

1 **RESEARCH ARTICLE**

2 **Dissipation of the herbicide active ingredient glyphosate in natural water samples**
3 **in the presence of biofilms**

4
5 Szandra Klátyik^{a*}, Eszter Takács^a, Mária Mörtl^a, Angéla Földi^b, Zsuzsa Trábert^b, Éva
6 Ács^b, Béla Darvas^a and András Székács^a

7
8 ^a *Agro-Environmental Research Institute, National Agricultural Research and*
9 *Innovation Centre, Budapest, Hungary;* ^b *MTA Centre for Ecological Research, Danube*
10 *Research Institute, Budapest, Hungary*

11
12
13 Dissipation of the herbicide active ingredient glyphosate by microbial communities and by
14 physical sorption on the surface of biofilms and solid particles in water was investigated in
15 natural waters in Hungary. To assess combined effects, glyphosate was applied in its pure form
16 (glyphosate isopropylammonium salt) and in preparation Roundup Classic[®] formulated with
17 polyethoxylated tallowamines (POEA). Standing and running surface water samples were
18 originated from Lake Balaton and River Danube between early May and mid-June of 2015.
19 Natural biofilms, grown on glass substrates fixed to AKK-1 type carrier buoy, were obtained
20 from the same locations. The kinetics of dissipation of glyphosate was investigated for 5 weeks,
21 under controlled laboratory conditions in aquaria containing natural water (15 L), with or
22 without the presence of mostly algal biofilms, with water exchange from the original locations
23 every week. The concentration of glyphosate was measured, upon chemical derivatisation with
24 9-fluorenylmethyloxycarbonyl chloride and solid phase extraction, by high-performance liquid
25 chromatography combined with UV-VIS absorbance detection or tandem mass spectrometry.
26 The quantity and the biofilm structure of algal biomass upon exposure to pure or formulated
27 glyphosate was determined by *in vivo* fluorimetry and by scanning electron microscopy. The
28 presence of POEA affected the dissipation of glyphosate, and dissipation profiles also differed
29 in the investigated natural water samples with or without the presence of biofilms. The results
30 indicate that glyphosate is capable to modify the structure of the algal community and to induce
31 increased secretion of extracellular polymeric substances matrix in the biofilms assessed.

32
33 **Keywords:** glyphosate; dissipation; biofilm; Roundup Classic; POEA

34 *Correspondence author:* Szandra Klátyik, tel.: +36 70 9311456, e-mail address:
35 sz.klatyik@cfri.hu

36
37 E-mail addresses of all Authors:

38 Szandra Klátyik sz.klatyik@cfri.hu
39 Eszter Takács e.takacs@cfri.hu
40 Mária Mörtl m.mortl@cfri.hu
41 Angéla Földi foldi.angela@okologia.mta.hu
42 Zsuzsa Trábert trabert.zsuzsa@okologia.mta.hu
43 Éva Ács acs.eva@okologia.mta.hu
44 Béla Darvas b.darvas@cfri.hu
45 András Székács a.szekacs@cfri.hu

46
47
48 **1. Introduction**

49 Various pesticide active ingredients and formulations used in intensive agriculture exert
50 high direct or mediated impact on the environment, especially in surface waters via their
51 leaching, drifting, surface run-off from treated sites, foliar spray and unintended
52 overspray and may pose hazards to the drinking water bases as well [1,2]. The appearance
53 of the worldwide used active ingredient glyphosate in surface water is a globally observed

*Corresponding author. Email: sz.klatyik@cfri.hu

54 phenomenon because of its good solubility in water and widespread use. The water
55 solubility of glyphosate is 11.6 g L^{-1} (25°C), while degradation half-life (DT_{50}) in water
56 is between 28 and 91 days (photodegradation excluded) [3]. Significant differences were
57 detected in glyphosate contamination all over the world. Although several studies report
58 levels of contamination at about $0.01 \text{ } \mu\text{g L}^{-1}$, i.e. near to the limit of detection (LOD)
59 [4,5], the average contamination level in surface water has been found between $100\text{-}200$
60 $\text{ } \mu\text{g L}^{-1}$ [6,7], and actual levels can reach up to $5200 \text{ } \mu\text{g L}^{-1}$ [8] in regions, where the use of
61 glyphosate-based pesticide formulations is substantial due to the cultivation of genetically
62 modified glyphosate-resistant crops. The concentrations of glyphosate in surface waters
63 in the European Union (EU) is between 0.05 and $4.7 \text{ } \mu\text{g L}^{-1}$ as reported in several studies
64 [4,9,10]. In the United States of America (USA), the accepted maximum level of for
65 glyphosate residues in drinking water is $700 \text{ } \mu\text{g L}^{-1}$ [11], while $0.1 \text{ } \mu\text{g L}^{-1}$ in the EU [12].
66 The acceptable maximum level of glyphosate (among all pesticide residues) is $1.0 \text{ } \mu\text{g L}^{-1}$
67 in the EU [13].

68 The half-life of glyphosate in environmental matrices is strongly influenced by
69 factors such as microbial activity. Glyphosate is rapidly adsorbed onto sediment particles
70 depending on the metal content of the sediment phase, and is gradually degraded into its
71 main metabolite, aminomethylphosphonic acid (AMPA). After 28 day post treatment
72 glyphosate and AMPA were detectable in surface water samples derived from an
73 estuarine pond, in contrast to the sediment samples, which did not contain the investigated
74 compounds [14].

75 Various co-formulants and additives used in pesticide formulations have traditionally
76 been considered as inactive/inert ingredients in pesticide formulations. However, these
77 substances are deliberately applied to modify the physical/chemical characteristics of the
78 active ingredient(s) in formulations, and several studies confirmed, that the formulating
79 agents, particularly polyethoxylated tallowamines (POEA), a complex combination of
80 homologs of different aliphatic moieties and ranges of ethoxylate units [15], exert their
81 own toxicity or affect the toxicity of the active ingredients [16,17]. Therefore,
82 comparative studies among pure active ingredients and their formulated products are of
83 increasing importance.

84 Biofilm development on natural or artificial solid surfaces in water media play a
85 particularly important role in the biogeochemical cycles, dynamics of the aquatic
86 ecosystems and biodegradation of pollutants in natural waters [18,19]. Biofilms are
87 compact communities of photoautotrophic (algae) and heterotrophic microorganisms
88 (bacteria, fungi, protozoa) embedded in their extracellular polymeric substance (EPS)
89 secretions [20]. EPS consists of proteins, polysaccharides, lipids, lectins, nucleic acids,
90 etc., and can serve as sorption sites [21]. The EPS matrix is a dynamic system,
91 responsible for the structure and morphology of the biofilms by filling and forming the
92 space between the algal cells [22]. The structure of the EPS matrix is significantly
93 stronger in the presence of various cations resulting in interactions with exposed carboxyl
94 groups on the EPS, formation of macromolecule networks, and increased viscosity or
95 gelation. The EPS matrix plays an important role in the protection of microbes against
96 physical-chemical stresses [23] and the sorption of toxic organic contaminants (e.g.
97 chlorophenols and polyaromatic hydrocarbons [24], atrazine, diclofop-methyl [25] or
98 organic pollutants BTX [26]), and additionally it concentrates nutrients [27]. Increased
99 production and secretion of the EPS matrix can be interpreted as stress responses of the
100 biofilms to different adverse effects [28,29]. Accumulation of various metal ions (e.g.
101 Cd^{2+} , Cr^{3+} , Cu^{2+}) by biofilms has been confirmed [30]: the sorption capacity of the
102 biofilms can be attributed to chelate or complex formation of the EPS matrix with various
103 cations, and the uptake of cations by bacteria and alga species in biofilms. Furthermore,

104 the binding capacity of the EPS matrix is significantly influenced by the pH of the water
105 and its physical stage (dissolved, slime or gel state) [31]. Biofilms are widely used for
106 monitoring studies, due to their sessile way of life; their rapid response to environmental
107 changes (because of their short life cycle); their microbial community consist of high
108 number of species with different sensitivity for various environmental effects; and the
109 easy way of sampling it [32,33]. The EPS matrix can trap nutrients from water for the
110 microorganisms in biofilms [34], and present a highly reactive surface area for sorption
111 and metabolism of chemical compounds [25]. In turn, biofilms can take part in the
112 adsorption, biodegradation and decomposition of the contaminants [35].

113 The aim of this study was to investigate and compare the dissipation of glyphosate
114 in pure and formulated forms in freshwater samples originated from Lake Balaton and
115 River Danube, with and without the presence of natural freshwater biofilms. Dissipation
116 was investigated as the biodegradation of glyphosate by microbial activities and physical
117 sorption on the surface of biofilms and solid particles of water samples.

