Inorganic Chemistry

pubs.acs.org/IC Article

Triapine Analogues and Their Copper(II) Complexes: Synthesis, Characterization, Solution Speciation, Redox Activity, Cytotoxicity, and mR2 RNR Inhibition

- 4 Iuliana Besleaga, Iryna Stepanenko, Tatsiana V. Petrasheuskaya, Denisa Darvasiova, Martin Breza,
- 5 Marta Hammerstad, Małgorzata A. Marć, Alexander Prado-Roller, Gabriella Spengler,
- 6 Ana Popović-Bijelić, Eva A. Enyedy,* Peter Rapta,* Anatoly D. Shutalev,* and Vladimir B. Arion*



Cite This: https://doi.org/10.1021/acs.inorgchem.1c01275



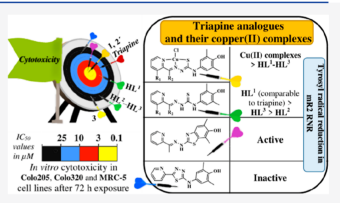
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7 **ABSTRACT:** Three new thiosemicarbazones (TSCs) $\mathrm{HL}^1\mathrm{-HL}^3$ 8 as triapine analogues bearing a redox-active phenolic moiety at the 9 terminal nitrogen atom were prepared. Reactions of $\mathrm{HL}^1\mathrm{-HL}^3$ 10 with $\mathrm{CuCl}_2\cdot 2\mathrm{H}_2\mathrm{O}$ in anoxic methanol afforded three copper(II) 11 complexes, namely, $\mathrm{Cu(HL}^1)\mathrm{Cl}_2$ (1), $[\mathrm{Cu(L}^2)\mathrm{Cl}]$ (2'), and 12 $\mathrm{Cu(HL}^3)\mathrm{Cl}_2$ (3), in good yields. Solution speciation studies 13 revealed that the metal-free ligands are stable as $\mathrm{HL}^1\mathrm{-HL}^3$ at pH 14 7.4, while being air-sensitive in the basic pH range. In dimethyl 15 sulfoxide they exist as a mixture of E and E isomers. A mechanism 16 of the E/E isomerization with an inversion at the nitrogen atom of 17 the Schiff base imine bond is proposed. The monocationic 18 complexes $[\mathrm{Cu(L}^{1-3})]^+$ are the most abundant species in aqueous 19 solutions at pH 7.4. Electrochemical and spectroelectrochemical



studies of 1, 2′, and 3 confirmed their redox activity in both the cathodic and the anodic region of potentials. The one-electron reduction was identified as metal-centered by electron paramagnetic resonance spectroelectrochemistry. An electrochemical oxidation pointed out the ligand-centered oxidation, while chemical oxidations of HL¹ and HL² as well as 1 and 2′ afforded several two-electron and four-electron oxidation products, which were isolated and comprehensively characterized. Complexes 1 and 2′ showed an antiproliferative activity in Colo205 and Colo320 cancer cell lines with half-maximal inhibitory concentration values in the low micromolar concentration range, while 3 with the most closely related ligand to triapine displayed the best selectivity for cancer cells versus normal fibroblast cells (MRC-5). HL¹ and 1 in the presence of 1,4-dithiothreitol are as potent inhibitors of mR2 ribonucleotide reductase as triapine.

28 INTRODUCTION

29 Thiosemicarbazones (TSCs) are known as biologically active 30 compounds with a broad spectrum of pharmacological 31 properties, including anticancer activity. 1—4 These properties 32 can be modulated by a coordination to physiologically relevant 33 metal ions. 5,6 In addition, as versatile ligands, TSCs have 34 tunable electronic and steric properties, which may have a 35 favorable effect on their pharmacological profile. $^{7-10}$ α -N-36 Heterocyclic TSCs such as 2-formylpyridine TSC (FTSC) and 37 5-hydroxy-2-formylpyridine TSC were reported to possess 38 anticancer activity several decades ago, 11,12 and further 39 optimization resulted in the most well-known TSC, 3-40 aminopyridine-2-carboxaldehyde TSC (triapine). Triapine 41 was tested in more than 30 clinical phase I and II trials and 42 currently is involved in a triapine-cisplatin-radiation combina-43 tion therapy in a phase III trial. 13 Because of the documented 44 side effects (e.g., methemoglobinemia) of triapine and its 45 unfavorable pharmacokinetic profile (e.g., short plasma halflife), 14 the development of novel TSCs with improved 46 pharmaceutical properties and an established mechanism of 47 action is of high research interest. Notably, two other TSCs, 48 namely, di-2-pyridylketone 4-cyclohexyl-4-methyl-3-thiosemi 49 carbazone (DpC) and 4-(2-pyridinyl)-2-(6,7-dihydro-8(5H)- $_{50}$ quinolinylidene)-hydrazide (COTI-2), are currently undergoing a phase I evaluation as chemotherapeutic agents. $_{8,15}^{8,15}$

The iron-containing ribonucleotide reductase (RNR) is $_{53}$ considered as one of the main targets for triapine and related $_{54}$ α -N-pyridinecarboxaldehyde TSCs. $_{16-19}$ This enzyme cata- $_{55}$ lyzes the reduction of ribonucleotides to deoxyribonucleotides, $_{56}$

Received: April 26, 2021



57 and it is particularly important in rapidly dividing cells, such as 58 tumor cells, virally infected cells, and invading bacteria. All 59 these cells share similar properties, such as high proliferation 60 rates, quickly spreading within the host, and aggressive disease 61 progression. 20 A sustained proliferation requires an increased 62 de novo nucleotide synthesis for DNA replication, making 63 RNR targeting a relevant strategy in the treatment of 64 cancer. 21,22 RNRs are free radical-containing proteins. One 65 way to control and modulate their reactivity is via quenching 66 the catalytically essential tyrosyl radical Y- located in the small 67 RNR subunit (R2 or NrdB). 23,24 The radical scavengers and 68 iron-chelating ligands, which are able to destroy the diferric-69 tyrosyl radical cofactor, with the aim to inhibit R2 RNR, are 70 widely investigated in anticancer research.²⁵ In the case of 71 triapine, it has been suggested that the intracellularly formed, 72 highly potent, redox-active iron complex either leads to 73 reactive oxygen species (ROS) formation, which are then 74 responsible for tyrosyl radical quenching, or that the iron 75 complex itself is able to directly reduce the tyrosyl radical. 16 76 Besides triapine, several other R2 RNR inhibitors such as 77 hydroxyurea, 3,4-dihydroxybenzohydroxamic acid (Didox), 78 and 3,4,5-trihydroxybenzamidoxime (Trimidox) have entered 79 clinical trials. 26 Among other potential tyrosyl radical 80 quenchers, p-alkoxyphenols (i.e., p-methoxyphenol, p-ethox-81 yphenol, p-propoxyphenol, and p-allyloxyphenol) and pyro-82 gallol as well as 4-mercaptophenol were identified.²⁷⁻ 83 mechanism of RNR inhibition by the p-alkoxyphenols and 84 pyrogallol was investigated by both experimental techniques 85 (electron paramagnetic resonance (EPR) and UV-visible 86 (UV-vis) spectroscopy) and theoretical tools (molecular 87 docking and molecular dynamics simulations). Among the 88 aminophenols several compounds were tested as anticancer 89 agents, for example, the nonsteroidal anti-inflammatory drug 90 N-acetyl-p-aminophenol (acetaminophen), which showed 91 antimelanoma activity to prooxidant glutathione (GSH) 92 depletion by the 3-hydroxy-1,4-quinone-imine-metabolite. ^{29,30} 93 Fenretinide (a synthetic retinoid derivative) was introduced in 94 clinical trials for the treatment of breast, bladder, renal, and 95 neuroblastoma malignancies due to its antioxidant activities via 96 scavenging radicals.

It is also worth noting that a coordination to copper(II) may significantly augment the cytotoxic activity of TSCs. ^{6,10} 99 Copper(II) as an essential trace element is redox-active, 100 biocompatible, and less toxic than nonendogenous heavy 101 metals. The redox metabolism of cancer cells is different from 102 that of healthy cells and is characterized by increased copper 103 levels in an intracellular environment. 32,33 Moreover, it was 104 recently suggested that the copper(II) TSC complexes, rather 105 than any metal-free TSCs or their cellular metabolites, are 106 responsible for the biological effects in vitro and in vivo. One 107 of the reasons for the increased antiproliferative activity of 108 copper(II) complexes of TSCs and the selectivity for cancer 109 cells is considered to be the redox cycling between two 110 oxidation states $(Cu^{2+} \leftrightarrow Cu^{+})$ in a biologically accessible 111 window of potentials (from -0.4 to +0.8 V vs normal 112 hydrogen electrode (NHE)) and ROS generation. 6,34 In this 113 context it is also remarkable that a copper-redox cycle 114 mechanism was found to be responsible for the oxidation of 115 phenolic compounds leading ultimately to reactive oxygen-116 dependent DNA damage.³⁵ The same authors suggested that 117 singlet oxygen or a singlet oxygen-like entity (e.g., a copper-118 peroxide complex) rather than the free hydroxyl radical plays a 119 role in DNA damage. 35 At the same time it is worth noting that

the idea that an efficient redox cycling of copper(II,I) 120 complexes with thiosemicarbazones can be involved in the 121 anticancer mechanism has been recently challenged 36 by 122 showing that the most resistant to reduction copper(II) 123 thiosemicarbazonates were the most cytotoxic. In addition, the 124 complexes can also dissociate fast, if the thiosemicarbazone has 125 different affinities to copper(II) and copper(I) and can lose the 126 competition for copper(I) to metallothioneins (MT) and 127 glutathione (GSH). 37

With this background in mind we aimed at (i) attachment of 129 a phenolic moiety at atom N4 of thiosemicarbazide, (ii) 130 investigation of solution speciation, complex formation 131 reactions of new TSCs with copper(II) in solution, and 132 synthesis of copper(II) complexes, (iii) investigation of the 133 reduction/oxidation of TSCs containing this potentially redox 134 active group, namely, the 4-aminophenolic unit, and copper- 135 (II) complexes thereof by electrochemical and spectroelec- 136 trochemical techniques and by using chemical oxidants, for 137 example, O2, p-benzoquinone (PBQ), 2,3-dichloro-5,6-dicya-138 no-1,4-benzoquinone (DDQ), and phenyliodine(III) diacetate 139 (PIDA), as two-electron/two proton acceptors and Ag₂O, 140 along with an analysis of the reversibility of the oxidation 141 process and the number of participating electrons, (iv) 142 identification of the effects of phenolic unit and coordination 143 to copper(II) on the redox activity and cytotoxicity in vitro as 144 well as on the mR2 RNR inhibition and estimation of their 145 potency to act as reductants for a tyrosyl radical with an 146 apparent redox potential of +1000 \pm 100 mV versus NHE.³⁸ 147

In this work we report on the synthesis of new triapine 148 derivatives HL^1-HL^3 , which contain a potentially redox-active 149 4-aminophenolic unit, and of copper(II) complexes $Cu(HL^1)$ - 150 Cl_2 (1), $[Cu(L^2)Cl]$ (2'), and $Cu(HL^3)Cl_2$ (3) (Chart 1). 151 cl

Chart 1. TSCs and Their Copper(II) Complexes Studied in This Work a

"Underlined labels/numbers indicate compounds studied by SC-XRD. The five-coordination of copper(II) in 1 and 3 has not been confirmed by X-ray crystallography.

The solution behavior of the new TSCs (HL^1-HL^3), the 152 mechanism typical for TSC E/Z isomerization, and the 153 stability and redox properties of both the metal-free ligands 154 and copper(II) complexes (1, 2', 3) were also investigated by 155 UV—vis spectrophotometry and UV—vis/EPR spectroelectro- 156 chemistry and density functional theory (DFT) calculations. In 157 addition, the two- and four-electron oxidation products $HL^{1a'}$ 158 and $HL^{1a''}$, respectively, were prepared both electrochemically 159

Chart 2. Oxidation Products of HL¹ and HL² and Copper(II) Complexes with Oxidized Ligands^a

1,3,4-Thiadiazole (TDA) core

1,2,4-Triazole-3-thione (TAT) core

Diphenol (DP) species

Benzo[d]thiazol-6-ol/one (BTA) core

OH

OH

$$R_1$$
 R_2
 $R_1 = H, R_2 = H$
 $R_1 = H, R_2 = Me$
 $R_1 = H, R_2 = Me$

Copper(II)-BTA complexes

"Underlined labels/numbers indicate compounds studied by SC-XRD, while the italic L denotes an oxidized ligand.

160 and by chemical oxidation and used in a complex formation 161 with copper(II). Several oxidation products of HL² (HL^{2b}, 162 HL2e, HL2c', and HL2c") were prepared by using different 163 oxidation agents. Likewise, copper(II) complexes with oxidized 164 ligands 4-6 were obtained (see Chart 2 and Scheme 1). The 165 isolated compounds were characterized by analytical and 166 spectroscopic methods (one-dimensional (1D) and two-167 dimensional (2D) NMR, UV-vis, IR), electrospray ionization 168 (ESI) mass spectrometry (MS), cyclic voltammetry (CV), and 169 single-crystal X-ray diffraction (SC-XRD). The anticancer 170 activity of the TSCs (HL1-HL3), their oxidized products 171 (HL^{1a'}, HL^{1a''}, and HL^{2c'}·CH₃COOH), and the copper(II) 172 complexes (1, 2', and 3) was tested against two human cancer 173 cell lines (doxorubicin-sensitive Colo205 and the multidrug-174 resistant Colo320 human colonic adenocarcinoma) and 175 normal human embryonal lung fibroblast cells (MRC-5) 176 along with their mR2 RNR inhibiting ability, and the results 177 are discussed.

■ EXPERIMENTAL SECTION

Chemicals. 2-Formylpyridine, 2-acetylpyridine, and CuCl₂·2H₂O 179 were purchased from commercial suppliers and used without further 180 purification. 3-(tert-Butoxycarbonyl)amino-2-formylpyridine and 4- 181 (4-hydroxy-3,5-dimethylphenyl)thiosemicarbazide were synthesized 182 as reported previously. 35,40 KCl, KOH, HCl, and dimethyl sulfoxide 183 (DMSO) were obtained from Reanal. GSH, 2-morpholinoethane- 184 sulfonic acid (MES), and 2-[4-(2-hydroxyethyl)piperazin-1-yl]- 185 ethanesulfonic acid (HEPES) were purchased from Sigma-Aldrich 186 and used without further purification. Copper(II) stock solution was 187 prepared by the dissolution of CuCl₂ in water, and its concentration 188 was determined by complexometry with ethylenediaminetetraacetic 189 acid (EDTA). The stock solutions of HL¹-HL³ were prepared on a 190 weight-in-volume basis dissolved in DMSO.

2-Formylpyridine 4-(4-hydroxy-3,5-dimethylphenyl)- 192 thiosemicarbazone (HL¹·0.5H₂O). 2-Formylpyridine (0.09 mL, 193 0.95 mmol) was added to 4-(4-hydroxy-3,5-dimethylphenyl)- 194 thiosemicarbazide (200 mg, 0.95 mmol) in ethanol (12 mL), heated 195 at 85 °C for 2 h, concentrated, and left for crystallization at 4 °C. The 196

c2s1

178

Scheme 1. Oxidation Products of HL¹ and HL² along with Those of Copper(II) Complexes^a

$$\begin{array}{c} C_{uCl_{2}\cdot 2H_{2}O} \\ H_{2}^{N} \\ N_{N} \\ N_{N$$

^aThe bottom left panel shows the oxidants used.

