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Title: "Reactions of Carbonyl Oxide with Aldehydes: Accurate Electronic Structure Methods, Kinetic Insights, and Atmospheric Implications"

Author(s): Chaolu Xie and Bo Long

Responses in blue.

RC1

Xie and Long have used a diverse combination of computational methods to study the reaction of carbonyl oxide (CH_2OO) with a series of aldehydes, and implemented key results in a chemistry-transport model to assess the atmospheric implications. Overall, the manuscript is well written, and the combination of methods used is appropriate to the task. I'm thus happy to recommend publication, subject to some very minor revisions.

Questions to the authors and/or suggestions for further improvement:

1) The authors perform their most accurate calculations for reaction 1, $\text{H}_2\text{COO} + \text{HCHO}$, and also use these results to "anchor" predictions on other reactions (as per the last term in their equation 3). However, as noted in the discussion on page 20, the computational results at 296 K are still 7.3 times higher than the most recent experimental rate. Can the authors comment on possible reasons for this? According to Figure 4, the experimental and computational temperature dependence is also quite different - can this be used to narrow down the source of the discrepancy? Also, given the so-far unresolved discrepancy, the repeated use of phrases like "quantitative accuracy", "underscoring the reliability of our calculated results" (and so on) is perhaps a bit overstated, maybe rephrase a few of these instances.

Author response: We thank the reviewer for raising this important point regarding the discrepancy between theoretical and experimental rate coefficients for the $\text{CH}_2\text{OO} + \text{HCHO}$ reaction. We agree that a noticeable deviation exists between theoretical predictions and available experimental data. At present, it is not entirely clear whether this discrepancy originates primarily from experimental uncertainties or from remaining limitations in the theoretical description.

From the theoretical side, we have carefully examined all factors that could significantly influence the predicted kinetics: Geometric optimization and frequency calculations have been performed using the currently most precise method—CCSD(T)F12a/cc-pVTZ-F12, and the singlet-point energy were characterized using high-level coupled-cluster methods with systematic treatment of basis set convergence and higher-order correlation effects. The computed energetics approach the practical limit of coupled-cluster theory and are expected to be close to the Full CI limit within chemical accuracy. Zero-point energy corrections and thermal contributions were consistently included. All accessible reaction pathways, including pre-reactive complexes, post-reactive complexes, and possible barrierless addition channels, were explored. No energetically competitive alternative channels were identified that would substantially alter the overall rate constant. The rate coefficients were calculated using dual-level strategy which including canonical and variational transition state theory where appropriate, including tunneling, recrossing, anharmonicity, and torsional anharmonicity corrections. Despite this comprehensive treatment, the discrepancy with experiment persists.

We have added the discussion in the revised version "We therefore consider two plausible

explanations: The experimental determination of CH₂OO kinetics may introduce systematic uncertainties. Alternatively, subtle dynamic effects beyond conventional transition state theory (e.g., non-statistical dynamics or complex-forming behavior) may play a role and require further investigation.” in page 21.

2) Please add a reference to the MW2-F12.L scheme when it is mentioned on page 7.

Author response: Thank you for your comments. We have now added the appropriate reference describing the MW2-F12.L scheme in the Supplement.

3) In the discussion on section 2.1., please briefly describe the main differences between W3X-L and GMMQ.L4, and if possible comment on the origin of the 0.24 kcal/mol deviation. The latter goes up to CCSDTQ while the former goes to CCSDT(Q), but given the very small CCSDTQ - CCSDT(Q) contribution this is probably NOT the major source of the deviation (as already noted by the authors on page 14 - basically I'm asking them to elaborate a bit on the “differences between the MW2-F12.L and W2X components” mentioned there).

Author response: Thanks for your comments. The difference between W2X and MW2-F12.L have been listed in Table S7 and S8 in the Supplement. We have rephrased the description in the revised article “We further compared GMMQ.L4 with the W3X-L composite method (Chan and Radom, 2015) for reaction R1. Although both protocols include identical post-CCSD(T) contributions, GMMQ.L4 employs the MW2-F12.L component, whereas W3X-L is based on W2X. Detailed comparisons are provided in Tables S1, S7, and S8. The observed deviation of 0.24 kcal mol⁻¹ indicates that W3X-L does not achieve quantitatively reliable barrier heights for this system. Our analysis shows that this discrepancy primarily originates from the difference between MW2-F12.L and W2X. Specifically, MW2-F12.L includes HF energies, Δ CCSD and Δ (T) correlation contributions, core-valence (Δ (C+V)) corrections, and scalar relativistic (Δ (C+R)) effects, all evaluated with larger basis sets. In contrast, W2X comprises analogous HF, Δ CCSD, Δ (T), and Δ (C+R) terms, but these are computed using smaller basis sets. The calculated results the difference of 0.24 kcal mol⁻¹ comes from the Δ (C+V) and Δ (C+R) terms, which differ by 0.19 kcal/mol and 0.12 kcal/mol, respectively.” in page 7.