118 119 **2. Experimental**

120 121 **2.1. Standards and reagents**

122 Glyphosate isopropylammonium (IPA) salt was received from Lamberti SpA (Albizzate,
123 Italy). Herbicide formulation Roundup Classic[®] (Monsanto Europe S.A./N.V.) [36] was
124 purchased from public commercial source. The main chemical characteristics of the
125 selected active ingredient, glyphosate-based herbicide and the surfactant POEA used in
126 Roundup Classic[®] can be found in Table 1. According to its Material Safety Data Sheet
127 (MSDS), Roundup Classic[®] contains 41.5% glyphosate IPA salt and 15.5% POEA, both
128 ingredients unequivocally identified by their Chemical Abstracts Service (CAS) Registry
129 Numbers (see Table 1). The authorization of Roundup Classic[®], formulated by Monsanto
130 Europe S.A. was cancelled its POEA content (see its MSDS) in Hungary at December
131 2016 [36,37]. All other chemicals, including analytical standards of glyphosate,
132 derivatising agent 9-fluorenylmethyl chloroformate (FMOC-Cl), organic solvents
133 acetonitrile (ACN), methanol (MeOH), dichloromethane, as well as phosphate and borate
134 buffers, aqueous formic acid and ammonium acetate for HPLC analyses and
135 glutaraldehyde for fixation for scanning electron microscopy were obtained from Sigma-
136 Aldrich Co. LLC (St. Louis, MO, USA). Analytical standards were ≥ 97.5 % purity. Solid
137 phase extraction was carried out using Strata-X Polymeric SPE cartridge (Phenomenex,
138 Torrance, USA) (volume of 3 mL, 200 mg sorbent).

139 140 **2.2. Experimental setup**

141 142 **2.2.1. Determination of dissipation in natural water samples**

143 Dissipation of glyphosate active ingredient was investigated in its pure (glyphosate IPA
144 salt) and formulated form (Roundup Classic[®] herbicide formulation) in surface waters of
145 two origins. Freshwater samples were originated from Lake Balaton (Tihany Bay –
146 46.914190, 17.892916, Tihany, Hungary), the largest standing water body in Europe and
147 River Danube (Green Island – 47.481641, 19.057645, Budapest, Hungary) the second
148 longest, navigable river of Europe. Water quality of the collected samples was
149 characterised by pH of 8.4-5.54 and 8.1-8.2, and conductivity of 650-700 and 715-755
150 $\mu\text{S cm}^{-1}$ for Lake Balaton and River Danube, respectively. The kinetics of dissipation
151 investigated under laboratory conditions in aquaria containing natural water (15 L) with
152 water exchange every week. During the experiments, the water in the aquaria was slowly
153 stirred (to assure oxygen dissolution), temperature-controlled ($22\pm 2^\circ\text{C}$) and illuminated

154 (L:D = 15:9, daily light program 6-9 hrs $5.4 \mu\text{mol m}^{-2} \text{s}^{-1}$ (photosynthetically active
155 radiation, PAR) (400 lux), 9-18 hrs $13.5 \mu\text{mol m}^{-2} \text{s}^{-1}$ (PAR) (2000 lux), 18-21 hrs 5.4
156 $\mu\text{mol m}^{-2} \text{s}^{-1}$ (PAR) (400 lux); XiLong White T8[®]). Illuminance (lux) was determined by
157 Light Meter MS-86 (Dostmann, Wertheim-Reicholzheim, Germany), PAR was
158 determined by Coherent[®] Field Max (Edmund Optics, Barrington, NJ, USA). For spiking,
159 pure glyphosate IPA salt and POEA-formulated glyphosate (Roundup Classic[®]) were
160 added to the aquaria containing original natural water samples, resulting in an initial
161 glyphosate concentration of $100 \mu\text{g L}^{-1}$ of the glyphosate IPA salt (equivalent to $74.1 \mu\text{g}$
162 L^{-1} glyphosate acid), corresponding to the lower range of average contamination levels
163 reported in surface waters [6,7].
164

165 2.2.2. *Determination of dissipation in presence of biofilms*

166 To determinate the dissipation in the presence of biofilms, natural biofilms were grown
167 on glass substrates (plates of dimensions: 23 cm x 9 cm, thickness: 3 mm, one side smooth
168 (untreated) and one side sand blasted) fixed on AKK-1 (originated from Cséffán, Darvas
169 & Pasaréti) type carrier buoys immersed for 6 weeks in Lake Balaton and River Danube
170 placed at the same location, where water sampling was regularly performed later
171 (described above) between early May and mid-June of 2015. Prior to the outplacement of
172 the carrier buoys, the orientation and intensity of waves, and the possibilities for
173 protection and the reach of the location were assessed. The AKK-1 buoy includes four
174 algal deposition rack units (containing no any metal or plastic elements) with 5 glass
175 plates in each unit, vertically submerged into the water (at a depth of 20-30 cm). After a
176 6-week colonization period, the glass substrates were placed into glass aquaria (without
177 any plastic elements) under laboratory conditions. Each aquarium contained 15 L water
178 from the original location of the buoy, and water parameters ($22 \pm 2 \text{ }^\circ\text{C}$, L:D = 15:9,
179 stirring) were controlled. Five biofilm substrates with sand blasted and smooth surface
180 sides were placed into each aquarium (the sixth substrate was used for further analytical
181 and microscopic evaluations). Control units in aquaria without glyphosate (pure or
182 formulated) treatment were applied during the experiments. The algal deposition units
183 were placed in the same position, and the order of the substrates was not modified in the
184 aquaria. The water in the aquaria was changed weekly, with water of original locations,
185 where the biofilm had been developed. The dissipation was investigated in case of both
186 glyphosate forms (formulated and pure active ingredient), and identical initial glyphosate
187 concentrations ($100 \mu\text{g L}^{-1}$ of glyphosate IPA salt, equivalent to $74.1 \mu\text{g L}^{-1}$ glyphosate
188 acid) were applied at the beginning of the experiments and upon each weekly water
189 exchange.
190

191 2.3. *Analytical methods*

192 193 2.3.1. *Sampling*

194 Dissipation of glyphosate was determined daily in freshwater samples originated from
195 Lake Balaton and River Danube, therefore 15 mL water sample was collected every day
196 from each aquarium during the experiment. In the presence of biofilms, dissipation was
197 investigated on the basis of sample collection performed daily during the first week (the
198 first sample taken in 30 minutes after glyphosate application), and weekly during each
199 further water exchange. The samples were frozen at -24°C until sample preparation and
200 measurement [38,39].
201

202 2.3.2. *Sample preparation*

203 Water samples (5 mL) were derivatised with 250 μL of FMOCCl (0.5 mM) and 0.3 ml
204 of borate buffer (pH 9) [40]. Upon 1 min of vigorous shaking, the solution was incubated
205 at room temperature for 1 hour. The excess amount of FMOCCl was removed by
206 extracting the reaction mixture three times, each with 1 ml of dichloromethane. The
207 aqueous phase separated was subjected to solid phase extraction (SPE) to concentrate the
208 samples for HPLC-UV analysis [41]. Cartridges (Strata-X sorbent, 33 μm , 200 mg;
209 Phenomenex, Torrance, USA) were conditioned by the addition of 5 mL of MeOH, then
210 5 mL of distilled water, and finally 5 mL of phosphate buffer (pH=3). Subsequently, the
211 derivatised water samples (5 mL) were added, the cartridges were washed with 3 mL of
212 distilled water, and were air-dried. The analytes were eluted with 3.5 mL of methanol, the
213 eluate was evaporated and redissolved in 0.5 mL of the initial eluent of the HPLC
214 analysis, and was filtered through a 0.45 μm hydrophilic polytetrafluoroethylene syringe
215 filter (FilterBio PFTE-L) purchased from Labex Ltd. (Budapest, Hungary). Derivatised
216 samples were not subjected to SPE prior to LC-MS/MS measurements.

217

218 2.3.3. Analytical determination

219 Glyphosate concentration of water samples was analysed by HPLC-UV using an
220 optimised analytical method reported elsewhere with fluorescent detection [40,42].
221 Negative samples in HPLC-UV (under LOD: 5 $\mu\text{g L}^{-1}$) were further analysed by LC-
222 MS/MS. HPLC-UV analyses of the investigated compounds were performed on a
223 Younglin YL9100[®] HPLC system equipped with an YL9150 autosampler. Glyphosate
224 were separated on a Chromegabond WR C18 column (150 mm \times 4.6 mm, i.d. 3 μm) (ES
225 Industries, Berlin, Germany) at 40 $^{\circ}\text{C}$. UV detector signals were recorded at $\lambda = 260 \text{ nm}$.
226 External calibration was based on the results obtained for 7 standard solutions in the range
227 of concentrations between 5 and 150 $\mu\text{g L}^{-1}$. Calibration solutions were prepared from a
228 stock solution by dilution with acetonitrile:buffer (10 mM ammonium acetate in water,
229 pH=6.0). The eluent flow rate was 0.7 mL min^{-1} with gradient elution. Initial eluent (1:9
230 = A:B eluents, A = 100% acetonitrile, B =10 mM sodium acetate buffer water) was
231 increased to 90% A at 6 min, maintained for 3 min, and then returned to initial
232 composition in a min and equilibrated for 3 min. The injection volume was 30 μL .