197 yellow solid was filtered off, washed with cold ethanol, and dried in 198 vacuo. Yield: 253 mg, 86.1%. Anal. Calcd for C₁₅H₁₆N₄OS·0.5H₂O 199 ($M_r = 309.39$): C, 58.23; H, 5.54.; N, 18.11; S, 10.36; Found: C, 200 57.91; H, 5.45; N, 17.92; S, 10.43%. Positive ion ESI-MS for 201 $C_{15}H_{16}N_4OS$ (MeCN/MeOH+1% H_2O): m/z 301.11 $[HL^1+H]^+$, 202 323.09 [HL¹+Na]⁺, 339.07 [HL¹+K]⁺, negative ion ESI-MS: m/z203 299.10 $[HL^1-H]^{-1}$. ¹H NMR (600 MHz, DMSO- d_6 , E isomer) δ , 204 ppm: 11.86 (s, 1H, H₉), 10.00 (s, 1H, H₁₁), 8.57 (d, J = 4.4 Hz, 1H, 205 H_6), 8.43 (d, J = 8.0 Hz, 1H, H_3), 8.22 (s, 1H, H_{18}), 8.16 (s, 1H, H_7), 206 7.82 (td, J = 7.8, 1.2 Hz, 1H, H₄), 7.37 (m, 1H, H₅), 7.02 (s, 2H, ²⁰⁷ H₁₃+H₁₇), ^{2.17} (s, 6H, H₁₉+H₂₀). ¹³C NMR (151 MHz, DMSO-d₆, E 208 isomer) δ , ppm: 176.55 (C₁₀), 153.31 (C₂), 151.10 (C₁₅), 149.27 209 (C_6), 142.51 (C_7), 136.43 (C_4), 130.18 (C_{12}), 126.26 ($C_{13}+C_{17}$), 210 124.10 (C_5), 123.84 ($C_{14}+C_{16}$), 120.54 (C_3), 16.62 ($C_{19}+C_{20}$). ¹⁵N 211 NMR (61 MHz, DMSO- d_6 , E isomer) δ , ppm: 325.04 (N₈), 315.07 212 (N_1) , 174.22 (N_9) , 128.93 (N_{11}) . IR (attenuated total reflectance

(ATR), selected bands, $\tilde{\nu}_{\text{max}}$): 3107.39, 2950.74, 1531.05, 1477.88, 213 1428.74, 1201.82, 1105.17, 926.54, 862.73, 761.37, 682.25 cm⁻¹. 214 UV–Vis (MeOH), λ_{max} , nm (ε , M⁻¹ cm⁻¹): 243 sh, 328 (3516). 215 Single crystals of $\text{HL}^1 \cdot \text{C}_2 \text{H}_5 \text{OH}$ suitable for X-ray data collection were 216 obtained from the mother liquor.

2-Acetylpyridine 4-(4-hydroxy-3,5-dimethylphenyl)- 218 **thiosemicarbazone** (HL²·0.2H₂O). 2-Acetylpyridine (0.21 mL, 219 1.91 mmol) was added to 4-(4-hydroxy-3,5-dimethylphenyl)- 220 thiosemicarbazide (269 mg; 1.27 mmol) in ethanol (8 mL), heated 221 at 85 °C overnight, concentrated, and left for crystallization at 4 °C. 222 The obtained light yellow precipitate was filtered off, washed with 223 cold ethanol, and dried in vacuo. Yield: 271 mg, 67.0%. Anal. Calcd 224 for $C_{16}H_{18}N_4OS\cdot0.2H_2O$ (M_r = 318.01): C, 60.43; H, 5.83; N, 17.62; 225 S, 10.08. Found: C, 60.47; H, 5.8; N, 17.55; S, 10.13%. Positive ion 226 ESI-MS for $C_{16}H_{18}N_4OS$ (M_r = 314.41) (MeCN/MeOH+1% H_2O): 227 m/z 315.13 [HL²+H]⁺, 337.11 [HL²+Na]⁺, negative ion ESI-MS: m/z 228

229 *z* 313.11 [HL²-H]⁻. ¹H NMR (600 MHz, DMSO- d_6 , *E* isomer) δ, 230 ppm: 10.46 (s, 1H, H₉), 9.94 (s, 1H, H₁₁), 8.59 (d, *J* = 4.7 Hz, 1H, 231 H₆), 8.54 (d, *J* = 8.1 Hz, 1H, H₃), 8.22 (s, 1H, H₁₈), 7.79 (td, *J* = 7.8, 232 1.7 Hz, 1H, H₄), 7.39 (dd, *J* = 7.2, 4.9 Hz, 1H, H₅), 7.02 (s, 2H, 233 H₁₃+H₁₇), 2.44 (s, 3H, H₇), 2.17 (s, 6H, H₁₉+H₂₀). ¹³C NMR (151 234 MHz, DMSO- d_6 , *E* isomer) δ, ppm: 177.36 (C₁₀), 154.59 (C₂), 235 151.12 (C₁₅), 148.54 (C₇), 148.43 (C₆), 136.34 (C₄), 130.36 (C₁₂), 236 126.33 (C₁₃+C₁₇), 124.00 (C₅), 123.83 (C₁₄+C₁₆), 121.18 (C₃), 237 16.63 (C₁₉+C₂₀), 12.31 (C₇·). ¹⁵N NMR (61 MHz, DMSO- d_6 , *E* 238 isomer) δ, ppm: 312.94 (N₈), 310.61 (N₁), 168.53 (N₉), 129.34 239 (N₁₁). IR (ATR, selected bands, $\tilde{\nu}_{max}$) (cm⁻¹): 3386.87, 3187.76, 240 1531.57, 1478.45, 1309.19, 1182.40, 1032.57, 942.48, 778.97, 652.93. 241 UV−Vis (MeOH), λ_{max} nm (ε , M⁻¹ cm⁻¹): 316 (2842), 407 sh. 242 Single crystals of **HL²** suitable for X-ray data collection were obtained 243 from the mother liquor.

3-Amino-2-formylpyridine 4-(4-hydroxy-3,5-245 dimethylphenyl)thiosemicarbazone (HL³·0.25H₂O). To a sol-246 ution of 3-(tert-butoxycarbonyl)amino-2-formylpyridine (210 mg, 247 0.95 mmol) and 4-(4-hydroxy-3,5-dimethylphenyl)thiosemicarbazide 248 (200 mg, 0.95 mmol) in a mixture of ethanol/water 3:1 (8 mL) was 249 added dropwise 12 M HCl (0.19 mL, 2.28 mmol). This solution was 250 stirred at room temperature for 1 h to give Boc-HL3·HCl 251 ($C_{20}H_{25}N_5O_3S \cdot HCl$, positive ion ESI-MS for $C_{20}H_{25}N_5O_3S$ ($M_r =$ 252 415.51) (MeCN/MeOH+1% H_2O): m/z 416.18 [Boc-HL³+H]⁺, 253 negative ion ESI-MS: m/z 414.02 [Boc-HL³-H]⁻). The Boc-254 deprotection of HL³ was completed at 85 °C for 7 h with monitoring by ESI-MS (positive ion ESI-MS for $C_{15}H_{17}N_5OS$ ($M_r = 315.39$) (MeCN/MeOH + 1% H₂O): m/z 316.12 [HL³+H]⁺, 338.11257 $[HL^3+Na]^+$, negative ion ESI-MS: m/z 314.11 $[HL^3-H]^-$). After 258 ethanol evaporation, the solution was neutralized with a saturated 259 solution of NaHCO₃ (pH = 8). The precipitate was collected and 260 dried in vacuo. Yield: 267 mg, 87.9%. Anal. Calcd for C₁₅H₁₇N₅OS· $261\ 0.25H_2O\ (M_r=319.90)$: C, 56.31; H, 5.51; N, 21.89; S, 10.02. ²⁶² Found: C, 56.33; H, 5.34; N, 21.68; S, 10.29%. ¹H NMR (600 MHz, 263 DMSO- d_6 , E isomer) δ , ppm: 11.47 (s, 1H, H₉), 9.70 (s, 1H, H₁₁), 264 8.39 (s, 1H, H₇), 8.21 (s, 1H, H₁₈), 7.85 (dd, J = 4.3, 1.4 Hz, 1H, H₆), 265 7.15 (dd, J = 8.3, 1.2 Hz, 1H, H₄), 7.08 (dd, J = 8.3, 4.3 Hz, 1H, H₅), 266 6.92 (s, 2H, H₁₃+H₁₇), 6.49 (s, 2H, H₃), 2.16 (s, 6H, H₁₉+H₂₀). ¹³C 267 NMR (151 MHz, DMSO- d_6 , E isomer) δ , ppm: 176.13 (C_{10}), 151.17 268 (C₁₅), 149.23 (C₇), 143.99 (C₃), 137.25 (C₆), 132.97 (C₂), 130.59 269 (C_{12}), 126.88 ($C_{13}+C_{17}$), 124.52 (C_{5}), 123.83 ($C_{14}+C_{16}$), 122.34 270 (C_{4}), 16.63 ($C_{19}+C_{20}$). ¹⁵N NMR (61 MHz, DMSO- d_{6} , E isomer) δ , 271 ppm: 321.53 (N₁), 312.8 (N₈), 174.57 (N₉), 126.69 (N₁₁), 71.10 272 (N_{3'}). IR (ATR, selected bands, $\tilde{\nu}_{max}$ cm⁻¹): 3456.59, 3347.73, 273 3142.99, 3002.80, 1615.50, 1547.68, 1512.07, 1299.63, 1248.47, 274 1189.77, 1143.84, 861.56, 796.22, 685.36. UV–Vis (MeOH), λ_{max} 275 nm (ε , M⁻¹ cm⁻¹): 299 (1374), 375 (2220), 448 sh. Single crystals of 276 HL³ suitable for X-ray data collection were obtained from the mother 277 liquor.

278 SYNTHESIS OF THE COPPER(II) COMPLEXES

 $Cu(HL^{1})Cl_{2}\cdot 0.5H_{2}O$ (1·0.5H₂O). $CuCl_{2}\cdot 2H_{2}O$ (128 mg, 280 0.75 mmol) was added to HL¹ (225 mg, 0.75 mmol) in anoxic 281 methanol (10 mL) in a Schlenk tube and stirred at room 282 temperature under argon for 10 min. The reaction mixture was 283 allowed to stand at 4 °C overnight. The dark green precipitate 284 was filtered off under argon, washed with anoxic methanol, and dried in vacuo. Yield: 294 mg, 88.4%. Anal. Calcd for 286 $C_{15}H_{16}N_4OSCuCl_2\cdot 0.5H_2O$ ($M_r = 443.84$): C, 40.59; H, 287 3.86; N, 12.62; S, 7.22. Found: C, 40.73; H, 3.59; N, 12.63; S, 288 7.19%. Positive ion ESI-MS for C₁₅H₁₆N₄OSCuCl₂ (MeCN/ 289 MeOH+1% H₂O): m/z 362.03 [Cu(HL¹)²⁺-H]⁺, negative ion 290 ESI-MS: m/z 395.99 [Cu(HL¹)Cl⁺-2H]⁻. IR (ATR, selected 291 bands, $\tilde{\nu}_{\text{max}}$): 3480.77, 2989.07, 1610.63, 1479.59, 1269.25, ²⁹² 1229.98, 1189.75, 1025.69, 774.69, 665.85 cm⁻¹. UV–Vis 293 (MeOH), λ_{max} , nm (ε , M⁻¹ cm⁻¹): 280 (16 800), 376 sh, 422 294 (18 160). Crystals of $[Cu(L^1)Cl]\cdot CH_3OH$ (1'·CH₃OH) ($M_r =$ 295 398.37) suitable for X-ray diffraction study were grown from an \sim 20-fold-diluted reaction mixture in a Schlenk tube under 296 argon upon standing at 4 °C. A recrystallization of [Cu(HL¹)- 297 Cl₂] (1) in methanol in air afforded a minor amount of X-ray 298 diffraction-quality crystals of [Cu(L¹c′)Cl] (4).

 $[Cu(L^2)Cl] \cdot 0.5H_2O$ (2'·0.5H₂O). CuCl₂·2H₂O (129 mg, 300 0.76 mmol) was added to a solution of HL² (238 mg, 0.76 301 mmol) in anoxic methanol (10 mL) in a Schlenk tube. The 302 reaction mixture was stirred at room temperature under argon 303 for 10 min and then allowed to stand at 4 °C overnight. The 304 greenish-brown precipitate was filtered off under argon, 305 washed with anoxic methanol, and dried in vacuo. Yield: 316 306 mg, 98.8%. Anal. Calcd for $C_{16}H_{17}N_4OSCuCl\cdot 0.5H_2O$ ($M_r = 307$ 421.40): C, 45.60; H, 4.31; N, 13.30; S, 7.61. Found: C, 45.74; 308 H, 4.03; N, 13.42; S, 7.56%. Positive ion ESI-MS for 309 $C_{16}H_{17}N_4OSCuCl$ (MeCN/MeOH+1% H_2O): m/z 376.04 310 $[Cu(L^2)]^+$, negative ion ESI-MS: m/z 410.00 $[Cu(L^2)Cl-H]^-$. 311 IR (ATR, selected bands, $\tilde{\nu}_{max}$): 3341.84, 3223.12, 1609.18, 312 1547.35, 1483.22, 1452.56, 1303.41, 1202.82, 1019.61, 846.14, 313 701.29 cm⁻¹. UV–Vis (MeOH), λ_{max} , nm (ε , M⁻¹ cm⁻¹): 277 314 (11 835), 316 sh, 421 (12 953). Crystals of $[Cu(L^2)Cl]$ (2') 315 suitable for X-ray diffraction study were obtained from an ~20- 316 fold-diluted reaction mixture under argon in a Schlenk tube at 317 4 °C.

Cu(HL³)Cl₂·0.25H₂O (3·0.25H₂O). CuCl₂·2H₂O (114 mg, 319 0.67 mmol) was added to HL³ (210 mg, 0.67 mmol) in anoxic 320 methanol (10 mL) in a Schlenk tube and stirred at room 321 temperature under argon for 10 min. The reaction mixture was 322 allowed to stand at 4 °C overnight. The green precipitate was 323 filtered off under argon, washed with anoxic methanol, and 324 dried in vacuo. Yield: 285 mg, 93.6%. Anal. Calcd for 325 $C_{15}H_{17}N_5OSCuCl_2\cdot 0.25H_2O$ ($M_r = 454.35$): C, 39.65; H, 326 3.88; N, 15.41; S, 7.06. Found: C, 39.58; H, 3.79; N, 15.21; S, 327 6.98%. Positive ion ESI-MS for C₁₅H₁₇N₅OSCuCl₂ (MeCN/ 328 MeOH+1% H₂O): m/z 377.04 [Cu(HL³)²⁺-H]⁺, negative ion 329 ESI-MS: m/z 411.00 [Cu(HL³)Cl⁺-2H]⁻. IR (ATR, selected 330 bands, $\tilde{\nu}_{\text{max}}$): 3422.07, 3340.63, 1647.85, 1569.29, 1480.67, 331 1223.63, 1185.74, 1023.07, 718.76, 660.61 cm⁻¹. UV-Vis 332 (MeOH), λ_{max} , nm (ε , M⁻¹ cm⁻¹): 262 (19 564), 288 333 (17 425), 462 (23 514). Crystals of [Cu(L3)Cl]·CH3OH, 334 $(3' \cdot CH_3OH)$ ($M_r = 413.38$) suitable for an X-ray diffraction 335 study were grown from an ~20-fold-diluted reaction mixture in 336 a Schlenk tube under argon at 4 °C.

Details about the synthesis and characterization of oxidized 338 thiosemicarbazones and their copper(II) complexes, X-ray data 339 collection and refinement (Tables S1—S3), elemental analysis, 340 UV—vis titrations, kinetic measurements, lipophilicity deter- 341 mination, spectroelectrochemical studies, in vitro cell studies, 342 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl-tetrazolium bromide 343 (MTT) assays, and tyrosyl radical reduction in mouse R2 RNR 344 protein as well as computational details are given in the 345 Supporting Information (Sections 1 and 2).

■ RESULTS AND DISCUSSION

Synthesis and Characterization of HL¹-HL³. The new 348 TSCs HL¹-HL³ were obtained by Schiff base condensation 349 reactions of 4-(4-hydroxy-3,5-dimethylphenyl)- 350 thiosemicarbazide 40 with the corresponding aldehyde (HL¹, 351 HL³) or ketone (HL²) in boiling ethanol (HL¹, HL²) or 352 ethanol/water (3:1, HL³) in the absence (HL¹ and HL²) or in 353 the presence of 12 M HCl (HL³). The hydrochloric acid in 354 this latter case was used for Boc-deprotection of the 355 intermediate Boc-HL³. This deprotection reaction was 356 monitored by ESI-MS (disappearance of peaks attributed to 357

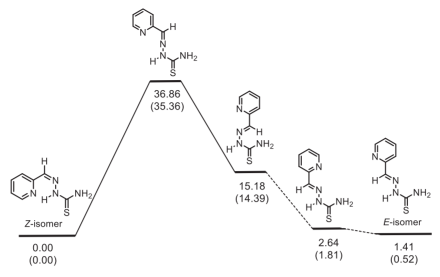


Figure 1. Electronic energy and Gibbs free energy profiles (in kcal/mol) for the transformation of the most stable conformer of (Z)-2-formylpyridine thiosemicarbazone in DMSO solution. Free energies (in parentheses) at 298 K and 1 atm.

