

**Title: Novel Seleno-Hydantoin Palladium(II) Complex – Antimigratory, Cytotoxic and Prooxidative Potential on Human Colon HCT-116 and Breast MDA-MB-231 Cancer Cells**

Running title: Seleno-Hydantoin and its Pd(II) Complex Biological Activity

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**Abstract**

Selenium and palladium containing compounds separately exert multifunctional effects on cells. While selenium containing compounds usually exert antioxidative properties, palladium(II) containing compounds are cytotoxic and prooxidative. Here we investigated biological effects of bicyclic seleno-hydantoin Cis-7a-Ethyl-5-methyl-5-phenylselanyl-methyl-tetrahydro-pyrrolo[1,2-c]imidazole-1,3-dione (Hid-Se), and its palladium(II) complex, trans-bis-(Cis-7a-Ethyl-5-methyl-5-phenylselanyl-methyl-tetrahydro-pyrrolo[1,2-c]imidazole-1,3-dionato) palladium(II) chloride ((Hid-Se)<sub>2</sub>Pd) on human colon HCT-116 and breast MDA-MB-231 cancer cell lines. Hid-Se and (Hid-Se)<sub>2</sub>Pd showed prooxidative and cytotoxic character. In all performed experiments (Hid-Se)<sub>2</sub>Pd proved to be more active, i.e. this substance exerted greater prooxidative effect, cytotoxicity and influence on cell migration potential. Even though Hid-Se and (Hid-Se)<sub>2</sub>Pd enhanced migration of HCT-116 cells, very important feature of these substances is the strong antimigratory potential on metastatic MDA-MB-231 cells.

Keywords: Cancer; Cell Migration; Oxidative stress; Palladium; Selenium

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1 **Novel Seleno-Hydantoin Palladium(II) Complex – Antimigratory, Cytotoxic and**  
2 **Prooxidative Potential on Human Colon HCT-116 and Breast MDA-MB-231 Cancer**  
3 **Cells**

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13 **Short title:** Bicyclic Seleno-Hydantoin and its Pd(II) Complex Biological Activity

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24 **Abstract**

25 Selenium and palladium containing compounds separately exert multifunctional effects on  
26 cells. While selenium containing compounds usually exert antioxidative properties,  
27 palladium(II) containing compounds are cytotoxic and prooxidative. Here we investigated  
28 biological effects of bicyclic seleno-hydantoin *Cis-7a-Ethyl-5-methyl-5-phenylselanylmethyl-*  
29 *tetrahydro-pyrrolo[1,2-c]imidazole-1,3-dione* (**Hid-Se**), and its palladium(II) complex, *trans-*  
30 *bis-(Cis-7a-Ethyl-5-methyl-5-phenylselanylmethyl-tetrahydro-pyrrolo[1,2-c]imidazole-1,3-*  
31 *dionato) palladium(II) chloride* (**(Hid-Se)<sub>2</sub>Pd**) on human colon HCT-116 and breast MDA-  
32 MB-231 cancer cell lines. **Hid-Se** and **(Hid-Se)<sub>2</sub>Pd** showed prooxidative and cytotoxic  
33 character. In all performed experiments **(Hid-Se)<sub>2</sub>Pd** proved to be more active, i.e. this  
34 substance exerted greater prooxidative effect, cytotoxicity and influence on cell migration  
35 potential. Even though **Hid-Se** and **(Hid-Se)<sub>2</sub>Pd** enhanced migration of HCT-116 cells, very  
36 important feature of these substances is the strong antimigratory potential on metastatic  
37 MDA-MB-231 cells.

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39 **Keywords:** Cancer, Cell Migration, Oxidative stress, Palladium, Selenium

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## 47 **Introduction**

48 Cancer (*malignant neoplasia*) represents a large disease group, which is characterized by  
49 unregulated cell differentiation and formation of malignant tumors, which may invade other  
50 parts of the body affecting the basic physiological functions (Craig A. Almeida 2010). In the  
51 field of cancer research the most of chemotherapeutic protocols are created with the aim of  
52 possessing the cytotoxic effects (Angelis et al. 2013; Blagosklonny and Fojo 1999). However,  
53 as primary tumors are significantly different from their metastatic analogues, the effect of a  
54 chemotherapeutic is not same on these types of tumors. Metastatic cells exhibit increased  
55 motility and increased cell migration from tumor stroma to other parts of the body (Polyak  
56 and Weinberg 2009). Compared to traditional chemotherapy and radiotherapy methods, there  
57 are numerous studies which indicate anti-metastatic effects (Deb et al. 2014; Eckhardt et al.  
58 2012).