233

234 Water samples with glyphosate content below the LOD (5 $\mu\text{g L}^{-1}$) were subjected to
235 liquid chromatography–tandem mass spectrometry (LC-MS/MS) [39,41] on a Thermo-
236 Finnigan TSQ-20003 Quantum Discovery MAX (Thermo Electron Corp., San Jose,
237 USA) liquid chromatograph (LC) equipped with a triple quadrupole mass spectrometer
238 with electrospray ionization (ESI). Compounds were separated on a Kinetex XB-C18
239 column (2.1 mm \times 100 mm, i.d. 5 μm) (Phenomenex, Torrance, CA, USA, purchased
240 from Gen-Lab Ltd, Budapest, Hungary) at 25 $^{\circ}\text{C}$. Gradient elution was conducted with at
241 flow rate of 0.2 mL min^{-1} . Aqueous formic acid (0.1%, eluent A) and acetonitrile (eluent
242 B) were used as eluents. Prior to the measurements, both eluents were filtered through
243 regenerated cellulose filters (0.2 μm). The composition of the eluents was changed in time
244 as follows: 0 min 3% B, 2 min 3% B, 10 min 50% B, 15 min 3% B, 25 min 3% B.
245 Experiments were conducted in positive and negative ionization modes. The LOD of the
246 method was 1 ng L^{-1} .

246

247 2.4. Biological experiments

248

249 2.4.1. Sampling procedure

250 Prior to the location of the AKK-1 carrier buoy and 6-week biofilm colonization period
251 1 cm x 1 cm sand blasted glass plates were fixed on the biofilm glass substrates, and the
252 developed biofilms were used for the electron microscopic examination of the biofilms.

253 The collection of the biofilm samples were performed after completion of the biofilm
254 development period and at the end of the experiment.

255

256 2.4.2. *Sample preparation*

257 Biofilm samples were fixed prior to the scanning electron microscopy (SEM). During the
258 fixation of the biofilm samples using 10 mL of 5 % glutaraldehyde solution for 3 hours
259 at room temperature (20 °C), followed by two washing steps using 10 mL of 0.2 M
260 phosphate buffer for 10 min. The fixed biofilm samples were stored at -80 °C until
261 lyophilisation performed by Christ Alpha 1-4 LSC[®] (Osterode, Germany). During
262 lyophilisation, the duration of the main freeze-drying was 20 hours (1.025 mbar, -56 °C)
263 followed by 4-hour final drying (0.825 mbar, -56°C) [43]. The lyophilised samples were
264 fixed onto a stub using double-sided carbon tape followed by coating with gold by a
265 rotary-pumped sputter coater (Quorum Q150 R S[®], London, England).

266

267 2.4.3. *Biological determination*

268 The effects of active ingredient glyphosate and formulation Roundup Classic[®] on algal
269 biomass of biofilms were determined with bbe Moldaenke BenthosTorch[®]
270 (Schwentinental, Germany) algae torch instrument based on real-time measurement of
271 benthic algal concentrations by *in situ* quantification of chlorophyll-a fluorescence and *in*
272 *vivo* fluorescence of algal cells. During the measurement, algal cells are excited by LEDs
273 at different wavelengths and emit red fluorescence light. The algal biomass is calculated,
274 on the basis of the quantity of chlorophyll-a content of different algae, using the intensity
275 of chlorophyll fluorescence. The concentration of different algae was expressed in the
276 unit of μg chlorophyll-a cm^{-2} . The measuring range of the instrument is 0-15 μg
277 chlorophyll-a cm^{-2} [44]. However according to Kahlert and McKie, the use of
278 BenthosTorch[®] for determination of the relative contribution of different algal group to
279 benthic algal biomass is recommended only with cautious evaluation [45]. To assess the
280 accuracy of the algal biomass determination, chlorophyll-a content was determined from
281 the biofilm using the corresponding standardised protocol [46], and the two methods
282 (spectrophotometric and *in situ* fluorometric determination of chlorophyll-a) were
283 compared to each other in the 1-50 $\mu\text{g mL}^{-1}$ concentration range. Moreover, in our
284 experiments, the results were used for comparative purposes, therefore, the rates of the
285 three algae taxa (green algae, cyanobacteria and diatom) studied were evaluated with
286 results from SEM considered. The composition of the algae community of biofilms and
287 their structural transformations, as well as the intensity of EPS formation were visualised
288 from 15 randomly selected fields of each samples by SEM performed by Zeiss EVO MA
289 10[®] scanning electron microscope operated at 10 kV and 8.5 mm distance using SE
290 detector. Changes in algal biomass in response to exposure to the chemicals studied were
291 determined, but biomasses of untreated biofilms were also measured as negative controls
292 in each sampling interval. Control units were incubated in aquaria under the same
293 conditions as the treatment groups, but without glyphosate (pure or formulated) treatment.
294 Determinations were conducted on the sand blasted and smooth surface of glass substrates
295 as well in triplicates. On both sides of the substrate the identical sampling sites of 9.62
296 cm^{-2} were measured in every two weeks, and total and relative biomass values were
297 calculated. Standard deviations (SD) of biomass values between the sampling sites on the
298 individual sides, glass substrates and rack units were determined.

299

300 2.5. *Statistical analysis*

301 Decomposition of glyphosate in pure and formulated forms in natural waters was assessed
302 by sampling in triplicates, and each sample subjected to chemical analysis in triplicates.

Standard calibration for quantitative determination of glyphosate has also been carried out in triplicates at each concentration level. Experiments of exposure of biofilms to pure and formulated glyphosate were performed in quadruplicates by separately immersing five glass plates with biofilms into natural waters spiked with glyphosate or Roundup Classic[®]. Corresponding control experiments without treatment with glyphosate have also been carried out in quadruplicates. Algal biomass was determined on each glass plate in two spots (9.62 cm² each) on each side of the plate, with even geometrical distribution along the plate and identical setup throughout the experiment in each treatment group. Thus, overall 20 parallel fluorometric determinations were carried out for each time points of each treatment. Extraction for spectrophotometric measurement of chlorophyll content was carried out in triplicates at each concentration level. Effects of various treatments were statistically evaluated by one-way ANOVA (Statistica[®] software, StatSoft, Tulsa, USA) followed by Tukey *post hoc* test for comparisons between groups ($p \leq 0.05$).

3. Results and discussion

3.1. Pesticide residue analysis in surface water

The retention time in the HPLC separation was 6.71 min for glyphosate. An LOD, defined as analyte concentrations corresponding to a signal level of signal/noise ratio of 3, of the developed HPLC-UV analytical method was 5 $\mu\text{g L}^{-1}$. The percentage recovery at a spiking level 100 $\mu\text{g L}^{-1}$ of the glyphosate IPA salt (equivalent to 74.1 $\mu\text{g L}^{-1}$ glyphosate acid) was found to be 83.5 \pm 6.0% for glyphosate. Glyphosate concentrations above 5 $\mu\text{g L}^{-1}$ reported in this manuscript correspond to analyses by HPLC-UV. In the rare cases, when glyphosate concentrations fell below 5 $\mu\text{g L}^{-1}$, water samples were analysed by LC-MS/MS.

The pesticide contamination status of the natural water bodies at both sampling locations was investigated weekly during the biofilm formation and sampling periods, and no detectable amounts of glyphosate residues were found. During the colonisation period of biofilms in river Danube, metolachlor (up to 1 $\mu\text{g L}^{-1}$) was detected for a longer period, and occasionally terbutylazine and dimethenamid also occurred (up to 1 $\mu\text{g L}^{-1}$). In mid-July, chlorpyrifos appeared (2-4 $\mu\text{g L}^{-1}$) in the water samples until the end of the sampling period. In contrast, no pesticide residues in the water samples from Lake Balaton were detected during the colonisation period, but later the presence of chlorpyrifos (2-4 $\mu\text{g L}^{-1}$) was detected at the same concentration range as seen in river Danube.

3.2. Effects of pure and formulated glyphosate on algal biomass and composition of biofilms

The *in situ* fluorometric algae torch was found a reproducible method for the determination of chlorophyll-a content in biofilms, as the surface density of chlorophyll-a detected highly correlated with corresponding chlorophyll-a concentrations measured by the ISO standard method of spectrometric determination of the chlorophyll-a concentration in water quality assessment [46]. Chlorophyll-a surface densities and concentrations highly correlated ($R^2 = 0.9996$) with each other in the concentration range of 1-50 $\mu\text{g mL}^{-1}$ of chlorophyll-a.

