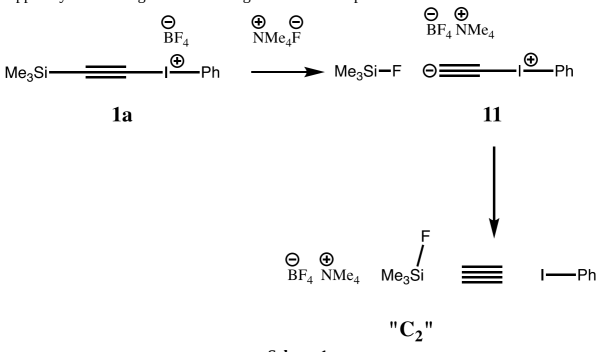
A Thermodynamic assessment of the reported room-temperature chemical synthesis of C₂

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Abstract: Chemical formation of C_2 from an alkynyl iodonium salt precursor is calculated to be strongly endo-energetic, in contrast to the reported chemical synthesis and trapping which is facile at room temperatures. If C_2 is to be formed at such temperatures, a mechanism to counter the unfavourable thermodynamic energies must be identified.

The room-temperature chemical synthesis of C_2 was first reported in the form of a pre-print¹ and has now appeared as a full paper.² The core of the article asserts at its simplest that a transient intermediate **11** formed as $C \equiv C - I^+$ -Ph by treating precursor **1a** with a source of fluoride anion can fragment to singlet C_2 and I-Ph at ambient or low temperatures (Scheme 1). The so-generated C_2 can then be trapped in a variety of ways which are highly suggestive of this putative intermediate. Iodonium species are indeed known in the literature as alkynylation reagents,³ albeit with a proposed mechanism of action for inserting $C \equiv C$ into molecules that does not involve free C_2 . Most of the trapping experiments in the present article are reported in solution, with an implied assertion that singlet C_2 as a discrete species is insufficiently reactive to be captured by solvent rather than by a chemical trap. One experiment is claimed to produce C_2 gas, but in this case the implication is that the C_2 is insufficiently reactive to be trapped by the fritted glass filter through which it must pass.



Scheme1.

One can subject the reaction sequence in Scheme 1 to a reasonableness check based on bond dissociation energies (BDEs). A more quantitative assessment is available through higher level quantum mechanics. The authors themselves have not commented² on this aspect in their

current article, which is based on purely experimental aspects. Addressing firstly the thermodynamics of the core equilibrium;

 $C \equiv C - I^+ - Ph \rightleftharpoons C \equiv C + I - Ph$

Estimates of the experimentally derived BDE of the C-I bond in C=C-I⁺-Ar iodonium salts are in the region of 70-80 kcal/mol.⁴ When the iodonium C-I bond cleaves, it is directly replaced by a fourth bond as represented by C \equiv C, the BDE of which is experimentally estimated at the much lower value of ~17 kcal/mol.⁵ When allowance is made for a gain of ~10 kcal/mol of free energy resulting from increase in entropy, this implies that around 43-53 kcal/mol of bond energy must be recovered by the formation and trapping at ambient temperatures of C₂ itself.

To assess this aspect more quantitatively, the ω B97XD/Def2-SVPD density functional method, with solvation energies estimated using a continuum method set for dichloromethane, has been applied to the reaction shown in Scheme 1 (with NMe₄ replacing NBu₄).⁶ This suggests that the relative free energies Δ G₂₉₈ of **1a**, **11** and the assemblage labelled "C₂" (C₂ + I-Ph + Me₃SiF + Me₄N⁺BF₄⁻) are 0.0, 0.1 and +68.2 kcal/mol respectively. The computed energetics of C₂ itself were calibrated against the two consecutive bond dissociation reactions

 $\mathrm{HC}{\equiv}\mathrm{CH} \rightarrow \mathrm{HC}{\equiv}\mathrm{C}^{\scriptscriptstyle\bullet} + \mathrm{H}^{\scriptscriptstyle\bullet} \rightarrow \mathrm{C}{\equiv}\mathrm{C} + 2\mathrm{H}^{\scriptscriptstyle\bullet}$

for which the thermochemistry has been determined in the gas phase.⁵ This calibration suggests that the relative energy of $C \cong C$ itself is too high by ~28 kcal/mol when computed using the ω B97XD functional and the Def2-SVP basis set. If this correction is applied to the ω B97XD results, then the computed free energy ΔG_{298} of the reaction $1a \rightarrow "C_2"$ is reduced from +68.2 to ~+40 kcal/mol. This is in broad agreement with the simple argument advanced above from experimentally based BDEs (43-53 kcal/mol).

A further, simplified, model at the CCSD(T)/Def2-TZVPPD/SCRF=dichloromethane level was computed, the dissociation of Me-I⁺-C=C⁻ \rightarrow Me-I + C=C⁷ A solvation model is essential, since the ionic reactant is expected to be substantially stabilized by solvation compared to the non-ionic reaction products. At this level of theory, the energy of C₂ itself is computed to be too stable by ~4.6 kcal/mol. With this correction applied, the overall reaction free energy emerges as ΔG_{298} +47.1 kcal/mol, again in the range 43-53 kcal/mol. The former value corresponds to a half-life of a unimolecular reaction (Eyring theory) of ~10¹⁸ hours at 298K.

