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Graph-Based Analysis of Ethylene Glycol Decomposition on a Palladium Cluster

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ABSTRACT: The ethylene glycol, CH₂OH–CH₂OH, decomposition mechanism, occurring on a subnanometric palladium cluster shaped by 12 atoms, was investigated by means of density functional theory. Different reaction routes were identified leading to H₂ and CO. The whole reaction network was analyzed, framing the results within the graph theory. The possible decomposition pathways were discussed and compared, allowing one to draw a whole picture of all the parallel, possibly competitive, routes that starting from CH₂OH–CH₂OH originate H₂ and CO.

INTRODUCTION

Substrates recognizable as sustainable source of carbon and deriving from biomass raw materials have attracted in the past few years the attention of the scientific community. The interest was determined also thanks to the possibility to produce both environment-friendly hydrogen as well as chemicals of potential interest.1–4 Generally, biomass-based feedstocks contain large amount of oxygenates, and these compounds mainly steer the reactivity of these processable materials. Studies on biomass conversion, demonstrated the efficacy of Pt- and Pd-supported catalysts, in the reforming processes hence in the consequent production of hydrogen.5–7 Generation of H₂ from oxygenates proceeds through a combination of C–C bond scissions and dehydrogenation reactions to form hydrogen and carbon monoxide on the catalyst surface. The total amount of obtainable hydrogen is generally increased, coupling the reforming with the water gas shift reaction.

In order to maximize the production of H₂, the control of C–C bond cleavage is highly advantageous. This means that all the side reactions deriving from the C–O bond scission have to be suppressed to avoid alkane formation while dehydrogenation and decarbonylation should be promoted.8,9 Various exper-imental and theoretical studies concerning the decomposition of alcohols and polyols10 on Pd surfaces are reported in the literature.11–19 As an example, Shekar and Barteau found that ethanol decomposes on both Pd(111) and (110) surfaces via dehydrogenation and decarbonylation steps, without any evidence of C–O scissions.20 Alcohol decomposition on the Pt-group metals was believed to start by the O–H bond breaking, which would form an alkoxide intermediate.21 However, according to more recent studies, the favored cleavage should be that involving the C–H bond.22 In agreement with these experimental evidence, Li et al.,23 investigating the ethanol decomposition on Pd(111) by means of periodic DFT calculations, found that the mechanism involving a C–H breaking as first stage should be preferred while decarbonylation would seem to be favored if compared to C–O bond breaking.

Yudanov and co-workers also focused on C–O bond cleavage on Pd nanoparticles, identifying it as a slow side process in the decomposition of methanol.24 Sutton and Vlachos furthermore suggested, according to the results of their periodic DFT studies, that ethanol activation is remarkably consistent on metal surfaces, the initial step being either an O–H breaking (on Co, Ni, Rh, and Ru surfaces) or a C–H scission (on Pd and Pt surfaces). The initial activation is then followed by a series of relatively facile dehydrogenation reactions until C–C cracking occurs. The more oxophilic is the metal (e.g., Co and Ru), the more ethane is produced, but it is likely only a minor product.25

Although important advances in understanding the chemistry of several biomass-model molecules on idealized metal surfaces were made, the impact of low coordinated sites26,27 and of characteristic geometric and electronic features caused by the subnanometric sizes of the catalyst received little attention. Since these aspects could give rise to variations in the reaction mechanism, as reported by Mehmood et al.,28 in a study dealing with the methanol decomposition on small Pd clusters, it is very useful to rationalize the effect of the sizes of the catalyst.
that indeed might result in different catalytic activities and selectivities.

In this study, in view of disclosing elementary processes involved in the reformation of light polyols, ethylene glycol (CH₂OH–CH₂OH) was selected as a multifunctional oxygenated model species, and its decomposition on a subnanometric Pd cluster was investigated by means of DFT. Studies dealing with the reactivity of this molecule are of interest both to enhance practical goals of hydrogen and syngas production and to provide fundamental insights into the underlying competition among C−H, O−H, C=C, and C−O bond cleavages in polyols.

The size of the catalytic system above, involving a small Pd cluster and simple C₂ species, allowed one to explore by computational techniques almost all the reactive routes associated with the ethylene glycol decomposition. This determined the generation of a large grid of possible events, useful to deepen the energetic ordering of different reaction pathways that in principle would let to isolate the whole reaction mechanism, involving many and complexly tangled molecular events. So, to effectively analyze the decomposition path, it is necessary to accurately characterize not only the elementary steps involved in the reaction network but also their hypothetical sequences, in the large ensemble of their possible combinations. For this purpose, within a graph theory (GT) based approach, in this work, all the pathways starting from CH₂OH=CH₂OH and leading to CO and H₂ were analyzed so as to draw a picture of the competitive routes leading to hydrogen production on a subnanometric Pd cluster.

