IR-Spectroscopic and X-ray-Structural Study of Vinyl-Type Carbocations in Their Carborane Salts

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ABSTRACT: The butylene carbocation in its salts with anions CHB\textsubscript{11}\textsubscript{I}F\textsubscript{1}\textsuperscript{−} and CHB\textsubscript{11}Cl\textsubscript{1}\textsuperscript{−} forms isomers CH\textsubscript{2}≡C−CH\textsubscript{2}−CH\textsubscript{3} (I) and CH\textsubscript{2}≡C≡CH−CH\textsubscript{3} (II), which were characterized here by infrared (IR) spectroscopy and X-ray diffraction analysis. The strongest influence on the structure of the cations is exerted by the electron-donor substituents to which the electron-donor substituents are attached, meaning that the donation of electron density from the Cl atom quenches the positive charge on the C≡C bond through which the C≡C bond more strongly, and the C≡C stretch frequency increases so much that it even exceeds that of neutral alkene analogues by 35–65 cm\textsuperscript{−1}. An explanation is given for the finding that upon stabilization of the vinyl cations by polyatomic substituents such as silylium (SiMe\textsubscript{3}) and t-Bu groups, the stretching C≡C frequency approaches the triple-bond frequency. Namely, the scattering of a positive charge on these substituents enhances their donor properties so much that the electron density on the C≡C bond with a weakened charge becomes much higher than that of neutral alkenes.

1. INTRODUCTION

Vinyl carbocations are known as a class of reactive intermediates important for organic synthesis\textsuperscript{1} and have been studied extensively.\textsuperscript{2−12} They are considered highly reactive and therefore uncontrollable intermediates that are difficult to investigate. Nonetheless, research by Mayr and co-workers shows that the reactivity of vinyl cations has been exaggerated.\textsuperscript{3} It is even weaker than that of diarylcarbenium cations, some of the stabllest trisubstituted cations. Recently, it was demonstrated that the benzylic carbocation has carbine-like reactivity: it acts as a strong protonating agent, thereby converting into a carbine molecule, whose high reactivity can be perceived as the reactivity of the carbocation.\textsuperscript{4} The same conclusions have been reached by Niggemann and Gao: the high reactivity of vinyl cations is a myth: they are converted to reactive intermediates with carbine-like reactivity.

Stabilization of vinyl cations by nucleophilic substituents decreases their reactivity and facilitates studies in this field. Thus, the use of alkyl,\textsuperscript{6−9} aryl,\textsuperscript{10−11} and other substituents,\textsuperscript{7,12−14} that effectively delocalize the positive charge on themselves has made it possible to obtain the salts of vinyl cations that are stable at room temperature. Their study by X-ray, infrared (IR), and NMR spectroscopy has revealed one unusual property:\textsuperscript{7,11,12} electron density on the C≡C bond, to which the electron-donor substituents are attached, significantly exceeds that of neutral alkene molecules. It seems that the positive charge on the C≡C bond promotes a strong increase in its electron density supplied by substituents. An explanation for this finding has not been reported.

Recently, vinyl-type isobutylene\textsuperscript{15} and propylene carbocation salts\textsuperscript{16} were obtained; they appear to be thermally stable up to 150 °C. Their investigation by X-ray crystallography and IR spectroscopy indicates that they strongly differ from stabilized cations by their νC≡C frequencies, which are significantly lower than those of the neutral alkenes (by ~160–180 cm\textsuperscript{−1}), which can be expected. Vinyl cations are strongly affected by the anionic environment. For example, the crystal lattice stabilizes the most unstable (according to quantum-chemical prediction\textsuperscript{17}) isobutylene isomer (CH\textsubscript{3})\textsubscript{2}C≡C=H, or else, in amorphous salts, it ionically pairs with the anion. Another peculiarity of vinyl cations is that the calculated frequency of their C≡C stretch is completely inconsistent with an experimental one: the difference reaches 360 cm\textsuperscript{−1} for the propylene cation\textsuperscript{16} and 330 cm\textsuperscript{−1} for the isobutylene...
cation.\textsuperscript{15} The same unacceptable discrepancies between the calculated and experimental IR frequencies have been noted for saturated alkane carbocations.\textsuperscript{18} Nevertheless, quantum-chemical calculations with the same basis set are quite applicable to stabilized vinyl cations with substituents on which the positive charge is effectively dissipated.\textsuperscript{7,24}

In the present work, we report the preparation of salts of other isomers of vinyl cation C\textsubscript{3}H\textsubscript{7}+, including monochlorinated cation C\textsubscript{2}H\textsubscript{5}Cl\textsuperscript{+}. As counterions, carbaborane anions CHB\textsubscript{11}F\textsubscript{11}\textsuperscript{−} and CHB\textsubscript{11}Cl\textsubscript{11}\textsuperscript{−} were chosen because of their extreme inertness.\textsuperscript{19} A comparison of the features of IR-spectroscopic properties and structures of isomers of carbocations C\textsubscript{2}H\textsubscript{5}Cl\textsuperscript{+}, C\textsubscript{3}H\textsubscript{7}+, and C\textsubscript{3}H\textsubscript{6}+ allows one to clearly see the general picture of the properties of vinyl carbocations (with the help of the literature data), both nonstabilized and stabilized ones. Quantum-chemical calculations were not carried out in this work owing to their inapplicability to nonstabilized vinyl cations\textsuperscript{15,16} and because of easy interpretation of their most characteristic IR bands without calculations.

2. RESULTS

2.1. Chloronium Cation as a Precursor for Obtaining Vinyl Carbocations. Injection of 1,4-dichlorobutane vapors into an IR cell reactor with a thin film of HCHB\textsubscript{11}Cl\textsubscript{11} acid (hereafter abbreviated as H(Cl\textsubscript{11})) led to their interaction involving HCl elimination (Figure 1). Upon completion of the reaction, the characteristic IR spectrum of the acid disappeared and the absorption intensity of gaseous HCl reached a maximum. That is, the known reaction of chloroalkanes with superacid proceeds via the first stage\textsuperscript{20} according to eq 1

\[
\text{C}_2\text{H}_4\text{Cl}_2 + \text{H(Cl}_{11}\right) \rightarrow \text{C}_2\text{H}_4\text{Cl}^+(\text{Cl}_{11}^-) + \text{HCl}
\]  