358 [Boc-HL³+H]⁺ and [Boc-HL³-H]⁻ ions) and completed at 85 359 °C after 7 h, with yields ranging from 67 to 88%. The 360 formation of HL¹-HL³ was confirmed by ESI mass spectra, 361 which showed peaks assigned to ions [HL1-3+H]+, 362 [HL¹⁻³+Na]⁺, and [HL¹⁻³-H]⁻. One- and two-dimensional 363 NMR spectra were in agreement with the expected structures 364 for HL^1-HL^3 of C_1 molecular symmetry. In addition, the 365 spectra indicated the presence of E and Z isomers in DMSO-366 d_6 , which is typical for thiosemicarbazones, ^{41–43} with a 367 significant predominance of E isomers (E/Z = 23:1, 17:1, 368 and 31:1 for HL¹-HL³, respectively). The assignment of E 369 and Z isomers was based on NMR spectra, including ¹H, ¹H 370 nuclear Overhauser effect spectroscopy (NOESY), which are 371 presented in more detail in the Supporting Information (see 372 also Schemes S1 and S2 and Tables S4-S6). It is noteworthy 373 that, in contrast to the E isomers of HL^1-HL^3 , their Z isomers 374 can form an intramolecular hydrogen bond between the 375 pyridine nitrogen and the NH-N group hydrogen, resulting in 376 an increase in the relative stability of these conformers. Indeed, 377 the DFT B3LYP/6-311++G (d,p) calculations for E- and Z-378 HL¹ in a DMSO solution (the polarizable continuum model 379 (PCM) solvation model) showed that the most stable 380 conformer of Z-HL¹ lies lower in energy than the most stable 381 conformer of E-HL¹ ($\Delta E = 1.45 \text{ kcal/mol}$; $\Delta G = 0.76 \text{ kcal/mol}$ 382 mol at 298 K and 1 atm). The calculations also demonstrate 383 that E- and Z-HL² are very close in thermodynamic stability 384 ($\Delta E = 0.90 \text{ kcal/mol}$ in favor of Z-HL², $\Delta G = 0.00 \text{ kcal/mol}$), and E-HL³ is slightly more stable than Z-HL³ ($\Delta E = 0.84 \text{ kcal/}$ 386 mol, $\Delta G = 0.86$ kcal/mol), which can be explained by the 387 presence of an intramolecular hydrogen bond between the 3-388 NH₂ group and the aldimine nitrogen in E-HL³. Thus, the 389 formation of HL¹-HL³ with a large predominance of the E 390 isomers indicates that the reactions proceed under a kinetic 391 control. By using DFT B3LYP/6-311++G(d,p) calculations to 392 understand the interconversion between E and Z isomers of 2-393 formylpyridine and thiosemicarbazones as model compounds 394 we found out that an isomerization involving a tautomeric shift 395 of the thioamide N2H proton to the pyridine nitrogen 396 followed by a rotation around the formed C-N1 bond, as 397 proposed previously, 44 is not favored energetically (see the 398 Supporting Information for details). We believe that the most plausible Z/E isomerization pathway in thiosemicarbazones 399 and semicarbazones involves an inversion at the imine 400 nitrogen. The intrinsic reaction coordinate (IRC) analysis 401 for one of the aforementioned model compounds revealed that 402 the found transition state connects the desired minima. 403 However, the calculation data obtained show (for more details 404 see the Supporting Information) that the Gibbs free energy 405 barrier for the conversion of the most stable conformer of the 406 Z isomer into the E isomer is relatively high ($\Delta G = 35.2 \text{ kcal}/407 \text{ mol}$ in the gas phase, 35.4 kcal/mol in DMSO solution) 408 (Figure 1), which rejects the possibility of an interconversion 409 ft between the isomers at room temperature.

The redox activity of HL^1 – HL^3 in the anodic region was 411 validated by cyclic voltammetry (vide infra). Their behavior as 412 reductants is also relevant for quenching the tyrosyl radical in 413 the mR2-protein. Therefore, attempts to perform an oxidation 414 of HL^1 and HL^2 by electrolysis and by chemical oxidation were 415 undertaken.

Oxidation of TSCs. The oxidation of different organic 417 molecules with p-benzoquinone derivatives is well-documented 418 in the literature. ⁴⁶ The reaction of HL^1 with DDQ ($2e^-/2H^+$ 419 E° = 0.887 V vs NHE in an acidic 0.1 M aqueous solution of p- 420 TsOH) ⁴⁷ in a 1:1 molar ratio resulted in two-electron and 421 four-electron oxidative cyclizations with the major formation of 422 $HL^{1a'}$ (60.9%) accompanied by a minor generation of $HL^{1a''}$ 423 (<5%), both containing a 1,3,4-thiadiazole ring (Chart 2, 424 Scheme 1). The formation of the 1,3,4-thiadiazole ring occurs 425 via a nucleophilic attack of the sulfur atom to the carbon atom 426 of the aldimine bond of HL^1 as evidenced by frontier 427 molecular orbitals with the highest occupied molecular orbital 428 (HOMO) and lowest unoccupied molecular orbital (LUMO) 429 located at opposite sides of the molecule (Figure 2).

The use of a double amount of DDQ led to the formation of 431 the four-electron oxidation product $HL^{1a''}$ in 71.6% yield. The 432 electrolysis of HL^1 at 1000 mV in CH_3CN versus Ag/AgCl 433 resulted in the same oxidation products (vide infra). Both 434 compounds were characterized by ESI mass spectra, which 435 showed peaks at m/z 299.17 $[HL^{1a'}+H]^+$, 321.16 436 $[HL^{1a'}+Na]^+$, 297.18 $[HL^{1a''}+H]^+$, 319.20 $[HL^{1a''}+Na]^+$, and 437 296.94 $[HL^{1a'}-H]^-$. The more sterically hindered ketimine 438 carbon atom in 439 was expected to reduce the likelihood of 439

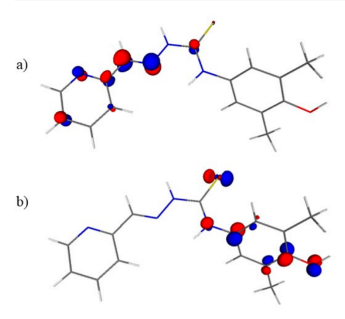


Figure 2. Frontier orbitals in HL^1 : (a) LUMO and (b) HOMO drawn at 0.1 au isosurface.

440 the 1,3,4-thiadiazole ring formation. The reaction of HL² with 441 DDO in a 1:1 molar ratio in methanol led to a decomposition 442 of the TSC with a formation of an unidentified species. When 443 PBQ, a weaker oxidant $(2e^{-}/2H^{+} E^{\circ} = 0.643 \text{ V vs NHE in an})$ 444 acidic 0.1 M aqueous solution of p-TsOH) than DDQ, was 445 used, 47 a two-electron oxidative cyclization with the formation 446 of a 1,2,4-triazole-3-thione ring (TAT group, HL2b) occurred, 447 accompanied by a desulfurization of HL² and a conversion into 448 a diphenolic species HL^{2e} (DP group). 48 The formation of 449 HL^{2b} was confirmed by ESI mass spectra, where peaks 450 corresponding to $[HL^{2b}+H]^+$ (m/z 313.25), $[HL^{2b}+Na]^+$ (m/z451 335.14), and $[HL^{2b}-H]^{-}$ (m/z 310.99) were present. We 452 suppose that the initial step of the reaction of HL² with PBQ 453 involves a one-electron oxidation of HL² favored by the 454 character of the HOMO of HL2 (see Figure S1) along with a 455 NH deprotonation to give a highly conjugated N/S-centered 456 free radical (see Scheme S8 in Supporting Information). This 457 radical intermediate transforms into triazole HL^{2b} in two steps 458 or undergoes a fragmentation affording 4-isothiocyanato-2,6-459 dimethylphenol. The phenol reacts with HL2 via an SE2 460 mechanism to form the corresponding thioamide followed by a 461 radical-promoted intermolecular transformation into indole 462 HL^{2e} according to a Fukuyama-like indole synthesis⁴⁹ (for a 463 more detailed discussion of the oxidation of HL² with PBQ see 464 the Supporting Information).

Other oxidation agents (lead tetraacetate, phenyliodine(III) diacetate (PIDA) with $E^{\circ} = +1.70 \text{ V}$ vs Fc/Fc⁺ in ACN, and for silver(I) oxide) for N-alkyl(aryl)-aminocarbonyl-4-aminophedes nols, were also used in an attempt to obtain the desired oxidation products with a 1,4-benzoquinone imine moiety (see also Scheme S3, its accompanying explanation, and Figure S2 in the Supporting Information). The exposure of HL² to 1 equiv of PIDA furnished the two-electron oxidized product $HL^{2c'}$ and traces of the four-electron oxidized species $HL^{2c''}$. As for $HL^{1a'}$ and $HL^{1a''}$, the use of a double amount of oxidant resulted in $HL^{2c''}$ as the main oxidation product. ESI mass spectra showed peaks at m/z 313.21, 310.98 attributed to $HL^{2c''}+H$ the $HL^{2c''}+H$ as well as 311.12, 309.01 assigned to $HL^{2c''}+H$ in $HL^{2c''}-H$ in line with the loss of two $HL^{2c'}$

or four ($HL^{2c''}$) protons when compared to original TSC HL^2 479 (315.13 [HL^2+H]⁺, 313.11 [HL^2-H]⁻).

Characterization of Oxidized Organic Compounds by 481 NMR Spectroscopy. The formation of a 1,3,4-thiadiazole- 482 ring in $HL^{1a'}$ and $HL^{1a''}$ by an oxidation of HL^1 resulted in the 483 disappearance of peaks of the aldimine CH proton (H₇) and 484 NH (H₉) in $HL^{1a'}$ and $HL^{1a''}$ as well as of the signal of NH 485 (H₁₁) in $HL^{1a''}$. The formation of a 1,4-benzoquinone imine 486 moiety in $HL^{1a\prime\prime}$ was confirmed also by the absence of the OH 487 signal, which resonates at 8.08-8.22 ppm in HL1-HL3, HL1a' 488 (see Scheme S4 and Tables S4-S6 in the Supporting 489 Information). The ring-closure reaction resulted in a downfield 490 shift of the resonance signal of carbon C7, which was directly 491 involved in the 1,3,4-thiadiazole ring formation. The 492 quaternary carbon C_7 in $HL^{1a\prime}$ and $HL^{1a\prime\prime}$ resonates at 493 158.40 and 169.98 ppm, respectively, whereas the aldimine CH 494 carbon atom C₇ in HL¹ resonates at 142.51 ppm. Analogously, 495 the involvement of the sulfur atom in the 1,3,4-thiadiazole ring 496 led to a downfield shift of the signal of the carbon atom C₁₀ 497 (C=S) to 166.77 ppm in $HL^{1a'}$ and to 171.58 ppm in $HL^{1a''}$ 498 when compared to 176.55 ppm in HL¹.

The four-electron oxidation of $\mathbf{HL^{1}}$ to $\mathbf{HL^{1a''}}$ with the 500 formation of the imine N(11) = C(C12) bond resulted in 501 strong downfield shift of the resonance signal of carbon C_{12} of 502 1,4-benzoquinone moiety of $\mathbf{HL^{1a''}}$ (162.21 ppm) when 503 compared to that of carbon C_{12} of phenolic moiety in $\mathbf{HL^{1}}$ 504 $\mathbf{HL^{3}}$, $\mathbf{HL^{1a'}}$ (130.18–132.53 ppm). In addition, the formation 505 of the carbonyl C(15) = O(18) bond in $\mathbf{HL^{1a''}}$ has a strong 506 effect on the resonance of carbon atom C_{15} , which is strongly 507 downfield-shifted to 187.14 ppm when compared to that in 508 $\mathbf{HL^{1}} - \mathbf{HL^{3}}$ and $\mathbf{HL^{1a''}}$ at 148.97–151.17 ppm. Remarkable 509 shifts of resonance signals for other atoms of the 1,4-510 benzoquinone moiety in $\mathbf{HL^{1a''}}$ in comparison to the phenolic 511 moiety in $\mathbf{HL^{1}} - \mathbf{HL^{3}}$ and $\mathbf{HL^{1a''}}$ were also noticed (see the 512 Supporting Information and Scheme S5 therein).

The formation of the benzothiazole ring in $HL^{2c\prime}$ is 514 evidenced by the presence in the ¹H NMR spectrum of one 515 singlet of the CH group and two singlets of methyl groups of 516 an unsymmetrical phenolic moiety with the intensity ratio of 517 1:3:3 as well as by one NH signal at 11.76 ppm in comparison 518 with a number of signals in the spectrum of HL^2 (1(NH)/ 519 $1(NH)/2(CH)/6(CH_3)$). Of the two proposed tautomers for 520 $HL^{2c'}$ (A (N(11)H) and B (N(9)H); see Scheme S6 in the 521 Supporting Information) the formation of the *E* isomer of form 522 **B** in DMSO- d_6 was evidenced by the cross-peak between 523 protons of methyl (H₇) and NH (H₉) groups in the ¹H, ¹H 524 NOESY spectrum. The DFT B3LYP/6-311++G(d,p) calcu- 525 lations showed that the E isomer of tautomer A is less stable 526 than the E isomer of tautomer **B** in a DMSO solution ($\Delta E = 527$ 1.58 kcal/mol; $\Delta G = 1.01$ kcal/mol at 298 K and 1 atm). We 528 found that, in contrast to HL^1-HL^3 , the E/Z isomerization 529 was observed for HL^{2c'}. As expected in case of HL^{2c'} 530 CH₃COOH, where nitrogen atom N₁ of the pyridine ring is 531 protonated and prevents the hydrogen-bond formation 532 between H_9 and N_1 , which is present in the Z isomer of 533 $HL^{2c'}$, only one set of signals attributed to the E isomer was 534 found. The neutral species HL^{2c'} in DMSO-d₆ and MeOH-d₄ 535 is present as the E isomer, which converts slowly into the Z 536 isomer. The process is solvent-dependent. The E/Z equili- 537 brium was reached in 6 d with a molar ratio of E/Z isomers of 538 7.2:1 (DMSO- d_6) and 3:1 (MeOH- d_4) (see Figure S3 in the 539 Supporting Information). The Z isomer of HL^{2c} in DMSO- d_6 540 is characterized by the downfield-shifted proton NH(9) due to 541

542 the hydrogen bond to the pyridine nitrogen atom and 543 resonates at 15.00 ppm (the same proton of the E isomer of 544 $HL^{2c'}$ is seen at 11.58 ppm). The Z/E isomerization of $HL^{2c'}$ 545 was also studied in MeOH-d₄ and methanol by ¹H NMR and 546 UV-vis spectroscopy reaching 1:3.6 molar ratio in 14 d 547 according to NMR spectra (for optical spectra difference see 548 Figure S4). The carbon atom of the methyl group $(C_{7'})$ in the 549 E isomers of HL^{2c'}•CH₃COOH and HL^{2c'} resonates at 12.55 sso and 12.56 ppm, respectively, whereas in the Z isomer of HL^{2c} 551 it resonates at 21.72 ppm. Note that these chemical shifts are 552 consistent with those calculated for E- and Z-HL^{2c} (8.29 and 553 23.26 ppm, respectively) by the gauge-independent atomic 554 orbital (GIAO) method at the WC04/6-311+G(2d,p) level of sss theory using the DFT B3LYP/6-311++G(d,p) optimized 556 geometries (DMSO solution, the PCM solvation model). A 557 similar difference in chemical shifts of the CH₃ group was also observed for the E (12.31 ppm) and Z isomers (21.73 ppm) of 559 HL^2 . The DFT calculation also demonstrated that E and Z 560 isomers of HL^{2c'} have a quite similar stability in a DMSO solution ($\Delta G = 0.11 \text{ kcal/mol}$ in favor of the *E* isomer; 298 K, $_{562}$ 1 atm). As expected, the pyridine ring carbon atom C_3 is also 563 sensitive to the hydrogen-bond formation between H₉ and N₁ 564 in the Z isomer of $HL^{2c'}$. The C_3 signal in the latter is 565 markedly shifted (124.08 ppm) in comparison to C₃ in the E 566 isomer (119.65 ppm). A full assignment of resonances was 567 possible only for HL2c'·CH3COOH (the three quaternary 568 carbons C₁₂, C₇, and C₁₇ were identified according to ¹H, ¹³C 569 HMBC; see Figure S5 in the Supporting Information).

The two-electron oxidation of $HL^{2c'}$ to $HL^{2c''}$ with the formation of the quinone moiety is accompanied by the downfield shift of the resonance signal of carbon C_{15} at 184.43 ppm in comparison to that of C_{15} in $HL^{2c'}$ ·CH₃COOH at 148.14 ppm, in E- $HL^{2c'}$ at 148.15 ppm, and in E- $HL^{2c'}$ at 148.39 ppm. The lack of the NH signal confirms the formation of the imine N(9)=C(C10) bond (see Scheme S7 and Tables S77 S4 and S5 in the Supporting Information).

