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# **RESEARCH ARTICLE**

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# Genotoxicity of selected pharmaceuticals, their binary mixtures, and varying environmental conditions – study with human adenocarcinoma cancer HT29 cell line

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#### ABSTRACT

Pharmaceutical residues are present in the environment in mixtures and their adverse effects may also result from interactions that occur between compounds. Studies presented in this work focus on genotoxicity of pharmaceuticals from different therapeutic groups in mixtures and in individual solutions impacted with different environmental conditions assessed using comet assay (alkaline approach). Binary mixtures of pharmaceuticals (in different concentration ratios) and in individual solutions impacted with pH change (range from 5.5 to 8.5) or addition of inorganic ions, were incubated with HT29 cells and after 24 h time period cells were tested for the presence of DNA damage. To estimate whether mixtures act more (synergistic) or less (antagonistic) efficiently Concentrations Addition (CA) and Independent Action (IA) approaches were applied followed by a calculation of the Model Deviation Ratio (MDR) to determine deviation from the predicted values. Addition of inorganic ions mainly reduced their genotoxicity. Diclofenac s. was the most susceptible to potassium, fluoride, and bromide ions. Change of the pH of pharmaceutical solutions had significant impact on genotoxicity of diclofenac s. and fluoxetine h. Among mixtures, more commonly observed interactions were synergistic ones, exactly twenty-five cases (ten pairs containing chloramphenicol or oxytetracycline h.) and ten cases of antagonism (four for pairs containing chloramphenicol or fluoxetine h.). The results obtained indicate that interactions between tested compounds occur frequently and can lead to DNA damage. This topic especially concerning in vitro tests using cells is still rare, however, it should not be neglected.

# **ARTICLE HISTORY**

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#### **KEYWORDS** Drugs residues; drugs mixtures; cytotoxicity; aenotoxicity: MTT: comet assav

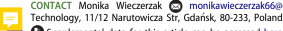
#### 1. Introduction

Environmental samples may contain variety of harmful substances just to mention heavy metals, plasticizers, polyaromatic hydrocarbons (PAHs), polychlorinated biphenyls (PCBs), nanomaterials, and pharmaceuticals residues (Huang et al. 2014, Ávila and García 2015, Bocchi et al. 2016). The latter is introduced into the environment due to the increasing activity of pharmaceutical industry, hospitals and households, also as a result of using veterinary medicines, agriculture, aguaculture, and animal husbandry (Gaw et al. 2014, Li 2014, Sui et al. 2015).

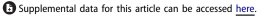
Pharmaceutical residues in the environment create a threat of long-term exposure, in particular, because drugs rarely undergo full elimination in classic water treatment processes and are difficult to degrade, what contributes to their accumulation in ecosystems (Eggen et al. 2010). Many conducted studies have confirmed that drug residues can cause adverse effects in single and multicellular organisms for example non-steroidal anti-inflammatory drugs (NSAIDs) are particularly toxic to aquatic organisms such as algae and fish (Cuthbert et al. 2016, Zanuri et al. 2017). Presence of hormone-like compounds in the environment may lead to

endocrine disorders and reproductive issues in organisms, including mammals, birds or fish (Li 2014).

Until recently most of toxicological studies have concentrated mainly on determining the harmfulness of individual compounds, however, in reality, living organisms generally are exposed to mixtures of chemical compounds originating from e.g., environmental pollution, the effects of which (including effects at the DNA levels) are difficult to predict (Parasuraman 2011). In the literature, there are few examples of toxicity of drug and other compounds mixtures studies and such as those described in this publication regarding research on human cells are even rarer. Just to mention research conducted by Chen et al. (2015) (performed with mixtures of often used herbicides and insecticides) showed that they are much more toxic towards soil organisms than solutions of single compounds (Chen et al. 2015); also  $17\alpha$ -ethinylestradiol commonly used in i.e., birth control pills in a mixture with and rogen  $17\beta$ -trenbolone (used in veterinary and illegally in doping) affects the process of differentiation of gonads in zebrafish and leads to sexual dysfunction in a significant way (Touber et al. 2007, Nasuhoglu et al. 2012, Orn et al. 2016). There are also few examples of studies,



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which show that drug mixtures present in the environment 117 can be harmful to human cells, e.g., Pomati et al. (2006) observed in their studies that the drugs mixture of atenolol, bezafibrate, carbamazepine, ciprofloxacin, cyclophosphamide, 120 furosemide hydrochlorothiazide, ibuprofen, lincomycin, ofloxacin, ranitidine salbutamol, and sulfamethoxazole (at the environmentally relevant exposure level) inhibited the growth of human embryonic cells (Pomati et al. 2006). 124

Determining the toxic impact of chemical mixtures in rela-125 tion to individual compounds is a difficult, complex and often 126 debatable subject, yet an accurate prediction of the overall 127 toxicity is very important for environmental risk assessment 128 129 and due to the fact that significant amounts of pharmaceuticals are now detected in drinking water, this knowledge also 130 contributes to the estimation of the impact on human health 131 132 (Caban et al. 2015).

