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TECHNICAL NOTE

Criminalistics

Clarifying the complex chemistry of cobalt(II) thiocyanate-based tests for cocaine using single-crystal X-ray diffraction and spectroscopic techniques

Raychelle Burks PhD^{1,2}  | Lars Öhrström PhD² | Francoise M. Amombo Noa PhD^{2,3} 

¹Department of Chemistry, American University, Washington, District of Columbia, USA

²Department of Chemistry and Chemical Engineering, Chalmers University of Technology, Gothenburg, Sweden

³AstraZeneca, Mölndal, Sweden

Correspondence

Raychelle Burks, Department of Chemistry, American University, 4400 Massachusetts Ave NW, Washington, DC 20016, USA.

Email: burks@american.edu

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Abstract

Cobalt(II) thiocyanate-based tests are routinely used to screen cocaine products, with the formation of a blue species interpreted as a positive response. An array of other organic bases has been identified as false positives – including well-documented cocaine product adulterant lidocaine and its salt. False positives prompt continued test development, though improvements are hindered by unresolved product structures and reaction pathways. Toward greater clarity, cobalt(II) thiocyanate reactions with cocaine hydrochloride, along with lidocaine and its salt, were investigated using multiple analytical techniques. Reactions involving cocaine hydrochloride yielded glassy, amorphous blue material while reactions of lidocaine hydrochloride monohydrate produced larger, needle-like crystals whose structure was determined via single-crystal X-ray diffraction to be an ion pair (Hlidocaine^+)₂[Co(SCN)₄]²⁻·H₂O. While the blue precipitate isolated from reactions involving cocaine hydrochloride was unsuitable for crystallographic structure determination, comparative ultraviolet–visible, attenuated total reflectance infrared, and Raman spectroscopic analysis – along with elemental analysis – supports that this solid is comprised of a comparable ion pair (Hcocaine^+)₂[Co(SCN)₄]²⁻. Pink crystals isolated from lidocaine reaction vessels were identified as coordination compounds *cis*-[CoL₂(SCN)₂] and *trans*-[CoL₂(SCN)₂] where L = lidocaine, while pink crystals from both cocaine hydrochloride and lidocaine hydrochloride monohydrate reaction vessels were the coordination polymer *trans*-[Co(H₂O)₂(SCN)₂]·H₂O. The results presented herein enable reaction optimization to favor a desired product, whether ion pair or coordination species.

KEYWORDS

attenuated total reflection infrared spectroscopy (ATR-IR) spectroscopy, cobalt thiocyanate, cocaine, color tests, lidocaine, presumptive tests, Raman spectroscopy, Scott's test, ultraviolet–visible (UV–vis) spectroscopy, X-ray diffraction

Highlights

- Product structures and reaction pathways of cobalt(II) thiocyanate-based tests are clarified.
- Hallmark blue products of cobalt(II) thiocyanate tests are ion pairs.
- Three pink products were isolated, each a unique coordination compound.

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1 | INTRODUCTION

The straightforward operational nature of many color tests may imply that their underlying chemistry is simplistic and exhaustively explored. In reality, color test reaction mechanisms can be quite complex with the elucidation of the chemical identity of their telltale-colored species an area of ongoing research activity [1–3]. The reactions of cobalt(II) thiocyanate typify the complexity of color tests and metal ion coordination chemistry. Cobalt(II) thiocyanate solutions have a long history as a colorimetric reagent for the determination of alkaloids and their salts, along with the wider category of organic nitrogen-containing bases (hereafter ‘organic bases’) and their salts [4–10]. Arguably, the most widespread forensic use of cobalt(II) thiocyanate reagents is for screening suspected samples of cocaine (base or salt form), with these controlled substances indicated by the formation of a blue species upon exposure of a suspect material with test reagent(s).

Challengingly, numerous organic bases produce blue reaction species when exposed to cobalt(II) thiocyanate solutions such as lidocaine, procaine, diphenhydramine, ephedrine, phencyclidine, fentanyl, α -pyrrolidinopentiphenone, and/or their salts [11–17]. Structures of cocaine and lidocaine are shown in Figure 1. Lidocaine is showcased herein as this local anesthetic is both a long-noted false-positive and oft-encountered adulterant in cocaine products [3, 18, 19]. The litany of false positives for cobalt(II) thiocyanate-based tests has prompted numerous revisions in pursuit of increased selectivity, while retaining ease-of-use and screening speed. Updates to cobalt(II) thiocyanate tests range from the chemical (e.g., Scott's test) to the computational (e.g., incorporating multivariate image analysis) [3, 5]. Stymying continued test improvement efforts are unsettled product structures and cobalt(II) thiocyanate reaction mechanisms [1, 3, 20]. Ahead of a cobalt(II) thiocyanate test solution being introduced to organic base samples, the complexity of the test solution is worthy of note and continued exploration.

Solid $[\text{Co}(\text{SCN})_2]$ is brown-yellow in color and a coordination polymer with a bridging Co-SCN motif [21, 22]. The dissolution of

brown-yellow $[\text{Co}(\text{SCN})_2]$ in both the classic test (2% w/v cobalt (II) thiocyanate in distilled water) and in Scott's test reagent #1 (1% w/v cobalt (II) thiocyanate in a 1:1 v/v solution of glycerine and 10% v/v acetic acid) results in a pink solution [11, 23]. This pink color indicates that the predominant species formed are octahedral (six-coordinated) cobalt complexes of the type $[\text{Co}(\text{H}_2\text{O})_n(\text{SCN})_m]^{2-m}$ with $n+m=6$, and a critical evaluation of the corresponding stability constants has been published [24, 25]. Charged complexes such as $[\text{Co}(\text{H}_2\text{O})_2(\text{SCN})_4]^{2-}$ can be balanced by species such as $[\text{Co}(\text{H}_2\text{O})_6]^{2+}$ and compounds with multiple cobalt centers (i.e., polynuclear species) [26]. More violet-hued species and solutions indicate the presence of low concentrations of blue tetrahedral species such as $[\text{Co}(\text{SCN})_4]^{2-}$, or similar tetrahedral species, along with the aforementioned pink octahedral complexes [2, 20, 27–29]. Like other transition metals, the color of cobalt(II) complexes may indicate the number and geometry of surrounding ligands, and for Co(II), the difference between tetrahedral and octahedral species is especially striking (Figure 1) [2, 30]. Upon exposure of pink cobalt(II) test solutions to select organic bases, blue species form that either precipitate and/or color the bulk solution depending on solvent conditions. The structure of these blue species remains unresolved or unconfirmed for a variety of analytes, including cocaine and its salt, though key work has provided structural insights.

A variety of researchers have posited the structure of the telltale blue species as an ion pair with the general formula $(\text{HB}^+)_2[\text{Co}(\text{SCN})_4]^{2-}$, where protonated organic base (HB^+) is paired with an anionic, tetrahedral Co(II) complex [2, 4, 5, 7, 9, 20, 27]. While the International Union of Pure and Applied Chemistry's *Nomenclature of Inorganic Chemistry* [31] directs that ion charges not be denoted in overall neutral species, we have included charges to emphasize ion pairs. Ion pairs such as $(\text{HB}^+)_2[\text{Co}(\text{SCN})_4]^{2-}$ are stabilized by supramolecular interactions, with such interactions finding use in organic synthesis [32]. While a host of such ion pair complexes are insoluble in aqueous media, many are soluble in a variety of organic solvents [10, 33, 34]. Such differential solubility has made ion-pair complex formation a popular extraction strategy [35–37], such as in Scott's test. Toward clarifying the blue species resulting from Scott's and other cobalt(II) thiocyanate-based tests, Oliver et al. recently worked with the citrate salt of the organic base diethylcarbamazine, publishing the crystal structure of an ion pair $(\text{Hdiethylcarbamazine}^+)_2[\text{Co}(\text{SCN})_4]^{2-}$ resulting from a 1:2 mixture of the citrate salt and $\text{K}_2[\text{Co}(\text{SCN})_4]$ [38]. Beyond tetrahedral coordination complexes and an ion pair, previous researchers propounded the hallmark blue species resulting from Co(II) reactions with organic base – specifically cocaine – are octahedral, six-coordinate complexes with one or two bis-chelating cocaine ligands [1, 17, 39].

