can be formed in carbon dioxide-containing interstellar ices. **1** and its isomer 2-hydroxyethyl hydrogen carbonate **(18**) were formed in interstellar model ices of carbon dioxide and ethylene glycol upon exposure to ionizing radiation at low temperatures of 5 K. Utilizing tunable photoionization reflectron time-of-flight mass spectrometry (PI-ReTOF-MS) along with isotopic labeling experiments, **1** and **18** were detected in the gas phase during the TPD. The overall reaction energy leading to **1** from carbon dioxide and **16** (reactions (1), (3)and(4)) is endoergic by 35 kJ mol−1 as compensated by the kinetic energy of the incident electrons (*41, 48*), thus highlighting the necessity of non-equilibrium chemistry in its formation. Carbon dioxide and **16** are abundant in the ISM and have both been detected in molecular clouds such as Orion-KL — the star-forming region of the Orion Nebula (*25, 49*), the presence of **1** and **18** in interstellar environments is therefore plausible. Thus, the hitherto astronomically unobserved **1** and **18** represent promising candidates for future astronomical searches via gas-phase rotational emissions exploiting telescopes such as the Atacama Large Millimeter/submillimeter Array (ALMA). Once formed within interstellar ices in cold molecular clouds, **1** and **18** can be incorporated into accreting planetoids, asteroids, and comets (*50*). Ultimately, at least a fraction of these molecules can be delivered to planets like the early Earth and act as an exogenous source of biomolecules and their feedstocks (*6*). In fact, detailed analyses of meteorites such as Murchison revealed that not only sugar-related organic compounds including **1**, but also amino acids and dipeptides can be embedded in meteoritic matter and survive the harsh conditions in space and impact on the Earth (*1, 51, 52*). Therefore, a firm detection of **1** in the ISM in conjunction with rigorous modeling studies of reaction pathways to form prebiotic biorelevant molecules (Fig. 1) in deep space is of particular importance to our understanding of the role of sugar acids for the molecular complexity and synthesis of key organic molecules linked to the Origin of Life (*35, 53, 54*).

**Materials and Methods**

**Experimental**. Experiments were carried out in a hydrocarbon-free stainless steel chamber under ultrahigh vacuum conditions maintained at pressures of a few 10−11 Torr by magnetically levitated turbomolecular pumps (Osaka, TG1300MUCWB and TG420MCAB), which are backed by a dry scroll pump (XDS35i, BOC Edwards) (*55*). A polished silver substrate (12.6 mm × 15.1 mm) for ice deposition was attached to a two-stage closed-cycle helium refrigerator (Sumitomo Heavy Industries, RDK-415E) that can be freely rotated and translated vertically. The chemical samples used in the experiment were carbon dioxide-18O2 (C18O2, Sigma-Aldrich, 95 atom % 18O), carbon dioxide-13C (13CO2, Sigma-Aldrich, 99 atom % 13C), carbon dioxide (CO2,Airgas, 99.999%), deuterated ethylene glycol-d4 (HOCD2CD2OH, CDN isotopes, 98.9 atom % D), ethylene glycol-13C2 (HO13CH213CH2OH, Sigma-Aldrich, 99 atom % 13C) and ethylene glycol (HOCH2CH2OH, Sigma-Aldrich, anhydrous, 99.8%). Ethylene glycol sample was stored in borosilicate vial that was connected to a high vacuum chamber at pressures of a few 10−8 Torr and subjected to several freeze-pump-thaw cycles to remove residual atmospheric gases. Ices were prepared by passing carbon dioxide gas and ethylene glycol vapor through separate glass capillary arrays directed at the wafer cooled to 5.0 ± 0.3 K. Partial pressures were 2 × 10−8 Torr for carbon dioxide and 4 × 10−9 Torr for ethylene glycol. Ice thickness was measured during the deposition via laser interferometry (*56*); a photodiode was used to record variations in the intensity of a helium-neon laser (CVI Melles Griot, 25-LHP-230; 632.8 nm) reflected from the ice and silver substrate resulting from thin-film interference. The concentration-weighted average index of 1.32 ± 0.09 was used to derive the thickness of the carbon dioxide−ethylene glycol ices from the refractive indexes of amorphous carbon dioxide ice (n = 1.21) at 10 K (*57*) and that of ethylene glycol ice (n = 1.43) (*58*). Ice thicknesses were determined to be 750 ± 80 nm (table S6) by taking into account the densities of carbon dioxide (0.98 g cm−3) (*57*) and ethylene glycol (1.1 g cm−3) (*59*). To monitor the changes of ices *in situ* before and after the deposition, a Fourier transform infrared (FTIR) spectrometer (Thermo Electron, Nicolet 6700) was utilized in the range of 6000 − 500 cm−1 with 4 cm−1 spectral resolution. The ratio of carbon dioxide to ethylene glycol in the ice was determined to be 2.1 ± 0.7:1 using the integrated infrared absorptions of carbon dioxide at 3701 cm−1 (ν1 + ν3, 1.8 × 10−18 cm molecule−1) and 2278 cm−1 (ν3(13CO2), 6.8 × 10−17 cm molecule−1) (*27*) and the absorption bands of pure ethylene glycol ice with known thickness (fig. S9).

