Extremely Bent Cyanide Coordination at a Preorganized Dinickel Site and Assembly of a Starlike Nonanuclear Complex from the Constrained Dinickel Building Blocks

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An extremely bent cyanide coordination at a dinickel scaffold is reported. Preorganization of two nickel ions is achieved by means of a compartmental dinucleating pyrazolate ligand L^- , setting up a bimetallic coordination pocket with constrained metal-metal separation. The mixed-spin dinickel(II) complex $[LNi₂(CN)(MeCN)](ClO₄)₂$ (**1**) has been characterized by X-ray diffraction. The MeCN bound to the high-spin nickel(II) ion can be removed or replaced by other ligands, e.g., by the cyanide ligand of a tetracyanonickelate(II) moiety to give the starlike nonanuclear complex $\{[LN]_2(CN)]_4[N(CN)_4]\}(CO_4)_6$ (2) that contains four of the constrained pyrazolate-based dinickel(II) fragments grouped around a central tetracyanonickelate(II) unit, as revealed by X-ray crystallography. Spectral and electrochemical properties of 1 and 2 are reported, and the formation of reduced mixed-valent Ni^INi^{II} species is investigated by IR and UV/vis spectroelectrochemistry.

Introduction

Current interest in oligonuclear transition metal complexes is often inspired by the prospect of distinct reactivity patterns or interesting physicochemical properties that may arise from the cooperative action of the adjacent metal centers.^{1,2} With regard to bimetallic reactivity much effort has been devoted to the design of elaborate dinucleating ligand matrixes that provide two coordination compartments in order to hold two metal ions in suitable proximity.3 Multidentate pyrazole derivatives have proven to be valuable ligand scaffolds for this purpose, as the metal-metal separation and individual coordination environments can be selectively altered by appropriate changes of chelating donor sidearms attached to the bridging heterocycle. $4-7$ The constrained preorganization of two metal ions in such pyrazolate-based dinuclear complexes has allowed the observation of unusual binding modes of small molecules within the bimetallic pocket, e.g., the enforced side-on π -interaction of a nitrile unit with a high-spin nickel (II) site.⁸

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In pursuit of special physicochemical phenomena arising from the cooperativity of several metal ions, such as magnetic ordering or electronic interactions, simple and tight-binding bridging ligands are employed preferentially for the construction of extended and ordered metal arrays. The bidentate cyanide ligand has played a major role in this field, as it is able to facilitate long-range electron transfer and to mediate magnetic exchange interactions. $9-12$ In addition, it generally forms extraordinarily stable complexes that allow for the construction of two- and three-dimensional arrays of cyanide-based coordination units.¹²

In the present contribution we combine the use of a multidentate compartmental ligand platform and of cyanide chemistry to (i) enforce an extremely bent cyanide binding mode within a predefined bimetallic pocket and (ii) assemble an unusual starlike nickel complex of higher nuclearity from such bimetallic building blocks. Structural features of these novel systems are reported, in particular the most acute $C \equiv N-M$ angles known

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to date for μ_2 - κ C: κ N-cyanide (111.2-112.9°), and redox phenomena mediated by the cyanide bridges are studied.

Experimental Section

General Procedures and Methods. All manipulations were carried out under an atmosphere of dry nitrogen by employing standard Schlenk techniques. Solvents were dried according to established procedures. HL was synthesized according to the reported method;^{4a} all other chemicals were used as purchased. Microanalyses: Mikroanalytische Laboratorien des Organisch-Chemischen Instituts der Universität Heidelberg. IR spectra: Perkin-Elmer 983G; recorded as KBr pellets. MS spectra: Finnigan MAT 8230 (FAB) or Finnigan TSQ 700 (ESI). UV/vis spectra: Perkin-Elmer Lambda 19. Magnetic measurement: Bruker Magnet B-E 15 C8, field controller B-H 15, variable temperature unit ER4111VT, Sartorius micro balance M 25 D-S. Experimental susceptibility data were corrected for the underlying diamagnetism. Cyclic voltammetry: PAR equipment (potentiostat/galvanostat 273), in 0.1 M NBu₄PF₆ MeCN solution. Potentials in volts on glassy carbon electrode, referenced to the SCE at ambient temperature. Spectroelectrochemistry: Self-constructed OTTLE cell comprising a Pt-mesh working and counter electrode and a silver wire as pseudo reference electrode sandwiched in between the CaF₂ windows of a conventional liquid IR cell. The working electrode is positioned in the center of the spectrometer beam with all other parts of the cell made nontransparent to the incident beam by means of an absorbing tape.25

CAUTION! Although no problems were encountered in this work, transition metal perchlorate complexes are potentially explosive and should be handled with proper precautions.