Greater structural clarity of reaction species and reaction mechanisms is ‘essential’ for the continued use and improvement of color tests [40]. These tests are a vital part of the presumptive-to-confirmatory workflow for identifying seized drug material, even in our era of portable instruments [15]. Toward the continued advancement of cobalt(II) thiocyanate-based tests, we evaluated the reaction products resulting from classic and Scott's tests screening of cocaine

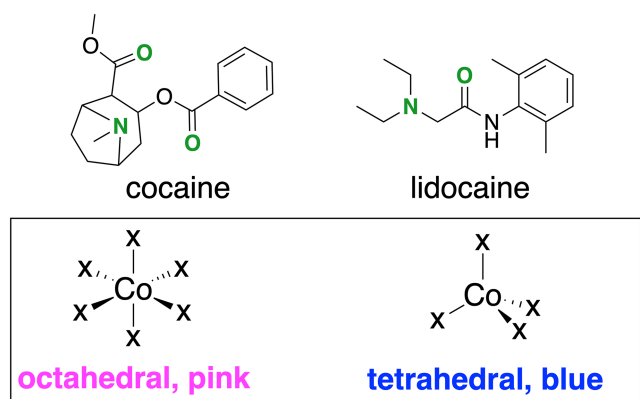


FIGURE 1 Top: Structures of cocaine and lidocaine. Bottom: cobalt(II) tetrahedral and octahedral ligand geometries with corresponding color of appearance.

hydrochloride, lidocaine hydrochloride monohydrate, and lidocaine using single-crystal X-ray diffraction along with ultraviolet-visible (UV-vis), attenuated total reflectance infrared (ATR-IR), and Raman spectroscopy with select material subjected to elemental analysis. Lidocaine and its salt were included due to this drug's aforementioned role as cobalt(II) thiocyanate-based tests false positive and cocaine product adulterant. Probing the reaction species further expands our knowledge of these widely employed tests, enabling their continued refinement and general color test innovation.

2 | MATERIALS AND METHODS

2.1 | Materials

Cobalt(II) thiocyanate, cocaine hydrochloride, lidocaine hydrochloride monohydrate, lidocaine, acetic acid, glycerine, concentrated hydrochloric acid, chloroform, and ethyl acetate were purchased from various chemical supply companies such as Sigma-Aldrich (St. Louis, MO) at technical grade or better and were used as received without further purification. Distilled water was used for all aqueous solutions. The use of standard laboratory equipment and glassware is described in the subsequent subsections.

2.2 | Cobalt (II) thiocyanate testing

Classic cobalt (II) thiocyanate test reagent and Scott's test reagents were prepared and administered as detailed in [11, 23], respectively. The former reagent is a 2% w/v cobalt (II) thiocyanate in distilled water and is hereafter referred to as 'classic test'. Scott's test is a trio of reagents (#1–#3) prepared as follows: (#1) 1% w/v cobalt (II) thiocyanate in a 1:1 v/v solution of glycerine and 10% v/v acetic acid, (#2) concentrated HCl, and (#3) chloroform. All presumptive testing and reagent storage was in a temperature-controlled indoor laboratory (~72°F or 22°C), being mindful of observed temperature effects on the sensitivity of cobalt (II) thiocyanate tests [20, 41]. Care was taken with regard to the w/v ratio of analyte to reagents, given the known complications excessive analyte amounts relative to test reagents can elicit [13]. Similarly, the ratio of Scott's test reagent #1 to #2 was carefully monitored as excessive addition of HCl can result in the formation of a blue species that is likely tetrachlorocobaltate (II), CoCl_4^{2-} , which is soluble in polar solvents [33, 42–46]. Analyte and reagent amounts were scaled in tandem for larger experimental yields as done in reference [3]. Samples of cocaine hydrochloride, lidocaine hydrochloride monohydrate, and lidocaine were subjected to both classic and Scott's test. For the salts, precipitates were not anticipated for Scott's test due to the enhanced solubility of reaction species in test reagents. The water insolubility of lidocaine [47] was expected to impact its reactivity in all reagent solutions. A total of thirty-six tests were run, and all resulting solutions were evaluated visually, including with the assistance of Zeiss Stemi SV6 microscope,

for precipitate that could be evaluated using either single-crystal or powder x-ray diffraction.

2.3 | Single-crystal X-ray diffraction

All crystallography work was done at the Chalmers Materials Analysis Laboratory (CMAL). Crystals obtained from test solutions – or grown in organic solvents like chloroform as specified – were selected and mounted on a nylon loop on an XtaLAB Synergy R, HyPix diffractometer. The crystal was kept at a steady $T = 100.0(7)\text{K}$ during data collection. Data reduction, scaling, and absorption corrections were performed using CrysAlisPro (Rigaku, V1.171.41.119a, 2021). The structures were solved with the ShelXT [48] structure solution program using the Intrinsic Phasing solution method and by using Olex2 [49] as the graphical interface. The model was refined with version 2016/6 of ShelXL 2016/6 using Least Squares minimization. Further details can be found in the supplementary material, and the CCDC entries 2235446–2235447 and 2235449–2235450 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +441,223 336,033. Structure searches discussed within were conducted among the more than 1 million structures in the Cambridge Structural Database (CSD) version 5.44, April 2023 [50].

2.4 | UV-vis, IR, and Raman spectroscopy

For spectra collection, cocaine hydrochloride and lidocaine hydrochloride monohydrate samples that provided positive results upon addition of the classic test reagent were used. For UV-vis analysis, ethyl acetate was added directly to selected positive test vials as in [51], with the resulting blue solution transferred to a new vial where additional ethyl acetate was added to generate suitably dilute solutions. Solutions were passed through 0.45 μm polytetrafluoroethylene (PTFE) membrane filters prior to UV-vis analysis using a Varian Cary® 50 UV-Vis spectrophotometer.

All ATR-IR and Raman spectroscopic work was done at CMAL. For ATR-IR and Raman analysis, blue solid material from representative classic test reaction vessels of cocaine hydrochloride and lidocaine hydrochloride was transferred to glass slides for solvent evaporation in a hood with no additional sample preparation prior to spectra collection. ATR-IR spectra were collected using a Bruker VERTEX 70v with ATR accessory, while Raman spectra were collected using an Oxford Instruments' alpha300 Raman Imaging Microscope with 10x, 50x, and 100x objective magnifications. Samples were measured using a 532nm laser with accompanying 600 grooves/millimeter grating was employed. To maximize Raman peak clarity and intensity while maintaining precipitate integrity, the

laser power, accumulations, and measurement time were varied for each sample analyzed.

2.5 | Elemental analysis

Elemental analysis (C, N, and S) was conducted at CMAL using an Elementar vario MICRO cube instrument. Blue solid material from a representative classic test reaction vessel of cocaine hydrochloride was transferred to a separate container and dissolved in chloroform. This solvent was allowed to evaporate and the resulting blue solid was wrapped in a small square of tin foil prior to placement in the instrument. All estimated element percentages were determined using instrument software and subsequently compared to theoretical elemental compositions using the analysis function in the ChemDraw software (version 20.1.0.112, PerkinElmer Informatics, 2021).

