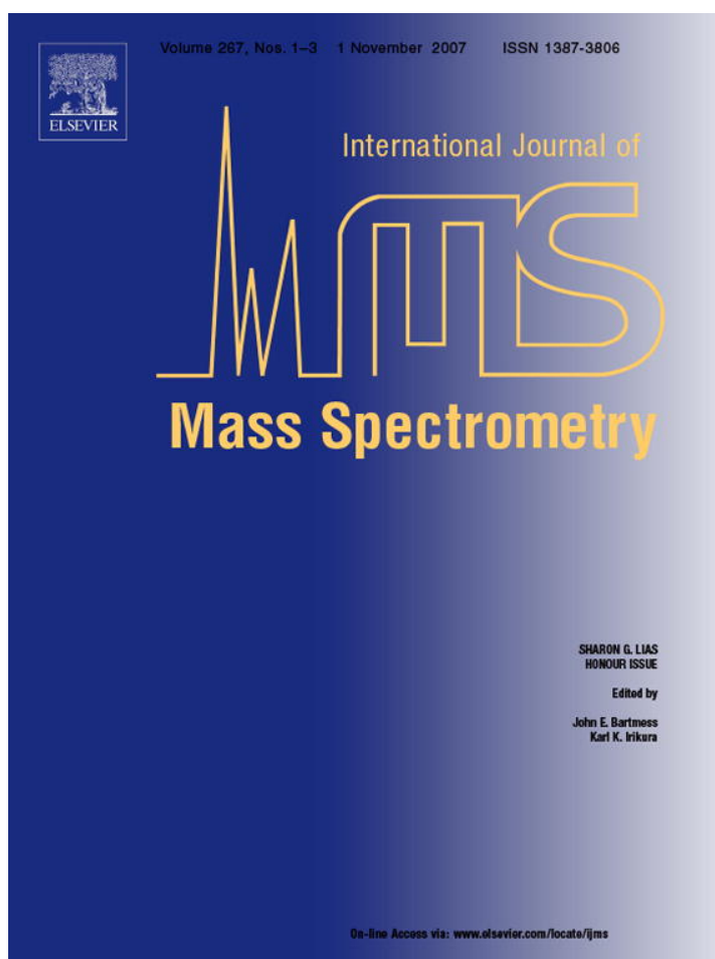


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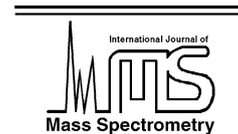


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Transfer of F⁺ versus fluoronium metathesis Competing reactivities of the gaseous fluoroformyl cation

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Received 11 October 2006; received in revised form 26 January 2007; accepted 20 February 2007

Available online 23 February 2007

Abstract

Reactions of unsaturated hydrocarbons with fluoroformyl cation (CFO⁺) and of ketene and diketene with CFO⁺ and CF₃⁺ have been studied in a Flowing Afterglow-Selected Ion Flow Tube (FA-SIFT) in helium at room temperature. Net F⁺ transfer from CFO⁺ to the neutral reactants, which is observed in the majority of cases studied, can be explained in terms of formation of a covalent adduct followed by expulsion of CO. Other product ions include the products of net metathesis, >C=C< + CFO⁺ → >C=C=O + ⁺F=C<, which are observed in nearly half of the hydrocarbons investigated. Reaction of cyclopentadiene with CFO⁺ produces *inter alia* two isomeric C₅H₅⁺ ions, neither of which appears to have the same reactivity, as does the C₃H₅⁺ ion produced by reaction of propargyl cation (C₃H₃⁺) with acetylene. Given that the number of plausible C₅H₅⁺ structures is not very great, one of the ions derived from cyclopentadiene is inferred to have the structure of the cyclopentadienyl cation. The reaction of CFO⁺ with ketene gives F⁺ transfer, yielding FCH₂C≡O⁺, as the major product channel. By contrast, the reaction of CFO⁺ with diketene gives no observable F⁺ transfer. All observed products can be rationalized as coming from initial adduct ions.