The IR spectrum of the resultant salt contains an intense band at 581 cm\textsuperscript{−1} typical for the C−Cl−C group of the chloronium ion.\textsuperscript{20,21} Two types of chloronium cations can arise under these conditions. The first one emerges if the C\textsubscript{2}H\textsubscript{4}Cl\textsuperscript{+}
cation formed according to eq 1 interacts with a molecule of C\textsubscript{2}H\textsubscript{4}Cl\textsubscript{2} to form di-(chlorobutyl)chloronium

\[
\text{C}_4\text{H}_8\text{Cl}^+(\text{Cl}_{11}^-) + \text{C}_2\text{H}_4\text{Cl}_2 \rightarrow (\text{C}_4\text{H}_8\text{Cl})_2\text{Cl}^+
\]

On the other hand, chloronium cations R−Cl\textsuperscript{−}−R with more than three carbon atoms in R decompose at room temperature with the release of HCl and carbocation formation.\textsuperscript{21} Therefore, this pathway should be excluded. The second type of chloronium cation arises if the formed C\textsubscript{2}H\textsubscript{4}Cl\textsuperscript{+} cation cyclizes to (CH\textsubscript{2}CH\textsubscript{2})\textsubscript{2}Cl\textsuperscript{+}. Such a cation should be close in stability to diethylchloronium (CH\textsubscript{2}CH\textsubscript{2})\textsubscript{2}Cl\textsuperscript{+}, which is stable at room temperature but will decompose when heated. Indeed, heating of the chloronium salt obtained in the IR cell reactor to 150 °C destroyed the cation with the disappearance of the 582 cm\textsuperscript{−1} band of chloronium group C−Cl−C and the emergence of a characteristic absorption pattern of gaseous HCl in the IR spectrum (Figure 1, black spectrum). The intensity of its absorption reached 90% of that of HCl generated by reaction 1. This finding proves that chloronium cation (CH\textsubscript{2}CH\textsubscript{2})\textsubscript{2}Cl\textsuperscript{+} actually forms, which decomposes with increasing temperature with the release of one HCl molecule and the emergence of the C\textsubscript{2}H\textsubscript{4}Cl\textsuperscript{+} carbocation

\[
\text{C}_4\text{H}_8\text{Cl}^+(\text{Cl}_{11}^-) \rightarrow \text{C}_2\text{H}_7^+(\text{Cl}_{11}^-) + \text{HCl}
\]  

Chloronium salt C\textsubscript{2}H\textsubscript{4}Cl\textsuperscript{+}(Cl\textsubscript{11}−) can be obtained in a weighable amount by direct interaction of a small amount of liquid 1,4-dichlorobutane with an acid powder (wetting without an excess), followed by washing with a small volume of cold dichloromethane (DCM) and drying in vacuum. The attenuated total reflectance (ATR) IR spectrum of the obtained white powder of the salt did not differ from that obtained in the IR cell reactor.

In the same way, we prepared a chloronium salt from the CHB\textsubscript{11}F\textsubscript{11}− anion (hereafter abbreviated as {F\textsubscript{11}−}) using the H(F\textsubscript{11})\textsuperscript{−} acid. Its IR spectrum with the most characteristic band ν\textsubscript{as}C−Cl−C at 584 cm\textsuperscript{−1} is consistent with that of the salt of the {Cl\textsubscript{11}−} anion (Table S1).
Both salts of carbocations $\text{C}_2\text{H}_8\text{Cl}^+$ and $\text{C}_2\text{H}_4^+$ proved to be soluble in DCM. Crystals were grown from these solutions, which were studied by X-ray crystallography.

2.2. Chlorinated Vinyl Carboxtation. We were unable to obtain crystals of the chloronium salt for X-ray-structural analysis. Nonetheless, the composition and schematic representation of the chloronium cation structure can be determined from a set of IR-spectroscopic assays and other experimental data. The main arguments in favor of the proposed composition and structure of the chloronium cation—which follow from the IR-spectroscopic data and the conditions for the preparation of its salts—are

1) the cation is formed by the elimination of anion Cl$^-$ from the $\text{C}_2\text{H}_4\text{Cl}_2$ molecule by reaction 1;

2) its IR spectrum contains the band of the asymmetric stretching vibration of the bridged Cl atom at 581 cm$^{-1}$, which is characteristic of the chloronium group $\text{C}--\text{Cl}^-$;

3) when the salt is heated to 150 °C, the cation decomposes according to eq 2 with the release of one HCl molecule, and the absorption band of the chloronium group disappears in the IR spectrum;

4) in the frequency range of stretching CH vibrations, four bands are present (Figure 1). They correspond to two nonequivalent CH$_2$ groups: the two frequencies at 3050 and 2971 cm$^{-1}$ virtually match the bands $\nu_{\text{CH}_2}\text{Cl}_2\text{(Cl)} = 3038 \text{ cm}^{-1}$ and $\nu_{\text{CH}_2}\text{(Cl)} = 2974 \text{ cm}^{-1}$ of the CH$_2$--(Cl$^+$) group in the crystalline salt of diethylchloronium, (CH$_2$CH$_2$)$_2\text{Cl}^+\text{Cl}^-\text{(Cl)}$; and two others (at 2986 and 2945 cm$^{-1}$) are typical for the frequencies $\nu_{\text{CH}_2}\text{Cl}$ and $\nu_{\text{CH}_2}$ of aliphatic CH$_2$ groups.

These arguments are enough to assert that the cation under study is cyclic butylichloronium

$$\text{CH}_2\text{CH}_2\text{CH}_2\text{Cl}^+\text{CH}_2\text{CH}_2$$.

The chloronium salt is well soluble in DCM. Incubation of the solution above the hexane vapor causes the crystals to appear. The yield of crystals from this solution is low, and their formation takes several days, which may be an indirect sign that the crystals derive not from the initial compound but rather from the products of its secondary reactions.

X-ray structure of the crystals revealed that this is a $\text{C}_2\text{H}_8\text{Cl}^+\text{(Cl)}$ salt with discrete cations and anions. The cationic part of the salt consists of two crystallographically nonequivalent C4 cations containing a Cl atom. One of the cations is more disordered. Therefore, to analyze the structure of the cations, we utilized additional information following from the IR spectra of these crystals.

