



Communication

Room-temperature conversion of ethane and the mechanism understanding over single iron atoms confined in graphene

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ABSTRACT

The catalytic conversion of ethane to high value-added chemicals is significantly important for utilization of hydrocarbon resources. However, it is a great challenge due to the typically required high temperature (> 400 °C) conditions. Herein, a highly active catalytic conversion process of ethane at room temperature (25 °C) is reported on single iron atoms confined in graphene via the porphyrin-like N₄-coordination structures. Combining with the operando time of flight mass spectrometer and density functional theory calculations, the reaction is identified as a radical mechanism, in which the C–H bonds of the same C atom are preferentially and sequentially activated, generating the value-added C₂ chemicals, simultaneously avoiding the over-oxidation of the products to CO₂. The in-situ formed O–FeN₄–O structure at the single iron atom serves as the active center for the reaction and facilitates the formation of ethyl radicals. This work deepens the understanding of alkane C–H activation on the FeN₄ center and provides the reference in development of efficient catalyst for selective oxidation of light alkane.

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Dehui Deng's Group for Two-dimensional (2D) Materials & Energy Catalysis. Our research focuses on the development of 2D material-based catalyst and their applications in catalytic conversion of energy molecules, mainly including: (1) Designing and synthesis of advanced 2D materials and their heterostructures, such as graphene, MoS₂, LDHs, for efficient catalytic activities. (2) Thermochemical, electrochemical and thermo-electrical coupled catalysis toward highly efficient activation and conversion of energy molecules, such as O₂, H₂, H₂O, CO, CO₂, CH₄ and CH₃OH. (3) Computational and theoretical modeling of energy catalysis systems to provide fundamental understandings of catalytic activation and reaction mechanism for catalyst optimization and development.

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Ethane as one abundant component of natural gas, shale gas and combustible ice, the exploitation has been in rapid growth in recent years [1]. The value-added C₂ chemicals converted from ethane, such as ethanol, acetic acid and ethene, are widely used as fuels, plastic building block, chemical feedstock [2–6], thus the catalytic conversion of ethane is an important process. However, the high bond energy of C–H (409 kJ/mol) makes the process extremely difficult. High temperature (> 400 °C) [7–9] conditions are typically required in the conventional industrial process, which usually brings side-effects such as over-oxidation of the products and carbon deposition on the catalyst.

Recently, it was reported that ethane can be partially oxidized by H₂O₂ in strong acidic medium at 50–180 °C using complexes with metal cores of V, Mn, Fe, Mo, Ru, Pd, Os, and Pt as homogeneous catalysts [10–14]. In addition, the Fe or Cu modified zeolite catalysts exhibited superior performance in ethane partial oxidation at 50 °C using H₂O₂ as oxidant in heterogeneous catalysis

