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Study of the influence of pulsed electric field pre-treatment on quality parameters of sea bass during brine salting

This is the final peer-reviewed author's accepted manuscript (postprint) of the following publication:

*Published Version:*

Study of the influence of pulsed electric field pre-treatment on quality parameters of sea bass during brine salting / Crototova J.; Tappi S.; Genovese J.; Rocculi P.; Laghi L.; Dalla Rosa M.; Rustad T.. - In: INNOVATIVE FOOD SCIENCE & EMERGING TECHNOLOGIES. - ISSN 1466-8564. - ELETTRONICO. - 70:June 2021(2021), pp. 102706.1-102706.12. [10.1016/j.ifset.2021.102706]

*Availability:*

This version is available at: <https://hdl.handle.net/11585/828249> since: 2021-07-16

*Published:*

DOI: <http://doi.org/10.1016/j.ifset.2021.102706>

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This is the final peer-reviewed accepted manuscript of:

Janna Crotova, Silvia Tappi, Jessica Genovese, Pietro Rocculi, Luca Laghi, Marco Dalla Rosa, Turid Rustad,

*Study of the influence of pulsed electric field pre-treatment on quality parameters of sea bass during brine salting,*

Innovative Food Science & Emerging Technologies, Volume 70, 2021, 102706,

ISSN 1466-8564

The final published version is available online at:

<https://doi.org/10.1016/j.ifset.2021.102706>

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# Innovative Food Science and Emerging Technologies

## Study of the influence of pulsed electric field pre-treatment on quality parameters of sea bass during brine salting

--Manuscript Draft--

<b>Manuscript Number:</b>	IFSET_2020_230R1
<b>Article Type:</b>	PEF Treatment – R & D
<b>Keywords:</b>	LF-NMR; Pulsed electric field; sea bass; brine salting; Water distribution
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<b>Abstract:</b>	<p>Pulsed electric field (PEF), as an emerging technique, has recently gained increased popularity in food processing and preservation. However, applications in the seafood industry are still scarce. In the present study, sea bass samples were subjected to PEF pre-treatment prior to brine salting to verify the possible acceleration of the brining rate, increasing the salt uptake and ensuring the homogeneous salt distribution in the muscle. The applied intensity of the current was set at 10 and 20 A (corresponding to a field strength of 0.3 and 0.6 kV/cm) prior to sea bass salting in brine with 5 and 10% salt concentration, respectively. The results have shown that PEF pretreatment could effectively shorten the brine salting time compared to control samples (from 5 to 2 days), or increase the salt uptake up to 77%, ensuring at the same time its homogenous distribution in the muscle. However, myofibrillar protein solubility was significantly reduced in PEF pretreated samples. At the same time, no significant differences in water holding capacity and water activity between PEF pre-treated and untreated samples were found during the whole salting period. Freezable water was influenced by PEF application, but the effect was significant only at the lowest salt concentration during the first period of the salting process. Industrial relevance: PEF-assisted brining appears a promising technology in the fish processing industry due to its efficacy in reducing the salt brining time, increasing the mass transfer and enhancing the diffusion of brine into the muscle to ensure the homogeneous distribution of salt in it. The increased salt uptake of the PEF-treated samples compared to control samples shows future potentiality of using PEF prior to salting in the fish processing industry.</p>
<b>Response to Reviewers:</b>	

## **Highlights**

- PEF pre-treatment allowed to shorten brining times in sea bass fillets
- NaCl uptake was increased in seabass fillets compared to untreated samples
- Water state and distribution was only slightly affected by PEF treatment
- Reduction of myofibrillar protein solubility during brining was observed

1 **Study of the influence of pulsed electric field pre-treatment on quality**  
2 **parameters of sea bass during brine salting**