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Keywords: Ion–molecule reaction; Halonium metathesis; Cyclopentadienylum; Ketene; Trifluoromethyl cation

1. Introduction

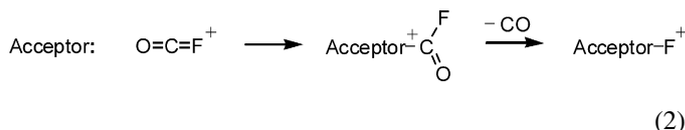
Ionized fluorine, F⁺, is among the strongest Lewis acids known to chemistry. In theory, F⁺ should attach exothermically to every neutral in its electronic ground state except for helium and neon [1]. In practice, isolated F⁺ reacts predominantly by charge transfer [2], because of the high ionization energy of the fluorine atom (17.42 eV) [3].

Reagents that transfer an F⁺ equivalent to anions have found a useful role in synthetic chemistry [4]. F⁺ donation to neutrals, producing fluorinated cations, requires more vigorous electrophiles. Noble gas adducts (such as KrF⁺) [5], adducts of inorganic diatomics (such as FN₂⁺) [6], and coordinatively saturated perfluoro cations (such as BrF₆⁺) [7] have the power to do this. In the gas phase XeF⁺ passes F⁺ to gaseous molecules in competition with Xe⁺ transfer [8]. The F⁺ adduct of acetylene subsequently acts as a Brønsted acid, except where proton trans-

fer is endothermic (as in the reaction with ethylene), in which case it transfers F⁺ [9].



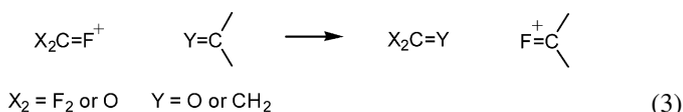
The conventional view of F⁺ transfer is represented by Eq. (1) [1]. A lone pair on the acceptor displaces the donor via an interchange mechanism. This paper describes the chemistry of the fluoroformyl cation, CFO⁺, as an F⁺ donor in competition with other reactions. The nature of this competition suggests the possibility that CFO⁺ transfers F⁺ via other pathways, such as ion–molecule association followed by expulsion of CO, as Eq. (2) represents. Additional alternatives can also be envisaged, as will be discussed below.



Fluoroformyl cation has proved elusive in condensed phases, where initial evidence for its observation has been reinterpreted [10]. But CFO⁺ is a well-characterized species in the gas phase,

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where its structure confirms that it is isoelectronic with carbon dioxide [11]. Needless to say, the carbon of fluoroformyl cation exhibits strong Lewis acidity.



Fluoroformyl cation undergoes fluoronium metathesis [12,13], a reaction long known for gaseous perfluorinated alkyl cations [14–21]. Eq. (3) depicts this metathesis reaction schematically, where transposition of F^+ for oxygen or a methylene group takes place via a 4-membered cyclic transition state. The results presented below, demonstrating that fluoronium metathesis and F^+ transfer compete with one another, raise the question of whether both reactions stem from a common intermediate.

2. Experimental

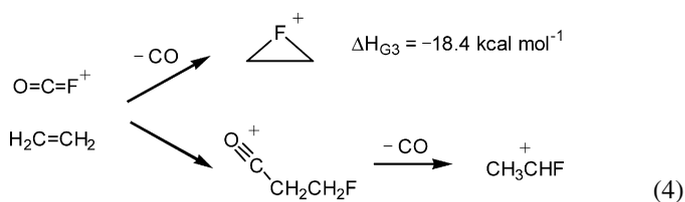
Ion–molecule reactions were examined using a Flowing Afterglow-Selected Ion Flow Tube (FA-SIFT) apparatus, which has been elsewhere described [22]. The carrier gas was helium at a flow tube pressure of 0.46 Torr, and all reactions were studied at room temperature. Reported relative ion abundances $>4\%$ of the total product ionization are rounded to the nearest 5%. Reaction rate constants were extracted from observed ion abundances using published procedures [23]. CFO^+ ions (m/z 47) were generated by plasma discharge through gaseous oxalyl fluoride (FCOCOF, Oakwood Products, Inc.) in the ion source of the SIFT. Ketene was produced by pyrolysis of diketene and stored in a glass bulb at liquid nitrogen temperature when not in use. G3 calculations were performed using the GAUSSIAN03 program suite.