3 | RESULTS AND DISCUSSION

Upon screening of cocaine hydrochloride and lidocaine hydrochloride monohydrate samples via the classic test, blue precipitate was readily observed. As anticipated for samples subjected to Scott's test, no collectable blue precipitate was observed, though expected solution color changes were noted. Pink precipitates were isolated from reaction vessels containing cocaine hydrochloride, lidocaine hydrochloride, and lidocaine samples screened by the classic test. In the sections directly following, evaluation of said blue and pink precipitates are discussed.

3.1 | Blue precipitates

Markedly different blue precipitates were observed upon exposure of cocaine hydrochloride and lidocaine hydrochloride monohydrate

to the classic test reagent. Images of representative precipitates are inserted in Figures 2 and 3. Cocaine hydrochloride reactions yielded glassy, amorphous blue material unsuitable for single-crystal X-ray diffraction. Crystallization efforts with either chloroform or ethyl acetate resulted in the same glassy, amorphous solid as dissolved in said solvents. Lidocaine hydrochloride monohydrate reactions yielded larger needle-like blue crystals whose structure was straightforwardly elucidated by single-crystal X-ray diffraction to be the ion pair $(HL^+)_2[Co(SCN)_4]^{2-}(H_2O)$ where HL^+ is the cationic form of lidocaine referred to as lidocainium (Figure 4). Our low-temperature structure determination of $(HL^+)_2[Co(SCN)_4]^{2-}(H_2O)$ has considerably higher precision than an earlier room temperature structure of the ion pair, CSD code YEPHIK [52]. A space-filling drawing and a plot of the asymmetric unit are available in the Supplemental Information (SI), Figures S1 and S2. As illustrated in Figure 4, adjacent ion pairs interact through neighboring lidocainium ions via double hydrogen bonds between their $C=O \cdots H-N_{amine}$, forming a well-known supramolecular heterosynthon motif denoted R2,2(10) as ten atoms are linked in a supramolecular ring [53]. This notable interaction is not a feature of lidocaine monohydrate hydrochloride structures with codes LIDOCN and LIDOCN0 in the CSD. While the formation of the R2,2(10) motif is available to lidocainium, the cationic form of cocaine (cocainium) is unable to engage in such hydrogen bonding which may contribute to the more disordered nature of precipitate formed upon classic test screening of cocaine hydrochloride.

Of the more than one million structures in the CSD [50], seven are cocaine or cocainium featuring compounds being either the free base or ion pairs. Three ion pairs are of particular interest here as each is associated with forensic applications. The long-used standard microcrystal test for cocaine featuring a 5% gold chloride ($HAuCl_4$) that results in the asymmetric ion pair $(Hcocaine^+)[AuCl_4]^-$, CSD code SETLOT, with the chloride ions arranged about Au(III) in a distorted square planar geometry [54, 55]. Laussmann et al. prepared the green appearing $(Hcocaine^+)_2[Cu(NCS)_4]$, CSD code HAGSEQ, with thiocyanate groups arranged in a distorted square

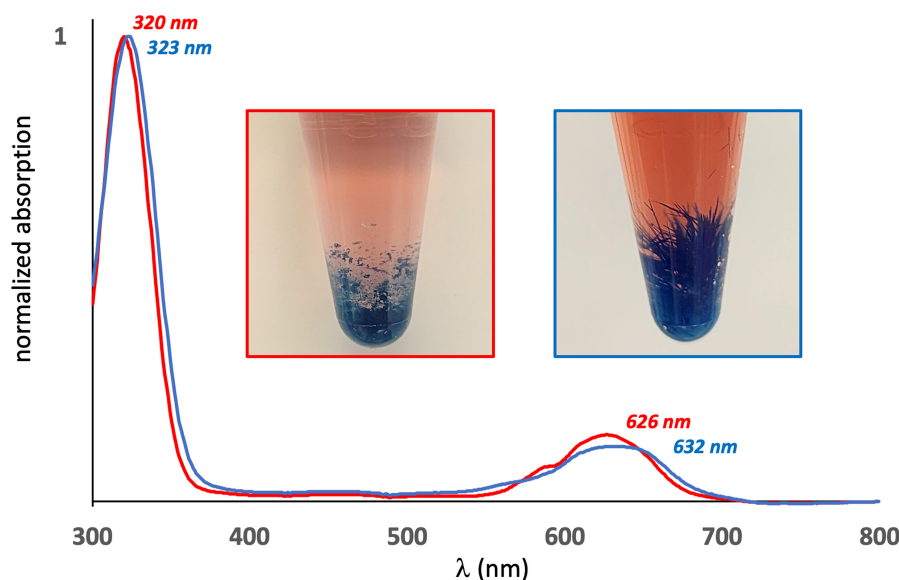
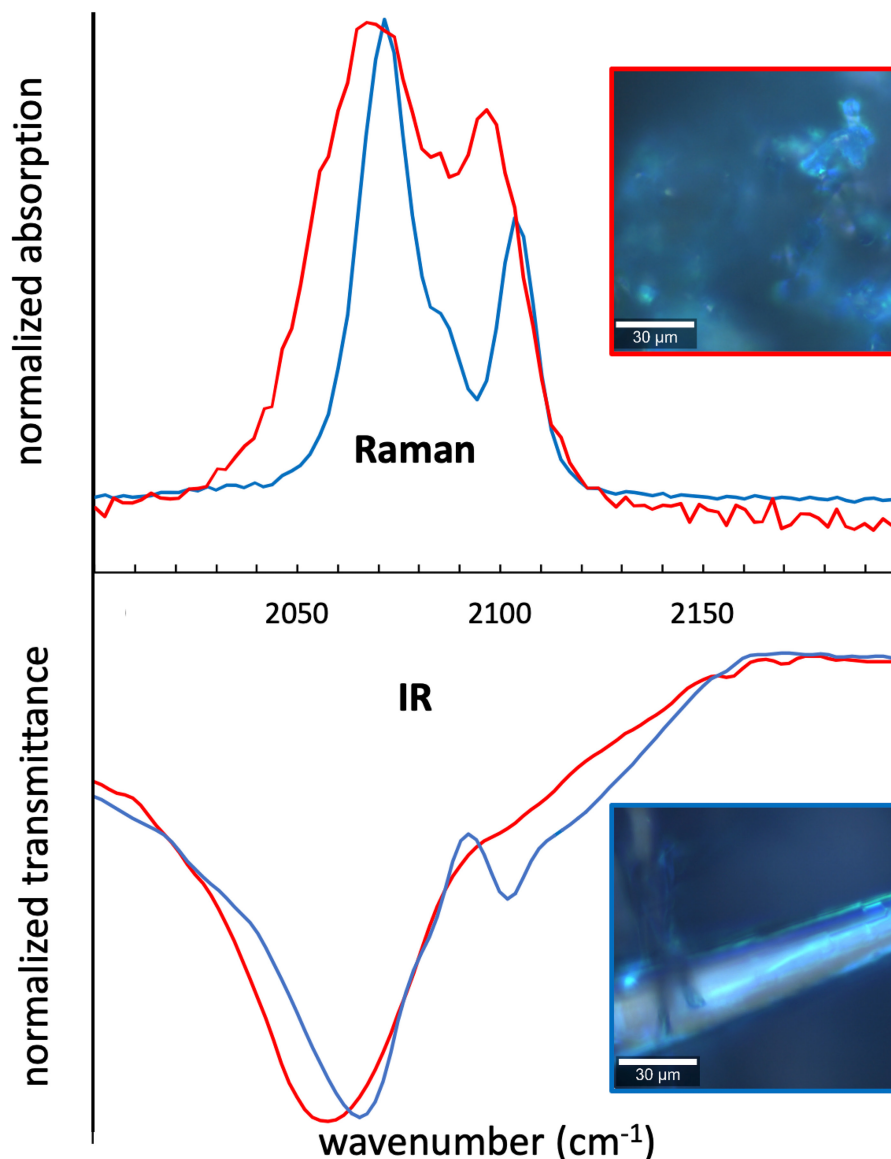


FIGURE 2 UV-vis spectra of blue precipitate retrieved from representative cocaine hydrochloride (—) and lidocaine monohydrate hydrochloride (—) reaction vessels dissolved in ethyl acetate. Image inserts of sampled reaction vessels show the markedly different appearance of each product.