Figure 2 shows an ATR IR spectrum of the single crystal flattened under pressure on the surface of a diamond of the ATR accessory. Sequentially recorded spectra indicated their time dependence. The first one contains two intense bands at 1710 and 1680 cm$^{-1}$, which can belong only to C=C stretch. With successive recordings at short intervals (tens of seconds), the intensity of the 1710 cm$^{-1}$ band decreased until it disappeared, and the 1680 cm$^{-1}$ band increased accordingly. Therefore, the crystalline salt contains two isomers of the vinyl cation, which we will designate as A (with $\nu\text{C}--\text{C}$ at 1680 cm$^{-1}$) and B (with $\nu\text{C}--\text{C}$ at 1710 cm$^{-1}$). Isomer B is stabilized by the crystal lattice and, upon its destruction, transforms into isomer A, which is more stable in the amorphous phase. Subtracting the spectrum of cation A from the spectrum of the mixture of cations A and B, we can isolate the spectrum of cation B (Figure 1, green).

The spectra of isomers A and B are very different from that of the starting chloronium salt. They each contain one band of C=Cl stretch at 768 and 733 cm$^{-1}$, respectively, which are characteristic of C--Cl bands. In the frequency range of stretching C--H vibrations, three narrow bands are observed: at 2959, 2928, and 2872 cm$^{-1}$, typical of CH$_2$ groups of alkanes. The spectrum of isomer A contains a very broad absorption pattern in the frequency range of C--H stretching (2930 cm$^{-1}$) and bending (1427 cm$^{-1}$; Figure 2) vibrations, which is a sign of fast proton exchange at the IR spectroscopy time scale, known for alkane carbocations. Therefore, it can be expected that the isomer contains CH$_3$ or CH groups, whose protons are involved in fast exchange.

Thus, according to the IR spectra, the cation of the crystalline salt $\text{C}_2\text{H}_8\text{Cl}^+\text{(Cl)}$ consists of two isomers A and

![Figure 2. ATR IR spectrum (with ATR correction) of the crystal salt of C$_2$H$_8$Cl$^+$Cl$_{1+}$ after X-ray-structural analysis, as recorded within the first minute after crushing on the ATR accessory (red) and repeating the recording in 3–4 min (blue). The isomer B spectrum, obtained by subtracting the "blue" spectrum from the "red" spectrum is green. Broad intense absorption at ~1400 cm$^{-1}$ is shaded in light blue.](https://doi.org/10.1021/acs.omega.2c03025)
B, which have nonequivalent C=\text{C} and C–Cl bonds and contain nearly equivalent CH3 groups (Figure 3).

Let us go back to the X-ray structure of the C3H8Cl+ cation. It does show the superposition of the structures of the two isomers A and B (Figure 3). The structure of A can be determined rather reliably (Figure 3b): it contains the −\text{C}=\text{CH}_2 group with a shortened double bond (1.33 Å) on which the positive charge is mainly concentrated. Isomer B is more disordered due to the crystallographic overlap of its two orientations, as presented in Figure 3c. This situation causes a strong increase in the ellipsoid of the Cl atom and to its apparent bridge type.

In the B cation, the chlorine atom is attached directly to the middle \text{C}=\text{C} bond. This arrangement leads to a stronger extinguishing of the positive charge on this bond and a greater weakening of the C–Cl bond, as compared with those of A, where the Cl atom is more distant from the −\text{C}″=\text{CH}_2 moiety. Cation A also contains CH and CH$_2$ groups, which can participate in fast proton exchange on the IR time scale.

We attempted to obtain crystals of the salt of cations A and B with the {F$_{11}^-$} anion by recrystallizing the chloronium salt C$_3$H$_8$Cl$^+$({F$_{11}^-$}) from a DCM solution under exactly the same conditions where the crystals of cations A and B with the {Cl$_{11}^-$} anion were obtained. Next, the crystals were grown. Nevertheless, X-ray diffraction analyses showed that they are an ionic salt, Cat$^+$({F$_{11}^-$}), with a carbocation that does not contain a chlorine atom. Therefore, we discuss this compound below.

2.3. X-ray Analysis of Vinyl-Type Butylene Carbocations in the Crystal Phase. The crystals growing from a DCM solution of C$_3$H$_7^+$-{Cl$_{11}^-$} salt appeared rather quickly (1–2 days) with a good yield. The X-ray crystal structure uncovered an ionic compound with gross formula C$_3$H$_7^+${Cl$_{11}^-$} (Table S4) and a cationic moiety, which represents three crystallographically overlapping structures (Figure 4). This means that in one cell formed by the anions, the cation isomer can be in three different orientations.

Separation of differently oriented isomers indicated that one of them has a trans-configuration, and we will denote it as I (Figure 5). The other two structures have a cis-configuration. Because they are very similar in geometry, we will designate them as IIa and IIb (Figure 5). The middle CC bond in the trans- and cis-isomers is noticeably shorter than two terminal CC bonds (Table S6) and should be attributed to a double bond. One of the two middle C atoms with the highest positive charge on it (C2', C2, and C4, Figure 5) should have sp$^3$ hybridization with an angle close to 180° (the same is true for A and B cations). By contrast, the experimental angle turned out to be much smaller. This issue will be discussed later in the paper.

We already noted that our attempt to obtain the crystalline chloronium salt C$_3$H$_7^+$({F$_{11}^-$}) from its DCM solution failed. Crystals were grown from this solution for a long time with a low yield, which may indicate their formation from the products of side reactions. The X-ray diffraction data showed that this is an ionic compound with gross formula C$_3$H$_7^+$({F$_{11}^-$}) (Table S4) and a chlorine-free organic moiety. The independent part of the unit cell contains the cation C$_3$H$_7^+$ (which will be denoted as III) and the neutral 2-butenec molecule, both in cis-configuration (Figure 6, the H atoms and the anion are not shown). Positions of the cation and neutral molecule are diversified by the center of symmetry. This adds two more positions, giving rise to real disordering shown in Figure 7. The location of four organic moieties with two {F$_{11}^-$} anions in the unit cell is presented in Figure 8.

The geometry is very similar between cation III and 2-butenec (within the margin of error of determination, Figure 9). Due to the high R-factor because of disorder of the crystal structure it is not possible to localize the H atom on C2 or C2* atoms. Nonetheless, geometric calculation gives a better R-factor when the H atom is positioned on the C2* atom instead of the C2 atom. Therefore, we will assume that the C2* atom (with all C* atoms) belongs to 2-butenec and the second entity is cation III. The geometric parameters of isomer III and 2-butenec are given in Figure 9 and Table S7.
3. DISCUSSION

3.1. Effect of the Environment on Vinyl Carbocations.

Vinyl cations are affected by the basicity of the anions and the order of their localization in the immediate environment.