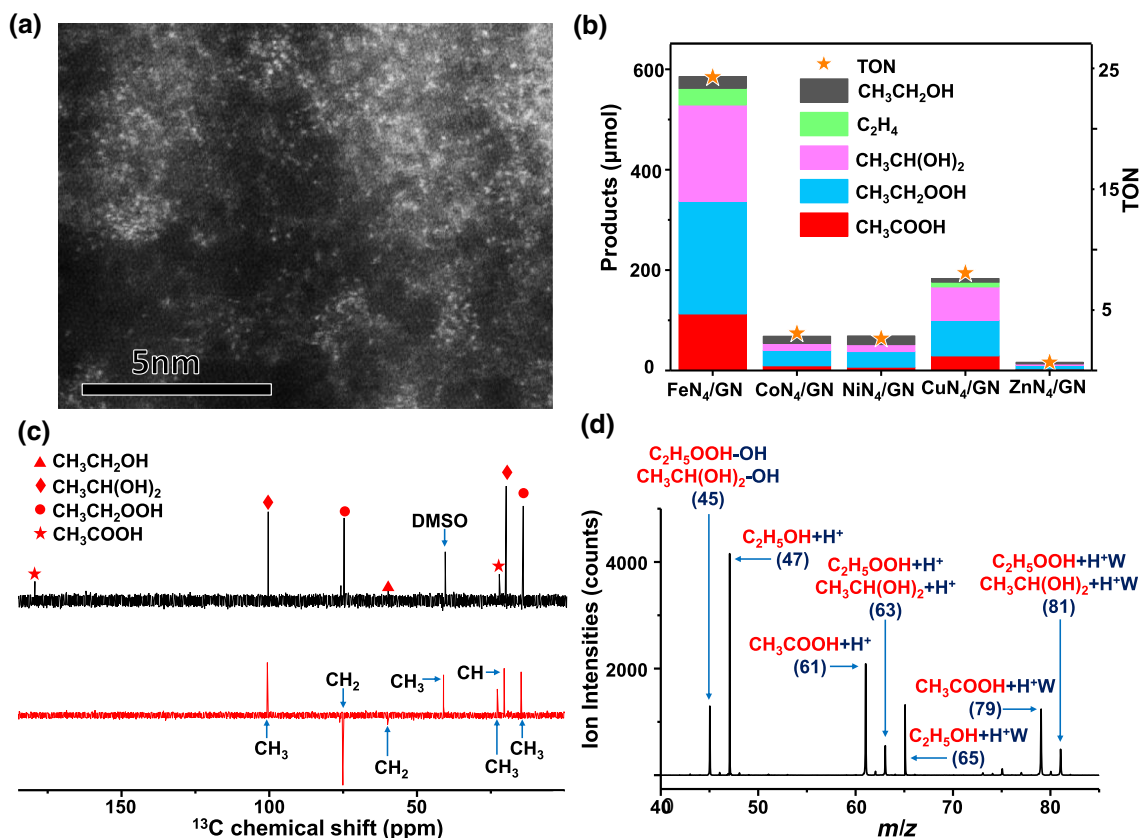


Fig. 1. The structure and catalytic performance of MN_4/GN . (a) HAADF-STEM image of FeN_4/GN . (b) The products of different MN_4/GN (Fe, Co, Ni, Cu, Zn) for ethane oxidation. (c) ^{13}C NMR, ^{13}C DEPT-135 spectra, and (d) TOF-MS spectrum obtained from the reaction of ethane oxidation by FeN_4/GN catalyst. The symbol "W" represents one water molecule. The TOF-MS intensity has been deducted from the background signal that using N_2 as the reaction gas. Reaction conditions in (a–d): 50 mg catalyst, 10 mL H_2O_2 (15%), 1.6 MPa ethane, in a steel autoclave with a Teflon inner vessel (100 mL) at 25 °C for 10 h.

[15–17]. However, achieving ethane conversion at room temperature (25 °C) remains a great challenge.

Herein, we report that single-atom iron with a N_4 coordination [18,19] confined in graphene matrix [20] is highly active for catalytic conversion of ethane to C_2 components (acetic acid (CH_3COOH), ethyl hydroperoxide (CH_3CH_2OOH), 1,1-ethanediol ($CH_3CH(OH)_2$) and ethene (CH_2CH_2)) at room temperature. Besides, the reaction route of ethane oxidation on the catalyst is proposed based on operando time of flight mass spectrometer (TOF-MS), nuclear magnetic resonance (NMR) and density functional theory (DFT) calculations, which indicate that ethane can be easily activated over the in-situ generated O- FeN_4 -O active site along a radical mechanism and is finally converted to various value-added products via different free radicals intermediates, such as $\cdot CH_2CH_3$, $\cdot CH(OH)CH_3$ and $\cdot COCH_3$.

A series of 3d transition metal- N_4 (MN_4 , $M=Fe, Co, Ni, Cu$ and Zn) centers confined in the matrix of graphene nanosheets (MN_4/GN) were prepared according to our previous reported methods by high-energy ball-milling the mixture of metal phthalocyanines and graphene nanosheets [21–23]. The sub-angstrom resolution high angle annular dark field scanning transmission electron microscopy (HAADF-STEM) image displayed that single iron atoms were distributed homogeneously in graphene nanosheets (Fig. 1a). The structure of FeN_4 was confirmed by extended X-ray absorption fine structure (EXAFS) spectroscopy and low-temperature scanning tunneling microscopy (LT-STM) in our previous work [21].

The ethane oxidation reaction was carried out in an autoclave at room temperature (25 °C) under 1.6 MPa using MN_4/GN as the catalyst and H_2O_2 as the oxidant after the optimization of the H_2O_2 concentration, reaction pressure, and catalyst amount (Figs. S1–S3).

