Dioxygen Activation Kinetics over Distinct Cu Site Types in Cu-CHA Zeolites

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ABSTRACT: Cu-exchanged zeolites activate dioxygen to form active sites for partial methane oxidation (PMO), nitrogen oxide decomposition, and carbon monoxide oxidation. Apparent rates of O2 activation depend both on the intrinsic kinetics of distinct Cu site types and the distributions of such sites within a given zeolite, which depend on the density and arrangement of the framework Al atoms. Here, we use hydrothermal synthesis methods to control the arrangement of framework Al sites in chabazite (CHA) zeolites and, in turn, the distinct Cu site types formed. Time-resolved in situ resonance Raman spectroscopy reveals the kinetics of O2 adsorption and activation within these well-defined Cu-CHA materials and the concomitant structural evolution of copper-oxygen (CuOx) complexes, which are interpreted alongside Cu(I) oxidation kinetics extracted from in situ X-ray absorption spectroscopy (XAS). Raman spectra of several plausible CuOx species simulated using density functional theory suggest that experimental spectra (λex = 532 nm) capture the formation of mono(μ-oxo)dicopper species (ZCuOCuZ). Transient experiments show that the timescales required to form CuOx structures that no longer change in Raman spectra correspond to the durations of oxidative treatments that maximize CH3OH yields in stoichiometric PMO cycles (approximately 2 h). Yet, these periods extend well beyond the timescales for the complete conversion of the initial Cu(I) intermediates to their Cu(II) states (<0.3 h, reflected in XANES spectra), which demonstrates that CuOx complexes continue to evolve structurally following rapid oxidation. The dependence of ZCuOCuZ formation rates on O2 pressure, H2O pressure, and temperature are consistent with a mechanism in which ZCuOH reduce to form ZCu+ sites that bind molecular oxygen and form ZCu-O2 intermediates. Subsequent reaction with proximate ZCu+ form bridging peroxo dicopper complexes that cleave O-O bonds to form ZCuOCuZ in steps facilitated by water. These data and interpretations provide evidence for the chemical processes that link rapid and kinetically irrelevant Cu oxidation steps (frequently probed by XAS and UV-Vis spectroscopy) to the relatively slow genesis of reactive Cu complexes that form CH3OH during PMO. In doing so, we reveal previously unrecognized complexities in the processes by which Cu ions in zeolites activate O2 to form active CuOx complexes, which underscore the insight afforded by judicious combinations of experimental and theoretical techniques.

1.0 INTRODUCTION

Copper ions exchanged onto Al-substituted zeolites facilitate various oxidation reactions and redox cycles, including NO decomposition,1-2 CO oxidation,3-4 NOx (x = 1, 2) selective catalytic reduction using NH3,5-9 and the partial oxidation of CH4 to form CH3OH in stoichiometric10-17 and catalytic18-19 cycles. In the context of partial methane oxidation (PMO), Cu(I) ions reduce to their Cu(II) states at high temperatures (e.g., 723 K in He,14, 17, 20-21 523 K in 5% CO20, 22) prior to exposure to extended O2 activation treatments (e.g., 723 K, > 2 h) that form active multinuclear Cu(II) species (CuOx)1-19. At lower temperatures, these CuOx intermediates react readily with CH4 to form surface methoxy-derived species (e.g., complete consumption within 0.5 h at 473 K), which desorb as CH3OH upon contact with H2O in vapor or liquid phases. The dependence of methanol yields (per Cu) on reaction conditions (e.g., reactant pressure, temperature, time) has been well studied;11, 14, 17, 23-24 however, comparatively few investigations describe the series of steps responsible for the activation of O2 molecules or the kinetics and formation of active CuOx complexes.11, 22, 24

Nearly two decades of research have sought to identify the CuOx species that form upon O2 activation treatments of Cu-exchanged zeolites and produced numerous proposals (e.g., bis(μ-oxo)dicopper,1-10 mono(μ-oxo)dicopper,1-13, 16, 23-24 μ-(η2:η2)peroxo dicopper,25 trans-η1,2-peroxo dicopper,13, 23 bis(μ-hydroxyl)dicopper26, mono(μ-oxo)tricopper27-29). Initial reports from Groothaert et al. combined in situ extended X-ray absorption fine structure (EXAFS), UV-Vis-near-IR, and electron paramagnetic
resonance (EPR) spectroscopies with comparisons to dicopper metalloenzymes to assign a 22,700 cm$^{-1}$ UV-Vis absorbance feature observed on O$_2$-activated Cu-ZSM-5 to a bis-(μ-oxo)dicopper species. Solomon and coworkers utilized resonance Raman spectroscopy and DFT from which they concluded a bent mono-(μ-oxo)dicopper forms upon O$_2$ activation over Cu-ZSM-5, and Cu-MOR. Recently, van Bokhoven and coworkers examined the formation of PMO active sites in MOR and MAZ zeolites and proposed that proximal [CuOH]$^+$ species form bridged Cu-oxo type species using X-ray powder diffraction, DFT, and XAS. Additionally, Pappas et al. studied Cu-MOR based on Cu-normalized stoichiometric CH$_3$OH yields (O$_2$ activation: pure O$_2$, 723 K, 1 h; CH$_4$ introduction: 91.2 kPa CH$_4$ balance He, 473 K, 4 h; CH$_3$OH extraction: equimolar H$_2$O and He, 408 K, 0.5 h) that correlated linearly with one-third of the total Cu content, consistent with structures identified within in situ EXAFS spectra during O$_2$ activation. Taken together, this series of contributions presents compelling evidence for multiple and seemingly contradictory conclusions regarding the identity of Cu$_x$O$_y$ species formed upon O$_2$ activation over Cu-zeolites, which remains an important area for continued investigation.