Synthesis and Characterization of Copper(II) Com-579 **plexes.** The reaction of HL^1-HL^3 with $CuCl_2\cdot 2H_2O$ in 580 anoxic methanol under an argon atmosphere to preclude an 581 eventual oxidation of the ligands by air oxygen in a 1:1 molar 582 ratio at room temperature afforded green-brown solids of the formulas $Cu(HL^1)Cl_2(1)$, $[Cu(L^2)Cl](2')$, and $Cu(HL^3)Cl_2(1)$ 584 (3) in almost quantitative yields. The formation of these 585 copper(II) complexes was confirmed by elemental analyses 586 and ESI mass spectra. The latter showed peaks attributed to 587 $[Cu(L^{1,3})-H]^+$, $[Cu(L^{1,3})Cl-H]^-$, or $[Cu(L^2)]^+$ and $[Cu-L^2]^+$ 588 (L²)Cl-H]⁻. XRD-quality single crystals of $[Cu(L^{1-3})Cl]$ 589 (1'-3') were grown from diluted by a factor of ca. 20 reaction 590 mixtures under argon upon standing at 4 °C. Under these 591 conditions the deprotonation of ligands HL1 and HL3 592 occurred. Attempts to crystallize 1, 2', and 3 in air failed, 593 most likely because of an occurring oxidation of complexes by 594 O₂.

Synthesis of the Copper(II) Complexes with Oxidized 596 Ligands. Upon a prolonged standing of a methanolic solution 597 of Cu(HL¹)Cl₂ (1) in air, a minor amount of crystals of 598 [Cu(L¹c)Cl] (4) formed, in which the ligand underwent an 599 oxidative dehydrogenation along with the intramolecular 600 cyclization via a C-S coupling reaction between phenolic 601 carbon and thione group into a five-membered thiazole ring, as 602 confirmed by SC-XRD (vide infra). Some rare examples of 603 thiosemicarbazone cyclization with the benzothiazole ring 604 formation due to a coordination to copper(II) were recently

reported. S2,53 A direct complex formation reaction between the 605 prepared benzo [d] thiazol-6-ol $HL^{2c'}$ and copper (II) chloride 606 produced $[Cu(HL^{2c'})Cl_2]$ (6) under an inert atmosphere. The 607 same reaction in air was accompanied by a further oxidation of 608 $HL^{2c'}$ with the formation of benzo [d] thiazol-6-one $(HL^{2c''})$ 609 bound to copper (II). Complex 6 was characterized by the 610 positive ion ESI mass spectrum with a peak at m/z 374.08 611 attributed to $[Cu(L^{2c'})]^+$, whereas the product obtained by an 612 oxidation in air revealed a peak at m/z 373.06 assigned to 613 $[Cu^1(HL^{2c''})]^+$. The peak at m/z 373.06 was also seen when 614 the reaction mixture of $HL^{2c''}$ with $CuCl_2 \cdot 2H_2O$ was subjected 615 to an ESI MS measurement.

The reactions of copper(II) with the oxidized TSCs, namely, 617 1,3,4-thiadiazole-containing species $HL^{1a'}$ and $HL^{1a''}$, were 618 monitored by ESI-MS experiments. When CuCl₂·2H₂O was 619 allowed to react with $HL^{1a'}$ and $HL^{1a''}$ in a 1:1 molar ratio, ESI 620 mass spectra of the reaction mixtures indicated the formation 621 of complexes with metal-to-ligand stoichiometry of 1:2, 622 namely, $[Cu(HL^{1a\prime})_2]^+$ and $[Cu(HL^{1a\prime\prime})_2]^+$, respectively. 623 Interestingly, under varied reaction conditions (different 624 solvents, air atmosphere, and varied temperature and reaction 625 time, see details in Table S7) the synthesis of copper(II) 626 complex of HL^{1a'} resulted in a sequential oxidation of the two 627 ligands, and several oxidized products could be identified based 628 on ESI-MS peaks as $[Cu(HL^{1a'})_2]^+$ (m/z 659.16), [Cu-629 ($HL^{1a'}$)($HL^{1a''}$)]⁺ (m/z 657.13), $[Cu(HL^{1a''})_2]^+$ (m/z 630 655.18), $[Cu(HL^{1a'})(CH_3CN)]^+$ (m/z 402.10), $[Cu(HL^{1a''})$ - 631 (CH_3CN)]⁺ (m/z 400.10). Moreover, attempts of the 632 chromatographic separation of the obtained compounds (on 633 SiO₂ with MeOH as eluent) led to a new species $[Cu(HL^{1a})]$ - 634 $(HL^{1d})]^+$ (m/z 537.15), in which one already oxidized ligand 635 $HL^{1a'}$ in $[Cu(HL^{1a''})_2]^+$ lost the phenolic moiety. The 636 complex formation of HL1a" in MeOH under heating at 50 637 °C resulted in two species $[Cu(HL^{1a''})(HL^{1d})]^+$ (m/z 537.15) 638 and $[Cu(HL^{1d})_2]^+$ (m/z 419.08), whereas under prolonged 639 heating (36 h) only $[Cu(HL^{1d})_2]^+$ was detected, and the 640 formation of complex [Cu(HL^{1d})₂Cl₂] (5) was confirmed by 641 SC-XRD.

The potentially redox-active TSC ligands $(HL^1, (L^2)^-$, and 643 HL^3) in 1, 2', and 3 proved to react slowly with oxygen in air. 644 Indeed, ESI mass spectra of methanolic solutions of 1, 2', or 3 645 after a prolonged standing in air showed peaks with m/z 646 shifted by 2 amu to lower masses in agreement with an 647 oxidative dehydrogenation required for the formation of two- 648 electron oxidation products.

To finally determine the redox status of the 4-aminophenolic 650 moiety, the configurations adopted by the metal-free ligands in 651 the solid state and their protonation level in copper(II) 652 complexes SC-XRD studies were performed.

X-ray Crystallography of the Metal-Free Ligands 654 HL^1-HL^3 and Copper(II) Complexes 1'-3'. The results 655 of X-ray diffraction studies of TSCs $HL^1\cdot C_2H_5OH$, HL^2 and 656 HL^3 are presented in Figure 3, while those of $[Cu(L^1)Cl]^{\cdot}$ 657 f3 CH_3OH (1'·CH₃OH), $[Cu(L^2)Cl]$ (2'), and $[Cu(L^3)Cl]^{\cdot}$ 658 CH_3OH (3'·CH₃OH) are in Figure 4. The $HL^1\cdot C_2H_5OH$ 659 f4 crystallized in the triclinic centrosymmetric space group $P\overline{1}$, 660 while HL^2 and HL^3 crystallized in the monoclinic space groups 661 $P2_1/c$ and $P2_1/n$, respectively. All three metal-free ligands 662 adopt an E configuration in terms of the nomenclature used for 663 the α -N-heterocyclic thiosemicarbazones 41 with the imine 664 nitrogen in the s-trans position to the sulfur atom and the 665 pyridine N1 atom. All TSCs crystallized in the thione form 666 with the C7-S bond length of 1.6839(15), 1.683(4) and 667

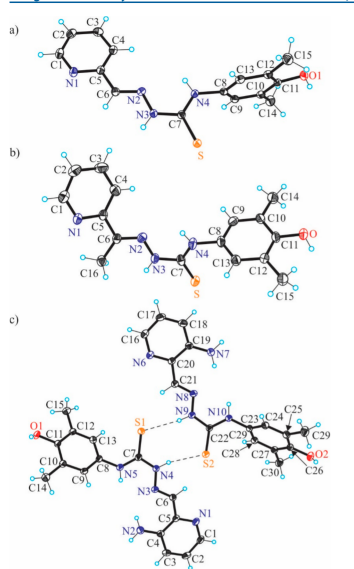


Figure 3. ORTEP views of ${\rm HL}^1{\rm - HL}^3$ with thermal ellipsoids at the 50% probability level. Selected bond distances (Å) and torsion angles (deg): (a) ${\rm HL}^1{\rm :}~{\rm C6-N2}~1.280(2),~{\rm N2-N3}~1.3701(18),~{\rm N3-C7}~1.357(2),~{\rm C7-S}~1.6839(15),~{\rm C7-N4}~1.331(2),~{\rm N4-C8}~1.442(2),~{\rm C11-O1}~1.3780(19);~{\rm \Theta_{C7-N4-C8-C13}}~88.7(2);~{\rm (b)}~{\rm HL}^2{\rm :}~{\rm C6-N2}~1.287(4),~{\rm N2-N3}~1.374(4),~{\rm N3-C7}~1.363(4),~{\rm C7-S}~1.683(4),~{\rm C7-N4}~1.326(4),~{\rm N4-C8}~1.446(4),~{\rm C11-O}~1.370(4);~{\rm \Theta_{C7-N4-C8-C13}}~78.4(4);~{\rm (c)}~{\rm HL}^3{\rm :}~{\rm C4-N2}~1.361(3),~{\rm C6-N3}~1.288(2),~{\rm N3-N4}~1.385(2),~{\rm N4-C7}~1.343(2),~{\rm C7-S1}~1.695(2),~{\rm C7-N5}~1.342(3),~{\rm N5-C8}~1.430(2),~{\rm C11-O1}~1.380(2);~{\rm \Theta_{C7-N5-C8-C13}}~52.5(3).$

668 1.695(2) Å, respectively. The distribution of electron density 669 in the dimethylphenolic moiety is typical for aromatic systems. 670 The C11–O bond length of 1.3780(19), 1.370(4), and 671 1.380(2) Å, respectively, is also characteristic for phenols. 672 The molecules of the three proligands are not planar. The 673 strong deviation of the phenolic unit from the mean plane of 674 the thiosemicarbazone fragment can be estimated by a 675 comparison of the torsion angle $\Theta_{\text{C7-N4-C8-C13}}$ of 88.7(2) 676 and 78.4(4)° in the first two structures (Figure 3a,b) and 677 $\Theta_{\text{C7-N5-C8-C13}}$ and $\Theta_{\text{C22-N10-C23-C28}}$ of 52.5(3) and 54.2(3)° 678 in two crystallographically independent molecules of HL³ 679 (Figure 3c).

680 In contrast to the structures of HL¹ and HL², the 681 asymmetric unit of HL³ consists of two molecules associated

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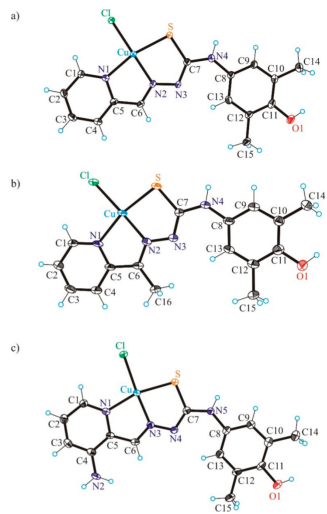


Figure 4. ORTEP views of 1'-3' with thermal ellipsoids at the 50% probability level. Selected bond distances (Å), bond angles (deg) and torsion angles (deg) in 1': Cu-N1 2.005(2), Cu-N2 1.962(2), Cu-S 2.2325(7), Cu-Cl 2.2507(7), C11-O1 1.370(4); N1-Cu-N2 81.77(9), N2-Cu-S 84.07(7), $\Theta_{\text{C7-N4-C8-C13}}$ -0.8(5); in 2': Cu-N1 2.022(4), Cu-N2 1.952(4), Cu-S 2.2636(16), Cu-Cl 2.2215(15), C11-O1 1.370(6); N1-Cu-N2 80.76(17), N2-Cu-S 84.46(12), $\Theta_{\text{C7-N4-C8-C13}}$ -2.1(8); in 3': Cu-N1 2.025(2), Cu-N3 1.961(2), Cu-S 2.2432(8), Cu-Cl 2.2636(8), C11-O1 1.374(4); N1-Cu-N3 81.58(10), N3-Cu-S 83.40(7), $\Theta_{\text{C7-N5-C8-C13}}$ 9.4(5).

in a centrosymmetric dimer via hydrogen-bonding interactions, 682 namely, N4–H···S2 [N4–H4 = 0.88 Å, H4···S2 = 2.48 Å, 683 N4···S2 = 3.3243(17) Å] and N9–H···S1 [N9–H9 = 0.88°, 684 H9···S1 = 2.47 Å, N9···S1 = 3.3341(17) Å]. A similar 685 centrosymmetric association was recently reported for 686 acetylpyrazine 4-N-phenyl thiosemicarbazone. 687

The copper(II) complexes $1'\cdot CH_3OH$ and $3'\cdot CH_3OH$ 688 crystallized in the monoclinic centrosymmetric space group 689 $P2_1/c$, while 2' crystallized in the triclinic centrosymmetric 690 space group $P\overline{1}$ without any cocrystallized solvent. The 691 copper(II) adopts a square-planar coordination geometry in 692 all three structures (Figure 4). The thiosemicarbazones act as 693 tridentate monoanionic ligands binding to copper(II) via a 694 pyridine nitrogen atom, an azomethine nitrogen atom, and a 695 thiolate sulfur atom. The fourth coordination site in all 696 complexes is occupied by the chlorido coligand. Pertinent 697 bond distances and bond angles are quoted in the legend to 698

699 Figure 4. The same coordination geometry of a copper(II) 700 bound by a monoanionic thiosemicarbazone and a mono-701 dentate coligand was reported for [CuCl(mPip-FTSC-H)]-702 0.15CH₃OH, ⁵⁵ [Cu(L₁)(μ -Cl)]Cl, and [Cu(L₂)(μ -Cl)]Cl-703 H₂O, where ligands L₁ and L₂ represent 3-methyl-5-oxo-1-704 phenyl-3-pyrazolin-4-carboxaldehyde and 5-oxo-3-phenyl-3-705 pyrazolin-4-carboxaldehyde TSC, respectively. ⁵⁶

A comparison of the Cu(II) to TSC ligand bond lengths in 707 1' with those in the copper(II) complex with pyridine-2-708 carboxaldehyde thiosemicarbazone⁵⁷ (Cu-N1 = 2.034(4), 709 Cu-N2 = 1.975(3), Cu-S = 2.278(1) Å) shows that these are 710 statistically significantly shorter in 1'. This difference is 711 probably due to the formation of centrosymmetric associates 712 via intermolecular interactions with the shortest contact Cu···Si $_{713} = 2.760(2)$ Å and not due to the presence of a phenolic moiety 714 at N4. The interatomic repulsions in the copper(II) complex 715 with a 4 + 1 coordination geometry are expected to be stronger 716 than those in 1', in which the Cu(II) is four-coordinate. In 717 another complex $[CuLCl]_2[Cu(pic)_2]$ (with HL = pyridine-2-718 carboxaldehyde thiosemicarbazone and pic = pyridine-2carboxylate), in which the intermolecular contacts are over 3 Å, 720 the Cu(II) to TSC bond distances are shorter and very similar 721 to those in 1' (Cu-N1 = 2.005(8), Cu-N2 = 1.942(9), Cu-S $722 = 2.264(3) \text{ Å}).^{48}$ The metric parameters in the copper(II)-723 ligand chromophore of [Cu(triapine-H)Cl](H₃O)Cl (Cu-N_{pv} 724 = 2.031(8), Cu-N_{hydrazine} = 1.937(9), Cu-S = 2.281(3) and 725 Cu-Cl = 2.2493(5) Å) are statistically the same as those in 3′, 726 except Cu–S, which is by ca. 0.04 Å (>12 σ) shorter in 3' than 727 in the copper(II) complex with triapine. This is likely due to 728 different protonation states of the ligands in the two 729 complexes, even though the authors described the triapine 730 ligand in its copper(II) complex as a monoanion with an extra 731 proton at a cocrystallized water molecule. S

Note that the organic ligands in all three complexes are 733 almost planar in contrast to the situation described previously 734 for the metal-free ligands. The value of the torsion angle 735 $\Theta_{\text{C7-N4-C8-C13}}$ for 1'•CH₃OH and 2' (Figure 4a,b) increased 736 from -88.7(2) and -78.4(4)° in HL¹ and HL² to -0.8(5) and 737 -2.1(8)°, respectively. Analogously, the torsion angle 738 $\Theta_{\text{C7-N5-C8-C13}}$ of 52.5(3) in HL³ becomes of 9.4(5)° in 3'• 739 CH₃OH upon coordination to copper(II).

As for the metal-free TSCs, the phenolic moiety remained intact in all three complexes, namely, in its original oxidation state. The distribution of electron density over the aromatic phenolic ring is well-comparable to that in the TSCs.