MODELS AND COMPUTATIONAL DETAILS

All the reported calculations were performed within the density functional theory employing the Gaussian09 code. As exchange-correlation functional the M06L developed by Zhao et al. was selected since it was explicitly parametrized to have good performance for both main-group elements and transition metal chemistry. For the Pd atom the cc-pVDZ-PP effective core potential joined with its associated basis set was employed, whereas for the H, C, O atoms, the DZVP2 one was selected.

It was already shown that the fragmentation of hydrogen on a subnanometric palladium cluster lowers the number of unpaired electron of the whole system, therefore, since the number of hydrogen atoms residing on the cluster is variable depending on the progress of the decomposition reaction, the geometries of the reagent, intermediate, product and transition state (TS) species were optimized with three different spin multiplicities (three, five, and seven). The search of TSs was performed with the Berny algorithm and all the stationary points were characterized by inspecting the vibrational frequencies calculated within the harmonic approximation. All the energy values reported include the vibrational zero point contribution and, where adsorption energies are discussed, the BSSE correction according to the counterpoise method was always applied. Activation barrier for a given elementary step was finally calculated taking as reference state the intermediate species preceding the TS associated with the considered molecular event.

A Pd₁₂ cluster was selected as the catalyst. This was obtained following a growing process applied to a Pd₉ cluster, which already was extensively studied with respect to its electronic, structural and catalytic features. In details, to the Pd₉ cluster, characterized by D₅h symmetry and quintet multiplicity, were added three Pd atoms in the convex faces then reoptimized. The most stable form of the resulting Pd₁₂ had geometrical properties sensibly changed with respect to those of the initial guess, showing D₂d symmetry and septet spin multiplicity. Pd₁₂ has three nonequivalent sites, with six (6c), five (5c) and four (4c) nearest neighbors, respectively, in turn delineating two triangular shaped hollows, namely H₁(5c,4c,6c) and H₂(5c,6c,6c). The cluster structure and the nonequivalent sites are shown in Figure 1.

Figure 1. Pd₁₂ cluster structure. The three nonequivalent sites having coordination number six (Pd⁶c), five (Pd⁵c), and four (Pd⁴c) are pinpointed.

The whole reaction mechanism was framed and analyzed within a graph theory approach, so according to a GT formalism, nodes and edges of the graph were defined as the a given species to another one respectively. When the C−C bond is present, an intermediate species is named reporting first the node number that identifies its position in the graph, followed by the type of cleavage involved in its formation: C and O individuate C=H and O=H bond cleavage, respectively, whereas C−O does the same for the C−O bond cleavage. The node numbers are not connected with a decomposition process, as the former are randomly attributed. On the contrary, the sequence of capital letters, present in the acronym of a given intermediate species, identifies the number of elementary steps as well as the sequence of reactions necessary to produce that species. When the C−C bond is not present in the intermediate, the node number is followed by the produced fragments, closed in brackets and separated by a vertical bar.

It is important to underline that, molecular diffusion on the cluster has to occur for the fulfillment of some surface events. However, considering the very low activation barrier of these events with respect to the other elementary steps considered in this investigation, it was assumed that diffusion could take place freely, therefore, this was not included within the graph-based analysis. In details, the protocol followed for the reaction network analysis can be divided into three steps: (i) ab-initio calculation of the activation barriers and preserving of the possible molecular events, (ii) organization of the graph representing the final reaction network, and (iii) analysis of the possible pathways. Preserving of the molecular events was ruled by the assumption that one activation barrier related with other activation barriers and exceeding, one of the latter by ca. 30 kJ mol⁻¹ correspond to a non competitive process. This assumption, which is justified considering the negative exponential dependence between the activation barrier and the kinetic constant of a given process, actually allowed us to exclude a lot of processes, enormously decreasing the involved intermediates and molecular events leading from...
connections number of the graph network. Once the events considered as feasible were isolated, the final graph was built. The analysis of the resulting network was performed considering that, as stated in the frame of the GT approach, a walk is an alternating succession of vertices and edges that starts with a vertex and ends with a vertex and a path is a walk in which no vertex occurs more than once. By use of a homemade python-based routine, forty-one simple paths of different length connecting the reactant and the product were found. The different length pathways found were ordered, taking into account the total energy content (TEC), defined as the summation of all the activation barriers involved in the development of a given route.

RESULTS AND DISCUSSION

The ethylene glycol decomposition mechanism on a subnanometric cluster was investigated, involving dehydrogenation and both C–C and C–O bond cleavages. Atomic hydrogens produced along with the dehydrogenation process were assumed to easily diffuse on the Pd$_{12}$ cluster$^{31}$ hence every time a couple of chemisorbed hydrogen atoms was produced, it was removed from the metal cluster hypothesizing an easy desorption following a quick meeting of the same surface species. All the elementary reactions are in the following commented, highlighting which among the considered processes is or is not included in the final graph.

Devising the Graph: Description of Nodes and Edges.