59 Modern medicine is still largely focused on the application of platinum-containing (Jevtic et  
60 al. 2014; Kalinowska-Lis et al. 2008) and palladium-containing (Matovic et al. 2013; Sabo et  
61 al. 2004) chemotherapeutic cytotoxic agents. On the other hand, selenium has proven to be an  
62 important supplement for the suppression of the side effects of chemotherapy (Brodin et al.  
63 2015; Markovic et al. 2011; Mut-Salud et al. 2016; Schroeder et al. 2004). Also, selenium-  
64 containing compounds exert inhibition of migration of cancer cells (Chen et al. 2013; Zec et  
65 al. 2012; Liu et al. 2015). Combining palladium and selenium in the same structure could  
66 provide a novel substance with benefits from the both elements. Substances, presented in this  
67 study combine bicyclic hydantoin structure and selenium in ligand, and additionally  
68 palladium(II) in complex of mentioned ligand. The goal of this study was to examine the  
69 cumulative effect of a hydantoin structure, selenium and palladium on cancer cells.  
70 Hydantoins are widely used compounds in cancer studies (Hmuda et al. 2014; Mudit et al.  
71 2009). Moreover, bicyclic hydantoins are also studied (Ananda Kumar et al. 2009; Basappa et

72 al. 2009). Although seleno-organic compounds (Bugarcic et al. 2015; Kosaric et al. 2014) and  
73 palladium(II) complexes of hydantoins are investigated (Kushev et al. 2002; Varbanov et al.  
74 2010), according to our knowledge there are no records regarding antitumor activities of  
75 bicyclic seleno-hydantoins and their Pd(II) complexes.

76 In our investigations we used human adherent colorectal cancer cell line, HCT-116 and  
77 human mammary gland breast carcinoma metastatic cells, isolated from lung pleura, MDA-  
78 MB-231. Besides the tissue source difference of these two cell lines, one of their most  
79 significant differences is metastatic potential. While HCT-116 cells originate from primary  
80 tumor, MDA-MB-231 cells are of metastatic origin. Thus, in this study, in addition to  
81 investigation of effects of cytotoxicity and impact on redox status, we have chosen also to  
82 investigate the effects of tested substances on migratory potential of cells.

83

## 84 **Experimental**

### 85 *Material and methods*

#### 86 **Synthesis of Hid-Se And Its Palladium(II) Complex (Hid-Se)<sub>2</sub>Pd**

87 Synthesis of ligand, a seleno derivate of fused bicyclic hydantoin (*cis*-7a-ethyl-5-methyl-5-  
88 phenylselanyl-methyl-tetrahydro-pyrrolo[1,2-c]imidazole-1,3-dione), is shown in Scheme 1  
89 and reported recently (Šmit and Pavlović 2015). In the reaction of the ligand with PdCl<sub>2</sub>, a  
90 complex with [PdCl<sub>2</sub>(ligand)<sub>2</sub>] stoichiometry was obtained.

91 A solution of 17.7 mg (0.1 mmol) PdCl<sub>2</sub> in acetonitrile (15 mL) was heated under reflux for 2  
92 hours. Yellow colored solution was cooled to room temperature and 70.3 mg (0.2 mmol) of  
93 ligand (Hid-Se) dissolved in acetonitrile (2.5 mL) was added dropwise. Reaction mixture was

94 stirred during 24 hours. Resulting yellow powder was filtered off and dried. Yield 60.9 mg  
95 (69%).

96 The structure of the complex was confirmed on the basis of elemental analyses and spectral  
97 data.

98 IR (KBr)  $\nu_{\max}$ : 3419, 3160, 3055, 2970, 1767, 1701, 1576, 1403, 1385, 1241, 1097, 922, 791,  
99 740, 687, 647  $\text{cm}^{-1}$ .  $^1\text{H}$  NMR (200 MHz, DMSO- $d_6$ )  $\delta$  0.78 (t,  $J = 7.3$  Hz, 3H), 1.41 (s, 3H),  
100 1.58-1.69 (m, 1H), 1.73 (q,  $J = 7.3$  Hz, 2H), 1.97-2.13 (m, 3H), 3.54 (d,  $J = 12.5$  Hz, 1H),  
101 3.75 (d,  $J = 12.5$  Hz, 1H), 7.20-7.41 (m, 3H), 7.47-7.65 (m, 2H), 10.77 (bs, 1H).  $^{13}\text{C}$  (50 MHz,  
102 DMSO- $d_6$ )  $\delta$  7.5, 29.2, 30.0, 30.8, 35.8, 39.6, 64.2, 74.9, 127.2, 129.5, 130.6, 131.8, 156.8,  
103 176.5. Anal. calc. for  $\text{C}_{32}\text{H}_{40}\text{Cl}_2\text{N}_4\text{O}_4\text{PdSe}_2$  C: 43.68, H: 4.58, N: 6.41%; found C: 43.36, H:  
104 4.40, N: 6.76%.