X-ray Crystallography of Oxidized Products. The 745 results of X-ray diffraction studies of oxidized organic species 746 $\rm HL^{1a'}$, $\rm HL^{1a''}$, $\rm HL^{2b}$, $\rm HL^{2e}$, and $\rm HL^{2c''}$ •0.5CHCl $_3$ are displayed 747 in Figure 5 and Figure S6, while those of copper(II) complexes 748 with oxidized ligands 4-6 are shown in Figure 6 and Figure 749 S7. The oxidized species $HL^{1a'}$ and $HL^{1a''}$ crystallize in the 750 monoclinic space groups $P2_1/n$ and Cc, respectively. The 751 molecule HL^{1a'} is almost planar, while in HL^{1a''} the moiety at 752 N4 slightly deviates from planarity. The dihedral angle 753 $\Theta_{\text{C7-N4-C8-C13}}$ is of 5.8(3)°. Both contain a thiadiazole five-754 membered ring. The distribution of electron density in them is 755 very similar. In contrast, the bond length distribution in the 756 aryloxide moiety is quite different. In the two-electron oxidized 757 product HL1a' the distribution of electron density is in 758 agreement with that of the 3,5-dimethyl-1,4-aminophenolic 759 moiety, while in the four-electron oxidized species HL1a" the 760 electron density agrees with that of the 3,5-dimethyl-1,4-761 benzoquinone imine unit (see legend to Figure 5a,b). In

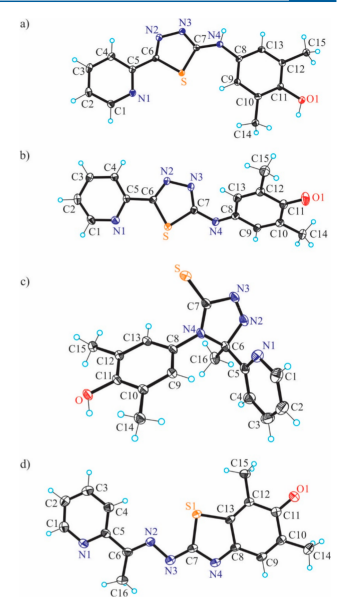


Figure 5. ORTEP views of two-electron and four-electron oxidized species of (a) $HL^{1a'}$ and (b) $HL^{1a''}$, as well as of products that resulted from an oxidation of HL^2 , namely, of (c) HL^{2b} and (d) HL2c"·0.5CHCl3. Selected bond distances (Å) and torsion angles (deg) in (a) HL^{1a}': C6-N2 1.3029(17), N2-N3 1.3739(15), C6-S 1.7405(14), C7-S 1.7382(13), C8-C9 1.3878(19), C9-C10 1.4003(18), C10-C11 1.3927(19), C11-C12 1.4006(19), C12-C13 1.3922(19), C11–O1 1.3820(16); $\Theta_{C7-N4-C8-C9}$ 1.0(2); in (b) HL^{1a}": C6-N2 1.305(3), N2-N3 1.382(2), C6-S 1.727(2), C7-S 1.734(2), C8-C9 1.458(3), C9-C10 1.341(3), C10-C11 1.480(3), C11-C12 1.491(3), C12-C13 1.342(3), C11-O1 1.226(3); $\Theta_{\text{C7-N4-C8-C13}}$ 5.8(3); in (c) HL^{2b} : C6-N2 1.485(2), N2-N3 1.247(2), N3-C7 1.472(2), C7-S 1.6465(18), C7-N4 1.325(2), N4-C6 1.479(2), C11-O 1.3728(18), N4-C8 1.4409(19); in (d) $HL^{2c''} \cdot 0.5CHCl_3$: C6-N2 1.306(4), N2-N3 1.394(4), N3-C7 1.296(4), C7-N4 1.388(4), N4-C8 1.311(4), C8-C9 1.444(4), C9-C10 1.340(4), C10-C11 1.488(5), C11-O1 1.234(4), C11-C12 1.496(4), C12-C13 1.349(4), C13-C8 1.461(4), C13-S1

particular, the C11–O1 bond length in these two compounds 762 is quite different at 1.3820(16) and 1.226(3) Å, respectively. 763 The X-ray diffraction structure of HL^{2b} confirmed the two- 764 electron oxidation of the original ligand HL^2 and the formation 765

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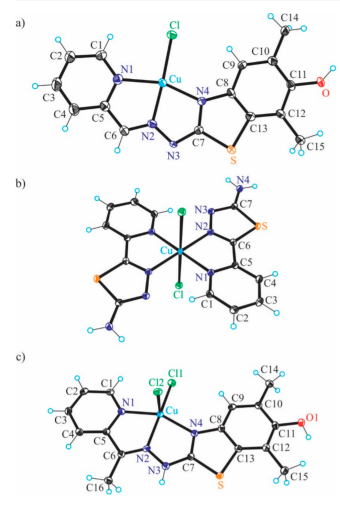


Figure 6. ORTEP views of $[Cu(L^{1c'})Cl]$ (4), $[Cu(HL^{1d})_2Cl_2]$ (5), and $[Cu(HL^{2c'})Cl_2]$ (6) with thermal ellipsoids at the 50% probability level. Selected bond distances (Å) and bond angles (deg) in (a) 4: Cu-N1 2.054(4), Cu-N2 1.956(4), Cu-N4 2.001(3), Cu-Cl 2.2575(12), C11-O 1.370(5); N2-Cu-N1 80.02(15), N2-Cu-N4 78.59(14); in (b) 5: Cu-N1 2.0384(11), Cu-N2 2.0089(11), Cu-Cl 2.8116(3), N2-Cu-N1 99.32(4); in (c) 6: Cu-N1 2.0329(16), Cu-N2 1.9854(16), Cu-N4 2.0358(16), Cu-Cl1 2.2100(5), C11-O 1.374(2); N2-Cu-N1 78.00(16), N2-Cu-N4 79.31(6).

766 of the TAT ring, while that of $HL^{2c\prime\prime}$ confirmed the further 767 two-electron oxidation of $HL^{2c\prime\prime}$. The bond-length distribution 768 in the molecule of $HL^{2c\prime\prime}$ indicates the presence of the 769 benzo[d]thiazol-6-one moiety. The double-bond character of 770 N3–C7 1.296(4) indicates the formation of this four-electron

oxidation product from the two-electron oxidation product 771 $HL^{2c'}$ by the loss of two electrons and two protons.

The X-ray diffraction study of 4 (Figure 6a) revealed that 773 the ligand underwent an oxidative dehydrogenation accom- 774 panied by the intramolecular cyclization via a C-S coupling 775 reaction between a phenolic carbon and a thione group into a 776 five-membered thiazole ring instead of the expected oxidative 777 dehydrogenation (two-electron oxidation accompanied by the 778 loss of two protons) of the 3,5-dimethyl-1,4-aminophenol unit 779 with formation of a 3,5-dimethyl-1,4-benzoquinone imine 780 moiety (see Chart 2, Scheme 1). This intramolecular sulfur 781 arylation resulted in the change of coordination mode, so that 782 the thioether sulfur atom with diminished electron-donor 783 properties is not involved in the coordination to copper(II). 784 This is in agreement with the coordination chemistry of 785 isothiosemicarbazones,⁵⁹ which as a rule do not use a sulfur 786 atom for coordination to first-row transition metals. In this 787 context, it is worth mentioning that the binding of 788 isothiosemicarbazones to zinc(II) and copper(II) via a 789 thioether sulfur atom has been documented quite recently, 60 790 when bulkier than chlorido coligands, for example, iodido and 791 bromido, were involved in a coordination to the metal. 792 Complex 4 might be one of the products of the oxidation of 793 copper(II) complexes over time in methanol by air oxygen. 794 Some rare examples of a thiosemicarbazone cyclization with 795 the thiazole ring formation due to the coordination to 796 copper(II) were recently reported (iminodiacetate-thiosemi- 797 carbazones and N-phenylthiosemicarbazones). 52,53,61 The new 798 ligand obtained by the intramolecular cyclization in Cu(HL¹)- 799 Cl₂ belongs to the class of biologically active substituted 2-800 hydrazinylbenzothiazoles, which showed anticancer activity 801 themselves as well as upon coordination to different 802 metals. 62-65 Two molecules of complex 4 are associated into 803 a centrosymmetric dimer via two intermolecular μ -chlorido 804 bridges as shown in Figure S7.

The molecular structure of **5** shown in Figure 6b indicates a 806 strongly tetragonally distorted six-coordinate geometry of 807 copper(II), in which two pyridine-thiadiazole ligands act as a 808 bidentate and occupy the equatorial sites in a *trans* mutual 809 arrangement and two quite weakly bound chlorido coligands in 810 axial positions. Taking into account the interatomic Cu–Cl 811 separation (2.8116(3) Å) the complex can also be described as 812 square-planar.

As in 4, the coordinated ligand in 6 acts as a tridentate and 814 binds to copper(II) via atoms N1, N2, and N4. However, while 815 4 is square-planar, 6 is very close to square-pyramidal (τ_5 = 816 0.16). The organic ligand is monoanionic in 4, while neutral 817 in 6. An additional coordination of chlorido coligand(s) 818 counterbalances the 2+ charge of the central atom.

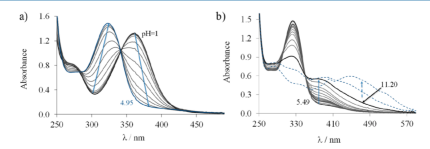


Figure 7. UV–Vis absorption spectra recorded for proligand HL¹ in the pH ranges of (a) 1.00–4.95 and (b) 5.49–11.82. $c_{\rm HL}$ = 50 μ M; 30% (v/v) DMSO/H₂O; I = 0.1 M (KCl); T = 25 °C.

Table 1. pK_a Values Determined by UV-Vis Titrations in 30% (v/v) DMSO/H₂O and log $D_{7.4}$ (n-Octanol/Water) Values of the TSCs HL^1-HL^3 and Their Complexes^a

	method	HL^1	HL^2	HL^3
pK_a (PyH ⁺)	UV-vis	3.01 ± 0.01	3.59 ± 0.02	3.95 ± 0.04
pK_a (NNH)	UV-vis	10.55 ± 0.01	11.08 ± 0.02	nd
$\logD_{7.4}$ (proligand)	partitioning	$+1.30 \pm 0.03$	$+2.1 \pm 0.1$	$+1.67 \pm 0.01$
$logK'_{5.9}$ (complex)	EDTA displacement	9.67 ± 0.01	nd^b	9.78 ± 0.01
$\log D_{7.4}$ (complex)	partitioning	-0.40 ± 0.06	nd^b	-0.42 ± 0.03
$k_{\rm obs}~({\rm min}^{-1})~({\rm complex})$ in 30% DMSO	UV-vis (with GSH)	0.033 ± 0.004	nd^b	0.035 ± 0.004
$k_{\rm obs}~({\rm min}^{-1})~({\rm complex})~{\rm in}~60\%~{\rm DMSO}$	UV-vis (with GSH)	0.021 ± 0.001	too slow ^c	0.024 ± 0.004

"Conditional stability constants (log $K'_{5,9}$) of the complexes determined by UV–vis EDTA displacement studies in 30% (v/v) DMSO/H₂O and rate constants (k_{obs}) obtained for the redox reaction of the complexes with GSH (pH = 7.4 (50 mM HEPES); $c_{\text{complex}} = 25 \, \mu\text{M}$; $c_{\text{GSH}} = 1.25 \, \text{mM}$ in 30% and); $c_{\text{complex}} = 12.5 \, \mu\text{M}$; $c_{\text{GSH}} = 600 \, \mu\text{M}$ in 60% (v/v) DMSO/H₂O) { $T = 25 \, ^{\circ}\text{C}$; $I = 0.1 \, \text{M}$ (KCl)}. Not determined (nd) due to the bad solubility of the complex under the conditions. Rate constant could not be determined due to the very slow redox reaction.

To understand the difference in protonation states and reactivity of the originally prepared complexes and those solutions equilibrium studies were performed on the ligands and their copper(II) complexes.

25 SOLUTION EQUILIBRIUM STUDIES

Proton Dissociation Processes and Lipophilicity of 827 the Ligands. Proton dissociation constants (pK_a) of drug 828 molecules indicate the actual protonation state and the charge 829 at a given pH, and therefore pK_a are important parameters that 830 affect the pharmacokinetic properties as well. The N-terminally 831 monosubstituted TSCs HL^1-HL^3 belong to the family of α -N-832 pyridyl TSCs; thus, they possess the pyridinium (PyH)+ and 833 the hydrazinic-NNH as proton dissociable groups besides the 834 phenolic moiety. Since these TSCs and their copper(II) 835 complexes have a limited water solubility, the equilibrium 836 studies were performed by UV-vis spectrophotometry in a 837 30% (v/v) DMSO/H₂O solvent mixture using relatively low concentrations (50 µM). Representative UV-vis spectra recorded for HL¹ at various pH values are shown in Figure 7a. On the basis of the spectral changes two well-separated 841 deprotonation processes were observed between pH 2 and 11. The first proton dissociation step taking place at pH < 5 is 843 accompanied by a blue shift, and the λ_{max} is shifted from 362 to 844 322 nm. This deprotonation step is attributed to the proton on 845 the pyridinium nitrogen (PyH⁺). Upon an increase of the pH a 846 new process occurred as evidenced by a new band in the range 847 of 350-450 nm (Figure 7b) and an isosbestic point at 350 nm, 848 namely, the deprotonation of the hydrazinic nitrogen. In the 849 strongly basic pH range (pH > 11.2) new broad bands appear 850 at 400-600 nm (Figure 7b) with irreversible spectral changes most likely due to an oxidation of the TSC by the air oxygen. Therefore, only two pK_a values could be determined (Table 853 1) based on the deconvolution of the UV-vis spectra recorded 854 at pH < 11.2 for HL¹ (molar absorbance spectra are seen in 855 Figure S8a) as the oxidation hindered the accurate 856 determination of the pK_a for the aromatic OH group. Two 857 p K_a values were computed for HL^2 from the UV-vis titration 858 data (Figure S9) as well; however, only one p K_a was obtained 859 in the case of HL³ (Table 1), namely, that for the 860 deprotonation of the PyH+, since the proton dissociation of 861 the hydrazinic nitrogen and the oxidation of the TSC were 862 partly overlapped. On the basis of the determined p K_a values, it 863 can be concluded that the presence of the electron-donating 864 methyl group in HL² results in a significant increase of both 865 p K_a values when compared to that of HL^1 . A similar behavior

was reported for the analogous 2-formylpyridine and 2- 866 acetylpyridine TSC in our previous work. The 67 The 67 The 67 The 67 The 67 PyH+ group was also increased significantly by the addition of 868 the electron-donating amine group at the pyridine ring, in 869 agreement with data reported previously for the FTSC and 870 triapine. 68 All proligands are air-sensitive in the strongly basic 871 pH range (pH > 11). Concentration distribution curves were 872 computed for them at pH < 11 (see Figure S8b for HL¹) 873 revealing that their neutral forms predominate at a 874 physiological pH.

The solution stability of the proligands was monitored at pH $_{876}$ 7.4 by spectrophotometry. The UV—vis spectra recorded over $_{877}$ 4 h revealed no measurable spectral changes, suggesting that $_{878}$ the oxidation of these proligands does not take place (or just $_{879}$ very slowly) in an aqueous solution at a physiological pH. $_{880}$ However, HL $_{2}$ showed a certain level of slow decomposition at $_{881}$ pH 1.5, namely, a $_{6\%}$ absorbance decrease at 354 nm in $_{882}$ (Figure S10), which is most likely the consequence of the less $_{883}$ extended conjugation in the molecule due to the cleavage of $_{884}$ the C=N Schiff base bond, as it was also reported for 2- $_{885}$ acetylpyridine TSC. $_{67}$ Thus, the rate of this acid-catalyzed $_{886}$ reaction is increased with the increasing number of methyl $_{887}$ groups present in the $_{\alpha}$ -N-pyridyl TSC.

Besides pK_a values, lipophilicity is also an important 889 pharmacological property of a drug, as it strongly influences 890 the ability of the compound to pass through biological 891 membranes. Therefore, distribution coefficients ($\log D_{7.4}$) were 892 determined using the shake-flask method in an n-octanol-893 buffered aqueous solution at pH 7.4 (Table 1). The $\log D_{7.4}$ 894 values indicate the moderate lipophilic character of the 895 proligands. The substitution at the end nitrogen atom of the 896 thosemicarbazide moiety and the presence of a methyl group at 897 the Schiff base bond induce a somewhat higher lipophilicity. 898 The presence of the phenolic moiety undoubtedly increases 899 the $\log D_{7.4}$ values compared to those of FTSC (+0.73), 67 900 AcTSC (+1.02) 67 and triapine (+0.85). 69

In summary, these TSCs are stable in their neutral form in a 902 quite broad pH range (including pH 7.4).