After the deposition, the ice mixtures were irradiated with 5 keV electrons (SPECS, EQ PU-22) at a 70° angle of incidence for 60 minutes at a current of 30 nA. Based on Monte Carlo simulations performed with the CASINO software suite (*60*), the irradiation conditions correspond to doses of 1.8 ± 0.3 eV molecule−1 for carbon dioxide and 3.1 ± 0.5 eV molecule−1 for ethylene glycol, respectively, simulating secondary electrons generated in the track of GCRs in cold molecular clouds aged around 5 × 106 years (*22*). The average penetration depth of electrons in CO2−HOCH2CH2OH ice was calculated to be 300 ± 60 nm based on Monte Carlo simulations using CASINO 2.42 (*60*). The penetration depth is notably less than the ice thickness (750 ± 80 nm), preventing electron-initiated interactions between the ice and the silver substrate. The changes in the spectrum of the ices were monitored by the FTIR spectrometer before, during, and after irradiation. After irradiation, the ices were subjected to temperature-programmed desorption (TPD) and heated from 5 K to 320 K at 0.5 K minute−1. Subliming molecules were photoionized 2 mm above the ice surface by pulsed 30 Hz vacuum ultraviolet (VUV) light at 11.10 eV or 8.77 eV, which was generated through resonant four-wave mixing (ωVUV = 2ω1 ± ω2) schemes using Xenon as a nonlinear medium (table S7). The VUV photons were generated via sum frequency generation (2ω1 + ω2; 11.10 eV) and difference frequency generation (2ω1 − ω2; 8.77 eV) with the 249.628 nm (ω1; dye laser, Sirah Lasertechnik, Cobra-Stretch) and 1064 nm (ω2; Nd:YAG laser, Spectra-Physics, Quanta Ray PRO 270-30) (*12*). A biconvex lithium fluoride lens (Korth Kristalle, R1 = R2 = 131 mm) was used in an off-axis geometry to spatially separate the VUV light from other laser beams. The ions resulting from VUV photoionization were mass-analyzed via reflectron time-of-flight mass spectrometry (ReTOF-MS) and detected with a dual microchannel plate (MCP) detector in the chevron configuration (Jordan TOF Products). The MCP signal was amplified with a preamplifier (Ortec, 9305), discriminated (Advanced Research Instruments Corp., F100-TD) and recorded by a multichannel scaler (FAST ComTec, MCS6A, 30 Hz). For each recorded mass spectra, the accumulation time of ion signals was 2 minutes (3600 sweeps) and the ion arrival time was recorded to 3.2 ns accuracy. Recorded TPD profiles were corrected for variations in the VUV flux throughout each experiment, which was monitored during TPD via a faraday cup (*61*). Additional experiment was also performed without irradiation (blank) at 11.10 eV for carbon dioxide−ethylene glycol ice, and no ion signal at *m/z* = 106 was observed.

**Computational**.The adiabatic ionization energies (IEs) and relative energies are computed as zero-point vibrational energy (ZPVE) corrected adiabatic differences in the energies of the neutral and radical-cation geometries. B3LYP/aug-cc-pVTZ was used to optimize the geometries and compute the harmonic vibrational energies in Gaussian16 (*62-66*). Explicitly correlated coupled-cluster singles, doubles, and perturbative triples (CCSD(T)-F12b) employing the cc-pVTZ-F12 basis set as available within MOLPRO 2020.1 program (*67-71*) produced single-point energies of C3H6O4 isomers at the optimized B3LYP/aug-cc-pVTZ geometries. These CCSD(T)-F12b energies along with the B3LYP ZPVEs produce the IEs and relative energies (table S5). The CCSD(T)-F12b/cc-pVTZ-F12 single points energy of PBE0/aug-cc-pVTZ and ωB97XD/aug-cc-pVTZ optimized structures with anharmonic vibrational analysis at the DFT level were calculated for **1** and **18**; the shifts in IE and relative energy are shown in table S8. This approach has demonstrated a good correlation with experiments in previous work (*72*). The computed Cartesian coordinates and harmonic and anharmonic vibrational frequencies are listed in table S9.