3

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29 **Abstract**

30 Pulsed electric field (PEF), as an emerging technique, has recently gained increased popularity in  
31 food processing and preservation. However, applications in the seafood industry are still scarce.  
32 In the present study, sea bass samples were subjected to PEF pre-treatment prior to brine salting  
33 to verify the possible acceleration of the brining rate, increasing the salt uptake and ensuring the  
34 homogeneous salt distribution in the muscle. The applied intensity of the current was set at 10 and  
35 20 A (corresponding to a field strength of 0.3 and 0.6 kV/cm) prior to sea bass salting in brine with  
36 5 and 10% salt concentration, respectively. The results have shown that PEF pretreatment could  
37 effectively shorten the brine salting time compared to control samples (from 5 to 2 days), or  
38 increase the salt uptake up to 77%, ensuring at the same time its homogenous distribution in the  
39 muscle. However, myofibrillar protein solubility was significantly reduced in PEF pretreated  
40 samples. At the same time, no significant differences in water holding capacity and water activity  
41 between PEF pre-treated and untreated samples were found during the whole salting period.  
42 Freezable water was influenced by PEF application, but the effect was significant only at the lowest  
43 salt concentration during the first period of the salting process.

44

45 **Industrial relevance:**

46 PEF-assisted brining appears a promising technology in the fish processing industry due to its  
47 efficacy in reducing the salt brining time, increasing the mass transfer and enhancing the diffusion  
48 of brine into the muscle to ensure the homogeneous distribution of salt in it. The increased salt  
49 uptake of the PEF-treated samples compared to control samples shows future potentiality of using  
50 PEF prior to salting in the fish processing industry.

51

52

53 **Keywords:** pulsed electric field, brine salting, sea bass, water distribution, LF-NMR

54

55

## 57 **1. Introduction**

58 Fish is a highly perishable raw material where deterioration caused by biochemical phenomena  
59 and microorganisms begin soon after slaughtering. Proper handling and preservation practices are  
60 therefore needed to prolong the shelf life of the product (Nagarajarao, 2016).

61 Salting is one of the oldest preservation methods used for long time storage of fish. Salted pelagic  
62 fish was well known to the old civilizations including the ancient Greeks and the Romans, the  
63 Vikings and other populations that lived on the shores of the Mediterranean Sea and the Atlantic  
64 Ocean. Today, a variety of salted pelagic fish products including sardines, anchovies, sea bass,  
65 *bacalao*, herring i.e., as well as Scandinavian dried and salted cod called *klippfisk*, literally "cliff-  
66 fish", are produced under the common name of "salted fish products" and marketed in many  
67 countries of the Mediterranean and the North Sea regions. Due to a fairly good market price and  
68 high palatability, these product commodities have become popular and highly appreciated in  
69 Europe and the USA. Along with the changes of lifestyle and growing consumer demands towards  
70 ready-to-eat, healthy and tasty foods, lightly salted fish products are currently gaining more and  
71 more popularity (Fan, Luo, Yin, Bao, & Feng, 2014).

72 Salting is one of the simplest methods of preserving large quantities of fish from spoilage. Salt is  
73 usually used at concentrations high enough to preserve the fish. Salting can be also used as a  
74 preliminary operation in smoking, drying and cooking processes helping to improve sensory  
75 parameters and increase the shelf-life of the final product (Bras & Costa, 2010). Salt can interact  
76 with proteins to increase hydration and water holding capacity of fish muscle thus improving its  
77 textural parameters. Increasing the water holding capacity of fish muscle helps to decrease cooking  
78 loss, thereby enhancing the tenderness and juiciness of the final product. Sodium chloride (NaCl),  
79 the common salt, is the main ingredient used in fish salting. It acts as a preservative by dehydration  
80 and osmotic pressure inhibiting bacterial growth and deactivating enzymes. Even at low  
81 concentrations, NaCl possesses some preservative action (Lupín, Boeri, & Moscardar, 1981). Other  
82 substances such as herbs, spices, sugar or antioxidants can also be used in the fish salting process  
83 to improve sensory attributes of the product, modify flavor and reduce shrinkage after salting. The  
84 conventional fish salting methods include dry-salting and wet-salting. During dry salting, the salt  
85 (traditionally sodium chloride) and other ingredients from the curing mixture (sugars and spices)