The excessive H_2O_2 might break some of FeN_4 active centers, causing the decreasing of conversion (Fig. S1), and therefore a moderate H_2O_2 concentration was used in this study. The reaction products are analyzed by NMR, TOF-MS and gas chromatography (GC). The catalytic activity of various MN_4/GN samples is presented in Fig. 1(b). As shown in Fig. 1(b), FeN_4/GN exhibits the optimum activity for ethane oxidation compared with other metal centers according to the amounts of products and turnover number (TON). The GC characterization results confirm the generation of C_2H_4 in gas phase and the absence of over-oxidized products (CO_2) (Fig. S4). After carefully analyzing the liquid products by various characterization techniques including the 1H NMR (Fig. S5), ^{13}C DEPT-135 (distortionless enhancement by polarization transfer) (Fig. 1c), ^{13}C NMR (Fig. 1c), and HSQC-NMR (heteronuclear single quantum coherence) (Fig. S6), we found that the ethane can be transformed to C_2 components, i.e. the CH_3CH_2OOH , CH_3COOH , $CH_3CH(OH)_2$ and CH_3CH_2OH at room temperature [15]. Moreover, the structure of these products was also confirmed by the TOF-MS (Fig. 1d). The peaks of CH_3COOH and CH_3CH_2OH can be easily marked. However, the signals of CH_3CH_2OOH and $CH_3CH(OH)_2$ are overlapped due to the same molecular weight. These results are consistent with the NMR data. In addition, considering the catalyst itself contains the carbon sources, we also carried out the contrast test under the N_2 condition (Fig. S7), while the C_2 products were not observed in the 1H NMR data. It illustrates that the C_2 components come from the ethane rather than the catalyst itself. In addition, the stability tests reveal that the FeN_4/GN can keep the 60% catalytic activity after six cycles, demonstrating its good stability (Fig. S8).

Furthermore, we used the operando TOF-MS to track the reaction process over the FeN_4/GN sample (Fig. S9). The liquid phase

