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N-Coordinated tellurium(II) and telluronium(IV) cations: synthesis, structure and hydrolysis†

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A set of N-coordinated tellurium(II) compounds containing either C,N-chelating ligands **CN^R** (where **CN** = 2-(RN=CH)C₆H₄, R = tBu or Dipp; Dipp = 2,6-iPr₂C₆H₃) or N,C,N pincer ligands **NCN^R** (where **NCN** = 2,6-(RN=CH)₂C₆H₄, R = tBu or Dipp) were synthesized. In the case of C,N-chelated compounds, the reaction of **CN^{Dipp}Li** with Te(dtc)₂ (where dtc = Et₂NCS₂) in a 1 : 1 molar ratio smoothly provided the carbamate **CN^{Dipp}Te(dtc)** which upon treatment with 2 eq. of HCl provided the chloride **CN^{Dipp}TeCl**. In contrast, the analogous conversion of **NCN^RLi** with Te(dtc)₂ surprisingly furnished ionic bromides **[NCN^RTe]Br** as a result of the exchange of dtc by Br coming from nBuBr present in the reaction mixture. Furthermore, the reaction of **CN^{Dipp}TeCl** or **[NCN^RTe]Br** with silver salts AgX (X = OTf or SbF₆) provided the expected tellurium cations **[CN^{Dipp}Te]SbF₆** and **[NCN^RTe]X**. To further increase the Lewis acidity of the central atom, the oxidation of selected compounds with 1 eq. of SO₂Cl₂ was examined yielding stable compounds **[CN^{tBu}TeCl₂]X** and **[NCN^{tBu}TeCl₂]X**. The oxidation of the Dipp substituted compounds proved to be more challenging and an excess of SO₂Cl₂ was necessary to obtain the oxidized products **[CN^{Dipp}TeCl₂]SbF₆** and **[NCN^{Dipp}TeCl₂]SbF₆**, which could solely be characterized in solution. Compounds **[CN^{tBu}TeCl₂]OTf** and **[NCN^{tBu}TeCl₂]OTf** were shown to undergo a controlled hydrolysis to the corresponding telluroxanes. All compounds were studied by multinuclear NMR spectroscopy in solution and for selected compounds solid state ¹²⁵Te NMR spectroscopy and single-crystal X-ray diffraction analysis were performed. The Lewis acidity of the studied cations was examined by the Gutmann–Beckett method using Et₃PO as the probing agent. The Te–N chalcogen bonding situation of selected compounds has also been examined computationally by a set of real-space bonding indicators.

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Introduction

Tellurium can be called the maverick of the chalcogen group,¹ being an element with very rich organometallic and coordination chemistry that provides various bonding possibilities as well as remarkable flexibility among its oxidation states.¹ Not surprisingly, organotellurium compounds have found widespread applications as precursors for materials, in organic syn-

thesis and catalysis.² Among these compounds, organotellurium cations certainly represent a prominent compound class that have recently gained considerable attention.³ In fact, two classes of these species are recognized based on the oxidation state of the tellurium, *i.e.* divalent organotellurenum(II) [R₂Te]⁺ and tetravalent organotelluronium(IV) [R₄Te]⁺ cations. The uncoordinated naked tellurium cations (Fig. 1A) are unknown due to very high reactivity caused mainly by the inherent electron deficiency of the tellurium atom; therefore, the presence of an external Lewis base (LB) is crucial for their stabilization.³ The utilization of inter-molecularly coordinated LBs has proved to be highly useful in this regard (Fig. 1B).⁴ Amongst the intramolecularly coordinating ligands, N,C,N-pincer type ligands have shown to be versatile in the stabilization of the Te center (Fig. 1C).⁵

In contrast, the organotelluronium(IV) cations provide more possibilities regarding the number of carbon bonded ligands, thus [R₃Te]⁺, [R₂TeX]⁺ and [RTeX₂]⁺ species can be distinguished.^{6–8} A structurally fully characterized dihaloorganotelluronium cation [RTeX₂]⁺ was unknown for a long time and only recently the first example has been synthesized by

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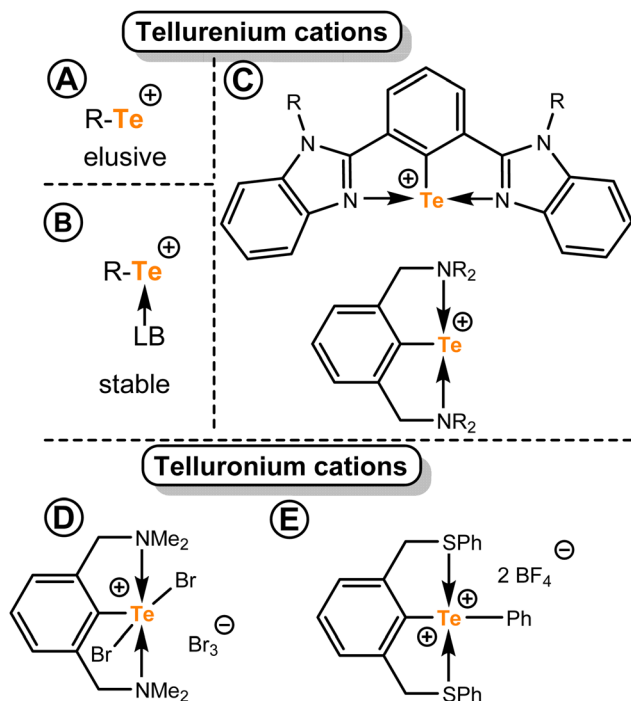


Fig. 1 Relevant examples of organotellurium cations for this study.

using an N,C,N-pincer ligand (Fig. 1D).⁹ The S,C,S-pincer ligands¹⁰ (Fig. 1E) and carbenes¹¹ even allowed the isolation of diorganotellurium(II) dications. Noteworthy, the tellurium cations of both types have gained increasing attention with regard to their utilization in catalysis or ion recognition.¹²

Organotelluroxanes, bearing at least one Te–C bond, constitute another interesting class of tellurium compounds that are usually synthesized by a hydrolysis of respective organotellurium halides.¹³ For example, a controlled base hydrolysis in the case of divalent R_2TeX results in the formation of oxygen bridged $(\text{R}_2\text{Te})_2(\mu\text{-O})$ telluroxanes.¹³ The chemistry of tetravalent telluroxanes is more diverse. The hydrolysis of R_2TeX_2 should lead to diorganotelluroxide R_2TeO , but these compounds mostly exist in their aggregated – dimeric or oligomeric forms $[\text{R}_2\text{TeO}]_n$.^{13,14} The monomeric species with a terminal Te–O bond could be isolated either using the support of a C,N-chelating ligand¹⁵ or by the protonation of the terminal oxygen by a strong Brønsted acid.^{7a,16} The oxidation of diorganotellurium compounds R_2Te also provided interesting dicationic oligotelluroxanes $[\text{R}_2\text{Te}^+ - [\text{R}_2\text{TeO}]_n - \text{O} - \text{Te}^+\text{R}_2][\text{X}]_2$ (where $\text{X} = \text{OTf}$, $n = 1-4$).¹⁷ The hydrolysis of monoorganotellurium(IV) compounds R_2TeX_3 is even more complicated, furnishing various products depending on the condensation and aggregation steps,¹⁸ but even monomeric organotellurinic acid $\text{R}_2\text{Te}(\text{O})(\text{OH})$ could be isolated.¹⁹ By analogy, a unique dimer of a organotelluronic acid $[\text{R}_2\text{Te}(\text{O})(\text{OH})_3]_2$ could be isolated first using sufficient steric shielding,²⁰ but later on related tellurium(IV) compounds could be obtained also with C,N-chelating ligands.^{19,21} Finally, diorganotellurones $\text{R}_2\text{Te}(\text{O})_2$ with two terminal Te–O bonds were isolated after oxidation of the starting R_2Te with NaIO_4 .²²

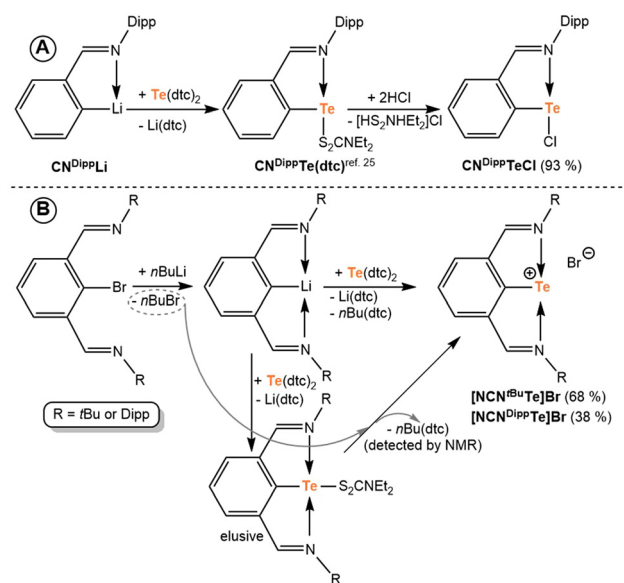
We have recently reported the synthesis of a set of tellurium cations stabilized by a C,N-chelating ligand CN^{tBu} with weakly coordinating anions.²³ We have also shown that in the case of the $[\text{CB}_{11}\text{H}_{12}]^-$ carborane counter-anion, a remarkable B–H bond activation by the tellurium centre is accessible.²⁴

The aim of this study is to further develop this family of promising N-coordinated organotellurium(II) cations $[\text{R}_2\text{Te}]^+$ and to examine the possibility of the isolation of organotelluronium(IV) cations $[\text{R}_2\text{TeCl}_2]^+$. For this purpose, either C,N-chelating ligands CN^{R} (where $\text{CN} = 2-(\text{RN}=\text{CH})\text{C}_6\text{H}_4$, $\text{R} = \text{tBu}$ or Dipp; Dipp = 2,6- $\text{iPr}_2\text{C}_6\text{H}_3$) or N,C,N-pincer ligands NCN^{R} (where $\text{NCN} = 2,6-(\text{RN}=\text{CH})_2\text{C}_6\text{H}_3$, $\text{R} = \text{tBu}$ or Dipp) were selected. These ligands enable us to follow the influence of the R group attached to the imino-function and the number of these donor moieties. The description of a controlled hydrolysis of selected dichloroorganotelluronium cations is also included.

Results and discussion

Synthesis

The recently presented synthetic protocol developed for $\text{CN}^{\text{tBu}}\text{TeCl}$,²⁵ including the conversion of $\text{CN}^{\text{R}}\text{Li}$ with $\text{Te}(\text{dte})_2$ (where $\text{dte} = \text{Et}_2\text{NCS}_2$) in a 1 : 1 molar ratio followed by the treatment with 2 eq. of HCl, was successfully applied for the synthesis of $\text{CN}^{\text{Dipp}}\text{TeCl}$ (Scheme 1A). In contrast, the treatment of $\text{NCN}^{\text{R}}\text{Li}$ with $\text{Te}(\text{dte})_2$ surprisingly led to the isolation of ionic bromides $[\text{NCN}^{\text{R}}\text{Te}]\text{Br}$. Importantly, both compounds precipitated directly from the reaction mixtures as the least soluble compound pointing to the fact that their ionic nature plays a crucial role (*vide infra*). The auto-ionization of both compounds is obviously caused by the presence of the second



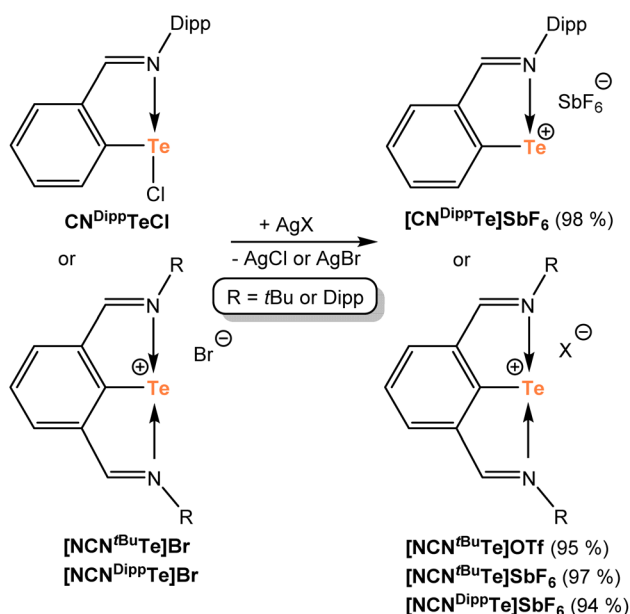
Scheme 1 Synthesis of starting compounds along with the proposed mechanism for the formation of $[\text{NCN}^{\text{R}}\text{Te}]\text{Br}$.