Synthesis of 1. A solution of HL (250 mg, 0.48 mmol) in ethanol (30 mL) was treated with 1 equiv of KO*^t* Bu (54 mg) and 2 equiv of $[Ni(H₂O)₆](ClO₄)₂$ (350 mg, 0.96 mmol). After stirring for 30 min, [NEt4]CN (73 mg, 0.47 mmol) was added in one portion. The reaction mixture was stirred for a further 1 h and was filtered. Evaporation of all volatile material under reduced pressure afforded a green residue, which was redissolved in acetonitrile. Layering the solution with diethyl ether gradually gave green crystals of **¹**'1.4MeCN (300 mg, 69%). IR (KBr, cm⁻¹): 2964 (m), 2877 (w), 2284 [w, $ν$ (C=N)_{MeCN}], 2245 [w, *ν*(C=N_{)MeCN}], 2106 [m, *ν*(C=N_{)CN}], 1461 (s), 1383 (m), 1305 (w), 1088 [vs, *ν*(ClO4)], 795 (m), 739 (m), 623 (s). Drying the product under dynamic vacuum causes the IR spectrum to change as follows and yields **1**[′]: IR (KBr, cm⁻¹): 2971 (m), 2874 (w), 2096 [m, *ν*(C≡N)_{CN}], 1468 (s), 1384 (m), 1300 (w), 1090 [vs, *ν*(ClO4)], 792 (m), 737 (m), 623 (s). MS (FAB+) $[m/z]$ (relative intensity)]: 762 (100) $[LNi₂(CN)$ - $(CIO₄)$ ⁺, 663 (75) [LNi₂(CN)]⁺. Anal. Calcd for C₃₀H₆₁Cl₂N₉Ni₂O₈ [LNi2(CN)(ClO4)2, **1**′]: C, 41.69; H, 7.11; N, 14.59. Found: C, 41.16; H, 7.22; N, 13.62.

Synthesis of 2. The synthesis is similar to the procedure described for **1** starting from HL (250 mg, 0.48 mmol), but almost 2 equiv of [NEt4]CN (145 mg, 0.93 mmol) was added. Purification was achieved by diffusion of diethyl ether into a solution of the product in acetone/ wet methanol (1:1), which yielded $2.0.5H₂O$ as greenish-yellow crystals (280 mg, 68% with respect to HL). IR (KBr, cm⁻¹): 2962 (m), 2869 (w), 2159 [m, *ν*(C≡N)], 2107 [m, *ν*(C≡N)], 1518 (w), 1453 (m), 1376 (m), 1341 (w), 1309 (w), 1094 [vs, *ν*(ClO4)], 794 (m), 739 (m), 622 (s). MS (FAB+) $[m/z]$ (relative intensity)]: 1592 (35) $\{[LNi_2(CN)]_2 Ni(CN)_{4}(ClO_{4})$ ⁺, 764 (60) [LNi₂(CN)(ClO₄)]⁺, 689 (65) [LNi₂(CN)₂]⁺, 663 (100) [LNi2(CN)]+. MS (ESI+) [*m*/*^z* (relative intensity)]: 1592 (5) $\{[LNi_2(CN)]_2Ni(CN)_4(CIO_4)\}^+$, 1519 (10) $\{[LNi_2(CN)]_2Ni(CN)_4$ - $(CIO₄)$ -HNEt₂}⁺, 764 (20) [LNi₂(CN)(ClO₄)]⁺, 746 (90) {[LNi₂(CN)]₂-Ni(CN)₄}²⁺, 689 (100) [LNi₂(CN)₂]⁺. MS (ESI-) [*m*/*z* (relative intensity)]: 1790 (10) $\{[LNi_2(CN)]_2Ni(CN)_4(ClO_4)_3\}^-$, 962 (5) $[LNi_2(CN)(ClO_4)_3]^-$. Anal. Calcd for $C_{124}H_{244}Cl_6N_{40}Ni_9O_{24}$ ⁻0.5 H_2O : C, 43.43; H, 7.20; N, 16.34. Found: C, 43.15; H, 7.23; N, 15.95.