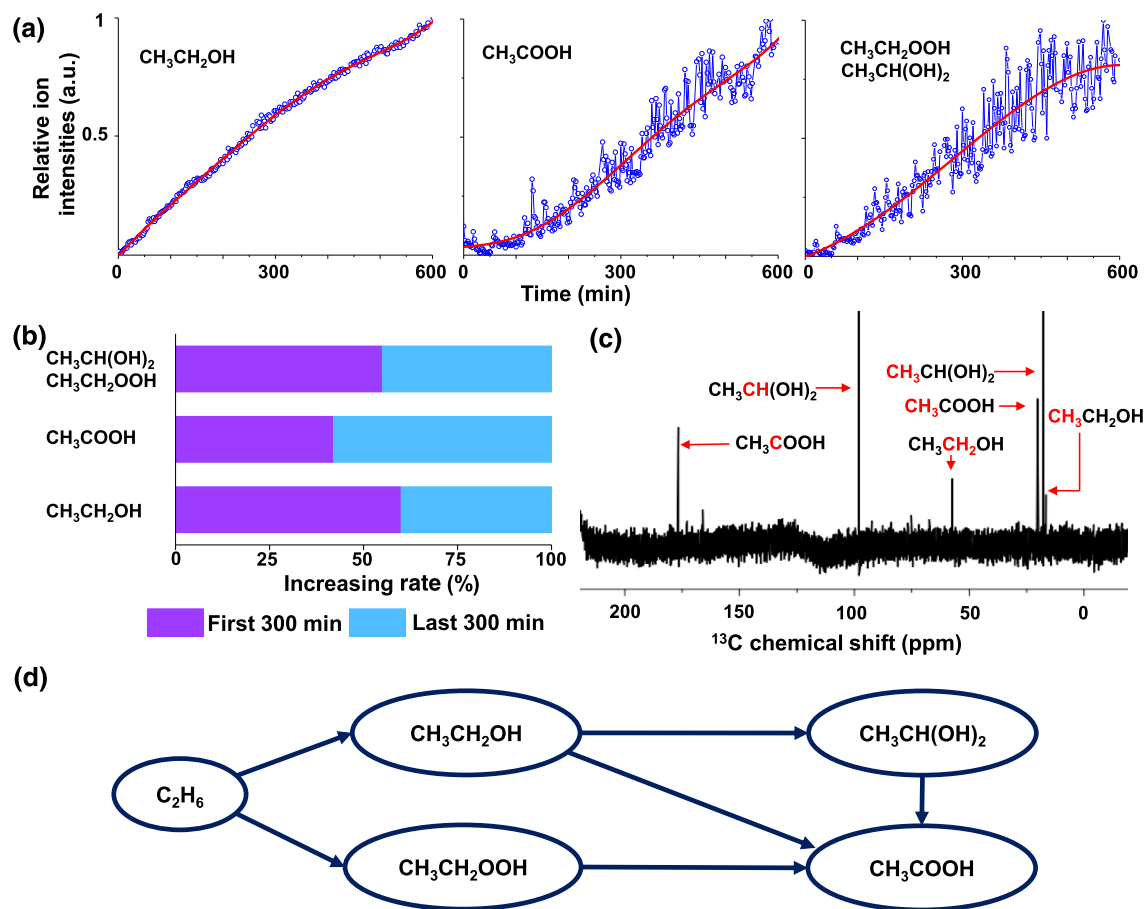


Fig. 2. Mechanism study of FeN_4/GN for ethane conversion. (a) The operando TOF-MS data obtained from C_2H_6 oxidation. (b) The increase rate of the products at 0–600 min, which corresponds to the relative intensity increment. (c) ^{13}C NMR result of $\text{CH}_3\text{CH}_2\text{OH}$ oxidation by FeN_4/GN . (d) A possible reaction path for ethane oxidation over FeN_4/GN catalyst. Reaction conditions in (a): 50 mg catalyst, 10 mL H_2O_2 (15%) at 25 °C for 10 h; in (c): 50 mg catalyst, 10 mL H_2O_2 (15%), 0.05 M $\text{CH}_3\text{CH}_2\text{OH}$, 1.6 MPa N_2 , in a steel autoclave with a Teflon inner vessel (100 mL) at 25 °C for 10 h.

products can be analyzed by TOF-MS in real time through the filter and capillary under the high pressure (typically 1.6 MPa). The increasing rate of different products can be observed during the reaction (Fig. 2a), which were further analyzed towards the increasing rate of the products in the first 300 min and last 300 min (Fig. 2b). The increasing rate of $\text{CH}_3\text{CH}_2\text{OOH}$, $\text{CH}_3\text{CH}(\text{OH})_2$ and $\text{CH}_3\text{CH}_2\text{OH}$ in the first 300 min is greater than in the last 300 min. On the other hand, most of CH_3COOH is produced in the last 300 min. Besides, the similar results were also observed under the different pressures (0.8 MPa, 1.2 MPa) in the operando TOF-MS studies (Fig. S10). These results suggest that ethane should be initially converted to $\text{CH}_3\text{CH}_2\text{OH}$ and $\text{CH}_3\text{CH}_2\text{OOH}$ via oxidation of single C–H bond of ethane followed by further oxidation of these products, leading to the formation of CH_3COOH and $\text{CH}_3\text{CH}(\text{OH})_2$. To confirm this hypothesis, we directly used ethanol as the reactant and obtained the CH_3COOH and $\text{CH}_3\text{CH}(\text{OH})_2$ (Figs. 2c and S11), which indicates that the ethane can be converted into CH_3COOH and $\text{CH}_3\text{CH}(\text{OH})_2$ via ethanol. Besides, through controlling the reaction time, it can be observed that the selectivity of $\text{CH}_3\text{CH}_2\text{OOH}$ and $\text{CH}_3\text{CH}_2\text{OH}$ decreases and the selectivity of CH_3COOH and $\text{CH}_3\text{CH}(\text{OH})_2$ increases during the reaction, which suggests that CH_3COOH and $\text{CH}_3\text{CH}(\text{OH})_2$ may come from $\text{CH}_3\text{CH}_2\text{OOH}$ and $\text{CH}_3\text{CH}_2\text{OH}$ (Fig. S12). According to the above experimental results, we propose one possible reaction route (Fig. 2d). The ethane is firstly converted to the $\text{CH}_3\text{CH}_2\text{OOH}$ and $\text{CH}_3\text{CH}_2\text{OH}$. The generated $\text{CH}_3\text{CH}_2\text{OH}$ will be further oxidized to $\text{CH}_3\text{CH}(\text{OH})_2$ and CH_3COOH . And the $\text{CH}_3\text{CH}_2\text{OOH}$ can also be converted to CH_3COOH .