86 are applied to the fish surface. Wet salting is performed by immersing the product into brine or  
87 injecting the brine directly into the fish muscle (Birkeland, Skåra, Bjerkgeng, & Rørå, 2003; Hall,  
88 2011). The concentration of salt in the brine affects the weight gain, water holding capacity and  
89 commercial quality of the end product (Nguyen, Thorarinsdottir, Gudmundsdottir, Thorkelsson,  
90 & Arason, 2010). Weight gain of salted fish products depends on the ability of the myofibril  
91 proteins to retain water inside the muscle affected by the salting procedures applied  
92 (Thorarinsdottir, Arason, Sigurgisladottir, Valsdottir, & Tornberg, 2011). The brining time usually  
93 varies from 2 to 10 days depending on the desired level of salt in the muscle. During immersion  
94 brining, fish is covered with brine for a period of time and held at a temperature between 0 to 4°C.  
95 In injection salting, the brine is injected into the fish fillet using a set of needles making this a  
96 faster method than immersion brining.

97 Myofibrillar proteins are of great importance for the functional properties of light-salted fish  
98 products, such as water holding capacity (WHC). It is well known that salting of fish alters protein  
99 extractability and thermal denaturation and aggregation of many muscle proteins (Nguyen,  
100 Thorarinsdottir, Gudmundsdottir, Thorkelsson, & Arason, 2010), which in turn affects the WHC.  
101 Salting also affects the proteolytic activity responsible for degradation of myofibrils and  
102 connective tissue proteins, as well as extra-cellular matrix (Thorarinsdottir, Arason,  
103 Sigurgisladottir, Valsdottir, & Tornberg, 2011). Thus, the influence of salting on the distribution  
104 of water within the muscle may be related to direct effects of salt on changes in structural  
105 components of the muscle (Thorarinsdottir, Arason, Geirsdottir, Bogason, & Kristbergsson, 2002;  
106 Larsen & Elvevoll, 2008). It is also assumed that the main components of fish muscle (proteins,  
107 lipids and salts) influence the arrangement of water molecules in a product matrix, thereby having  
108 an effect on the product quality and shelf-life (Pacetti et al., 2015). Therefore, it is important to  
109 study how the salt content and water distribution within the muscle may affect water holding  
110 capacity of the product. Low-field nuclear magnetic resonance (LF-NMR) has been employed in  
111 the food industry to study water mobility and distribution within the fish muscle (Løje, Green-  
112 Petersen, Nielsen, Jørgensen, & Jensen, 2007; Aursand, Gallart-Jornet, Erikson, Axelson, &  
113 Rustad, 2008). This technique has been suggested a tool for rapid and non-destructive analysis of  
114 water mobility and identification of intra-myofibrillar or extra-myofibrillar water components  
115 (Andersen and Jørgensen, 2004; Jensen, Jørgensen, Nielsen, & Nielsen, 2005; Løje, Green-  
116 Petersen, Nielsen, Jørgensen, & Jensen, 2007) in the muscle.