Solution Stability and Redox Properties of the 904 Copper(II) Complexes. The metal complexes often undergo 905 transformation processes upon dissolution, such as proto- 906 nation, deprotonation, or dissociation to a metal-free ligand 907 and metal ion depending on the pH, their concentration, and 908 the solution speciation. The knowledge of the actual chemical 909 form of the biologically active metal complexes in solution 910 close to physiologically relevant conditions is quite important 911 to elucidate the mechanism of action. Therefore, the solution 912

913 stability of the copper(II) complexes (Cu(HL¹)Cl₂, [Cu(L²)-914 Cl], and Cu(HL³)Cl₂) was studied by UV-vis spectropho-915 tometry. The simple α -N-pyridyl TSCs (e.g., triapine, FTSC) 916 generally form very stable monoligand copper(II) complexes, 917 and the species in which the monoanionic ligand is 918 coordinated via the (N_{pyridine},N,S⁻) mode predominates in a 919 wide pH range at a 1:1 metal-to-ligand ratio. 68 At lower pH 920 this type of complex is protonated, and thus the neutral ligand 921 is bound via (N_{pyridine},N,S) donor atoms, while a mixed 922 hydroxido complex with the $(N_{pyridine}, N, S^-)(OH)$ coordina-923 tion pattern is formed in the basic pH range. On the basis of 924 the close structural similarities between HL1-HL3 and the 925 listed TSCs with a simpler scaffold, the formation of the same 926 type of complexes is feasible. UV-Vis titrations were 927 performed with the complexes in a 30% (v/v) DMSO/H₂O 928 solvent mixture, and representative spectra are shown for 929 Cu(HL¹)Cl₂ in Figure 8. The spectra remain intact in a broad

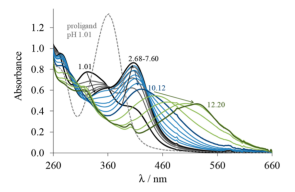


Figure 8. UV–Vis absorption spectra recorded for complex $Cu(HL^1)Cl_2$ in the pH range of 1.01–12.20 (solid lines) and for HL^1 at pH 1.01 (dashed gray line). $c_{complex/HL} = 50 \mu M$; 30% (v/v) DMSO/H₂O; I = 0.1 M (KCl); T = 25 °C.

930 pH range (2.7–7.6), and an absorption band is observed with 931 $\lambda_{\rm max}$ at 406 nm being typical for a S \rightarrow Cu charge transfer. This 932 finding indicates the dominant presence of only one kind of 933 complex, which is most probably the species with the 934 (N_{pyridine},N,S⁻) tridentate coordination mode. By decreasing 935 the pH the $\lambda_{\rm max}$ is hypsochromically shifted to 322 nm. The 936 presence of the isosbestic point at 362 nm implies that only 937 two species are involved in this equilibrium. As the spectrum 938 recorded at pH 1.01 significantly differs from that of the TSC, 939 this equilibrium corresponds to the protonation of the complex 940 at the noncoordinating hydrazinic nitrogen (Chart S1) rather 941 than to its dissociation to the free metal ion and ligand. This 942 process is not completed when the pH decreases to 1, and a

 pK_a value less than 1.5 could be estimated. When the pH is 943 increased, two overlapping processes are suggested to take 944 place at pH > 8 via the continuous bathochromic shift of the 945 absorption maximum, and pK_a values of 9.80 ± 0.01 and 11.02 946 ± 0.01 were computed. In this pH range most probably the 947 coordinated water molecule deprotonates, and a mixed 948 hydroxido complex is formed along with the deprotonation 949 of the phenolic group of the bound ligand. Similar spectral 950 changes were monitored for $Cu(HL^3)Cl_2$, and $pK_a < 1.5$ was 951 estimated for the process in the acidic pH range as well.

However, the formation of precipitate (significant baseline 953 elevation and absorbance decrease in the whole wavelength 954 range) at pH > 8 hindered the calculation of the proton 955 dissociation constants of the complexes from spectra collected 956 in this pH range. Unfortunately, during the titration of 957 $[Cu(L^2)Cl]$ the formation of a precipitate was observed 958 already at the acidic pH; thus, the deprotonation processes 959 could not be evaluated.

The copper(II)-TSC complexes are often redox-active 961 under physiological conditions, which has an impact on their 962 cytotoxicity. To investigate whether complexes $[Cu(L^1)]^+$, 963 $[Cu(L^2)]^+$, and $[Cu(L^3)]^+$ can be reduced by the most 964 abundant low molecular mass cellular reductant, GSH, 965 spectrophotometric measurements were performed on their 966 direct reaction under strictly anaerobic conditions at pH 7.4. 967 First, the assay was performed in the presence of 30% DMSO 968 using a 25 µM complex concentration. However, the limited 969 solubility of $[Cu(L^2)]^+$ did not allow the measurement. 970 Therefore, the assay was also performed in the presence of 971 60% DMSO at a lower (12.5 μ M) concentration for all the 972 three complexes. The spectral changes are shown in Figure 9 973 f9 for $[Cu(L^1)]^+$ and $[Cu(L^3)]^+$ complexes in the presence of a 974 large excess of GSH in 30% (v/v) DMSO/H₂O. After the 975 complexes were mixede with GSH, a well-detectable change is 976 observed due to the formation of ternary complexes via the 977 coordination of GSH as it was reported for several TSC 978 complexes. 70,71 Then the spectral changes show the 979 absorbance decrease at the $\lambda_{\rm max}$ of the S ightarrow Cu charge transfer 980 band of the complexes. The final spectra show a strong 981 similarity to those of HL^1 and HL^3 at $\lambda > 310$ nm suggesting 982 the release of the TSCs. However, in this case the reduction is 983 responsible for the liberation of the TSCs and copper(I), 984 which forms complexes with GSH (that is in high excess in the 985 sample). Copper(I) favors a tetrahedral coordination environ- 986 ment, while HL1 and HL3 as planar tridentate ligands cannot 987 satisfy these requirements and accommodate the cation. This 988 contradiction is a driving force for a complex destabilization, 989 especially in the presence of GSH, which can efficiently bind 990 copper(I). 64 In addition, a one-electron reduction increases the 991

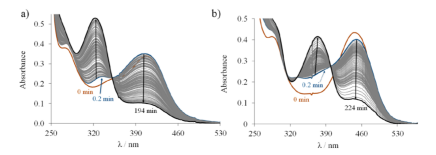


Figure 9. Time-dependent changes of the UV–vis spectra of (a) $\text{Cu}(\text{HL}^1)\text{Cl}_2$ and (b) $\text{Cu}(\text{HL}^3)\text{Cl}_2$ in the presence of 50 equiv of GSH at pH 7.4 under anaerobic conditions. $c_{\text{complex}} = 25 \ \mu\text{M}$; $c_{\text{GSH}} = 1.25 \ \text{mM}$; pH = 7.40; 30% (v/v) $\text{DMSO/H}_2\text{O}$; $I = 0.1 \ \text{M}$ (KCl); $T = 25 \ ^{\circ}\text{C}$.

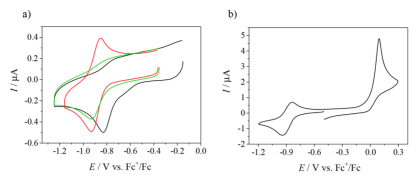


Figure 10. (a) Cyclic voltammograms of 0.5 mM 1 (black trace), 2' (red trace), and 3 (green trace) in DMSO/n-Bu₄NPF₆ at a GC working electrode at the scan rate of 100 mV s⁻¹; (b) comparison of the reduction and the oxidation peak of 2' (scan rate of 100 mV s⁻¹).

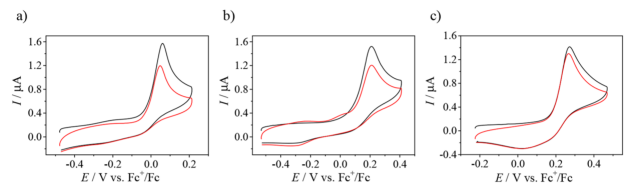


Figure 11. Cyclic voltammograms of 0.5 mM of (a) 1 and (b) the corresponding ligand in DMSO/n-Bu₄NPF₆ and of (c) 1 in MeOH/LiClO₄ at the GC working electrode, at scan rate of 100 mV s⁻¹.

992 basicity of the coordinated TSCs facilitating their protonation 993 and dissociation from the copper(I). Note, however, that the 994 process was reversible, as bubbling oxygen into the samples 995 regenerated the starting spectra. Complex $[Cu(L^2)]^+$ behaved 996 differently, as only minor spectral changes were seen upon 997 treatment with GSH in 60% (v/v) DMSO/H₂O (Figure S11b). From the measured absorbance-time curves rate constants (k_{obs}) were calculated (Table 1). Similar reduction 1000 rates for $[Cu(L^1)]^+$ and $[Cu(L^3)]^+$ complexes were obtained, 1001 and somewhat lower $k_{\rm obs}$ values were found in the presence of 1002 the higher fraction of DMSO. Notably, ascorbate, which is a 1003 weaker reducing agent compared to GSH and is found in 1004 higher concentration in the extracellular fluids, was not able to 1005 reduce these complexes under the same conditions. On the 1006 contrary, the more powerful reducing agent DTT could reduce $[Cu(L^1)]^+$, $[Cu(L^2)]^+$, and $[Cu(L^3)]^+$ in a very fast reaction. The reduction was complete within several seconds (at 12.5 μ M complex and 600 μ M DTT concentrations in the presence 1010 of 60% DMSO, Figure S11c,d). In this case, the reaction was 1011 reversible upon exposure to O_2 only for $[Cu(L^2)]^+$.

Overall, the solution equilibrium data provide further lol3 evidence that the complex $[Cu(L)]^+$ with the coordinated lol4 monoanionic ligand predominates in a wide pH range. In order to obtain a deeper insight into the observed behavior of both lol6 metal-free ligands and their copper(II) complexes in the lol7 presence of oxidants (atmospheric oxygen) and reductants lol8 (GSH and ascorbate) spectroelectrochemical investigations lol9 were also performed.

1020 **Electrochemistry and Spectroelectrochemistry.** Cyclic 1021 voltammograms of 1, 2', and 3 in DMSO/*n*-Bu₄NPF₆ recorded 1022 with a glassy carbon (GC) working electrode at a scan rate of 1023 100 mV s⁻¹ showed a redox activity in both cathodic and

anodic regions. Copper(II) undergoes an electrochemically 1024 irreversible or quasi-reversible reduction to copper(I) at $E_{pc} = 1025$ -0.83 V for 1 and -0.93 V versus Fc⁺/Fc for both 2' and 3 1026 (Figure 10a). Notably, the corresponding ligands are not 1027 f10 redox-active in the cathodic region (data not shown). An 1028 irreversible oxidation was observed for these complexes, which 1029 was identified as a two-electron oxidation of the TSCs with a 1030 release of two protons. A two-electron oxidation was confirmed 1031 by a comparison of the reduction peak (one-electron $Cu(II) \rightarrow 1032$ Cu(I) redox process) and the oxidation peak of 2' taken in 1033 equivalent amounts as shown in Figure 10b. In addition, an 1034 electrolysis of HL¹ at 1000 mV versus Ag/AgCl in CH₃CN in 1035 the presence of 0.2 M n-Bu₄NPF₆ generated a mixture of 1036 several products from which $HL^{1a'}$ and $HL^{1a''}$ were separated 1037 on silica. ESI-MS and ¹H NMR spectra were identical with 1038 those of the products obtained by an oxidation of HL1 with 1039 DDQ as mentioned previously.

The oxidation peak of the TSC ligand was observed at $E_{\rm pa}=1041+0.06~\rm V$ for 1 and 2' and at +0.04 V for 3, and it is negatively 1042 shifted in comparison to the corresponding metal-free ligands 1043 ($E_{\rm pa}=+0.21~\rm V$ for $\rm HL^1$, +0.24 V for $\rm HL^2$, and +0.18 V for $\rm HL^3$ 1044 (all vs Fc⁺/Fc at a scan rate of 100 mV s⁻¹)), as shown for 1 1045 and its corresponding metal-free ligand $\rm HL^1$ in Figure 11a,b, 1046 f11 respectively. There are also significant changes in the shape 1047 and intensity of cyclic voltammograms upon the second 1048 oxidation scan (see red traces in Figure 11a,b), which indicate 1049 a further oxidation of the products obtained after the first 1050 oxidation in DMSO, in line with the chemical oxidation of the 1051 compounds. Note that, in a proton-donating solvent, the 1052 potentials of both reduction and oxidation processes were 1053 shifted to the more positive values versus the internal potential 1054 standard Fc⁺/Fc, and additionally, a broad reduction peak 1055

1056 appeared during the reverse scan in the cyclic voltammogram 1057 at a strongly negatively shifted potential (Figure 11c). A 1058 distinct oxidation pattern of the corresponding voltammo-1059 grams in protic media is caused by the involvement of protons 1060 in the process in accordance with chemical oxidations 1061 discussed previously and the well-known reaction mechanism 1062 proposed for the quinone-like systems. ^{70,71}

Similar redox behavior was observed for the anodic 1064 oxidation of $HL^{1a'}$ in DMSO with several new redox-active 1065 species, which appeared upon the first and the second 1066 voltametric scans (Figure S12a). However, the oxidized 1,4-1067 benzoquinone imine species $HL^{1a''}$ can be reversibly reduced in 1068 the cathodic part (Figure S12b) with a voltammetric pattern 1069 characteristic for the electrochemistry of quinones in aprotic 1070 media. Moreover, EPR spectroelectrochemistry confirmed 1071 the formation of an anion radical at the first reduction peak 1072 (see inset in Figure S12b). A rich hyperfine splitting and a g-1073 value of 2.0046 points to the spin delocalization and 1074 contribution of heteroatom (presumably nitrogen) to the g-1075 value.

To support the assignment of the redox processes described previously, EPR/UV—vis spectroelectrochemical measurements were performed, and the results are shown for 1 in Figures 12 and 13. The UV—vis spectrum of 1 exhibits two absorption bands at 276 and 428 nm, where the first one is due to the absorption of the TSC ligand, while the second one can lose be attributed to the ligand-to-metal (S \rightarrow Cu) charge transfer (LMCT). T3,74 Upon the cathodic reduction of 1 in the region of the first reduction peak a new broad absorption band at 331

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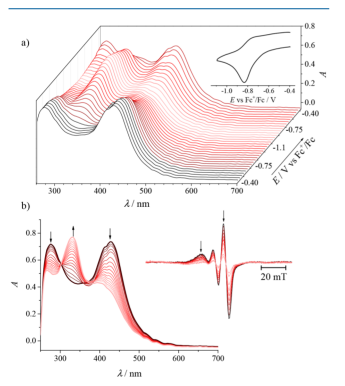


Figure 12. Spectroelectrochemistry of 1 in $n\text{-Bu}_4\text{NPF}_6/\text{DMSO}$ in the region of the first cathodic peak: (a) potential dependence of UV–vis spectra with the corresponding in situ cyclic voltammogram (Pt-microstructured honeycomb working electrode, scan rate of 5 mV s⁻¹); (b) evolution of UV–vis spectra in 2D projection upon forward scan. (inset) Evolution of EPR spectra measured at the first reduction peak using a Pt mesh working electrode.

nm appears with a simultaneous decrease of the initial optical 1085 bands at 276 and 428 nm via an isosbestic point at 302 nm 1086 (Figure 12). An analogous spectroelectrochemical response 1087 was observed for 2' as shown in Figure S13. This observation 1088 is different from that encountered by the reduction of the 1089 copper(II)-TSC complexes by GSH (vide supra), which led 1090 to the liberation of the ligand and formation of the copper (I) 1091 complex with GSH. In the spectroelectrochemical experiment 1092 in the absence of strong Cu(I) complexing agents, such as 1093 GSH, the TSC ligand may coordinate to Cu(I) and form a 1094 linear or tetrahedral complex. Upon the voltammetric reverse 1095 scan, a nearly full recovery of the initial optical bands was 1096 observed, which confirms the relatively good stability of 1097 cathodically generated Cu(I) complex with HL² and, thus, the 1098 chemical reversibility of this redox process. Rare examples of 1099 four- and three-coordinate copper($ar{I}$) complexes with poten- $_{1100}$ tially tridentate and bidentate thiosemicarbazones were 1101 reported previously. 75,76 The room-temperature X-band EPR 1102 spectrum of 1 showed a typical signal for d⁹ Cu(II) species, 1103 which decreased stepwise upon a cathodic reduction at the first 1104 cathodic peak. This is in line with the metal-centered reduction 1105 and formation of EPR-silent d¹⁰ Cu(I) species¹⁰ (see inset in 1106 Figure 12b). EPR spectra of 1, 2', and 3 measured in frozen n- 1107 Bu₄NPF₆/DMSO at 77 K show a characteristic axial symmetry 1108 $(g_{\parallel} > g_{\perp} > g_{e})$ implying a square-planar coordination and the 1109 presence of one dominating species in DMSO (Figure S14). 1110

The in situ cyclic voltammogram and simultaneously 1111 recorded evolution of UV-vis spectra upon an anodic 1112 oxidation of 1 in DMSO provide further evidence for the 1113 ligand-based irreversible oxidation. Spectral changes accom- 1114 panying the oxidation of 1 are shown in Figure 13. These 1115 changes are characteristic for the other two complexes 2' and 3 1116 as well. Note that, in the region of the first oxidation peak, new 1117 optical bands at 295 and 356 nm appear with a simultaneous 1118 decrease of the initial absorption with a maximum at 428 nm 1119 (Figure 13a). However, the product formed upon oxidation is 1120 not reduced back during the reverse voltammetric scan (Figure 1121 13b), indicating the chemical irreversibility of the redox 1122 process. In the EPR spectroelectrochemistry of 1 in DMSO/n- 1123 Bu₄NPF₆, no changes of the EPR signal were detected upon 1124 the oxidation at the first anodic peak, providing evidence of the 1125 two-electron oxidation process taking place on the TSC ligand. 1126

The remarkable stability of copper(II) complexes 1, 2′, and 1127 3 at a physiological pH, their moderate lipophilic character 1128 (log $D_{7.4} = -0.4$ to -0.42) and copper(II)/copper(I) redox 1129 activity ($E_{\rm red} = -0.83$ to -0.93 V vs Fc⁺/Fc) in a biologically 1130 relevant window of redox potentials (-0.4 to +0.8 V vs NHE 1131 or -1.04 to 0.16 V vs Fc/Fc⁺) prompted the investigation of 1132 their antiproliferative activity in cancer cell lines.