## 105 **Chemicals**

106 Dulbecco's Modified Eagle Medium (DMEM) and PBS were obtained from GIBCO,  
107 Invitrogen, USA. Foetal bovine serum (FBS) and trypsin-EDTA were from PAA (The Cell  
108 Culture Company, Pasching, Austria). Dimethyl sulfoxide (DMSO), Bovine Serum Albumin  
109 (BSA), 3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide (MTT), and nitro blue  
110 tetrazolium (NBT) were obtained from SERVA, Heidelberg, Germany. Polyvinyl alcohol  
111 mounting medium, N-1-naphthylethylenediamine dihydrochloride were purchased from Fluka  
112 chemie GMBH, Buchs, Switzerland. Sulfanilamide and sulphosalicylic acid were purchased  
113 from MP Hemija Belgrade, Serbia. 5,5'-dithio-bis(2-nitrobenzoic acid) was purchased from  
114 Sigma Chemicals Co., St Louis, MO, USA. Paraformaldehyde originated from Merck,  
115 Germany. All solvents and chemicals were of analytical grade. The stock solutions were  
116 prepared in 100% DMSO at final concentration of 100 mM. Subsequent dilution was  
117 performed with DMEM cell growing medium, where in the highest treatment concentration  
118 (500  $\mu\text{M}$ ) DMSO concentration was at 0.05%, which is non-toxic to investigated cells.

## 119 **Cell Preparation and Cultivation**

120 Human colorectal HCT-116 and breast MDA-MB-231 cancer cell lines were purchased from  
121 the American Tissue Culture Collection (Manassas, VA, USA). The cells were cultivated in  
122 75 cm<sup>2</sup> culture flasks under controlled conditions in a humidified atmosphere with 5% CO<sub>2</sub> at  
123 37 °C in DMEM serum supplemented with 10% FBS and 100 IU/mL penicillin and 100  
124 µg/mL streptomycin until reaching a confluency of about 80%. For MTT, NBT and Griess  
125 assays, 10<sup>4</sup> cells were seeded in 96-well plates in triplicates for each treatment concentration,  
126 while for GSH assay it was seeded 5x10<sup>4</sup> cells per well. For immunofluorescence  
127 determination of expression of iNOS protein 7x10<sup>4</sup> cells were seeded in triplicates in 6-well  
128 plates on glass coverslips (Thermo Scientific), while for migratory potential inhibition assay  
129 10<sup>5</sup> cells per well were seeded in transwell cell culture chamber.

## 130 **Determination of Cell Viability and Redox Status Parameters**

131 Cell viability was determined by MTT (Mosmann 1983) assay. Redox parameters: superoxide  
132 anion radical determined by NBT (Auclair and Voisin 1985), nitrites by Griess (Griess 1879)  
133 and glutathione by GSH assay (Baker et al. 1990). For investigation of influence of tested  
134 substances on cell viability and redox status we used standardized procedures briefly  
135 described in our previous papers (Kosaric et al. 2014; Petrovic et al. 2014; Petrovic et al.  
136 2015). For the purposes of redox status assays we used concentrations of 1, 10, 50 and 100  
137 µM, while IC<sub>50</sub> values are determined as a plot of % cytotoxicity versus sample  
138 concentrations. All results were expressed as µM.

## 139 **Inducible Nitric Oxide Synthase (iNOS) Protein Expression (Immunofluorescence** 140 **Microscopy)**

141 Inducible nitric oxide synthase (iNOS) protein expression in HCT-116 and MDA-MB-231  
142 cells was detected by immunofluorescence method (Javois 1999). Cells were cultured in 6-

143 well plates on glass coverslips. When cells were at about 80% confluence the supporting  
144 medium was removed and the cells were treated with medium containing tested substances at  
145 final concentration of 50  $\mu$ M. After 24 h medium was removed and cells were triple washed  
146 with PBS (pH 7.2). Next, the cells were fixed with 4% *p*-formaldehyde in PBS for 20 min at  
147 37 °C. After the fixation, the cells were washed three times with PBS and then permeabilized  
148 with methanol at -20 °C for 2 min, washed with PBS three times and blocked on non-specific  
149 binding sites using 1% Bovine Serum Albumin-BSA for 20 min. The fixed cells were  
150 incubated with 20  $\mu$ g/mL anti-iNOS specific primary antibody (RD Systems) for 1 h at 37 °C.  
151 Sample coverslips were then washed twice and incubated with anti-mouse secondary antibody  
152 conjugated with Alexa448 (Thermo Scientific) at a 1:200 dilution in PBS. DAPI was used to  
153 stain the cell nuclei (blue) at 1:1000 dilutions. Sample coverslips were washed twice and  
154 mounted on glass slide by polyvinyl alcohol mounting medium. The cell visualization was  
155 performed at Nikon inverted fluorescent microscope (Ti - Eclipse) at 600x magnification by  
156 using Nikon NIS-Elements Advanced Research software.