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**Competing interests:** The authors declare that they have no competing interests.

**Data and materials availability:** The data that support the findings of the current research are provided in the paper and/or the Supplementary Materials.

**Figures**

A diagram of chemical formulas

Description automatically generated

**Fig. 1.** **Formation of glyceric acid in interstellar ices.** The preparation of glyceric acid (**1**) in low-temperature ices containing carbon dioxide and ethylene glycol (**16**) is accomplished through energetic processing by galactic cosmic ray (GCR) proxies. This process involves carbon-carbon bond coupling via recombination of the hydroxycarbonyl radical (HOĊO, **11**) with the 1,2-dihydroxyethyl radical (HOĊHCH2OH, **17**). Glyceric acid (**1**) serves as a precursor for critical biomolecules including the proteinogenic amino acid serine (**4**), 2-methyl glyceric acid (**7**), and lactic acid (**15**). In contemporary biochemistry, glyceric acid further represents molecular building blocks of 2-phosphoglyceric acid (**2**) and 3-phosphoglyceric acid (**3**) via phosphorylation reactions, linking to the tricarboxylic acid cycle (top right) and Calvin cycle (bottom right), respectively.

A diagram of a molecule

Description automatically generated

**Fig. 2.** **Reaction scheme leading to three C3H6O4 (*m/z* = 106) isomers (1, 18, 19) in irradiated carbon dioxide–ethylene glycol ices.** Barrierless radical-radical reactions **11** plus **17** and **11** plus **20** produce **1** and **18**, respectively; tautomerization of **1** may lead to the enol **19**. The adiabatic ionization energies (IEs) are computed at the CCSD(T)-F12b/cc-pVTZ-F12//B3LYP/aug-cc-pVTZ level of theory including zero-point vibrational energies and are corrected by incorporating error (table S4). The insert (bottom right) compiles the computed IEs of isomers (black solid line) and ranges of the conformers (grey area) after error analysis. VUV photon energies (dashed lines) at 11.10 eV and 8.77 eV were used to photoionize subliming molecules during the TPD process.

A group of graphs showing different types of data

Description automatically generated with medium confidence

**Fig.** **3.** **PI-ReTOF-MS data as a function of temperature and mass-to-charge (*m/z*) ratios during the TPD of the carbon dioxide–ethylene glycol ices**. Data were recorded for the unirradiated (blank) CO2–HOCH2CH2OH ice at 11.10 eV (**A**), the irradiated (30 nA, 60 minutes) CO2–HOCH2CH2OH ice at 11.10 eV (**B**) and 8.77 eV (**F**), the irradiated C18O2–HOCH2CH2OH ice at 11.10 eV (**C**), the irradiated CO2–HOCD2CD2OH ice at 11.10 eV (**D**), and the irradiated 13CO2–HO13CH213CH2OH ice at 11.10 eV (**E**).

A graph of a chemical reaction

Description automatically generated with medium confidence

**Fig. 4.** **Ion signal during TPD of irradiated carbon dioxide–ethylene glycol (16) ices shown as a function of temperature.** (**A**) TPD profiles measured at 11.10 eV with irradiated (30 nA, 60 minutes) CO2–HOCH2CH2OH ice (*m/z* = 106), 13CO2–HO13CH213CH2OH ice (*m/z* = 109) and C18O2–HOCH2CH2OH ice (*m/z* = 110); (**B**) TPD profiles measured at 11.10 eV at *m/z* = 109 and *m/z* = 111 in irradiated (30 nA, 60 minutes) CO2–HOCD2CD2OH ice; (**C**) TPD profiles measured at 8.77 eV at *m/z* = 106 in CO2–HOCH2CH2OH ice irradiated with a low dose (30 nA, 60 minutes) and a higher dose (100 nA, 120 minutes).

**Supplementary Materials**

**This PDF file includes:**

Tentative assignments of C2H4O3 and C3H6O3 isomers

Figs. S1 to S9

Tables S1 to S9