Efficient Blue Phosphorescence in Gold(I)-Acetylide Functionalized Coinage Metal Bis(amidinate) Complexes

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Abstract: The synthesis of linear symmetric ethynyl- and acetylide-amidinates of the coinage metals is presented. Starting with the desilylation of the complexes \([\text{Me}_3\text{SiCC(NDipp)}_2\text{M}_2]\) (Dipp = 2,6-diisopropylphenyl) \((\text{M} = \text{Cu, Au})\) it is demonstrated that this compound class is suitable to serve as a versatile metalloligand. Deprotonation with n-butyllithium and subsequent salt metathesis reactions yield symmetric tetranuclear gold(I) acetylide complexes of the form \([\{(\text{PPh}_3)\text{AuCC(NDipp)}_2\}_2\text{M}_2]\) \((\text{M} = \text{Cu, Au})\). The corresponding Ag complex \([\{(\text{PPh}_3)\text{AuCC(NDipp)}_2\}_2\text{Ag}_2]\) was obtained by a different route via metal rearrangement. All compounds show bright blue or blue-green microsecond long phosphorescence in the solid state, hence their photophysical properties were thoroughly investigated in a temperature range of 20–295 K. Emission quantum yields of up to 41% at room temperature were determined. Furthermore, similar emissions with quantum yields of 15% were observed for the two most brightly luminescent complexes in thf solution.

Introduction

In the last decades, a great number of luminescent coinage metal complexes have been reported in the literature. The ligands applied for the synthesis of these complexes varied broadly from, for example, differently substituted phosphines, NHCs, to multidentate PN-type ligands. Another ligand group are ethynyl or acetylide containing compounds, which are highly attractive for several reasons: 1) their ability to realize \(\sigma\)- and \(\pi\)-coordination modes allows for the synthesis of multinuclear complexes with short metal–metal contacts;

2) their multidentate character and linearity are key factors for the formation of supramolecular structures;

3) the \(\text{C}^1\text{C}\) moiety is often involved in photophysical processes in organometallic compounds. The latter is related to intraligand or metal-to-ligand charge transfer to the \(\pi^*(\text{C}^1\text{C})\) orbitals upon photoexcitation, from which radiative relaxation can also occur. Concerning photoluminescence (PL) among coinage metal complexes, the phenomenon of metallophilicity often plays a significant role as well. Metallophilicity describes the tendency of metal cations to form very short metal–metal contacts, which occur mostly in the solid state. The metallophilic interactions may have a strong impact on the photophysical properties, however, the relations between the structure and the resulting properties are still a matter of investigations.

Dependent on the ligand, metallophilic interactions are classified into unsupported, semi-supported and fully supported ones. For example, monovalent coinage metal amidinate complexes are known to form dimeric species \("\text{L}_2\text{M}_2\)"\), exhibiting fully supported metal–metal contacts that go along with metallophilic interactions. Therefore, those complexes may show distinct PL depending on the amidinate substituents. Our group recently reported on the alkynyl substituted gold(I) amidinate \([\text{Me}_3\text{SiCC(NDipp)}_2\text{Au}_2]\) with an inherent aurophilic interaction indicated by an extraordinarily short gold–gold contact (Au1-Au1’ 2.7009(11) \(\text{Å}\)). Furthermore, this complex shows distinct PL in the solid state, as well as in tetrahydroduranne (thf). It has also been demonstrated that further reaction with \([\text{AuCl(tht)})\) (tht – tetrahydrothiophene) results in a desilylation, yielding an octanuclear species.