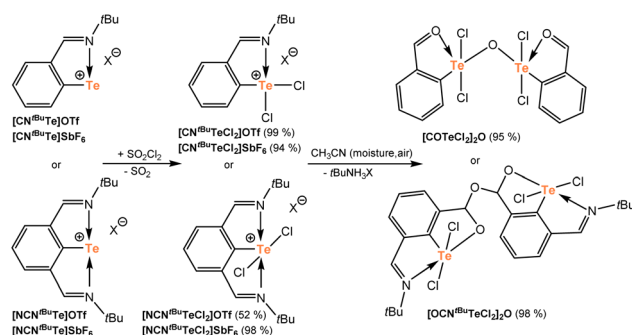
donor group in the pincer ligand. Although the dithiocarbamates $\text{NCN}^{\text{R}}\text{Te}(\text{dtc})$ are formed first, due to the presence of $n\text{BuBr}$ in the reaction mixture (as a result of *in situ* lithiation of the ligand) an exchange of dtc with Br occurs giving $[\text{NCN}^{\text{R}}\text{Te}]\text{Br}$ as isolable products. Low solubility helps in promoting this procedure (Scheme 1B), although the isolated yields are still rather moderate. This hypothesis has been clearly proven by the inspection of the mother liquors after the reactions, where the presence of $n\text{Bu}(\text{dtc})$ was established by NMR spectroscopy (Fig. S89–S91†) and the obtained data also agree with those published elsewhere.²⁶

The chloride or bromide in $\text{CN}^{\text{Dipp}}\text{TeCl}$ or $[\text{NCN}^{\text{R}}\text{Te}]\text{Br}$ can subsequently be substituted by less nucleophilic anions using silver salts AgX ($\text{X} = \text{OTf}$ or SbF_6 , Scheme 2) and the corresponding telluronium cations $[\text{CN}^{\text{Dipp}}\text{Te}]\text{X}$ and $[\text{NCN}^{\text{R}}\text{Te}]\text{X}$ can be isolated in quantitative yields as crystalline solids.

Finally, selected telluronium cations were oxidized by 1 eq. of SO_2Cl_2 producing rare examples of monoorganotelluronium (iv) cations $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{X}$ and $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{X}$ ($\text{X} = \text{OTf}$ or SbF_6) that were obtained as yellow solids in high yields (Scheme 3). In contrast, the oxidation of the Dipp substituted compounds $[\text{CN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$ and $[\text{NCN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$ was more challenging and an excess (5 to 10 eq.) of SO_2Cl_2 was necessary for complete oxidation. Furthermore, telluronium cations are highly sensitive towards even traces of moisture, which hampered the isolation of Dipp substituted compounds as pure solid samples. In the case of $[\text{CN}^{\text{Dipp}}\text{TeCl}_2]\text{SbF}_6$ and $[\text{NCN}^{\text{Dipp}}\text{TeCl}_2]\text{SbF}_6$, this hydrolysis always led to complicated mixtures of products. Interestingly, the *t*Bu species $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$ and $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$ can be hydrolyzed in a controlled manner, *e.g.* in wet acetonitrile, giving $[\text{COTeCl}_2]_2\text{O}$ and $[\text{OCN}^{\text{tBu}}\text{TeCl}_2]_2\text{O}$ in very high yields (Scheme 3), where $\text{CO} = 2-(\text{O}=\text{CH})\text{C}_6\text{H}_4$, $\text{OCN}^{\text{tBu}} = 2-(\text{O}-\text{CH})-6-(\text{tBuN}=\text{CH})\text{C}_6\text{H}_3$.



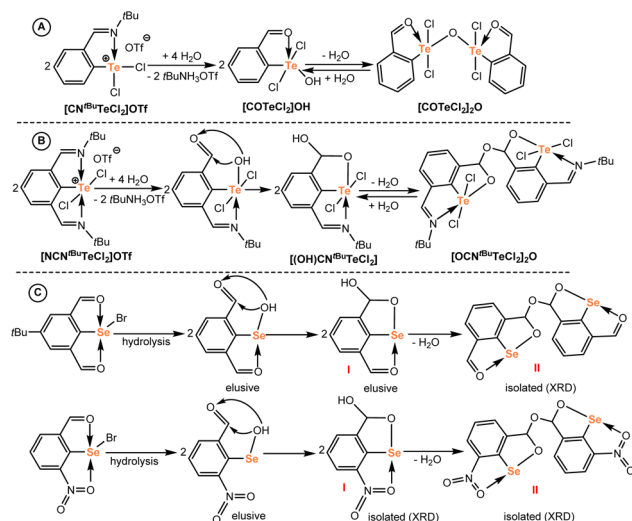
Scheme 2 Synthesis of tellurium(II) cations.



Scheme 3 Oxidation of tellurium(II) cations and subsequent hydrolysis.

Close inspection of the molecular structures of both hydrolyzed species revealed substantial structural differences (*vide infra* for sc-XRD analysis). In the case of $[\text{COTeCl}_2]_2\text{O}$, the oxygen bridges two tellurium atoms and its formation can be, thus, explained by a conventional hydrolysis of the pendant imino group to an aldehyde with a subsequent condensation of two tellurium hydroxides to give the final telluroxane $[\text{COTeCl}_2]_2\text{O}$ (Scheme 4A). Similar condensation reactions are well documented in the literature.^{18a,19}

In contrast, the oxygen atom in $[\text{OCN}^{\text{tBu}}\text{TeCl}_2]_2\text{O}$ bridges the tellurium fragments *via* two CH groups and no Te–O–Te linkage is formed. Considering the hydrolysis of the nitrogen function in the first step, the resulting Te–OH species does not undergo a classical condensation, but an attack of the Te–OH moiety to the $\text{CH}=\text{O}$ function seems to be feasible (Scheme 4B). Importantly, similar reactions have been recently well documented during an accidental hydrolysis of selenium (II) bromides (Scheme 4C).^{27,28} Attack of the Se–OH group leads to intermediates **I**, which dimerize to **II** with concomitant water elimination. In both the reported cases, the second *ortho*-position next to the selenium is occupied by a group con-



Scheme 4 Proposed mechanisms for the hydrolysis of telluronium(IV) cations (A and B) and comparison with the literature (C, ref. 27 and 28).

taining formally an oxygen donor atom and the corresponding molecular structures revealed non-negligible Se...O intramolecular interactions in all cases. Entirely the same situation is found in our case, when the intact imino-group may coordinate the central tellurium, thereby probably facilitating the Te-OH attack on the aldehyde function instead of a simple condensation toward the Te-O-Te dimer (Scheme 4B). Thus, the intermediate $[(\text{OH})\text{CN}^{\text{tBu}}\text{TeCl}_2]$ (where $(\text{OH})\text{CN}^{\text{tBu}} = 2\text{-(O-C(OH)H)-6-(tBuN=CH)C}_6\text{H}_3$), *vide infra* then condenses to the final product $[\text{OCN}^{\text{tBu}}\text{TeCl}_2]_2\text{O}$ as the least soluble compound similarly to the selenium analogues mentioned above.^{27,28}

Multinuclear NMR spectroscopy

All tellurium(II) compounds were characterized using high-resolution MALDI MS spectra (see the Experimental section and Fig. S99–S105†), but it was not possible for corresponding tellurium(IV) compounds due to their very high sensitivity to moisture; a similar problem was encountered with combustion analysis of the studied compounds. Nevertheless, all compounds were in detail investigated by ^1H , $^{13}\text{C}\{^1\text{H}\}$, $^{125}\text{Te}\{^1\text{H}\}$ and ^1H – ^{15}N HMBC NMR spectroscopy. The presence of the appropriate ligand was established mainly by diagnostic signals for the $\text{CH}=\text{N}$ group in the range of 8.83–10.31 ppm for $\delta(^1\text{H})$ and 159.4–171.1 ppm for $\delta(^{13}\text{C})$ (see Table 1). One signal was also obtained in 2D ^1H – ^{15}N HMBC experiments. Compounds substituted with *t*BuN groups showed signals with $\delta(^{15}\text{N}) = -86.7$ to -130.2 ppm, while for DippN substituted compounds values between -102.8 to -174.9 ppm were detected. Selected examples were also characterized by solid

state ^{125}Te NMR *vide infra*, where they are discussed together with solution ^{125}Te NMR data.

The hydrolysed products $[\text{COTeCl}_2]_2\text{O}$ and $[\text{OCN}^{\text{tBu}}\text{TeCl}_2]_2\text{O}$ exhibited only limited solubility in most organic deuterated solvents. $[\text{COTeCl}_2]_2\text{O}$ was reasonably soluble in thf-d8 that allowed us to obtain ^1H and $^{13}\text{C}\{^1\text{H}\}$ NMR spectra. Surprisingly, if non-dried thf-d8 is used, the spectra contained two sets of chemical shifts pointing to the presence of two compounds (*i.e.* $[\text{COTeCl}_2]_2\text{O}$ and $[\text{COTeCl}_2]\text{OH}$ *vide infra*) that both display shifts at 10.22 and 10.38 ppm (198.3 and 200.5 ppm) for the $\text{CH}=\text{O}$ group (Fig. S52 and S53†), respectively. Furthermore, the ^1H NMR spectrum also contained a singlet at 7.74 ppm that lacks any cross-peak in the 2D- ^1H – ^{13}C HSQC NMR spectrum corresponding to the OH group in $[\text{COTeCl}_2]\text{OH}$. Furthermore, the ^1H – ^1H EXSY NMR spectrum unambiguously proved a dynamic mutual exchange between both species and traces of present water (Fig. S55†). $^{125}\text{Te}\{^1\text{H}\}$ NMR spectra also showed two signals at 1342 ($[\text{COTeCl}_2]\text{OH}$) and 1405 ppm ($[\text{COTeCl}_2]_2\text{O}$). Based on these findings, we propose that upon dissolving of $[\text{COTeCl}_2]_2\text{O}$ in wet thf-D8 a partial hydrolysis of the Te–O–Te bridge occurred during the formation of $[\text{COTeCl}_2]\text{OH}$ (Scheme 4A). This assumption was verified by the addition of 10 μL of water to this sample that led to a complete disappearance of the signals attributable to the telluroxane $[\text{COTeCl}_2]_2\text{O}$ and only one set of signals for $[\text{COTeCl}_2]\text{OH}$ remained present (Fig. S56–S58†). The use of freshly dried thf-d8 almost suppressed the formation of $[\text{COTeCl}_2]\text{OH}$, but traces remain detectable in the NMR spectra most probably as a result of the moisture present in the original sample after hydrolytic synthesis of $[\text{COTeCl}_2]_2\text{O}$ (Fig. S59–S62†).