INHIBITION OF CELL VIABILITY AND APOPTOSIS 1134 ASSAY 1135

Cytotoxicity of the TSCs, Their Oxidized Products 1136 and Copper(II) Complexes. The in vitro cytotoxicity of the 1137 TSCs HL¹-HL³, copper(II) complexes Cu(HL¹)Cl₂, [Cu-1138 (L²)Cl], and Cu(HL³)Cl₂, and oxidized TSCs HL¹a², HL¹a², 1139 and HL²c²·CH₃COOH was tested in the doxorubicin-sensitive 1140 Colo205 and the multidrug-resistant Colo320 human colonic 1141 adenocarcinoma cell lines as well as in normal human 1142 embryonal lung fibroblast cells (MRC-5) by the colorimetric 1143 MTT assay. The data that resulted (expressed as the half-1144 maximal inhibitory concentration (IC₅0)) are collected in 1145

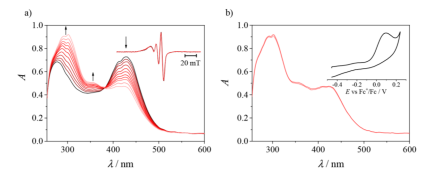


Figure 13. UV—Vis spectra measured simultaneously (a) upon anodic oxidation of 1 in the region of the first anodic peak (inset: time evolution of EPR spectra acquired at the first anodic peak) and (b) upon the back scan (inset: the corresponding in situ cyclic voltammogram).

1146 Table 2 and compared with those for triapine, doxorubicin, 1147 and CuCl₂.

Table 2. In Vitro Cytotoxicity (IC₅₀ Values in μ M) of Metal-Free Ligands HL^1-HL^3 , Copper(II) Complexes $Cu(HL^1)Cl_2$, $[Cu(L^2)Cl]$, and $Cu(HL^3)Cl_2$, and, of the Oxidized Species $HL^{1a'}$, $HL^{1a''}$, and $HL^{2c'}\cdot CH_3COOH$ in Colo205, Colo320, and MRC-5 Cell Lines after 72 h of Exposure

IC_{50} (μM)	Colo205	Colo320	MRC-5		
HL^1	>100	6.32 ± 0.49	>100		
HL^2	>100	>100	>100		
HL^3	48.2 ± 6.8	>100	>100		
$Cu(HL^1)Cl_2$	2.08 ± 0.12	2.21 ± 0.18	3.13 ± 0.17		
$[Cu(L^2)Cl]$	0.181 ± 0.039	0.159 ± 0.009	0.276 ± 0.049		
Cu(HL3)Cl2	26.6 ± 1.6	27.6 ± 1.6	>100		
$HL^{1a\prime}$	>25	>25	>25		
$HL^{1a''}$	>25	>25	>25		
HL ^{2c} ′⋅CH ₃ COOH	2.733 ± 0.059	0.188 ± 0.041	2.15 ± 0.10		
CuCl ₂	19.7 ^a	20.0 ^a	24.5 ^a		
triapine	3.34 ± 0.12	4.21 ± 0.46	10.2 ± 1.3		
doxorubicin	3.28 ^a	3.12 ^a	5.19 ^a		
^a Data are taken from ref 77.					

The metal-free ligands were either devoid of cytotoxicity or 1149 showed a weak response; only HL1 and HL3 revealed a 1150 somewhat higher activity against Colo320 and Colo205 cells, respectively, even though it was inferior to that of triapine. Notably, the copper(II) complexes are quite cytotoxic. So the 1153 effect of the copper(II) coordination is obvious in all cases. 1154 Low IC₅₀ values (0.16–2.2 μ M) were obtained for Cu(HL¹)-1155 Cl₂ and [Cu(L²)Cl] in both cancer cells (Colo205 and 1156 Colo320). To gain further insights into the cytotoxic behavior 1157 of the compounds, an apoptosis induction by lead compounds 1158 HL¹ and [Cu(L²)Cl] was investigated by a flow cytometry 1159 analysis of multidrug-resistant Colo320 cells stained with 1160 Annexin-V-FITC and propidium iodide (PI). The two 1161 compounds that displayed the highest cytotoxicity against 1162 this cell line were tested at two concentrations in the range of 1163 their IC₅₀ values. 12H-Benzophenothiazine (M627) and 1164 cisplatin were used as positive controls. The fluorescence of 1165 PI (FL3) was plotted versus Annexin-V fluorescence (FL1) as 1166 shown in Figure 14 for the positive controls and for the tested 1167 compounds at a chosen concentration. The percentage of the 1168 gated events regarding the early apoptosis, the late apoptosis 1169 and necrosis, and cell death is quoted in Table S8. According to these data, both compounds studied, HL^1 and $[Cu(L^2)Cl]$, 1170 can be considered as efficient apoptosis inducers.

The antiproliferative activity of $\hat{\bf 1}$ and ${\bf 2}'$ in the normal cells 1172 (MRC-5) was only slightly lower than in Colo205 and 1173 Colo320 cells, indicating a quite moderate selectivity for cancer 1174 cells. Complex ${\bf Cu(HL^3)Cl_2}$ was found to be less cytotoxic 1175 compared to the other two complexes tested, and the IC₅₀ 1176 values are similar to those of the copper(II) chloride, while the 1177 selectivity for cancer cells is obvious in this case (SI > 3). It is 1178 worth mentioning that the analogous α -N-pyridyl thiosemi- 1179 carbazones, that is, FTSC, AcTSC, and triapine, were reported 1180 to be cytotoxic in the low micromolar concentration range 1181 against several human cancer cells, the latter being the most 1182 potent among them (IC₅₀ values reported for triapine: 0.4–2.6 1183 μ M (in good agreements with the data quoted in Table 5), for 1184 FTSC: 1.9–10.6 μ M, for AcFTSC: 2.5–3.6 μ M in SW480, ³⁶ 1185 MES-SA, ³⁶ MES-SA/Dx5, ³⁶ HL60, ⁵⁸ 41M, ⁸⁰ SK-BR-3 ⁸⁰).

Their Cu(II) complexes were reported to possess a similar 1187 or even weaker cytotoxicity compared to the metal-free ligands, 1188 in contrast to complexes studied in the present work, which 1189 might indicate a distinct mode of action. It is also of note that 1190 the two-electron oxidized product HL^{2c'} revealed a superior 1191 antitumor activity in the two cancer cell lines over that of 1192 HL^{1a'} and HL^{1a''}. In agreement with this, closely related 2- 1193 formyl- and 2-acetylpyridine 2-benzothiazolyl hydrazones were 1194 shown to be potent cytotoxic drugs against a series of 17 1195 murine (e.g., L1210 lymphoid leukemia, P388 lymphocytic 1196 leukemia) and human cancer cells (e.g., HeLa uterine 1197 carcinoma, bone SOS, lung MB9812, lung A549). In addition, 1198 these compounds showed selectivity for the multidrug-resistant 1199 doxorubicin-selected uterine sarcoma cell line MES-SA/Dx5 1200 over parental or sensitive MES-SA cells.^{78,79}

Tyrosyl Radical Reduction in mR2 RNR. The TSCs 1202 HL¹-HL³ and their copper(II) complexes 1, 2', and 3 were 1203 found to effectively quench the tyrosyl radical in mR2 RNR in 1204 the presence of an external reductant (DTT). The time- 1205 dependent tyrosyl radical reduction in mR2 RNR by equimolar 1206 concentrations of TSCs and their respective copper(II) 1207 complexes, under reducing conditions, is shown in Figure 15. 1208 f15 The mR2 inhibition potency follows the order $HL^1 \approx triapine$ 1209 > HL³ > HL². The coordination to copper(II) was found to 1210 increase the tyrosyl radical quenching potential for all TSCs, 1211 which is in agreement with the observed lowering of IC₅₀ 1212 values in all cancer cell lines (Table 2). Complex 1 was shown 1213 to be as efficient as triapine, 17 reducing 100% of the tyrosyl 1214 radical in 3 min. Complexes 2' and 3 exhibited comparable 1215 reduction kinetics despite the fact that, among the investigated 1216 TSCs, HL² was found to be most inefficient. The favorable 1217

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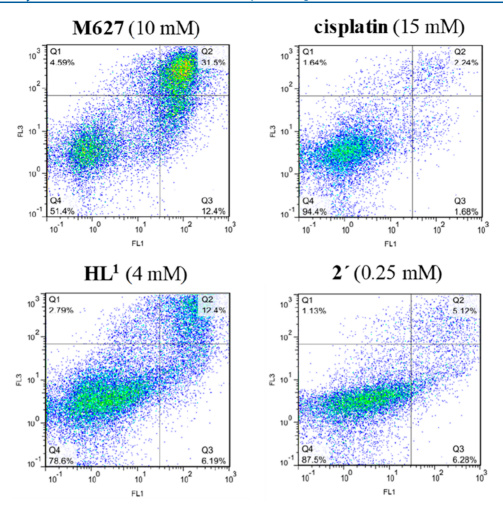


Figure 14. Quantification of apoptosis in Colo320 cells treated with HL^1 and 2' and M627 and cisplatin (as positive controls) using the Annexin-V/PI double staining assay. Colo320 cells were treated at the indicated concentration of the drugs. The dual parametric dot plots that combine the Annexin-V (FL1) and PI (FL3) fluorescence show the viable cell population in the lower-left quadrant Annexin-V-/PI- (Q4), the early apoptotic cells in the lower-right quadrant Annexin-V+/PI- (Q3), and the late apoptotic and necrotic cells in the upper-right quadrant Annexin-V+/PI+ (Q2). (Number of cells counted: 23 193 (M627), 20 262 (cisplatin), 33 193 (HL¹), and 19 312 (2')).

1218 impact of the copper(II) coordination on the HL² inhibitory 1219 activity is quite obvious, when the ability to quench the tyrosyl 1220 radical by HL² is compared to that of 2'. Interestingly, the two-1221 electron oxidized product of HL², namely, HL^{2c'}·CH₃COOH, 1222 is as potent as HL³ in the tyrosyl radical reduction.

The ability of HL^1-HL^3 and 1, 2′, and 3 to quench the 1224 tyrosyl radical correlates well with their first anodic redox 1225 potentials (0.82–0.88 V vs NHE) and (0.68–0.70 V vs NHE), 1226 respectively, which are well-compared with redox potential of 1227 hydroxyurea (+0.724 V), 8³ which reduced the tyrosyl radical in 1228 the R2 protein with an estimated redox potential of 1.0 ± 0.1 V 1229 vs NHE. 36 Note, however, that hydroxyurea, a well-known 1230 inhibitor of RNR and an anticancer drug, 8⁴ is a small molecule 1231 able to enter the hydrophobic R2 protein pocket, where the 1232 tyrosyl radical is buried. Finally, the two- and four-electron 1233 oxidized products of HL^1 , namely, $HL^{1a'}$ and $HL^{1a''}$, do not 1234 have an effect on the tyrosyl radical in the absence of DTT 1235 and, interestingly, cause an increase in the radical content in 1236 the presence of DTT (Figure S15).

 $_{1237}$ It has previously been shown that the radical content in mR2 $_{1238}$ may be slightly increased in the presence of DTT, as the result $_{1239}$ of the so-called radical reconstitution reaction, $_{17,85}$ in which the $_{1240}$ DTT-reduced diiron center in the reaction with molecular

oxygen is spontaneously oxidized through a series of 1241 intermediate states, generating the active Fe(III)- $^{2-}$ -Fe- 1242 (III)/ $^{2-}$ -Fe- 1242 (III)/ $^{2-}$ -Fe- 243 HL $^{1a'}$ and HL $^{1a''}$ (in reducing conditions) is much greater 1244 than that observed for DTT, providing evidence that the 1245 formation of the active iron/radical site in mR2 is more 1246 efficient when the DTT—reduced form of mR2 is oxidized by 1247 HL $^{1a'}$ or HL $^{1a''}$, than by molecular oxygen only.

Consistent with enzyme inhibition studies, which revealed a $_{1249}$ potent inhibition of mR2 RNR, compounds HL^1 , 1, and 2′ $_{1250}$ were found to increase the population of the S-phase in SW480 $_{1251}$ cells.

Cell Cycle Arrest. The perturbation effects of $10~\mu\text{M}$ HL¹, 1253 1, and 2' on the cell cycle progression of SW480 cells when 1254 compared to negative control are shown in Figure 16 and 1255~fi 16 Table S9, while the effects of 0, 1.0, and $10~\mu\text{M}$ are presented 1256 in Figure S16. It can be noted that the population of S-phase 1257 cells increased after an incubation with HL¹ (37.1%), complex 1258 1 (44.0), and 2' (46.5) compared with the negative control 1259 (29.8%). Gemcitabine (GC), a positive control, showed a 1260 canonical G1/S-phase arrest at the concentration of 0.01 μM 1261 with 26.8% of cells in the G1 phase and 62.3% of cells in the S 1262 phase compared to the negative control with 49.1% of cells in

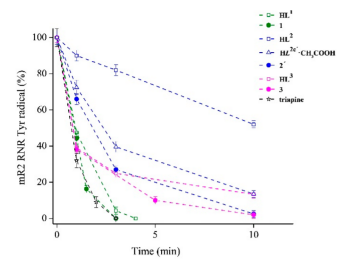


Figure 15. Tyrosyl radical reduction kinetics in mouse R2 RNR protein by TSCs HL^1 , HL^2 , HL^3 , and their corresponding copper complexes 1, 2′, and 3 as well as by the two-electron oxidized product of HL^2 ($HL^{2c'}$ ·CH₃COOH), in the presence of an external reductant, measured at 30 K by EPR spectroscopy and compared to triapine. The samples contained 20 μ M mR2 in 50 mM HEPES buffer, pH 7.60/100 mM KCl, 20 μ M compound in 1% (v/v) DMSO/H₂O, and 2 mM DTT.

1264 the G1 phase and 29.8% of cells in the S phase (Figure S17). 1265 An increase in the population of the S-phase cells by ca. 20% 1266 has been reported for a series of triapine analogues at 1267 concentrations from 0.25 to 5.0 μ M. ⁸⁰ The S-phase arrest is 1268 characteristic for cells treated with triapine. ⁸¹

These data indicate that there is a correlation between the 1270 ability of the compounds tested to inhibit R2 RNR and their 1271 ability to induce an S-phase arrest. Nevertheless, the inhibition 1272 of RNR does not appear to be the main mechanism underlying 1273 the antiproliferative activity of both TSCs studied herein and 1274 their copper(II) complexes.