The ability of Cu species to bind O$_2$, cleave the O-O bond, and form distinct Cu$_x$O$_y$ structures depends strongly upon the spatial and crystallographic distribution of the Al atoms within the zeolite and the fraction of this distribution associated with Cu ions. The Al distribution in a given zeolite depends on its bulk Si/Al ratio and framework topology, and the conditions used for its synthesis, which can collectively lead to significant Cu site heterogeneity among experimental studies. Divalent copper complexes are particularly sensitive to non-uniformities in the arrangement of AlO$_4$ sites, because they either require charge compensation from two AlO$_4$ centers or coordination to anionic ligands if only one AlO$_4$ center is present. As stated by Borfecchia et al., the differences between the many proposals for active Cu$_x$O$_y$ sites for stoichiometric PMO cycles likely reflect complexities both among the zeolite topologies studied (e.g., ZSM-5, MOR, CHA) and among the varied reaction protocols used (e.g. O$_2$ activation conditions, CH$_4$ reaction conditions).

The high symmetry chabazite (CHA) framework provides a model material to study the nature of O$_2$ activation, because CHA contains a single crystallographically unique tetrahedral site. Moreover, CHA zeolites can be synthesized intentionally with precise control over the relative proximity of framework Al atoms. For example, Dussel and coworkers used this capability to demonstrate on Fe-CHA zeolites that paired Al atoms in the 6-membered ring (MR) (i.e., in second- or third-nearest neighbor configurations) stabilize the PMO active site (α-Fe) and correlate to CH$_3$OH yields (per Fe) from PMO cycles. Well-defined Cu-CHA materials with controlled speciation of Cu ions can be synthesized wherein 6-MR paired Al sites (2 Al per 6-MR) exchange Cu$^{2+}$ ions (Z$_2$Cu), while 6-MR isolated framework Al atoms (1 Al per 6-MR) nominally exchange [CuOH]$^+$ species (ZCuOH). Upon O$_2$ activation of Cu-CHA, a fraction of ZCuOH species form binuclear O$_2$-bridged structures observed within in situ UV-Vis spectra, while Z$_2$Cu sites do not. These findings agree with reports Z$_2$Cu sites do not participate in PMO, however, remain elusive.

Here, we use time-resolved in situ resonance Raman spectroscopy to measure the spectrokinetics of O$_2$ activation over well-defined Cu-CHA materials that contain predominantly Cu either at 6-MR isolated or paired Al sites, or mixtures thereof. In situ Raman spectroscopy demonstrates that mono-(μ-oxo)dicopper complexes form over extended periods (2 – 6 h) similar to those needed to maximize CH$_3$OH yields, which suggests the Cu$_x$O$_y$ complexes detected by Raman correspond to the Cu(II) species responsible for PMO. In contrast, in situ XAS measurements indicate that Cu(I) converts to Cu(II) intermediates (Cu-O$_2$; e.g., μ-$\eta^1$-$\eta^1$-peroxo dicopper) on much shorter timescales (< 0.3 s) and, therefore, the Cu(I) to Cu(II) transformation detected by XAS may signify the formation of a precursor to PMO-active sites. The combination of DFT+U computed Raman intensities, steady-state Raman spectra, and $^{18}$O-labeling suggest ZCuOH sites activate O$_2$ to form mono-(μ-oxo)dicopper species. Rates of formation for the mono-(μ-oxo)dicopper complex are measured as a function of O$_2$ (5 – 42 kPa O$_2$) and H$_2$O pressure (0 – 1 kPa H$_2$O) and temperature (648 – 773 K) to probe the series of chemical transformations that occur during O$_2$ activation. These rates depend on the isotopic substitution of water (k$_{H}$/k$_{D}$ ≳ 2), which suggests that proton transfer may mediate O-O bond cleavage to form mono-(μ-oxo)dicopper species. Collectively, these findings provide evidence for the molecular processes that determine the rates at which PMO-active sites form by interactions with O$_2$ and H$_2$O reactants.

### 2.0 MATERIALS AND METHODS

#### 2.1 Raman Spectroscopy

Resonance Raman spectra were obtained on a spectrometer (Renishaw, inVia) equipped with a 532 nm laser. Cu-CHA samples were first pelletized and sieved to retain ~2 mm particles, which were loaded into a temperature-controlled gas-phase reaction cell (Linkam, CCR1000). Cu-CHA samples were first treated to either 723 K in He (Airgas, Ultra-zero grade; 50 cm$^3$ min$^{-1}$; 0.33 K s$^{-1}$) and held for 2 h or to 523 K in 5% CO (Airgas, 99.999%); balance He; 50 cm$^3$ min$^{-1}$ total flow rate; 0.33 K s$^{-1}$) for 1 h. After pretreatments in CO, the sample chamber was purged with He (50 cm$^3$ min$^{-1}$) for 0.5 h at 523 K. The sample was then heated to a desired temperature in He (0.33 K s$^{-1}$). Once at the desired O$_2$
activation temperature (648 – 773 K), O2 (Airgas, 99.999%) diluted in He was introduced to the sample to maintain a volumetric flow rate of 50 cm³ min⁻¹ and H2O (17.8 MΩ cm) or D2O (Sigma Aldrich, 99.9% D atom) was introduced using a syringe pump (KD Scientific, Legato 100) through a liquid-injection port. Simultaneously, spectra (0.1 s, 25 co-added spectra) were obtained using line-scan mode with a long 50x objective, such that the power density was approximately 0.4 mW μm⁻². All Raman spectra were referenced to the 520 cm⁻¹ feature of Si(111). Control experiments which varied the power of the laser (0.04 – 0.8 mW μm⁻²) and time delays between laser exposure indicate that the measured spectra are not affected by localized heating induced from the laser. In our hands, the CO gas must be purified using a combination of an in-line moisture and O₂ trap (Supelco), as well as Cu turnings (Sigma Aldrich, 99%) that were heated to 553 K. We observed a discoloration of the Cu-CHA surface, which indicates contamination by metal carbonyls, when these traps were not in place. Other contaminants were removed from the O₂ and He streams using inline moisture/hydrocarbon and moisture/hydrocarbon/oxygen combination purifying traps (Supelco), respectively. Consequently, we assume that the partial pressure of H₂O is <10⁻⁴ Pa (in the absence of intentionally added water vapor), based upon specifications from the gas-trap manufacturers.