$[\text{OCN}^{\text{tBu}}\text{TeCl}_2]_2\text{O}$ exhibited good solubility in dmsO-d6 (anhydrous) only and the obtained ^1H and $^{13}\text{C}\{^1\text{H}\}$ NMR spectra agreed well with the proposed structure. The presence of the intact $\text{CH}=\text{NtBu}$ group is reflected by the observation of the signal at 9.14 ppm (157.3 ppm), while the chemical shift at 7.3 ppm (100.7 ppm) corresponds to the CH group involved in the bridge between both tellurium fragments. The $^{125}\text{Te}\{^1\text{H}\}$ NMR spectrum showed one signal at 1381 ppm (Fig. S64–S66†). The utilization of undried dmsO-d6 resulted in the appearance of the second set of signals that was tentatively assigned to $[(\text{OH})\text{CN}^{\text{tBu}}\text{TeCl}_2]$ pointing to a reversible hydrolysis of the present bridge (Scheme 4B); but in contrast to $[\text{COTeCl}_2]_2\text{O}$, the full conversion to $[(\text{OH})\text{CN}^{\text{tBu}}\text{TeCl}_2]$ was not obtained even after 8 days, with about 15% of intact $[\text{OCN}^{\text{tBu}}\text{TeCl}_2]_2\text{O}$ still present (see Fig. S67–72†).²⁹ The ^1H NMR spectrum of $[(\text{OH})\text{CN}^{\text{tBu}}\text{TeCl}_2]$ contained two singlets at 6.88 and 9.06 ppm corresponding to CHO and $\text{CH}=\text{N}$ groups ($\delta(^{13}\text{C}) = 99.9$ and 156.7 ppm) respectively, along with a broad one at 7.30 ppm for a new OH group (no cross-peak in 2D- ^1H , ^{13}C HSQC NMR). The $^{125}\text{Te}\{^1\text{H}\}$ NMR spectrum also revealed a new signal at 1374 ppm that is, however, only very slightly shifted in comparison to the starting $[\text{OCN}^{\text{tBu}}\text{TeCl}_2]_2\text{O}$ (*cf.* 1381 ppm), indicating a rather similar bonding situation around the tellurium atom (Fig. S68†). Note that the cleavage of the Te–O–Te bridge ongoing from $[\text{COTeCl}_2]_2\text{O}$ to $[\text{COTeCl}_2]$

Table 1 Relevant ^1H , ^{13}C , ^{15}N and ^{125}Te NMR data of studied compounds in CD_2Cl_2 , CDCl_3 or CD_3CN . Chemical shifts are given in [ppm]

Compound	$\text{CH}=\text{N}$ group $\delta(^1\text{H})$	$\delta(^{125}\text{Te})$ $\delta(^{13}\text{C})$	$\delta(^{15}\text{N})^a$	
Covalent CN-chelated tellurium(II) compounds				
$\text{CN}^{\text{tBu}}\text{TeCl}_2^b$	8.58	159.6	–93.4	1259
$\text{CN}^{\text{Dipp}}\text{TeCl}_2$	9.16	164.7	–121.4	1392
Ionic CN-chelated tellurium(II) compounds				
$[\text{CN}^{\text{tBu}}\text{Te}]\text{OTf}^c$	9.53	163.8	–123.0	1753
$[\text{CN}^{\text{tBu}}\text{Te}]\text{SbF}_6^c$	9.54	165.5	–130.2	1897
$[\text{CN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$	9.37	171.1	–174.9	2113
Ionic NCN-chelated tellurium(II) compounds				
$[\text{NCN}^{\text{tBu}}\text{Te}]\text{Br}$	10.27	160.7	–91.2	1394
$[\text{NCN}^{\text{tBu}}\text{Te}]\text{OTf}$	9.75	160.0	–88.8	1400
$[\text{NCN}^{\text{tBu}}\text{Te}]\text{SbF}_6$	9.57	159.4	–89.5	1409
$[\text{NCN}^{\text{Dipp}}\text{Te}]\text{Br}$	10.31	166.5	–119.8	1522
$[\text{NCN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$	9.60	165.9	–117.4	1557
CN-chelated tellurium(IV) compounds				
$\text{CN}^{\text{tBu}}\text{TeCl}_3^b$	8.86	163.4	^d	1177
$[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$	9.27	168.2	–86.8	1373
$[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{SbF}_6$	9.27	168.6	–87.1	1399
$[\text{CN}^{\text{Dipp}}\text{TeCl}_2]\text{SbF}_6$	8.83	170.3	–102.8	1428
NCN-chelated tellurium(IV) compounds				
$[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$	9.45	163.2	–86.7	1180
$[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{SbF}_6$	9.40	163.2	–87.0	1179
$[\text{NCN}^{\text{Dipp}}\text{TeCl}_2]\text{SbF}_6$	9.22	168.2	–112.9	1308

^a Determined by ^1H , ^{15}N HMBC experiments. ^b From ref. 23. ^c From ref. 25. ^d Not measured.



OH was accompanied by more pronounced downfield shift by 63 ppm. Unfortunately, all attempts to crystallize either $[\text{COTeCl}_2]\text{OH}$ or $[(\text{OH})\text{CN}^t\text{BuTeCl}_2]$ remained unsuccessful. Nevertheless, their existence in solution supported the proposed mechanism for the hydrolysis of telluronium cations shown in Scheme 4.

Solid state structures

Molecular structures determined by sc-X-Ray diffraction analysis are shown in Fig. 2–5 and the relevant geometrical parameters are compiled in Table 2.

The Te(1)–N(1) distance in $\text{CN}^{\text{Dipp}}\text{TeCl}$ of 2.249(4) Å (*cf.* $\Sigma_{\text{cov}}(\text{Te}–\text{N}) = 2.07$ Å,³⁰ Fig. 1) is slightly elongated in comparison to that in CN^tBuTeCl (2.203(2) Å),²⁵ but the former remains exclusively monomeric, while the latter forms a dimer *via* very weak Te–Cl...Te contacts (3.579 Å, *cf.* $\Sigma_{\text{cov}}(\text{Te}–\text{Cl}) = 2.35$ Å (*ref.* 30)) that can be probably ascribed to the steric effects of the Dipp group. The Te(1)–N(1) distance becomes significantly shorter (2.0645(16) Å) upon abstraction of the chlorine atom in $[\text{CN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$ (Fig. 3) and is comparable to its *t*Bu counter-

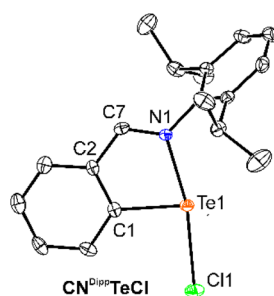


Fig. 2 ORTEP drawings of molecular structures of $\text{CN}^{\text{Dipp}}\text{TeCl}$. The thermal ellipsoids are given with 30% probability and hydrogen atoms are omitted for clarity.

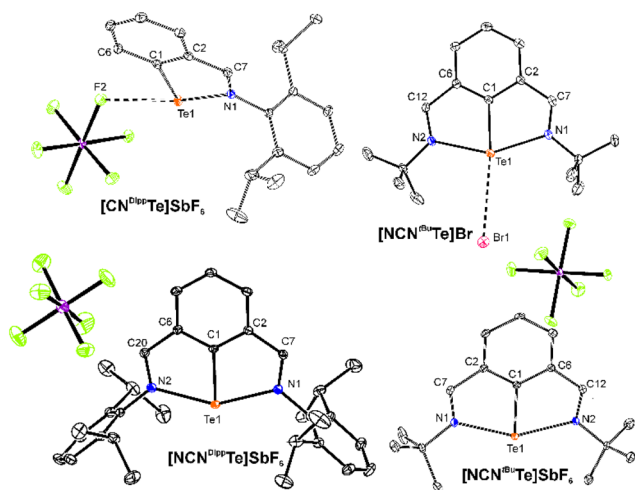


Fig. 3 ORTEP drawings of molecular structures of telluronium(iv) cations. The thermal ellipsoids are given with 30% probability and hydrogen atoms are omitted for clarity.

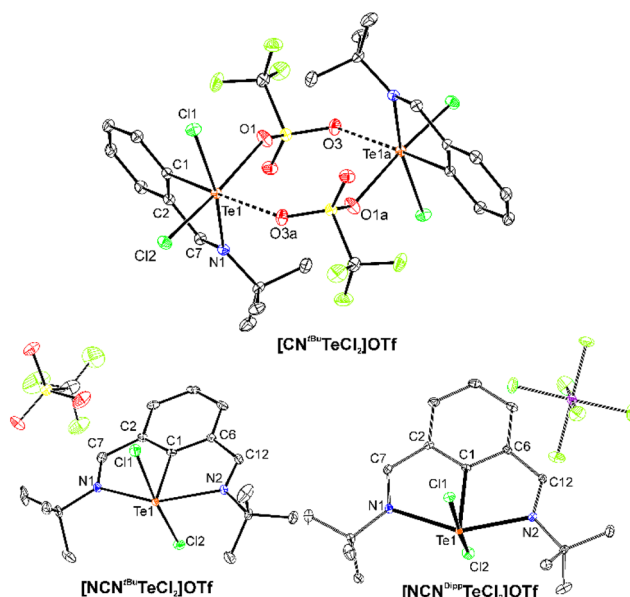


Fig. 4 ORTEP drawings of molecular structures of telluronium(iv) cations. The thermal ellipsoids are given with 30% probability and hydrogen atoms are omitted for clarity.

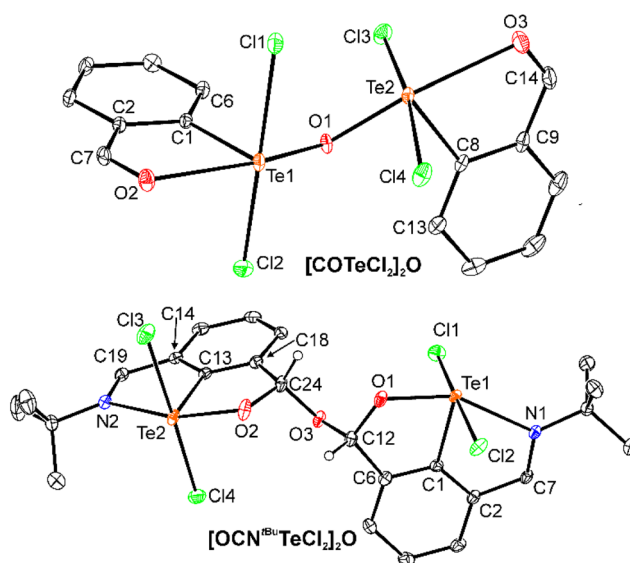


Fig. 5 ORTEP drawings of molecular structures of compounds $[\text{COTeCl}_2]_2\text{O}$ and $[\text{OCN}^t\text{BuTeCl}_2]_2\text{O}$. The thermal ellipsoids are given with 30% probability and hydrogen atoms except for CH group in the latter are omitted for clarity. Selected bond lengths [Å] for $[\text{COTeCl}_2]_2\text{O}$: Te(1)–C(1) 2.124(3), Te(1)–O(1) 1.9526(17), Te(1)–O(2) 2.5474(18), Te(1)–Cl(1) 2.5607(7), Te(1)–Cl(2) 2.4411(8), Te(2)–C(8) 2.117(2), Te(2)–O(1) 1.9432(18), Te(2)–O(3) 2.569(2), Te(2)–Cl(3) 2.4487(7), Te(2)–Cl(4) 2.5440(8). For $[\text{OCN}^t\text{BuTeCl}_2]_2\text{O}$: Te(1)–C(1) 2.070(3), Te(1)–N(1) 2.483(2), Te(1)–O(1) 1.997(2), Te(1)–Cl(1) 2.4861(14), Te(1)–Cl(2) 2.5375(14), Te(2)–C(13) 2.060(3), Te(2)–N(2) 2.456(2), Te(2)–O(2) 2.006(2), Te(2)–Cl(3) 2.4928(14), Te(2)–Cl(4) 2.5445(14), O(3)–C(12) 1.418(3), O(3)–C(24) 1.424(3), O(1)–C(12) 1.424(4), O(2)–C(24) 1.417(3).