Due to identical geometric arrangement of the algae rack units containing 6 racks each, total production rate of biomass grown on the AKK-1 type buoy was not statistically different among rack units for Lake Balaton and River Danube, respectively. Thus, differences in glyphosate concentration among treatment groups were not due to the initial biomass, but to the condition, whether glyphosate was applied in its pure or

353 formulated form. Effects in biomass production were determined on identical surface
354 dimensions among the 6 glass substrates. Higher biomass values were measured on the
355 edge of glass plates and on the terminal plates. Maximum relative SD (SD%) of the
356 average biomass content among sampling sites were 35% and 40%, for Lake Balaton and
357 River Danube, respectively. However, commensurable biomass results, significantly not
358 different from each other, were determined among rack units in the case of both surface
359 water sources. Average biomass production on the 2-2 rack units (used in this dissipation
360 experiment) after the colonization period (before treatments with two form of glyphosate)
361 were 2.26 and 2.13 μg chlorophyll-a cm^{-2} for River Danube and 3.21 and 3.32 μg
362 chlorophyll-a cm^{-2} for Lake Balaton.

363 On-going spontaneous changes in the algal community and the structure of the
364 biofilms from River Danube in response to the various treatments were observed by algal
365 biomass measurement and microscopic analysis, while such alterations were not observed
366 in the corresponding control units. Biofilms originated from River Danube continued to
367 grow under laboratory conditions, unlike those from Lake Balaton (see below). Exposure
368 to glyphosate alone occurred to slightly promote biomass production. This is not
369 unreasonable, as it has been reported that glyphosate at low concentrations (0.01 to 5 mg
370 P L^{-1}) may serve as a source of phosphate and nutrients for certain biofilm community
371 components [47], and/or may trigger pathways for the synthesis of metabolites and
372 proteins [48,49], which can result in increased biomass growth. At higher concentrations
373 (8 mg L^{-1}), however, it inhibits the colonization of algae [50]. Upon treatments with
374 POEA-formulated glyphosate (Roundup Classic[®]), the initial biomass decreased in the
375 first 2 weeks in both surface waters. Average relative biomass values were 2.04, 2.14 and
376 1.50 μg chlorophyll-a cm^{-2} for algae grown on glass substrates in River Danube for the
377 control and the glyphosate and POEA-formulated glyphosate treatments, respectively.
378 After 2 weeks, biomass in River Danube started to increase.

379 In contrast, initial biomass from Lake Balaton decreased continuously during the
380 five-week experimental period not only under treatments with pure and POEA-
381 formulated glyphosate, but in the control experiment as well from the second week on, as
382 indicated by *in situ* fluorimetry and SEM images. These biofilms were rich in small, tube-
383 building, algivorous chironomid larvae; *Procladius choreus*, *Tanytus punctipennis* and
384 *Chironomus balatonicus* being the most abundant at the Tihany Peninsula [51,52]. The
385 emergence of these larvae, especially *Procladius* species occurred to be essential for the
386 subsistence of the biofilms, and in cases of lacking emergence, the biofilms collapsed in
387 the aquaria in two weeks. After the two-week incubation period, 2.65, 2.82 and 2.30 μg
388 chlorophyll-a cm^{-2} were determined for the control and the treatment groups with pure
389 and formulated glyphosate, respectively.

390 SEM analysis indicated considerable changes in biofilm structure. Realignment
391 of the biofilms was typical, and glyphosate-sensitive species were replaced by tolerant
392 ones like filamentous green algal species (Figure 1). The realignment of biofilms and the
393 effects of glyphosate on the microbial community structure in freshwater were observed
394 in other studies as well [50,53]. The electron microscopic analysis also indicated
395 increased production of the EPS matrix, relative to the corresponding negative controls,
396 in each treatment group. Visual analysis of the ESM images suggested an intensive EPS
397 production for exposure to POEA-formulated glyphosate. This phenomenon can be
398 attributed to the protective mechanism of bacteria and algae to eliminate and reduce the
399 effects of contaminants [23,28,29]. Additionally, glyphosate can affect the metabolic
400 processes of bacteria and algae simultaneously, resulting in an enhanced production of
401 the EPS matrix as response to physical, chemical and biological stress factors [28,29]
402 (Figure 2).

403

404 **3.3. Dissipation of pure and formulated glyphosate in natural water samples** 405 **without the presence of biofilms**

406 Differences were observed between pure and POEA-formulated glyphosate levels (Figure
407 3). Significantly higher initial concentrations were measured (30 min after the addition of
408 $100 \mu\text{g L}^{-1}$ of the glyphosate IPA salt (equivalent to $74.1 \mu\text{g L}^{-1}$ glyphosate acid) in water
409 samples originated from River Danube for formulated glyphosate treatment due to the
410 presence of formulating agent POEA. A possible mechanism involved in this process can
411 be that the surfactant suppressed the physical adsorption of glyphosate on the solid-liquid
412 surfaces (e.g. glass materials of aquaria, solid phase and floating particles in water
413 samples) [54].

414 Degradation of glyphosate was not detected in water samples from Lake Balaton,
415 the level of glyphosate stagnated at 90 and $100 \mu\text{g L}^{-1}$ in case of the pure and POEA-
416 formulated active ingredient, respectively. Therefore, the observed changes in
417 concentration are likely to be due to absorption or accumulation in the tissue of the
418 biofilm. In contrast, the concentration of glyphosate in River Danube, after an initial rapid
419 decrease, reached a constant level approximately at the concentration of $60 \mu\text{g L}^{-1}$.

420 According to our results, the environmental fate and degradation of glyphosate can
421 be different in various natural water matrices, as the processes may be influenced by the
422 presence of the formulating agents, the composition of the microbial communities, and
423 the physical and chemical parameters of the water phase [14,55].

424

425 **3.4. Dissipation of pure and formulated glyphosate in the presence of biofilms**

426 Differences were observed between the reduction of pure and POEA-formulated
427 glyphosate levels in the presence of biofilms. Similar effects of the formulating agent
428 POEA on initial glyphosate concentrations (30 min after the addition of $100 \mu\text{g L}^{-1}$ of the
429 glyphosate IPA salt (equivalent to $74.1 \mu\text{g L}^{-1}$ glyphosate acid) as described in Section
430 3.2 (Figures 4-7). However, the presence of the biofilm resulted in further decreases of
431 glyphosate levels, likely due to the adsorption capacity [24-26] of the EPS matrix
432 produced by microbial activity of the biofilms. When pure glyphosate was applied, after
433 an immediate (within 30 minutes) steep drop, glyphosate concentration remained stagnant
434 during the first week at 15 and $80 \mu\text{g L}^{-1}$ for River Danube and Lake Balaton, respectively.
435 When applied in formulation, glyphosate concentrations decreased similarly, but less
436 instantaneously likely due to the surfactant effect of POEA, possibly facilitating the
437 maintenance of the active ingredient molecules in solution.

438

439 **3.4.1. River Danube**

440 The phytotoxic effects of glyphosate, particularly if enhanced by a formulating agent,
441 may have contributed to the observed decrease of the algal biomass relative to the
442 untreated control. Moreover, the gradual increase in glyphosate concentrations detected
443 after repeated weekly addition of $100 \mu\text{g L}^{-1}$ of pure glyphosate IPA salt (equivalent to
444 $74.1 \mu\text{g L}^{-1}$ glyphosate acid) is likely to be due to saturation of the sorption sites in the
445 EPS matrix in the biofilm. By the fourth week, the total biomass increased, accompanied
446 by significant decreases in glyphosate concentration, possibly due to the utilization of
447 glyphosate from water as a nutrient by tolerant algal species (Figure 4) [34,48].

448

449 When glyphosate was applied in a formulated form, the treatment resulted in a rapid
450 gradual decrease of the concentration of glyphosate during the first week in the presence
451 of high biomass. The treatment resulted in a decrease in the algal biomass, relative to the
452 untreated control, within 2 weeks. Possible factors contributing to this trend are the
453 phytotoxic effect of the formulation and the increased production of the EPS matrix

453 observed in a qualitative estimation based on the SEM images. The measured level of
454 glyphosate was stagnant upon weekly additions of glyphosate. From the third week on,
455 gradually increasing glyphosate concentrations were detected likely due to the saturation
456 of the sorption sites in the EPS matrix (Figure 5). Similarly to the treatment with pure
457 glyphosate, the algal biomass increased by the fourth week. Despite the lower
458 bioavailability of glyphosate in water, tolerant algal species occurred utilising glyphosate
459 as a nutrient from the EPS matrix.