The spectral contributions of the independent species formed during O₂ activation of Cu-CHA were obtained by multivariate curve resolution-alternating least squares (MCR-ALS) performed in the WiRE™ software package. Iterative fitting procedures, involving up to five components, were used to describe 99.9% of the data. In all cases, greater than 90% of the variance of the measured spectra was described by a single component, while the other extracted spectra primarily represent changes in the baseline during the course of the experiment related to shifts in the sample position or focus over the course of many hours (Section S8).

2.2 X-Ray Absorption Spectroscopy

X-ray absorption spectroscopy experiments were performed at the Advanced Photon Source (APS), Argonne National Laboratory in Lemont, Illinois, in sector 10 MR-CAT (Materials Research Collaborative Access Team). The insertion device beamline at sector 10 (10-ID) was used for in situ experiments. A Cu metal foil reference spectrum (edge energy of 8979 eV) was measured simultaneously with each sample spectrum collected to calibrate the X-ray beam for spectral measurements at the Cu K-edge. All sample spectra were analyzed in WinXAS and normalized using first and third order polynomials for background subtraction of the pre- and post-edges, respectively. The standards used for linear combination fitting (LCF) of the XANES spectra were Cu-CHA dehydrated in 21 kPa O₂ (balance He) at 723 K and Cu₂O (Sigma Aldrich), for Cu(II) and Cu(I), respectively, as shown in Figure S9.

Auto-reduction in inert helium (99.999% UHP), flowed through an oxygen/moisture trap (Matheson, MTRP-0019-XX) at a flow rate of 50 cm³ min⁻¹, was studied at 723 K (0.167 K s⁻¹) until the XANES spectra stopped changing (sample dependent, up to 2 h). For the oxidation treatment, the sample was held in oxygen (21 kPa in balance He) at a total flow of 50 cm³ min⁻¹ while increasing the temperature from ambient to 723 K (0.167 K s⁻¹) and holding at temperature for up to 2 h.

2.3 Partial Methane Oxidation

Cu-CHA samples were first pretreated in He (UHP, Indiana Oxygen, 50 cm³ min⁻¹) at 723 K (0.167 K s⁻¹) for 2 h followed by activation in dry air (21 kPa O₂, Zero grade air, 50 cm³ min⁻¹) at 723 K for various times (0 – 15 ks) and then cooled to ambient temperature in He (Indiana Oxygen, 99.999%, 50 cm³ min⁻¹). Methane (20 kPa, Indiana Oxygen, 99.999%) with balance of helium (81 kPa) was introduced and heated to reaction temperature of 473 K at 0.167 K s⁻¹ and held for 0.5 h. Prior to methanol extraction, the system was purged with He (0.1 h, 50 cm³ min⁻¹) and then the flow was switched to 2.5 kPa H₂O in a He carrier (30 cm³ min⁻¹) at 473 K for 1 h. An online mass selective detector (Agilent, 5973N) was used to quantify methanol (m/z = 31) using Ar (Indiana Oxygen, 99.999%) as a calibration standard (m/z = 40), and to monitor potential side products including CO, CO₂, CH₂O, and C₂H₅O (m/z = 28, 44, 29, 45, respectively).

2.4 Density Functional Theory Calculations

Calculations were performed using the Vienna ab initio Simulation Package (VASP)30 version 5.4.4. For all calculations, only considered the Γ point was considered when sampling the first Brillouin zone. Periodic spin-polarized DFT+U calculations were performed with a 400 eV cutoff energy, a plane-wave basis, and the PBE functional coupled with a semi-empirical D3 dispersion correction with Becke–Johnson damping39-40 on a CHA supercell containing 12 T-sites. Electron-ion interactions were described with projector augmented wave (PAW)41 method. For the +U calculations, the U value for the d-orbitals of Cu was set to 6.0 eV based on a recent benchmarking study by Chen et al.42 Lowest energy Cu dimer structures were sampled over 150 ps 300 K ab initio molecular dynamics (AIMD) simulations with NVT ensemble with Nosé–Hoover thermostat.20 Unique local minima for each Cu dimer motif were identified by optimizing 400 structures (equally spaced in time). The structures were relaxed until all forces were smaller than 0.01 eV/Å, with a convergence criterion of 10⁻⁶ eV for energies inside self-consistent-field (SCF) cycles. Raman intensity calculations followed the same protocol described by Liang et al.43 For Cu dimer structures with multiple unique local minima, spectra were Boltzmann-averaged to compute the final spectra reported in Figures 3 and S15:

\[
\langle I \rangle = \frac{\sum I_i e^{-E_i/k_B T}}{\sum e^{-E_i/k_B T}}
\]

where \(\langle I \rangle\) is the Boltzmann averaged Raman intensity for a Cu dimer type (e.g., IV), \(I_i\) is the simulated spectra of the \(i^{th}\)
local minimum within that dimer type (e.g., IV-1), $k^B$ is the Boltzmann constant, $E_i$ is the energy spectra of the $i^{th}$ local minimum within that type, and $T$ is the temperature. Figure S15 reports the computed spectra for each Cu dimer and local minima. All unique local minima structures used to compute frequencies are provided in the SI, Section S7.