Table 2 Relevant bond lengths in molecular structures of studied compounds given in [Å]

Compound	N1–Te1	N2–Te1	Other important contacts ^a
Covalent CN-chelated tellurium(II) compounds			
CN ^{tBu} TeCl ^b	2.203(2)	—	Te–Cl...Te 3.579
CN ^{Dipp} TeCl	2.249(4)	—	—
Ionic CN-chelated tellurium(II) compounds			
[CN ^{tBu} Te]OTf ^c	2.113(1)	—	Te...O(OTf) 2.500
[CN ^{tBu} Te]SbF ₆ ^c	2.073(2)	—	Te...F(SbF ₆) 2.687
[CN ^{Dipp} Te]SbF ₆	2.0645 (16)	—	Te...F(SbF ₆) 2.666
Ionic NCN-chelated tellurium(II) compounds			
[NCN ^{tBu} Te]Br	2.286(4)	2.303(4)	Te...Br 3.609
[NCN ^{tBu} Te]SbF ₆	2.259(2)	2.259(2)	—
[NCN ^{Dipp} Te]	2.259(2)	2.2596 (19)	—
CN-chelated tellurium(IV) compounds			
CN ^{tBu} TeCl ₃ ^b	2.286(1)	—	Te–Cl...Te 3.506(3)
[CN ^{tBu} TeCl ₂]OTf	2.308(3)	—	Te...O(intra-OTf) 2.523 Te...O (inter-OTf) 3.042
NCN-chelated tellurium(IV) compounds			
[NCN ^{tBu} TeCl ₂]OTf	2.281(4)	2.311(4)	Te–Cl...Te 3.699
[NCN ^{tBu} TeCl ₂]SbF ₆	2.300(5)	2.294(5)	Te–Cl...Te 3.620

^a Contacts include either intermolecular contacts of the type Te–X...Te or contacts between the cation and the anion. ^b From ref. 23. ^c From ref. 25. ^d Two closely related molecules present and data for only one are displayed.

part [CN^{tBu}Te]SbF₆ (2.073(2) Å).²³ The Te(1) atom is di-coordinated with a weak chalcogen Te...F bond with the counter-anion. As mentioned above, the utilization of the pincer ligands resulted in the autoionization even in the case of the bromide [NCN^{tBu}Te]Br, where the Br(1) atom again forms a weak chalcogen bond with Te(1) atom (3.609 Å, cf. $\Sigma_{\text{cov}}(\text{Te–Br}) = 2.50$ Å (ref. 30)). In the case of hexafluoroantimonates [NCN^{tBu}Te]SbF₆ and [NCN^{Dipp}Te]SbF₆, the tellurium cation remains without a similar contact. The Te(1) atom is three-coordinated with a T-shaped geometry and the Te–N distances span over a narrow interval of 2.259(2)–2.303(4) Å, which are elongated in comparison with C,N chelated cations (Table 2) but are well comparable to other N,C,N-pincer ligand stabilized tellurium cations (2.225–2.392 Å).^{5b–d}

The molecular structures of the tellurium(IV) cations differ significantly based on the used ligand. In the case of [CN^{tBu}TeCl₂]OTf, the tellurium atom adopts an octahedral array (Fig. 4), where C(1)/N(1) and Cl(1)/Cl(2) atoms are mutually *cis* coordinated. The Te(1)–N(1) distance 2.308(3) Å is similar to that in CN^{tBu}TeCl₃ 2.286(1) Å,²⁵ while the Te(1)–Cl(1/2) bonds 2.4528(8)/2.3350(9) Å, respectively, correspond to single bonds albeit the first one is a bit elongated in comparison to $\Sigma_{\text{cov}}(\text{Te–Cl}) = 2.35$ Å.³⁰ The triflate coordinates to the Te(1) atom with the bond length Te(1)–O(1) 2.523(2) Å that is significantly elongated in comparison to $\Sigma_{\text{cov}}(\text{Te–O}) = 2.35$ Å (ref. 30) and the octahedron is completed by the intermolecular contact with a second triflate (Te(1)–O(3a) 3.042(3) Å), thereby forming a dimer. In the case of [NCN^{tBu}TeCl₂]OTf and [NCN^{tBu}TeCl₂]SbF₆, the presence of the pincer ligand again supported the formation of ionic species and both OTf

and SbF₆ are situated outside the coordination sphere of the central atom. The Te(1) atom is captured by the pincer ligand with the Te–N distances ranging from 2.281(4) to 2.311(4) Å and these values are shorter than those established for the only structurally characterized analogue, *i.e.* [2,6-(Me₂NCH₂)₂C₆H₄TeBr₂][Br₃] 2.372(2) and 2.435(2) Å.⁹ The chlorine atoms are coordinated mutually in *trans* positions, which is in contrast to [CN^{tBu}TeCl₂]OTf and the Te–Cl bond lengths are in the interval of 2.4532(16)–2.5147(17) Å, again a bit elongated in comparison with $\Sigma_{\text{cov}}(\text{Te–Cl}) = 2.35$ Å.³⁰ The shape of the coordination polyhedron can be described as a distorted square-pyramid with the pincer *ipso*-carbon atom in the apical position. The closest intermolecular, but negligible, contact with the chlorine atom (3.699 and 3.620 Å, respectively) from the adjacent molecule is not considered.

The structures of the hydrolysis products [COTeCl₂]₂O and [OCN^{tBu}TeCl₂]₂O are shown in Fig. 5. The molecular structure of the former proved the hydrolysis of the imino-*t*BuN=CH function to the benzaldehyde framework while the condensation of the Te–OH groups produced a central –Cl₂Te(μ-OTe)Cl₂– linkage. Similar hydrolytic products were obtained upon basic hydrolysis of neutral tellurium(IV) complexes ArTeCl₃ (Ar = 2-pyC₆H₄¹⁹ or 8-Me₂NC₁₀H₆^{18a}). The coordination polyhedron of the central atoms is best described as a distorted square-pyramidal. The Te1/2–O2/3 distances 2.5474(18)/2.569(2) Å, respectively, are elongated in comparison with $\Sigma_{\text{cov}}(\text{Te–O}) = 1.99$ Å,³⁰ but still indicate intramolecular Te...O chalcogen interactions. Both Te(1) and Te(2) atoms are bridged by the O(1) atom and the bond lengths of 1.9526(18) and 1.9432(18) Å correspond well to the single-bond³⁰ and are also comparable to [ArTe(Cl₂)₂](μ-O) (Ar = 2-pyC₆H₄¹⁹ 1.969(3) and 1.963(3) Å or 8-Me₂NC₁₀H₆^{18a} 1.97(1) and 1.96(1) Å). The Te(1)–O(1)–Te(2) bonding angle of 123.96(13)° also approaches reported values for the abovementioned analogues 124.9(2)° and 125.4(5)°,^{18a} respectively. Each of the two tellurium atoms is further coordinated by two chlorine atoms and in each case one of the Te–Cl bonds is a bit longer than the other (*cf.* 2.4411(8) vs. 2.5607(7) for Te(1) and 2.4487(8) vs. 2.5440(8) Å for Te(2)).

The structure of [OCN^{tBu}TeCl₂]₂O differs significantly from that of its counterpart [COTeCl₂]₂O as mentioned above. Two tellurium fragments are connected *via* the C(12)–O(3)–C(24) bridge and both bond lengths correspond to single bonds 1.418(3) and 1.424(3) Å (ref. 30) (bonding angle 113.5(2)°), while the geometry around carbon atoms C12 and C24 is tetrahedral corresponding to the sp³ hybridization. Both tellurium atoms Te(1) and Te(2) form single bonds with O(1) and O(2) atoms (1.997(2) and 2.006(2) Å, respectively) and each is further coordinated by one nitrogen atom N(1) or N(2) (2.483(2) and 2.456(2) Å). The distorted square pyramidal array around the central atoms is completed by two chlorine atoms (the range of bond lengths 2.4861(14)–2.5445(14) Å) and *ipso*-carbon atoms of the pincer ligands in the axial position. The geometrical framework in [OCN^{tBu}TeCl₂]₂O is only rarely found for heavier p-block elements. With the exception of two selenium compounds^{27,28} depicted in Scheme 4, only pure organic compounds were reported.³¹



¹²⁵Te NMR spectroscopy and the Gutmann–Beckett method

All ¹²⁵Te NMR chemical shifts of the studied compounds along with those of relevant earlier published compounds are compiled in Table 1. It is to be noted that the values of the ¹²⁵Te NMR chemical shifts of compounds containing the CN^{Dipp} ligand are always downfield shifted in comparison with that of CN^{tBu} compounds, which is evidently a consequence of the electron withdrawing aromatic Dipp group in the former. For selected tellurium(II) compounds, solid state ¹²⁵Te NMR spectroscopy has also been examined to gain insight into the relationship between solution and solid state structure.

Regarding the C,N-chelated compounds, it becomes obvious that the abstraction of the chloride ion from the neutral precursors CN^{tBu}TeCl (1259 ppm) and CN^{Dipp}TeCl (1392 ppm) by weakly coordinating anions resulted in a significant downfield shift of the signals in [CN^{tBu}Te]X (X = OTf 1753 and SbF₆ 1897 ppm) and [CN^{Dipp}Te]SbF₆ (2113 ppm). The latter represents the highest obtained value among reported compounds pointing to its high Lewis acidity (*vide infra* further discussion). In the case of pincer complexes, the ¹²⁵Te NMR chemical shifts are found in a narrow interval *i.e.* 1394–1409 ppm for [NCN^{tBu}Te]X (X = Br, OTf and SbF₆) and 1522–1557 ppm for [NCN^{Dipp}Te]X (X = Br or SbF₆) pointing to the fact that all compounds form practically identical ionic pairs in solution even in the case of bromides. These values also suggest a significant shielding of the central atom in comparison with the C,N-chelated analogues due to the presence of the second donor functionality.

In the solid state, the $\delta_{\text{iso}}(^{125}\text{Te})$ NMR chemical shifts for the starting CN^{Dipp}TeCl (1403 and 1349 ppm, two independent molecules in the unit cell) are close to the value found in solution (1392 ppm). In the case of CN^{tBu}TeCl, $\delta_{\text{iso}}(^{125}\text{Te}) = 1340$ ppm is shifted in comparison with solution (1259 ppm) most probably indicating the absence of the weak intermolecular Te...Cl contact in solution.²⁵ In contrast, $\delta_{\text{iso}}(^{125}\text{Te})$ values of tellurium cations [CN^{tBu}Te]X (X = OTf 1742 and SbF₆ 1895 and 1842 ppm, two independent molecules in the unit cell) and [NCN^{tBu}Te]SbF₆ (1388 ppm) are all close to the values found in solution proving an analogous structure in both phases.

Looking to solution $\delta(^{125}\text{Te})$ of tellurium(IV) compounds, the formal abstraction of one of the chlorides in neutral CN^{tBu}TeCl₃ (1177 ppm)²⁵ resulted in a downfield shift, but it is significantly less pronounced in comparison to the abovementioned tellurium cations, *cf.* [CN^{tBu}TeCl₂]X (X = OTf 1373 and SbF₆ 1399 ppm) and [CN^{Dipp}TeCl₂]SbF₆ (1428 ppm). Again, the presence of the second ligand arm in pincer compounds helps to shield the tellurium atom more efficiently based on the obtained $\delta(^{125}\text{Te})$, *cf.* [NCN^{tBu}TeCl₂]X (X = OTf 1180 and SbF₆ 1179 ppm) and [NCN^{Dipp}TeCl₂]SbF₆ (1308 ppm).

The Gutmann–Beckett method,³² even with its shortcomings regarding steric effects and (or) “Pearson hardness”,³³ is still used as a gauge of Lewis acidity that enables a reasonable and straightforward scaling within one class of compounds,

being often applied to p-block elements as well.³⁴ We used this method to shed more light on the Lewis acidity of the studied compounds using Et₃PO as the probing agent (Table 3). As we are aware that ³¹P NMR chemical shifts of both Et₃PO and its complexes with Lewis acids are solvent dependent, we refrain from calculating the exact values of Gutmann–Beckett acceptor numbers and rather compare $\Delta\delta(^{31}\text{P})$, defined as $\Delta\delta(^{31}\text{P}) = \delta(^{31}\text{P})_{\text{obs}} - \delta(^{31}\text{P} \text{ of Et}_3\text{PO})$, within the set of compounds and with relevant examples known from the literature. All samples were analysed in dichloromethane-d₂ as the mostly used solvent, except for [NCN^{tBu}TeCl₂]OTf and [NCN^{tBu}TeCl₂]SbF₆ that are completely insoluble and acetonitrile had to be used instead. The Dipp substituted telluronium cations are also not included as they are not obtainable as pure compounds (*vide supra*).

