Supplementary Information: Unraveling the Mystery of “Tech Red” – a Volatile Technetium Oxide

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Single Molecule Simulation Methods

Simulations were performed with the Q-Chem 4.0 package1 using the spin-unrestricted generalized gradient approximation (GGA) functional of Perdew, Burke, and Ernzerhof (PBE)2 with Grimme’s -D33 semi-empirical dispersion correction. The simulations employed the def2-TZVP triple-ζ valence with polarization basis set that uses a small core effective core potential (ECP) to describe the Tc core ([Ar]+3d10) electrons.4,5 The ECP was parameterized to account for scalar relativistic effects. The exchange-correlation functional was evaluated on an Euler-Maclaurin-Lebedev quadrature grid with 75 angular points and 302 radial points per atom. All simulations were performed without consideration for molecular or electronic symmetry. All initial guesses were open shell with broken spin symmetry to avoid trapping the solution in a closed-shell saddle point. The energy convergence tolerance was 10⁻⁹ Eh (Hartree) with an integral cutoff of 10⁻¹⁴ Eh. Structures were optimized with the convergence criterion of a maximum gradient component of 10⁻⁴ Eh/Å and a maximum atomic displacement of 3x10⁻⁴ Å. Vibrational frequencies were evaluated by numeric second derivatives using the analytic gradient. Zero point vibrational energies determined from frequency analysis were included to compare the relative energetics. Previously, we demonstrated that PBE-D33 is very predictive for the structure of Tc2O7,6,7 however pure GGAs are known to produce several spurious TD-DFT charge transfer states and produce larger errors for the excitation energies of non-local charge transfer excited states.8,9 The inclusion
of exact exchange via a long-range correction has been shown to remedy this problem,\textsuperscript{10} and for that reason the excitation energies are computed here with TD-DFT using long-range corrected PBE (LRC-ωPBE) with $\omega = 0.3a_0$.\textsuperscript{8,11}

**Relative Stabilities, Bonding, and Comparative TD-DFT Spectra**

Table S1: Relative stabilities as heat of formation per atom (kcal/mol) compared to an isolated Tc atom, O\textsubscript{2} molecule, and H\textsubscript{2} molecule for each of the lowest energy structures of each stoichiometry and motif. The spin multiplicity of the lowest energy solution is indicated in parentheses.

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Figure S1: Example of a Tc-Tc $\sigma$ bonding orbital
Figure S2: TD-DFT predicted adsorptions for HTcO₄, Tc₂O₇, TcO₃, and TcO₂. A thin line denoting 510 nm has been overlaid.

Figure S3: TD-DFT predicted absorption excitations for the species in Figure 1: purple – 0-br, red – 1-br, blue – 2-br, green – 3-br. The experimental curve (scaled by ¼) for the red product is overlaid along with a thin line denoting 510 nm.¹²
The triplet state of 1-br Tc$_2$O$_5$ is the most stable state of Tc$_2$O$_5$, however a singlet state of the molecule was also simulated. The singlet and triplet spin solutions of 1-br Tc$_2$O$_5$ differ by only 0.17 kcal/mol (triplet is lower) and have similar electronic and molecular structures: a Tc-Tc bond (see Figure S1), an unpaired $d$ electron per Tc, and a Tc-Tc distance of 2.61 and 2.65 Å for the singlet and triplet, respectively. They also have similar predicted spectra, except the target feature is shifted down to 484 nm for the singlet and is a quarter as intense, which confirms that the triplet state is the more likely candidate of the two. Several other functionals were evaluated to test the accuracy of the predicted spectra for 1-br Tc$_2$O$_5$, as may be seen in Figure S4. Each of the functionals predicts a similar spectrum, albeit shifted by at most 0.15 eV, which is a reasonable value for the expected error for predicting $d$→$d$ transitions in transition metal compounds. Therefore, we will use ±0.15 eV as a reasonable confidence interval for the predicted spectra in this work.