Surrounded by weakly basic \( \text{F}^{-} \) anions, the \( \text{C}_4\text{H}_7^{+} \) cation forms only cis-isomer III. The shortest eight \( \text{C} \cdots \text{F} \) distances between the C atoms of the isomer and the F atoms of the anions are similar and are in fact identical to those formed by neutral 2-butene (Figure 10). This means that the weak basicity of the anionic environment contributes to the increased uniformity of its interaction with the cations and to the uniformity of charge scattering over the cation.

The stronger interaction of the \( \text{C}_4\text{H}_7^{+} \) cation with the more basic \( \text{Cl}^{-} \) anions diminishes the homogeneity of this interaction: the CH\( \cdots \)CH moieties of isomers I and IIa,b are the same because the interactions of their C atoms with the Cl atoms of neighboring anions (C\( \cdots \)Cl distances) are identical. Nonetheless, the same interaction of their second part CH\( \cdots \)C\( ^{\text{\textcircled{C}}} \) with the environment is different (Figure 10). This state of affairs enhances the asymmetry of the cations, and the CH\( \cdots \)C\( ^{\text{\textcircled{C}}} \) angle can take two values, thereby leading to the formation of the trans-I isomer.

Isomers I–III all have one feature: the bond angle at the sp\(^1\) carbon atom with the highest positive charge is much smaller (130\(^{\circ}\)–109\(^{\circ}\)) than the 180\(^{\circ}\) angle at the sp\(^3\) C atoms in neutral hydrocarbons. It can be assumed that the reason is the enhanced interaction of the sp\(^1\) carbon atom with one or two fluorine or chlorine atoms of the nearest anion, but this is not the case. The \( \text{C}_{\text{sp}^1} \cdots \text{F} / \text{Cl} \) distances are similar to those of other carbon atoms in the isomers (Figure 10). That is, this property is an intramolecular feature of carbocations. The same feature (the angle at sp\(^1\) C atom less than 180\(^{\circ}\)) is also true for cations A and B.

X-ray crystallography determines only the carbon skeleton of cations. The positions of H atoms in structures I–III presented in Figures 5 and 10 are the result of a calculation based on the fact that, according to X-ray diffraction data, the C\( ^{=\text{C}} \) bond is located in the middle of the nearly symmetric molecule, and in the IR spectra in the CH stretch frequency range, only absorption of CH\(_3\) groups is observed. Nonetheless, the high similarity of geometrical parameters of cation III with those of neutral 2-butene (Figure 10) suggests that a H atom must be attached to the C2 atom in III. It also follows from the similarity of the geometry of cations II and III that their two middle C atoms with sp\(^1\) hybridization are nearly equivalent. That is, they must contain the C\( ^{\text{\textcircled{C}}} \cdots \text{H} \) group and their structure can be depicted as shown in Scheme 1.

In the a structure, the H atom looks like a bridge: it quickly migrates between two potential holes located near the C atoms, as in proton disolvates L\(^{\text{\textcircled{L}}} \cdots \text{H} \cdots \text{L} \), where L is the base molecule. The IR spectrum of II contains intense low-frequency absorption at \( \sim 1400 \text{ cm}^{-1} \) (Figure 11), which indicates the presence of fast proton dynamic effects in the cation (in the spectrum of III, this frequency range overlaps with intense absorption of the anion) and can indirectly confirm the presence of a bridging proton with a double-well
proton potential. The bridging proton promotes the formation of cis-isomers.

The geometry of the carbon skeleton of trans-isomer I can be explained satisfactorily by structure **I**. Nonetheless, because the details of its IR spectrum, overlapping with the spectra of isomers **IIa** and **IIb** cannot be seen, this structure can neither be confirmed nor refuted.

IR spectra of cations A and B also contain very strong and broad absorption at 1427 cm\(^{-1}\) (Figure 2), indicating a fast proton exchange at the IR time scale that is characteristic of a bridging proton. Therefore, for cation A, a similar structural scheme can be proposed as for cations II and III (Scheme 2). The positions of the C and Cl atoms in the B cation were determined with greater error; therefore, for this cation, we will not discuss the situation with a fast proton exchange and the possibility of the formation of a bridging proton.

It should be noted that a linear isomer CH\(_3\)–C\(\equiv\)CH\(_2\) with an angle at the central sp\(^1\) carbon atom of 180° does not yield broad intense absorption at 1400 cm\(^{-1}\) in the IR spectrum. Its presence in the IR spectra can probably serve as an indicator of a bridging proton in carbocations, as shown in Schemes 1 and 2.
The fact that, in crystalline salts, there are no shortened $\cdot \cdot \cdot \cdot$ F/Cl distances between cations and anions and all of the distances are similar to or exceed the corresponding van der Waals distances ($r_C + r_{Cl} = 3.53 \text{ Å}$) \(^{24}\) indicates that the cations are in uniform anionic surroundings (UAS) with strong dispersion of a positive charge over them. This observation supports their symmetric structure with a bridged proton (Scheme 1). They will be hereafter designated as UAS cations.

For all studied salts of carbocations, two sets of $\nu C\equiv C$ stretch frequencies can be distinguished, which belong to two types of isomers of cations $C_4H_7^+$ and $C_3H_5^+$. The first set is UAS cations of the crystal salts with a broad band of the C$\equiv C$ stretch at $\sim 1490 \text{ cm}^{-1}$ (Figure 12). Their frequencies are almost independent of the basicity of the anionic environment, $\{Cl_{11}^-\}$ or $\{F_{11}^-\}$ (Figure 11, Table 1); being very weak, basicity differs significantly between these anions.\(^{21}\) Moreover, the C$\equiv C$ stretches of the middle group of isomers I–III are very similar to those of the terminal $H-C\equiv C$ groups of UAS vinyl cations $^{15,16}$ $H-C\equiv CH_2$ and $H-C\equiv C(CH_3)_2$ (Table 1). That is, the middle and terminal C$\equiv C$ bonds of the UAS cations hardly differ in electron density.