FIGURE 3 Distinctive ATR IR and Raman $\text{C}\equiv\text{N}$ stretching band region around 2100cm^{-1} of blue precipitate retrieved from representative cocaine hydrochloride (—) and lidocaine monohydrate hydrochloride (—) reaction vessels. Inserted images collected during Raman analysis provide magnified views of the precipitate resulting from reactions involving cocaine hydrochloride (—) versus lidocaine monohydrate hydrochloride (—).



planar geometry around Cu(II) [37]. The third complex of note is the yellow cocaine salt with Erdmann's anion, $\text{trans}[\text{Co}(\text{NH}_3)_2(\text{NO}_2)_4]^-$, CSD code OSEFIE, which features two cocainium ions and two cobalt(III) containing Erdmann's anions in the asymmetric unit [56]. The paucity of cocainium containing ion pairs in the CSD may, in some part, reflect the commonality of non-crystalline precipitates such as we observed for samples of cocaine hydrochloride exposed to the classic test. While our crystallization efforts employing either chloroform or ethyl acetate did not yield crystalline material, we recognize that other crystallization methods may do so.

UV-vis, ATR-IR, and Raman spectroscopic analysis of blue precipitates was performed to ascertain if a comparable ion pair comprised the glassy, amorphous blue solid produced by reactions involving cocaine hydrochloride. Representative UV-vis spectra of solubilized blue species resulting from classic test screening of lidocaine hydrochloride monohydrate and cocaine hydrochloride samples are shown in Figure 2. These spectra agree with those presented in [51], with the peaks around 320nm most likely $\pi\text{-}\pi^*$ transitions of the $\text{C}\equiv\text{N}$

group and broader peaks around 630nm more complicated, consisting of overlapping bands corresponding to d-d transitions [57]. Similar results were noted with chloroform (data not shown), though the enhanced solubility of all precipitates in ethyl acetate [51] resulted in spectra with higher signal to noise. Given the color of the precipitates and their resulting solutions, peaks within the red zone of the UV-vis regions are expected. As seen in Figure 2, similarities about between the UV-vis spectra of the studied blue reaction species, suggesting similar structures. IR and Raman analysis of blue precipitates provided further structural insights.

An overview of the vibrational spectroscopy of thiocyanate complexes can be found in [58]. Critical for the examination of such complexes are the distinctive $\text{C}\equiv\text{N}$ stretching bands around 2100cm^{-1} [37, 51, 58]. Figure 3 highlights this distinctive wavenumber area for ATR-IR and Raman spectra of representative blue precipitates resulting from classic test screening of lidocaine hydrochloride monohydrate and cocaine hydrochloride. Though Figure 3 image insets show the dissimilarity in blue precipitate

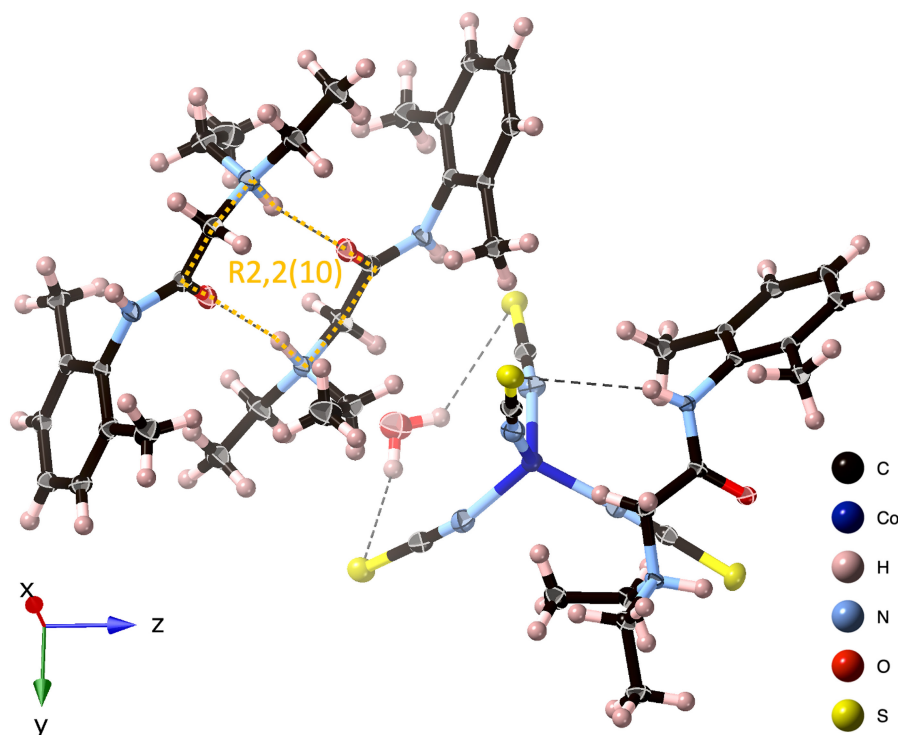


FIGURE 4 Crystal structure of $(\text{HL}^+)_2[\text{Co}(\text{SCN})_4]^{2-}(\text{H}_2\text{O})$ where HL^+ = lidocainium. The supramolecular heterosynthon motif R2,2(10) between neighboring lidocainium ions incorporated in adjacent ion pairs is highlighted in yellow. Also illustrated are the hydrogen bonds between (i) water and $[\text{Co}(\text{SCN})_4]^{2-}$ and (ii) amide hydrogen and $[\text{Co}(\text{SCN})_4]^{2-}$. Displacement ellipsoids shown at 50%. See also Figures S1 and S2.

appearance, similarities in nitrile stretching bands indicate similar ion pair complexes. Peak shifting around the 2100cm^{-1} reflects the sensitivity of nitrile groups to their local electrostatic and hydrogen-bonding environment [59]. Full ATR-IR and Raman spectra of collected blue precipitates, along with lidocaine hydrochloride monohydrate and cocaine hydrochloride are provided in SI Figures S3 and S4, respectively. Considering SI Figure S3, we note the bands associated with protonated amines around 2500cm^{-1} [60] are much weaker for both cobalt-containing compounds compared to either analyzed drugs' salt. Considering the elucidated structure for the lidocainium containing ion pair (Figure 4), the diminished protonated amine peaks are likely due to substantial changes in hydrogen bonding in $(\text{HL}^+)_2[\text{Co}(\text{SCN})_4]^{2-}\cdot\text{H}_2\text{O}$ compared to lidocaine monohydrate hydrochloride. For the blue precipitate produced by classic test screening of cocaine hydrochloride, the near vanishing of protonated amine peaks in the spectrum of the precipitate versus the salt also likely indicates changing hydrogen bond interactions.