Herein, we describe a reaction protocol for the controlled desilylation of the \(-\text{C}^1\text{C}^1\text{SiMe}_3\) group to obtain the versatile luminescent building block \([\text{HC}^1\text{C}^1\text{CC(NDipp)}_2\text{Au}_2]\) for subse-
sequent reactions such as lithiation with a strong base. The efficiency of this protocol is ascribed to the kinetic stabilization of the metal core by the bulky 2,6-diisopropylphenyl (Dipp) substituents. The controlled metalation of the ethynyl end group allows for the preparation of multinuclear complexes, which represent potential building blocks for the synthesis of heterometallic metallopolymers. In addition, the access is extended to the other coinage metals copper and silver. In this manner, small molecules with promising photophysical characteristics can be further derivatized. Subsequently, the auration of the ethynyl moiety by a one-pot reaction with KHMDS and [AuCl(PPh3)] was tested, but it was not possible to obtain the desired product. Therefore, a stepwise approach was chosen. In the first step, complex 1 was reacted with two equivalents of n-butyllithium at −78 °C to yield the lithium acetylide [(thf)3LiCC(NDipp)2]Au2] (2) (Scheme 1). Interestingly, since compound 1 shows a cyan colored luminescence in solution, the deprotonation towards complex 2 can be easily monitored by the immediate change of the luminescence to azure after addition of the base, using a conventional UV-lamp (λexc ≈ 365 nm). Full deprotonation of compound 1 was proved by the 1H NMR spectrum of 2, in which, in contrast to the spectrum of 1, no resonances for the terminal alkyne protons were detected at δ = 2.64 ppm. In the 13C(1H) NMR spectrum, the resonances for the alkyne moieties at δ = 89.1 and 75.8 ppm are significantly shifted to lower ppm compared to the analogue resonances of the starting compound (δ = 109 and 97.3 ppm). Furthermore, two characteristic bands for the stretching modes of the terminal alkyne at ν(C–H) = 3276 cm⁻¹ and ν(C≡C) = 2114 cm⁻¹ arise in the IR spectrum of 1. The latter is also observed as the signal with the highest intensity in the Raman spectrum of 1 at ν(C≡C) = 2113 cm⁻¹. This mode is shifted by 53 cm⁻¹ to smaller wavenumbers in comparison to the parent compound. The purity of compound 1 was confirmed via elemental analysis, as well as HR-ESI mass spectrometry. Thereby, the compound was detected in form of its molecule peak [M]+ at m/z = 1168.4932.

The structural integrity of compound 1 in the solid state was finally verified by single crystal X-ray analysis (Figure 1). All bond lengths and angles are similar to the starting material, for example, the gold–gold distance (Au1–Au1)̃ = 2.7050(11) Å as well as the carbon–carbon triple bond (C2–C3 1.182(5) Å) are nearly identical to the observed values for [(Me3SiCC≡CC(NDipp)2]Au2] (Au1–Au1 = 2.7009(11) Å and C2–C3 = 1.184(6) Å). Compound 1 is a versatile precursor for the synthesis of multinuclear metal complexes because the terminal alkyne function can be further derivatized.

Results and Discussion

As starting point, the previously published compound [(Me3SiCC≡CC(NDipp)2]Au2] was used, which was desilylated with caesium fluoride in a 2:1 stoichiometric mixture of absolute thf and methanol to yield the desired complex [(HC≡CC(NDipp)2]Au2] (Scheme 1). In almost quantitative yield (Scheme 1).

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Quite unusually, no agglomeration of the lithium acetylide moieties is observed. In fact, there is no report of a crystallographically investigated compound with an isolated end-on Li–C coordination, so far. The C2-C3-Li1 angle of 175.6(7)° is close to the ideal value of 180° for the end-on coordination. Nevertheless, the C3-Li1 distance of 2.06(2) Å is comparable to other lithium acetyrides.\[14\]

The structural parameters of the central gold-containing stretched hexagonal motif show no significant deviations in comparison to the molecular structure of compound 1, for example, the gold–gold distance is in the same range (Au1-Au1' 2.6837(5) Å). However, the carbon–carbon triple bond length C2-C3 of 1.217 cm is slightly shorter than the determined values for the starting material 1.