Inspection of the $\Delta\delta(^{31}\text{P})$ values summarized in Table 3 revealed that neutral tellurium(II) compounds, not surprisingly, exhibited limited shift difference (entries 2 and 3); similarly the N,C,N-chelated telluronium cations (entries 7–11) showed only negligible influence as the Lewis acidity of the central atom is effectively compensated by the pincer ligands. In contrast, in the case of C,N-coordinated telluronium cations (entries 4–6), $\Delta\delta(^{31}\text{P})$ in dichloromethane-d₂ between 19.0 and 22.0 ppm indicate remarkable Lewis acidity as these values approach that of the well-established borane B(C₆F₅)₃ with $\Delta\delta(^{31}\text{P}) \sim 26$ ppm.^{34i,j} Regarding the telluronium cations bearing pincer ligands (entries 15 and 16), the $\Delta\delta(^{31}\text{P})$ is surprisingly quite different depending on the counter-anion. It could be speculated that the forced utilization of acetonitrile-d₃ may play a role in this phenomenon. In contrast, the C,N-chelated telluronium cations (entries 12 and 13) showed impressive $\Delta\delta(^{31}\text{P})$ values of 37.9 and 45.5 ppm, for [CN^{tBu}TeCl₂]OTf and [CN^{tBu}TeCl₂]SbF₆ respectively. These values are well comparable *e.g.* to those reported for silylium ions (39–45 ppm),^{34a,c} neutral super Lewis acidic silanes^{34e,f,m}

Table 3 $\delta(^{31}\text{P})_{\text{obs}}$ [ppm] of mixture of Et₃PO and studied compounds (1 : 5 molar ratio, dichloromethane-d₂); $\Delta\delta(^{31}\text{P}) = \delta(^{31}\text{P})_{\text{obs}} - \delta(^{31}\text{P}, \text{Et}_3\text{PO})$ [ppm]

Entry	Compound	$\delta(^{31}\text{P})_{\text{obs}}$	$\Delta\delta(^{31}\text{P})$
1	Pure Et ₃ PO	50.7	0
2	CN ^{tBu} TeCl	50.8	0.1
3	CN ^{Dipp} TeCl	50.7	0
4	[CN ^{tBu} Te]OTf	69.7	19.0
5	[CN ^{tBu} Te]SbF ₆	70.3	19.6
6	[CN ^{Dipp} Te]SbF ₆	72.7	22.0
7	[NCN ^{tBu} Te]Br	51.4	0.8
8	[NCN ^{tBu} Te]OTf	54.4	3.7
9	[NCN ^{tBu} Te]SbF ₆	51.1	0.4
10	[NCN ^{Dipp} Te]Br	51.5	0.8
11	[NCN ^{Dipp} Te]SbF ₆	54.6	3.9
12	[CN ^{tBu} TeCl ₂]OTf	88.6	37.9
13	[CN ^{tBu} TeCl ₂]SbF ₆	96.2	45.5
14	Pure Et ₃ PO ^a	50.4	0
15	[NCN ^{tBu} TeCl ₂]OTf ^a	66.2	15.8
16	[NCN ^{tBu} TeCl ₂]SbF ₆ ^a	54.6	4.2

^a Measured in acetonitrile-d₃ due to the solubility problems.



(~35 ppm) or organofluorophosphonium salts (~35 ppm).³⁵ This fact also well corresponds to their extremely high affinity to hydrolysis and make it challenging to utilize them as Lewis acids in catalysis.

DFT computations

Density functional theory (DFT) computations were conducted on the compounds $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{SbF}_6$ and $[\text{CN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$ as well as on the cations $[\text{NCN}^{\text{tBu}}\text{Te}]^+$, $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]^+$ and $[\text{NCN}^{\text{Dipp}}\text{Te}]^+$. In order to gain insights into the N–Te bonding, a set of real-space bonding descriptors have been studied. According to the atoms-in-molecules³⁶ analysis, N–Te bond critical points (bcp) have been observed in all studied cases, showing characteristics of polar covalent bonding contributions. The effect of the Dipp-substituent on the nature of the N–Te bond was studied by comparing the compound pairs $[\text{CN}^{\text{R}}\text{Te}]\text{SbF}_6$ ($\text{R} = t\text{-Bu}^{23}$ and Dipp) and $[\text{NCN}^{\text{R}}\text{Te}]^+$ ($\text{R} = t\text{-Bu}$ and Dipp) and revealed no substantial differences in the N–Te bonding (Table S2†). The additional steric effects imposed by the Dipp group are also discernible in the AIM picture, as additional $\text{CH}\cdots\text{Te}$ bcps are observed, which are absent in the respective $t\text{-Bu}$ species (Fig. S93–S98†).

In contrast, single (CN) or double (NCN) N-coordination to the Te-atom has a significant effect on the (individual) N–Te chalcogen bonds. Key parameters of the AIM analysis gave rise to smaller electron densities ($\rho_{\text{bcp}}(r)$) and Laplacians ($\nabla^2(\rho_{\text{bcp}}(r))$) at the bcp for $[\text{NCN}^{\text{R}}\text{Te}]^+$ ($0.51\text{--}0.56 \text{ e } \text{\AA}^{-3}/3.0\text{--}3.3 \text{ e } \text{\AA}^{-5}$) compared to $[\text{CN}^{\text{R}}\text{Te}]\text{SbF}_6$ ($0.73\text{--}0.77 \text{ e } \text{\AA}^{-3}/4.7\text{--}5.0 \text{ e } \text{\AA}^{-5}$), indicating a stronger coordination of the single N-ligand in the latter (Table S1†). This is also revealed in the kinetics ($G/\rho_{\text{bcp}}(r)$) and total energies ($H/\rho_{\text{bcp}}(r)$) over ρ values. Both parameters are closer to zero in $[\text{NCN}^{\text{R}}\text{Te}]^+$ than in $[\text{CN}^{\text{R}}\text{Te}]\text{SbF}_6$, suggesting weaker ionic and covalent bonding contributions in the pincer type compounds (Table S1†). However, it should be noted that these pincer ligands lead to a more balanced coordination in $[\text{NCN}^{\text{R}}\text{Te}]^+$, whereas in $[\text{CN}^{\text{R}}\text{Te}]\text{SbF}_6$ there is a stronger (N–Te) and weaker (F–Te) coordination. This is also reflected upon inspection of the Wiberg bond indices, NLMO/NPA bond orders and the delocalization indices for the N–Te interaction (Table S6†), which show smaller values for $[\text{NCN}^{\text{R}}\text{Te}]^+$ compared to $[\text{CN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$. Interestingly, the electron population of the disynaptic ELI-D³⁷ $V_2(\text{N},\text{Te})$ bonding basin is less affected by the single or double coordination, showing populations of 2.48 e ($[\text{CN}^{\text{tBu}}\text{Te}]\text{SbF}_6^{23}$), 2.58 e ($[\text{CN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$), $2.49/2.58 \text{ e}$ ($[\text{NCN}^{\text{tBu}}\text{Te}]^+$) and $2.48/2.49 \text{ e}$ ($[\text{NCN}^{\text{Dipp}}\text{Te}]^+$). The non-covalent interaction (NCI) index³⁸ is a powerful tool in unravelling weak, non-covalent bonding contributions and for the N–Te bonds in $[\text{NCN}^{\text{Dipp}}\text{Te}]^+$ clear red-colored shaped rings are observed, which is barely visible in the single N–Te bond of $[\text{CN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$, indicating the stronger covalent bonding contributions in the latter (see Fig. S95†). Interestingly, based on the AIM analysis there is only a little effect on the doubly coordinating N–Te interaction upon chlorination from $[\text{NCN}^{\text{tBu}}\text{Te}]^+$ to $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]^+$. The $\rho_{\text{bcp}}(r)$ and $H/\rho_{\text{bcp}}(r)$ values are very similar in both compounds and the slight decrease of the Laplacians ($\nabla^2(\rho_{\text{bcp}}(r))$) and

$G/\rho_{\text{bcp}}(r)$ values in $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]^+$ points to a minimal reduction of ionic bonding contributions.

NBO³⁹ analysis reveals no distinct Te–N bonding orbitals, but second order perturbation theory gives rise to $\text{LP}(\text{N}) \rightarrow \text{LV}(\text{Te})$ or $\text{LP}(\text{N}) \rightarrow \text{RY}(\text{Te})$ donation of a total of $E_2 = 103.73\text{--}119.73 \text{ kcal mol}^{-1}$ for the pincer ligands $[\text{NCN}^{\text{R}}\text{Te}]^+$ (Table S4†), which is in agreement with similar NCN-stabilized $\text{Te}(\text{II})$ cations.^{5d,9}

For $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{SbF}_6$, we investigated two possible structural isomers: one with a linear Cl–Te–Cl linkage and another with a nearly rectangular Cl–Te–Cl linkage. The gas-phase structure of the latter form is energetically favoured by $-51.24 \text{ kJ mol}^{-1}$ and resembles also the experimentally obtained geometry (Fig. 4, although with OTf^- as anion). Interestingly, the bonding situation in both isomers differs significantly. With a rectangular Cl–Te–Cl linkage the N–Te bonding can be compared to the N–Te interaction in the pincer compounds of the $[\text{NCN}^{\text{tBu}}\text{Te}]^+$ type with $\rho_{\text{bcp}}(r)$ values and respective Laplacians of $0.51 \text{ e } \text{\AA}^{-3}/2.2 \text{ e } \text{\AA}^{-5}$. Also the $G/\rho_{\text{bcp}}(r)$ and $H/\rho_{\text{bcp}}(r)$ values are very similar to the ones in $[\text{NCN}^{\text{tBu}}\text{Te}]^+$. The isomer with a linear Cl–Te–Cl linkage shows stronger N–Te coordination as observed *e.g.* by higher $\rho_{\text{bcp}}(r)$ and Laplacian values ($0.64 \text{ e } \text{\AA}^{-3}/2.8 \text{ e } \text{\AA}^{-5}$), but are smaller compared to the $\text{Te}(\text{II})$ species $[\text{CN}^{\text{R}}\text{Te}]\text{SbF}_6$. The F–Te interaction in the rectangular Cl–Te–Cl form is also weakened in comparison to the form with a linear Cl–Te–Cl linkage. The effects in both isomers on the F–Te and N–Te interaction can also be observed in the NCI. The incomplete red-colored ring around the N–Te in the linear Cl–Te–Cl form (Fig. 6, right) points towards stronger covalent interactions compared to the clearly visible ring-shape in the rectangular form (Fig. 6, left). Furthermore, the considerably weaker F–Te interaction in the rectangular form is reflected by the blue disk-shaped area on the F–Te axis in the NCI (Fig. 6, left).

Interestingly, the NBO analysis of the rectangular form shows a 3-center, 4-electron Cl–Te–N bond, with Cl–Te/Te–N contributions of 65.0/35.0% (Table S5†).

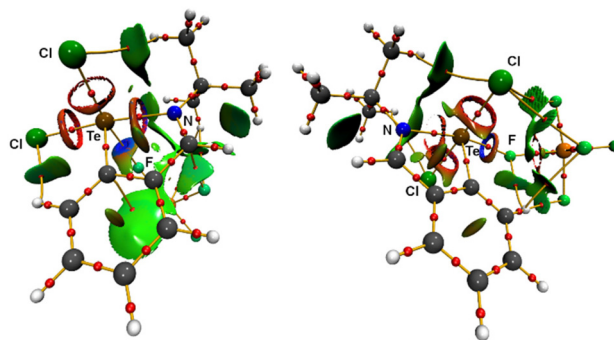


Fig. 6 AIM molecular graph of $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{SbF}_6$ with rectangular Cl–Te–Cl linkage (left) and linear Cl–Te–Cl linkage (right) with bond critical points as red spheres and bond paths in orange as well as NCI iso-surfaces at $s(r) = 0.5$ colour coded with $\text{sign}(\lambda_2)\rho$ in a. u. Blue surfaces refer to attractive forces and red to repulsive forces. Green indicates weak interactions.



Conclusions

We have demonstrated the potential of both C,N- and N,C,N-chelating ligands, containing imino-donor function(s), for the stabilization of tellurium and rare examples of dichlorotelluronium cations. The combined solution and solid-state ^{125}Te NMR study showed that the structure revealed by sc-X-ray diffraction analysis is retained in solution in the case of tellurium cations. Two dichlorotelluronium cations underwent a controlled hydrolysis with a concomitant condensation producing either a telluroxane with the $-\text{Te}(\text{Cl})_2-\text{O}-\text{Te}(\text{Cl})_2-$ bridge or, in the case of the pincer compound, the condensation occurred at the ligand arm.

Furthermore, the Gutmann–Beckett method was applied for a straightforward assessment of the Lewis acidity of the reported cations. Based on the obtained results, it became obvious that pincer ligands significantly suppress the acidity of the tellurium atom due to the presence of the second nitrogen donor atom. In contrast, the C,N-chelated telluronium cations showed values comparable to $\text{B}(\text{C}_6\text{F}_5)_3$, thereby indicating their potential in Lewis acid induced activation of various substrates. This is consistent with recently reported B–H bond activation of a carborane cage²⁴ and the field will be developed in the future. More importantly, the C,N-chelated telluronium cations possess remarkable Lewis acidity as one can deduce from the very high $\Delta\delta(^{31}\text{P})$ values obtained by the Gutmann–Beckett method. Unfortunately, the extremely high propensity of these compounds to hydrolysis may hamper their further reasonable application.

Our next steps will be directed towards the extension of the applicability of C,N-chelated telluronium cations in the activation of E–H bonds (E = B or Si) and building of more robust C,N-chelated telluronium cations. In particular, the latter as Lewis acids hold remarkable potential for further investigation of various bond activations or catalysis.