A representative sample of blue precipitate produced by classic test screening of cocaine hydrochloride was further investigated via C, N, S elemental analysis as detailed in Section 2. It is important to note the potential inclusion of chloroform, CHCl_3 , into solubilized material even after subsequent solvent evaporation. The inclusion of CHCl_3 can make the element percentages deviate 1%–2% from the expected values. We found more than 17,000 CHCl_3 -incorporating compounds reported in the CSD. Elemental analysis of the specified sample revealed 49.37% C, 8.61% N, and 13.66% S with a sulfur-to-nitrogen ratio of 1.59. These percentages are consistent with an ion pair $(\text{Hcocaine}^+)_2[\text{Co}(\text{SCN})_4]^{2-}$ having 0.3 CHCl_3 molecules per Co, a species with calculated percentages of 49.16% C, 8.98% N, 13.70% S and a sulfur-to-nitrogen ratio of 1.53. In contrast, our experimentally determined C, N, and S percentages are incompatible

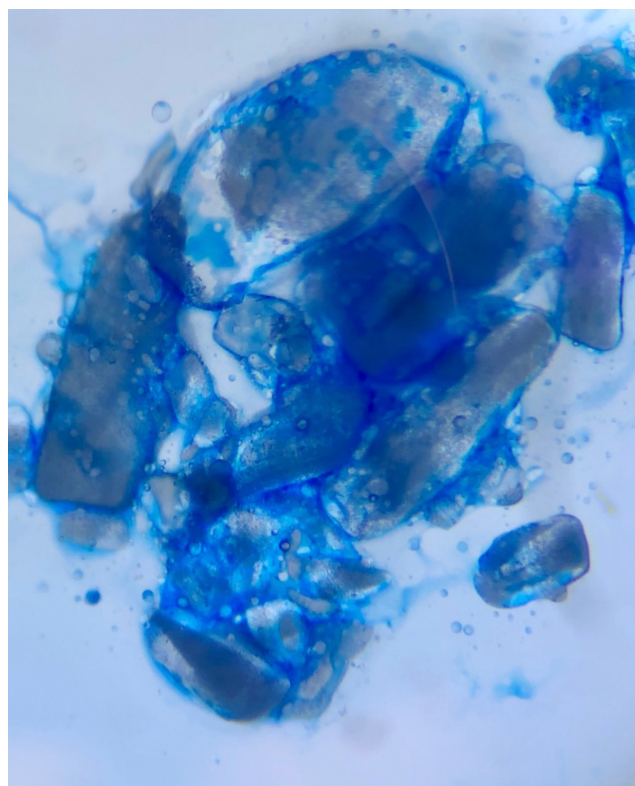


FIGURE 5 Image of a representative sample of lidocaine after classic test screening. The blue liquid-like layer is likely the lidocainium (HL^+) containing ion pair $(\text{HL}^+)_2[\text{Co}(\text{SCN})_4]^{2-}(\text{H}_2\text{O})$ localized at the drug-reagent interface.

with $[\text{Co}(\text{cocaine})_2(\text{SCN})_2]$ – where cocaine is non-chelating – which has calculated percentages of 55.31% C, 7.17% N, 8.20% S and a sulfur-to-nitrogen ratio of 1.14.

Lidocaine is noted as water-insoluble [47], which impacts reactivity within aqueous test reagents. For select lidocaine samples exposed to the classic test reagent and all lidocaine samples exposed to Scott's test reagent #1, the formation of a blue-colored layer upon solid lidocaine crystals was observed. Evaluation of these layered samples under magnification revealed a blue layer between the lidocaine solid surface and the bulk reagent solution. A representative image is shown in Figure 5. Based on analysis detailed in this subsection's previous paragraphs, we suggest the blue liquid-like layer is likely the lidocainium containing ion pair localized at the drug-reagent interface.

3.2 | Pink precipitates

Pink precipitates of a crystalline appearance were observed and collected from select classic test reaction vessels containing cocaine hydrochloride, lidocaine hydrochloride, and lidocaine. Three distinct pink-appearing compounds – designated herein as A, B, and C – were identified (Figure 6). The structure of A crystals, which were analyzed by single-crystal X-ray diffraction directly after classic test screening of lidocaine, was revealed to be $cis\text{-}[\text{CoL}_2(\text{SCN})_2]$ where L=lidocaine binding to cobalt in a bidentate fashion via tertiary amine N and the carbonyl O. Interestingly, the structure of crystals B that were retrieved approximately 8 weeks after classic test screening of lidocaine was determined to be $trans\text{-}[\text{CoL}_2(\text{SCN})_2]$. Based on a solid-state general rule [61] saying that intermolecular interactions are maximized when molecular entities come closer together, we posit the *trans* product to be the thermodynamic product as the calculated density from the crystal structure is 1.318 g/cm^3 , compared to the kinetic product *cis* with a calculated density of 1.255 g/cm^3 .

To the best of our knowledge, we are the first to report the structure of $trans\text{-}[\text{CoL}_2(\text{SCN})_2]$ where L=lidocaine, while room temperature structural elucidation of $cis\text{-}[\text{CoL}_2(\text{SCN})_2]$ was previously reported by Tabrizi et al. (CSD code XUTYUI) in their work probing the bio-activity of cobalt(II) and nickel(II) complexes of lidocaine [62]. There was an absence of comparable *cis* or *trans* compounds in reaction vessels containing cocaine hydrochloride or lidocaine monohydrate hydrochloride. For both salts, this absence may be explained by the tertiary amine site being unavailable for coordination with cobalt(II) due to protonation. Crystals C retrieved from cocaine hydrochloride and lidocaine hydrochloride monohydrate samples screened via the classic test were determined to be the coordination polymer $trans\text{-}[\text{Co}(\text{H}_2\text{O})_2(\text{SCN})_2]\cdot\text{H}_2\text{O}$. An earlier room temperature structure determination of $trans\text{-}[\text{Co}(\text{H}_2\text{O})_2(\text{SCN})_2]\cdot\text{H}_2\text{O}$ was detailed in [22].

3.3 | Structures and reactivity

Though the structures of lidocaine and cocaine are markedly different (Figure 1), they share a common feature of nonbonding electrons pairs available for Lewis acid-base interactions with Co(II) via tertiary amine and carbonyl groups [3, 19]. As discussed by de Jong et al., [19] these shared features may explain similar appearing blue species interpreted as positive cobalt(II) thiocyanate bases test results for an array of organic bases. Beyond reactive species with both tertiary amine and carbonyl groups, de Jong et al. noted that analytes observed to yield false positive results without a carbonyl (e.g., diphenhydramine) all have an electron-donating tertiary amine group. In this context, it is important to distinguish between (i) ion pairs formed by the combination of a protonated Lewis base such as lidocainium or cocainium and Co(II) containing anions like