460

461 3.4.2. Lake Balaton

462 Biofilms formed in Lake Balaton resulted different dissipation patterns of glyphosate
463 than those seen for River Danube. The phytotoxic effect of glyphosate or Roundup
464 Classic[®] herbicide formulation resulted in a continuous decrease in the biomass during
465 the five-week experimental period. Compared to the degradation without the presence of
466 biofilms, lower concentrations of glyphosate were detected in the first week possibly
467 attributed to chelate or complex formation with the EPS matrix [31]. After the first week
468 during the weekly, repeated addition of pure glyphosate into the aquaria, the
469 concentration of glyphosate stabilised at the same level as observed in the first week, but
470 on the fifth week the concentration ($62.3 \mu\text{g L}^{-1}$) of the spiked glyphosate dose was
471 significantly reduced 30 minutes after the addition (Figure 6).

472 Upon treatment with POEA-formulated glyphosate, the initial decline in glyphosate
473 concentration during the first week was less rapid as observed with pure glyphosate. Upon
474 repeated addition of formulated glyphosate, the entire dose ($100 \mu\text{g L}^{-1}$ of pure glyphosate
475 IPA salt (equivalent to $74.1 \mu\text{g L}^{-1}$ glyphosate acid) applied was detected in the water
476 samples 30 minutes after treatment until the fourth week, when the level of glyphosate
477 detected slightly dropped ($86.5 \mu\text{g L}^{-1}$) (Figure 7). This is expected to result from an
478 increased stress response of the algal community to the exposure to Roundup Classic[®],
479 potentially resulting in an increased EPS matrix production.

480

481 4. Conclusion

482 Among studies on pesticide formulating agents only a few investigate the effects of
483 surfactants on the environmental fate of the active ingredients. Our results demonstrate
484 that dissipation of glyphosate can be different in various natural waters, and additionally
485 highly depends on the presence of the formulating agents, the composition of the
486 microbial communities exposed, as well as the physical and chemical parameters of the
487 water phase. Dissipation profiles of given glyphosate forms were different in natural
488 water samples investigated without or in the presence of biofilms. Worldwide detectable
489 water contamination by glyphosate can modify the structure of the algal communities in
490 freshwater biofilms, and may induce increased stress response in them. Tests used for
491 authorisation and environmental risk assessment of the active ingredients and their
492 formulations are based on DT_{50} values determined in distilled water under laboratory
493 conditions. However, several data and our results suggest that a revision of the applied
494 DT_{50} values and determination of habitat-specific data are needed to be used in the
495 environmental risk assessment of the pesticide active ingredients and their formulations.

496

497 Acknowledgement

498 The authors thank to Péter Bohus (Lamberti SpA, Albizzate, Italy) for the sample of glyphosate IPA salt;
499 to Gyula Pasaréti and Tamás Cséffán (Agro-Environmental Research Institute, National Agricultural
500 Research and Innovation Centre) for their contribution in the development of the AKK-1 carrier buoy; to
501 Prof. Gyula Záray and Pál Hofmann (Eötvös Loránd University, Green Island, Budapest, Hungary), to
502 János Győri and Géza Dobos (Balaton Limnological Institute, Centre for Ecological Research, HAS,
503 Tihany) for the outplacement of the buoys on River Danube and Lake Balaton, respectively; to Ottó Etlér

504 (C Mobil Labor Kft, Zalakomár, Hungary) for the LC-MS/MS determinations; as well as to Csilla Magor,
505 Judit Juracsek, Balázs Magyarósy and Dániel Takács for their technical support.

506

507 **Disclosure statement**

508 No potential conflict of interest was reported by the authors.

509

510 **Funding**

511 This work was supported by the Hungarian Scientific Research Fund [OTKA K109865] and the Hungarian
512 Ministry of Agriculture [FM AD002].

513

514 **References**

- 515 [1] K.R. Solomon and D.G. Thompson, *J. Toxicol. Environ. Health B Crit. Rev.* **6**, 289
516 (2003). doi: 10.1080/10937400306468
- 517 [2] I. Hanke, I. Wittmer, S. Bischofberger, C. Stamm and H. Singer, *Chemosphere* **81**,
518 422 (2010). doi: 10.1016/j.chemosphere.2010.06.067
- 519 [3] C. MacBean, editor, *The Pesticide Manual*, 16th Edition (The British Crop
520 Protection Council, Brighton, 2012).
- 521 [4] J. Kjaer, P. Olsen, M. Ullum and R. Grant, *J. Environ. Qual.* **34**, 608 (2005). doi:
522 10.2134/jeq2005.0608
- 523 [5] R.H. Coupe, S.J. Kalkhoff and P.D. Capel, *Pest. Manag. Sci.* **68**, 16 (2012). doi:
524 10.1002/ps.2212
- 525 [6] J.C. Feng, D.G. Thompson and P.E. Reynolds, *J. Agricult. Food Chem.* **38**, 1110
526 (1990). doi: 10.1021/jf00094a045
- 527 [7] P.J. Peruzzo, A.A. Porta and A.E. Ronco, *Environ. Pollut.* **156**, 61 (2008). doi:
528 10.1016/j.envpol.2008.01.015
- 529 [8] W.M. Edwards, G.B. Triplett and R.M. Kramer, *J. Environ. Qual.* **9**, 661 (1980).
530 doi: 10.2134/jeq1980.00472425000900040024x
- 531 [9] M. Mörtl, Gy. Németh, J. Juracsek, B. Darvas, L. Kamp, F. Rubio and A. Székács,
532 *Microchem. J.* **107**, 143 (2013). doi: 10.1016/j.microc.2012.05.021
- 533 [10] A. Székács, M. Mörtl and B. Darvas, *J. Chem.*, **2015**, Article ID 717948 (2015).
534 doi: 10.1155/2015/717948
- 535 [11] United States Environmental Protection Agency (US EPA), Occurrence Estimation
536 Methodology and Occurrence Findings Report for the Six-Year Review of Existing
537 National Primary Drinking Water Regulations, US EPA Report No. EPA-815-R-
538 03-006, 2003.
- 539 [12] European Parliament and Council, *Offic. J. Eur. Comm.* **L330**, 32 (1998).
- 540 [13] European Parliament and Council, *Offic. J. Eur. Comm.* **L129**, 23 (1976).
- 541 [14] M.T. Tsui and L.M. Chu, *Chemosphere* **71**, 439 (2008). doi:
542 10.1016/j.chemosphere.2007.10.059
- 543 [15] D. Tush, K.A. Loftin and M.T. Meyer, *J. Chromatogr. A* **1319**, 80 (2013). doi:
544 10.1016/j.chroma.2013.10.032
- 545 [16] J.M. Brausch and P.N. Smith, *Arch. Environ. Contamin. Toxicol.* **52**, 217 (2007).
546 doi: 10.1007/s00244-006-0151-y
- 547 [17] I. Székács, Á. Fejes, Sz. Klátyik, E. Takács, D. Patkó, J. Pomóthy, M. Mörtl, R.
548 Horváth, E. Madarász, B. Darvas and A. Székács, *Internat. J. Biol., Vet. Food*
549 *Engineer.* **87**, 213 (2014).
- 550 [18] M. Schorer and M. Eisele, *Water Air Soil Pollut.* **99**, 651 (1997). doi:
551 10.1007/BF02406904
- 552 [19] G. Lear, editor, *Biofilms and Bioremediation: Current Research and Emerging*
553 *Technologies* (Caister Academic Press, New Zealand, 2016).
- 554 [20] W.G. Characklis and K.C. Marshall, *Biofilms* (John Wiley & Sons, New York,
555 1990).