Adsorption of C$_2$ Species on Pd$_{12}$. Ethylene glycol adsorbed on the palladium cluster is identified by the N0 node of the graph. This is the source vertex used to evaluate all the simple paths, which characterize the whole ethylene glycol decomposition network. It is well-known that although twenty-seven ethylene glycol conformers exist the representative conformational space of the species is actually formed by just ten structures.$^{36}$ According to Prasanta et al.$^{37}$ the representative ensemble ranges in-between an energy span of ca. 15 kJ mol$^{-1}$. In order to investigate the adsorption process of the CH$_2$OH–CH$_2$OH molecules on the Pd$_{12}$ cluster it was decided to consider only one conformer per each of the two limit staggered configurations (i.e. gauche and anti): the first (C$^9_2$) retains the intramolecular hydrogen bond while the other (C$^{*}_2$) not. For the sake of comparison, in vacuum, C$^9_2$ is 10.7 kJ mol$^{-1}$ more stable than C$^{*}_2$, at the here level of theory while Prasanta et al., which employed the B3LYP exchange-correlation functional joined with the 6-311++G** basis set, calculated an energy difference for these two conformers (Gg’ and tT in the cited work) of 9.0 kJ mol$^{-1}$. When approaching the cluster, the hydroxyl groups of the CH$_2$OH–CH$_2$OH species might interact with one or more among three nonequivalent atop sites. Thus, to analyze the adsorption process, the C$^9_2$ species was placed with one –OH group pointing, in turn, to a 4c, 5c, and 6c site (see Figure 1). Among these configurations, that involving the 5c site resulted to be the most stable (those involving 4c and 6c sites actually were 5.7 and 32.9 kJ mol$^{-1}$ higher in energy). Moreover, the adsorption energies of the C$^9_2$ and C$^{*}_2$ conformers on the 5c site were found to be –83.4 and –64.5 kJ mol$^{-1}$, respectively. The activation barrier for the conversion of the gauche conformer into the anti one was calculated, showing that the process on the Pd$_{12}$ cluster requires at least 36.2 kJ mol$^{-1}$ and that C$^9_2$ is 29.7 kJ mol$^{-1}$ less stable than C$^{*}_2$, very likely because of the breaking of an intramolecular hydrogen bond in the former surface system. The activation barrier associated with the rotation on Pd$_{12}$ hence is, in someway, affordable therefore it is possible to infer that the gauche conformer could transform into the anti one. This could be important because the possible occurrence of two different conformations would guarantee the proximity to the surface catalyst of different fragments of the same polyol. As a consequence, different molecular events might occur, starting from the two different geometrical arrangements.

CH$_2$OH–CH$_2$OH Species: Node N0. Being aware that for highly hydrogen-saturated species the C–C bond scission is quite unlikely, this elementary step was not considered in the analysis of the initial stages of the reaction mechanism. Thus, due to the specific symmetry of CH OH–CH OH just three bond breaking events involving C–H, O–H and C–O have to be considered. All the intermediates formed in these processes are reported in Figure 2. It was found that the C–H

![Figure 2](image-url)
required to produce N3−C−O from N0 equal to 202.5 kJ mol⁻¹.

The results above confirm that dehydrogenation on palladium is more feasible than C−O cleavage, being the formation of alkoxide species favored. The results here reported are in line with previous experimental investigations, pointing out that for the transition metals belonging to the platinum-group the dissociative adsorption of alcohols leading to alkoxides is almost ubiquitous. Nevertheless, for the sake of completeness, it should be recalled that in a previous theoretical study, dealing with decomposition of alcohols on Pd(111), it was reported that the first step, occurring along with reforming, is the cleavage of C−H, being the dehydrogenation of the hydroxyl fragment at all not competitive. In this study, Li et al. for the ethanol C−H cleavage on a smooth Pd surface calculated an activation barrier value comparable to the one here reported i.e., 113.4 vs 116.4 kJ mol⁻¹ whereas for the O−H breaking the two outcomes were very different. For the subnanometric cluster, the activation barrier is indeed sensibly lower i.e., 84.1 vs 160.2 kJ mol⁻¹. Despite the different substrates and theoretical approaches used to get them, it is reasonable to infer, considering the large disagreement found, that there could exist a connection between the subnanometric size of the metal particle and the easier hydroxyl dehydrogenation. In addition, a similar trend was also found in our recent work, dealing with Pd₆-catalyzed formic acid decomposition: in particular, it was demonstrated that the activation barrier for the O−H bond breaking is considerably lower on subnanometric clusters than on a Pd(111) smooth surface. On the contrary the activation barrier for the C−H bond cleavage was not affected by the size and shape of the catalyst.

Comparing the activation barrier values characterizing the processes discussed above, it is possible to state that the steps leading to N1−C and N3−C−O are not competitive and have to be excluded from the reaction network landscape, leaving only the N2−O transformations as those to be further analyzed.