3.0 RESULTS AND DISCUSSION

3.1 Disparate Timescales for Cu Oxidation and Cu_xO_y Formation Observed via X-ray Absorption and Raman Spectroscopy

The thermodynamics of copper oxidation and reduction depends on Cu site speciation, and in turn, the zeolite composition and framework Al arrangement, as shown by the DFT-derived phase diagrams reported by Paolucci et al.\textsuperscript{7} All Cu ions at 6-MR paired sites are in their Cu(II) states at 723 K (21 kPa O\textsubscript{2}), while Cu ions at 6-MR isolated Al sites may reside as mixtures of Cu(II) and Cu(I) states at 723 K (21 kPa O\textsubscript{2}). The Cu speciation also influences observed auto-reduction behavior at 673 K (1 Pa O\textsubscript{2} in balance He), as ZCu sites remain Cu(II) while ZCuOH sites thermodynamically prefer the Cu(I) state.\textsuperscript{14} Elementary steps to complete ZCuOH auto-reduction events likely require two [CuOH]$^+$ within close enough proximity to form binuclear Cu-oxo species,\textsuperscript{44} which suggests that spatially distant ZCuOH sites will not auto-reduce.\textsuperscript{14} This interpretation is consistent with Raman spectra ($\lambda_{ex} = 532$ nm, 18,800 cm$^{-1}$) collected in He at 723 K that do not show discernible features for binuclear Cu-oxo structures, (Figure 1a; spectrum at 0 ks solely reflects background fluorescence), which if formed presumably have auto-reduced.

The Cu-CHA samples studied here will be referred to as Cu-CHA-I, Cu-CHA-P or Cu-CHA-M where I, P, and M contain Cu exchanged at predominantly 6-MR isolated Al sites, 6-MR paired Al sites, or mixtures thereof, respectively. All Cu-CHA samples exhibit UV-Vis absorption features between 8,000 and 20,000 cm$^{-1}$ after O\textsubscript{2} activation at 723 K (Figures S3 - S5), which are consistent with the Cu speciation expected within Cu-CHA materials that contain predominantly ZCuOH sites, Z\textsubscript{2}Cu, or mixtures thereof.\textsuperscript{20}

Figure 1a shows \textit{in situ} resonance Raman spectra of Cu-CHA-I during O\textsubscript{2} activation, demonstrating that more than 7
ks (~2 h) at 723 K is required for CuOx to reach their steady-state structures. These spectra were obtained with an excitation wavelength of 532 nm to achieve resonance Raman spectra. To capture the transient changes in the number of Raman-active species, we continuously collect Raman spectra during O2 activation. Multivariate curve resolution-alternating least squares (MCR-ALS) analysis allows us to recover the changes in spectral contributions over time from each kinetically and spectroscopically distinguishable species. For each sample, we find that a single component (i.e., spectrum) and concentration profile describes the time-resolved measurements. If two or more components are modeled, the additional spectra resemble changes within the spectral baseline or random fluctuations in the measurements (See Section S8) and do not contain features consistent with reported CuOx complexes (vide infra) or the CHA framework.

During O2 activation treatments at 723 K (21 kPa O2, 80 kPa He), the formation of Raman-active CuOx species occurs over a period of 12 ks (Figure 1b), while the near-complete oxidation of Cu(I) to Cu(II) approaches a steady-state within 1 ks as monitored by X-ray absorption near edge spectroscopy (XANES; Figure 1d). In both cases, the rate \( r_{\text{app}} \) of O2 activation (Raman) and Cu(I) oxidation (XANES) follows pseudo-first order kinetics:

\[
r_{\text{app}} = k_{\text{app}}[\text{Cu(I)}] \tag{2}
\]

where \( k_{\text{app}} \) is the apparent rate constant and \([\text{Cu(I)}] \) is the molar density of Cu(I) ions in the Cu-CHA sample. Within Cu-CHA-I, the rate constant for Cu(I) oxidation \( k_{\text{app,XANES}} = 1.96 \text{ ks}^{-1} \) is 6-times greater than the rate constant to form the CuOx species observed by Raman spectroscopy \( k_{\text{app,Raman}} = 0.31 \text{ ks}^{-1} \). Similarly, the rate constants for Cu(I) oxidation within Cu-CHA-M (Figure S14; \( k_{\text{app,XANES}} = 4.01 \text{ ks}^{-1} \)) are 40-fold greater than rate constant obtained from transients measured by Raman \( k_{\text{app,Raman}} = 0.10 \text{ ks}^{-1} \). In situ EXAFS after 1 ks reveal that Cu sites in Cu-CHA-M (Table S6) and Cu-CHA-I (Table S4) are three-coordinate. These spectral features are distinct from those observed in Raman (Figure 2) and EXAFS (four-coordinate Cu; Table S5) of Cu-CHA-P that contain predominantly Cu(II) before and after contact with O2 (Figure S11) as expected for Z2Cu sites that are PMO-inactive.