117 The migration of salt from brine to fish matrix is generally quite slow. Different brining methods  
118 have previously been tested to accelerate salt transport through the product, for instance high  
119 intensity ultrasound brining and marinating (Chemat, Zill-e-Huma, & Khan, 2011; Turhan,  
120 Saricaoglu, & Oz, 2013), pulsed vacuum brining (Andres, Rodrigues-Barona, Barat, & Fito, 2002),  
121 and vacuum tumbling (Mathias, Jittinandanana, Kenney, & Kiser, 2003; Esaiassen et al., 2004).  
122 Pulsed electric field (PEF), as an emerging technology, has great potential to contribute to  
123 improved salting of fish products through enhanced diffusion of salt into the fish muscle  
124 (Hafsteinsson Gudmundsson Arnarson Jonsson, & Siguroardottir, 2000). However, to our  
125 knowledge, no studies have so far been published on PEF applications for salting of fish. Even  
126 though the concept of PEF was introduced to the food industry about 50 years ago, this technique  
127 can be still considered an emerging technology due to the recent developments related to microbial  
128 inactivation applications and improvement of mass transfer through cell disruption (Gómez et al.,  
129 2019). In general, PEF technique applies high voltage pulses of short duration to food placed  
130 between two electrodes, resulting in specific structural modifications of the tissue including the  
131 disruption of cell membrane (Barba et al., 2015). Under the application of the high electric field  
132 pulses, the membrane permeability is increasing due to either enlargement of existing pores or  
133 generation of new ones (Gómez et al., 2019). This concept was previously applied in the seafood  
134 industry with the aim of enhancing water holding capacity of fish and tenderization of shellfish  
135 products (Klonowski, Heinz, Toepfl, Gunnarsson, & Þorkelsson, 2006). PEF has also been  
136 suggested as a promising technique for accelerating mass transfer which could potentially be used  
137 as a pre-treatment in the fish drying process (Gómez et al., 2019).  
138 Therefore, the main aim of the present study was to investigate whether the PEF pre-treatment can  
139 be applied to accelerate the brining process and ensure a uniform distribution of salt within the  
140 muscle of fish, evaluating mass transfer kinetic and, in parallel, water state and distribution. The  
141 study aims also at investigating the effect of PEF pre-treatment on quality parameters of sea bass  
142 during salting. It is well known that PEF may affect the extractability and aggregation of proteins,  
143 since electroporation within the muscle tissue can result in chemical modifications by the  
144 formation of free radicals which can further alter the structure of proteins and the intermolecular  
145 forces (Gudmundsson & Hafsteinsson, 2001; Zhao, Sun & Tiwari, 2019). Therefore, this research  
146 also investigated the effect of different PEF pre-treatments on protein functionality by evaluating  
147 water holding capacity and protein solubility.

148

## 149 **2. Materials and Methods**

150

### 151 **2.1. Materials**

152 Sea bass (*Dicentrarchus labrax*) were supplied by Tagliapietra e Figli s.r.l. (Venice, Italy) in May  
153 2019. The day after catch, the fish were delivered to Economia del Mare (Cesenatico, Italy) where  
154 they were gutted, filleted and de-skinned. The sea bass fillets were placed on ice in Styrofoam  
155 boxes and transported to the CIRI-Agrifood laboratory in Cesena (Italy), where the experiment  
156 was carried out **in the same day**. Commercial salt ‘Sale alimentare di Sicilia’ from Italkali s.r.l.  
157 (NaCl ~98%) was used for brines preparation.

158

### 159 **2.2. PEF pre-treatment and brine salting**

160 Sea bass fillets were cut into small pieces ( $8.3 \pm 0.2$  g each) with the dimensions of length  $2.3 \pm$   
161  $0.2$  cm, width  $3.1 \pm 0.4$  cm and height  $1.3 \pm 0.5$  cm.

162 Prior to salting, the obtained sea bass pieces were subjected to PEF pre-treatment, performed using  
163 a lab scale PEF unit Mod. S-P7500 delivering a maximum output current and voltage of 60A and  
164 8kV, respectively (Alintel, Bologna, Italy). The generator provides monopolar rectangular-shape  
165 pulses and adjustable pulse duration (5-20  $\mu$ s), pulse frequency (50-500 Hz) and total treatment  
166 time (1-600 s). The treatment chamber (50 mm length x 50 mm width x 50 mm height) consisted  
167 of two parallel stainless-steel electrodes (3 mm thick) with a 47 mm fixed gap. Output voltage and  
168 current were monitored using a PC-oscilloscope (Picoscope 2204a, Pico Technology, UK). Sea  
169 bass pieces were treated at room temperature in tap water delivering  $n = 1000$  pulses at fixed pulse  
170 width ( $10 \pm 1$   $\mu$ s), frequency (100 Hz), repetition time ( $10 \pm 1$  ms) and selecting two different  
171 current intensities, 10A and 20A, corresponding to values of electric field strengths of 0.3 and 0.6  
172 kV/cm **and specific energy input of  $0.25 \pm 0.01$  and  $1.01 \pm 0.03$  kJ/kg**, respectively. The process  
173 parameters were chosen on the basis of preliminary experimental trials.

174 The sea bass pieces were randomly distributed into the three experimental groups (two PEF-treated  
175 and one control samples) and salted by immersion into a brine with two different salt (NaCl)  
176 concentrations in tap water (5% and 10% (w/w)) and in closed plastic containers (500ml) each  
177 containing a ratio of 4 to 1 w/w brine/fish. Five independent replicates were considered for each

178 sample type and for each sampling time. The salting process was carried out in a cold room at 0-  
179 4°C for 2, 5 and 8 days according to the experimental plan displayed in **Table 1**.