For the synthesis of the desired coinage metal functionalized compounds, the lithium acetylide 2 was reacted in salt metathesis reactions with coinage metal halides. First, the in situ generated complex 2 was reacted with two equivalents of chloro(triphenylphosphine)silver(I) to give the tetranuclear gold complex [(PPh3)AuCC(NDipp)2]2Ag2 (3) (Scheme 1). Recrystallization from dichloromethane/n-pentane resulted in a colorless microcrystalline solid. The 1H NMR spectrum of 3 is comparable to the parent compound, except that the proton resonances of the triphenylphosphine moieties are observed as a separate multiplet in the range of δ = 7.52–7.34 ppm. In comparison to compound 2, the alkylene carbon resonances in the 13C(1H) NMR spectrum are detected at higher ppm values, i.e. at δ = 155.3 and 123.7 ppm. Both resonances are detected as doublets with coupling constants of 1JCC = 145.9 Hz (Au–C) and 3JCP = 32.2 Hz (Au–CP), respectively. The phosphine resonance in the 31P(1H) NMR spectrum appears at δ = 41.2 ppm and therefore at the usual chemical shift for Au(PPh3) substituted alkynes.\[19\] In the IR spectrum of compound 3, a broad weak band is observed for the asymmetric combination of the Cs–C stretching at ν(Cs–C) = 2109 cm\(^{-1}\). The analogue symmetric combination is observed in the Raman spectrum at ν(Cs–C) = 2117 cm\(^{-1}\) as the band with the highest intensity. Finally, single crystal X-ray diffraction revealed that the initially desired tetranuclear gold complex 3 was obtained (Figure 3, left). In analogy to the previously discussed compounds, the structural parameters for the central motif show no significant changes.

Consecutively, we felt challenged to synthesize the acetylide of the lighter congeners of gold. Therefore, two equivalents of chloro(triphenylphosphine)silver(I) were added to in situ generated compound 2. After a similar workup as for the tetranuclear gold complex 3, to our surprise, [(PPh3)AuCC(NDipp)2]2Ag2 (4) was obtained as a white microcrystalline solid. Single crystal X-ray diffraction of 4 revealed a metal rearrangement, hence the silver ions of the metal precursor displaced the gold atoms in the amidinate pocket of compound 2, while the gold ions are consecutively coordinated by the acetylide moieties and the phosphines (Figure 3, right).

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Molecular structure of compound 1 in the solid state. Hydrogen atoms, except for the terminal alkyne protons, are omitted for clarity. Structural parameters are given in the Supporting Information.

![Figure 2](https://example.com/figure2.png)

**Figure 2.** Molecular structure of compound 2 in the solid state. Hydrogen atoms are omitted for clarity. Structural parameters are given in the Supporting Information.
The metal–metal contact (Ag1-Ag1’ 2.7042(5) Å) in compound 4 is nearly identical to the contact in compound 3 (Au1-Au1’ 2.7040(5) Å). The Au1-P1 (2.2767(10) Å) and Au1-C3 (2.006(4) Å) distances, as well as the C3-Au1-P1 (176.26(11)°) angles are also in the same range as for complex 3.

The 1H, 13C{1H}, as well as the 31P{1H} NMR spectra of the silver complex are very similar to the spectra of compound 3, which partly is a result of the metal rearrangement. The stretching mode of the alkyne moieties is also quite similar and appears at ν(C=C) = 2109 cm⁻¹ in the IR and Raman spectra of 4.

In order to complete the series of the tetranuclear coinage metal complexes, we synthesized the digold–dicopper complex [((PPh₃)Au=C=C(NDipp))₂]Cu₂(7), with two copper atoms in the central amidinate pocket (Scheme 1). Since a metal rearrangement of copper and gold, as observed for the access of 4, is not possible, a route similar to the synthesis of 3 was chosen. Therefore, the dicopper complex [(Me₅SiC≡CC(NDipp))₂]Cu₂ (5) was synthesized in a similar fashion as [(Me₅SiC≡CC(NDipp))₂]Au₂ (6), and was consecutively desilylated to yield [(HC≡CC(NDipp))₂]Cu₂ (6). Due to the occurrence of side reactions during the deprotection with CsF, the desilylation was realized using potassium carbonate to obtain compound 6 in nearly quantitative yield and high purity. In situ lithiation and subsequent metathesis reaction with two equivalents of [AuCl(PPh₃)] yielded compound 7 in a total yield of 49% over three steps. All copper containing compounds (5–7) have been fully characterized including HR-ESI mass spectrometry and single crystal X-ray diffraction.