X-ray Crystallography. The measurements were carried out on a Nonius Kappa CCD diffractometer using graphite-monochromated Mo $K\alpha$ radiation. All calculations were performed using the SHELXT PLUS software package. Structures were solved by direct methods with the SHELXS-97 and refined with the SHELXL-97 program.¹³ Atomic coordinates and thermal parameters of the non-hydrogen atoms were refined in fully or partially anisotropic models by full-matrix least-

Table 1. Crystal Data and Refinement Details for Complexes **1** and **2**

	$1.1.4$ MeCN	2.0.5H ₂ O
formula	$C_{32}H_{64}C_{12}N_{10}Ni_2O_8$ 1.4MeCN	$C_{124}H_{244}C_{16}N_{40}Ni_9O_{24}$ 0.5H ₂ O
$M_{\rm r}$	962.68	3420.49
cryst size (mm)	$0.40 \times 0.25 \times 0.15$	$0.20 \times 0.15 \times 0.05$
cryst syst	orthorhombic	triclinic
space group	$P2_12_12_1$	P ₁
a(A)	13.717(3)	20.593(4)
b(A)	14.054(3)	20.793(4)
c(A)	24.374(5)	21.034(4)
α (deg)	90	90.79(3)
β (deg)	90	95.78(3)
γ (deg)	90	110.32(3)
$V(A^3)$	4699(1)	8392(1)
$\rho_{\rm{calcd}}$ (g cm ⁻³)	1.361	1.344
Ζ	4	2
$F(000)$ (e)	2043	3604
T(K)	200	200
$\mu(Mo\ K\alpha)$ (mm ⁻¹)	0.972	1.151
scan mode	ω	ω
hkl range	-12 to 17, -14 to 18,	-28 to 28, -28 to 28,
	-24 to 31	-29 to 29
2θ range (deg)	$3.3 - 55.0$	$2.7 - 59.6$
measd reflns	23592	90732
unique reflns	10783	46588
obsd reflns $[I \geq 2\sigma(I)]$	9031	20687
refined params	602	1854
resid electron dens $(e \, \mathring{A}^{-3})$	$0.68/-0.41$	$1.66/-1.14$
R ₁	0.044	0.083
wR2 (refinement on F^2)	0.122	0.269
goodness-of-fit	1.005	1.000

Chart 1

squares calculations based on $F²$. In general the hydrogen atoms were placed at calculated positions and allowed to ride on the atoms to which they are attached. For **1**, MeCN solvent molecules are included at two positions in the unit cell, each one having an occupancy of 70%. The only moderate quality of the structural analysis of **2** mainly results from severe disorder of some of the perchlorate counteranions. Also, most of the residual electron density is located in the vicinity of the perchlorate. Part of the perchlorate O atoms could not be refined anisotropically. Data for **2** have been collected to rather high 2*θ* values in order to obtain a reasonable reflex/parameter ratio. Table 1 compiles the data for the structure determinations.

Results and Discussion

Synthesis and Structural Characterization. The dinucleating pyrazole ligand HL is employed in the present work.^{4a} Its anionic form, L^- (Chart 1), has previously been shown to bind two metal ions in its adjacent coordination compartments, setting up a central bimetallic pocket in which additional ligands or substrate molecules may be held within the grip of the two metals.6,8,14 As enforced by the particular length of the ligand

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Figure 1. View of the molecular structure of the cation of **1** (40% probability ellipsoids). In the interest of clarity all hydrogen atoms have been omitted.

Chart 2

sidearms, the metal-metal separation in complexes of L^- is too large for allowing small ligands like OH- to form a monatomic secondary bridge within the bimetallic pocket.⁶ On the other hand, molecular models suggest that incorporation of two-atom bridges that prefer linear coordination, e.g., the cyanide ion in M-CN-M fragments, is strongly disfavored due to the geometrical constraints imposed by the dinuclear scaffold.

Addition of 1 equiv of [NEt4]CN to an in situ prepared dinickel(II) complex of L^- in acetonitrile afforded a green solution, from which greenish-yellow crystals of the new complex **1** could be isolated (Chart 2). Binding of cyanide was indicated by FAB mass spectrometry, showing major signals for $[LNi_2(CN)(ClO_4)]^+$ and $[LNi_2(CN)]^+$ with the expected isotopic distribution patterns. The molecular structure of **1** was revealed by X-ray crystallography, the result of which is depicted in Figure 1. Selected interatomic distances and bond angles are listed in Table 2.

As anticipated, the two nickel ions in **1** are nested within their respective coordination compartments and are spanned by the pyrazolate moiety. The cyanide ligand is housed within the bimetallic pocket, thereby inducing a pronounced asymmetry of the dinuclear framework. Assignment of the C and N atoms of the cyanide was based on the clearly different quality of the two crystallographic models for both alternative orientations, and on the geometric characteristics discussed in the following: as expected for the strong-field cyanide ligand, the Ni1 ion bound to the cyanide C is found in a square planar environment, which leaves one of the respective ligand sidearms uncoordinated and dangling. In contrast, Ni2 bound to the cyanide N is six-coordinate due to the attachment of an additional MeCN solvent molecule in an exo position cis to the cyanide and trans to the pyrazolate nitrogen atom. Magnetic susceptibility measurements on a powdered sample of **1** confirm the presence of a mixed-spin complex with only one high-spin nickel(II) ion $(S = 1)$ per bimetallic entity.