4) The discussion in section 2.1 mainly concerns the convergence of results with respect to the level of correlation (highest number of excitations) in the coupled cluster method. This is understandable, as this aspect is the most novel part of the work. However, I note that all the “post-CCSD(T)” corrections are computed with very modest basis sets. (Again, understandable given the demonstrated rapid basis-set convergence of these corrections). Nevertheless, a brief recap of the employed basis set extrapolation (presumably performed at lower levels of theory in the MW2-F12.L scheme) could be helpful to readers. How large basis sets are used to extrapolate e.g. the HF or CCSD or CCSD(T) energies? This question is also related to point 1 above - what remaining error sources could possibly explain a discrepancy of a factor of 7.3...?

Author response: Thanks for your comments. We have added Table S7 to the Supplement, which provides comprehensive details of the MW2-F12.L scheme, including the specific basis sets used for each energy component. Specifically, we have highlighted that MW2-F12.L utilizes significantly larger basis sets compared to the W2X protocol, whilst also accounting for nuclear charge corrections. The 7.3-fold difference may not originate from the basis set used in this work. We have performed the discussion

“We further compared GMMQ.L4 with the W3X-L composite method (Chan and Radom, 2015) for reaction R1. Although both protocols include identical post-CCSD(T) contributions, GMMQ.L4 employs the MW2-F12.L component, whereas W3X-L is based on W2X. Detailed comparisons are provided in Tables S1, S7, and S8. The observed deviation of 0.24 kcal mol⁻¹ indicates that W3X-L does not achieve quantitatively reliable barrier heights for this system. Our analysis shows that this discrepancy primarily originates from the difference between MW2-F12.L and W2X. Specifically, MW2-F12.L includes HF energies, Δ CCSD and Δ (T) correlation contributions, core–valence (Δ (C+V)) corrections, and scalar relativistic (Δ (C+R)) effects, all evaluated with larger basis sets. In contrast, W2X comprises analogous HF, Δ CCSD, Δ (T), and Δ (C+R) terms, but these are computed using smaller basis sets. The calculated results the difference of 0.24 kcal mol⁻¹ comes from the Δ (C+V) and Δ (C+R) terms, which differ by 0.19 kcal/mol and 0.12 kcal/mol, respectively.” In page 7.

5) “Precreation” on line 240 (page 15) should presumably be “pre-reaction”.

Author response: Thanks for your comments. We have corrected this problem.

6) “Barrierless barrier process” on line 340 (page 22) should presumably read just “Barrierless process”.

Author response: Thanks for your comments. We have corrected this problem.

7) Do the “base-version model simulations” mentioned on line 420 (page 23) refer to the GEOS-CHEM simulations performed in this study, or to something else? Please clarify.

Author response: Thanks for your comments. The “base” model using default setting were clarified in 2.5. Atmospheric modeling section.

8) Please explain why the four specific regions/areas in Table 6 were selected. Are these representative for various chemical regimes in the atmosphere, or what is the reasoning?

Author response: Thanks for your comments. These four regions were selected owing to their significant concentration differences in CH₂OO/OH.

9) Moderate (6-12%) reductions in “gas-phase sulfate” (presumably meaning gas-phase sulfuric acid, as sulfate ions are not even stable in the in the gas phase - I do understand the phrasing may originate from the GEOS-CHEM model) were observed in Arctic/sub-Arctic regions during night due to CH₂OO depletion. This is interesting - but to assess the implications better, it would be good to know what the absolute H₂SO₄ concentrations or production rates are in these conditions. A 10% reduction in a number that is already too small to matter is not very impactful, while a 10% reduction of a substantial number is much more important. Please elaborate on this.

Author response: Thanks for your comments. The PH₂SO₄ here refers to gas-phase sulphate, which in GEOS-Chem primarily originates from the gas-phase oxidation of SO₂. Its maximum concentration globally can reach 10⁸, and we have depicted its concentration in the base version in Figure S10. We also mentioned in the revised article “We found that the concentration of gas-phase sulfate can reach 10⁸ molecules cm⁻³ in Mexico region in Figure S10.” On page 26.

RC2

The order-of-magnitude difference between experimental and theoretical rate constants for the $\text{CH}_2\text{OO} + \text{HCHO}$ reaction is a big question whose resolution is highly significant. The theoretical rate constants presented in this manuscript clearly agree better with experiment than previous theoretical predictions (Figure 4), but it would be worthwhile to suggest at least one reason for the remaining discrepancy.