CH₂OCH₂OH Species: Node N2. Since N2−O shows two non equivalent C−H bonds, it is possible to hypothesize three dehydrogenation steps (C−H₁, C−H₂, O−H) as well as one C−O cleavage. Atom numbering and intermediates produced in these molecular events are reported in Figure 2. The process that involves the C−H₁ bond cleavage has an activation barrier of 34.7 kJ mol⁻¹ and produces a glycolaldehyde molecule, in the following identified as N₄−OC₁. This species interacts with the Pd₁₂ cluster via the oxygen of the carbonyl moiety in a η₁ configuration. The occurrence of this binding mode (determining a calculated O−Pd distance equal to 2.17 Å) was already inferred by Manus and colleagues who, analyzing a HREEL vibrational spectrum of glycolaldehyde adsorbed on Pd(111), reported that the ν(CO) stretching at 1715 cm⁻¹ is slightly perturbed with respect to the one of the gaseous species. There is convergence of results also regarding the reactivity of this species. In the study above it was, in fact, hypothesized that

Figure 3. Schematic mechanism including the geometries of the intermediates deriving from the N6-OO species.
The aldehydic species in order to react has to change the binding mode from $\eta_1$ to $\eta_2$. In agreement, this binding mode change was also observed, in the present work, for the transition state leading from N4–OC1 to N8–OOC$^*$. The activation barrier for the rupture of the C–H2 bond to form N5–OC2 resulted 99.2 kJ mol$^{-1}$; being the produced surface species chemisorbed via two stronger, C–Pd$^{6c}$ (2.09 Å), O–Pd$^{5c}$ (2.07 Å), and one weaker, O$_2$–Pd$^{6c}$ (2.65 Å), interactions. The minimum amount of energy required to break the second O–H bond, originating the ethylenedioxy species, N6–OO, was equal to 75.6 kJ mol$^{-1}$. The binding mode of this dioxy-species involves interactions of both the oxygen atoms with the metallic cluster, one simple (O–Pd$^{5c}$ 1.98 Å) and one having a bridge arrangement (O–Pd$^{5c}$ 2.21 and O–Pd$^{6c}$ 2.16 Å). This result is in agreement with Griffin et al., whose experimental studies on the reforming of CH$_2$OH–CH$_2$OH on Pd(111), suggested that this process is always characterized by the occurrence of surface species interacting by both the oxygen atoms. For the sake of completeness, it should be said that the previous species, in accordance with the solely energetic criterion should have been excluded because of the limit of the 30 kJ mol$^{-1}$; however, considering the experimental findings above and the fact that the barrier exceeds of few kJ mol$^{-1}$ the limit, the further decomposition of N6–OO was in any case investigated. The C–O cleavage of N2–O was also analyzed and the corresponding activation barrier, producing N7–OC–O, resulted 181.4 kJ mol$^{-1}$. In agreement with this energy value, the C–O rupture on Pd catalysts is always reported in the literature as highly unfavorable with respect of both the dehydrogenation and the C–C scission. As a consequence, the C–O rupture event was not further investigated to characterize the title reaction mechanism.

In summary, after the first dehydrogenation, which leads to the formation of an alkoxide species, the reaction mechanism shows a bifurcation, generating two branches that originate from ethylenedioxy (see Figure 3) and glycolaldehyde (see Figure 4) species. For sake of clarity, at first all the elementary steps deriving from the N6–OO species are, in the next eight sections, discussed and then, in the following, those related to the N4–OC1 intermediate.

CH$_2$OCH$_2$O Species: Node N6. Given the high symmetry of the ethylenedioxy molecule only the C–C and one C–H bond rupture were considered as possible. To break the C–H bond, however, a local diffusion step of N6–OO is necessary. This guarantees the right arrangement of the surface ethylenedioxy on the cluster, see the N6–OO(a) → N6–OO(b) step in Figure 3. Associated to the rearrangement an activation barrier of 17.8 kJ mol$^{-1}$ was found. The corresponding TS shows a change in the binding mode of one of the two oxygens involved, that proceeds from a bridge surface interaction to an atop one. The right orientation could be actually reached also by rotating of 180° the OCCO torsion angle. This process, however, is well-known to be highly unfavored, since a strong...
Once N6–OO reached by diffusion the correct arrangement to break the C–H bond, the minimum amount of energy necessary to cleave it and produce N8–OOC resulted 35.3 kJ mol$^{-1}$. This species interacts with the cluster via one Pd–O bond bridge (O–Pd$^{6c}$ 2.25 Å, O–Pd$^{6c}$ 2.13 Å), and one atop, C–Pd$^{6c}$ bond (2.44 Å). Finally, the energy barrier to get two formaldehyde molecules, N9(CH$_2$O)/CH$_2$O, from N6–OO by means of a C–C cleavage, was found to be equal to 50.1 kJ mol$^{-1}$. According to the energetic range into which the activation barriers so far reported are included, all these processes were considered as feasible, and therefore further analyzed for unravelling the title species reaction network.