- 556 [21] I.W. Sutherland, *Microbiology* **147**, 3 (2001). doi: 10.1099/00221287-147-1-3
- 557 [22] Z. Lewandowski, P. Stoodley, S. Altobelli and E. Fukushima, *Water Sci. Technol.*
- 558 **29**, Article ID 223e229. (1994).
- 559 [23] J. Wingender, T.R. Neu and H.-C. Flemming, in *Microbial Extracellular Polymeric*
- 560 *Substances. Characterization, Structure and Function*, edited by J. Wingender,
- 561 T.R. Neu, H.-C. Flemming (Springer, Berlin, 1999).
- 562 [24] E. Antusch, J. Sauer, C. Ripp and H.H. Hahn, *Gas Wasser Abwass.* **75**, 1010 (1995).
- 563 [25] J.R. Lawrence, G. Kopf, J.V. Headley and T.R. Neu, *Canad. J. Microbiol.* **47**,
- 564 Article ID 634e641 (2001). doi: 10.1139/w01-061
- 565 [26] R. Spath, H.-C. Flemming and S. Wuertz, *Wat. Sci. Tech.* **37**, 207 (1998). doi:
- 566 10.1016/s0273-1223(98)00107-3
- 567 [27] C. Freeman, P.J. Chapman, K. Gilman, M.A. Lock, B. Reynolds and H.S. Wheeler,
- 568 *Hydrobiologia* **297**, 61 (1995). doi: 10.1007/BF00033502
- 569 [28] D.S. Domozych, *Int. J. Plant. Sci.* **168**, 763 (2007).
- 570 [29] H.-C. Flemming and J. Wingender, *Nat. Rev. Microbiol.* **8**, 623 (2010). doi:
- 571 10.1038/nrmicro2415
- 572 [30] F.G. Ferris, S. Schultze, T.C. Witten, W.S. Fyfe and T.J. Beveridge, *Appl. Environ.*
- 573 *Microb.* **55**, 1249 (1989).
- 574 [31] A.W. Decho, *Cont. Shelf. Res.* **20**, 1257 (2000). doi: 10.1016/S0278-
- 575 4343(00)00022-4
- 576 [32] P.V. McCormick and J. Cairns, *J. Appl. Phycol.* **6**, 509 (1994). doi:
- 577 10.1007/bf02182405
- 578 [33] M. Mages, M. Ovári, W. von Tümpling and K. Kröpfl, *Anal. Bioanal. Chem.* **378**,
- 579 1095 (2004). doi: 10.1007/s00216-003-2291-5
- 580 [34] N. Das, L.V.G. Basak, J.A. Salam, M.E.A. Abigail, *J. Microbiol. Biotechnol. Res.*
- 581 **2**, Article ID 783e790 (2012).
- 582 [35] I. Bohuss, T. Rékasi, Sz. Szikora, K. Barkács, Gy. Zárny and É. Ács, *Microchem.*
- 583 *J.* **79**, 201 (2005). doi: 10.1016/j.microc.2004.08.001
- 584 [36] Monsanto Europe S.A., Material Safety Data Sheet of Roundup Classic®.
- 585 [http://www.sdslibrary.monsanto.com/Lists/MSDS%20Library/DispForm.aspx?ID=1203\(2015\)](http://www.sdslibrary.monsanto.com/Lists/MSDS%20Library/DispForm.aspx?ID=1203(2015))
- 586 [accessed in English version 10.06.2017];
- 587 <http://www.sdslibrary.monsanto.com/MSDS%20Datashet/7ce97326-c83a-4983-8b0a-8156ffa779de/Roundup%20Classic-9543CLPhu-hu.pdf> (2015) [accessed in
- 588 Hungarian version 17.06.2017]
- 589
- 590 [37] National Food Chain Safety Office (2016). Authorizations of some glyphosate-
- 591 based herbicides are cancelled. (in Hungarian) <http://portal.nebih.gov.hu/-/szamos-glifozat-keszitmeny-engedelye-visszavonasra-kerul> [accessed in 17.06.2017]
- 592
- 593 [38] H. Kylin, *Chemosphere* **90**, 1821 (2013). doi: 10.1016/j.chemosphere.2012.09.020
- 594 [39] J.D. Byer, J. Struger, P. Klawunn, A. Todd and E. Sverko, *Environ. Sci. Technol.*
- 595 **42**, 6052 (2008). doi: 10.1021/es8005207
- 596 [40] T.V. Nedelkoska and G.K.C. Low, *Anal. Chim. Act.* **511**, 145 (2004). doi:
- 597 10.1016/j.aca.2004.01.027
- 598 [41] I. Hanke, H. Singer and J. Hollender, *Anal. Bioanal. Chem.* **391**, 2265 (2008). doi:
- 599 10.1007/s00216-008-2134-5
- 600 [42] M. Küsters and M. Gerhartz, *J. Sep. Sci.* **33**, 1139 (2010). doi:
- 601 10.1002/jssc.200900556
- 602 [43] D. Anda, G. Büki, G. Krett, J. Makk, K. Márialigeti, A. Eröss, J. Mádl-Szőnyi and
- 603 A.K. Borsodi, *Acta. Microbiol. Immunol. Hung.* **61**, 329 (2014). doi:
- 604 10.1556/AMicr.61.2014.3.7
- 605 [44] bbe Moldaenke, BenthosTorch <

- 606 moldaenke.de/en/products/chlorophyll/details/benthotorch.html> (2017) [accessed
607 17.06.2017].
- 608 [45] M. Kahlert and B.G. McKie, Environ. Sci. Process. Impact **16**, 2627 (2014). doi:
609 10.1039/c4em00326h
- 610
- 611 [46] International Organization for Standardization, *ISO 10260:1992 Water quality --*
612 *Measurement of biochemical parameters -- Spectrometric determination of the*
613 *chlorophyll-a concentration* (International Organization for Standardization,
614 Geneva, 1992), p. 6.
- 615 [47] H. Qiu, J. Geng, H. Ren, X. Xia, X. Wang and Y. Yu, J. Hazard. Mater. **248-249**,
616 172 (2013). doi: 10.1016/j.jhazmat.2012.12.033
- 617 [48] M.A. Saxton, E.A. Morrow, R.A. Bourbonniere and S.W. Wilhelm, J. Great Lakes
618 Res. **37**, 683 (2011). doi: 10.1016/j.jglr.2011.07.004
- 619 [49] C. Wang, X. Lin, L. Li and S. Lin, PLoS One **11**, e0151633 (2016). doi:
620 10.1371/journal.pone.0151633
- 621 [50] M.S. Vera, L. Lagomarsino, M. Sylvester, G.L. Pérez, P. Rodríguez, H. Mugni, R.
622 Sinistro, M. Ferraro, C. Bonetto, H. Zagarese and H. Pizzaro, Ecotoxicology **19**,
623 710 (2010). doi: 10.1007/s10646-009-0446-7
- 624 [51] A. Specziár, Ann. Limnol. – Int. J. Lim. **44**, 181 (2008). doi: 10.1051/limn:2008002
- 625 [52] D. Árvai, M. Tóth, H. Horváth, S.A. Nagy and A. Specziár, Hydrobiologia **742**, 249
626 (2015). doi: 10.1007/s10750-014-1989-z
- 627 [53] G.L. Pérez, A. Torremorell, H. Mugni, P. Rodríguez, M. Solange Vera, M. do
628 Nascimento, L. Allende, J. Bustingorry, R. Escaray, M. Ferraro, I. Izaguirre, H.
629 Pizarro, C. Bonetto, D.P. Morris and H. Zagarese, Ecol Appl., **17**, 2310 (2007). doi:
630 10.1890/07-0499.1
- 631 [54] L. Ming, D. Fengpei, C. Chong, L. Baoying and Z. Xihai, J. Dispersion. Sci. Techn.
632 **37**, 213 (2016). doi: 10.1080/01932691.2015.1039022
- 633 [55] L.G. Goldsborough and D.J. Brown, Environ. Toxicol. Chem. **12**, 1139 (1993). doi:
634 10.1002/etc.5620120702
- 635

636 **Table 1.**
 637 **Chemical characteristics of the selected plant protection product, active ingredient and surfactant**
 638

Substance ^a	Chemical or product name	Chemical structure	CAS No. ^b	Concentration in formulation	Type of formulation
PPP	Roundup Classic [®]		–	–	liquid
a.i.	glyphosate isopropylammonium (IPA) salt		38641-94-0	41.5%	liquid
surfactant	polyethoxylated tallowamines (POEA)		61791-26-2	15.5%	liquid

639 ^a PPP: plant protection product; a.i.: active ingredient

641 ^b CAS No.: Chemical Abstracts registry number

642

643

644 Figure legends:

645

646 Figure 1. Occurrence of filamentous green algae (*indicated by arrow*) in natural biofilms
647 from River Danube, due to treatment, visualised by scanning electron microscopy. A:
648 Control biofilm without green algae (as verified by fluorimetry). B: The characteristic
649 filaments of green algae occurring upon exposure to POEA-formulated glyphosate-based
650 herbicide.

651

652 Figure 2. Increased production of EPS matrix (*indicated by arrow*) in natural biofilms
653 from River Danube, due to treatment, visualised by scanning electron microscopy. A:
654 Control biofilm with smooth EPS layer. B: Intensive EPS formation upon exposure to
655 POEA-formulated glyphosate-based herbicide.

656

657 Figure 3 Dissipation of the IPA salt of glyphosate in pure form (*hollow markers*) and in
658 preparation Roundup Classic® (*filled markers*) in water samples from River Danube (□/■)
659 and Lake Balaton (◇/◆). Glyphosate concentrations were detected with HPLC-UV.

660

661 Figure 4 Dissipation of pure glyphosate (□) in water samples from River Danube in the
662 presence of biofilms, depicting glyphosate concentrations (■) in 30 minutes after each
663 repeated glyphosate addition (▼). Arrows indicate concentration changes due to
664 dissipation (*solid lines*) or reagent addition (*dashed lines*). Biomass levels in the treatment
665 group (*open black columns with solid line*) and the untreated control (*open grey columns
666 with dashed line*) are indicated. Corresponding algal composition (*pie diagrams below
667 each column, treatment group in the upper and control in the lower row*) show the
668 biomass proportion of cyanobacteria (*dark grey*), diatom (*black*) and green (*light grey*)
669 algae.