In the literature phenomena of interactions between com-133 pounds in mixtures have been described long time ago; the 134 definitions of synergy (interaction of two or more substances 135 that mutually reinforce each other's actions) and antagonism 136 (reduction of activity as a result of interactions between 137 compounds) as well as models with which the type of these 138 interactions can be determined are described (Fent et al. 2006, 139 González-Pleiter et al. 2013, Wieczerzak et al. 2015, 2016b). 140

Concentration Addition (CA) and Independent Action (IA) 141 models are commonly used in ecotoxicological studies, to cal-142 culate the predicted toxicity values of mixture based on the 143 toxicity of individual pharmaceutical solutions. Both models 144 are based on different assumptions about active sites and 145 modes of actions (MOA), still, especially in ecotoxicological 146 studies, this type of knowledge is frequently incomplete 147 (Drescher and Boedeker 1995). CA models are more frequently 148 applied than IA although the last one is more accurate (if all 149 assumptions of the model are fulfilled, which sometimes 150 is problematic), still, in most of the cases, differences are 151 negligible (Belden et al. 2007). 152

Studies described in this paper aimed at estimation of 153 genotoxicity of selected pharmaceuticals (which are present 154 in environment at different concentration levels) towards 155 HT29 cell line studied in binary mixtures of selected pharma-156 ceuticals in individual solutions impacted with pH change or 157 addition of inorganic ions (at environmentally stated levels). 158 Information on physicochemical properties, medical applica-159 tions, levels of concentrations found in the environment and 160toxicity of pharmaceuticals selected for studies are listed in 161 Table 1. 162

In order to eliminate the risk of using cytotoxic concentra-163 tion of pharmaceutical solutions in the genotoxicity studies, 164 Thiazolyl Blue Tetrazolium Bromide (MTT) test was performed, 165 which assesses cell metabolic activity. Concentrations deter-166 mined in the MTT test were used in the next stage of geno-167 toxic studies, which have been carried out with alkaline 168 comet assay (to determine magnitude of DNA damage) 169 (Koss-Mikołajczyk et al. 2015). In order to determine the 170 nature of the phenomena observed the Model Deviation 171 Ratio (MDR) was used, which compare predicted values (cal-172 culated using CA and IA) and experimental results 173 et al. 2016a, 2016b, 2018b, Szczepańska 174 (Wieczerzak et al. 2018).

Application of such research methodology gives not only information on the cytotoxic concentrations of the studied drugs but also allows for the assessment of observable interactions occurring in the mixture of selected drugs and the influence of environmental conditions such as coexistence of inorganic ions and pH changes in pharmaceutical solutions to their genotoxicity towards cells. Studies presented in this publication are a continuation of research on the impact of pharmaceutical mixtures and environmental conditions on single and multicellular organisms and supplement the knowledge obtained in earlier studies on Vibrio fischeri, Sorghum bicolor and Saccharomyces cerevisiae (Wieczerzak et al. 2016a, 2016b, 2018a, 2018b).

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# 2. Experimental

#### 2.1. Chemicals, reagents, and apparatus

#### 2.1.1. Model compounds and ions

Model substances selected for the study: diclofenac sodium salt (diclofenac s.) (CAS no. 153907-79-6), chloramphenicol (CAS no. 56–75-7), oxytetracycline hydrochloride (oxytetracycline h.) (CAS no. 2058-46-0), fluoxetine hydrochloride (fluoxetine h.) (CAS no. 56296-78-7), estrone (CAS no. 53-16-7), ketoprofen (CAS no. 22071-15-4), progesterone (CAS no. 57-83-0), gemfibrozil (CAS no. 25812-30-0), and androstenedione (CAS no. 63–05-8) were purchased from Sigma Aldrich (Germany) and were of analytical purity grade (>99%). Diazepam (CAS no. 439–14-5, purity grade >99%) was purchased from LGC Standards (UK). Inorganic ions in the salt form (KCI (CAS no. 7447–40-7), NH<sub>4</sub>CI (CAS no. 12125–02-9), NaF (CAS no. 7681–49-4), NaBr (CAS no. 7647–15-6)) were purchased from Avantor Performance Materials S.A. (Poland) while NaCl (CAS no. 7647–14-5) from Sigma Aldrich (Germany). pH values were adjusted with concentrated solutions of NaOH (CAS no. 1310-73-2) or HCI (CAS no. 7647-01-0) purchased from Avantor Performance Materials S.A. (Poland). All purchased ions were analytical purity grade >99%.

#### 2.1.2. MTT test and comet assay reagents

MTT (CAS no. 298-93-1, biological grade), fetal bovine serum (FBS), penicillin, and streptomycin solution, McCoy's 5a (Modified) Medium, DMSO (CAS no. 67-68- 5, biological grade), H<sub>2</sub>O<sub>2</sub> (CAS no. 21–67-63), N<sub>2</sub>EDTA (CAS no. 6381–92-6, biological grade), trypsin-EDTA solution (sterile-filtered), NaCl (CAS no. 7647-14-5), NaOH (CAS no. 1310-73- 2), Trizma<sup>®</sup>-base (CAS no. 77–86-1, biological grade), Trizma<sup>®</sup> hydrochloride (CAS no.1185–53-1, biological grade), Triton<sup>™</sup> X-100 (CAS no. 92046–34-9, biological grade), low (LMP) and normal melting (NMP) points agarose, trypsin-EDTA and SYBR<sup>®</sup> Green I nucleic acid gel stain were purchased from Sigma-Aldrich (Germany). Sterile serological pipette (5, 10 and 25 ml), cell cultures bottles, UltraFine<sup>™</sup> tips, coverslips, microscope slides, 10-ml syringes, sterile centrifuge tubes, and filters were purchased from VWR (Poland). All reagents were of analytical grade purity or better, in the case of reagents for microbiological purposes. Water was purified with a Milli-Q<sup>®</sup> system from Millipore (Germany).