3. Results

Product ion distributions and rate constants for reaction of CFO^+ with various un-saturated hydrocarbons are summarized in Table 1. CFO^+ was not observed to react at all with carbon monoxide (rate constant $k < 2.5 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$). The extent of clustering with hydrocarbons was barely detectable, $\leq 2\%$ of the ion–molecule reaction products (except for styrene). In the case of acetylene, F^+ donation dominates, forming $C_2H_2F^+$ (m/z 45), which exceeds simple adduct formation (m/z 73) by approximately 50:1. Some loss of HF from the adduct ion (m/z 53) is also seen, but only about four times as abundant as the adduct ion itself. The $C_2H_2F^+$ ion reacts with a second acetylene molecule with HF loss to give $C_4H_3^+$ (m/z 51), which then clusters with up to two more acetylene molecules.

The rate constant for CFO^+ with acetylene may be compared with that of CF_3^+ with acetylene, which proceeds entirely by adduct formation and obeys second-order kinetics with a rate constant of $5.5 \times 10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ (slightly slower than reported from a previous SIFT study at 300 K (which was performed at a different carrier gas pressure) [20], but in

agreement with the value from a low-energy ion beam experiment [21]). F^+ donation also dominates the reaction of CFO^+ with ethylene. Because the ion–molecule reaction product from C_2H_4 has the same nominal mass as CFO^+ itself, deuterated ethylene was examined, as Table 1 summarizes. Two stable isomers of the resulting ion have been demonstrated experimentally [24]: the 1-fluoroethyl cation and the bridged fluoriranium ion (which is 28 kcal mol^{-1} less stable [24–29] and which rearranges to the more favorable structure when collisionally activated [29]). The interchange mechanism symbolized by Eq. (1) above would predict initial formation of the latter structure, as the upper pathway in Eq. (4) represents. Alternatively, CFO^+ could add to the double bond with concomitant transfer of fluorine, as the lower pathway depicts. Subsequent loss of CO with a simultaneous 1,2-hydrogen shift would produce the more stable isomer. In contrast to ethylene, heavier alkenes yield product ions from metathesis in competition with F donation.



The thermochemistry of ion–molecule reactions can be gauged using G3 theory. The net reaction of CFO^+ with ethylene to give $CH_3CH=F^+$ (corresponding to the lower pathway in Eq. (4)) is calculated to be exothermic by $-46.8 \text{ kcal mol}^{-1}$. The empirical ΔH has a large uncertainty, owing to variations in the reported heat of formation of gaseous CFO^+ , for which the experimental values range from $160 \pm 11 \text{ kcal mol}^{-1}$ [30] to $178 \pm 2 \text{ kcal mol}^{-1}$ [31]. Given that the heat of formation of $CH_3CH=F^+$ (based on the reported heat of formation and proton affinity of vinyl fluoride [3]) is $159 \text{ kcal mol}^{-1}$, the thermochemistry predicted by the G3 calculation would correspond to a value of $167 \text{ kcal mol}^{-1}$ for $\Delta_f H(CFO^+)$. As summarized for the upper pathway in Eq. (4), G3 theory predicts a substantially smaller exothermicity for the simple F^+ transfer that produces the bridged fluoriranium ion.

The reactions of other hydrocarbons with the fluoroformyl cation are treated below. In addition to the ion–molecule chemistry of CFO^+ with unsaturated hydrocarbons, the reactions of ketene and diketene are also reported and compared with those of CF_3^+ .