### 157 **Migratory Potential Inhibition Assay**

158 Principle of migratory assay is based on potential of investigated substance to inhibit or  
159 promote cell migration. Cells are placed on the upper layer of a cell permeable membrane  
160 with the medium containing tested substance. After incubation period, the cells which are  
161 migrated through the membrane were stained and counted. Migration of HCT-116 and MDA-  
162 MB-231 cancer cells through 8  $\mu$ m filter pores was followed using Transwell cell culture  
163 chamber (ThinCert with translucent membranes, Greiner Bio-One GmbH, Germany)  
164 (Entschladen et al. 2005). Firstly, cells that are cultured in DMEM medium with 10% FBS  
165 were twice washed with PBS and resuspended in the serum free DMEM medium. After  
166 centrifugation at 800 rpm for 3 min the  $10^5$  cells per well were resuspended in 500  $\mu$ L of  
167 serum free DMEM medium with addition of **Hid-Se** and **(Hid-Se)<sub>2</sub>Pd** and plated in 24-well



168 plates onto the upper compartment of the chamber. The final concentrations of substances  
169 were 10 and 100  $\mu$ M resp. In the lower compartment of the chamber 750  $\mu$ L of complete  
170 medium (DMEM with 10% FBS as chemoattractant) was added. The control cells have not  
171 been treated with the substances. After 24 h of incubation at 37 °C and 5% CO<sub>2</sub> cells were  
172 washed three times with PBS and fixed with 4% *p*-formaldehyde for 20 minutes at room  
173 temperature, followed by washing three times with PBS. After fixation, the cells were  
174 carefully and thoroughly removed with cotton swab. Remains of cells at the lower part of the  
175 Transwell membrane were stained with 0.1% of Crystal Violet in 200 mM MES buffer, pH  
176 6.0 for 10 minutes at room temperature. After washing, and drying at room temperature  
177 overnight, membranes were cut and placed in another 24-well plate. Crystal Violet stained  
178 membranes were resuspended in 100  $\mu$ L 10% acetic acid and 80  $\mu$ L of stained acetic acid  
179 solution was moved to a 96-well microtiter plate with subsequent reading of optical density at  
180 595 nm at Microplate reader.

## 181 *Statistics*

182 All experimental data were expressed as mean  $\pm$  standard error (SE). Biological activity  
183 assays are performed in triplicate for each dose. Statistical significance was determined using  
184 the one-way ANOVA test for multiple comparisons. A p value < 0.05 was considered as  
185 significant. The magnitude of correlation between variables was done using SPSS (Chicago,  
186 IL) statistical software package (SPSS for Windows, version 17, 2008). The IC<sub>50</sub> values were  
187 calculated from the dose curves by a computer program (CalcuSyn).

188

## 189 **Results**

### 190 *Cytotoxic Effects*

191 The cytotoxicity of investigated substances was determined by MTT assay. The results  
192 were expressed graphically in Figure 1 and as IC<sub>50</sub> values presented in Table 1.  
193 Evaluation on HCT-116 cells revealed that investigated substances exhibit cytotoxic  
194 character. **Hid-Se** induced decreasing of cell viability in dose and time depended  
195 manner, with significant toxicity 72 h from treatment. On the other hand, **(Hid-Se)<sub>2</sub>Pd**  
196 exhibits more significant cytotoxic effect acutely, i.e. after 24 h. Based on the obtained  
197 results, **(Hid-Se)<sub>2</sub>Pd** is more cytotoxic to HCT-116 cells than **Hid-Se**. On MDA-MB-  
198 231 cells, investigated substances showed cytotoxic character exhibited in dose and  
199 time dependent manner, i.e. with increasing the substance concentration cell viability  
200 decreased which is more evident 72 h from treatment. After 24 h from treatment the  
201 both substances exhibited no cytotoxic effect, while after 72 h the cytotoxic effect was  
202 obvious. The more significant effect was obtained with **(Hid-Se)<sub>2</sub>Pd**. From the results  
203 explaining cytotoxic character one could observe that HCT-116 cells are more  
204 sensitive to the investigated substances than MDA-MB-231 cells.

## 205 *Redox Status*

### 206 **Superoxide Anion Radical (O<sub>2</sub><sup>•-</sup>) Concentration**

207 Superoxide anion radical, O<sub>2</sub><sup>•-</sup> is an important indicator of reactive oxygen species (ROS)  
208 level (Hancock et al. 2001). O<sub>2</sub><sup>•-</sup> was determined by spectrophotometric NBT assay, and  
209 results of measurement are depicted in Table 2. Measurement on HCT-116 cells revealed that  
210 **Hid-Se** and **(Hid-Se)<sub>2</sub>Pd** induced significant change in O<sub>2</sub><sup>•-</sup> content.