Figure S4: The TD-DFT UV-Vis absorption spectra of 1-br Tc$_2$O$_5$ evaluated with several functionals. The experimental curve (scaled by ¼) for the red product is overlaid along with a thin line denoting 510 nm.
Extended Systems

To further test the favorability oligomer formation, canonical ensemble (NVT) plane-wave DFT molecular dynamics simulations were performed with 10 Tc$_2$O$_5$ monomers at a liquid-like density of 1.6 g/mL, about half the density of Tc$_2$O$_7$(s). Plane-wave density functional theory (PW-DFT)$^{17,18}$ canonical ensemble (NVT) molecular dynamics simulations were performed with the Vienna ab initio simulation package (VASP) version 5.4.1 using the PBE-D3$^{2,3}$. The projector augmented wave (PAW) pseudo-potentials formulated for PBE GW calculations were used to represent the ionic cores, and the valence configurations were 4s$^2$4p$^6$5s$^2$5d$^5$ for Tc and 2s$^2$2p$^4$ for O.$^{19}$ The Γ-point alone was used to represent the first Brillouin zone. The plane waves were cut off at 600 eV. The energy convergence criterion for an SCF cycle is $10^{-6}$ eV. The molecular dynamics used a time step of 2 fs and ran for 10 ps. The simulation volume was a cube with a 12 Å side length, and 5 different initial guesses for the placement of 10 optimized bent corner sharing Tc$_2$O$_5$ molecules were used. The initial velocities were generated from a randomly generated Maxwell-Boltzmann distribution around the input temperature of 300 K, and the Nose-Hoover thermostat was used to control the temperature. Along the trajectories stacking and extended chain formation was observed, and edge-sharing Tc connections and bonding of a Tc to a neighboring molecule’s bridging oxygen was also observed. Example snapshots taken from the trajectories exhibiting these oligomerization motifs may be seen in Figure S5. While the time scale, 10 ps, and simulation setup is insufficient to make any concrete conclusions, it does appear that a condensed Tc$_2$O$_5$ would have a very diverse, frequently muting self-reactivity which could easily explain why no solid, crystalline form of ‘tech red’ has ever been successfully isolated unlike Re$_2$O$_5$,$^{20-23}$ which is still known to disproportionate into +4 and +7 species.$^{24}$
Figure S5: Snapshots taken from NVT-MD simulations of 10 1-br Tc₂O₅ molecules in a (12 Å)³ cube displaying several possibilities for self-reactivity of Tc₂O₅ molecules.

Figure S6: TD-DFT predicted UV-Vis absorption spectra for identified dimers and trimers of 1-br Tc₂O₅. The experimental curve (scaled by ¼) for the red product is overlaid along with a thin line denoting 510 nm.¹²
Functional group additions away from basic oxides

Figure S7: The structures, heat of formation per atom, and TD-DFT predicted UV-Vis spectra of the 6 lowest energy conformers of 1-br Tc$_2$O$_5$ + 2OH. The experimental curve (scaled by 1/4) for the red product is overlaid along with a thin line denoting 510 nm. All structures are singlets.

Figure S8: The structures, heat of formation per atom, and TD-DFT predicted UV-Vis spectra of the 6 lowest energy conformers of 1-br Tc$_2$O$_5$ + 2H. The experimental curve (scaled by 1/4) for the red product is overlaid along with a thin line denoting 510 nm. All structures are singlets.
Figure S9: The structures, heat of formation per atom, and TD-DFT predicted UV-Vis spectra of the 5 lowest energy conformers of 2-br Tc$_2$O$_6$ + 2H. The experimental curve (scaled by $\frac{1}{4}$) for the red product is overlaid along with a thin line denoting 510 nm. $^{12}$ d) is a triplet, all the rest are singlets.

Figure S10: The structures, heat of formation per atom, and TD-DFT predicted UV-Vis spectra of the 6 lowest energy conformers of 1-br Tc$_2$O$_6$ + H$_2$O. The experimental curve (scaled by $\frac{1}{4}$) for the red product is overlaid along with a thin line denoting 510 nm.$^{12}$ e) and f) are triplets, all the rest are singlets.
Figure S11: The structures, heat of formation per atom, and TD-DFT predicted UV-Vis spectra of the 6 lowest energy conformers of 2-br TcO₄ + H₂O. The experimental curve (scaled by ¼) for the red product is overlaid along with a thin line denoting 510 nm. d) is a quintet, a,e,f) are triplets, c,d) are singlets.

Figure S12: The structures, heat of formation per atom, and TD-DFT predicted UV-Vis spectra of the 6 lowest energy conformers of 2-br TcO₆ + H₂O. The experimental curve (scaled by ¼) for the red product is overlaid along with a thin line denoting 510 nm. All structures are singlets.
References


(22) Borisova, L. V.; Ryabchikov, D. I.; Yarinova, T. I. Rhenium Compounds in Sulfuric Acid