2017, Ren et al. 2017, Warkus and Marikawa 2018).	kus and marikawa	2010).			
Analyte/CAS no.	M <sub>w</sub> /logP	Medical application	Action at the cellular level	Concentrations detected in environment	Observed toxicity towards cells/ single or multicellular organisms
Diclofenac sodium/ 15307-76-6	318.13/0.6	Nonsteroidal anti-inflammatory drug.	A potent inhibitor of COX-2, which reduces the synthe- sis of prostaglandins, prostacyclin and thromboxane- compounds associated with inflammation.	2900 and 2100 ng/L lake and river water <sup>a</sup> 2100 ng/L effluent wastewater <sup>a</sup>	Teratogenic to <i>Xenopus</i> embryos <sup>a</sup> . Inhibition of oxidative phosphorylation (ATP synthesis) in rats <sup>a</sup> .
Ketoprofen/22071-15-4	254.28/3.12	Nonsteroidal anti-inflammatory drug.	Reversibly inhibits COX-1 and COX-2 enzymes, which reduces the production of prostaglandins.	120-341 ague water 128-184 ng/L influent 5-57 ng/g sediment and seware	Mitochondrial toxicity in yeast Saccharomy cescerevisiae Toxic for embryonic stages of zehrafish
Androstenedione/63- 05-8	286.41/2.75	Male and female sex hormone precursor: testosterone, estrone and estradiol.	Acts as a weak androgen receptor agonist.	5 ng/L surface water 0.1 ng/g sediment	Carcinogenic in B6C3F1 mice.
Progesterone/57-83-0	314.46/3.87	Female steroid hormone, used to complement the decrease of naturally occurring progesterone to prevent miscarriage, it prevents menstrual cycle disorders.	Genomic pathway: Regulates expression of specific genes by binding to specific nuclear receptors (PRA and PRB) upon entry into the cell. Non-genomic pathway: Induces rapid growth of intracellular $Ca^{2+}$ concentra- tions, and reculates the flow of Na <sup>+</sup> and Cl <sup>-</sup> ions.	2.4 ng/g sediment 26 ng/L surface water	Apoptosis of insulin-secreting cells.
Estrone/53-16-7	270.37/3.13	Steroid estrogen with estradiol-like effects.	complex ns. It has one indu I COX ac	<1–5 ng/L surface water 65 ng/L river water 28 ng/L surface water	No data.
Chloramphenicol/56- 75-7	323.13/1.14	Antibacterial and bacteriostatic agent.	Prevents chain lengthening by inhibiting peptidyltrans- ferase activity of bacterial ribosomes	5800–47400 ng/L sediment and sewage 3000 ng/L wastewater	Cytotoxic to myeloma cells.
Oxytetracycline h./79- 57-2	460.43/-0.9	Natural antibiotic with bacteriostatic and antiallergic effect.	Inhibits protein synthesis by blocking the aminoacyl- tRNA binding at site A on the bacterial ribosome. This prevents the introduction of new amino acids into the growing peptide chain. Mammalian cells are less susceptible to tetracycline, despite the fact that tetracycline binds to prokaryotic and eukaryotic ribo- somal subunits (30S and 40S sites respectively).	200–5700 ng/L sediment and sewage <sup>c</sup> 564.30 ng/L effluent and 337.81 ng/L influent wastewater	Inhibition of expression of phase I and II detoxification genes in swimming crab ( <i>Portunus</i> <i>trituberculatus</i> ) larvae <sup>b</sup> . Acute toxicity to <u>Pseudokirchneriella</u> subcapitat <sup>b</sup> .
Diazepam/439-14-5	284.74/2.82	Psychotropic drug, anticonvulsant, sedative and anxiolytic.	Benzodiazepines are positive allosteric modulators of type A GABA receptors. GABA receptors are selective ion channels for Cl <sup>-</sup> ions. Binding of benzodiazepines to the GABA receptor complex increases the chloride ions' conductivity through the membrane of neur- onal cells. Diazepan seems to work in the limbic, hill and hvoothalamic areas.	335 ng/L surface water LOD-0.21 ng/L surface water	Inhibits proliferation and activation of T lymphocytes isolated from rat main lymphoid organ. Oxidative damage in the liver and gill tissues catfish <i>Clarias gariepinus</i> .
Fluoxetine h./56296- 78-7	345.79/1.8	Used mainly in the treatment of depressive and obsessive- compulsive disorders.	Fluoxetine is a selective serotonin reuptake inhibitor (SSRI).	1.7 to 1244 ng/L effluent waste water <sup>b</sup> LOD-2.7 ng/L surface water <sup>b</sup>	Induce an early necrosis in human peripheral blood lymphocytes and Jurkat cells <sup>c</sup> . Inhibition of canonical Wnt sig- naling and reduction of cellular proliferation <sup>c</sup> .
Gemfibrozil/25812-30-0	250.33/4.77	It normalizes blood fats, lowers LDL cholesterol and triglycerides, improves HDL cholesterol.	An <i>α</i> -receptor (PPAR) activator, a nuclear receptor that participates in the metabolism of carbohydrates and fats, as well as the differentiation of adipose tissue.	6 ng/g sediment and sewage >1000 ng/L	Genotoxic to gilthead seabream (Sparus aurata).

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Table 2. Main concentration values of pharmaceuticals solution used in MTT test and the concentrations selected for the comet test.

Analyte	Stock concentration of solution [mol/L]	Concentration (C2) of pharmaceutical solutions selected for further studies with comet assay [mol/L]					
Diclofenac s.	0.0064	0.00066					
Oxytetracycline h.	0.0047	0.00021					
Chloramphenicol	0.00058	0.00006					
Fluoxetine h.	0.0008	0.00045					
Analyte	Stock concentration of solution [mol/L]	Maximal tested concentration of pharmaceutical					
		solutions with effect $\pm$ SD					
Ketoprofen	0.00295	Concentration could not be calculated					
		(max. tested conc. 0.00074 mol/L					
		tested with effect 99.0 $\pm$ 1.3%)					
Gemfibrozil	0.0006	Concentration could not be calculated					
		(max. tested conc. 0.00015 mol/L					
		tested with effect $112.0 \pm 1.3\%$ )					
Estrone	0.00094	Concentration could not be calculated					
		(max. tested conc. 0.00024 mol/L					
		tested with effect $78.0 \pm 2.6\%$ )					
Diazepam	0.00088	Concentration could not be calculated					
		(max. tested conc. 0.00022 mol/L					
		tested with effect $94.4 \pm 9.0\%$ )					
Androstenedione	0.0008	Concentration could not be calculated					
		(max. tested conc. 0.0002 mol/L					
	0.0000	tested with effect $123.5 \pm 1.0\%$ )					
Progesterone	0.0008	Concentration could not be calculated					
		(max. tested conc. 0.0002 mol/L					
		tested with effect $111.6 \pm 1.0\%$ )					