DFT calculations were carried out to further understand the ethane conversion process. A model of FeN_4 structure confined in the matrix of graphene with the central Fe atom coordinated with four N atoms has been built to simulate the FeN_4/GN catalysts. In the aqueous H_2O_2 environment, FeN_4 is easy to adsorb two O atoms and form the O– FeN_4 –O structure (Fig. 3a), which is the active site for C–H bond cleavage [21,23]. The structure of O– FeN_4 –O has been proved by X-ray absorption fine structure spectroscopy (XAFS) and Mössbauer spectra in our previous work [21]. The ethane activation is along a radical mechanism. As shown in Fig. S13, the C–H bond of C_2H_6 is stretched on the O– FeN_4 –O site, through a transition state (TS) forming a $\cdot\text{CH}_2\text{CH}_3$ radical, which will bind with the OH group on the surface to generate $\text{CH}_3\text{CH}_2\text{OH}$, and the left FeN_4 –O structure will react with the H_2O_2 in the solution to cycle back to the O– FeN_4 –O active structure. The activation barrier is only 0.48 eV, which means the ethane activation can be performed in mild conditions (Fig. 3b). As shown in Fig. 3(a), the $\cdot\text{CH}_2\text{CH}_3$ radical may also bind with one $\cdot\text{OOH}$ group in the H_2O_2 solution to form $\text{CH}_3\text{CH}_2\text{OOH}$ or break another C–H bond to form CH_2CH_2 .

The generated $\text{CH}_3\text{CH}_2\text{OH}$ molecules can be further activated on the O– FeN_4 –O site. As shown in Figs. S14 and S15, compared with the CH_3 group (barrier: 0.56 eV), the CH_2 group is more easily to be activated (barrier: 0.20 eV) to form $\text{CH}_3\text{CH}(\text{OH})_2$ radical, which was confirmed by the electron paramagnetic resonance experiments (Figs. S16 and S17). The $\text{CH}_3\text{CH}(\text{OH})_2$ radical can also be dehydrated to generate CH_3CHO . Moreover, the last C–H bond

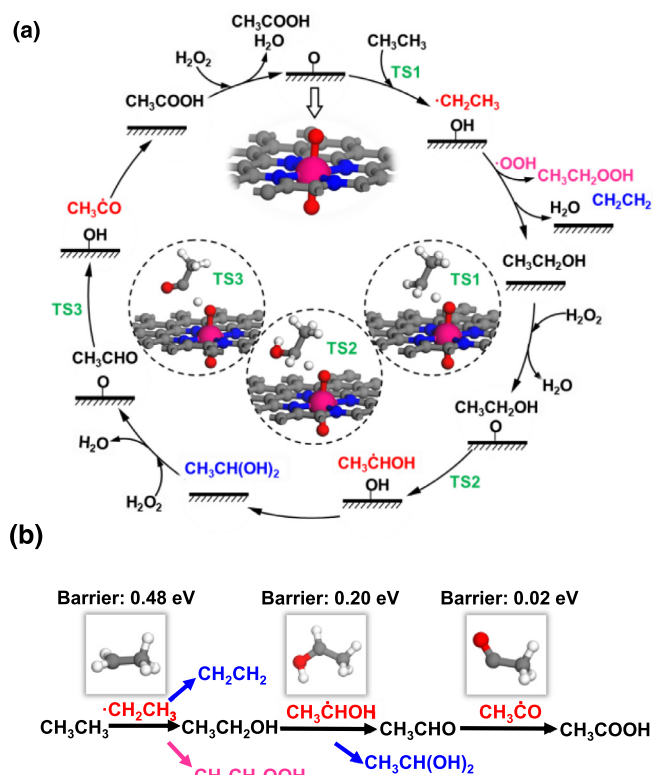


Fig. 3. Proposed mechanism for C_2H_6 activation on the $O-FeN_4-O$ active site. (a) Reaction cycle with the reactants (black), products (black, blue and pink), the radical intermediates (red) and the atomic structures of the active site and transition states TS1–TS3. (b) Reaction pathway with atomic structure of intermediates and the corresponding activation barrier.

on the C atom of CH_3CHO can be very easily activated (barrier: 0.02 eV) to form a further oxidation product CH_3COOH , which explains that almost no aldehyde products were detected. The DFT results agree well with the experimental results.

In conclusion, we report a FeN_4/GN catalyst with high activity for the conversion of ethane to C_2 chemicals at room temperature. Through a series of experiments including operando TOF-MS and DFT calculations, we found that the $O-FeN_4-O$ structure formed in aqueous H_2O_2 is able to convert ethane to ethyl radical in mild conditions. The ethyl radicals will bind with the $\cdot OH$ and $\cdot OOH$ groups to form CH_3CH_2OH and CH_3CH_2OOH . Furthermore, through activating the C–H bonds on the same C atom, CH_3CH_2OH could be further converted to $CH_3CH(OH)_2$ and CH_3COOH . These findings provide the reference for understanding and designing highly efficient catalysts for C–H bond activation and conversion.

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Supplementary material

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.jechem.2019.04.003.

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