In the CH stretch frequency region, IR spectra of I/II and III show only the absorption of the CH$_3$ groups, and that of the CH$_3$ group should be at $\sim 1400 \text{ cm}^{-1}$ if its H atom is bridged (Figure 11). The CH$_3$ frequencies are very close to those of neutral alkenes,\(^{22}\) suggesting that the positive charge

Figure 11. ATR IR spectra (with ATR corrections) of cations I/II (green) and III (red) in their crystal salts with anions $\{Cl_{11}^-\}$ and $\{F_{11}^-\}$, respectively, and the IR spectrum (in transmission mode) of amorphous salt $C_4H_7^+\{Cl_{11}^-\}$ (black). To illustrate the spectrum of the $\{F_{11}^-\}$ anion, an IR spectrum of the Cs$\{F_{11}\}$ salt is given (gray). Two intense bands of the adsorbed CD$_2$Cl$_2$ molecules that were not removed by evacuation after we washed the amorphous salt with $d_2$-DCM are marked with an asterisk. Intense bands with unspecified frequencies belong to the anion. Broad intense absorption in the region $\sim 1400 \text{ cm}^{-1}$ is shaded in green.
amorphous (black) phases. The latter was obtained by evaporation of
cations in the crystal phase (dark green) and in IPs in the amorphous phase
in both IPs and UAS conditions in an amorphous salt. (e) CH
within the crystal salt.

The spectra of this group differ from those of UAS cations not only by the increased frequency of the C=C stretch (by 70–
100 cm$^{-1}$) but also by the presence of new bands at 3112 and
3056 cm$^{-1}$, which are highly characteristic for the CH$_2$ = group
(Figure 11, Table 2).

This means that the other isomer of the C$_3$H$_7^+$ cation with the
terminal CH$_3$ = (C) group arises in the amorphous phase; we
notated it as IV

$$\text{CH}_2 = \text{C}^+ - \text{CH}_2 - \text{CH}_3$$

(Formation of the methyl allyl cation CH$_3$ − CH$\equiv$ CH$\equiv$ CH$_2$ is
not possible because its $\nu_{\text{CH}}$ frequency must be close to
that of the allyl cation, 1303 cm$^{-1}$, with CH stretch frequencies
below 3000 cm$^{-1}$.$^{16}$)

We have already encountered such phenomena when investigating
isobutylene cation HC$\equiv$C(CH$_3$)$_2$ and propylene cation C$_3$H$_7^+$ in amorphous salts with the {Cl$_{11}^-$} anion.$^{15,16}$ The reason for the
difference between their spectra and the spectra of crystalline salts is that in the amorphous
phase, the anionic environment of the cation is disordered.
This situation favors the interaction of the cation with one of
the anions, thereby producing an ion pair (IP) in which a
cation charge is shifted to the site oriented toward the anion.

C=C stretch frequencies of isomer IV (and frequencies of
its =CH$_3$ group) match those of the CH$_3$=C$^+$−CH$_4$ cation in
the crystalline salt with {Cl$_{11}^-$} (Figure 12d,e), which was
studied by X-ray crystallography.$^{16}$ In other words, cation IV in
the amorphous phase, and cation CH$_3$=C$^+$−CH$_3$ in the
crystalline phase, form identical IPs with the {Cl$_{11}^-$} anion.
Therefore, if we analyze the structure of the IP of a propylene
cation in a crystal phase in more detail than we did earlier in
ref 16, it will be possible to acquire information about the
structure of the IP formed by IV in the amorphous phase.

The C$_3$H$_7^+$[Cl$_{11}^-$] crystal contains two types of linear
isomers of CH$_2$=C$^+$−CH$_4$, which differ in the character of
interaction with the anions of the environment. In one isomer,
all C···Cl distances exceed 3.642 Å (i.e., the sum of van der
Waals radii r$_C$ and r$_{Cl}$: 3.53 Å), and this is the UAS cation.
The second cation has a shortened C···Cl distance, 3.189 Å, which
indicates the formation of IPs by this cation. Let us consider
the second case.

The X-ray structure of the ion-paired cation looks symmetric
due to the crystallographic overlap of oppositely oriented
cations, as displayed in Figure 13. This arrangement leads to
equalization of a shortened C1/C3···Cl(anion) distance and bond lengths between atoms C1, C2, and C3.

in the IP, the emergence of a C1/C3···Cl contact between the
≡CH$_3$ group of the cation and the Cl atom of the anion
inevitably lengthens the opposite C(H$_3$)···Cl distances, which
should exceed 3.642 Å. Then, for the average value of the C1/
C3···Cl distances to match the experimentally determined
3.189 Å, the length of the short contact should be less than
2.73 Å. The latter value can be used as an estimate of the upper
limit of the contact length in the IPs. Then, the structure of similar IPs formed by the butylene cation in an amorphous salt
can be schematically presented as in Figure 14.

The similarity of the IR spectra between cations IV and
CH$_2$=C$^+$−CH$_3$ is manifested not only in the matching of their
C=C stretch frequency but also in the identity of their spectra
in the region of CH stretch vibrations (Figure 11): all $\nu_{\text{CH}}$ of
CH$_2$ and CH$_3$ groups are lowered by ~100 cm$^{-1}$ as compared
with those of the neutral alkenes or UAS cations. For the ion-

Figure 12. IR spectra in the C=C stretch frequency range: (a) UAS
cations I/II with the {Cl$_{11}^-$} anion in the crystalline (brown) and
amorphous (black) phases. The latter was obtained by evaporation of
the solution from which the crystals were grown. The dotted
spectrum is the difference between the black and brown spectra. (b)
UAS cation III with the {F$_{11}^-$} anion in the crystal phase. (c)
(CH$_3$)$_2$C=C−H cation with the {Cl$_{11}^-$} anion under UAS conditions
in the crystal phase (dark green) and in IPs in the amorphous phase
(light green).$^{15}$ (d) CH$_3$=C−CH$_2$CH$_3$ cation with the {Cl$_{11}^-$} anion
in both IPs and UAS conditions in an amorphous salt. (e) CH$_3$=C
= CH$_2$ cation with the {Cl$_{11}^-$} anion in both IP and UAS conditions
within the crystal salt.$^{16}$
Table 1. Some Indicative C–H and C=C Stretch Frequencies of the UAS Cations

| cation or neutral analogue | phase | CH stretches for \(==\text{CH}_2\) or \(==\text{CH}^+\) | C=C stretches | \(\Delta \nu \text{C}==\text{C}^+\) |
|---------------------------|-------|---------------------------------|--------------|------------------|
| \(\text{CH}_2==\text{C}==\text{CH}_2\) | solid | 3112 3056                         | 1589\text{a} 1576 | 1561 56          |
| \(\text{CH}_2==\text{CH}==\text{CH}_2\) | gas   | 3090 3008                         | 1645 66       | 1565 66–77       |
| \(\text{CH}_2==\text{C}==\text{Cl}\) | solid | 3104 2990                         | 1589 1576     | 1642 66–77       |
| \((\text{CH}_3)_2\text{C}==\text{C}^+\) | gas   | 3087 2980                         | 1556 104      | 1660 104         |
| \((\text{CH}_3)_2\text{C}==\text{CH}_2\) | solid | 3040                             |               |                  |
| \((\text{CH}_3)_2\text{C}==\text{CH}_2\) | gas   | 3088 2980                         |               |                  |

\text{aThe} most intense bands are underlined. \(\text{bNot determined.}\) The difference between C=C stretching frequencies of neutral alkenes and their corresponding cationic carbocations.