The NMR spectra for compound 5 are in full agreement with the measured spectra for [(Me₅SiC≡CC(NDipp))₂]Au₂, for example, the alkyne moiety in the 1H NMR spectrum is detected at similar resonances (δ = 105.6 and 95.3 ppm (5) vs. δ = 109 and 97.3 ppm). Also in consistence, no ν(C≡C) vibration is observed in the IR spectrum, while it appears as the vibration mode with the highest intensity in the Raman spectrum of 5 at similar wavenumbers (ν(C≡C) = 2162 cm⁻¹ (5) vs. ν(C≡C) = 2167 cm⁻¹ in [(Me₅SiC≡CC(NDipp))₂]Au₂). The molecular structure of 5 in the solid state (Figure S39) shows, in agreement with the smaller ionic radius of the copper(I) ion, an even shorter metal–metal contact (Cu1-Cu2 2.4573(7) Å) than for [(Me₅SiC≡CC(NDipp))₂]Au₂ (Au1-Au1’ 2.7009(11) Å), which is in the usual range for cuprophilic interactions.[21]

In analogy to the desilylated gold(I) complex 1, a separate singlet resonance at δ(C₆D₆) = 1.79 ppm is detected in the 1H NMR spectrum of 6, which can be assigned to the terminal alkyne protons. In comparison to the resonance in the 1H NMR spectrum of 1 (δ(CDCl₃) = 2.64 ppm), it is significantly shifted to higher ppm values which likely results from the less polar solvent C₆D₆. In good agreement with the previously discussed results, a similar shift of the alkyne carbon atoms (δ = 87 and

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**Figure 3.** Molecular structures of compounds 3 (left), 4 (right), and 7 (below) in the solid state. Hydrogen atoms are omitted for clarity. Structural parameters are given in the Supporting Information.
74.8 ppm) is observed in the $^{13}$C($^1$H) NMR spectrum of 6 in comparison to the spectrum of 5.

Furthermore, the two characteristic bands for the stretching modes of the terminal alkyne at $\nu(C-H) = 3267$ cm$^{-1}$ and $\nu(C=C) = 2109$ cm$^{-1}$ arise in the IR spectrum of 6 as well.

Compared to 5, the band of the alkyne stretching mode $\nu(C=C)$ at 2107 cm$^{-1}$ in the Raman spectrum of 6 is shifted by about 55 cm$^{-1}$ to smaller wavenumbers. This is in agreement with the observations on the analogue gold(I) complexes (shift=$53$ cm$^{-1}$). A metal–metal contact (Cu1-Cu1 2.5319(12) Å), which is comparable to the value for 5, and 2.5319(12) /C138), which is comparable to the value for about 55 cm$^{-1}/C0$ (shift)

Greater central structural parameters in the molecular structure of 6 in the solid state solely show small deviations as well.

Moreover, the spectroscopic data for 7 is also in full agreement with the corresponding gold(I) and silver(I) complexes 3 and 4. Thus, the $^1$H NMR spectrum of 7 shows the characteristic separate multiplet in the range of $\delta=7.50–7.28$ ppm for the aromatic phosphate protons, while a singlet resonance is observed in the $^{13}$C($^1$H) NMR spectrum ($\delta=41.5$ ppm). This resonance is in the range of 3 and 4.

The molecular structure of compound 7 reveals a metal–metal contact of Cu1–Cu2 2.4717(12) Å, which is nearly the same as for complex 5. Thus, alterations at the terminal alkyne function do not significantly influence the central $M_3$ core.