**CHOCH$_2$O Species: Node N8.** For the N8–OOC intermediate two nonequivalent C–H ruptures were evaluated. One of them led to glyoxal, N12–OOC$^*$, that showed a binding mode, involving four links with the Pd$_{12}$ cluster. Among these, two were formed by both the carbon atoms with the same 6c palladium site (C–Pd$^{6c}$ 2.34 and 2.45 Å) and two by both the oxygen atoms, which singly were connected to a 5c and a 6c site (O–Pd$^{6c}$ 2.17 Å, O–Pd$^{6c}$ 2.21 Å). This process was related to a small activation barrier, namely 16.6 kJ mol$^{-1}$. The other C–H cleavage produced N13–OCCC, an intermediate characterized by one Pd–O–Pd bridge (O–Pd$^{5c}$ 2.16 Å, O–Pd$^{6c}$ 2.20 Å) and one terminal CO moiety bridging two Pd atoms (C–Pd$^{5c}$ 2.14 Å, C–Pd$^{6c}$ 2.10 Å). The activation barrier for producing the N13–OCCC intermediate was equal to 32.3 kJ mol$^{-1}$. Further decompositions were investigated for both the intermediates formed through the two dehydrogen-ations just discussed.

**CH$_2$O + CH$_2$O Fragments: Node N9.** Both the formaldehyde molecules show the carbonyl moiety interacting with a 6c and a 5c site of the Pd cluster (O–Pd$^{6c}$ 2.09 Å, C–Pd$^{5c}$ 2.15 Å). This binding mode is very similar to that characterizing the interaction geometry of a single CH$_2$O molecule adsorbed on a smooth Pd(111) surface, as reported by Lim et al. (O–Pd 2.11 Å, C–Pd 2.15 Å). Considering the molecular geometry of the CH$_2$O couple as a reasonable descriptor of lateral interactions, it is possible to argue that the coadsorbed CH$_2$O fragments slightly influence each other. The cleavage of the first C–H bond to form a CHO species, N10(CH$_2$O)(CHO), requires the overcoming of an activation barrier of 43.2 kJ mol$^{-1}$. The structural rearrangements occurring during the dehydrogenation do not affect the binding mode of the other CH$_2$O molecule placed on the Pd cluster, confirming as not influential the lateral interactions. The bond between the formyl species and the Pd cluster is characterized by two different bonds: one C–Pd in a bridge arrangement (C–Pd$^{4c}$ 2.03 Å, C–Pd$^{6c}$ 2.03 Å) and one atop O–Pd (O–Pd$^{5c}$ 2.20 Å).

**CH$_2$O + CHO Fragments: Node N10.** The second consecutive C–H rupture shows an activation barrier of 9.9 kJ mol$^{-1}$ and produces a coadsorbed CO molecule, N11–(CH$_2$O)(CO). According to Lim et al., a formaldehyde H abstraction on a Pd(111) surface requires an activation barrier of ca. 38 kJ mol$^{-1}$ while the dehydrogenation of the formyl species of ca. 60 kJ mol$^{-1}$. The findings here presented would suggest an opposite behavior, however, it is hard to state if this has to be related to the Pd cluster size or to other more subtle reasons.

**CH$_2$O + CO Fragments: Node N11.** The CH$_2$O dehydrogenation to initially produce a CHO species coadsorbed with a carbon monoxide molecule already present on the palladium cluster, N16(CH$_2$O)(CO), and then a second coadsorbed CO molecule, N26(CO)(CO), were analyzed, finding the related activation barriers 40.4 and 13.2 kJ mol$^{-1}$, respectively.

**CHOCHO Species: Node N12.** As glyoxal is a symmetric molecule, only one C–H and one C–C bond breaking process was analyzed. Their activation barriers were 64.7 and 75.7 kJ mol$^{-1}$ for producing either N14–OOC$^*$GC or two CHO fragments coadsorbed on the cluster, N15(CH)(CHO), respectively. The N14–OOC$^*$GC species shows a terminal CO moiety bridging a 5c and a 6c site (C–Pd$^{5c}$ 2.10 Å, C–Pd$^{6c}$ 2.14 Å). On the contrary, the interaction geometry of the adsorbed formyl fragment is very similar to that previously reported for the decomposition of formaldehyde. Considering the values of the activation barriers calculated for these two molecular events, both the produced intermediates were further considered in the analysis.

**COCH$_2$O Species: Node N13.** The analysis of the N13–OCCC reactivity would require the investigation of both a decarbonylation process and of a C–H bond cleavage. The latter, however, was not evaluated for two main reasons: due to the COCH$_2$O surface arrangement (i) any among its C–H bonds was close enough to the cluster to be cleaved and (ii) the strong interaction with the metal sites did not allow any easy molecular rearrangements. The minimum amount of energy to form CO and formaldehyde coadsorbed on the Pd cluster, N11(CH$_2$O)(CO), starting from N13–OCCC was just 24.3 kJ mol$^{-1}$.