Experimental

General consideration

All syntheses were performed under an argon atmosphere using standard Schlenk techniques except for the hydrolysis of $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$ and $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$. All used solvents were dried using MD7 Pure Solv instrument (Innovative Technology, MA, USA), degassed and stored in Young-valve containers over potassium mirror or molecular sieves (3 Å). Deuterated solvents were dried by standard procedures and stored over molecular sieves. Procedures for the synthesis of all already published compounds are referenced in the main text. All other chemicals were purchased from commercial companies and used as delivered. ^1H , $^{13}\text{C}\{^1\text{H}\}$, and $^{125}\text{Te}\{^1\text{H}\}$ NMR spectra were recorded on a Bruker 400 or Bruker 500 spectrometers, using a 5 mm tunable broadband probe or the cryoprobe Prodigy. Chemical shifts in the ^1H and $^{13}\text{C}\{^1\text{H}\}$ NMR spectra were referenced to the residual solvent signals (CDCl_3 : $\delta(^1\text{H}) = 7.27$ ppm, $\delta(^{13}\text{C}) = 77.23$ ppm, CD_2Cl_2 : $\delta(^1\text{H}) = 5.32$ ppm, $\delta(^{13}\text{C})$

$= 54.00$ ppm, acetonitrile- d_3 : $\delta(^1\text{H}) = 1.94$ ppm, $\delta(^{13}\text{C}) = 1.39$ ppm, dmsO-d_6 : $\delta(^1\text{H}) = 2.50$ ppm, and $\delta(^{13}\text{C}) = 39.51$ ppm). ^{15}N NMR spectra are referenced to external neat nitromethane [$\delta(^{15}\text{N}) = 0.0$ ppm]. All ^{15}N chemical shifts were obtained *via* 2D ^{15}N , ^1H HMBC experiment. The ^{125}Te NMR chemical shifts are referenced to an external CDCl_3 solution of Ph_2Te_2 [$\delta(^{125}\text{Te}) = 422$ ppm relative to Me_2Te]. Solid-state NMR spectra of ^{125}Te (Fig. S89†) were acquired on a JEOL 600 MHz spectrometer under magic-angle-spinning conditions at 18 kHz MAS frequency without active temperature regulation. Samples were packed into 3.2 mm zirconia rotors under an inert atmosphere and spun using N_2 . Spectra were acquired using direct excitation experiment (90° pulse of 3.5 us) with direct FID detection. Typical pre-acquisition delay was set 120 s. The isotropic shift was identified by comparison with spectra acquired at a slower MAS rate and referenced using an external sample of solid $\text{Te}(\text{OH})_6$ with $\delta(^{125}\text{Te}) = 692.2$ and 685.6 ppm (two independent molecules in the unit cell).⁴⁰ High-resolution MALDI MS spectra were measured with a MALDI mass spectrometer LTQ Orbitrap XL (Thermo Fisher Scientific, Bremen, Germany) equipped with a nitrogen UV laser (337 nm, 60 Hz). The LTQ Orbitrap instrument was operated in positive-ion mode over a normal mass range (m/z 50–2000) with 100 000 resolution at $m/z = 400$. The survey crystal positioning system (survey CPS) was set for the random choice of shot position by automatic crystal recognition. *trans*-2-[3-(4-*tert*-Butylphenyl)-2-methyl-2-propenylidene]malononitrile (DCTB) was used as a matrix. Mass spectra were averaged over the whole MS record for all measured samples.

The bulk purity of tellurium(II) compounds was established by high-resolution MALDI MS spectra in combination with multinuclear NMR spectroscopy (see the ESI†). Unfortunately, telluronium cations could be characterized by multinuclear NMR spectroscopy only (see the ESI†) because their high sensitivity toward hydrolysis hampered our attempts to get satisfactory high-resolution MS spectra or combustion analysis.

Syntheses of starting compounds

Synthesis of $\text{CN}^{\text{Dipp}}\text{TeCl}$. 1.03 g (1.91 mmol) of $\text{CN}^{\text{Dipp}}\text{Te}$ (dte) was dissolved in CH_2Cl_2 (50 mL) and 3.82 mL (3.82 mmol) of 1 M solution of HCl in diethylether. A yellow precipitate formed and the reaction mixture was stirred overnight and then evaporated *in vacuo*. The obtained solid was extracted with toluene (2×20 mL) and the yellow extract was crystallized at -20°C . Yellow crystals were collected by decantation and dried *in vacuo*. Yield 760 mg, 93%, m.p. 211–215 $^\circ\text{C}$. ^1H NMR (500 MHz, CDCl_3): δ 1.13 [6H, d, $^3J_{\text{H,H}} = 6.7$ Hz, iPr- CH_3]; 1.24 [6H, d, $^3J_{\text{H,H}} = 6.7$ Hz, iPr- CH_3]; 2.70 [2H, sept, $^3J_{\text{H,H}} = 6.7$ Hz, iPr-CH]; 7.27 [2H, d, $^3J_{\text{H,H}} = 7.5$ Hz, Ar-H]; 7.39 [1H, t, $^3J_{\text{H,H}} = 7.5$ Hz, Ar-H]; 7.58 [1H, t, $^3J_{\text{H,H}} = 7.4$ Hz, Ar-H]; 7.69 [1H, t, $^3J_{\text{H,H}} = 7.3$ Hz, Ar-H]; 8.11 [1H, d, $^3J_{\text{H,H}} = 7.7$ Hz, Ar-H]; 8.80 [1H, d, $^3J_{\text{H,H}} = 8.0$ Hz, Ar-H]; 9.16 [1H, s, $\text{CH}=\text{N}$] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (100.61 MHz, CDCl_3): δ 24.4 [iPr- CH_3]; 25.1 [iPr- CH_3]; 28.6 [iPr-CH]; 124.2 [Ar-C]; 126.9 [Ar-C]; 129.0 [Ar-C]; 132.3 [Ar-C]; 133.1 [Ar-C]; 133.8 [Ar-C]; 135.1 [Ar-C]; 138.9 [Ar-C]; 142.3 [Ar-C]; 144.1 [Ar-C]; 164.7



[CH=N] ppm. ^{15}N NMR (via ^{15}N , ^1H HMBC, CDCl_3): δ -121.4 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CDCl_3): δ 1392 ppm. ^{125}Te NMR (solid): δ_{iso} 1349 and 1403 ppm (two independent molecules in the unit cell). HRMS (MALDI) m/z calc. for $\text{C}_{19}\text{H}_{22}\text{N}^{130}\text{Te}$: 394.0809 $[\text{M} - \text{Cl}]^+$, found 394.0805.

Synthesis of $[\text{NCN}^{\text{tBu}}\text{Te}]\text{Br}$. 2.00 g (6.19 mmol) of $\text{NCN}^{\text{tBu}}\text{Br}$ was dissolved in thf (70 mL), cooled to -80°C and 2.48 mL (6.20 mmol) of 2.5 M solution of $n\text{BuLi}$ in hexane was added dropwise. The resulting red solution was stirred for 1 h at this temperature and then added to a solution of $\text{Te}(\text{dte})_2$ 2.63 g (6.20 mmol) in thf (50 mL). The reaction mixture was stirred overnight at r.t. and during this time a yellow precipitate formed. This was collected by filtration, washed with thf (2 mL), hexane (10 mL) and dried *in vacuo* to give $[\text{NCN}^{\text{tBu}}\text{Te}]\text{Br}$ as pale-yellow powder. Single crystals could be obtained *via* crystallization from saturated dichloromethane solution. Yield 1.9 g, 68%, m.p. $237\text{--}240^\circ\text{C}$. ^1H NMR (500 MHz, CD_2Cl_2): δ 1.71 [18H, s, tBu-CH_3]; 7.91 [1H, t, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 8.76 [2H, d, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 10.27 [2H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.61 MHz, CD_2Cl_2): δ 32.4 [tBu-CH_3]; 63.3 [tBu-C]; 129.3 [Ar-C]; 135.5 [Ar-C]; 136.6 [Ar-C]; 136.8 [Ar-C]; 160.7 [CH=N] ppm. ^{15}N NMR (via ^{15}N , ^1H HMBC, CD_2Cl_2): δ -91.2 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CD_2Cl_2): δ 1394 ppm. HRMS (MALDI) m/z calc. for $\text{C}_{19}\text{H}_{22}\text{N}^{130}\text{Te}$: 373.0918 $[\text{M} - \text{Br}]^+$, found 394.0913.

Synthesis of $[\text{NCN}^{\text{Dipp}}\text{Te}]\text{Br}$. 1.01 g (1.90 mmol) of $\text{NCN}^{\text{Dipp}}\text{Br}$ was dissolved in thf (25 mL) and cooled to -80°C and 0.76 mL (1.90 mmol) of 2.5 M solution of $n\text{BuLi}$ in hexane. The reaction mixture was stirred for 1 h at this temperature and then it was added to a solution of $\text{Te}(\text{dte})_2$ 0.81 g (1.90 mmol) in thf (20 mL). The resulting red-brown solution was stirred overnight at r.t. and concentrated to *ca.* 10 mL that caused precipitation of a yellow solid that was collected by filtration at 0°C . This solid was washed with thf (5 mL) and hexane (10 mL) and dried *in vacuo* giving $[\text{NCN}^{\text{Dipp}}\text{Te}]\text{Br}$ as a yellow powder. Yield 475 mg, 38%, m.p. $329^\circ\text{C}(\text{dec.})$. ^1H NMR (500 MHz, CDCl_3): δ 1.16 [12H, d, $^3J_{\text{H,H}} = 6.9$ Hz, iPr-CH_3]; 1.23 [12H, d, $^3J_{\text{H,H}} = 6.9$ Hz, iPr-CH_3]; 2.61 [4H, sept, $^3J_{\text{H,H}} = 6.9$ Hz, iPr-CH]; 7.26 [4H, d, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 7.39 [2H, t, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 8.18 [1H, t, $^3J_{\text{H,H}} = 7.7$ Hz, Ar-H]; 9.44 [2H, d, $^3J_{\text{H,H}} = 7.7$ Hz, Ar-H]; 10.31 [2H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.6 MHz, CDCl_3): δ 24.3 [iPr-CH_3]; 24.8 [iPr-CH_3]; 28.8 [iPr-CH]; 124.5 [Ar-C]; 129.6 [Ar-C]; 130.4 [Ar-C]; 134.5 [Ar-C]; 138.6 [Ar-C]; 140.2 [Ar-C]; 141.3 [Ar-C]; 141.6 [Ar-C]; 166.5 [CH=N] ppm. ^{15}N NMR (via ^{15}N , ^1H HMBC, CDCl_3) δ -119.8 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CDCl_3) δ 2113 ppm. HRMS (MALDI) m/z calc. for $\text{C}_{32}\text{H}_{39}\text{N}_2^{130}\text{Te}$: 581.2170 $[\text{M} - \text{Br}]^+$, found 581.2161.

General procedure for the synthesis of tellurium(II) cations

The particular tellurium precursor was mixed with 1 eq. of the corresponding silver salt AgX ($\text{X} = \text{OTf}$ or SbF_6) and dichloromethane was added to this mixture. The reaction mixture was stirred for 1 h while the reaction flasks were protected from ambient light by an aluminum foil. Then, the precipitated AgX ($\text{X} = \text{Cl}$ or Br) was filtered and the filtrate was evaporated and

the resulting solid was washed with hexane to give the target tellurium(II) cations. For particular loadings, yields and experimental data, see below.