Importantly, the yields of CH3OH (per mol of Cu) increase monotonically from 0 to ~10 ks over Cu-CHA-I (Figure 1b), and \( k_{\text{app}} \) values measured via Raman (0.31 ks\(^{-1}\)) and PMO (0.37 ks\(^{-1}\)) are in close agreement, suggesting that the features within these Raman spectra reflect PMO-relevant active sites proposed previously to be binuclear Cu-oxo site types that require longer timescales to form. A non-negligible amount of methanol forms on Cu-CHA-I following a rapid oxidation of Cu(I) to Cu(II) before and after contact with O2 (Figure S11) as expected for Z2Cu sites that are PMO-inactive.

The identity of Raman-Active CuOx Species Formed Over ZCuOH

![Figure 2](Image)

**Figure 2.** In situ steady-state Raman spectra (\( \lambda_{\text{ex}} = 532 \text{ nm} \)) of Cu-CHA samples containing predominantly Z2Cu sites (Cu-CHA-P), ZCuOH sites (Cu-CHA-I), or mixtures thereof (Cu-CHA-M) after O2 activation (21 kPa O2, 80 kPa He, 723 K). Samples were pretreated in He at 723 K for 2 h prior to the introduction of O2. All spectra have been normalized to the most-intense feature (~450 cm\(^{-1}\)) and are vertically offset for clarity.

The identity of the PMO- and Raman-active CuOx species must be determined to generate a meaningful model that describes its genesis during O2 activation. Figure 2 shows steady-state spectra contain several significant vibrational features between 300 – 1,300 cm\(^{-1}\). The sharp features at 330 and 475 cm\(^{-1}\) correspond to the \( \nu(T-O-T) \) mode of the
Figure 3. In situ steady-state Raman spectra ($\lambda_{ex} = 532$ nm) of Cu-CHA-I (top) and Cu-CHA-M (bottom) after O$_2$ activation (21 kPa O$_2$, 80 kPa He, 723 K) with natural abundance $^{16}$O$_2$ (black), $^{18}$O$_2$ (red), or $^{16}$O$_2$ with 1 kPa of H$_2$O (blue). b) Optimized geometries of Cu dimers I (orange, trans-peroxo dicopper), II (green, $\mu$-(η$^2$:η$^2$)peroxo dicopper), III (purple, bis(µ-hydroxyl) dicopper), IV (blue, mono-µ-oxo dicopper), V (red, mono-µ-oxo dicopper), and VI (teal, mono-µ-oxo dicopper), where dicopper species IV, V, and VI differ in the arrangement of Al atoms within the 8-membered ring. c) The corresponding simulated Raman-active vibrations for each of the Cu dimers and monomers. Colors within panel c) are consistent with the colored boxes for each Cu dimer in b).

The identity of the species observed during O$_2$ activation by Raman spectroscopy was evinced through a combination of isotopic labeling and computed Raman spectra of plausible chemical species. Isotopic labeling experiments with $^{18}$O$_2$ were performed by treating Cu-CHA-I and Cu-CHA-M at 723 K in flowing $^{18}$O$_2$ (21 kPa $^{18}$O$_2$, 80 kPa He) to identify vibrational shifts to aid in the identification of reactive Cu$_x$O$_y$ species formed. Both Cu-CHA-I and Cu-CHA-M possess Raman scattering features at 603 cm$^{-1}$ when either O$_2$ or $^{16}$O$_2$ is used (Figure 3). Cu-CHA-M activated in O$_2$ under dry conditions, however, possesses an additional feature around 580 cm$^{-1}$ previously attributed to trans-µ-1,2-peroxo dicopper(II). Consequently, these data suggest that the presence of H$_2$O deliberately added to 16O$_2$ or present in trace amounts within 18O$_2$ aid in the structural conversion of trans-µ-1,2-peroxo dicopper(II) to the species that exhibits a Raman scattering at 603 cm$^{-1}$. Notably, these findings differ from those of Lobo and coworkers, who observed an isotopic shift (Δ$^{18}$O$_2$) of 24 cm$^{-1}$ for the vibration at 603 cm$^{-1}$ was assigned to ν(O-O) of trans-µ-1,2-peroxo dicopper(II). Here, the lack of shift in the Raman spectra of 18O$_2$-activated Cu-CHA zeolites seems to exclude this possibility. Consequently, the broad features centered around 800 and 1100 cm$^{-1}$ (Figure 2) appear to reflect framework ν(Si-O) and νas(Si-O) modes that are strongly scattered based upon preferential excitation (i.e., resonance) of Cu$_x$O$_y$ species they stabilize.
To aid in spectral assignments, we used DFT+U to model frequently discussed Cu dimer structures (Figure 3b) and calculate their Raman spectra (Figure 3c). For each optimized structure, the change in polarizability was calculated along the normal modes of the simulated vibrations to determine the frequencies of Raman-active vibrations (see Section 2.4 for full details). The precise assignment of a complex whose calculated molecular vibrations describe the Raman features in Figures 1 and 3a (i.e., at 603 cm⁻¹) requires that several criteria be met. First, the CuₓOᵧ complex must exist as Cu(II), which excludes the possibility of bis-μ-oxo dicopper and trimeric Cu-oxo complexes. Second, the CuₓOᵧ intermediate must absorb 532 nm (18,800 cm⁻¹; Figures S3 and S5) light to provide the resonance Raman effect. Third, the candidate dimer structures modeled must possess Raman scattering features within 20 cm⁻¹ of the experimentally measured vibrations, which is the range of deviations between experimental and theoretical frequencies commonly reported for similar materials. Fourth, the simulated vibration near 603 cm⁻¹ must not contain significant Cu-O or O-O bond deformations (e.g., stretching modes of superoxo or peroxo complexes), as these vibrations would produce a significant Δ18O₂.