3.1. Propene and 2-methyl-1-butene

The major product from higher homologues of ethylene is net hydride transfer to give $[M - 1]^+$ ions or fragments resulting therefrom. In the case of propene, the F^+ adduct ion is observed at m/z 61. Previous studies of this ion suggest that it corresponds to the $(CH_3)_2CF^+$ isomer, regardless of the structure that may have been initially formed [16,17].

Propene is well known to undergo metathesis with CF_3^+ to give CH_3CHF^+ , m/z 47 [22]. Our experiments give the same rate constant and product distribution for that reactant ion as

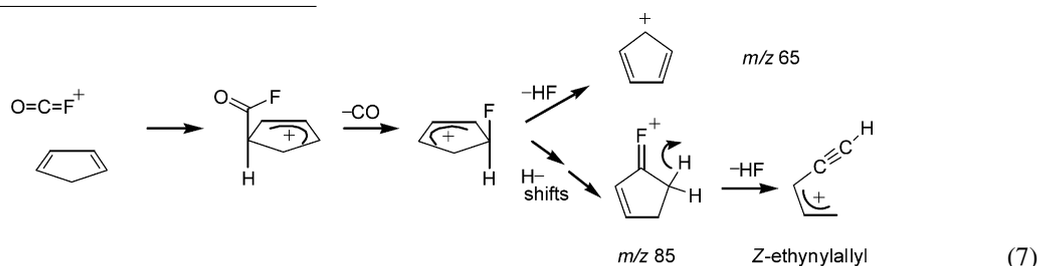
3.4. Vinylacetylene and 1,3-butadiene

Both of these conjugated systems exhibit competition among F^+ donation, metathesis, and net hydride abstraction. The metathesis of vinylacetylene gives a fluorinated propargylic ion, $HC\equiv CCHF^+$ (m/z 57) as a minor product. Of the products listed in Table 1, the $[M - 1]^+$ ion (m/z 51) reacts fastest with the parent neutral. The $HC\equiv CCHF^+$ ion reacts nearly as rapidly, while the ion from F^+ donation (m/z 71) reacts much more slowly.

In contrast to vinylacetylene, half of the ions from reaction of 1,3-butadiene with CFO^+ come from metathesis: $CH_2=CHCHF^+$ (m/z 59). Comparison with the reaction of 1,3-butadiene with CF_3^+ helps interpret the subsequent chemistry of the ion–molecule reaction products. CF_3^+ reacts with butadiene with a rate coefficient of $1.1 \times 10^{-9} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ to yield the m/z 59 metathesis ion as approximately 80% of the

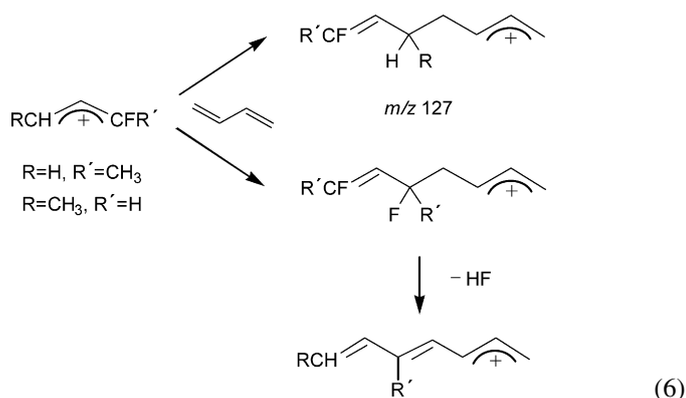
3.5. Cyclopentadiene

Little or no adduct ion is seen from reaction of cyclopentadiene with CFO^+ . The net result of metathesis with CFO^+ is to sever a double bond, attaching CO to one sp^2 center and F^+ to the other. Metathesis with cyclopentadiene therefore yields $O=C=CHCH_2CH=CHCHF^+$, an isomer of the adduct; hence, an upper bound of 1:50 can be placed on the ratio of metathesis to F^+ donation. F^+ donation leads to the dominant product ion, m/z 85. Because hydrogen transpositions occur rapidly within cyclic cations, we infer that this $C_5H_6F^+$ has the same structure as the ion produced by metathesis of 2-cyclopentenone with CF_3^+ [13], drawn in the lower pathway of Eq. (7). DFT calculations predict that ion to correspond to the minimum energy for that formula [13].