**CHOCHO Species: Node N14.** Starting from N14–OOC$^*$GC, the C–C and C–H bond cleavages were investigated. The C–C breaking, i.e., the decarbonylation process, showed an activation barrier of 28.2 kJ mol$^{-1}$ whereas the C–H hydrogen abstraction 68.8 kJ mol$^{-1}$. The former transformation produces CO coadsorbed with CHO, N16(CH)(CO), whereas the latter, the dehydrogenated N25–OOC$^*$CC fragment.

**CHOCH$_2$OH Species: Node N4.** For glycolaldehyde (N4–OC1), four molecular events were considered, namely C–H2, C–H1, C–C and O–H breaking (for the numbering of the hydrogen atoms, see Figure 4). The first three processes generating the species N17–OC12, N18–OC1C1 and N19(CH$_2$O)(CHO) showed activation barrier values equal to 45.7, 84.1, and 122.8 kJ mol$^{-1}$, respectively. The O–H scission can take place only if changes in the glycolaldehyde interactions with the cluster occur because its local configuration does not allow any interaction of the hydroxyl hydrogen atom with the metallic sites. This is likely due to the presence of an intramolecular hydrogen bond and to the structural rigidity caused by the carbonyl fragment interactions with the Pd cluster. The hydroxyl oxygen atom repositioning, from 6c to 5c site is on the other hand, a process requiring the overcoming of 26.2 kJ mol$^{-1}$ and giving a product essentially iso-energetic with the starting species. The new configuration of glycolaldehyde on Pd$_{12}$ shows the correct position for determining the O–H bond breaking, which actually occurs with an energy barrier equal to 73.4 kJ mol$^{-1}$. The intermediate obtained, N8–OOC$^*$, however, cannot undergo the C–H2 scission, since both its H atoms point outward the cluster. An additional diffusion process must be thus invoked. This shows an activation barrier of 31.8 kJ mol$^{-1}$ and converts N8–OOC$^*$ into N8–OOC. According to the activation barriers, further decomposition processes were investigated for the N17–OC1C2 intermediate and the connection between glycolaldehyde and the N8–OOC species, through a combined concurrence of diffusion steps and dehydrogenation, was considered as feasible.
Figure 5. Whole decomposition mechanism of CH$_2$OH$\rightarrow$CH$_2$OH to CO and H$_2$: points where one hydrogen molecule is assumed to leave the cluster are shown in squares; red arrows indicate pathways excluded during the presieving phase; energy barriers (expressed in kJ mol$^{-1}$) for bond scissions, formation and diffusions, in blue, refers to the reactant transformation. Dangling lines indicate bonds between fragments and Pd sites; interactions of chemically saturated atoms with the cluster are not shown.

CHOCHOH Species: Node N17. The intermediate produced from the C$\equiv$H bond cleavage, N17$\rightarrow$OC1C2, could be further decomposed by breaking C$\equiv$H1, C$\equiv$H2, C$\equiv$C, and O$\equiv$H bonds. The produced surface species and activation barriers are N20$\rightarrow$OC1C2C1, N21$\rightarrow$OC1C2C2, N22(CHOH(CHO)), and N12$\rightarrow$OCCg and 75.2, 107.1, 107.0, and 79.2 kJ mol$^{-1}$, respectively. In agreement with the values of the activation barriers found, only the conversion of N17$\rightarrow$OC1C2 either into N20$\rightarrow$OC1C2C1 or N12$\rightarrow$OCCg was further considered.

COCHOH and CO + CHOH Species: Node N20 and N23. The N20$\rightarrow$OC1C2C1 intermediate can undergo three different bond breaking events, namely C$\equiv$H2, C$\equiv$C, and O$\equiv$H, that lead to two new intermediates N24$\rightarrow$OC1C2C1C2, N23(CHOH(CHO)) and the already found N14$\rightarrow$OCCgC species. The related processes are characterized by activation barriers equal to 94.3, 74.1, and 57.9 kJ mol$^{-1}$. Because of these values, just the reactions leading to N23(CHOH(CHO)) and to N14+OCCgC should be practicable. Furthermore, the N23(CHOH(CHO)) through dehydrogenation could be converted into the already met N16(CHO(CO)) fragment, overcoming an activation barrier of 19.6 kJ mol$^{-1}$.