Among the complexes depicted in Figure 3b, mono(μ-oxo) dimer species (IV, V, and VI) most closely satisfy these criteria. These complexes exhibit strong absorbance between 8,000 – 20,000 cm⁻¹ that do not show significant deformation of the Cu-O bonds (Supporting Information, Videos). In comparison, our calculations show that Cu dimers I – III (Figure 3b) do not have significant UV-Vis absorbance features near 18,800 cm⁻¹ and do not possess Raman features near 603 cm⁻¹ that correspond to vibrations with significant Cu-O or O-O bond deformations (which is inconsistent with Δ18O₂ = 0). Consequently, the CuₓOᵧ species that form over Cu-CHA-I and Cu-CHA-M over long time scales (0 – 15 ks) are active for PMO and that are responsible for the 603 cm⁻¹ feature.
likely correspond to mono(μ-oxo)dicopper intermediates. We note, however, that the precise and unambiguous assignment of the 603 cm⁻¹ Raman scattering feature to a multinuclear vibration of mono(μ-oxo)dicopper species remains difficult as DFT+U calculations are performed off-resonance and do not account for the preferential excitation of vibrations due to induced electronic transitions. Consequently, we tentatively assign the species responsible for the observed Raman feature at 603 cm⁻¹ and for PMO activity as a mono(μ-oxo)dicopper species but do not exclude any of the plausible isomers (species IV, V, or VI). For brevity, we refer to mono(μ-oxo)dicopper as ZCuOCuZ in the following sections.

### 3.3 Mechanistic Interpretations of Mean-Field O₂ Activation Kinetics

Observations presented earlier in this report suggest that rates of ZCuOCuZ formation over Cu(I) ions likely depend on O₂ and H₂O pressure with a functional form resembling

\[
r_{\text{app}} = k_{\text{form}} P_{O_2}^{a} P_{H_2O}^{b} [Cu(I)]
\]

where \(k_{\text{form}}\) is the rate constant for ZCuOCuZ formation, \(P_i\) is the partial pressure of species \(i\), and \(a\) and \(b\) represent the power-law dependence of rates on \(P_{O_2}\) and \(P_{H_2O}\), respectively, and \([Cu(I)]\) is the number of Cu(I) sites. The conversions of \(O_2\) and \(H_2O\) remain differential (< 1%) throughout the formation of ZCuOCuZ complexes, because these reactants are introduced with molar flowrates that greatly exceed the consumption of Cu(I) ions within the sample over the relevant timescales for ZCuOCuZ formation (1 - 150 ks). As such, the rate of ZCuOCuZ formation can be stated in a form that reflects a pseudo-first-order dependence on [Cu(I)]

\[
r_{\text{app}} = k_{\text{app}} [Cu(I)]
\]

where \(k_{\text{app}}\) is the apparent rate constant for ZCuOCuZ formation and implicitly contains the dependence of \(O_2\) activation rates on \(P_{O_2}\) and \(P_{H_2O}\).

Figures 4c and 4f show that values of \(k_{\text{app}}\) over Cu-CHA-I and Cu-CHA-M samples exhibit two distinct kinetic regimes at low (648 – 673 K) and high (685 – 773 K) temperature. The Cu-CHA-P control sample prepared to contain predominantly \(Z\) Cu sites was omitted from this analysis because this sample does not facilitate PMO. At low temperatures, rates of ZCuOCuZ formation do not depend on \(P_{O_2}\) and \(P_{H_2O}\).

Moreover, ZCuOCuZ formation rates exhibit a normal kinetic isotope effect \((k_{\text{app},D}/k_{\text{app},D}) = 2 - 3.2, 21\) kPa \(O_2, 0.5\) kPa \(H_2O\) or \(D_2O\) at both 648 and 723 K, which in conjunction with the functional dependence of formation rates on \(P_{H_2O}\) suggests that \(H_2O\) participates in the activation of \(O_2\). Comparisons between rate dependences for ZCuOCuZ formation at low (< 673 K) and high (> 685 K) temperatures suggest that Cu ions are saturated with \(O_2\)-derived intermediates at low temperatures, while Cu sites are not occupied by an \(O_2\)-derived species at high temperatures. ZCuOCuZ formation rates depend similarly on reactant pressures, temperature, and the isotopologue of water used for both Cu-CHA-I and Cu-CHA-M samples, which strongly suggests that the kinetically competent species probed via Raman spectroscopy within these samples are the same (i.e., corresponding to ZCuOCuZ formation from \(Cu^+\) species).