# 2.1.3. Cell lines

HT29 (human colon adenocarcinoma, obtained from American Type Culture Collection, Manassas, USA) were grown in a monolayer culture at 37 °C in a humidified atmosphere of 5% CO<sub>2</sub> in the Smart cell incubator (Heal Force, China) in McCoy's 5a (modified) medium, supplemented with 10% FBS and antibiotic (1% penicillin-streptomycin) in a culture flask. The medium was changed twice a week. Single cell suspensions were prepared with a trypsin-EDTA solution (diluted ten times) and finally re-suspended in McCoy's 5a (modified) medium, supplemented with serum and antibiotics (all activities took place in aseptic environment). Cellular studies were conducted due to courtesy of Professor Agnieszka Bartoszek from the Department of Chemistry, Technology, and Biotechnology of Food at Gdańsk University of Technology.

# 2.2. Methods

#### 2.2.1. MTT

MTT test was performed for individual solution of pharmaceuticals mainly to avoid using concentration of pharmaceuticals with could prove to be cytotoxic during the comet assay. The cells were seeded in 96-well tissue culture plates (about  $20 \times 10^3$  cells/well in 150 µL) and allowed to settle for 24 h at 37 °C after this period cells were treated for 24 h with 50 µL of different concentrations of pharmaceutical solution in distilled waters. Due to the fact that some of the studied compounds were found to react with PBS, this buffer either in the MTT test or in the comet assay was not used, instead purified water was used as control or in dilutions. After 24 h of the incubation time, 50  $\mu$ L of MTT solution (4 g/L) was added to each well in the plate and cells were left for another 3 h at 37 °C (up to this point all operations were performed under aseptic conditions). Finally, the medium was carefully removed from wells, and formazan crystals formed

were dissolved in  $50\,\mu\text{L}$  of DMSO added to each well. The absorption of the resultant solutions was determined at 540 nm wavelength with TECAN Infinite M200 plate reader (Tecan Group Ltd., Switzerland) (Koss-Mikołajczyk et al. 2015). The cytotoxicity was expressed as growth inhibition of cells exposed on tested plant samples compared to control of non-treated cells whose growth was regarded as 100%. Each dilution of pharmaceuticals was studied in triplicates on two plates in separate repetition performed at weekly intervals. Concentrations of pharmaceutical solutions studied in the MTT test ranged from 1 to 25% (v/v) of stock concentration listed in Table 2. Drugs concentration values to perform comet assay studies were selected as 30% cytotoxicity decrement in relation to the control if the cytotoxicity of studied solution was significant (refer to Table 2). The data were analyzed using Q-Dixon test to eliminate major errors followed by t-Student comparison test. The statistical significance was set to \*p < 0.05, \*\*p < 0.01, \*\*p < 0.001 and are presented in the supplementary materials (Supplementary Figures 1-2).

# 2.2.2. Comet assay

Epithelial colon cancer cells were grown in a monolayer culture at 37°C in a humidified atmosphere of 5% CO<sub>2</sub> in McCoy's 5a (modified) medium, supplemented with 10% fetal bovine serum and antibiotic (1% penicillin-streptomycin) in a culture flask. The medium was changed twice a week. Single cell suspensions were prepared with a trypsin-EDTA solution (diluted 10 times) and finally re-suspended in McCoy's 5a (modified) medium, supplemented with serum and antibiotics.

The cells from the culture flask were seeded in 24-well tissue culture plates (about  $200 \times 10^3$  cells/well in 1800 µL) and allowed to settle for 24 h at 37 °C, after this period cells were treated for 24 h with 200 µL of different concentrations of pharmaceutical solutions diluted in deionized water (different concentrations of pharmaceuticals solution were used at 429

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different stages of the study, which will be reported when
discussing the individual results) each sample was performed
in three repetitions.

After 24 h incubation period medium from cell surface was 468 carefully removed and cells were trypsinized (200 µL of trypsin 469 was added to each well) in order to detach the cells from well 470 wall. Afterward to the suspension of the cells in each well 471 1800 µL of fresh McCoy's medium was added, then the cell 472 suspension was pipetted and 1 mL was transferred to 1.5 mL 473 volume vortex tubes (up to this point of procedure all opera-474 tions were performed under aseptic conditions). Vortex tubes 475 were centrifuged (1000 rpm, 5 min, 4°C) the supernatant was 476 removed, and then 1 mL of PBS was added to the tubes and 477 centrifuged again. The supernatant was removed again in such 478 a way to leave about 100 µL of buffer. Cell suspensions were 479 pipetted and 50  $\mu$ L of each suspension was mixed with 150  $\mu$ L 480 of 0.5% low melting point (LMP) agarose in PBS maintained at 481 37 °C. 40 µL agarose with cells was transferred two times on 482 glass microscope slides pre-coated with 1% normal melting 483 point (NMP) agarose. Gels were covered with a coverslip and 484 kept on ice for 5 min. Two gels per glass slide were prepared. 485 For the purpose of this study, protocol for the exposure of 486 slides, which was previously described by (Lah et al. 2005) and 487 later developed by (Durgo et al. 2009, Mihaljevic et al. 2011) 488 was adopted. Following the treatment the cells subjected to 489 lysis solution (2.5 mol/L NaCl, 0.1 mol/L Na<sub>2</sub>EDTA, 10 mmol/L 490 Trizma<sup>®</sup>-base, pH 10 and 10% Triton X-100, min. 1 h of incuba-491 tion, 4°C, in the darkness). The comet assay was performed 492 under alkaline conditions (pH >13.3) (Fairbairn *et al.* 1995), as 493 described in (Kazimirova et al. 2012) with some modifications. 494 After lysis, slides were placed in a horizontal electrophoresis 495 tank and DNA was allowed to unwind for 20 min in electro-496 phoretic buffer (0.3 mol/L NaOH, 1 mmol/L Na<sub>2</sub>EDTA, pH >13.3, 497 4°C) before electrophoresis was performed for 20 min at 26 V 498 (300 mA). After neutralization by washing gels three times, 499 each for 5 min with 0.4 mol/L Trizma<sup>®</sup>-base (pH 7.5) at 4°C, 500 and then with water and dried at room temperature. After 501 staining the slides with SYBR GREEN I solution, the comets 502 were detected and quantified as described below. 503