Author response: Thanks for your comments. We thank the reviewer for raising this important point regarding the discrepancy between theoretical and experimental rate coefficients for the $\text{CH}_2\text{OO} + \text{HCHO}$ reaction. We agree that a noticeable deviation exists between theoretical predictions and available experimental data. At present, it is not entirely clear whether this discrepancy originates primarily from experimental uncertainties or from remaining limitations in the theoretical description.

From the theoretical side, we have carefully examined all factors that could significantly influence the predicted kinetics: Geometric optimization and frequency calculations have been performed using the currently most precise method—CCSD(T)F12a/cc-pVTZ-F12, and the singlet-point energy were characterized using high-level coupled-cluster methods with systematic treatment of basis set convergence and higher-order correlation effects. The computed energetics approach the practical limit of coupled-cluster theory and are expected to be close to the Full CI limit within chemical accuracy. Zero-point energy corrections and thermal contributions were consistently included. All accessible reaction pathways, including pre-reactive complexes, post-reactive complexes, and possible barrierless addition channels, were explored. No energetically competitive alternative channels were identified that would substantially alter the overall rate constant. The rate coefficients were calculated using dual-level strategy which including canonical and variational transition state theory where appropriate, including tunneling, recrossing, anharmonicity, and torsional anharmonicity corrections. Despite this comprehensive treatment, the discrepancy with experiment persists.

We have added the discussion in the revised version “We therefore consider two plausible explanations: The experimental determination of CH_2OO kinetics may introduce systematic uncertainties. Alternatively, subtle dynamic effects beyond conventional transition state theory (e.g., non-statistical dynamics or complex-forming behavior) may play a role and require further investigation.” in page 21.

The authors make solid arguments for why CH_2OO ought to be include alongside OH and HO_2 as significant aldehyde oxidants in the atmosphere. The prediction that fluorinated aldehydes react with CH_2OO at near the collision limit is highly significant.

The new electronic structure approaches that allow for the achievement of sub-kcal/mol accuracy for systems with eight non-hydrogen atoms is an impressive methodological advance.

The authors report the useful result that post-CCSD(T) corrections (*i.e.* very high levels of electron correlation) are not necessary for accurate reaction barriers as fluoro or longer alkyl substituents are added to aldehydes.

Author response: Thanks for your comments.

Scientific quality

1). On p. 4, the authors mention the common use of structure-reactivity relationships such

as those proposed by Jenkin et al. It would be good to circle back to them at some point in the manuscript, as the reliability of these estimation methods is an important issue for the atmospheric chemistry community.

Author response: Thanks for your comments. We agree that while the structure-reactivity relationships (SARs) developed by Jenkin et al. are widely used for their efficiency in large-scale models. We have rephased the sentence in the revised version “To address these gaps, atmospheric models have effectively utilized rate constants derived from empirical structure–reactivity relationships (SRRs)—such as those proposed by Jenkin et al. (Jenkin et al., 2018)—which provide a practical and robust framework for large-scale modeling. Given the inherent complexity of computing atmospheric kinetics, these empirical methods remain a primary tool for estimation.” in page 4.

2). Related to ways that the presentation quality can be improved (see below), the authors should briefly explain why their methodology is better than the methods of Chan and Radom. Specifically, the authors should briefly explain why W3X-L is not sufficient for describing post-CCSD(T) effects (p. 7). Also, why is the MW2-F12.L energy corrected stepwise by T-(T) and (Q)-T terms (equation 1), while the W2X energy can be corrected by one term from the (T) level to the (Q) level (equation 3)?

Author response: Thanks for your comments. The discrepancy of 0.24 kcal/mol highlights that for the $\text{CH}_2\text{OO} + \text{HCHO}$ reaction, the basis set requirements for the CCSD(T) base energy and subsequent high-order corrections are critical. GMMQ.L4 addresses this by leveraging the MW2-F12.L scheme to provide a more robust CCSD(T) foundation than the W2X component of W3X-L due to larger basis set listed in Table S7 and S8. The stepwise framework enables the T-(T) and (Q)-T terms to utilize larger basis sets to approach the CBS limit, whereas the Q-(Q) are handled with a meticulously selected smaller basis set to maintain computational feasibility. Additionally, our inclusion of full CCSDTQ rather than the perturbative CCSDT(Q) ensures that the post-CCSD(T) effects are described with higher theoretical fidelity. We have rephased the description in the revised version “We further compared GMMQ.L4 with the W3X-L composite method (Chan and Radom, 2015) for reaction R1. Although both protocols include identical post-CCSD(T) contributions, GMMQ.L4 employs the MW2-F12.L component, whereas W3X-L is based on W2X. Detailed comparisons are provided in Tables S1, S7, and S8. The observed deviation of 0.24 kcal mol⁻¹ indicates that W3X-L does not achieve quantitatively reliable barrier heights for this system. Our analysis shows that this discrepancy primarily originates from the difference between MW2-F12.L and W2X. Specifically, MW2-F12.L includes HF energies, ΔCCSD and $\Delta(\text{T})$ correlation contributions, core–valence ($\Delta(\text{C}+\text{V})$) corrections, and scalar relativistic ($\Delta(\text{C}+\text{R})$) effects, all evaluated with larger basis sets. In contrast, W2X comprises analogous HF, ΔCCSD , $\Delta(\text{T})$, and $\Delta(\text{C}+\text{R})$ terms, but these are computed using smaller basis sets. The calculated results the difference of 0.24 kcal mol⁻¹ comes from the $\Delta(\text{C}+\text{V})$ and $\Delta(\text{C}+\text{R})$ terms, which differ by 0.19 kcal/mol and 0.12 kcal/mol, respectively.” in page 7.