211 After 24 h **Hid-Se** induced decreasing of O<sub>2</sub><sup>•-</sup> in low concentrations, while higher  
212 concentrations significantly increased O<sub>2</sub><sup>•-</sup>. On the other hand, 72 h from treatment **Hid-Se**  
213 decreased O<sub>2</sub><sup>•-</sup> content. Similar decreasing was observed with **(Hid-Se)<sub>2</sub>Pd** in both treatment  
214 periods. **(Hid-Se)<sub>2</sub>Pd** exerted stronger reduction of O<sub>2</sub><sup>•-</sup> content than **Hid-Se** on HCT-116

215 cells. While tested substances, in general, decreased  $O_2^{\bullet-}$  in HCT-116 cells, on MDA-MB-231  
216 cells they acted as prooxidants, i.e. induced significant increasing in  $O_2^{\bullet-}$  content 24 and 72 h  
217 from treatment. This effect was stronger after 24 h, where **(Hid-Se)<sub>2</sub>Pd** again showed greater  
218 influence. Comparison of these two cell lines showed that  $O_2^{\bullet-}$  production was greater on  
219 MDA-MB-231 cells.

## 220 **Nitrites (NO<sub>2</sub><sup>-</sup>) Concentration**

221 Concentration of nitrites indicates quantity of nitrosonium ions (NO<sup>+</sup>) in anaerobic conditions.  
222 In water, the final product of aerobic phase reaction between nitrogen oxide (NO) and O<sub>2</sub> is  
223 nitrogen dioxide (NO<sub>2</sub>), which quickly combines with excess of NO, forming nitrogen  
224 trioxide (N<sub>2</sub>O<sub>3</sub>) that is hydrolyzed to nitrites. So, nitrite concentration may indicate level of  
225 NO and other reactive nitrogen species (RNS) in cells (Lancaster 2006). Results of  
226 measurement of nitrite level are presented in Table 3. On HCT-116 cells in treatment with  
227 **Hid-Se** it was observed decreasing of nitrites, except in the highest concentration which  
228 induced significant increasing. **(Hid-Se)<sub>2</sub>Pd** significantly induced increasing of nitrites. On  
229 MDA-MB-231 cells after 24 h **Hid-Se** in lower concentrations induced decreasing, while  
230 higher concentrations induced increasing of nitrite concentration. After 72 h change of nitrites  
231 was not significant as after 24 h. **(Hid-Se)<sub>2</sub>Pd** predominantly induced significant increasing of  
232 nitrite level. Similarly to NBT test, measurement of nitrite concentration indicated MDA-MB-  
233 231 cells as more sensitive to investigated substances when compared to HCT-116 cells.

## 234 **Inducible Nitric Oxide Synthase (iNOS) Protein Expression**

235 Nitric oxide synthase is a protein which induces the synthesis of NO from L-arginine (Liu and  
236 Gross 1996). In order to investigate the possible origin of the increase of nitrites we  
237 considered examination of the expression of iNOS protein. On HCT-116 cells 24 h from  
238 treatment **Hid-Se** in concentration of 50 μM decreased expression of iNOS protein (Figure 2).

239 Similarly, in the same measuring conditions **Hid-Se** significantly decreased nitrite content  
240 (Table 3). On the other hand, **(Hid-Se)<sub>2</sub>Pd** significantly increased iNOS protein expression,  
241 which also could be related to the nitrite production (Table 3). On MDA-MB-231 cells both  
242 substances induced significant increasing of iNOS protein expression.

243

#### 244 **Reduced Glutathione (GSH) Concentration**

245 Glutathione is an important tripeptide responsible for neutralizing of ROS/RNS and thus for  
246 maintaining the redox equilibrium in eukaryotic cell (Pompella et al. 2003). Table 4  
247 represents the effects of investigated substances on GSH level in HCT-116 and MDA-MB-  
248 231 cells. On HCT-116 we observed a moderate effect on GSH level change, i.e. **Hid-Se** did  
249 not induced any statistically significant changing of GSH concentration, while cells treated  
250 with **(Hid-Se)<sub>2</sub>Pd** slightly increased GSH level. This increasing was slightly higher after 24 h.  
251 On MDA-MB-231 cells it was observed significant increasing in GSH level with both  
252 substances. It could be concluded that greater increasing in GSH for **(Hid-Se)<sub>2</sub>Pd** is followed  
253 by greater increasing in ROS/RNS induced with this substance. Thus, cells respond to  
254 changing in redox equilibrium by lesser or greater induction in GSH production.

#### 255 **Migratory Potential Inhibition**

256 To evaluate influence of tested substances on migratory potential we examined the migration  
257 of cells through Transwell membrane assembled in 24-well plate. The size of pores on  
258 Transwell membrane is 8.0  $\mu\text{m}$ , i.e. small enough compared to cancer cells. Thus,  
259 investigated cells influenced by substances, could only actively migrate through the pores in  
260 greater or smaller extent. Influence of tested substances on migratory potential of HCT-116  
261 cells is less significant when compared to the results obtained with MDA-MB-231 cells  
262 (Figure 3). Investigated substances induced increasing of HCT-116 motility. **Hid-Se** in