For quantitative analysis of nuclear DNA damage, the slides 504 were viewed at 50× magnification with an epifluorescence 505 microscope Zeiss Imager Z2. Microscopic images of comets were 506 taken with digital camera (CoolCube1) connected to computer, 507 and the comets were scored using Metafer 4 Software. The data 508 were analyzed by Q-Dixon test to eliminate major errors and 509 were expressed as the mean values with the standard deviations 510 (means  $\pm$  SD) of the three independent experiments and were 511 used as numeric values for model studies (Kudłak et al. 2016). In 512 order to systematize the research sequence, the test procedure 513 is schematically shown in Figure 1. 514

The concentration levels of the model compounds solu-515 tions (diclofenac s., oxytetracycline h., chloramphenicol, and 516 fluoxetine h., refer to Figure 2) calculated based on results of 517 the MTT test, are summarized in Table 2. Concentrations of 518 inorganic ions selected for genotoxicity studies correspond to 519 the environmental concentrations and were selected based 520 on previous research on surface water samples collected 521 from waters bodies receiving waters from sewage treatment 522 plants in Poland i.e., Na<sup>+</sup> C = 1.09 mmol/L, K<sup>+</sup> C = 0.13 mmol/ L,  $NH_4^+$  C = 0.042 mmol/L, Cl<sup>-</sup> C = 1.41 mmol/L, F<sup>-</sup> C = 0.022 mmol/L, Br<sup>-</sup> C = 0.013 mmol/L (Kudłak *et al.* 2016).

#### 2.3. Calculation of MDR values

In order to determine whether the genotoxicity towards HT29 (if so: how and to what extent) depends on the concentration of compounds in the binary mixture, each of the compounds was tested at three concentrations levels. Concentrations (C2) determined during the MTT test (exactly 30% of the toxic concentration) was reduced by 50% (C1) and increased by 50% (C3). The binary mixtures were constructed in such a way that C1 concentration of first substance as mix with C1, C2 or C3 concentration of first substance with C1, C2 or C3 concentration of first substance with C1, C2 or C3 concentration of second substance with C1, C2 or C3 concentration of second substance setc. All concentrations of studied compounds are summarized in Table 2.

In this study, the combined toxicological effect of a mixture of pharmaceuticals and the impact of environmental conditions toward HT29 cells was assessed both with the CA and IA models using Equations 1 and 2, respectively (Kienzler *et al.* 2016):

$$EC_{X_{Mix}} = \left(\sum_{i=1}^{n} \frac{P_i}{EC_{x_i}}\right)^{-1}$$
(1)

where:

- *ECx<sub>mix</sub>* is the  $x_{mix}$  effect cause by total concentration of the mixture of studied chemicals (components) (Expected value).
- *p<sub>i</sub>* indicates the fraction of component *i* in the mixture, calculated on the basis of the concentration of *i* component in the mixture;
- n indicates the number of components in the mixture;
- *ECx<sub>i</sub>* indicates the *x<sub>i</sub>* effect caused by component *i* at given studied concentration in the mixture,

$$E(C_{mix}) = 1 - \prod_{i=1}^{n} (1 - E(c_1))$$
(2)

where:

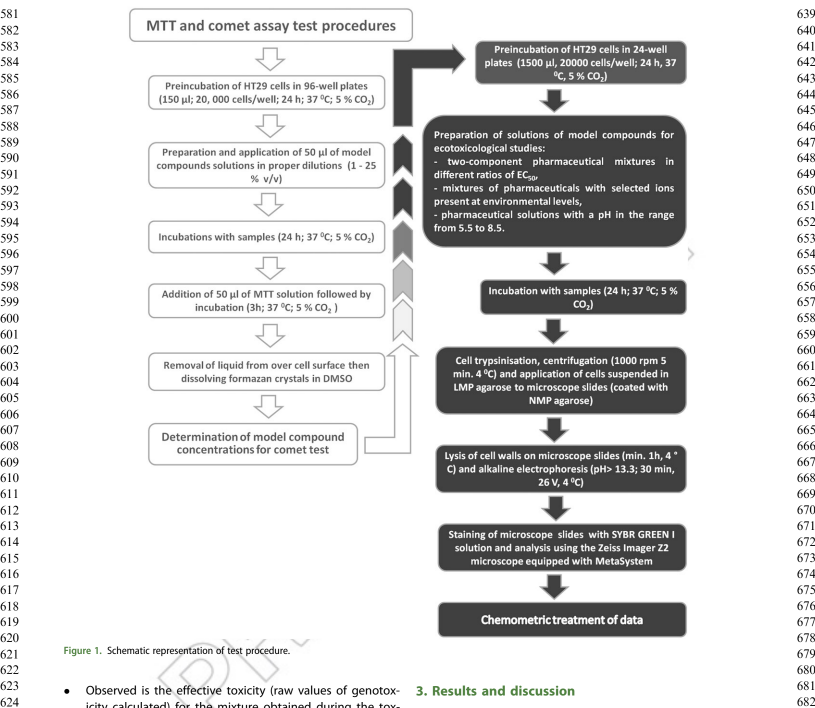
- *EC<sub>mix</sub> is* overall effect expressed as a fraction of maximal possible effect of a mixture of *i* chemical, (Expected value).
- c<sub>i</sub> indicates the concentration of component i in the mixture;
- *n* indicates the number of components in the mixture;
  - $E(c_i)$  indicates the effect of component *i*, applied separately.