Synthesis of $[\text{NCN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$. 1.02 g (2.38 mmol) of $\text{NCN}^{\text{Dipp}}\text{TeCl}$ and 0.82 g of AgSbF_6 gave $[\text{NCN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$ as yellow-orange powder. Single-crystals could be obtained by layering of a saturated dichloromethane solution with hexane. Yield 1.47 g, 98%, m.p. $231\text{--}236^\circ\text{C}$. ^1H NMR (500 MHz, CD_2Cl_2): δ 1.20 [6H, d, $^3J_{\text{H,H}} = 6.8$ Hz, iPr-CH_3]; 1.25 [6H, d, $^3J_{\text{H,H}} = 6.8$ Hz, iPr-CH_3]; 2.55 [2H, sept, $^3J_{\text{H,H}} = 6.8$ Hz, iPr-CH]; 7.37 [2H, d, $^3J_{\text{H,H}} = 7.9$ Hz, Ar-H]; 7.58 [1H, t, $^3J_{\text{H,H}} = 7.9$ Hz, Ar-H]; 7.80 [1H, t, $^3J_{\text{H,H}} = 7.9$ Hz, Ar-H]; 7.89 [1H, t, $^3J_{\text{H,H}} = 8.0$ Hz, Ar-H]; 8.43 [1H, d, $^3J_{\text{H,H}} = 8.1$ Hz, Ar-H]; 8.61 [1H, d, $^3J_{\text{H,H}} = 8.0$ Hz, Ar-H]; 9.37 [1H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (100.61 MHz, CD_2Cl_2): δ 24.3 [iPr-CH_3]; 25.0 [iPr-CH_3]; 29.2 [iPr-CH]; 125.5 [Ar-C]; 129.4 [Ar-C]; 132.0 [Ar-C]; 132.5 [Ar-C]; 135.3 [overlap of two signals based on HSQC, Ar-C]; 136.7 [Ar-C]; 137.1 [Ar-C]; 143.8 [Ar-C]; 150.7 [Ar-C]; 171.1 [CH=N] ppm. ^{15}N NMR (via ^{15}N , ^1H HMBC, CD_2Cl_2) δ -174.9 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CD_2Cl_2) δ 1392 ppm. HRMS (MALDI) m/z calc. for $\text{C}_{19}\text{H}_{22}\text{N}^{130}\text{Te}$: 394.0809 $[\text{M} - \text{SbF}_6]^+$, found 394.0803.

Synthesis of $[\text{NCN}^{\text{tBu}}\text{Te}]\text{OTf}$. 354 mg (0.79 mmol) of $[\text{NCN}^{\text{tBu}}\text{Te}]\text{Br}$ and 202 mg of AgOTf gave $[\text{NCN}^{\text{tBu}}\text{Te}]\text{OTf}$ as yellowish powder. Single-crystals suitable for X-ray diffraction analysis could not be obtained at this moment. Yield 388 mg, 95%, m.p. $206\text{--}207^\circ\text{C}$. ^1H NMR (500 MHz, CD_2Cl_2): δ 1.71 [18H, s, tBu-CH_3]; 7.98 [1H, t, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 8.56 [2H, d, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 9.75 [2H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.61 MHz, CD_2Cl_2): δ 32.3 [tBu-CH_3]; 63.4 [tBu-C]; 121.6 [q, $^1J_{\text{F,C}} = 322$ Hz, CF_3]; 129.6 [Ar-C]; 135.3 [Ar-C]; 136.5 [Ar-C]; 136.8 [Ar-C]; 160.0 [CH=N] ppm. $^{19}\text{F}\{^1\text{H}\}$ NMR (376.5 MHz, CD_2Cl_2) δ -78.8 ppm. ^{15}N NMR (via ^{15}N , ^1H HMBC, CD_2Cl_2) δ -88.8 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CD_2Cl_2) δ 1400 ppm. HRMS (MALDI) m/z calc. for $\text{C}_{19}\text{H}_{22}\text{N}^{130}\text{Te}$: 373.0918 $[\text{M} - \text{CF}_3\text{O}_3\text{S}]^+$, found 373.091.

Synthesis of $[\text{NCN}^{\text{tBu}}\text{Te}]\text{SbF}_6$. 298 mg (0.66 mmol) of $[\text{NCN}^{\text{tBu}}\text{Te}]\text{Br}$ and 227 mg of AgSbF_6 gave $[\text{NCN}^{\text{tBu}}\text{Te}]\text{SbF}_6$ as yellowish powder. Single-crystals could be obtained by layering a saturated dichloromethane solution with hexane. Yield 389 mg, 97%, m.p. $268\text{--}272^\circ\text{C}$. ^1H NMR (500 MHz, CD_2Cl_2): δ 1.71 [18H, s, tBu-CH_3]; 8.01 [1H, t, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 8.45 [2H, d, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 9.57 [2H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.61 MHz, CD_2Cl_2): δ 32.3 [tBu-CH_3]; 63.5 [tBu-C]; 129.7 [Ar-C]; 135.2 [Ar-C]; 136.3 [Ar-C]; 137.1 [Ar-C]; 159.4 [CH=N] ppm. ^{15}N NMR (via ^{15}N , ^1H HMBC, CD_2Cl_2) δ -89.5 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CD_2Cl_2) δ 1409 ppm. ^{125}Te NMR (solid): δ_{iso} 1388 ppm. HRMS (MALDI) m/z calc. for $\text{C}_{19}\text{H}_{22}\text{N}^{130}\text{Te}$: 373.0918 $[\text{M} - \text{SbF}_6]^+$, found 373.0912.

Synthesis of $[\text{NCN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$. 438 mg (0.66 mmol) of $[\text{NCN}^{\text{Dipp}}\text{Te}]\text{Br}$ and 228 mg of AgSbF_6 gave $[\text{NCN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$ as yellow-orange powder. Single-crystals could be obtained by layering of saturated dichloromethane solution with hexane at 6°C . Yield 506 mg, 94%, m.p. $249\text{--}254^\circ\text{C}(\text{dec.})$. ^1H NMR (500 MHz, CD_2Cl_2): δ 1.18 [12H, d, $^3J_{\text{H,H}} = 6.9$ Hz, iPr-CH_3]; 1.20 [12H, d, $^3J_{\text{H,H}} = 6.9$ Hz, iPr-CH_3]; 2.62 [4H, sept, $^3J_{\text{H,H}} =$



6.9 Hz, *i*Pr-CH]; 7.32 [4H, d, $^3J_{\text{H,H}} = 7.9$ Hz, Ar-H]; 7.46 [2H, t, $^3J_{\text{H,H}} = 7.9$ Hz, Ar-H]; 8.24 [1H, t, $^3J_{\text{H,H}} = 7.7$ Hz, Ar-H]; 8.44 [2H, d, $^3J_{\text{H,H}} = 7.7$ Hz, Ar-H]; 9.60 [2H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.6 MHz, CD_2Cl_2): δ 24.5 [iPr-CH₃]; 24.6 [iPr-CH₃]; 29.2 [iPr-CH]; 125.0 [Ar-C]; 130.1 [Ar-C]; 130.7 [Ar-C]; 134.7 [Ar-C]; 138.8 [Ar-C]; 139.0 [Ar-C]; 142.0 [Ar-C]; 142.8 [Ar-C]; 166.9 [CH=N] ppm. ^{15}N NMR (*via* ^{15}N , ^1H HMBC, CD_2Cl_2) δ -117.4 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CD_2Cl_2): δ 1557 ppm. HRMS (MALDI) *m/z* calc. for $\text{C}_{32}\text{H}_{39}\text{N}_2^{130}\text{Te}$: 581.2170 [$\text{M} - \text{SbF}_6$]⁺, found 581.2156.

Oxidation of tellurium(II) cations by SO_2Cl_2

Synthesis of $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$. 702 mg (1.61 mmol) of $[\text{CN}^{\text{tBu}}\text{Te}]\text{OTf}$ was dissolved in CH_2Cl_2 (15 mL) and 130 μL (1.61 mmol) of neat SO_2Cl_2 was added with stirring. The originally yellow solution became colorless and the reaction mixture was stirred for 20 min. The reaction mixture was evaporated *in vacuo* to give $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$. Single-crystals were obtained by slow cooling of saturated warm dichloromethane solution. Yield 816 mg, 99%, m.p. 176 °C. ^1H NMR (500 MHz, CD_3CN): δ 1.65 [9H, s, *t*Bu-CH₃]; 7.97 [1H, t, $^3J_{\text{H,H}} = 7.7$ Hz, Ar-H]; 8.05 [1H, t, $^3J_{\text{H,H}} = 7.8$ Hz, Ar-H]; 8.13 [1H, d, $^3J_{\text{H,H}} = 7.8$ Hz, Ar-H]; 8.66 [1H, d, $^3J_{\text{H,H}} = 7.9$ Hz, Ar-H]; 9.27 [1H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.61 MHz, CD_3CN): δ 30.5 [*t*Bu-CH₃]; 65.2 [*t*Bu-C]; 120.9 [q, $^1J_{\text{F,C}} = 319$ Hz, CF₃]; 132.0 [Ar-C]; 135.1 [Ar-C]; 135.5 [Ar-C]; 136.5 [Ar-C]; 137.1 [Ar-C]; 143.8 [Ar-C]; 168.3 [CH=N] ppm. $^{19}\text{F}\{^1\text{H}\}$ NMR (376.5 MHz, CD_2Cl_2) δ -79.2 ppm. ^{15}N NMR (*via* ^{15}N , ^1H HMBC, CD_3CN) δ -86.8 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CD_3CN) δ 1373 ppm.

Synthesis of $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{SbF}_6$. 405 mg (0.77 mmol) of $[\text{CN}^{\text{tBu}}\text{Te}]\text{SbF}_6$ was dissolved in CH_2Cl_2 (50 mL) and 75 μL (0.93 mmol) of neat SO_2Cl_2 was added with stirring. A colorless precipitate formed and the reaction mixture was stirred for 30 min. The white solid was collected by filtration and dried *in vacuo* to give $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$. Single-crystals suitable for X-ray diffraction analysis could not be obtained at this moment. Yield 431 mg, 94%, m.p. 191 °C(dec.). ^1H NMR (500 MHz, CD_3CN): δ 1.64 [9H, s, *t*Bu-CH₃]; 8.03 [1H, t, $^3J_{\text{H,H}} = 7.8$ Hz, Ar-H]; 8.10 [1H, t, $^3J_{\text{H,H}} = 7.7$ Hz, Ar-H]; 8.18 [1H, d, $^3J_{\text{H,H}} = 7.8$ Hz, Ar-H]; 8.68 [1H, d, $^3J_{\text{H,H}} = 7.9$ Hz, Ar-H]; 9.27 [1H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.61 MHz, CD_3CN): δ 30.4 [*t*Bu-CH₃]; 65.5 [*t*Bu-C]; 132.7 [Ar-C]; 135.8 [Ar-C]; 136.0 [Ar-C]; 136.8 [Ar-C]; 137.5 [Ar-C]; 141.7 [Ar-C]; 168.6 [CH=N] ppm. ^{15}N NMR (*via* ^{15}N , ^1H HMBC, CD_3CN) δ -87.1 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CD_3CN) δ 1399 ppm.

Synthesis of $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$. 926 mg (1.78 mmol) of $[\text{NCN}^{\text{tBu}}\text{Te}]\text{OTf}$ was dissolved in CH_2Cl_2 (25 mL) and 144 μL (1.78 mmol) of neat SO_2Cl_2 was added with stirring. The originally yellow solution turned to a white suspension that was stirred for an additional 24 h. The precipitate was collected by filtration and dried *in vacuo* to give $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$ as white powder. Single-crystals were grown by slow cooling of saturated warm dichloromethane solution. Yield 547 mg, 52%, m.p. 220 °C(dec.). ^1H NMR (400 MHz, CD_3CN): δ 1.73 [18H, s, *t*Bu-CH₃]; 8.16 [1H, t, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 8.42 [2H, d, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 9.54 [2H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR

(125.61 MHz, CD_3CN): δ 30.6 [*t*Bu-CH₃]; 66.8 [*t*Bu-C]; 121.5 [q, $^1J_{\text{F,C}} = 320$ Hz, CF₃]; 134.5 [Ar-C]; 136.0 [Ar-C]; 138.5 [Ar-C]; 145.8 [Ar-C]; 163.2 [CH=N] ppm. $^{19}\text{F}\{^1\text{H}\}$ NMR (376.5 MHz, CD_3CN) δ -79.2 ppm. ^{15}N NMR (*via* ^{15}N , ^1H HMBC, CD_3CN) δ -86.7 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CD_3CN) δ 1180 ppm.

Synthesis of $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{SbF}_6$. 108 mg (0.18 mmol) of $[\text{NCN}^{\text{tBu}}\text{Te}]\text{SbF}_6$ was dissolved in CH_2Cl_2 (15 mL) and 16 μL (0.2 mmol) of neat SO_2Cl_2 was added with stirring. The originally yellow solution turned to white suspension that was stirred for an additional 30 min. The precipitate was collected by filtration and dried *in vacuo* to give $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{SbF}_6$ as white powder. Single-crystals were obtained by slow cooling of saturated warm dichloromethane solution. Yield 118 mg, 98%, m.p. 266 °C(dec.). ^1H NMR (500 MHz, CD_3CN): δ 1.72 [18H, s, *t*Bu-CH₃]; 8.16 [1H, t, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 8.40 [2H, d, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 9.40 [2H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.61 MHz, CD_3CN): δ 30.7 [*t*Bu-CH₃]; 66.9 [*t*Bu-C]; 134.5 [Ar-C]; 136.1 [Ar-C]; 138.6 [Ar-C]; 146.0 [Ar-C]; 163.2 [CH=N] ppm. ^{15}N NMR (*via* ^{15}N , ^1H HMBC, CD_3CN) δ -87.0 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CD_3CN) δ 1179 ppm.