ions, with virtually all of the remainder corresponding to loss of HF from the CF_3^+ adduct (m/z 103). The m/z 59 ion then goes on to react with another molecule of butadiene to give the further adduct (m/z 113) and the adduct minus HF (m/z 93) in nearly equal proportions. The same products are observed from the subsequent reaction of the m/z 59 from CFO^+ and butadiene.

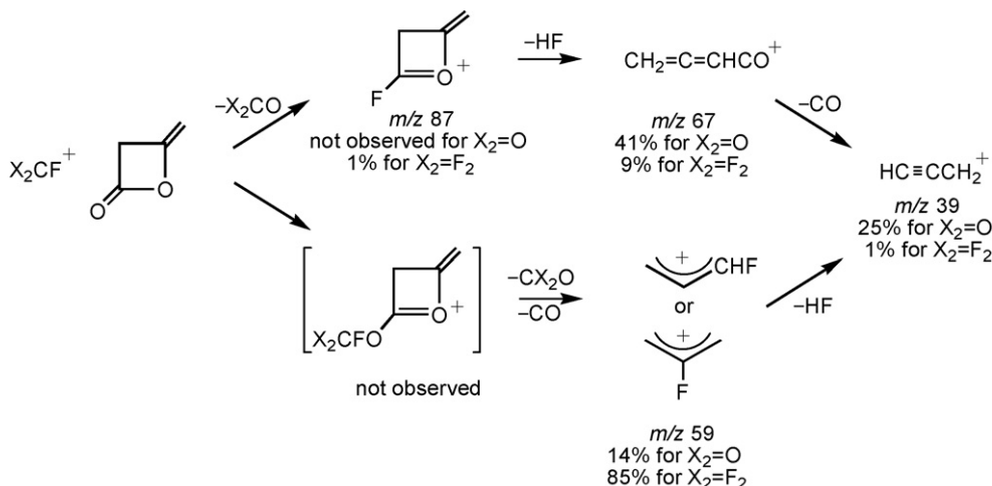
F^+ transfer from CFO^+ to butadiene gives m/z 73, which reacts with another molecule of butadiene to give a second adduct (m/z 127). The $[M - 1]^+$ from butadiene also adds to a molecule of butadiene (m/z 107) at roughly the same rate. A variety of constitutional isomers can be envisaged for the m/z 73 ion. DFT calculations predict that the pair of fluoroallylic cations drawn to the left in Eq. (6) represent the most stable ones [12]. If, indeed, the F^+ adduct of butadiene possesses either of these structures (or a mixture of the two), the stability of the m/z 127 adduct of a second butadiene molecule suggests that the fluorinated end of the m/z 71 is not adding to the butadiene terminus. Otherwise, one would expect prompt loss of HF to generate a cation with extended conjugation, as the lower reaction in Eq. (6) depicts.



The $[M - 1]^+$ ion from cyclopentadiene, m/z 65, can arise from $C_5H_6F^+$ via either of two pathways for HF expulsion, as Eq. (7) depicts. A 1,2-elimination would give the cyclopentadienyl cation, as the upper pathway represents. Alternatively, hydrogen shift can lead to the fluoroallylic m/z 85 ion drawn in Eq. (7) (the global minimum on the $C_5H_6F^+$ potential energy surface [13]), which can then eliminate HF with ring opening to give the Z-ethynylallyl cation.