To add further insights, CCSD(T)/Def2-TZVPP model studies^{8,9} were conducted in which the I⁺-Ph leaving group is replaced by what must be the ultimate leaving group He⁺, itself formed by radioactive decay of tritium. Here, unlike the C-I bond, the BDE of the C-He⁺ bond is tiny (~1 kcal/mol) and its replacement by C₂ does indeed then result in a reasonably exo-energic equilibrium ($\Delta\Delta G_{298}$ -42.2 kcal/mol), augmented again by entropy gain. This serves as a reminder that C₂ itself is a very high energy species.

How can a reaction shown in Scheme 1 and generating the proposed free C₂ overcome a reaction endo-energicity $\Delta\Delta G_{298}$ of +47 kcal/mol, the most accurate estimate obtained by the computations reported here. Eyring theory tells us that at 298K, reactions with respectively a half-life of 1 minute and 1 hour correspond to free energy barriers of 20.0 or 22.5 kcal/mol,

significantly lower than the range of energies predicted above. There are several possibilities to consider.

- Firstly, an as yet un-identified mechanism recovers the energy. Is it possible that sufficient enthalpy can be recovered by say reorganisation of ionic lattice energies so that the resulting free energy barrier $\Delta G_{298}^{\ddagger}$ could promote a sufficiently rapid (half-life <1 hour) reaction at room temperatures? This would allow C₂ to be trapped by another species, but this would compete with ~barrierless reverse trapping by PhI. If so, by what type of mechanism could this recovered energy then be concentrated directly into the carbon-iodine bond in order to cleave it?
- That the range of thermochemical values for the reaction obtained by the three models above is wrongly predicted to be too high by 20-25 kcal/mol.
- Alternatively, one might consider that free C₂ itself is not produced, but instead some other species which must account for the results of the trapping reactions. A possible check on the gas-phase trapping would be to condense whatever species emerges from the dry reaction flask onto a cold-finger at liquid helium temperatures in an argon matrix and subject this directly to spectroscopic (Raman or other) analysis as an alternative to chemical trapping.

If the formation and trapping of C_2 by the chemical reaction sequence shown above can indeed be independently confirmed, then this leaves us with a fascinating chemical challenge of how an otherwise apparently excessively endo-energic reaction can be promoted to viability. No solution to that perplexing dilemma is offered here.

Temperature Chemical Synthesis of C₂. *ChemRxiv*, Preprint., **2019**, DOI: <u>10.26434/chemrxiv.8009633.v1</u> ² K. Miyamoto, S. Narita, Y. Masumoto, T. Hashishin, T. Osawa, M. Kimura, M. Ochiai & M. Uchiyama, *Nature Comm*, **2020**, DOI: 10.1038/s41467-020-16025-x

www.academia.edu/37089575/PROPERTIES_OF_ATOMS_RADICALS_AND_BONDS

⁶ H. S. Rzepa, Double, triple or quadruple? Does a room-temperature chemical synthesis of C₂ resolve the nature of the bond?, *Imperial College Research Data Repository*, **2019**, DOI: <u>10.14469/hpc/5610</u>

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³ T. Kitamura, M. Hasan Morshed, S. Tsukada, Y. Miyazaki, N. Iguchi, D. Inoue, Alkynylation of Benzotriazole with Silylethynyliodonium Triflates. Regioselective Synthesis of 2-Ethynyl-2H-benzotriazole Derivatives, *J. Org. Chem.* **2011**, *76*, 8117-8120, DOI: <u>10.1021/jo2015467</u>

⁴ T. L. Cottrell, The Strengths of Chemical Bonds, 2d ed., Butterworth, London, 1958; B. deB. Darwent, National Standard Reference Data Series, National Bureau of Standards, no. 31, Washington, **1970;** S. W. Benson, J. *Chem. Educ.* **1965**, *42*, 502-518. DOI: 10.1021/ed042p502; J. A. Kerr, *Chem. Rev.* **1966**, *66*, 465-500. DOI: 10.1021/cr60243a001 Compilation at

⁵ S. Shaik, D. Danovich, B. Braida, P. C. Hiberty, The Quadruple Bonding in C₂ Reproduces the Properties of the Molecule. *Chem. Euro. J.*, **2016**, *22*, 4116-412. DOI: <u>10.1002/chem.201600011</u> and Table 2 in D. Danovich, P. C. Hiberty, W. Wu, H. S. Rzepa and S. Shaik, "The Nature of the 4th Bond in the Ground State of C₂: The Quadruple Bond Conundrum" *Chem. Euro. J.*, **2014**, **20**, 6220–6232. DOI: 10.1002/chem.201400356

⁷ H. S. Rzepa, A Thermodynamic assessment of the reported room-temperature chemical synthesis of C₂, *Imperial College Research Data Repository*, **2020**, DOI: <u>10.14469/hpc/7185</u>

⁸ H. S. Rzepa, Startling bonds: revisiting C≣N+, v*ia* the helium bond in N≡C-He+. *Chemistry with a Twist* (*Blog*), **2019**, DOI: <u>10.14469/hpc/5696</u>

⁹ H. S. Rzepa, Diatomics with eight valence-electrons: formation by radioactive decay, *Chemistry with a Twist* (*Blog*), **2019**, DOI: <u>10.14469/hpc/5736</u>