Table 2. CH and C=C Stretch Frequencies of an Ion-paired Cation with the \(\{\text{Cl}_{11}^-\}\) Anion in Comparison with Those of Their Neutral Analogues

| cation or neutral analogue | phase | CH stretches for \(==\text{CH}_2\) or \(==\text{CH}^+\) | C=C stretches | \(\Delta \nu \text{C}==\text{C}^+\) |
|---------------------------|-------|---------------------------------|--------------|------------------|
| \(\text{CH}_2==\text{C}==\text{CH}_2\) | solid | 3112 3056                         | 1589\text{a} 1576 | 1561 56          |
| \(\text{CH}_2==\text{CH}==\text{CH}_2\) | gas   | 3090 3008                         | 1645 66       | 1565 66–77       |
| \(\text{CH}_2==\text{C}==\text{Cl}\) | solid | 3104 2990                         | 1589 1576     | 1642 66–77       |
| \((\text{CH}_3)_2\text{C}==\text{C}^+\) | gas   | 3087 2980                         | 1556 104      | 1660 104         |
| \((\text{CH}_3)_2\text{C}==\text{CH}_2\) | solid | 3040                             |               |                  |
| \((\text{CH}_3)_2\text{C}==\text{CH}_2\) | gas   | 3088 2980                         |               |                  |

\text{aThe} most intense bands are underlined. \(\text{bNot determined.}\) The difference between C=C stretching frequencies of neutral alkenes and their corresponding cationic carbocations.

Figure 13. Schematic representation of a linear \(\text{CH}_2==\text{C}==\text{CH}_2\) cation ion-paired with the \(\{\text{Cl}_{11}^-\}\) anion in the crystal salt. The carbon skeleton of the cation was determined by X-ray crystallography and is shown by gray balls. Crystallographic superposition of two oppositely oriented structures of \(\text{C}_9\text{H}_7^+\) leads to the alignment of the C=C bonds and C==Cl distances.

\[\text{(Anion)}\text{Cl}^\cdots\text{C}=\text{C}^\cdots\text{Cl(Anion)}\]

\[\text{long (Anion)}\text{Cl}^\cdots\text{C}=\text{C}^\cdots\text{Cl(Anion)}\]

\[\text{short (Anion)}\text{Cl}^\cdots\text{C}=\text{C}^\cdots\text{Cl(Anion)}\]

\[\text{average 3.168 A}\]

\(\text{long C=Cl distance 3.068 A}\)

\(\text{short C=Cl distance 3.068 A}\)

The components of crystallographic overlapping

Experimental image

Figure 14. Schematic representation of the structure of the IPs arising in the amorphous salt \(\text{C}_9\text{H}_7^+\{\text{Cl}_{11}^-\}\).

\[\text{(Anion)}\text{Cl}^\cdots\cdots\text{CH}_2==\text{C}==\text{CH}_2==\text{CH}_3\]

IV

paired \(\text{CH}_2==\text{C}==\text{CH}_2\) cation, this feature has been explained by the involvement of the \(\text{CH}_3\) group in weak hyperconjugation with the 2p\(_x\) orbital of the charged central carbon atom.\(^{26}\) In the case of cation IV, such an explanation is hardly acceptable. For vinyl-type carbocations, the nature of the hyperconjugation effect probably requires clarification.

The absorption bands of stretching C=C vibrations of cations IV in the IP consist of three components (Figure 12d, Table 2). A possible reason is that anions \(\{\text{F}_{11}^-\}\) and \(\{\text{Cl}_{11}^-\}\) have three sites of halogen atoms with different basicity: \(a < b < c\) (Figure 15). This has been experimentally proven by means of protonated carbon monoxide, \(\text{H}==\text{CO}\), as a test cation: it is attached to three sites—\(a\), \(b\), and \(c\)—forming IPs, thereby leading to the splitting of its \(\nu\text{CH}\) and \(\nu\text{CO}\) frequencies into three components.\(^{26}\)

The same can happen when vinyl cations form IPs. The X-ray structure shows that the \(\text{CH}_3==\text{C}==\text{CH}_2\) cation is attached to the most basic site \(c\) of the anion. Nonetheless, different rotational orientations of the anion are possible, allowing the cation, albeit less likely, to come into contact with sites \(a\) and \(b\), which will cause the observed splitting of the \(\nu\text{C}==\text{C}\) band into three components.

The influence of the inhomogeneous basicity of the Cl atoms of the \(\{\text{Cl}_{11}^-\}\) anions on the absorption bands of the C=C stretch in the UAS cations is weaker and can broaden them. Indeed, the IR spectra of UAS cations show one broad intense band \(\nu\text{C}==\text{C}\) at \(\sim 1490\text{ cm}^{-1}\). Nonetheless, weak bands are also observed at \(1513\) and \(1537\) cm\(^{-1}\) (Figure 12). Their nature was revealed by the following experiment. A drop of the amorphous salt \(\text{C}_9\text{H}_7^+\{\text{Cl}_{11}^-\}\) in DCM, from which crystals of the salt of isomers I/II were grown, was placed on the surface of the crystal of ATR accessory and evaporated. The IR spectrum of the resulting amorphous film shows two C=C stretch bands: one is the same as that of UAS isomers I/II at \(1490\text{ cm}^{-1}\) and the second at \(1537\text{ cm}^{-1}\) belongs to another cation. By subtraction of the spectrum of isomers I/II from this spectrum, the band at \(1490\text{ cm}^{-1}\) can be well compensated and the band at \(1537\text{ cm}^{-1}\) can be isolated (Figure 12a) along with the full spectrum across the entire frequency range. It hardly differs from the spectrum of I/II, except for the band of the C=C stretch and moreover does not contain the characteristic bands of the \(==\text{CH}_2\) group. Consequently, the amorphous film comprises only isomers of UAS-type cations: the most symmetrical I/II, and a second cation, more distorted in a less homogeneous anionic environment; it approaches the formation of the IP but does not reach it. Therefore, the presence of a weak band at \(1537\) cm\(^{-1}\)
cm$^{-1}$ in the spectra of crystals can be explained either by the fact that the surface of the crystals is contaminated with a dried initial solution, or by the finding that the crystal contains inclusions of UAS isomers in a more inhomogeneous anionic environment.