**Scheme 1.** A series of plausible chemical transformations that describe ZCuOH reduction and \(O_2\) activation to form ZCuOCuZ species. \(Z\) denotes a framework Al atom. Step 3 represents ZCuOH auto-reduction and is described by two hypothetical steps (steps 3a and 3b). The \([H_2O]\) within step 6 represents the catalytic role of \(H_2O\) in \(O-O\) bond dissociation. ZCuOCuZ represents the Raman-active intermediate, and a ^ atop an arrow denotes a kinetically-relevant step.

\[
\begin{align*}
2 ZCuOH & \rightarrow ZCu(OH)_2CuZ \quad (1) \\
ZCu(OH)_2CuZ & \rightarrow ZCuOCuZ + H_2O \quad (2) \\
ZCuOH & \rightarrow ZCu^+ + OH^- \\
ZCuOCuZ^* & \rightarrow ZCu^+ + O^2^- \\
ZCu^+ + O_2 & \rightarrow ZCuO_2^- \\
ZCuO_2^- + Cu^+ & \rightarrow ZCuOCuZ \\
ZCuO_2CuZ & \rightarrow ZCuOCuZ + O^2^- \\
[\text{[H}_2\text{O]}] & \rightarrow ZCuOCuZ + O^2^- \\
\end{align*}
\]

Scheme 1 shows a plausible series of chemical transformations that describe the formation of ZCuOCuZ species upon contact between \(O_2\) and Cu(I) ions within Cu-CHA materials. Two ZCuOH moieties react to form a bis(μ-hydroxy)dicopper species (ZCu(OH)_2CuZ; step 1), which then dehydrates to form ZCuOCuZ species (step 2). Alternatively, treatments in He (or CO, vide infra) auto-reduce some fraction of ZCuOH sites to form ZCu^- species (step 3).44 These reactions are exergonic by chemisorbing molecular \(O_2\) to form ZCuO_2^- intermediates (step 4). Reaction between ZCuO_2^- and a proximate ZCu^+ forms a bridging dicopper peroxide complex (ZCuO_2CuZ; step 5).52 These ZCuO_2CuZ species may be active intermediates for PMO or may undergo O-O bond cleavage in the absence (step 6a) or presence of \(H_2O\) (step 6b) to form ZCuOCuZ and release an equivalent of \(O^2^-\). These ZCuOCuZ complexes appear to form irreversibly, because the Raman features attributed to this species (580 cm⁻¹) persist for hours under flowing \(\text{He}\) after activation with \(O_2\) in agreement with previous reports.23 Considering the processes described by Scheme 1, the rapid oxidation of Cu(I) to Cu(II) observed by XAS (Figure 1d) likely corresponds to step 4 or step 5.

To examine whether auto-reduction (steps 1 - 3, Scheme 1) possibly contributes to the formation of ZCuOCuZ, we investigated the influence of different reductive pretreatments of ZCuOH sites. Auto-reduction treatments of ZCuOH species in He at 723 K do not fully reduce all Cu(II) complexes present to Cu(I) (Figure 1).14, 20, 22 Rather, treatments in He yield a combination of ZCu^+, ZCuO_2Cu^-, and ZCu species that do not auto-reduce. The Raman active ZCuOCuZ
species seem unlikely to form via the dehydration of ZCu(OH)₂CuZ (step 2, Scheme 1), because no Raman features appear at elevated temperatures in He (Figure 1a). We also used Raman spectromicroscopy to measure rates of ZCuOCuZ formation over Cu-CHA-I samples treated in carbon monoxide (5 kPa CO, 96 kPa He, 523 K), which fully reduces multinuclear Cu₆O₆ sites to ZCu⁺. Following these CO pretreatments, we observe ZCuOCuZ formation rates that depend on O₂ pressure, temperature, and D₂O/H₂O (Section S8) similarly as Cu-CHA-I instead auto-reduced in He. These comparisons demonstrate that step 1 and step 2 are not kinetically relevant for the formation of the ZCuOCuZ complex after a He auto-reduction treatment. Finally, we must note that this sequence of reactions (steps 1 – 3, Scheme 1) does not involve gaseous O₂; therefore, these steps do not explain the dependence of ZCuOCuZ formation rates on the pressure of O₂ (Fig. 4a and 4d). Consequently, the subsequent and slower formation of the Raman active ZCuOCuZ complexes must reflect steps 6a or 6b if O-O bond rupture limits rates.

To test if our series of hypothetical steps can account for the observed rates of ZCuOCuZ formation, we derive a simple rate expression based on steps 4 – 6 in Scheme 1. In this model, the rate of ZCuOCuZ formation is given by

\[ \frac{d[ZCuOCuZ]}{dt} = k_6[ZCuO₂CuZ] \]  

where \( k_6 \) represents the rate constant for step 6, and \([a]\) represents the number of a given species (i.e., if it is a surface species, such as ZCuOCuZ) or the activity of the gas-phase species (e.g., \( O_2 \)). Under these reaction conditions (i.e., low pressures and high temperatures), the activities are equal to the partial pressure. Step 6b implies that ZCuOCuZ formation involves H₂O, which we rationalize as either solvation of the transition state for the O-O bond rupture or evidence that proton transfer facilitates this event. Invoking the pseudo-steady state assumption on \([ZCuO₂CuZ]\) and \([ZCuO₂]\) species simplifies equation 5 to yield

\[ \frac{d[ZCuOCuZ]}{dt} = k_4[ZCu⁺][O₂] \]  

The expression for \([ZCu⁺]\) arises from a site balance over all likely surface intermediates present in steps 4, 5, and 6

\[ [L] = [ZCu⁺] + [ZCuO₂] + [ZCuO₂CuZ] \]  
\[ [L] = [ZCu⁺](1 + \frac{k_4}{k_5}[O₂] + \frac{k_5}{k_6}[O₂][ZCu⁺]) \]  

where \([L]\) represents the total number of Cu ions that form Raman-active intermediates and the three terms within the parentheses of equation 8 correspond to the relative coverages of ZCu⁺, ZCuO₂, and ZCuOCuZ species, respectively. Substitution of equation 8 into equation 6 yields

\[ \frac{d[ZCuOCuZ]}{dt} \frac{1}{[L]} = \frac{k_4}{k_5}[O₂] + \frac{k_5}{k_6}[O₂][ZCu⁺] \]  