263 concentration of 10  $\mu\text{M}$  showed no effect, while concentration of 100  $\mu\text{M}$  increased migration  
264 for about 15%. On the other hand **(Hid-Se)<sub>2</sub>Pd** increased cell migration in concentrations of  
265 10  $\mu\text{M}$  and 100  $\mu\text{M}$  for 60 and 80% resp. MDA-MB-231 cells are influenced in a great extent,  
266 especially in concentration of 100  $\mu\text{M}$ . **Hid-Se** induced no statistically significant change in  
267 concentration of 10  $\mu\text{M}$ , but 100  $\mu\text{M}$  reduced migration for about 65%. **(Hid-Se)<sub>2</sub>Pd** in  
268 concentration of 10  $\mu\text{M}$  increased migration for 20%, while 100  $\mu\text{M}$  reduced migration for  
269 60%. All results presented in Figure 3 are recalculated in relation with the number of survived  
270 cells estimated in MTT assay.

271

## 272 **Discussion**

273 Results presented in this paper are the first investigation of biological activity of such fused  
274 bicyclic seleno-hydantoin and its Pd(II) complex on cancer cell lines. Cytotoxic character of  
275 palladium(II) complexes, hydantoins and seleno-derivatives is proved and well explained in  
276 numerous articles (Ananda Kumar et al. 2009; Basappa et al. 2009; Bugarcic et al. 2015;  
277 Hmuda et al. 2014; Kosaric et al. 2014; Kushev et al. 2002; Matovic et al. 2013; Mudit et al.  
278 2009; Sabo et al. 2004; Varbanov et al. 2010). Comparing with results obtained in this article,  
279 many of tested Pd(II) complexes, seleno-organic compounds, hydantoins and their  
280 combinations exerted considerable cytotoxicity on cancer cell lines. Such pronounced  
281 cytotoxicity in most cases is closely related to their prooxidative character. This point of view  
282 is important, but one of the most important features that also could be followed is the  
283 influence on potential of cancer cells to migrate. Our results suggest that these compounds  
284 possess potential to reduce migration of metastatic MDA-MB-231 cells.

285 At the beginning it was considered that palladium(II) complexes possess no anti-tumor  
286 properties. Compared to cisplatin, cispalladium, *cis*-[Pd(NH<sub>3</sub>)<sub>2</sub>Cl<sub>2</sub>] complex does not show

287 anti-tumor activity. Coordination chemistry of Pd(II) and Pt(II) is very similar whereas their  
288 structural and equilibrium behavior of the solutions are very similar (Bugarčić et al. 2015),  
289 but introduction of specific ligands in coordination with Pd(II) can result with great anti-  
290 tumor activities (Ulukaya et al. 2011, Divsalar et al. 2011). Compared to action of the  
291 cisplatin presented in our previous article (Petrovic et al. 2015), our results showed that **Hid-**  
292 **Se** exerts lower cytotoxic effects, while **(Hid-Se)<sub>2</sub>Pd** showed greater effect 24 h after  
293 treatment on HCT-116 cells and lower but comparable effect on MDA-MB-231 cells. From  
294 the presented results it could be concluded that **Hid-Se** and especially **(Hid-Se)<sub>2</sub>Pd** exert  
295 cytotoxic character. More significant effect showed **(Hid-Se)<sub>2</sub>Pd**, especially on HCT-116  
296 cells. Cytotoxicity is closely related to enhanced production of ROS/RNS, i.e. superoxide  
297 anion radical and nitrites. The produced  $O_2^{\bullet-}$  could be from many possible origins. One is that  
298 structures of investigated substances possess asymmetric electron densities that could provide  
299 electron(s) that could subsequently start radical chain reaction. Another source certainly could  
300 be explained by Fenton reaction inside the cells.  $H_2O_2$  created in cells easily reacts with iron  
301 yielding  $O_2^{\bullet-}$ . Besides iron, also many transition metals could catalyze Fenton reaction, such  
302 as palladium, platinum and rhodium (Halliwell and Gutteridge 2007). Nitrites, and thus the  
303 NO in cancer cell, also could be originated from a wide range of sources. One of possibilities,  
304 we intended to investigate was iNOS. This specific protein from the family of nitric oxide  
305 synthases (NOS) could be responsible for statistically significant formation of nitrites and also  
306 according to Xu et. al for the cytotoxicity in tumor cells (Xu et al. 2002). Our results follow  
307 strong relationship between iNOS synthesis induction/suppression and nitrite level in tested  
308 cells. According to showed analogy, we could suggest that these substances significantly  
309 influenced iNOS protein expression, and thus the increasing or decreasing of level of nitrites.  
310 On the other hand, NO (and thus nitrite) level greatly depends on  $O_2^{\bullet-}$  production. This could  
311 be explained by fact that  $O_2^{\bullet-}$  possess great affinity towards NO, forming peroxyntirites

312 (Ferrer-Sueta and Radi 2009). Also, the fact that investigated substances contain a lot of  
313 nitrogen atoms in their imidazole structure could offer a possibility that substances were  
314 metabolized by cells. As the human cell is very complex system, every source that impacts at  
315 least one of the intracellular parameters greatly influences the redox equilibria (Cordero and  
316 deMiguel 2012). Greater production of free radical species influenced greater production of  
317 glutathione, which is in agreement with the nature of GSH and its cellular defense role against  
318 disruption of redox equilibria. **Hid-Se** does not possess such a denominated prooxidative and  
319 cytotoxic character as **(Hid-Se)<sub>2</sub>Pd** does, thus the changing in GSH level influenced by **Hid-**  
320 **Se** is not as significant as with **(Hid-Se)<sub>2</sub>Pd**.