The spectra of amorphous and crystalline salts of vinyl cations containing an IP also show additional weak bands of C–C stretch vibrations of UAS cations (Figure 12). That is, in solid salts with IP cations, some portion of UAS cations is always in equilibrium with them.

Crystalline salts of C\(_4\)H\(_7\)\(^+\) cations contain only UAS cations, possibly with a small proportion of slightly distorted ones. Amorphous salts of cations C\(_4\)H\(_5\)^+ and C\(_6\)H\(_4\)^+ contain mainly IP cations with some proportion of UAS cations as well.

The frequencies of C–C stretches of the IP cations exceed those of the UAS cations by \(\sim 100\ \text{cm}^{-1}\), which means that the \(\pi\)-bonding is strengthened when some of the positive charge is transferred to the anion upon contact. In this regard, it is interesting to trace how the addition of an electron-donating substituent to the vinyl group affects the C=C bond.

**3.2. Effect of Nucleophilic Substituents on the Vinyl Cations.** The crystalline salt C\(_4\)H\(_7\)Cl\(^+\)(Cl\(_{11}^\ominus\)) contains two isomers A and B of the chlorobutylene cation. On the other hand, when the crystal lattice is destroyed, isomer B turns into A. The question arises whether isomer B is stabilized by the crystal lattice and then is a UAS isomer, and in the amorphous phase, whether isomer A is stabilized by ion pairing, and then is an IP isomer? Serious arguments in favor of such an assumption are the finding that the C\(^+\)=C bond in B is the middle one, which only UAS cations have, and during ion pairing, it relocates to the terminal site of the cation, converting the cation to isomer A (Scheme 3).

![Scheme 3. Transition of Cation B to A upon Destruction of the Crystal Lattice and Formation of IPs](https://example.com/scheme3)

The replacement of the H atom in UAS cations by a chlorine nucleophile raises the C=C stretching frequency by \(\sim 220\ \text{cm}^{-1}\), which almost twofold exceeds that when UAS cations are transferred to IP cations (Scheme 4). The same substitution in IP cations leads to a smaller increase in \(\nu\)C=C (by 90–115 cm$^{-1}$) because the Cl atom is at a greater distance from the C=C bond. For both isomers A and B, the \(\nu\)C=C exceeds that of the neutral alkene analogues by 35 and 65 cm$^{-1}$, respectively.

It is interesting how much the chlorine atom extinguishes the positive charge on the C=C bond? C–Cl stretch frequency of cation A (770 cm$^{-1}$) matches that of the CH\(_2\)Cl$^-$ groups of the di(chloromethyl)chloronium cation: (CH\(_2\)Cl)$_2$Cl$^-$ (768 cm$^{-1}$).\(^\text{20}\) Therefore, the positive charge on the Cl atom of A is similar to that on the terminal Cl atoms of the (CH\(_2\)Cl)$_2$Cl$^+$ cation, meaning that a smaller part of the charge is distributed to the Cl atom, and the charge remains mostly on the C=C bond.

The attachment of a nucleophile to a charged C\(^+=\)C bond results in a decrease of its charge and in a high-frequency shift of the C=C stretch to such an extent that it can even exceed the \(\nu\)C=C of neutral alkenes. This observation is in agreement with other reports on vinyl-type carbocations R’\(^+\)=CR$^2_1$ stabilized by electron-donating groups R’ and R’$^\ominus$.

![Scheme 4. Comparison of the UAS and IP Cations with Their Chlorovinyl Analogues B and A, Respectively](https://example.com/scheme4)

\(R' = \text{alkyl or aryl groups and two } \beta\text{-silyl substituents, which are not nucleophiles in neutral molecules, the positive charge on the } C=C \text{ bond is greatly reduced due to the combined influence of three substituents directly attached to the positively charged } C=C \text{ bond.}\)\(^\text{7,10,11,12}\) From NMR data, it follows that considerable delocalization of the charge onto the R’ substituents and the silyl groups occurs. According to the X-ray diffraction analysis, the C=C bond is shortened to 1.234 Å for \(R' = \text{cyclopropyl}^\ominus\) or 1.221 Å for \(R' = \text{Bu}^\ominus\), and the C=C stretching frequencies in the IR spectra increase significantly up to 1958 and 1987 cm$^{-1}$, respectively. These data are more consistent with a triple C=C bond than with a double C=C bond.

Not only do electron-donating substituents R’ significantly decrease the electron deficiency in the positively charged C=C group but also \(\beta\)-silyl groups promote quenching of the charge of the (C\(^+=\)C)$^\ominus$ bond owing to the involvement of C\(^\alpha\)–Si $\sigma$-bonds in hyperconjugation with the formally empty 2p orbital at the positively charged C\(^\alpha\) atom.\(^\text{7,10,11,12}\)

For small nonstabilized vinyl-type cations, C\(_4\)H\(_3\)^+ and C\(_6\)H\(_4\)^+, the positive charge is localized mainly on the C=C bond that naturally approaches one-and-a-half-bond status with a reduced C=C stretching frequency down to 1490 cm$^{-1}$. When vinyl-type cations are stabilized by substituents capable
of effectively dissipating a positive charge on themselves, their ability to donate electrons through σ-bonds, π-bonds, or the hyperconjugation effect is enhanced and nucleophilicity increases. This brings the C≡C linkage closer to the triple-bond status.

The peculiarity of the C≡C stretch frequency is in agreement with the results of density functional theory (DFT) calculations [at the B3LYP/6-311+G(d,p) level of theory] only for the stabilized vinyl cation.7,8,11,16 At the same time, DFT and MP2 calculations with the same basis set predicted for isobutylene (CH₃)₂C=CH⁺ and all studied isomers of the nonstabilized C₅H₅⁺ cation5,16 that the frequency of C≡C stretches exceeds that of neutral alkenes by ~200 cm⁻¹, while experimental ν=C≡C values are lower by ~160 cm⁻¹. This is a drastic mismatch, reaching 360 cm⁻¹. A similar mismatch between calculated (with different basis sets) and experimental frequencies for CH stretches of C−H bonds involved in hyperconjugation was reported for all alkane carbocations.18,23 It looks like the modern quantum-chemical calculations are consistent with experimental data for carbocations with effective charge dispersion on a large number of substituent atoms but are not applicable to nonstabilized carbocations with significant charge localization on the C≡C bond or a small number of C atoms. This issue remains to be resolved.