At low temperatures, the chemisorption of O₂ is facile, which suggests that an O₂-derived intermediate (e.g., ZCuO₂⁻) comprises the most-abundant reactive intermediate (MARI). Equation 9 collapses to a simpler form with this approximation

\[ \frac{d[ZCuOCuZ]}{dt} \frac{1}{[L]} = \frac{k_4}{k_5}[O₂] \]  

In contrast, the entropic losses incurred by O₂ chemisorption prevail at higher temperatures, which leads to ZCu⁺ as the MARI and reduces equation 9 to

\[ \frac{d[ZCuOCuZ]}{dt} \frac{1}{[L]} = k_4[O₂] \]  

Equations 10 and 11 quantitatively describe the O₂ dependence on the rates of ZCuOCuZ formation, while the complex and likely non-elementary role of water molecules is captured within \( k_6 \) (Scheme 1).

While Scheme 1 and equations 5 – 9 yield functional expressions that describe O₂ activation kinetics, we note that this series of chemical transformations arises largely from chemical intuition and experimental observations. The unambiguous identification of the elementary steps that lead to ZCuOCuZ formation requires a combination of currently inaccessible synthetic methods and DFT-calculated reaction trajectories. Furthermore, DFT-calculated reaction pathways to quantitatively evaluate the free energy landscape of O₂ activation may not yield meaningful results to interpret experimental data, because the values calculated are a strong function of the Al arrangement that influences the bimetallic Cu configurations chosen. Even nominally “single-site” Cu-CHA samples that contain only 6-MR isolated Al sites possess multiple types of Al-Al pair configurations in 8-MR windows that influence the energetics of bimetallic Cu-oxo species formed.

Collectively, these results present evidence for the involvement of the ZCuOCuZ complex as an active intermediate for PMO and a plausible series of chemical steps for O₂ activation over ZCuOH sites consistent with the observed dependence of O₂ activation on O₂ and H₂O pressure, protons, and temperature. The precise identification of bimetallic Cu species responsible for a specific chemistry (e.g., NO decomposition, CH₃OH synthesis), however, remains an important scientific challenge.

4.0 CONCLUSIONS

Cu-ion exchanged CHA zeolites that contain detectable quantities of ZCuOH activate molecular O₂ to form Cu₇O₇ species active for partial methane oxidation (PMO), nitrogen oxide decomposition, and carbon monoxide oxidation. Temporally-resolved resonance Raman spectroscopy (λex = 532 nm) evinces the genesis of reactive Cu₇O₇ species that form at ZCuOH sites via the activation of O₂. The population of these Raman-active species correlate strongly with increases in the yield of methanol from stoichiometric PMO, whereas, rates of the bulk oxidation of Cu(I) to Cu(II) proceed significantly faster as observed with X-ray absorption spectroscopy. These kinetic comparisons suggest that kinetically-relevant structural rearrangements (and O-O bond rupture) determine formation rates of the Cu₇O₇ complex responsible for methanol formation and that rates of Cu(I) oxidation do not correspond to the formation of the reactive intermediate.
The Raman active Cu-O2 complex possesses a distinct vibrational feature at 603 cm⁻¹ assigned to the deformational mode of the 8-membered ring of a mono(µ-oxo)dicopper intermediate (ZCuOCuZ), through comparisons of measured spectra with Raman scattering frequencies calculated via density functional theory. The rates of O2 activation of ZCuOH-containing CHA samples show two distinct kinetic regimes at low (< 673 K) and high (> 685 K) temperatures. At low temperatures, rates of O2 activation do not depend on O2 pressure, show a first-to-zeroth order kinetic dependence on H2O pressure, and exhibit a normal H2O/D2O KIE. At high temperatures, O2 activation rates increase linearly with O2 pressure, show a weak dependence on H2O pressure, and possess a normal KIE on H2O/D2O. These observations agree with a plausible mechanism in which a fraction of ZCuOH sites initially reduce to form ZCu⁺ sites and readily bind molecular O2 to form ZCuO2 intermediates. ZCu⁺ and ZCuO2 then react to form a ZCuO2CuZ complex. These ZCuO2CuZ intermediates undergo kinetically relevant O-O bond scission, which can be facilitated by water, to form ZCuO2CuZ species and O2⁻.

These data, methodologies, and interpretation provide a basis to understand how Cu ions within zeolites activate O2 and evolve to form species responsible for socially- and environmentally-important oxidation chemistries. The precise identification of the active species and the development of design principles that increase rates and selectivities remains a challenge. These goals motivate the development of synthetic methods to prepare model Cu-zeolites with more uniform Cu structures, innovative kinetic and spectroscopic tools to probe active Cu structures, and connections between experimental data and computational models to understand these complex systems.

ASSOCIATED CONTENT

Supporting Information. CHA synthesis methods, Cu ion exchange, UV-vis spectroscopy, Cu-CHA characterization, in situ X-ray absorption spectroscopy, methanol yields, simulated Raman spectra, in situ Raman spectrokinetics on CO-reduced Cu-CHA-I

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51. For example, over the course of 1 ks, the molar ratio of dioxygen to total Cu within the sample is >5,000.

52. Step 5 is assumed to be irreversible because the heterolytic cleavage of ZCuO$_2$CuZ to form ZCuO$_2$ and ZCu$^+$ is energetically disfavored.