While the N9-C30 bond length [1.152(4) Å] is in the normal range, other geometric parameters around the bridging cyanide

Figure 2. Scatter plot showing the correlation between the metalmetal distance $d(M-M)$ and the angle M-N-C for cyanide-bridged complexes obtained from a CSD search:¹⁶ (O) complexes with μ_2 -*κ*C: κ N-cyanide; (\bullet) PtCu complex reported by Falvello et al.;¹⁵ (\blacksquare) complexes **1** and **2**; (\triangle) complexes with μ_2 - κ C: η ²-C,N-cyanide.¹⁷

Table 2. Selected Interatomic Distances (Å) and Bond Angles (deg) for **1**

ັ			
$Ni1-N1$	1.845(3)	Ni2-N7	2.335(3)
$Ni1-N3$	1.945(3)	$Ni2-N8$	2.254(3)
$Ni1 - N4$	1.933(3)	$Ni2-N9$	2.198(3)
$Ni1-C30$	1.863(4)	$Ni2-N10$	2.056(3)
$Ni2-N2$	2.029(3)	$Ni1\cdots Ni2$	4.002
$Ni2-N6$	2.094(3)		
N1-Ni1-N3	83.9(1)	$N9-Ni2-N2$	91.8(1)
$N3 - N_11 - N_4$	89.4(1)	$N9-Ni2-N7$	97.4(1)
N4-Ni1-C30	99.3(1)	N9-Ni2-N8	97.3(1)
$C30-Ni1-N1$	88.3(1)	$N9 - Ni2 - N10$	88.7(1)
$N1-Ni1-N4$	164.2(1)	$N2-Ni2-N6$	80.3(1)
C30-Ni1-N3	170.9(1)	$N2-Ni2-N7$	89.4(1)
Ni1-C30-N9	162.1(3)	$N2-Ni2-N8$	94.2(1)
$C30-N9-Ni2$	111.2(2)	N6-Ni2-N7	82.1(1)
N9-Ni2-N6	172.1(1)	N6-Ni2-N8	84.0(1)
$N2-Ni2-N10$	177.0(1)	N6-Ni2-N10	99.2(1)
N7-Ni2-N8	164.8(1)	$N10-Ni2-N7$	87.6(1)
		$N10-Ni2-N8$	88.7(1)

ligand are quite unusual. The bond angle $Ni1-C30-N9$ is relatively small $[162.1(3)^\circ]$, and the angle Ni2-N9-C30 is extremely acute for a μ_2 - κ C: κ N-cyanide [111.2(2)^o], which apparently is enforced by the fixed metal-metal separation and the rigidity of the bimetallic scaffold. A different case of a severely bent cyanide bridging between a platinum(II) and a copper(II) has recently been observed by Falvello et al. (angle $C=N-Cu: 120.1(6)°$.¹⁵ In that compound, the two metal ions are not linked by any other ligand than the bent cyanide, and the deformation from linearity is traced to a collection of noncovalent intermolecular and intramolecular H-bonding interactions in the solid state.

The strong deviation of the situation encountered in **1** from the more conventional geometric patterns of cyanide binding is evidenced by a scatter plot showing the M'''M distances versus the $M-N-C$ angles for 454 $M-CN-M$ fragments obtained from a CSD search as depicted in Figure 2.16 Metalmetal distances for bridging *µ*2-*κ*C:*κ*N-cyanide are usually found in the range 4.75-5.75 Å, and angles $C \equiv N-M$ at the cyanide N generally lie above 120° (O; the PtCu complex reported by

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Table 3. Selected Interatomic Distances (Å) and Bond Angles (deg) for 2^a