Reaction Network Analysis. The reaction mechanism for the CH$_2$OH$\rightarrow$CH$_2$OH decomposition is reported in Figure 5. The C$\equiv$O bond breaking seems to be, in any case, an energy demanding process, suggesting that Pd$_{12}$ is not active in the alkane formation. The O$\equiv$H bond scission shows an activation barrier between 70 and 80 kJ mol$^{-1}$, and is likely to occur in the early stages of the decomposition process; this indicates a tendency of the Pd$_{12}$ cluster to be selective toward the formation of carbonylic, instead of enolic, compounds. These findings are confirmed by Griffin et al. through HREELS and TPD studies, pointing out that formation and transformation of glyoxal are among the main processes involved in the CH$_2$OH$\rightarrow$CH$_2$OH decomposition on Pd(111) planes. The C$\equiv$H bond shows very different scission energy barriers, namely in the range 10$\rightarrow$110 kJ mol$^{-1}$, and it is a process that has the chance to occur at every step along the C$_2$ species decomposition. The activation barrier in the C$\equiv$C bond
cleavage usually decreases with decreasing the hydrogenation degree of the involved species; this phenomenon is increased if decarbonylation occurred. In agreement, comparing the C–C cleavage in N13–OOC and N12–OCCg, both having the same hydrogenation degree but involving and not involving decarbonylation activation barriers equal to 24.3 and 75.7 kJ mol\(^{-1}\) were found. Activation barrier of 28.2 kJ mol\(^{-1}\) was also found for N14–OCCgC when decarbonylated.

Once obtained the whole reaction network, the trends associated with the variation of the spin states characterizing the systems during the reforming processes were analyzed. The multiplicity of the system could change for the elementary steps composing the ethylene glycol decomposition; however, the energy differences associated with the three analyzed spin states, i.e., triplet, quintet, and septet, generally resulted but with some remarkable exceptions in being small and, as a consequence, the emerging picture is too complex to be described by simple schemata. Still, some recurrent patterns for the surface species multiplicity variations as a function of the chemisorbed hydrogen quota on the Pd\(_{12}\) cluster could be found.

In Table 1 are reported all the intermediates species, related to the number of chemisorbed hydrogens on the cluster. For sake of clarity, at first, it is useful to analyze the species, whose C–C bond had not yet been broken: (i) apart from a few cases, species where the cluster is not hydrogenated showed more stable septet states than either quintet or triplet ones, which is in agreement with the most stable septet state of naked Pd\(_{12}\); (ii) there are eight singly hydrogenated cluster systems and in most cases showed quintet as the most stable spin state, which is in agreement with the results of D’Anna et al.,\(^{31}\) showing a lowering of two units in the spin multiplicity when a dehydrogenation process occurs on a fragment in which the C–C bond is not yet cleaved; (iii) seven species with two chemisorbed hydrogens were analyzed, not showing a clear trend, so different spin states seem indeed to be favored. However, excluding N6–OO, having spin states very close in energy, the septet state is by far the less stable.

When the C–C bond is cleaved different coadsorbed unsaturated fragments on the cluster are formed, very likely inducing a spin state change of the systems. No simple trends, anyway, emerge by analyzing the results obtained for these species. However, when the C–C bond is cleaved, irrespective of the number of hydrogen atoms adsorbed on the Pd cluster, clearly it comes into view that triplet and quintet states are close in energy, being the former usually more stable than the latter, at variance with the septet that is less stable.

On the whole, 25 intermediates were characterized and thirty-one molecular events analyzed, resulting the final graph formed by 22 edges and 17 nodes. The intermediates omitted in the final graph are showed in Figure 5 as products of elementary steps pointed by red arrows while the connected undirected graph of the considered species is reported in Figure 6a.

There are several analyses that could be applied in order to extract information on the relevance of the nodes in a graph and, among these, the ones calculating the centrality indexes in GT are able to provide a node rank, identifying the most relevant in a considered context. Betweenness centrality (b–c), in particular, is representative of the node centrality in a given network. Here was calculated, by using the Brandes algorithm,\(^{43}\) as implemented in the Gephi program.\(^{44}\) The metric of the b–c parameter is defined by the number of shortest paths from any vertex to all the others, which cross a given node. A large b–c of a node actually indicates a substantial influence of it on the item transfer ability through the network. Therefore, when framed within a chemical context, even without energetic information, b–c could help to identify preliminarily key intermediates own of a reaction mechanism on the grounds of the solely connections between intermediates. The b–c of a given node, n, is