180 At each sampling day, sea bass samples were randomly collected and analyzed. Changes in weight  
181 yield, water-holding capacity, water activity, freezable water by differential scanning calorimetry  
182 and water behavior and distribution inside the muscle by LF-NMR as affected by different PEF  
183 pre-treatment and salting parameters, were studied directly after each sampling day at the  
184 laboratories of the University of Bologna (Cesena, Italy). The remaining experimental samples  
185 from each treatment were frozen at -80°C and transported to Norwegian University of Science and  
186 Technology (Trondheim, Norway) for determination of water and salt content, pH and protein  
187 solubility.

188 Analyses were performed in 3-6 replicates for each sample as described in detail in the following  
189 section.

190

## 191 **2.3. Physico-chemical analyses**

### 192 *2.3.1 Mass transfer parameters*

#### 193 *Weight yield*

194 The fish samples were weighed raw and after each sampling day. The weight yield was determined  
195 with respect to the weight of the raw fillets as described by Thorarinsdottir, Arason, Bogason, &  
196 Kristbergsson (2004).

197

#### 198 *Water content*

199 Water content was determined by drying a sample of 2 g at 105 °C for 24 h to a constant weight,  
200 according to the official method (AOAC 2005). Finely chopped fish obtained from 5 individual  
201 pieces was mixed and analysed in triplicate.

202

#### 203 *Salt (NaCl) content*

204 Salt content in all sea bass samples was determined by titration according to AOAC 976.18 (1995).  
205 Briefly, the fish obtained by 5 different pieces was minced with a kitchen blender (Bosch 600W,  
206 Gerlingen, Germany), and 2 g of the resulting mince was weighed in a 150 ml glass beaker, filled  
207 with 80 ml warm distilled water (60°C) and mixed for 5 min until a homogeneous mixture was  
208 obtained. Then, 1 ml of 1M HNO<sub>3</sub> was added to the mixture, the electrode type AgCl 32 and

209 burette tip was placed in the solution, and the titration was performed with an automatic titrator  
210 (mod. TitroLine 7800, Xylem Analytics, Mainz, Germany). The analysis was performed in three  
211 replicates and the results were expressed in % salt as a mean value  $\pm$  SD.

212  
213 The total water and NaCl weight changes ( $\Delta M_t^O$ ,  $\Delta M_t^W$  and  $\Delta M_t^{NaCl}$ , respectively) of salted samples  
214 were determined with Eqs (1), (2) and (3) as follow:

$$215 \quad \Delta M_t^O = \frac{(M_t^O - M_0^O)}{M_0^O} \quad (1)$$

$$216 \quad \Delta M_t^W = \frac{(M_t^O \cdot x_t^W - M_0^O \cdot x_0^W)}{M_0^O} \quad (2)$$

$$217 \quad \Delta M_t^{NaCl} = \frac{(M_t^O \cdot x_t^{NaCl} - M_0^O \cdot x_0^{NaCl})}{M_0^O} \quad (3)$$

218 where  $M_t^O$  and  $M_0^O$  are the sea bass weights,  $x_t^W$  and  $x_0^W$  are the water weight fractions, and  $x_t^{NaCl}$   
219 and  $x_0^{NaCl}$  are the NaCl weight fractions, at sampling time  $t$  and before the salting process  $0$ ,  
220 respectively.

### 221 222 **2.3.2 Water state and mobility**

#### 223 **Water activity**

224 Water activity was measured with a Water Activity Meter mod. AQUALAB, (Decagon Devices,  
225 US). Briefly, the fish samples were cut into small pieces (0.2 x 0.2 cm) and introduced into sample  
226 holders prior to the analysis. Between measurements, the samples were covered with lids and  
227 protected with parafilm. For each of the experimental groups, four measurements were performed  
228 and the mean value  $\pm$  SD was calculated.