Photoluminescence properties

Compounds 1–7 are either colorless or off-white solids. All of them show azure, bright blue or blue-green photoluminescence (PL) both at low and ambient temperatures, with a moderate rise of PL intensity by decreasing the temperature from 295 to 20 K (Figures 4 and 5, see also the SI). The emission spectra show a similar vibronic structure, particularly distinct at low temperatures and for complex 4. In consistency with colorlessness, the PL excitation (PLE) spectra of 1–7 show the absorption onset below approximately 450 nm. The PL decay of 1–7 (upon nsec-pulsed excitation with a nitrogen laser at 337 nm) occurs on a microsecond time scale, thus clearly indicating that the PL is phosphorescence (Figure S53). The PL lifetimes decrease moderately upon raising the temperature to 295 K, thus roughly correlating with the decrease of PL intensity due to enhanced nonradiative relaxation. Figure 4 combines PL spectra of the dinuclear metal complexes 1, 5, and 6. The blue-green emission of the silylated copper-complex 5 is centered at approximately 520 nm and is similar to that of the gold-complex [Me$_3$SiC=CC(NDipp)$_2$]Au$_2$. In comparison, the PL emission and excitation spectra of the desilylated compounds 1 and 6 are blue-shifted by approximately 20 nm. In difference to 5, the reduction of PL intensity of 1 and 6 is more pronounced upon raising the temperature from 20 to 295 K. Correspondingly, PL quantum yields $\Phi$ (295 K) = 34, 16 and 10% were determined using an integrating sphere under ambient conditions for compounds 1, 5, and 6, respectively. There are also some differences in the PL decay. While it is monoeponential for 1 and 5, with lifetimes $\tau=56$ (24) and 220 (173) μsec at 20 K (295 K), respectively, the PL decay curve of 6 transforms from monoexponential at 20 K ($\tau=252$ μsec) to biexponential at ambient temperature (Figure S53).

Figure 5 shows the spectra of the tetranuclear complexes 2–4 and 7. All these complexes emit in the blue color region with the PL bands centered at approximately 450–490 nm. Hence, the PL of the tetranuclear metal complexes is blue-shifted in comparison to the dinuclear metal complexes 1, 5, and 6 by approx. 40 nm. At 20 K, a similar vibronic structure of the emissions, comparable to compounds 1, 5, and 6, is observed for 2, 3, and 7.

Likely due to a higher sample crystallinity, 4 exhibits particularly sharp vibronic features under cryogenic conditions, assigned to vibrational modes at approx. 1380 cm$^{-1}$ (probably a bisaminadate-based mode) and 2100 cm$^{-1}$ (C=C stretching) and their overtones (Figure S54). Upon raising the temperature from 20 to 295 K, the PL intensity decreases nearly linearly for

![Figure 4](image-url)

**Figure 4.** Left: normalized photoluminescence excitation (PLE) and emission (PL) spectra of the binuclear compounds 5, 1, and 6 at 20, 150, and 295 K. The PL/PLE spectra were excited/recorded at values between 350–407 nm, respectively (Table S3). Right: integrated PL intensities vs. temperature in the range of 20 to 295 K.

![Figure 5](image-url)

**Figure 5.** Left: normalized photoluminescence excitation (PLE) and emission (PL) spectra of the tetrametallic compounds 2, 3, 4, and 7 at 20, 150, and 295 K. The PL/PLE spectra were excited/recorded at values between 350–365/432–476 nm, respectively (Table S4). The PL spectral intensities of compounds 2 and 4 at 295 K were multiplied by a factor of 2 for clarity. Right: integrated PL intensities vs. temperature in the range of 20 to 295 K.
all tetrametallic complexes (Figure 5, right). The decrease is quite moderate for gold-gold and gold-copper compounds 3 and 7 and more pronounced for lithium-gold and gold-silver compounds 2 and 4. In line with this observation, the highest quantum yields $\eta(295 \text{ K}) = 40$ and 41% were found for 3 and 7, in comparison with 13 and 18% for 2 and 4, respectively, by using an integrating sphere and excitation at 350 nm (Table S4). The time scale of emission decay of the tetrametallic complexes, e.g., tens to hundreds of microseconds at 20 K is comparable to the binuclear compounds. However, the decay kinetics of the former is more complicated and follows biexponential curves both at low and ambient temperatures (Figure S53). The similarity of the PL properties of compounds 1–7 suggest common types of the emissive triplet excited states. These are likely mostly located on the Dipp-amidinate-alkynyl framework and moderately affected by variation of the coinage metals and coordination patterns across 1–7.