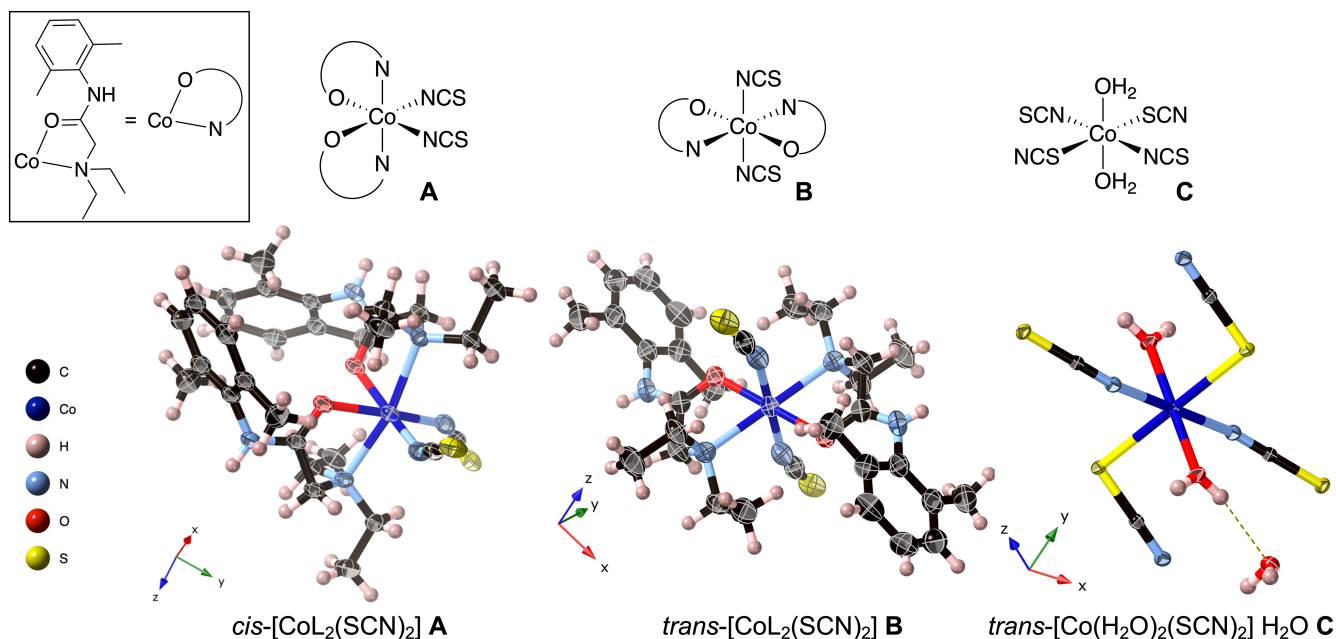


FIGURE 6 Crystal and line structures of $cis\text{-}[\text{CoL}_2(\text{SCN})_2]$ A, $trans\text{-}[\text{CoL}_2(\text{SCN})_2]$ B, and coordination polymer $trans\text{-}[\text{Co}(\text{H}_2\text{O})_2(\text{SCN})_2]\cdot\text{H}_2\text{O}$ C. For A and B, L=lidocaine. Displacement ellipsoids shown at 50%.

$[\text{Co}(\text{SCN})_4]^{2-}$ from (ii) compounds resulting from such bases forming a coordination bond to Co(II). In ion pairs, charged species are held together by electrostatic forces and intramolecular (supramolecular) interactions such as hydrogen bonds. Coordination compounds (complexes) result from bonds between a Lewis base and a metal ion such as Co(II).

For the blue ion pair $(\text{HL}^+)_2[\text{Co}(\text{SCN})_4]^{2-}$ where L = select reactive organic bases, it is the anion's tetrahedral geometry that is responsible for these tests' hallmark color (Figure 1). Considering the link between a blue appearance and tetrahedral geometry, two coordination compounds featuring organic bases as ligands are possibilities: $[\text{CoL}(\text{SCN})_2]$ and $[\text{CoL}_2(\text{SCN})_2]$. The first features a chelating base, while the second features a non-chelating base. We found no structures resembling $[\text{CoL}(\text{SCN})_2]$ in the CSD. The only structure resembling $[\text{CoL}_2(\text{SCN})_2]$ in the CSD is the green $[\text{Co}(4-(2,6\text{-Di-isopropylphenyl})\text{amino})-3\text{-methylpent-3-en-2-one})_2(\text{SCN})_2]$, code WUWUO, with a distorted tetrahedral geometry elucidated by Lugo and Richards [63] shown in SI Figure S5. Contrary to the fairly mild conditions of cobalt(II) thiocyanate-based tests, this compound was prepared in boiling toluene.

Moving to an octahedral coordination geometry about Co(II), we observed such species with the formation of the pink-appearing compounds *cis*- $[\text{CoL}_2(\text{SCN})_2]$ and *trans*- $[\text{CoL}_2(\text{SCN})_2]$ where L = lidocaine (Figure 6). While comparable species were not observed for cocaine hydrochloride and lidocaine hydrochloride monohydrate, reactions of these salts with classic test reagent did yield the pink coordination polymer *trans*- $[\text{Co}(\text{H}_2\text{O})_2(\text{SCN})_2]\cdot\text{H}_2\text{O}$ with an octahedral geometry (Figure 6). As stated previously, the seven cocaine or cocaineium featuring compounds in the CSD are either the free base or ion pairs. Thus, there are no structures of compounds with cocaine as a coordinating ligand deposited in the CSD.

The coordination compounds and ion pairs we observed for the evaluated cobalt(II) thiocyanate-based tests typify reactivity seen for Lewis acid metal ions. Specifically speaking of Co(II), it has fast ligand exchange kinetics so that in principle thermodynamic products should be obtained, but any type of coordination entity may be formed especially if trapped in a stable solid form [64]. In addition, which species will be thermodynamically most stable, and the corresponding reaction rates, will be critically dependent on the concentrations of all participating components and reaction conditions. There is a very facile conversion and co-existence of the tetrahedral $[\text{Co}(\text{SCN})_4]^{2-}$ (blue) and octahedral (pink) coordination geometries (Figure 1), in part explaining the performance of cobalt(II) thiocyanate based rapid color tests.

4 | CONCLUSION

A litany of organic bases (L) can participate in the Lewis acid–Lewis base interactions giving rise to ion pairs and coordination compounds observed in cobalt(II) thiocyanate reactions. Our work with cocaine hydrochloride, lidocaine hydrochloride monohydrate, and lidocaine greatly bolsters the conclusion that an ion pair of general

structure $(\text{HL})_2^+[\text{Co}(\text{SCN})_4]^{2-}$ is the origin of the hallmark blue color interpreted as a positive result for cobalt(II) thiocyanate-based tests. Single-crystal X-ray diffraction work identified $(\text{Hlidocaine}^+)_2([\text{Co}(\text{SCN})_4]^{2-})\cdot\text{H}_2\text{O}$ as the blue species produced for reactions involving lidocaine hydrochloride monohydrate, with the reasonable extension that this ion pair composes the blue layer around lidocaine crystals exposed to cobalt(II) thiocyanate-based reagents. While the glassy, amorphous blue precipitate resulting from reactions involving cocaine hydrochloride was unsuitable for crystallographic analysis, our comparative spectroscopic work supports its composition to be $(\text{Hcocaine}^+)_2[\text{Co}(\text{SCN})_4]^{2-}$. Co-existing with these blue ion pairs were pink coordination compounds, the structure of each determined by single-crystal X-ray diffraction. The coordination polymer *trans*- $[\text{Co}(\text{H}_2\text{O})_2(\text{SCN})_2]\cdot\text{H}_2\text{O}$ was isolated from cocaine hydrochloride and lidocaine hydrochloride monohydrate reaction vessels, while *cis*- $[\text{Co}(\text{lidocaine})_2(\text{SCN})_2]$ and *trans*- $[\text{Co}(\text{lidocaine})_2(\text{SCN})_2]$ were retrieved in lidocaine reaction vessels. This cataloging of products per analyte enables the optimization of appropriate reaction conditions to favor a desired product. In general for Co(II), different chemistry strategies would be required to optimize for an ion pair versus a coordination compound for each specific analyte.



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CONFLICT OF INTEREST STATEMENT

The authors report no conflicts of interest.

ORCID

Raychelle Burks  <https://orcid.org/0000-0003-4696-609X>
Francoise M. Amombo Noa  <https://orcid.org/0000-0001-8361-3432>

REFERENCES

- Philp M, Fu S. A review of chemical "spot" tests: a presumptive illicit drug identification technique. *Drug Test Anal.* 2018;10(1):95–108. <https://doi.org/10.1002/dta.2300>
- Bell S. *Drugs as physical evidence: seized drugs and their analysis.* Forensic chemistry. 2nd ed. Upper Saddle River, NJ: Pearson; 2013. p. 213–56.
- de Souza DM, de Moura Messias PJ, da Silva Santos I, Ramalho ED, Ferrari Júnior E, de Oliveira Moraes PA. Scott test associated with multivariate image analysis: a more selective alternative for cocaine research in forensic laboratories. *Forensic Sci Int.* 2022;335:111277. <https://doi.org/10.1016/j.forsciint.2022.111277>
- Nerin C, Garnica A, Cacho J. Indirect determination of alkaloids and drugs by atomic absorption spectrometry. *Anal Chem.* 1985;57(1):34–8. <https://doi.org/10.1021/ac00279a013>
- Schlesinger HL. Topics in the chemistry of cocaine. *Bull Narc.* 1985;37(1):63–78.
- Young JL. The detection of cocaine in the presence of novocaine by means of cobalt thiocyanate. *Am J Pharm.* 1931;103(12):709–10.