321 In addition to examination of the influence of tested substances on the viability and redox  
322 status of cells, testing the impact on migratory potential is also very important. A certain  
323 substance does not need to possess significant cytotoxic character, but if given substance  
324 significantly reduces the migratory potential then it may be considered as potentially  
325 interesting for further studies towards the synthesis the appropriate drug. Our results showed  
326 that **Hid-Se** and **(Hid-Se)<sub>2</sub>Pd** induced increasing in migration potential of HCT-116 cells. On  
327 the other hand, these substances greatly decreased the migration potential of metastatic MDA-  
328 MB-231 breast cancer cells. The occurrence that MDA-MB-231 cells migrate more intensive  
329 than HCT-116 cells is previously described (Wu et al. 2011), and the result indicating that  
330 MDA-MB-231 cells are migratory limited is promising. Also, it has been found that  
331 significant increase in ROS/RNS has direct impact on antimigratory effect on cancer cells  
332 (Urbich et al. 2002). It was found that moderate production of ROS/RNS promotes cancer  
333 migration (Nishikawa 2008), while overproduction possesses an opposite effect, i.e.  
334 significantly reduces cancer cell migration (Fini et al. 2008). Considering the cell viability  
335 MTT assay, one could observe that HCT-116 cells are more sensitive than MDA-MB-231  
336 cells, which is in agreement with our earlier findings (Kosaric et al. 2014; Petrovic et al. 2014;

337 Petrovic et al. 2015). But, redox status parameters measured and presented in this paper  
338 suggest that MDA-MB-231 cells are more susceptible to ROS/RNS increasing in treatment  
339 with investigated substances. We observed moderate ROS/RNS increasing in HCT-116 cells,  
340 and thus increasing in cell migration. In MDA-MB-231 cells lower concentration (10  $\mu$ M)  
341 predominantly increased ROS/RNS in less significant extent when it is compared to higher  
342 concentration (100  $\mu$ M). According to Nishikawa (Nishikawa 2008) and Fini (Fini et al.  
343 2008) we consider that moderate increasing of ROS/RNS induced with 10  $\mu$ M of substances  
344 promotes cell migration, while higher 100  $\mu$ M treatment induced great oxidative stress and  
345 thus decreasing in cell migration potential.

346 Evidence that these substances possess cytotoxic and antimigratory character (especially on  
347 metastatic MDA-MB-231 cells) appears to be promising in terms of considering the further  
348 serious examination *in vitro* and/or *in vivo*. The exact mechanism of this action still remains  
349 unclear and many studies is needed to be done to reveal influence of superoxide anion radical  
350 and other radical species on migration processes in origin-different cancer cells.

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354

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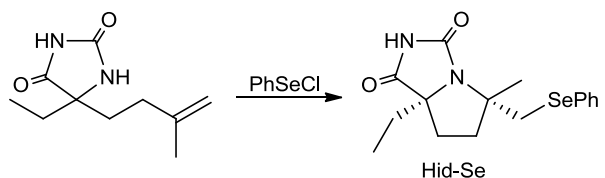
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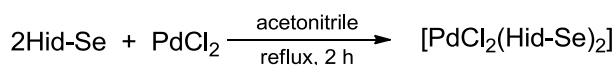
493

494 **Figure legends**

495 Scheme 1. Synthesis of Hid-Se and (Hid-Se)<sub>2</sub>Pd



496



497

498

499 Figure 1. The dose response curve of effect of **Hid-Se** and **(Hid-Se)<sub>2</sub>Pd** on HCT-116 and  
500 MDA-MB-231 cell lines after 24 and 72 h of exposure. All values are mean ± standard error.

501

502 Figure 2. Detection of iNOS protein expression in HCT-116 and MDA-MB-231 cells by  
503 immunofluorescence staining.

504 Cells are treated with 50 μM substances and iNOS protein expression was examined 24 h  
505 from treatment. Cell nuclei are DAPI stained to blue, while iNOS proteins are green (NIS-  
506 Elements Advanced Research Captures merged). All sections were examined by Nikon  
507 inverted fluorescent microscope (Ti-eclipse) at 600x magnification.

508

509 Figure 3. The influence of **Hid-Se** and **(Hid-Se)<sub>2</sub>Pd** on migratory potential of HCT-116 and  
510 MDA-MB-231 cells, after 24 h of treatment related to the number of viable cells.