***In situ* generation of $[\text{CN}^{\text{Dipp}}\text{TeCl}_2]\text{SbF}_6$.** 30 mg (0.05 mmol) of $[\text{CN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$ was dissolved in CH_2Cl_2 (5 mL) and 20 μL (0.25 mmol, 5 eq.) of neat SO_2Cl_2 was added and the mixture was stirred for 20 min. The reaction mixture was evaporated, dissolved in CD_2Cl_2 and directly analysed by NMR spectroscopy. All attempts to isolate $[\text{CN}^{\text{Dipp}}\text{TeCl}_2]\text{SbF}_6$ even using higher loading of starting compound always resulted only in material contaminated by unidentified products of hydrolysis.

^1H NMR (500 MHz, CD_2Cl_2): δ 1.20 [6H, d, $^3J_{\text{H,H}} = 6.7$ Hz, *i*Pr-CH₃]; 1.30 [6H, d, $^3J_{\text{H,H}} = 6.7$ Hz, *i*Pr-CH₃]; 2.84 [2H, sept, $^3J_{\text{H,H}} = 6.7$ Hz, *i*Pr-CH]; 7.37 [2H, d, $^3J_{\text{H,H}} = 7.8$ Hz, Ar-H]; 7.47 [1H, t, $^3J_{\text{H,H}} = 7.8$ Hz, Ar-H]; 8.11 [1H, t, $^3J_{\text{H,H}} = 7.8$ Hz, Ar-H]; 8.20 [1H, t, $^3J_{\text{H,H}} = 8.0$ Hz, Ar-H]; 8.25 [1H, d, $^3J_{\text{H,H}} = 7.5$ Hz, Ar-H]; 8.78 [1H, d, $^3J_{\text{H,H}} = 8.0$ Hz, Ar-H]; 8.83 [1H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.6 MHz, CD_2Cl_2): δ 24.2 [iPr-CH₃]; 25.9 [iPr-CH₃]; 30.0 [iPr-CH]; 125.7 [Ar-C]; 130.6 [Ar-C]; 134.5 [Ar-C]; 134.8 [Ar-C]; 136.2 [overlap of two signals based on HSQC, Ar-C]; 137.2 [Ar-C]; 138.1 [Ar-C]; 138.9 [Ar-C]; 142.9 [Ar-C]; 170.3 [CH=N] ppm. ^{15}N NMR (*via* ^{15}N , ^1H HMBC, CD_2Cl_2) δ -102.8 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CD_2Cl_2) δ 1428 ppm.

***In situ* generation of $[\text{NCN}^{\text{Dipp}}\text{TeCl}_2]\text{SbF}_6$.** 102 mg (0.13 mmol) of $[\text{NCN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$ was dissolved in CH_2Cl_2 (5 mL) and 100 μL (1.13 mmol, 10 eq.) of neat SO_2Cl_2 was added and the mixture was stirred for 30 min. The reaction mixture was evaporated, dissolved in CD_2Cl_2 and directly analysed by NMR spectroscopy. All attempts to isolate $[\text{NCN}^{\text{Dipp}}\text{TeCl}_2]\text{SbF}_6$ even using higher loadings of starting material always resulted in crude mixtures contaminated by unidentified products of hydrolysis.

^1H NMR (500 MHz, CD_2Cl_2): δ 1.28 [12H, d, $^3J_{\text{H,H}} = 6.7$ Hz, *i*Pr-CH₃]; 1.31 [12H, d, $^3J_{\text{H,H}} = 6.7$ Hz, *i*Pr-CH₃]; 3.20 [4H, sept, $^3J_{\text{H,H}} = 6.7$ Hz, *i*Pr-CH]; 7.42 [4H, d, $^3J_{\text{H,H}} = 7.8$ Hz, Ar-H]; 7.54 [2H, t, $^3J_{\text{H,H}} = 7.8$ Hz, Ar-H]; 8.33 [1H, t, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 8.79 [2H, d, $^3J_{\text{H,H}} = 7.6$ Hz, Ar-H]; 9.22 [2H, s, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.6 MHz, CD_2Cl_2): δ 25.0 [iPr-CH₃]; 25.6 [iPr-CH₃]; 30.1 [iPr-CH]; 126.1 [Ar-C]; 131.5 [Ar-C]; 132.8 [Ar-C];



136.1 [Ar-C]; 139.0 [Ar-C]; 140.6 [Ar-C]; 144.0 [Ar-C], 150.0 [Ar-C]; 168.2 [CH=N] ppm. ^{15}N NMR (*via* ^{15}N , ^1H HMBC, CD_2Cl_2) δ -112.9 ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, CD_2Cl_2) δ 1308 ppm.

Preparation of crystals of $[\text{COTeCl}_2]_2\text{O}$ and $[\text{OCNtBuTeCl}_2]_2\text{O}$

Hydrolysis of $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$. 185 mg of $[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$ was dissolved in acetonitrile (3 mL) and a drop of distilled water was added and after mixing the mixture was left to evaporate slowly (to *ca.* 1/3 of the original volume) to give yellow single crystals of $[\text{COTeCl}_2]_2\text{O}$ suitable for X-ray diffraction analysis. These crystals were washed with acetonitrile and dried *in vacuo*. Yield 108 mg, 95%, m.p. 153 °C. The NMR data for $[\text{COTeCl}_2]_2\text{O}$: ^1H NMR (500 MHz, thf-d_8): δ 7.88 [1H, t, $^3J_{\text{H,H}} = 7.3$ Hz, Ar-H]; 8.13 [1H, t, $^3J_{\text{H,H}} = 7.7$ Hz, Ar-H]; 8.31 [1H, d, $^3J_{\text{H,H}} = 7.3$ Hz, Ar-H]; 8.72 [1H, d, $^3J_{\text{H,H}} = 7.7$ Hz, Ar-H]; 10.38 [1H, d, $^4J_{\text{H,H}} = 1$ Hz, CH=O] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.61 MHz, thf-d_8): δ 130.4 [Ar-C]; 132.8 [Ar-C]; 134.9 [Ar-C], 136.6 [Ar-C]; 139.6 [Ar-C]; 151.5 [Ar-C]; 200.5 [CH=O] ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, thf-d_8) δ 1405 ppm.

The NMR data for $[\text{COTeCl}_2]\text{OH}$: ^1H NMR (500 MHz, $\text{thf-d}_8/\text{water}$): δ 7.75 [1H, t, $^3J_{\text{H,H}} = 7.4$ Hz, Ar-H]; 8.00 [1H, t, $^3J_{\text{H,H}} = 7.4$ Hz, Ar-H]; 8.17 [1H, d, $^3J_{\text{H,H}} = 7.3$ Hz, Ar-H]; 8.38 [1H, d, $^3J_{\text{H,H}} = 7.5$ Hz, Ar-H]; 10.22 [1H, s, CH=O] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.61 MHz, $\text{thf-d}_8/\text{water}$): δ 127.2 [Ar-C]; 132.3 [Ar-C]; 134.3 [Ar-C], 136.3 [Ar-C]; 138.1 [Ar-C]; 150.2 [Ar-C]; 198.3 [CH=O] ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, d_8/water) δ 1342 ppm.

Hydrolysis of $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$. 274 mg of $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$ was dissolved in acetonitrile (5 mL) and a drop of distilled water was added and after mixing the mixture was left to evaporate slowly (to *ca.* 1/3 of the original volume) to give colorless single crystals of $[\text{NCOTeCl}_2]_2\text{O}$ suitable for X-ray diffraction analysis. These crystals were collected by decantation, washed with acetonitrile and dried *in vacuo*. Yield 180 mg, 98%, m.p. 214 °C(dec.). ^1H NMR (500 MHz, dmsO-d_6): δ 1.54 [9H, s, tBu-CH_3]; 7.30 [1H, s, CH]; 7.77 [1H, d, $^3J_{\text{H,H}} = 7.4$ Hz, Ar-H]; 7.84 [1H, t, $^3J_{\text{H,H}} = 7.4$ Hz, Ar-H]; 8.05 [1H, d, $^3J_{\text{H,H}} = 7.3$ Hz, Ar-H]; 9.14 [1H, CH=N] ppm. $^{13}\text{C}\{^1\text{H}\}$ NMR (125.61 MHz, dmsO-d_6): δ 30.1 [tBu-CH_3]; 61.3 [tBu-C]; 100.7 [CH]; 130.5 [Ar-C]; 132.0 [Ar-C]; 132.8 [Ar-C], 132.9 [Ar-C]; 142.6 [Ar-C]; 143.3 [Ar-C]; 157.3 [CH=N] ppm. $^{125}\text{Te}\{^1\text{H}\}$ NMR (158 MHz, dmsO-d_6) δ 1381 ppm.

Computational methodology

Geometry optimizations of the isolated molecular structures were carried out using density functional theory (DFT) at the B3PW91/6-311+G(2df,p)^{41,42} level of theory using the Gaussian16⁴³ software package. For the Sb and Te atoms, effective core potentials (ECP28MDF) and corresponding cc-pVTZ basis sets⁴⁴ were used. Dispersion effects were modelled using Grimme's GD3BJ parameters.⁴⁵ Wavefunction files were used for a topological analysis of the electron density according to the Atoms-In-Molecules partitioning scheme³⁶ using AIMAll,⁴⁶ whereas DGRID⁴⁷ was used to generate and analyze the Electron-Localizability-Indicator (ELI-D)³⁷ related real-space bonding descriptors applying a grid step size of 0.05 a.u.

NBO³⁹ analysis was carried out using the NBO6 software.⁴⁸ The NCI grids were computed with NCIPLOT.⁴⁹ AIM, ELI-D and NCI figures are displayed using Multiwfn 3.8⁵⁰ and VMD.⁵¹

Crystallography

The X-ray data for colourless crystals of studied compounds were obtained at 150 K using an Oxford Cryostream low-temperature device with a Bruker D8-Venture diffractometer equipped with a Mo (Mo/K α radiation; $\lambda = 0.71073$ Å) microfocus X-ray ($\text{I}\mu\text{S}$) source, and a photon CMOS detector was used for data collection. The obtained data were treated with XT-version 2019/1 and SHELXL-2019/1 software implemented in APEX3 v2016.9-0 (Bruker AXS) system.⁵²

The frames for all complexes were integrated with the Bruker SAINT software package using a narrow-frame algorithm. Data were corrected for absorption effects using the Multi-Scan method (SADABS). The structures were solved and refined using the Bruker SHELXTL software package. Hydrogen atoms were mostly localized on a difference Fourier map; however, to ensure uniformity of treatment of crystals, most of the hydrogen atoms were recalculated into idealized positions (riding model) and assigned temperature factors $H_{\text{iso}}(\text{H}) = 1.2U_{\text{eq}}$ (pivot atom) or $1.5U_{\text{eq}}$ (methyl). H atoms in methyl, methine moieties and C-H of imine or located in aromatic rings were placed with C-H distances of 0.96, 0.97, 0.98 and 0.93 Å. In $[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$, there is an electron density hole between the tellurium atom and *ipso* carbon of the aromatic ring of $\sim 2.7 \text{ e}^- \text{ Å}^{-3}$, which has no chemical sense.

Crystallographic data for all structural analysis have been deposited with the Cambridge Crystallographic Data Centre, CCDC 2254918 ($[\text{CN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$), 2254919 ($[\text{CN}^{\text{Dipp}}\text{TeCl}]$), 2254920 ($[\text{COTeCl}_2]_2\text{O}$), 2254921 ($[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{SbF}_6$), 2254922 ($[\text{NCN}^{\text{tBu}}\text{TeCl}_2]\text{OTf}$), 2254923 ($[\text{NCN}^{\text{tBu}}\text{Te}]\text{SbF}_6$), 2254924 ($[\text{NCN}^{\text{tBu}}\text{Te}]\text{Br}$), 2254925 ($[\text{CN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$), 2254926 ($[\text{OCN}^{\text{tBu}}\text{TeCl}_2]_2\text{O}$) and 2254927 ($[\text{NCN}^{\text{Dipp}}\text{Te}]\text{SbF}_6$).†

Conflicts of interest

There are no conflicts to declare.

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