7. Stainier CL. Utilisation du thiocyanate de cobalt dans l'analyse des bases organiques [the use of cobalt thiocyanate in the analysis of organic bases]. *Il Farmacco*. 1974;29(3):119–35.
8. Deltombe J, Leboutte G, Rosier N. The application of compounds with a cobalt sulfocyanide base to the determination of alkaloids. *J Pharm Belg*. 1962;17:236–8.
9. Eisman M, Gallego M, Valcarcel M. Automatic continuous-flow method for the determination of cocaine. *Anal Chem*. 1992;64(14):1509–12. <https://doi.org/10.1021/ac00038a003>
10. Shahine S, Khamis S. The spectrophotometric and heterometric determination of some alkaloids as cobalthiocyanates. *Microchem J*. 1983;28(1):26–32. [https://doi.org/10.1016/0026-265X\(83\)90024-3](https://doi.org/10.1016/0026-265X(83)90024-3)
11. National Institute of Justice. Color test reagents/kits for preliminary identification of drugs of abuse, NIJ standard-0604.01. Washington, DC: U.S. Department of Justice, National Institute of Justice; 2000.
12. Kosecki PA, Brooke P, Canonico E. Fentanyl as a potential false positive with color tests commonly used for presumptive cocaine identification. *J Forensic Sci*. 2022;67(5):2082–8. <https://doi.org/10.1111/1556-4029.15090>
13. Tsumura Y, Mitome T, Kimoto S. False positives and false negatives with a cocaine-specific field test and modification of test protocol to reduce false decision. *Forensic Sci Int*. 2005;155(2–3):158–64. <https://doi.org/10.1016/j.forsciint.2004.11.011>
14. Tsujikawa K, Iwata YT, Segawa H, Yamamuro T, Kuwayama K, Kanamori T, et al. Development of a new field-test procedure for cocaine. *Forensic Sci Int*. 2017;270:267–74. <https://doi.org/10.1016/j.forsciint.2016.10.019>
15. Burks R, Winokur A. Presumptive chemical tests. In: Houck MM, editor. *Encyclopedia of forensic sciences*. 3rd ed. Oxford, UK: Elsevier; 2023. p. 263–77.
16. Alliston GV, Bartlett AF, De Faubert Maunder MJ, Phillips GF. An improved test for cocaine, methaqualone and methadone with a modified cobalt (II) thiocyanate reagent. *Analyst*. 1972;97(153):263–5. <https://doi.org/10.1039/an9729700263>
17. Oguri K, Wada S, Eto S, Yamada H. Specificity and mechanism of the color reaction of cocaine with cobaltous thiocyanate. *Japanese J Toxicol Environ Health*. 1995;41(4):274–9.
18. United Nations Office on Drugs and Crime. Cocaine—a spectrum of products, cocaine insights 2. Vienna, Austria: UNODC; 2021. p. 31.
19. de Jong M, Florea A, Eliaerts J, Van Durme F, Samyn N, De Wael K. Tackling poor specificity of cocaine color tests by electrochemical strategies. *Anal Chem*. 2018;90(11):6811–9. <https://doi.org/10.1021/acs.analchem.8b00876>
20. McGill JW, Dixon CA, Ritter D, Sides JD. Discovery of an interesting temperature effect on the sensitivity of the cobalt thiocyanate test for cocaine. *Microgram*. 2008;6:26–35.
21. Batten SR, Champness NR, Chen X-M, Garcia-Martinez J, Kitagawa S, Öhrström L, et al. Terminology of metal-organic frameworks and coordination polymers (IUPAC recommendations 2013). *Pure Appl Chem*. 2013;85(8):1715–24. <https://doi.org/10.1351/PAC-REC-12-11-20>
22. Cano FH, García-Blanco S, Laverat AG. The crystal structure of cobalt(II) thiocyanate trihydrate. *Acta Crystallogr B*. 1976;32(5):1526–9. <https://doi.org/10.1107/S0567740876005694>
23. United Nations Office on Drugs and Crime. Rapid testing methods of drugs of abuse. Vienna, Austria: UNODC; 1994. p. 102.
24. Zeltmann AH, Morgan LO. Ligand substitution processes in aqueous cobalt(II)-thiocyanate solutions. Nuclear magnetic resonances of oxygen-17 and nitrogen-14. *Inorg Chem*. 1970;9(11):2522–8. <https://doi.org/10.1021/ic50093a029>
25. Bahta A, Parker G, Tuck D. Critical survey of stability constants of complexes of thiocyanate ion. *Pure Appl Chem*. 1997;69(7):1489–548. <https://doi.org/10.1515/iupac.69.0563>
26. Aromí G, Batsanov AS, Christian P, Helliwell M, Parkin A, Parsons S, et al. Synthetic and structural studies of cobalt-pivalate complexes. *Chemistry*. 2003;9(20):5142–61. <https://doi.org/10.1002/chem.200304993>
27. Lockwood T-LE, Leong TX, Bliese SL, Helmke A, Richard A, Merga G, et al. idPAD: paper analytical device for presumptive identification of illicit drugs. *J Forensic Sci*. 2020;65(4):1289–97. <https://doi.org/10.1111/1556-4029.14318>
28. Hurst TK. An analysis of the chemistry of the Scott (Ruybal) test for cocaine [dissertation]. East Lansing, MI: Michigan State University; 2002.
29. Johnson CE. Distribution studies of cobalt thiocyanate complexes [dissertation]. East Lansing, MI: Michigan State University; 1958.
30. Cotton FA, Wilkinson G, Murillo C, Bochmann M. *Advanced inorganic chemistry*. 6th ed. New Delhi, India: Wiley India Pvt. Limited; 2007. p. 817–24.
31. Hartshorn RM, Hellwich K-H, Yerin A, Damhus T, Hutton AT. Brief guide to the nomenclature of inorganic chemistry. *Pure Appl Chem*. 2015;87(9–10):1039–49. <https://doi.org/10.1515/pac-2014-0718>
32. Gillespie JE, Fanourakis A, Phipps RJ. Strategies that utilize ion pairing interactions to exert selectivity control in the functionalization of C–H bonds. *J Am Chem Soc*. 2022;144(40):18195–211. <https://doi.org/10.1021/jacs.2c08752>
33. Cotton FA, Goodgame DML, Goodgame M. The electronic structures of tetrahedral cobalt(II) complexes. *J Am Chem Soc*. 1961;83(23):4690–9. <https://doi.org/10.1021/ja01484a002>
34. Magnell KR, Reynolds WL. Complexes of cobalt(II) with chloride and thiocyanate ions in dimethyl sulfoxide. *Inorganica Chim Acta*. 1972;6:571–4. [https://doi.org/10.1016/S0020-1693\(00\)91859-0](https://doi.org/10.1016/S0020-1693(00)91859-0)
35. Carson MC. Ion-pair solid-phase extraction. *J Chromatogr A*. 2000;885(1–2):343–50. [https://doi.org/10.1016/S0021-9673\(00\)00471-4](https://doi.org/10.1016/S0021-9673(00)00471-4)
36. Phadungcharoen N, Pengwanput N, Nakapan A, Sutitaphan U, Thanomklom P, Jongudomsombut N, et al. Ion pair extraction coupled with digital image colorimetry as a rapid and green platform for pharmaceutical analysis: an example of chlorpromazine hydrochloride tablet assay. *Talanta*. 2020;219:121271. <https://doi.org/10.1016/j.talanta.2020.121271>
37. Laussmann T, Grzesiak I, Krest A, Stirnat K, Meier-Giebing S, Ruschewitz U, et al. Copper thiocyanate complexes and cocaine—a case of “black cocaine”. *Drug Test Anal*. 2015;7(1):56–64. <https://doi.org/10.1002/dta.1658>
38. Oliver AG, Lockwood T-LE, Zinna J, Lieberman M. Bis(N,N-diethyl-4-methyl-4-piperazine-1-carboxamide) tetra-kis-(iso-thio-cyanato-κN)-cobalt(II), a model compound for the blue color developed in the Scott test. *Acta Crystallogr E Crystallogr Commun*. 2023;79(Pt 3):163–6. <https://doi.org/10.1107/S2056989023000981>
39. Haddoub R, Ferry D, Marsal P, Siri O. Cobalt thiocyanate reagent revisited for cocaine identification on TLC. *New J Chem*. 2011;35(7):1351–4. <https://doi.org/10.1039/C1NJ20234K>
40. Kovar K-A, Laudsun M. *Chemistry and reaction mechanisms of rapid tests for drugs of abuse and precursors chemicals*. Vienna, Austria: United Nations; 1989. Scientific and Technical Notes: v.89-51669.
41. Nuffield Foundation. The equilibrium between two coloured cobalt species. RSC Education. 2018. <https://edu.rsc.org/experiments/the-equilibrium-between-two-coloured-cobalt-species/1.article>. Accessed 16 Apr 2023.
42. United Nations Office on Drugs and Crime. Recommended methods for the identification and analysis of cocaine in seized materials. Vienna, Austria: UNODC; 2012. p. 20–1.
43. Deakin AL. A study of acids used for the acidified cobalt thiocyanate test for cocaine base. *Microgram J*. 2003;1(1–2):40–3.
44. Sadyrbaeva TZ. Separation of cobalt(II) from nickel(II) by a hybrid liquid membrane-electrodialysis process using anion exchange