670

671 Figure 5 Dissipation of formulated glyphosate (■) in water samples from River Danube
672 in the presence of biofilms, depicting glyphosate concentrations (■) in 30 minutes after
673 each repeated glyphosate addition (▼). Arrows indicate concentration changes due to
674 dissipation (*solid lines*) or reagent addition (*dashed lines*). Biomass levels in the treatment
675 group (*open black columns with solid line*) and the untreated control (*open grey columns
676 with dashed line*) are indicated. Corresponding algal composition (*pie diagrams below
677 each column, treatment group in the upper and control in the lower row*) show the
678 biomass proportion of cyanobacteria (*dark grey*), diatom (*black*) and green (*light grey*)
679 algae.

680

681 Figure 6 Dissipation of pure glyphosate (◇) in water samples from Lake Balaton in the
682 presence of biofilms, depicting glyphosate concentrations (■) in 30 minutes after each
683 repeated glyphosate addition (▼). Arrows indicate concentration changes due to
684 dissipation (*solid lines*) or reagent addition (*dashed lines*). Biomass levels in the treatment
685 group (*open black columns with solid line*) and the untreated control (*open grey columns
686 with dashed line*) are indicated. Corresponding algal composition (*pie diagrams below
687 each column, treatment group in the upper and control in the lower row*) show the
688 biomass proportion of cyanobacteria (*dark grey*), diatom (*black*) and green (*light grey*)
689 algae.

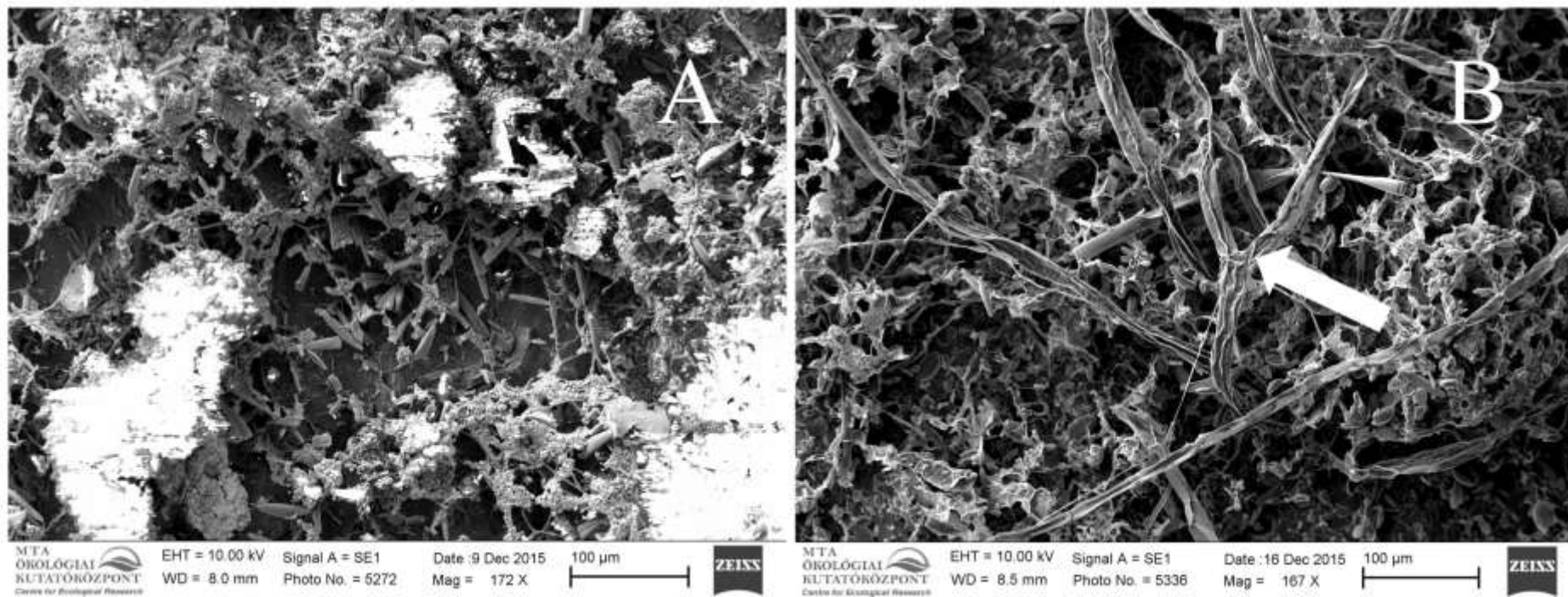
690

691 Figure 7 Dissipation of formulated glyphosate (◆) in water samples from Lake Balaton in
692 the presence of biofilms, depicting glyphosate concentrations (■) in 30 minutes after each
693 repeated glyphosate addition (▼). Arrows indicate concentration changes due to

694 dissipation (*solid lines*) or reagent addition (*dashed lines*). Biomass levels in the treatment
695 group (*open black columns with solid line*) and the untreated control (*open grey columns*
696 *with dashed line*) are indicated. Corresponding algal composition (*pie diagrams below*
697 *each column, treatment group in the upper and control in the lower row*) show the
698 biomass proportion of cyanobacteria (*dark grey*), diatom (*black*) and green (*light grey*)
699 algae.