In order to verify the difference between the predicted and observed effect, the Model Deviation Ratio (MDR) approach was applied, defined as shown in the Equation (3) (Kudłak *et al.* 2016, Wieczerzak *et al.* 2016b):

$$MDR = \frac{exp \ ected}{Observed} \tag{3}$$

where:

• Expected is the effective toxicity (raw values of genotoxicity calculated according to Equation (1) or (2) of the mixture predicted by the CA or IA model,



 Observed is the ellective toxicity (raw values of genotoxicity calculated) for the mixture obtained during the toxicity studies.

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In the case of studies on impact of pH change pharmaceuticals solutions, use of CA and IA modeling methods was impossible due to the fact that these solutions cannot be treated as a mixture of chemicals. Therefore, in this case, a simple ratio was used (described by the Equation (4).) to assess impact of pH change:

Effect ratio = (4) Effect observed for pharmaceuticals solutions with corrected pH Effect observed for pharamceuticals solutions without pH adjustment

Solutions of pharmaceuticals with set pH for which the value of the ratio was lower than one were less toxic and those with values above one were more toxic toward HT29 cells.

# **3.1.** Impact of pharmaceuticals solutions to cytotoxicity towards HT29 cell line

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Diclofenac s., oxytetracycline h., chloramphenicol, and fluoxetine h. have been shown to be significantly cytotoxic to HT29 cells. The most cytotoxic turned out to be chloramphenicol, which decreased the mitochondrial activity of cells by half at 0.088 mmol/L concentration (concentrations selected for further genotoxicity studies are summarized in Table 2). Other studies carried out by Parolini *et al.* (2011) at different cell from the zebra mussel *Dreissena polymorpha* (haemocytes, gills and digestive gland cells) showed that among the tested drugs (diclofenac, atenolol, carbamazepine) – diclofenac turned out to be very cytotoxic in the Trypan Blue Exclusion test and only slightly cytotoxic in the MTT test

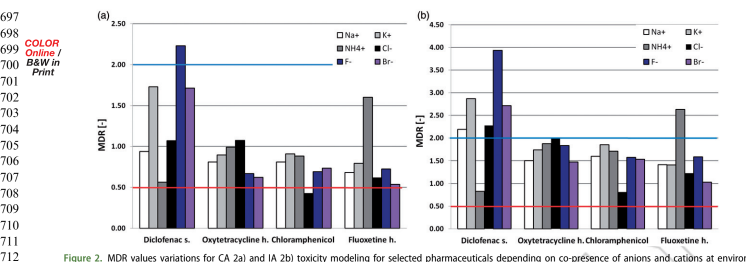


Figure 2. MDR values variations for CA 2a) and IA 2b) toxicity modeling for selected pharmaceuticals depending on co-presence of anions and cations at environmentally relevant levels.

715 (in the concentration range up to 10 mg/L), the second most toxic turned out to be gemfibrozil, which shows to be cyto-716 toxic to HT29 cells (Parolini et al. 2011). In case of drugs: 717 718 ketoprofen, gemfibrozil, estrone, diazepam, androstenedione, and progesterone no significant cytotoxic or genotoxic effect 719 was observed at concentrations levels that exceed solubility 720 equilibrium, therefore, no further studies with these substan-721 ces were conducted. Maximum reduction of mitochondrial 722 activity for described above substances was observed for 723 estrone it was 78% at the concentration 0.24 mmol/L in rela-724 725 tion to the control if the cytotoxicity of studied solution was 726 significant (maximum effects obtained in the MTT test for the highest concentrations of discussed substances studied are 727 728 summarized in Table 2). Raw results obtained for MTT 729 cytotoxicity of solutions of pharmaceuticals are presented in 730 supplementary materials: supplementary Figures 1-2.

### 3.2. Genotoxicity studies

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In this article, mixtures of pharmaceuticals at various concen-734 tration levels were tested on HT29 cells for the first time and 735 CA and IA modeling was used to evaluate the results obtained 736 and compare them with controls with individual compounds 737 solutions. The application of the CA and IA models alone does 738 not allow for the determination of possible interactions 739 between chemicals (in this case in pharmaceuticals) in mixture 740 and define their character. The deviation from the predicted 741 effect of CA and IA mode may be evidence for occurrence of 742 synergistic or antagonistic action. The mixtures with MDR >2.0 743 exhibit a high probability of antagonism while those with val-744 ues below 0.5 show synergistic character (Faust et al. 2000, 745 Backhaus and Faust 2012, Kienzler et al. 2016, Wieczerzak 746 et al. 2016b). In current research it was arbitrarily assumed 747 that MDR falling within 0.50-0.71 and 1.40-2.00 justify the 748 conclusion on, respectively, possible under- and overesti-749 mation of presented models. 750

# 7523.2.1. Impact of inorganic ions on genotoxicity towards753HT29 cell line of selected compounds

In case of impact of ion addition on toxicity of diclofenac s. one can notice a reduction in genotoxicity practically for all ions (see Figure 2 to compare), the highest MDR values were observed for bromide ions (for the CA model they were above 2.0 and for the IA model above 3.5 which indicates strong antagonism). The exceptions were ammonium ions, which clearly affected toxicity of diclofenac s., MDR values for this mixture were bit above 0.5. The MDR parameter for oxytetracycline h. varied depending on the model used, for the CA model only two values were in the overestimated range (fluorides and bromides), and for the IA model all values and MDR were in the underestimated range (chloride ions were closest to 2.0).