#### 229 230 **Differential scanning calorimetry (DSC)**

231 A differential scanning calorimeter (DSC) mod. Q20 (TA Instrument, Germany), equipped with a  
232 low- temperature cooling unit (TA-Refrigerated Cooling System90.) was used to assess freezable  
233 water content (FW, g/g of water) and to evaluate the effect of processing on protein denaturation.  
234 Temperature and melting enthalpy calibrations were performed with ion exchanged distilled water  
235 (mp 0.0°C) and indium (mp 156.60°C), while heat flow was calibrated using the heat of fusion of  
236 indium ( $\Delta H = 28.71$  J/g). For the calibration, the same heating rate and dry nitrogen gas flux of 50  
237 ml/min used for the analysis were applied. Each sample was weighed (about 15 mg) into a 50- $\mu$ L  
238 aluminum pan, sealed hermetically and frozen at -40°C. Frozen samples were then loaded into the

239 DSC instrument. The heating rate of DSC scans was 5°C/min over a range of -40 to 90°C. Empty  
240 aluminum pans were used as reference and for baseline corrections. Eight replications for each  
241 sample were performed and results were elaborated through PeakFit Software (SeaSolve Software  
242 Inc. Framingham, MA, USA).

243 The FW was determined as follows:

244

$$245 \quad FW = \frac{\Delta H_m}{\Delta H_w} \quad (4)$$

246

247 where  $\Delta H_w$  (325 J/g) is the latent heat of melting per gram of pure water at 0°C, and  $\Delta H_m$  (J/g) is  
248 the measured latent heat of melting of water per gram of sample obtained by the integration of the  
249 melting endothermic peak. FW was further related to the water content and expressed as grams  
250 per gram of water content (FW<sup>w</sup>).

251 PeakFit Software (SeaSolve Software Inc. Framingham, MA, USA) was used to analyse thermal data and  
252 obtain deconvoluted peaks and calculate relative melting enthalpy.

253

#### 254 ***LF-NMR***

255 **A 10 mm deep slice was cut from each sample, then cylinders (6 mm diameter) of about 400 mg**  
256 **were obtained with a cork borer.** Signals weighted by T2 were registered with the CPMG pulse  
257 sequence (Meiboom & Gill, 1958), using a Bruker mod. Minispec PC/20 spectrometer operating  
258 at 20 MHz. Each measurement consisted in 30K points, spaced 0.080 ms. Subsequent scans were  
259 separated by a recycle delay of 3.5 s. The specified interpulse spacing avoided sample overheat  
260 but allowed the observation of the protons with T2 higher than a few milliseconds. UPEN software  
261 (Borgia, Brown, & Fantazzini, 1998) allowed to obtain an overview of the protons T2 distributions  
262 (the relaxograms) by inverting the T2-weighted signals towards a semi-continuous distribution of  
263 exponential curves, according to Eq. (5):

$$264 \quad I(2\tau n) = \sum_{i=1}^M I_0(T_{2,i}) \exp(-2\tau n/T_{2,i}) \quad (5)$$

265 where  $2\tau$  is the CPMG interpulse spacing,  $n$  is the index of each CPMG point while  $I_0$  is the  
266 intensities of each T2 component extrapolated at  $t = 0$ , sampled logarithmically. As some  
267 components resulted as partially overlapped in the relaxograms from several samples, we observed  
268 them separately by fitting the T2-weighted signals to the sum of an increasing number of

269 exponential curves. An F-test showed that the optimum ratio between fitting ability and complexity  
270 of the model was reached for most samples with three exponentials. Six measurements were  
271 performed for each of the experimental sets.

272

### 273 ***2.3.3 Protein functionality***

#### 274 ***pH***

275 pH was measured at room temperature by inserting electrode directly into the sea bass mince (mod.  
276 MP-220 pH-meter, Mettler-Toledo, Hong Kong) according to Thorarinsdottir, Arason, Bogason,  
277 & Kristbergsson (2004). Prior to pH measurements, the pH meter was calibrated with standard  
278 buffer solutions. The measurements were performed at least in triplicate, and the mean value  $\pm$  SD  
279 was calculated.