In a separate series of experiments, $C_5H_6F^+$ was generated using CF_4 plus 2-cyclopentenone [13] in the afterglow source and injected into the SIFT. Collisions with the flow gas cause some of the ions to lose HF. The 7.9:1 mixture of $C_5H_6F^+$ and $C_5H_5^+$ was then reacted with allene in the SIFT. The former ion reacts slowly ($k < 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$), while the latter displayed biexponential decay, producing an adduct of $C_5H_5^+$ with allene, $C_8H_9^+$ (m/z 105). Approximately 55% of the $C_5H_5^+$ reacted with a rate coefficient of $6 \times 10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, while the remaining 45% reacted more slowly by a factor of 100. The $C_5H_6F^+$ ion also reacts with allene to give some adduct ion ($C_8H_{10}F^+$, m/z 125), as well as fragments from loss of C_2H_2 ($C_6H_8F^+$, m/z 99) or C_2H_4 ($C_6H_6F^+$, m/z 97), in approximately a 15:25:60 ratio.

In another series of experiments, 2-cyclopentenone was reacted with CFO^+ in the afterglow source, and the resulting $C_5H_5^+$ ion injected into the SIFT. Some $HC\equiv CCH_2^+$ was formed by collision, which is then observed to cluster rapidly with up to five molecules of acetylene, consistent with previous reports [32–34]. The $HC\equiv CCH_2^+$ ion also clusters with benzene, as do its adducts with one or two molecules of acetylene. In order to examine $C_5H_5^+$ formed from the 5-membered ring, the mixture of $C_3H_3^+$ and $C_5H_5^+$ (approximately a 4:6

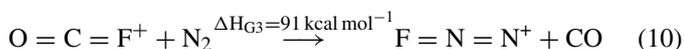


Scheme 1.

The hypothesis that metathesis takes place finds confirmation in the reaction of diketene with CF_3^+ , which goes with a rate coefficient $k = 3.4 \times 10^{-9} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$. Here a small amount of F^+ -for-O metathesis product is detected (m/z 87), in addition the $\text{C}_4\text{H}_3\text{O}^+$ ion from its loss of HF (m/z 67). Apparently the $\text{C}_4\text{H}_3\text{O}^+$ is not produced with as much internal energy as the ion from CFO^+ , since much less C_3H_3^+ is observed. The major ion (m/z 59) comes from expulsion of CF_2O and CO from the CF_3^+ adduct of the carbonyl. Expulsion of HF from this ion could provide an alternative pathway to the propargyl ion, as Scheme 1 summarizes.

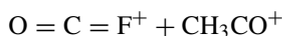
4. Discussion

In solution, F^+ donation takes place as Eq. (1) represents [1]. It is hard to imagine any alternative pathway by which a noble gas adduct or a coordinatively saturated perfluoro cation might transfer F^+ , except via nucleophilic attack at fluorine. The fluoroformyl cation, by contrast, is coordinatively unsaturated, and it can donate F^+ by first forming a covalent adduct at carbon, as Eq. (2) represents. The upper part of Eq. (7) above illustrates that pathway for cyclopentadiene. This hypothesis offers a new mechanism for F^+ donation.



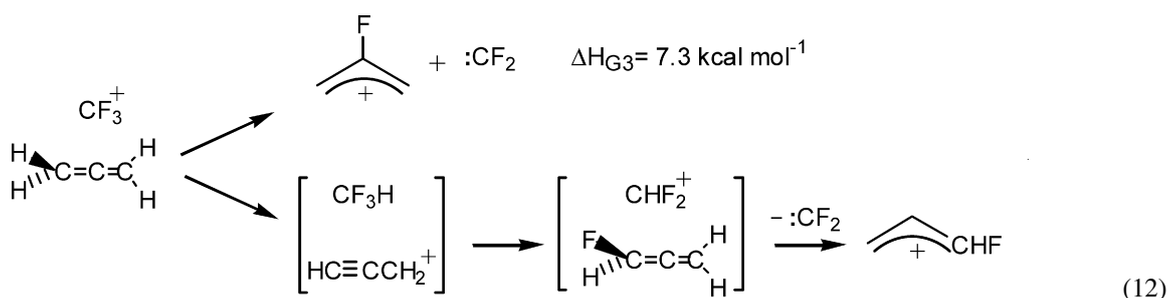
The experimental evidence for this mechanism is buttressed by G3 calculations. First of all, carbon monoxide binds F^+ much