In the case of chloramphenicol, only chloride ions significantly influenced its genotoxicity, resulting in an increase in the CA model, the MDR value decreased below 0.5 which means synergistic activity was observed (see Table 2). Surprising behavior is observable in case of studies on fluoxetine h.; ammonium ions show antagonistic impact being, most probably, result of shifting ion equilibrium. It is proven by fact that all anions seem to show synergistic impact on toxicity of fluoxetine h. (with CA model studies).

In earlier conducted by the authors test on Vibrio fischeri bacteria, the opposite values of the MDR parameter were observed, the addition of ions rather contributed to the increase of acute toxicity of the studied drugs, it may result from various cell structure, and other defense and repair mechanisms, however without additional testing it is only conjecture (Wieczerzak et al. 2018a). In the case of previously tested yeast Saccharomyces cerevisiae (also by the authors), ions had no significant effect on the endocrine potential tested in the XenoScreen YES/YAS test, the only compound susceptible to both anions and cations was fluoxetine h., ions present in its solution contributed to the increase of its agonistic properties against estrogen receptors (Wieczerzak et al. 2018a). Raw results obtained for genotoxicity of solutions of pharmaceuticals impacted with ions addition are presented (as % of DNA in the comet tail ± SD) in supplementary materials: Supplementary Table 1.

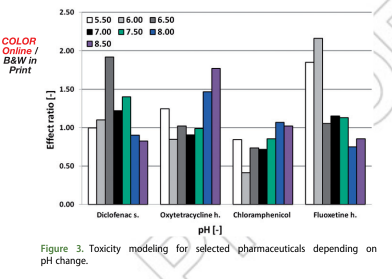
# 3.2.2. Impact of pH change on genotoxicity of solution of selected pharmaceuticals towards HT29 cell line

Synergic impact of pH change is noticeable in case of diclofenac, significant (>90%) increase of this drug's toxicity

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took place at pH of 6.50 (refer to Figure 3). In case of oxytetracycline, the synergy was observed in low and high pH values, namely at 5.50, 8.00, and 8.50 (increment by over 24, 46 and 77%, respectively). Chloramphenicol is not synergistically impacted by pH variations, only antagonistic influence was detected at 6.00–7.00 pH range. Low pH values act in a synergic way on genotoxicity fluoxetine (84 and 116% increase of DNA damage at 5.50 and 6.00, respectively), some tendency of antagonistic behavior is observed at pH >8.00. Raw results of genotoxicity of pharmaceuticals solution with ions and pharmaceuticals solution with ph adjustment (presented as % of DNA in the comet tail  $\pm$  SD) are summarized Supplementary Table 1.

Studies on the pH change of pharmaceutical solutions regarding both Vibrio fischeri and Saccharomyces cerevisiae did not show any effect on the test organisms, only oxytetracycline h. has a slight antagonistic effect on estrogen receptors under the influence of pH change to 8.5. HT29 cells seem to be more sensitive to pharmaceutical solutions that have changed pH (Wieczerzak et al. 2018a). Raw results obtained for genotoxicity of solutions of pharmaceuticals impacted with pH change are presented (as % of DNA in the comet tail ± SD) in supplementary materials: Supplementary Table 1.



# 3.2.3. Modeling of results of selected pharmaceuticals binary mixtures genotoxicity studies

Diclofenac s. impacts oxytetracycline h. in synergic way at medium concentration level of diclofenac h., some overestimation/synergy cases are detected at lowest and highest concentration levels of diclofenac (refer to Table 3 for details). The situation is different when diclofenac is present in mixture with chloramphenicol - more cases of underestimation/synergy are detected at medium, highest and lowest concentrations when CA model was applied. Similar situation is confirmed in case of diclofenac-fluoxetine although the lowest concentration of fluoxetine has antagonistic impact on genotoxicity of medium and highest concentration levels of diclofenac. Oxytetracycline h. is influenced by diclofenac at medium level concentration in a synergic manner what is confirmed with both models used. Lowest and highest levels of diclofenac affect oxytetracycline h. genotoxicity in antagonistic way. Interestingly mixture of chloramphenicol with oxytetracycline shows synergy in most cases (despite situation when oxytetracycline is present in mixture at highest concentration level with low and medium content of chloramphenicol). Situation of fluoxetine-oxytetracycline mixture is complex in its nature; one can say that antibiotic affect activity of fluoxetine at lowest concentration level in antagonistic manner. Other cases show most often synergic behavior. Chloramphenicol in mixture with oxytetracycline shows synergic impact in most cases with both models used. In case of mixture with fluoxetine synergy is observed only at lowest concentration level of fluoxetine and highest content of chloramphenicol - other cases seem to exhibit either antagonistic or overestimated behavior. Diclofenac at medium concentration level impact chloramphenicol from antagonistic to synergy with increasing content of chloramphenicol. Low content of diclofenac shows fully antagonistic impact on chloramphenicol at its highest concentration level. Fluoxetine is impacted by low concentration levels of diclofenac in synergic manner, only in case of lowest fluoxetine content with medium and high levels of diclofenac some overestimation and antagonism takes place. Medium and highest levels of oxytetracycline affect fluoxetine (at medium and highest concentration levels) in a synergic way.

Interactions observed between substances studied were mainly of synergistic character. Most cases of synergistic

Table 3. MDR values variations for CA and IA for mixtures of selected drugs at three concentration levels for each tested compounds (color coding: red: synergism; blue: antagonism; green: overestimation; yellow: underestimation).