- carriers. *Desalination*. 2015;365:167–75. <https://doi.org/10.1016/j.desal.2015.02.036>
45. Blitz-Raith AH, Paimin R, Cattrall RW, Kolev SD. Separation of cobalt(II) from nickel(II) by solid-phase extraction into Aliquat 336 chloride immobilized in poly(vinyl chloride). *Talanta*. 2007;71(1):419–23. <https://doi.org/10.1016/j.talanta.2006.04.017>
 46. Chang J-C, Ho W-Y, Sun I-W, Chou Y-K, Hsieh H-H, Wu T-Y. Synthesis and properties of new tetrachlorocobaltate (II) and tetrachloromanganate (II) anion salts with dicationic counterions. *Polyhedron*. 2011;30(3):497–507. <https://doi.org/10.1016/j.poly.2010.11.009>
 47. Zhou G, Wang B, Ding L, Dong J, Wang F, Feng C. Measurement and correlation of the solubility of lidocaine in eight pure and mixed solvents at temperatures from (292.15 to 332.15) K. *J Mol Liq*. 2017;242:168–74. <https://doi.org/10.1016/j.molliq.2017.06.110>
 48. Sheldrick GM. Crystal structure refinement with SHELXL. *Acta Crystallogr B*. 2015;71(Pt 1):3–8. <https://doi.org/10.1107/S2053229614024218>
 49. Dolomanov OV, Bourhis LJ, Gildea RJ, Howard JAK, Puschmann H. OLEX2: a complete structure solution, refinement and analysis program. *J Appl Cryst*. 2009;42(2):339–41. <https://doi.org/10.1107/S0021889808042726>
 50. Groom CR, Bruno IJ, Lightfoot MP, Ward SC. The Cambridge structural database. *Acta Crystallogr B Struct Sci Cryst Eng Mater*. 2016;72(Pt 2):171–9. <https://doi.org/10.1107/S2052520616003954>
 51. Conceição VN, Souza LM, Merlo BB, Filgueiras PR, Poppi RJ, Romão W. Study of Scott test using spectroscopic techniques: an alternative method for detecting cocaine hydrochloride and its adulterants in street drugs. *Quim Nova*. 2014;37(9): 1538–44. <https://doi.org/10.5935/0100-4042.20140240>
 52. Qayyas NNA, Sridhar M, Indira A, Prasad JS, Abdoh M. Crystal structure of bis(lignocainium) tetrathiocyanatocobaltate(II) hydrate, (C₁₄H₂₂ON₂)₂ Co(NCS)₄ (H₂O). *Z Kristallogr Cryst Mater*. 1994;209(11):918–9. <https://doi.org/10.1524/zkri.1994.209.11.918>
 53. Etter MC, MacDonald JC, Bernstein J. Graph-set analysis of hydrogen-bond patterns in organic crystals. *Acta Crystallogr B*. 1990;46(Pt 2):256–62. <https://doi.org/10.1107/s0108768189012929>
 54. Standard practice for microcrystal testing in forensic analysis for cocaine. <https://www.astm.org/e1968-19.html>. Accessed 27 Jul 2023
 55. Wood MR, Brettell TA, Lalancette RA. The gold(III) tetrachloride salt of L-cocaine. *Acta Crystallogr C*. 2007;63(2):m33–5. <https://doi.org/10.1107/S0108270106053108>
 56. Wood MR, Mikhael S, Bernal I, Lalancette RA. Erdmann's anion—an inexpensive and useful species for the crystallization of illicit drugs after street confiscations. *Chemistry*. 2021;3(2):598–611. <https://doi.org/10.3390/chemistry3020042>
 57. Nicholls D. The chemistry of iron, cobalt and nickel. Oxford, UK: Pergamon Press; 1975. p. 1090–2.
 58. Nakamoto K. Infrared and Raman spectra of inorganic and coordination compounds, part B: applications in coordination, organometallic, and bioinorganic chemistry. 6th ed. Hoboken, NJ: John Wiley & Sons; 2009. p. 120–32.
 59. Maienschein-Cline MG, Londergan CH. The CN stretching band of aliphatic thiocyanate is sensitive to solvent dynamics and specific solvation. *J Phys Chem A*. 2007;111(40):10020–25. <https://doi.org/10.1021/jp0761158>
 60. Smith BC. Organic nitrogen compounds V: amine salts. *Spectroscopy*. 2019. <https://www.spectroscopyonline.com/view/organic-nitrogen-compounds-v-amine-salts>. Accessed 3 Aug 2023.
 61. Price SL. Predicting crystal structures of organic compounds. *Chem Soc Rev*. 2014;43(7):2098–111. <https://doi.org/10.1039/c3cs60279f>
 62. Tabrizi L, McArdle P, Erxleben A, Chiniforoshan H. Nickel(II) and cobalt(II) complexes of lidocaine: synthesis, structure and comparative in vitro evaluations of biological perspectives. *Eur J Med Chem*. 2015;103:516–29. <https://doi.org/10.1016/j.ejmech.2015.09.018>
 63. Lugo AF, Richards AF. Ketiminato and ketimine Co, Eu, Cu and Fe complexes. *Inorganica Chim Acta*. 2010;363(10):2104–12. <https://doi.org/10.1016/j.ica.2010.02.019>
 64. Ma H, Wan C, Zewail AH. Dynamics of ligand substitution in labile cobalt complexes resolved by ultrafast T-jump. *Proc Natl Acad Sci USA*. 2008;105(35):12754–57. <https://doi.org/10.1073/pnas.0806869105>

SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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