more tightly than does molecular nitrogen. The G3 endothermicity of the transfer from CO to N_2 (shown in Eq (10)), predicts the F^+ affinity of CO to be $\geq 230 \text{ kcal mol}^{-1}$. In other words, the two isoelectronic, linear triatomic ions in Eq (10) should have different chemical properties. FN_2^+ donates F^+ to atomic xenon⁹ and fluorinates methane in solution [36], while CFO^+ ought to be completely inert towards those acceptors.



CFO^+ is a strong electrophile. G3 calculations on the hypothetical transfer of carbon monoxide between two cations shown in Eq. (11) predict a heat of formation of $\text{O}=\text{CFCO}^+$ that implies the attachment of CFO^+ to carbon monoxide should be exothermic by 19 kcal mol^{-1} . Failure to observe clustering of CFO^+ with CO under our SIFT conditions therefore suggests that, if adduct ions form between CFO^+ and other neutrals, they must then undergo unimolecular rearrangement in order for ion–molecule reaction products to be seen.

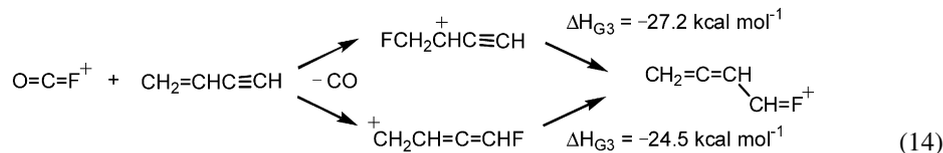
Not long ago, Tsuji et al. reported that gaseous CF_3^+ donates F^+ to allene [37]. They proposed that this reaction takes place via formation of an adduct ion, followed by expulsion of difluorocarbene. In part, their rationale derived from the fact that the major competing reaction is loss of HF from the CF_3^+ adduct ion. Since difluorocarbene is a high energy species, net F^+ transfer from CF_3^+ to a neutral molecule requires a very good acceptor.



(12)

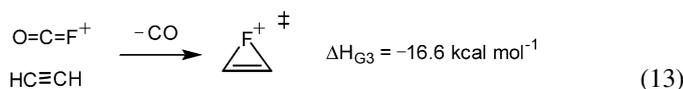
Thermochemical arguments support the contention that CF_3^+ does not donate F^+ via the mechanism of Eq. (1). If that were the case, then transfer would have to occur either to one of the terminal carbons (forming a vinyl cation) or else to the center carbon (forming the more stable allylic cation). The latter transfer

constitutional isomer on the $\text{C}_4\text{H}_4\text{F}^+$ potential energy surface corresponds to a fluorinated carbocation conjugated with an allenyl group, drawn to the right in Eq. (14). This ion (in its more favored *E*-geometry) could conceivably be formed by exothermic hydrogen shifts following F^+ transfer, as Eq. (14) portrays.



is depicted in the upper pathway in Eq. (12). G3 calculations predict this reaction, which produces the 2-fluoroallyl cation, to be significantly endothermic, as summarized for that pathway.

In order for CF_3^+ to donate F^+ to allene exothermically, the lowest energy structures must be formed (1-fluoroallyl cations, as depicted in the lower pathway of Eq. (12)). This reaction necessitates hydrogen rearrangement, either by means of a covalent adduct (as proposed by Tsuji et al. [37]) or via a succession of ion-neutral complexes, as the lower pathway of Eq. (12) represents. While the available experimental data do not differentiate between these alternative mechanisms, the fact that CF_3^+ is also reported to donate F^+ to propyne [37] is consistent with the same sequence of ion-neutral complexes as is drawn above for the reaction of allene.