				Diclof	enac s.														
		CA	IA	CA	IA	CA	IA												
	C [mmol/L]	C	1	C	2	C	3												
Oxytetracycline h.	C1	0.96	1.79	0.35	0.60	0.72	1.47	Oxyte	tracycli	ne h.									
	C2	0.79	1.51	0.54	0.97	0.98	2.06	ĊÁ	IÁ	CA	IA	CA	IA						
	C3	1.01	1.97	0.28	0.52	0.80	1.53	C1		C2		C3							
Chloramphenicol	C1	0.65	1.37	1.29	2.28	1.11	1.34	0.09	0.26	0.24	0.55	1.07	1.92	Chlora	ampher	nicol			
	C2	0.55	1.87	0.42	1.17	0.73	1.11	0.10	0.27	0.14	0.30	0.81	1.43	CA	IA	CA	IA	CA	IA
	C3	1.77	6.31	0.30	0.87	0.68	1.19	0.31	1.06	0.27	0.76	0.27	0.58	C1		C2		C3	
Fluoxetine h.	C1	0.19	0.36	1.68	3.02	1.67	3.21	0.65	1.39	1.34	2.74	0.85	1.62	0.91	1.94	0.67	1.71	0.33	0.82
	C2	0.43	0.81	0.35	<mark>0.66</mark>	0.48	1.01	0.85	1.52	0.91	1.71	0.20	0.38	1.49	2.46	1.09	2.07	0.65	1.24
	C3	0.42	0.88	0.44	1.20	0.95	1.64	0.74	1.45	0.26	0.52	0.60	1.13	2.10	3.20	0.64	1.12	0.72	1.28

Diclofenac s. C1: 0.330 mmol/L; C2: 0.660 mmol/L; C3: 0.990 mmol/L; Oxytetracycline h. C1: 0.105 mmol/L; C2: 0.210 mmol/L; C3: 0.315 mmol/L; Chloramphenicol C1: 0.029 mmol/L; C2: 0.059 mmol/L; C3: 0.0.088 mmol/L; Fluoxetine h. C1: 0.225 mmol/L; C2: 0.450 mmol/L; C3: 0.675 mmol/L

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		Synergism	Under-estimation	Over-estimation	Antagonism	80%	90%	95%	9
Ph						1.339	1.794	1.893	2
lons	CA	1	7	3	1	1.071	1.68	1.727	2
	IA	0	0	14	6	2.226	2.688	2.848	3
Mixtures CA	CA	20	9	4	1	0.994	1.325	1.674	1
	IA	5	6	15	9	1.925	2.41	3.084	4

interactions (ten) were observed for the mixture chloramphenicol with oxytetracycline h. and confirmed applicability of CA and IA as prediction models. The number of interaction cases increased with the decreasing amount of oxytetracycline h. in the mixture, these mixtures also had the lowest values of MDR. In the mixture of diclofenac s. and fluoxetine h., seven cases of synergism were observed with the lowest MDR value (0.2 - CA and 0.4 - IA) for the mixture of lowest concentrations of these compounds. In all cases of mixtures tested, some synergistic effects have been observed.

There were several (four) examples of antagonistic impact, in most cases for fluoxetine h. and chloramphenicol mixture. In all cases, there were ten antagonistic interactions observed. Raw results of genotoxicity of mixtures tested (presented as % of DNA in the comet tail  $\pm$  SD) are summarized in Supplementary Table 2.

In previous studies of mixtures made on Vibrio fischeri, no pairs, compounds that behaved synergistically or antagonistically were detected all values of the MDR parameter were between 0.5 and 2.0 for both models. Similarly, in the case of earlier testing of yeast, there was no significant effect on one of the test substances on the other, using CA and IA models and MDR parameters. It seems that this time the HT29 cell is more sensitive, it may be related to the fact that the comet test is a very sensitive test, thanks to which even a small influence of tested compounds can be pinched (Olive and Banáth 2006, Wieczerzak et al. 2016b).

# 4. Conclusions

Mixtures of chemicals may be genotoxic despite the fact that the same compounds do not possess such properties when act separately, additionally, under environmental conditions, they may undergo various chemical or physical transformations, which can affect the toxicity of chemical compounds. Residues of drugs always occur in mixtures with other pollutants in the environment, impact of these environmental mixtures on human and animal organisms is a huge gap in knowledge as far as the environmental fate of these compounds is in stake. The lack of cheap and green technology that eliminates this type of chemical pollution from sewage causes the human and animal organisms to be constantly exposed to residues of pharmaceuticals. In our studies, we determined that environmental conditions such as a change in the pH of the solution resulted in increased toxicity of diclofenac, oxytetracycline, and fluoxetine, but only in a limited range of pH, chloramphenicol showed lower genotoxicity at low pH ranges. All ions, in particular, potassium, fluoride, and bromide, contributed to the reduction of toxicity of the pharmaceuticals tested, however, only in the case of diclofenac

and fluoxetine antagonism was observed. The results indicate that synergism was mainly observed for mixtures of fluoxetine h. with diclofenac s. and chloramphenicol with oxytetracycline h. with fluoxetine h., chloramphenicol showed antagonistic interactions depending on the concentrations used.

The results obtained indicate that the environmental conditions can contribute to increase or decrease of drug's genotoxicity, which is worth considering when conducting ecotoxicological studies, because the results obtained in the laboratory may differ from phenomena actually occurring in the media due to the presence of other substances or additional factors. Table 4 specifies how many cases of synergism and antagonism were observed for both CA and IA models and the percentiles for the whole of the research stages that were counted.

Comet assay showed high sensitivity to 4 pharmaceuticals (out of ten studied) and the research was carried out on cell culture, organisms from higher organization level have defensive and repair mechanisms start when exposure to genotoxic compounds is detected, however, it does not mean that this type of chemicals can be ignored. This study provides an important basis for further investigation into examination of mixtures of chemical compounds and characterization of genotoxic mechanism of drug residues and they can be an introduction to more complex multi-component mixtures.

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