280

#### 281 ***Protein solubility***

282 Water and salt soluble proteins were determined in white muscle extracts according to a  
283 modification of the methods of Licciardello et al (1982), as previously described by Hultmann &  
284 Rustad (2002). The amount of proteins in the extracts was determined with BioRad protein assay  
285 after centrifugation at 8000 g and 4°C for 20 min, using gamma globulin as a standard. The  
286 analyses were run in triplicate and the mean value  $\pm$  SD was calculated.

287

#### 288 ***Water Holding Capacity (WHC)***

289 WHC of sea bass samples was measured according to the method described by Thorarinsdottir,  
290 Arason, Bogason, & Kristbergsson (2004), as follows. The minced samples were placed in  
291 centrifuge tubes and centrifuged at 200 g for 10 min (0–4 °C). The weight (g) of the fish pieces  
292 before and after the centrifugation was determined. WHC was expressed as the amount of released  
293 water divided by the original weight (g) of the sample before centrifugation. Four replicates were  
294 performed for each treatment group.

295

### 296 **2.4. Statistical analysis**

297 The data sets from the experiment were analyzed by Statistica 8.0 software (StatSoft, Tulsa, USA)  
298 **The effect of the parameters of PEF treatment (PEF), NaCl concentration (Salt) and brining time**  
299 **(Time) and their interaction on dependent variables was evaluated through the factorial Analysis**

300 of Variance (ANOVA). Statistical significance of the experimental data was verified using Tukey  
301 as post-hoc ( $p < 0.05$ ). To establish a relationship between certain parameters, Pearson correlations  
302 were calculated. Differences were considered significant at  $p < 0.05$ .

303

### 304 **3. Results and discussion**

305

#### 306 *3.1 Mass transfer parameters*

307 **Fig. 1** reports the total weight change (A), water (B) and salt uptake (C) mass fraction of control  
308 and PEF (0.3 and 0.6 kV/cm) treated sea bass samples during the brining process at 5% and 10%  
309 salt concentrations.

310 In control samples, weight increased between 24 and 26 % during the first 5 days of brining.  
311 However, on the last day of brine salting, the weight yield of control samples was reduced up to -  
312 0.13% and 2.56% for 5% and 10% salt concentration in the brine, respectively. The lowest weight  
313 yield in the control group on day eight may possibly be explained by an inhomogeneous salt  
314 distribution within the inner and outer parts of the fish muscle at the beginning of brining, leading  
315 to disintegration of the fish muscle pieces in the last part of the experiment, as previously showed  
316 by Thorarinsdottir, Arason, Bogason, & Kristbergsson (2004). Differently PEF treated samples  
317 showed a constant increase of weight during the entire brining period. While no significant  
318 differences were observed compared to the control until the 5<sup>th</sup> day of salting, on the 8<sup>th</sup> day all  
319 PEF treated samples (0.3 and 0.6 kV/cm) reached a weight gain of 28-32%.

320 The total water content in the sea bass samples varied from 73.9 to 88.7 % (w/w) during brine  
321 salting. In all samples, water uptake (**Fig. 1B**) was observed until the 5<sup>th</sup> day, when samples  
322 immersed in the 5% salt brine showed significantly higher values compared to samples in the 10%  
323 one. However, no differences were observed among the control and the PEF treated samples in  
324 each of the 2 groups (0.3 and 0.6 kV/cm). At the 8<sup>th</sup> day, the water uptake showed a drastic drop  
325 for both the control samples, as already observed with the total weight change. PEF treated samples  
326 in the 5% brine, did not show a further water uptake, while samples in the 10% brine showed a  
327 further increase. All PEF treated samples showed similar water fraction values at the end of the  
328 brining period.

329 Initial salt content of sea bass fillets was 0.01 g/100g. Salt weight fraction changes are reported in  
330 **Fig. 1C**. In control samples, an increase of salt content was observed until the 5<sup>th</sup> day, reaching

331 values of 0.03 and 0.07 that corresponded to 2.7 and 5.9 % of net salt content for the 5 and 10%  
332 brining respectively. Hence, as expected, the salt uptake was driven by concentration gradients  
333 between the muscle and brine, similarly to previous studies (Nguyen, Thorarinsdottir,  
334 Gudmundsdottir, Thorkelsson, & Arason, 2010). However, as observed for the weight and water  
335 uptake, on the last day of brining, the salt fraction decreased to values corresponding to 0.46 and  
336 2.05% for the 5 and 10% brining respectively.