| Table 1. Ethylene Glycol Decomposition Species in Absence (0H) or in the Presence of One (1H) or Two (2H) Co-Adsorbed Hydrogen Atoms on the Cluster\(^{a}\) |
|-----------------|-----------------|
|                 | E(2S + 1) (kJ mol\(^{-1}\)) |     |
|                 | 3               | 5   | 7 |
| 0H              |                 |     |   |
| N0              | 22.5            | 9.1 | 0.0 |
| N3–C–O         | 10.5            | 0.0 | 1.2 |
| N4–OC1         | 73.0            | 42.0| 0.0 |
| N5–OC2         | 16.9            | 4.7 | 0.0 |
| N6–OO          | 18.0            | 6.7 | 0.0 |
| N12–OCCgC      | 17.5            | 7.3 | 0.0 |
| N13–OCC        | 20.2            | 10.2| 0.0 |
| N20–OC1C2C1    | 10.7            | 0.0 | 11.2|
| N21–OC1C2C2    | 41.9            | 0.0 | 35.3|
| N25–OCCgCC     | 9.8             | 0.0 | 3.8 |
| N9(CH\(_2\)O(CH\(_2\)O) | 12.7            | 0.0 | 9.8 |
| N11(CH\(_2\)O(CO) | 9.4             | 0.0 | 46.9|
| N15(CHO(CHO)   | 6.7             | 0.0 | 41.7|
| N19(CH\(_2\)OH(CHO) | 10.2            | 0.0 | 37.8|
| N23(CO)(CHOH)  | 1.6             | 0.0 | 31.0|
| N26(CO)(CO)    | 0.0             | 5.6 | 39.3|
| N1–C           | 11.2            | 0.0 | 5.8 |
| N2–O           | 17.6            | 6.8 | 0.0 |
| N8–OOC         | 25.7            | 15.9| 0.0 |
| N14–OCCgC      | 8.4             | 0.0 | 36.1|
| N17–OC1C2      | 11.8            | 0.0 | 18.2|
| N18–OC1C1      | 17.5            | 0.0 | 18.8|
| N24–OC1C2C1    | 0.0             | 3.5 | 83.2|
| N10(CH\(_2\)O(CHO) | 0.0             | 6.1 | 32.9|
| N16(CO)(CHO)   | 0.0             | 8.0 | 71.6|
| N22(CO)(CHOH)  | 0.0             | 2.0 | 39.5|

\(^{a}\)For each species the energy difference E(2S+1) between triplet, quintet and septet state is reported.\(^{b}\)The energy difference of the triplet state of N25–OCCgCC is not reported because, irrespective of the initial guess, the system evolves into two separated CO molecules.
Figure 6. (a) Connected undirected graph of the reaction network: node size and color-gradient, from gray to blue, are related to the node betweenness centrality values. (b) Paths with the lowest TEC. Nodes are here individuated just by the first part of the species name.

\[ b-c(n) = \sum_{s \neq n \neq t}^{\sigma} \sigma_{st}(n) \]  

with \( \sigma_{st}(n) \) being the number of paths linking, for the shortest route, nodes \( s \) and \( t \) and the ones, among these, that cross \( n \).

In Figure 6, nodes are differently sized, according to their \( b-c \) values. From the connections properties of the nodes, it is evident that N8-OOC and N12-OOCCg are key intermediates of the title process, once more showing that glyoxal is fundamental in the ethylene glycol decomposition mechanism. Moreover, as already pointed out, 41 simple paths were found: 11 composed by 7 elementary steps as well as 14, 6, 8, and 2 with lengths 9, 11, 13, and 15, respectively (see Supporting Information).

These paths plainly represent the possible sequences of edges that connect the source node to the target one, without any energetic implication. In particular, in the title chemical framework, these are, nothing but, all the possible routes, given the intermediate set, to produce carbon monoxide starting from ethylene glycol. In the hypothesis that the total energy content could be used as an occurrence discriminating factor, characterizing different pathways, the latter were ordered by TEC values. The seven-cardinality sequences N0-N2-N4-N8-N12-N15-N16-N26 and N0-N2-N4-N8-N13-N11-N16-N26, as reported in Figure 6b, following the inference above, should be the most probable. These routes both include glycolaldehyde and the N8-OOC species. In the former, the glyoxal formation is followed by the C-C bond cleavage that would occur after its formation, again suggesting a kind of selectivity to the carbonylic compounds by the here considered Pd\(_{12}\) cluster. Instead in the second path, the C-C bond scission will occur by means of the N13-OOCC intermediate decarbonylation.

\section*{CONCLUSIONS}

Reaction networks are complex systems often composed by several interconnecting routes. In the present study various ethylene glycol, CH\(_2\)OH\(=\)CH\(_2\)OH, decomposition pathways occurring on a subnanometric Pd\(_{12}\) cluster were characterized by means of density functional theory calculations, aimed at pinpointing the energy of reagents, products, and intermediates.
as well as transition states linking them. A starting presieving phase, based on the analysis of elementary step activation barrier values, allowed us to exclude several molecular events because highly unprobable, hence to delineate a peculiar selectivity of the Pd$_{12}$ to the carboxylic intermediate formation, keeping completely out the enolic formation possibility. A graph theory, GT, based analysis, applied on events of comparable activation energies, allowed us to individuate the possible pathways starting from CH$_2$OH–CH$_2$OH leading to H$_2$ and CO. Glycolaldehyde and glyoxal seem to be cornerstones of the whole reaction mechanism while the C=C cleavage likely occurs only after the formation of scarcely hydrogenated intermediates. It was finally noted that GT centrality indexes, outlining the role of nodes within graphs, could be of interest also in the analysis of the tangled reaction mechanism.

**ASSOCIATED CONTENT**

* Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.7b00850.

Reaction paths potentially taking place, along with the ethylene glycol decomposition calculated by means of GT, optimized Cartesian coordinates of intermediates, and transition states characterizing the reaction network along with the energy and imaginary frequency of transition states (PDF)

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**Notes**

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