3). On p. 15, there is an assertion that the interconversion TS between C2a and C2b has a low barrier that leads to facile interconversion. However, the M11L TS_{2ISO} 0 K enthalpy is -3.98 kcal/mol, which is essentially identical to the enthalpy of TS2d and higher than the enthalpy of TS2c. Thus, it does not seem obviously valid to assume rapid interconversion of the two five-membered ring complexes.

Author response: Thanks for your comments. To further validate our assertion of “facile

interconversion”, we calculated the rate constants for these processes. Our kinetic analysis reveals that the rate constant for the interconversion via TS_{2ISO} is approximately two orders of magnitude larger than those for the addition pathways (TS_{2c} and TS_{2d}) at the relevant temperature. This significant kinetic acceleration confirms that the interconversion between C2a and C2b is indeed much faster than the addition reaction steps. We have listed the results in Table S27 in Supplement and add the discussion in the revised article “The five-membered ring species C2a and C2b readily interconvert, as the rate constant for the isomerization process is approximately two orders of magnitude larger than that of the addition reaction (Table S26).” in page 23.

4). There is a good justification for the use of M11L as the lower level of electronic structure theory for the direct dynamics calculations. The authors also have performed thorough benchmarking to validate more affordable electronic structure approaches applicable to larger molecules.

Author response: Thanks for your comments.

5). The authors should briefly describe, on p. 16 or in the Supplementary Information, how they determined the number of conformers for larger aldehydes.

Author response: Thanks for your comments. These conformers were generated using the Mst0r 2017 code by rotating the dihedral angles of all reactants and transition states. we have added presentation in the revised article “Conformers for each reactant and transition state were obtained by rotating the dihedral angles listed in Table S10.” In page 16.

6). The authors should briefly explain how the (small) falloff factor of 1.34 (p. 18) is determined for the pressure-dependent rate constants in Table S12.

Author response: Thanks for your comments. The falloff factor is from the ratio of is calculated as the ratio of 0.0316 bar (at 298 K) to 1000 bar. we have added presentation in the revised article “For example, the falloff factor calculated for the CH₂OO + HCHO reaction at 298 K and 0.0316 bar is 1.34 (Table S13). This factor, defined as the ratio of the rate constant at 1000 bar to that at 0.0316 bar, indicates only a weak pressure dependence for this system. This result is in excellent agreement with the findings reported by Luo et al. (Luo et al., 2023).” in Table S18.

7). The observation that, based on their atmospheric modeling, atmospheric CH₂OO concentrations are lowered far more than aldehyde concentrations (Figure 5) is helpful. It is also commendable that the authors refrain from overselling the atmospheric significance of their predicted rate constants.

Author response: Thanks for your comments.

Presentation quality

The authors provide a solid, concise introduction that does a good job of motivating the research.

The authors assume too much familiarity with the theoretical details of their research. This makes parts of the current version of the manuscript not approachable for the large majority of atmospheric chemists who read *Atmospheric Chemistry and Physics*. I recommend the following improvements:

(1). Brief definitions and/or explanations of topics like anharmonicity, re-crossing, the dual-level strategy for calculating rate constants, and the multi-structural anharmonic factor (equations 5 and 6).

Author response: Thanks for your comments. We have added relative explanation in Table A1 of Supplement.

(2). There needs to be a literature reference for the MW2-F12.L scheme, along with a brief description of what the scheme achieves.

Author response: Thanks for your comments. We have added the reference about MW2-F12.L scheme in Table S9 of Supplement.

(3). Equation 2, which presents the basis set extrapolation scheme, must be unpacked; the notation is obscure.

Author response: Thanks for your comments. We have added the explanation “with L=2 for cc-pVDZ and 3 for cc-pVTZ and VTZ(d).”

(4). There appears to be an error in the color scale for the rightmost map in Figure 5. I think the color scale is reversed: most of the world appears white, which should mean that there is virtually no change in predicted sulfate concentration in spite of the predicted faster rate constant for the $\text{CH}_2\text{OO} + \text{HCHO}$ reaction.

Author response: Thanks for your comments. We have amended this error in the revised article.