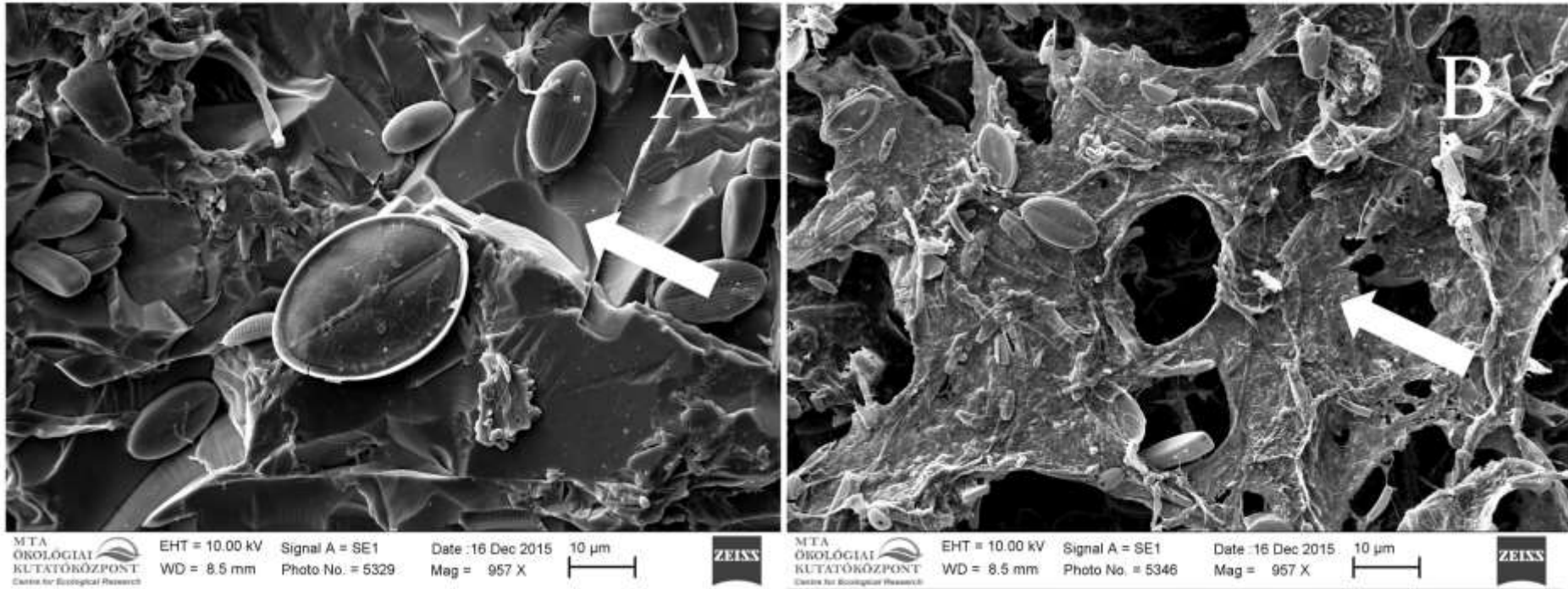
700

701 Figure 1
702



703
704
705

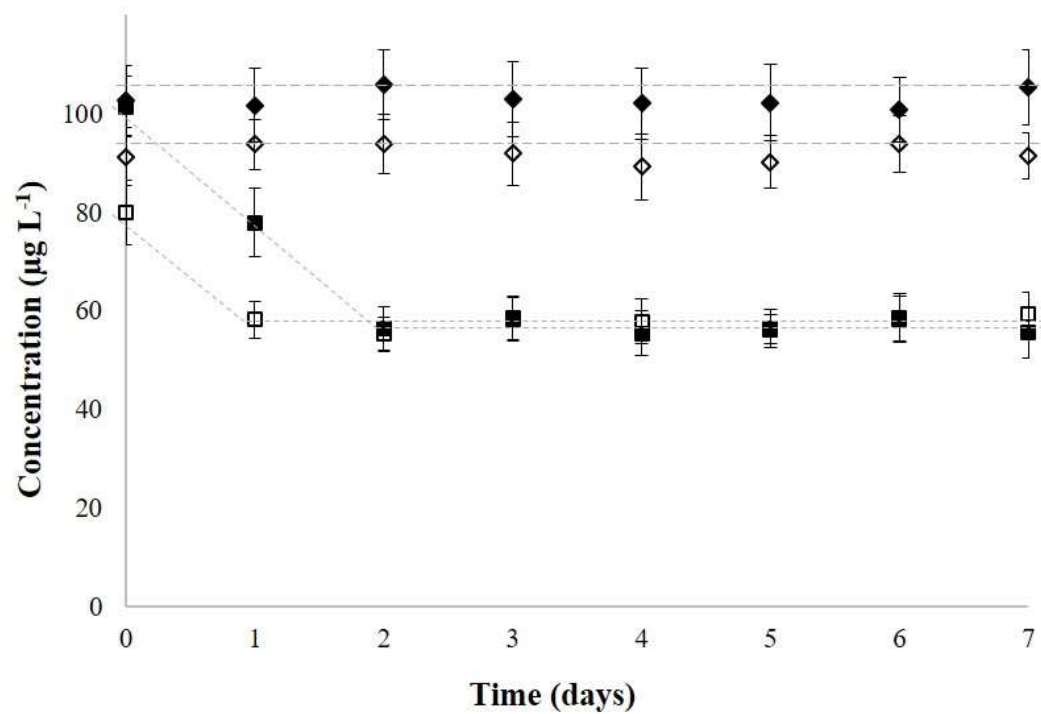
706 Figure 2
707



708
709
710

711 Figure 3

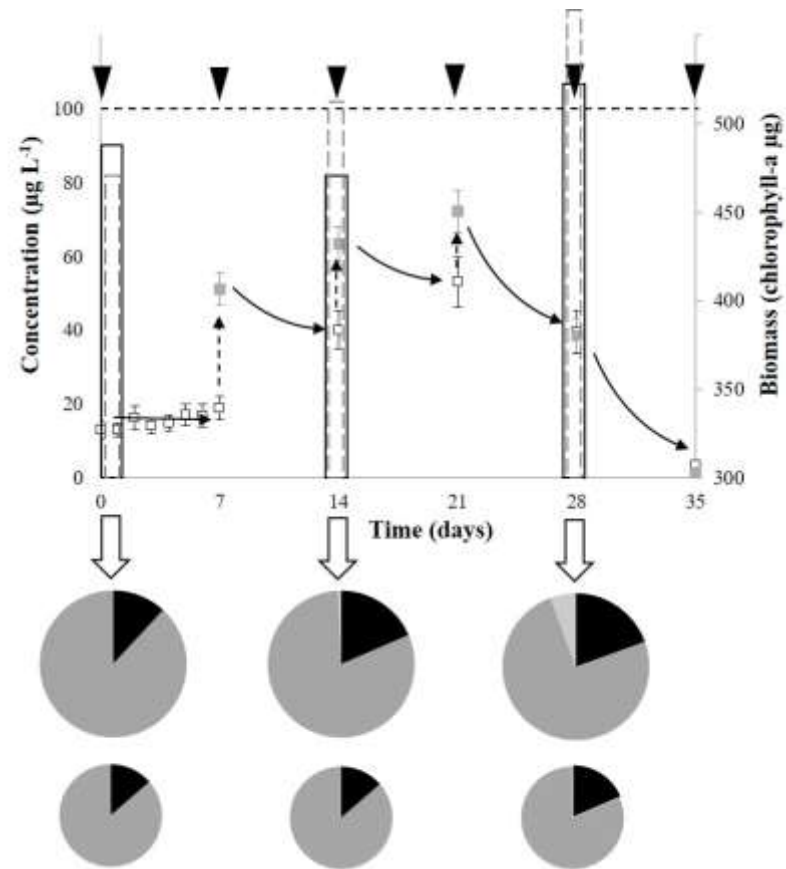
712



713

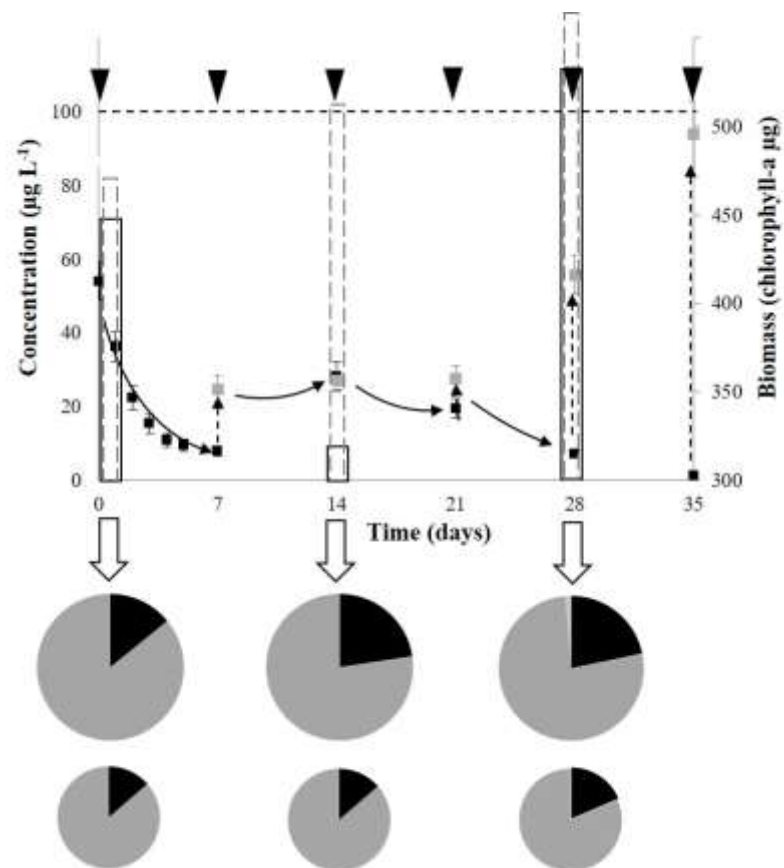
714

715 Figure 4



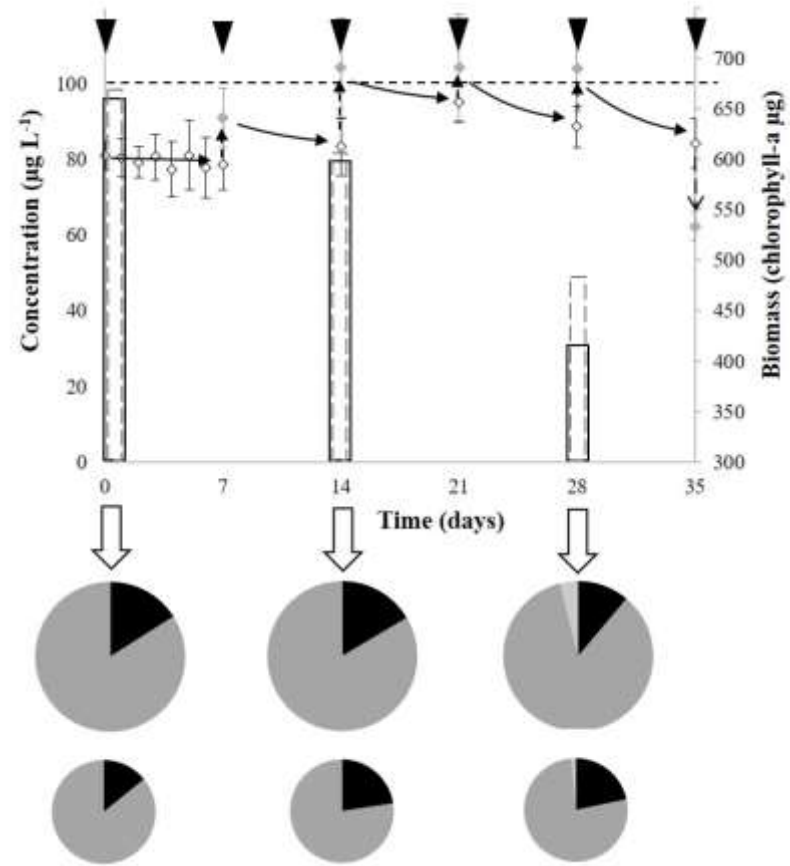
716
717
718

719 Figure 5
720



721
722

723 Figure 6



724
725
726

727 Figure 7