$Ni1-C30$ $C30-N9$ $Ni2-N9$ $Ni2-N10$ $N10 - C31$ $Ni9-C31$	1.853(6)/1.860(6)/1.862(6)/1.854(6) 1.158(7)/1.163(7)/1.156(7)/1.158(7) 2.209(5)/2.197(5)/2.199(5)/2.239(5) 2.038(5)/2.029(5)/2.042(5)/2.033(5) 1.144(7)/1.157(7)/1.146(7)/1.158(7) 1.881(6)/1.867(6)/1.875(6)/1.863(6)
$Ni1 \cdots Ni2$	4.022/3.997/4.024/4.033
$Ni2\cdots Ni9$	5.031/4.980/5.024/5.027
$Ni1 - C30 - N9$ $C30-N9-Ni2$ $Ni2-N10-C31$ $N10 - C31 - N19$	161.6(5)/160.7(5)/160.5(5)/160.7(5) 112.0(4)/111.2(4)/112.9(4)/112.0(4) 168.6(5)/166.0(5)/168.9(5)/168.8(4) 176.9(5)/171.5(6)/174.7(6)/177.3(5)

^a Atom numbering refers to the Ni1/Ni2 subunit, but the corresponding values for the Ni3/Ni4, Ni5/Ni6, and Ni7/Ni8 subunits are also listed.

Table 4. IR Absorptions in the C \equiv N Stretching Range (Values in cm^{-1})

1 ^a	2284, 2245, 2106
$1'$ a	2096
1 ^b	2112
$\mathbf{1}_{\mathrm{red}}^b$	1950 (sh), 1905
2^a	2159, 2107
2 ^b	2160, 2112
$2_{\rm red}$ ' b	2098, 1950 (sh), 1905

^a KBr pellet. *^b* In MeCN solution.

Falvello et al.¹⁵ is highlighted as a solid circle). In contrast, the Ni \cdots ^Ni distance in **1** is fixed at only 4.002 Å, thus causing the extreme bending of the Ni-C \equiv N-Ni fragment (\Box). Also included in Figure 2 are those values for the related complex **2** described below and for the few crystallographically characterized systems where μ_2 - κ C: η ²-C,N bridging of a cyanide ligand has been observed (A) .¹⁷ In these latter cases the metal ions that exhibit η^2 side-on π -bonding to the C=N cyanide bond are electron-rich, low-valent 4d/5d-transition metals well capable of *π*-back-bonding, and the bimetallic scaffolds feature short ^M-M distances due to metal-metal bonds, which is clearly different from the situation in **1**. Enforced by the geometric constraints of the pyrazolate ligand, **1** adopts geometric parameters en route to such rare $\eta^2 \pi$ -bonding of the bridging cyanide, although any side-on π -interaction is supposedly negligible because of the high-spin character and the relative "hardness" of the N-bound Ni2 ion. It should be noticed, however, that some weak side-on π -interaction with a high-spin nickel(II) ion could be enforced for a nitrile $C \equiv N$ bonded within the highly preorganized bimetallic pocket of a similar dinickel(II) system.8

The IR spectrum (KBr) of freshly prepared **1** shows a band for the ν (C=N) cyanide stretch at 2106 cm⁻¹ (Table 4). Its somewhat higher frequency compared to the band position for free cyanide is not unexpected and is usually attributed to the kinematic coupling effect that occurs upon attachment of two metal ions to both ends of a cyanide.18 The observed band position for the CN stretch is by no means unusual for a μ_2 *κ*C:*κ*N-cyanide, despite the unusual binding situation imposed by the bimetallic scaffold. This is in accordance with the highspin $+2$ oxidation state of the nearby six-coordinate nickel(II) ion, as already discussed above: even if the positioning of Ni2 appears suitable for interaction with an antibonding *π** orbital of the CN^- ligand [which should lower ν (C=N)], the relative "hardness" of the metal ion and the contracted character of its 3d shell prevent any significant π -back-bonding.

An additional pair of bands appearing at 2245 and 2284 cm-¹ in the IR spectrum of 1 results from the $\nu(C \equiv N)$ stretching vibration of the terminally bound acetonitrile. When powdered material of the isolated complex **1** was dried under vacuum, a slight color change from greenish-yellow to green was observed. The IR spectrum of the new green material **1**′ displays no bands of coordinated acetonitrile and only one cyanide stretch at 2096 cm^{-1} . Apparently, the acetonitrile ligand is only loosely bound, leading to its facile removal under reduced pressure and concomitant generation of an accessible coordination site at Ni2 (in the green compound **1**′, Ni2 presumably is five-coordinate similar to the situation in most other dinickel(II) complexes of the ligand L^{-}).¹⁴ This vacant coordination site in **1'** is now available for the binding of further cyanide building blocks, as demonstrated by the isolation of the starlike nonanuclear complex **2**.