511 The cells were treated with substances with concentrations of 10 and 100  $\mu\text{M}$ . All values are  
512 mean  $\pm$  SE, n=2, \*p < 0.05 as compared to control.



Fig. 2 [Download full resolution image](#)

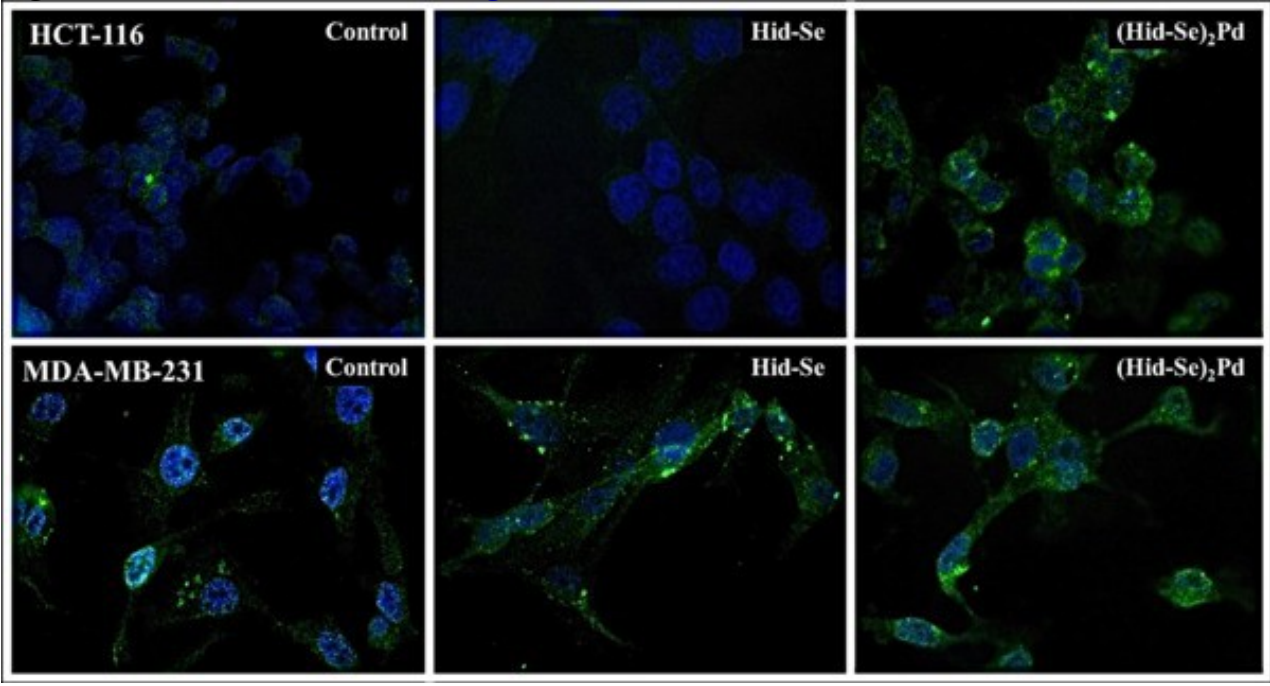


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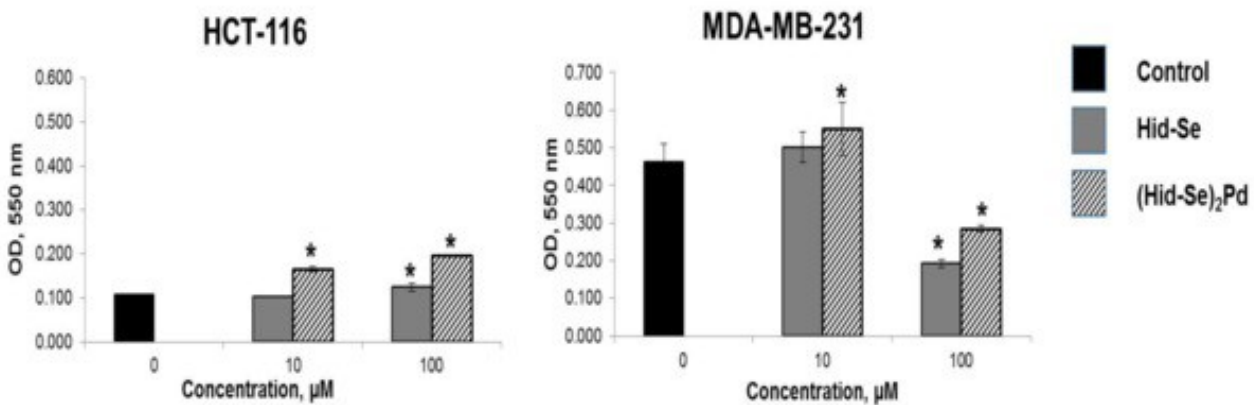


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