CFO^+ releases carbon monoxide when it transfers F^+ . From a thermochemical standpoint, it can therefore react with a much greater variety of acceptors than can CF_3^+ . For instance, as Eq. (13) summarizes, formation of the bridged fluorirenium ion from acetylene is calculated to be exothermic, even though that bridged ion represents a transition state ($\text{CH}_2=\text{C}=\text{F}^+$ being the only stable isomer). If formed, the bridged ion must rearrange to the stable isomer, which is more stable by 40 kcal mol^{-1} , according to G3 calculations.

Three mechanisms can therefore be envisaged for F^+ transfer from CFO^+ : interchange (Eq. (1)), adduct formation followed by rearrangement (Eq. (2)), and hydride abstraction followed by a sequence of ion-neutral complexes (as exemplified by Eq. (5)). The experimental evidence for Eq. (2) comes from the competition between F^+ transfer and metathesis in the C_4 hydrocarbons summarized in Table 1. For reactants with fewer carbons, F^+ transfer takes place, but no metathesis is seen. For reactants with ≥ 5 carbons, metathesis is observed, but no F^+ transfer. We ascribe the failure to detect F^+ transfer to these heavier reactants to the instability of the adducts with respect to unimolecular elimination of HF.

Vinylacetylene is the lightest hydrocarbon for which we observe metathesis in competition with F^+ transfer. If one assumes that rearrangement cannot occur concertedly with the interchange mechanism of Eq. (1), then F^+ must be donated to one end or the other, as represented in Eq. (14). Uggerud [38] has performed calculations showing that the lowest energy

Alternatively, hydride abstraction mechanism followed by interconverting ion-neutral complexes (as exemplified by Eq. (5)) is also possible, given that $[\text{M} - 1]^+$ ions are prominent ion-molecule reaction products. However, the occurrence of metathesis suggests a more economical option, namely that a single adduct ion accounts for all of the observed product ions.

The hydride abstraction mechanism pathway can be ruled out in at least one case. Thermochemical estimates show that this mechanism cannot be operating in the case of ketene, since the $[\text{M} - 1]^+$ ion, HC_2O^+ , has a very high heat of formation [39]. HC_2O^+ is calculated to exist as a ground state triplet [40], and G3 calculations gauge the triplet-singlet gap to be 20.3 kcal . G3 theory predicts the endothermicity of hydride abstraction from ketene by CFO^+ to be $39.3 \text{ kcal mol}^{-1}$, so great that it is hard to imagine that the energy liberated by a noncovalent ion-molecule association could be large enough to overcome the thermochemical barrier. The predominance of transfer to ketene must therefore be ascribed to interchange or decomposition of an adduct ion (or both). Since, as Eq. (8) summarizes, a covalent adduct is a necessary precursor to protonated carbon suboxide, F^+ transfer via an intermediate adduct ion represents a plausible mechanism.

5. Conclusions

Transfer of F^+ via a covalent adduct ion, Eq. (2), accounts for the diversity of products seen when the fluoroformyl cation reacts with a variety of unsaturated molecules. This mechanism posits a common intermediate, the adduct with CFO^+ , for two competing pathways, fluoronium metathesis versus F^+ transfer. Both of these reaction pathways are observed, for instance, when CFO^+ reacts with vinylacetylene or 1,3-butadiene. CFO^+ adduct ions with unsaturated hydrocarbons cannot, in many cases, be observed to any great extent under the reaction conditions (except in the case of styrene), because their formation is highly exothermic and they have low barriers for expulsion of HF, CO, or ketene. The hypothesis of an intermediate adduct ion unifies a large number of otherwise disparate experimental data.

Acknowledgement

This work was supported by NSF grant CHE0306515. P.F.W. acknowledges the award of a Marsden Postdoctoral Fellowship.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.ijms.2007.02.019.

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