The present work yielded another interesting result: in a DCM solution, chloronium cation C₅H₅⁺Cl⁺ (with the {F₁₁⁺} or {Cl₁₁⁺} cation-counterions) containing eight H atoms transforms into vinyl cations containing six H atoms and a double C≡C bond. This happens in two ways: (i) with a release of a H₂ molecule and transition to chlorinated vinyl cation A or B (Scheme 5), which crystallizes from the solution with the {Cl₁₁⁺} anion or (ii) with the release of an HCl molecule, thus producing vinyl cation III (Scheme 5), which crystallizes from the solution with anion {F₁₁⁻}.

These reactions are possible if the solvation of the chloronium cation with DCM leads to the opening of its ring with the formation of an unstable chlorobutyl cation, which decomposes into more stable vinyl cations with the release of H₂ or HCl. At room temperature, spontaneous decay of a saturated carbocation to vinyl with the release of H₂ requires more detailed research, and we will devote a separate publication to this phenomenon.

In solid salts, the chloronium cation is stabler because it is “solvated” by anions {Cl₁₁⁻} or {F₁₁⁻}. To decompose it, one has to raise the temperature to 140–150 °C. Then, it degrades with the release of HCl and the formation of cation IV (Scheme 6).

### 4. CONCLUSIONS

Vinyl-type butylene carbocation C₄H₅⁺ can form trans- and cis-isomers CH₃−C⁺═CH−CH₃ (I–III) in crystalline salts and isomer CH₂−C⁺═CH−CH₃ (IV) in amorphous salts. They exhibit the following properties:

- In crystals, isomers I–III are uniformly surrounded by F/Cl atoms of neighboring anions with C···F/Cl distances between the cation and anion not less than the sum of van der Waals radii rC and rF. This contributes to the greatest localization of the positive charge on the C≡C bond and a decrease in νC≡C up to ~1490 cm⁻¹ (i.e., by 160 cm⁻¹ compared to that of neutral alkenes), which approaches that of aromatic CC bonds.
- In amorphous phase, the disorder of the anionic environment of the cation drives the formation of contact IPs with asymmetric cation isomer IV. It binds to the anion through the CH₂═ group with the contact distance C···Cl not exceeding 2.73 Å, thereby leading to a decrease in the positive charge on the C≡C bond and an increase in its C≡C stretch by ~100 cm⁻¹, as compared to UAS cations. Intermediate states are also possible.
- Stabilization of cations C₅H₅⁺ by the replacement of the H atom with a Cl atom gives rise to isomers A (CH₁−CHCl−C⁺═CH₂) and B (CH₁−C≡Cl−C⁺═CH₂), both of which are present in crystals. Upon amorphization, isomer B transitions to isomer A owing to ion pairing. Donation of the electron density from the Cl atom to the C≡C bond leads to its strengthening and rC≡C rises by 90–115 cm⁻¹ (A) and 220 cm⁻¹ (B) relative to their nonchlorinated analogues, respectively.
- Stabilization of vinyl cations with polyatomic substituents such as alkyl, aryl, and silylum (R₅Si) groups induces a further significant increase in the C≡C stretching vibration up to 1958 cm⁻¹7,8,11,12. This value exceeds that for neutral alkenes by ~308 cm⁻¹. The effective scattering of the positive charge on these polyatomic substituents significantly improves their electron-donating ability. As a consequence, the charge on the C≡C linkage greatly wanes, and the electron density on it goes up so much that it approaches a triple-bond state.

Summing up, we can conclude that in this series of “in UAS—ionically paired—stabilized by the Cl atom—stabilized by polyatomic substituents”, the frequency of C≡C stretch (and bond strength) monotonically increases from 1490 to 1958 cm⁻¹ (by 470 cm⁻¹).
A specific feature of the chloronium cation $\text{C}_4\text{H}_4\text{Cl}^+$ is its ability to spontaneously decompose into the vinyl chloride cation $\text{C}_4\text{H}_3\text{Cl}^+$ (with release of $\text{H}_2$) or the butylene cation $\text{C}_6\text{H}_7^+$ (with release of $\text{HCl}$) in a DCM-based medium at room temperature. This means that, under solvation conditions in solutions, the stability of chloronium is inferior to that of vinyl cations. In solid salts, the chloronium is stronger, stabilized by “solvation” with anions $\{\text{Cl}^-\}$ or $\{\text{F}^-\}$, and decomposes at elevated temperatures of 140–150 °C to form the carbocation $\text{CH}_2=\text{C}^+\cdot\text{CH}_2\cdot\text{CH}_3$.

The thermal stability of vinyl cations in their carbocation salts up to at least 140 °C allows their salts to be stored for a long time under anaerobic conditions and used as stable reagents in synthetic chemistry.

5. EXPERIMENTAL SECTION

The carbonyl acids $\text{H}\{\text{Cl}^-\}$ and $\text{H}\{\text{F}^-\}$ were prepared as described previously. They are purified by sublimation at 150–160 °C under a pressure of 10⁻³ Torr on cold Si windows in a specially designed IR cell reactor, whose detailed description is given in ref.15. The formed thin translucent layer yielded a clearcut IR spectrum. Dry vapors of 1,4-dichlorobutane were injected anaerobically into the IR cell and their interaction with the acid was controlled by sequential recording of IR spectra at short time intervals. All sample handling was carried out in an atmosphere of argon ($\text{H}_2\text{O}$ concentration < 0.5 ppm) in a glovebox. To obtain crystals, a weighable amount of the studied salts was prepared via a direct synthesis via the GRAMMS/A1 (7.00) software from Thermo Fisher Scientific. A specific feature of the chloronium cation $\text{C}_4\text{H}_4\text{Cl}^+$ is its ability to spontaneously decompose into the vinyl chloride cation $\text{C}_4\text{H}_3\text{Cl}^+$ (with release of $\text{H}_2$) or the butylene cation $\text{C}_6\text{H}_7^+$ (with release of $\text{HCl}$) in a DCM-based medium at room temperature. This means that, under solvation conditions in solutions, the stability of chloronium is inferior to that of vinyl cations. In solid salts, the chloronium is stronger, stabilized by “solvation” with anions $\{\text{Cl}^-\}$ or $\{\text{F}^-\}$, and decomposes at elevated temperatures of 140–150 °C to form the carbocation $\text{CH}_2=\text{C}^+\cdot\text{CH}_2\cdot\text{CH}_3$.

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