Complex **2** was originally obtained when a slight excess of [NEt4]CN was employed in the synthesis of **1**, which obviously causes formation of the exceedingly stable $[Ni(CN)₄]^{2-}$ ion (*K*) $\approx 10^{31}$).¹⁹ Accordingly, 2 may also be prepared by treating 1 with the appropriate amount of a $[Ni(CN)₄]^{2-}$ salt. An oligonuclear structure of **2** was inferred from the FAB-MS and ESI-MS spectra, which revealed dominant peaks at $m/z = 1592$ corresponding to $[LNi_2(CN)]_2Ni(CN)_4(CIO_4)^+$ (both FAB-MS and ESI-MS) and at $m/z = 746$ corresponding to the dication $[LNi₂(CN)]₂Ni(CN)₄²⁺$ (only ESI-MS) with the expected isotopic distribution patterns. These findings already suggested the attachment of several $[LNi₂(CN)]²⁺ bimetallic fragments to one$ $[Ni(CN)₄]$ ²⁻ group in 2. Greenish-yellow single crystals suitable for X-ray crystallography could be obtained from acetone/wet methanol/diethyl ether. The molecular structure of the cation is shown in Figure 3, and selected interatomic distances and bond angles are listed in Table 3.

Complex 2 consists of four $[LNi₂(CN)]²⁺$ entities, each one similar to **1**, that are held together by a central $[Ni(CN)₄]$ ²⁻¹ unit. Instead of the acetonitrile ligand in **1**, a cyanide ligand of the central $[Ni(CN)₄]²$ unit is now bound to the six-coordinate nickel(II) ions of all pyrazolate-based dinuclear fragments via its nitrogen atom. Geometric parameters of the four central cyanide ligands are ordinary, with $d(C=N)$ in the range 1.144– 1.158 Å and angles C-N-Ni between 166.0° and 168.9° . In contrast, coordination of the outer four cyanide ligands deviates as severely from linearity as was found in **1** (compare Figure 2) with $d(C=N)$ in the range 1.156-1.163 Å and angles $C-N-$ Ni between 111.2° and 112.9°. The two types of cyanide ligands give rise to two IR absorptions at 2107 and 2159 cm⁻¹, where the former is assigned to the bent CN ligands by comparison with the findings for **1** (Table 4). The difference in stretching frequency for the two kinds of bridging CN ligands in **2** can be traced to their different environment: the bent cyanide ligands receive full back-bonding from their respective C-coordinated nickel ions, while the four linear CN ligands have to share the back-bonding ability of the central Ni9.

The $[Ni(CN)₄]²⁻$ unit has previously been employed as a bridging unit in multinuclear metal-cyanide systems.^{20,21}

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Figure 3. View of the molecular structure of the cation of **2** (50% probability ellipsoids). In the interest of clarity all hydrogen atoms, all dangling ligand sidearms, and all N-bound ethyl groups have been omitted.

Table 5. UV/vis Absorptions of the Complexes [Values in nm $(\epsilon$ in M^{-1} cm⁻¹)]

418 (250), 621 (20), 944 (25), 1107 (sh) 1 ^a 1' 414(265), 650(50) 314 (5100), 324 (sh), 356 (5600), 380 (sh), 442 (sh), $\mathbf{1}_{\mathrm{red}}^a$ 505 (740), 1125 (65), 1177 (70) 418 (1125), 644 (100), 955 (110) 2^a $2_{\rm red}$ ' a 805 (410), 1105 (385), 1185 (375)	
	312 (14000), 355 (12800), 381 (sh), 440 (sh), 520 (sh, 1340),

^a In MeCN. *^b* In acetone/MeOH (2:1).

However, very few detailed structures of discrete molecular complexes have been reported, in which the $[Ni(CN)₄]^{2-}$ is linked to four metal ions via all of its terminal cyanide-N atoms,21 and to the best of our knowledge the nonanuclear starlike topology of **2** involving three different types of nickel- (II) ions is unprecedented.

Spectroscopy and Electrochemistry. Electronic absorptions of the different nickel(II) chromophores cannot be unambiguously assigned due to multiple transitions and partial overlap of the various bands. UV/vis data for **1** and **2** are collected in Table 5. Spectra of **1** are highly solvent dependent with pronounced differences when recorded in either MeCN or acetone/methanol (2:1) solution. This is attributed to dissociation of the MeCN ligand in the latter solvent mixture, leaving the respective metal ion five-coordinate in **1**′. Band positions for **2** in acetone/methanol (2:1) are almost identical to those of complex **1** in MeCN solution, in accordance with the above interpretation. While any bands below 350 nm (including the ^d-d transition expected at [∼]320 nm for the tetracyanonickelate- (II) moiety in 2^{2^2} are obscured by intense $\pi \rightarrow \pi^*$ intraligand