For instance, the structurally closely related bi- and tetrametallic gold complexes 1 and 3 demonstrate quite similar PL properties, indicating a moderate effect of the gold atoms linearly coordinated by the acetylide moieties. On the other hand, the metals in the central amidinate pocket apparently affect the triplet (non)radiative relaxation rates: as expected for more efficient spin–orbit coupling associated with the gold atoms, the PL decay of 1 and 3 is notably faster in comparison to their copper-containing counterpart structures 6 and 7 (Figure S53). To further determine and compare the specific colors emitted by compounds 1–7, CIE coordinate diagrams at 295 K (Figure 6) and 20 K (Figure S55) were prepared.[22] The bimetallic complexes 1, 5, and 6 clearly emit in the cyan–green color range, while the PL colors of the tetrametallic complexes 3 and 7 are further shifted to the blue color range. Remarkably azure blue colored PL is observed for the Li$_2$Au$_2$ and Ag$_2$Au$_2$ complexes 2 and 4, respectively. At cryogenic temperatures, the emission colors are only slightly shifted (Figure S55).

Since complexes 3 and 7 show the highest PL quantum yields in the solid state, we further investigated their PL properties in thf solution. The observed emission spectra are quite similar to those in the solid state (Figures S49–S51). The PLE spectra well correspond to the respective absorption spectra in solution and qualitatively correspond to the PLE spectra in the solid state. Note that in the latter spectra weak absorption features appear overemphasized due to non-validity of Lambert-Beer law for the optically “thick” solid sample preparations. Finally, PL quantum yields of 15% were measured for both complexes in thf solution.

Conclusions

A straightforward synthetic protocol towards a full series of tetrametallic coinage metal acetylide-amidinate complexes $[\{\text{PPPh}_3\text{AuCl}_{2}\text{CC(NDipp)}_2\text{M}_2\}]$ (M = Cu, Ag, Au) is presented. We demonstrated that the ethynyl compound $[\{\text{HCC(NDipp)}_2\text{M}_2\}]$ (M = Cu, Au) serves as a versatile metalloligand. The gold compound even undergoes ligand exchange reactions when treated with silver precursors. The obtained metal complexes exhibit a combination of a very short intrinsic fully supported metallophilic interaction and two gold(IIItriph- nylphosphine moieties which are linearly aligned towards each other.

All metal complexes emit azure, bright blue or blue-green phosphorescence with quantum yields of up to 41% in the solid state under ambient conditions. Furthermore, a similar and efficient PL with quantum yields of 15% was observed for two exemplary complexes when dissolved in thf. Similar properties have also been observed for the previously characterized alkynyl substituted gold(I) amidinate and its octanuclear derivative.[15] Accordingly, the efficient PL, both in the solid state and in solution, appears to be a characteristic feature of the dimeric coinage metal amidinate structural motif.

Experimental Section

Experimental details are given in the Supporting Information (SI) which is available free of charge via the internet. The Supporting Information includes Materials, characterization methods, experimental procedures, full spectroscopic data for all new compounds, copies of $^1$H, $^13$C($^1$H), $^31$P($^1$H), $^2$Li($^1$H) NMR, IR and Raman spectra, elemental analyses, crystallographic data, additional photoluminescence and UV/Vis spectra.

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Two plus two is four: The synthesis of linear symmetric ethynyl- and acetylide-amidinates of the coinage metals is presented. Their photophysical properties were thoroughly investigated in a temperature range of 20–295 K. Short lifetimes and quantum yields of up to 41% were determined. Furthermore, the complexes show pronounced PL in THF solution as well.