ROS Generation. Since metal-free TSCs that enter the 1276 cells or are released from copper(I) complexes generated by a 1277 reduction of their copper(II) counterparts can react in the cells with iron(II), the redox activity of the $[Fe^{II}(L^1)_2]$ complex, prepared by the reaction of an anoxic aqueous solution of 1280 FeSO₄·7H₂O with a DMSO solution of HL¹ at a 1:2 molar 1281 ratio, was investigated by EPR spin-trapping experiments. To 1282 investigate whether this ferrous complex is able to generate 1283 ROS in the aqueous environment by a Fenton reaction, which 1284 is supposed to quench the tyrosyl radical of the mR2 enzyme, 1285 hydrogen peroxide was added into the system in the presence 1286 of 5,5-dimethyl-1-pyrroline N-oxide (DMPO) as the spin-1287 trapping agent. In the system containing H₂O₂, a four-line EPR 1288 signal characteristic for the OH-DMPO spin adduct was 1289 observed (Figure 17, black trace, EPR signal marked with 1290 circles)

Additionally a \cdot DMPO-OCH₃ spin adduct can be seen in the corresponding EPR spectrum as a consequence of the reaction of hydroxyl radicals with the DMSO solvent forming methyl radicals, which react with molecular oxygen resulting in the generation of peroxomethyl radicals serving as a source of DMPO-OCH₃ spin adducts (Figure 17, black trace, EPR 1297 signal marked with squares). Only a trace amount of carbon-1298 centered radicals was detected for Fe(II)/HL¹/DMPO in 1299 H₂O-DMSO in the absence of H₂O₂ (Figure 17, blue and red 1300 traces, EPR signal marked with stars). In this case DMSO acts

as a HO· scavenger, generating reactive carbon-centered 1301 radicals, which are trapped by DMPO. It is important to 1302 mention that no radicals were formed in the system of HL^1 / 1303 $H_2O_2/DMPO/H_2O-DMSO$ (not shown), which indicates the 1304 crucial role of the Fe(II) complex for ROS generation. 1305 Consequently complex $\left[Fe^{II}(L^1)_2\right]$ is redox-active in the 1306 Fenton reaction indicating the important role of the HL^1 1307 ligand for the observed antiproliferative activity against cancer 1308 cell lines and its ability to quench the tyrosyl radical in the 1309 mR2 protein. A direct reduction of the tyrosyl radical by 1310 iron(II) complexes with reported TSCs can also not be 1311 excluded. 16

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CONCLUSIONS

New triapine analogues bearing a redox-active p-amino- 1314 phenolic moiety and their copper(II) complexes have been 1315 synthesized and characterized by spectroelectrochemical and 1316 analytical techniques, which confirmed the noninnocent 1317 identity of the latter. The crystal structures of TSCs HL¹- 1318 HL³ and complexes [Cu(L¹⁻³)Cl] were studied by SC-XRD 1319 revealing the tridentate (N,N,S) coordination mode of the 1320 ligands. The presence of E and Z isomers of HL^1-HL^3 with a 1321 predominance of the first one in DMSO has been disclosed by 1322 1D and 2D NMR spectroscopy. These data along with DFT 1323 calculations on the model compound 2-formylpyridine TSC 1324 indicate that the Z/E isomerization involves an inversion at the 1325 aldimine nitrogen atom, rather than a tautomeric shift of the 1326 thioamide N2H proton to the pyridine nitrogen, followed by a 1327 rotation around the C-N1 bond as suggested previously. 44 1328 The relatively high Gibbs free energy barrier (~35.3 kcal/mol) 1329 for the Z/E conversion rules out the possibility of an 1330 isomerization at room temperature, in agreement with time- 1331 dependent NMR data.

A two-electron oxidative dehydrogenation of HL¹ by a 1333 reaction with 1 equiv of DDQ afforded the new species HL1a1 1334 containing a thiadiazole five-membered ring formed via a 1335 nucleophilic attack of a thione sulfur atom on an aldimine 1336 carbon atom. This is supported by frontier molecular orbitals 1337 (MOs) with the HOMO and LUMO located at opposite parts 1338 of the molecule of HL1. When 2 equiv of DDQ were used, a 1339 further two-electron oxidation coupled with a two-proton loss 1340 occurred at the 3,5-dimethyl-4-aminophenolic moiety to give 1341 the 3,5-dimethyl-1,4-benzoquinone imine unit in HL^{1a}". Also 1342 note that the coordinated ligand HL1 is able to form a thiazole 1343 five-membered ring in 4 via a sulfur attack on the carbon atom 1344 in position 2 or 6 of the 3,5-dimethyl-4-aminophenolic moiety. 1345 The arylated sulfur atom has lost the competition in binding to 1346 copper(II) for an end nitrogen atom due to the reduction of 1347 the electron-donating ability of the sulfur atom. The oxidation 1348 of HL² with PBQ in a 1:1 molar ratio furnished the two- 1349 electron oxidative cyclization product HL^{2b} and the diphenolic 1350 species HL^{2e}. A tentative mechanism of their formation is 1351 proposed. The pathway to HL^{2e} implies the formation of the 4- 1352 isothiocyanato-2,6-dimethylphenol intermediate. Treatments 1353 of HL2 with 1 and 2 equiv of PIDA afforded the two-electron 1354 oxidation product HL^{2c} and the four-electron oxidation 1355 product $HL^{2c''}$, respectively. In contrast to HL^1 – HL^3 , the $Z/_{1356}$ E isomerization was observed at room temperature for $HL^{2c'}$. 1357 The isolation and investigation of oxidation products of new 1358 TSCs was of interest also from the point of view of collecting 1359 spectroscopic data that might be useful for an eventual analysis 1360 of metabolites, which can be generated in vivo from the 1361 corresponding TSCs and their copper(II) complexes.

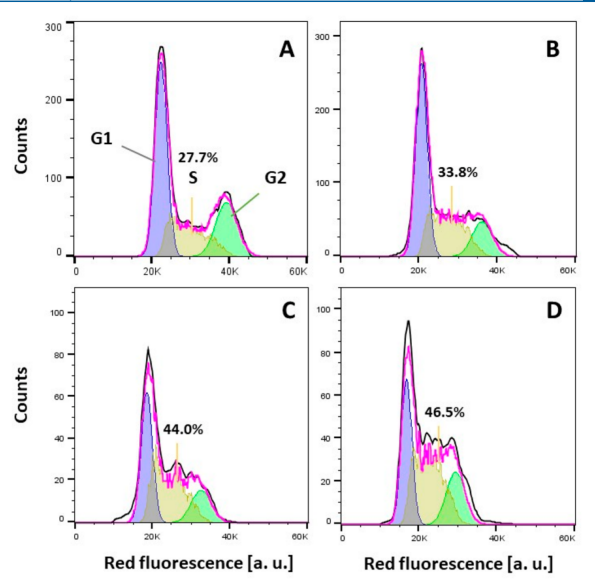


Figure 16. Flow cytometry analysis for a cell cycle distribution of SW480 cells induced by TSC HL^1 (B) and complexes 1 (C) and 2' (D) at the concentration of 10 μ M for 24 h compared to the negative control (DMSO) (A).

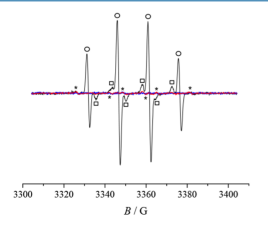


Figure 17. Experimental EPR spectra of $Fe(II)/HL^1/DMPO/H_2O_2$ in 5% (v/v) $H_2O-DMSO$ (black line), in the system of $Fe(II)/HL^1/DMPO$ in 5% (v/v) $H_2O-DMSO$ (blue line), and in 30% (v/v) $H_2O-DMSO$ (red line). Initial concentrations: $c(HL^1) = 0.2$ mM, $c(FeSO_4·7H_2O) = 0.1$ mM, c(DMPO) = 20 mM, $c(H_2O_2) = 10$ mM.

Solution equilibrium studies performed by UV-vis spec- 1363 trophotometry revealed the acidic p K_a values (3.01-3.95) of 1364 the pyridinium nitrogen and pK_a values greater than or equal 1365 to 10.55 for the hydrazinic-NNH and phenolic (PhOH) 1366 moiety of the metal-free ligands. The latter are neutral and 1367 stable at a physiological pH. However, they become air- 1368 sensitive upon deprotonation of the OH group in the basic pH 1369 range. The formation of high-stability monoligand copper(II) 1370 complexes was found in different protonation states in 1371 solution; namely, coordination via $(N_{pyridine},N,S)(H_2O)$, 1372 $(N_{pyridine}, N, S^{-})(H_{2}O)$, and $(N_{pyridine}, N, S^{-})(OH^{-})$ donor sets 1373 are probable. The complexes with a $(N_{pyridine}, N, S^-)(H_2O)$ 1374 coordination predominate in a wide pH range including pH 1375 7.4. Conditional stability constants determined for the 1376 $[Cu(L^1)]^+$ and $[Cu(L^3)]^+$ complexes by an EDTA UV-vis 1377 spectrophotometric competition experiment show the some- 1378 what higher stability of the $[Cu(L^3)]^+$ complex. The 1379 attachment of a phenolic moiety undoubtedly increases the 1380 lipophilicity of new Schiff bases and copper(II) complexes 1381 when compared to triapine and its copper(II) complex. The 1382 new complexes can be reduced by glutathione, the most 1383

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1384 abundant low molecular mass reducing agent in a cell, in a 1385 reversible redox reaction. According to the electrochemical 1386 studies complexes 1, 2', and 3 can undergo a redox process in a 1387 biologically accessible window (-0.4 to +0.8 V vs Fc^+/Fc). 1388 These findings suggest a possible role of the redox properties 1389 of the copper(II) complexes in their biological activity.

The metal-free ligands and several oxidized products showed 1391 no or only a moderate cytotoxicity against doxorubicin-1392 sensitive Colo205 and the multidrug-resistant Colo320 human 1393 colonic adenocarcinoma cell lines. Their copper(II) complexes 1394 revealed a high cytotoxic potency when compared to that of 1395 the corresponding metal-free ligands. [Cu(L²)Cl] showed the 1396 highest cytotoxic activity with IC50 values in the low 1397 micromolar concentration range and induced apoptosis, while Cu(HL³)Cl₂ has the highest selectivity for cancer cells over the normal fibroblast MRC-5 cells. The highest 1400 antiproliferative activity of $[Cu(L^2)Cl]$ is likely due to the more negative reduction potential when compared to those of 1402 1 and 3 and low reduction rate in reaction with GSH.³⁶ In 1403 addition, HL1-HL3 and their copper(II) complexes were 1404 found to efficiently quench the tyrosyl radical in mR2 RNR in 1405 the presence of DTT as an external reductant and increase the 1406 population of S-phase cells. The capacity of HL¹ to destroy the 1407 tyrosyl radical is almost identical with that of triapine, which is 1408 by the factor of 1000 a more potent R2 RNR inhibitor than 1409 hydroxyurea, a known clinical drug.¹⁷ Thus, the copper(II) 1410 complexes reported herein deserve further investigation as 1411 potential anticancer drugs.

1412 ASSOCIATED CONTENT

1413 Supporting Information

1414 The Supporting Information is available free of charge at 1415 https://pubs.acs.org/doi/10.1021/acs.inorgchem.1c01275.

> Synthesis of oxidized thiosemicarbazones and their copper(II) complexes, methods used for characterization of the compounds, atom labeling schemes used in the NMR resonances assignment, NMR and UV-vis spectra showing E/Z isomerization of TSCs in solution, 2D NMR spectra, UV-vis spectra of TSCs at different pH values and measured over time, tyrosyl radical kinetic behavior in absence and presence of DTT, crystal data and details of data collection, collected multinuclear NMR data, summarized ESI mass spectra, computational details (PDF)

1427 Accession Codes

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1428 CCDC 2074017-2074030 and 2074341 contain the supple-1429 mentary crystallographic data for this paper. These data can be 1430 obtained free of charge via www.ccdc.cam.ac.uk/data request/ 1431 cif, or by emailing data_request@ccdc.cam.ac.uk, or by 1432 contacting The Cambridge Crystallographic Data Centre, 12 1433 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

1434 AUTHOR INFORMATION

Email: anatshu@gmail.com

1435 Corresponding Authors

Vladimir B. Arion – Institute of Inorganic Chemistry, 1436 University of Vienna, A-1090 Vienna, Austria; o orcid.org/ 1437 0000-0002-1895-6460; Email: vladimir.arion@univie.ac.at 1438 Anatoly D. Shutalev -N. D. Zelinsky Institute of Organic 1439 1440 Chemistry, Russian Academy of Sciences, 119991 Moscow, Russian Federation; orcid.org/0000-0002-8038-8230; 1441

Complexes Research Group, University of Szeged, H-6720 1444 Szeged, Hungary; Email: peter.rapta@stuba.sk 1445 Eva A. Enyedy - Department of Inorganic and Analytical 1446 Chemistry, Interdisciplinary Excellence Centre and MTA-1447 SZTE Lendület Functional Metal Complexes Research 1448 Group, University of Szeged, H-6720 Szeged, Hungary; orcid.org/0000-0002-8058-8128; Email: enyedy@ 1450 chem.u-szeged.hu 1451 **Authors** 1452 Iuliana Besleaga – Institute of Inorganic Chemistry, University 1453 of Vienna, A-1090 Vienna, Austria 1454 Iryna Stepanenko - Institute of Inorganic Chemistry, 1455 University of Vienna, A-1090 Vienna, Austria 1456 Tatsiana V. Petrasheuskaya – Department of Inorganic and 1457 Analytical Chemistry, Interdisciplinary Excellence Centre and 1458 MTA-SZTE Lendület Functional Metal Complexes Research 1459 Group, University of Szeged, H-6720 Szeged, Hungary 1460 Denisa Darvasiova – Institute of Physical Chemistry and 1461 Chemical Physics, Faculty of Chemical and Food Technology, 1462 Slovak University of Technology in Bratislava, SK-81237 1463 Bratislava, Slovak Republic 1464 Martin Breza - Institute of Physical Chemistry and Chemical 1465 Physics, Faculty of Chemical and Food Technology, Slovak 1466 University of Technology in Bratislava, SK-81237 Bratislava, 1467 Slovak Republic; orcid.org/0000-0001-5995-0279 1468 Marta Hammerstad – Section for Biochemistry and Molecular 1469 Biology, Department of Biosciences, University of Oslo, NO- 1470 0316 Oslo, Norway 1471 Małgorzata A. Marć – Department of Inorganic and 1472 Analytical Chemistry, Interdisciplinary Excellence Centre, 1473 University of Szeged, H-6720 Szeged, Hungary; Department 1474 of Medical Microbiology and Immunobiology, University of 1475 Szeged, H-6720 Szeged, Hungary 1476 Alexander Prado-Roller – Institute of Inorganic Chemistry, 1477 University of Vienna, A-1090 Vienna, Austria 1478 Gabriella Spengler - Department of Medical Microbiology 1479 and Immunobiology, University of Szeged, H-6720 Szeged, 1480 Hungary; MTA-SZTE Lendület Functional Metal Complexes 1481 Research Group, University of Szeged, H-6720 Szeged, Hungary Ana Popović-Bijelić – Faculty of Physical Chemistry, 1484 University of Belgrade, 11158 Belgrade, Serbia; o orcid.org/ 1485 0000-0003-3121-2391 1486 https://pubs.acs.org/10.1021/acs.inorgchem.1c01275 1488 1489

Peter Rapta - MTA-SZTE Lendület Functional Metal

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Complete contact information is available at:

The authors declare no competing financial interest. 1490

ACKNOWLEDGMENTS

The financial support of the Austrian Science Fund (Grant No. 1492 I4729) and of the Russian Foundation for Basic Research 1493 (Grant No. 20-53-14002) is gratefully acknowledged. This 1494 work was also supported by the Lendület program of the 1495 Hungarian Academy of Sciences (LP2019-6/2019), the 1496 National Research, Development and Innovation Office- 1497 NKFIA through Project Nos. GINOP-2.3.2-15-2016-00038 1498 and FK 124240 and the and Ministry of Human Capacities, 1499 Hungary, Grant No. TKP-2020. T.V.P. is thankful for the 1500 support of Scholarship Foundation of the Republic of Austria 1501 (ICM-2019-14969). We are thankful to Dr. D. Dumitrescu for 1502

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1503 the collection of X-ray diffraction data for complex 2' at the 1504 XRD2 structural biology beamline, Elettra synchrotron.

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