## **Electronic Supplementary Information**

## Biliverdin-copper complex at physiological pH

Milena Dimitrijević, Jelena Bogdanović Pristov, Milan Žižić, Dalibor Stanković, Danica Bajuk-Bogdanović, Marina Stanić, Snežana Spasić, Wilfred Hagen, Ivan Spasojević<sup>\*</sup>

\*E-mail: : <u>redoxsci@gmail.com</u>



**Figure S1.** The stability of  $[BV]/[Cu^{2+}] = 1$  system in the presence of copper chelating agent bathocuproine in phosphate buffer (50 mM; pH 7.4). Green line - BV and Cu<sup>2+</sup> were incubated for 5 min before the addition of bathocuproine. Blue line - Cu<sup>2+</sup> was added to the buffer after BV and bathocuproine (arrow – absorbance line of bathocuproine complex with copper). It is important to note that bathocuproine is a non-innocent copper chelator. In the presence of bathocuproine, the reduction potential for the Cu<sup>2+</sup>/Cu<sup>1+</sup> couple is raised by approximately 500 mV, making Cu<sup>2+</sup> a powerful oxidant.<sup>1</sup> In the process, Cu<sup>2+</sup> oxidizes BV-Cu complex, resulting in BV degradation (note that BV-related absorance is completely lost (blue line)) and Cu release.



**Figure S2.** Changes in UV-Vis spectra of  $BV-Cu^{2+}$  systems prepared at high concentrations in 50 mM phosphate buffer, pH 7.4. Left:  $[BV]/[Cu^{2+}] = 1$ ;  $[BV] = [Cu^{2+}] = 0.3$  mM. Right:  $[BV]/[Cu^{2+}] = 2$ ; [BV] = 2mM;  $[Cu^{2+}] = 1$  mM. Aliquots were taken from each system and diluted to lower final concentrations (10 or 40  $\mu$ M, respectively), to allow spectra acquisition. It can be observed that the  $[BV]/[Cu^{2+}] = 1$  system underwent degradation within 10 min, whereas the  $[BV]/[Cu^{2+}] = 2$  system was relatively stable for 5 h.



**Figure S3.** Speciation diagrams of  $Cu^{2+}$  in phosphate buffer (50 mM) at two concentrations – 40  $\mu$ M (top) and 300  $\mu$ M (bottom). Diagrams were prepared in Hydra-Medusa Software, using the presented parameters.



**Figure S4.** Comparison of UV-Vis spectra of biliverdin (BV) in the absence and the presence of  $Cu^{2+}$  in phosphate buffer (50 mM; pH 7.4) and in DMSO. Spectra were recorded after 5 min incubation period.



**Figure S5. HESI-MS** spectrum (full scan mode) of the system with  $[BV] = 20 \ \mu M$  and  $[Cu] = 40 \ \mu M$  ([BV]/[Cu] = 0.5). Assignation: *m/z* 643, BV-Cu complex; *m/z* 264, propentdyopent; *m/z* 320, propentdyopent complex with copper.<sup>2</sup>



**Figure S6.** <sup>1</sup>H NMR spectra of biliverdin (0.3 mM) in DMSO-d6 in the absence or the presence of  $Cu^{2+}$  at equimolar concentration. The bottom spectrum was recorded in a copper-free system, with the equimolar amount of D<sub>2</sub>O as in experiments with copper. It can be observed that  $Cu^{2+}$  and D<sub>2</sub>O induced similar (but not identical) changes. Changes in NH signals could not be observed in the presence of D<sub>2</sub>O because of the chemical exchange. The peaks were assigned in accordance to previous reports.<sup>3</sup> The signals labeled with 4 come from two types of protons (- CH= and =CH<sub>2</sub>). The spectra were collected within 20 min after sample preparation.



**Figure S7.** Scan rate analysis of BV and BV-Cu complex in phosphate buffer (50 mM; pH 7.4). (a) Cyclic voltammograms of BV (0.4 mM) in absence or presence of Cu<sup>2+</sup> (0.2 mM) at the boron doped diamond electrode obtained at different scan rates (v = 0.025-0.5 V/s). (b) The dependence between anodic peak currents *I* at potentials  $E_{pal}$  (oxidation of BV), and  $E_{pa2}$  and  $E_{pa3}$  (oxidation of BV-Cu complex(es)) and  $v^{1/2}$ . Linear fit and  $R^2$  values are presented. (c) *D* for BV and BV-Cu complex(es). Randles–Sevick equation (in the box): *n*, number of transferred e

(1e<sup>-</sup> for all peak currents), *A*, area of the working electrode (0.0707 cm<sup>2</sup>); *C*, concentration of redox species in solution ([BV] = 0.4 mM; [BV-Cu] = 0.2 mM). Results are presented as means ( $\pm$  standard deviation) of measurements made at various v. All three *D* values were statistically different (p < 0.001; ANOVA with *post hoc* Duncan's test).

Line [cm <sup>-1</sup> ]	Assignment	References
1619	Lactam stretching	4
1470	C–C deformation, likely between rings	4,5
1443	Stretching CC, stretching CN	6
1393	CH3 asymmetric deformation	6
1362	CH3 deformation	6
1331	In plane bending CH(CH3)	6
1303	CH wagging	6
1254	Lactam ring	6
1179	C–H twisting	4
1101	Stretching C–C, stretching C–N	6
1003	Asymmetric CH <sub>3</sub> deformation	6
971	C–C stretching mixed with C–H rocking	4
954	Stretching C–C–O	7
844	Stretching ring	6
767	In plane ring deformation	6
717	Out of plane ring deformation	6
684	Out of plane bending C=O	6

**Table S1.** Raman spectral lines that were observed for BV (1 mM), using the  $\lambda = 532$  nm laser excitation line.

## References

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