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Figure 4. Cyclic voltammogram of complex $1 \cdot (ClO₄)₂$ (a, top) and 2 ['](ClO_{4)⁶} (b, bottom) recorded on a platinum electrode in MeCN containing 0.1 M NⁿBu₄PF₆ as supporting electrolyte at scan speed 200 mV s⁻¹. Indicated peak potentials are given in volts versus SCE.

features, all spectra are characterized by a strong band at 418 $(1, 2)$ or 414 $(1')$ nm. This is assigned to the spin-allowed d_{xy} \rightarrow d_{x} ²- y ² transition of the square planar LNi(CN) fragment, indicating significant destabilization of the antibonding $d_x^2-y^2$
critical $(d - d_x)$ apparation $\approx 24,000$ cm⁻¹) ²³ A modium orbital $(d_{xy}-d_{x^2-y^2})$ separation ≈ 24.000 cm⁻¹).²³ A medium-
intensity band at 650 nm observed for 1' is absent for both 1 intensity band at 650 nm observed for **1**′ is absent for both **1** and **2** and may thus be safely assigned to the five-coordinate nickel(II). In the case of **1** and **2**, a weak absorption at 621 or 644 nm, respectively, presumably arises from the ${}^3A_{2g} \rightarrow {}^3T_{1g}(F)$ transition of the six-coordinate nickel(II), while broad (and overlapping) bands of low intensity around 950 and 1110 nm are tentatively assigned to the ${}^3A_{2g} \rightarrow {}^3T_{1g}(P)$ transitions and to spin-forbidden processes of the square-planar low-spin nickel- (II).

Complexes **1** and **2** have been studied by cyclic voltammetry in MeCN solution in the potential range of $+1.8$ to -1.7 V versus the saturated calomel electrode (SCE). Anodic scans for both **1** and **2** reveal only ill-defined irreversible processes well above $+1.0$ V, thus implying that stable Ni^{III} states are inaccessible. In contrast, cathodic responses are more informative. The dinickel(II) complex **1** shows a well-behaved oneelectron-reduction wave at $E_{1/2} = -1.05$ V that is assigned to the formation of the Ni^INi^{II} mixed-valent state (Figure 4a). This process is fully reversible $[i_{pa}/i_{pc}]$ close to 1; $i_{pc}/v^{1/2} \approx$ constant; $\Delta E_p = 78$ mV with $\Delta E_p(Cp_2Fe/Cp_2Fe^+) = 71$ mV under the same experimental conditions], indicating structural integrity of the bimetallic framework. Further reduction occurs as a broad irreversible wave peaking at around -1.7 V. Continuing the sweep into this region causes disappearance of the anodic response corresponding to reoxidation of the mixed-valent species upon scan reversal. Considering that nickel(I) prefers low coordination numbers and soft π -acceptor ligands, we assume the low-spin nickel(II) in **1** to be the primary reduction site. This is also corroborated by the fact that no similar cathodic process has previously been observed for any of those dinickel-

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Figure 5. IR spectroscopic changes during gradual reduction of **1** (top) and **2** (bottom) in an OTTLE cell.

(II) complexes of HL that feature five- or six-coordinate metal centers within both coordination compartments.14

The nonanuclear complex **2** undergoes reduction at *E*^p red -1.18 V, which is assigned to $Ni¹Ni^{II}$ formation within the nyrazolate-based bimetallic subunits similar to the situation in pyrazolate-based bimetallic subunits similar to the situation in **1** (Figure 4b). However, in the case of **2** this redox process is only partially reversible and appears to consist of several oneelectron steps too closely spaced to give separate waves, suggesting little electronic communication between the redox centers within the four outer bimetallic building blocks. At somewhat more negative potentials the first composite wave is followed by another irreversible process centered at -1.56 V (scan speed 200 mV/s). Upon scan reversal an anodic peak at -0.34 V is found while the immediate reoxidation wave due to the first reduction events is strongly diminished. This feature is also detected if the sweep is reversed past the first composite wave, accounting for the only partial chemical reversibility. A continuous sweep between 0 and -1.38 V reveals no additional cathodic response due to the peak at -0.34 V. We assume a square scheme ECEC mechanism for these redox events of **2**: the hexacation 2 is first reduced to a dicationic species 2_{red} , which undergoes structural rearrangement forming **2red**′. This is reoxidized at -0.34 V, yielding an unstable hexacation $2'$ that quickly reisomerizes to give the starting complex **2**.

To gain further insight into the changes accompanying electron uptake and to characterize the putative mixed-valent Ni^INi^{II} species, the first reductions of 1 and 2 in MeCN were followed by IR and UV/vis spectroscopy in an OTTLE cell (Figures 5 and 6, Tables 4 and 5). Upon gradual electrolysis of **1**, the intensity of the cyanide stretching vibration at 2112 cm^{-1} decreases at the expense of a new band at 1905 cm^{-1} (featuring a shoulder at 1950 cm^{-1}). This large shift to lower frequency of around 200 cm⁻¹ reflects the increased π -back-bonding ability of the reduced Ni^I. Upon reduction of 2, the IR stretch assigned to the bent CN ligands within the peripheral dinickel entities experiences exactly the same shift as was observed for **1**. The CN stretch of the linear cyanide bridges that link the outer dinickel subunits to the central tetracyanonickelate is much less affected, shifting from 2160 to 2098 cm^{-1} . Since both different types of CN ligands are each N-bound to the same high-spin Ni^{II}, this provides clear evidence that reduction takes place at the peripheral four-coordinate low-spin Ni^{II} sites within the

Figure 6. UV/vis spectroscopic changes during gradual reduction of **1** (top) and **2** (bottom) in an OTTLE cell.

bimetallic pyrazolate-bridged entities (and likewise at the respective low-spin Ni^{II} in 1). Albeit smaller, the change of the IR stretching frequency of the inner, linear bridging CN ligands shows that electronic information is transmitted from the outer to the central four-coordinate nickel sites via the six-coordinate Ni^{II}.

In UV/vis spectroelectrochemistry the reduction of both **1** and **2** is accompanied by a dramatic increase in absorptivity (Figure 6). Since upon reoxidation of the reduced species the spectrum of the starting material is nearly restored, the absorption coefficients given are quite reliable. The most prominent feature in the spectra of the reduced complexes is a pair of very intense bands around 313 and 355 cm⁻¹ (log ϵ > 3.6 M⁻¹ cm⁻¹), which we tentatively assign to the $d\pi \rightarrow \pi^*(CN)$ or $d\pi \rightarrow \pi^*$ -(pyrazolate) MLCT transitions. In any electronically coupled, mixed-valent system one expects metal-to-metal intervalence charge-transfer (MMCT-IVCT) absorptions. There is ample literature precedent that bridging cyanide ligands allow for such electronic communication even in unsymmetrical dinuclear complexes.24 Due to the inherently different electronic properties of the low- and high-spin nickel sites the energy of the IVCT transition may be considerably higher than in symmetrical counterparts. Here, the most likely candidate for the IVCT band is the rather weak absorption at 505 nm for 1 ($\epsilon \approx 740 \text{ M}^{-1}$) cm^{-1}) and at 520 nm for 2, both being located at the low-energy portion of the much more intense MLCT bands. Spectral deconvolution gives a half-bandwidth ∆*ν*1/2 of around 7100 cm^{-1} for **1**, in good agreement with the value estimated from Hush's formula $[\Delta v_{1/2} = (2310v_{\text{max}})^{1/2} \approx 6800 \text{ cm}^{-1}]$ describing only moderately coupled class II mixed-valent systems. Additional low-intensity absorptions of the reduced species at around 1150 nm (see insets in Figure 6) clearly have much

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smaller bandwidths and absorptivities and presumably represent ^d-d type transitions.

Conclusions

A dinickel complex and a related nonanuclear array comprising four such bimetallic entities have been prepared and structurally characterized, both incorporating extremely bent cyanide bridges between two different six- and four-coordinated Ni^{II} sites. The two metal ions within the dinickel building blocks are preorganized at a well-defined distance by means of a compartmental ligand matrix. The mixed-spin dinuclear complex **1** features an accessible coordination site at the high-spin nickel- (II) ion, which allowed us to assemble the unique system **2** of higher nuclearity by linking four bimetallic building blocks via a central cyanometalate. Both **1** and **2** can be reduced electrochemically to give mixed-valent Ni^INi^{II} species. The structural features observed for **1** and for the starlike nonanuclear complex **2** demonstrate the ability of two cooperating metal ions to enforce unusual coordination modes by their synergetic action.

The unusually bent cyanide binding at the pyrazolate scaffold is obviously imposed by the dinuclear pincer effect, i.e., by the seizing of the small coligand within a highly preorganized bimetallic pocket. Such phenomena may also be encountered for the highly organized and constrained substrate binding within the protein active site pockets of natural metalloenzymes.

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Supporting Information Available: Crystallographic data, in CIF format, for **1** and **2**. A cyclic voltammogram and IR spectra for **1** and **2**. This material is available free of charge via the Internet at http://pubs.acs.org.

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