337 Following PEF pre-treatment, there was a general increase of the salt uptake in all samples at the  
338 end of the salting process. After two days, both 10 and 20A PEF (0.3 and 0.6 kV/cm) treated  
339 samples were significantly higher compared to their respective controls, while after 5 days, only  
340 the 10A sample and the 20A sample in the 5% brine. Salt concentration in PEF treated sea bass  
341 fillets increased slightly between the 5<sup>th</sup> and the 8<sup>th</sup> day, but, although samples treated at 10A (0.3  
342 kV/cm) showed an increasing trend, differences were not statistically significant. The higher salt  
343 weight fractions reached corresponded to a salt content in the samples of 4.47 and 6.84 g/100g for  
344 the 5 and 10% brining respectively, showing an increase of 77 and 35% compared to the highest  
345 salt content obtained in control samples at day five.

346 Applying PEF pretreatment allowed to reach a similar salt uptake after 2 days of brining, instead  
347 of 5 days in the control samples, thus reducing the time necessary for the process.

348 PEF has previously been shown to increase mass transfer in other animal and vegetable foods,  
349 such as ham, cured and salted meat, potato crisps, dried fruits etc. (Gómez et al., 2019).  
350 Electroporation is one of the several complex mechanisms attributed to this phenomenon. It was  
351 previously assumed that a greater number of pores in the muscle emerges with increasing the  
352 electric field intensity, which is why generally a mass transfer increase is obtained (Gómez et al.,  
353 2019). Electroporation has been shown to cause increased inter-myofibrillar spacing in fish and  
354 meat products (Gómez et al., 2019) which could aid mass transfer, thus increasing the salt uptake  
355 by the muscle. Therefore, we suggest that in the present study electroporation facilitated the salt  
356 uptake by the fish through increasing the extra-cellular spaces in the muscle serving as additional  
357 channels for diffusion of brine. Moreover, Klonowski, Heinz, Toepfl, Gunnarsson & Porkelsson  
358 (2006) found a more porous structure in cod fillets pre-treated by PEF, that might have aided the  
359 diffusion of salt. Even though this effect was observed with the application of a higher electric  
360 field strength (2kV/cm) compared to the ones applied in this present research (0.3-0.6kV/cm), it is  
361 possible that a change on the flesh structure might have happened.



362 The increase of salt concentration in the tissue results, especially at the level of myofibrils, in  
363 greater water absorption and swelling under certain conditions (Krasnow, Loss, Ahrens, & Fiore  
364 III, 2013). This phenomenon is linked to the action of Cl<sup>-</sup> chloride anions, which tend to associate  
365 with the positively charged groups of proteins. Positive charges are neutralized and therefore the  
366 repulsive force of negative charges increases. The intra-myofibrillary space expands due to the  
367 repulsive forces and a greater water retention capacity is determined. However, brines with a saline  
368 concentration above 10-15% can lead to an opposite effect, worsening the water retention capacity.  
369 In this case the salting-out phenomenon may occur: the ions in excess of Cl<sup>-</sup>, not being able to  
370 interact with the positive charges of the proteins already occupied by the other ions, interfere with  
371 them for the interaction with the water molecules, sequestering the solvation water and causing the  
372 loss of solubility and the precipitation of proteins (Aberoumand e Nejad, 2015; Kalra, Tugcu,  
373 Cramer, & Garde, 2001; Offer e Trinick, 1983). This phenomenon, however, was not observed in  
374 PEF treated samples by Klonowski, Heinz, Toepfl, Gunnarsson & Porkelsson (2006), although the  
375 final salt concentration was higher.

376 We hypothesize that, contrarily to control samples, PEF treatment **in the range of 0.3 and 0.6**  
377 **kV/cm** promoted a more homogeneous distribution of NaCl within inner and outer parts on the  
378 fish muscle due to formation of small pores in the muscle, facilitating the mass transfer and leading  
379 to enhanced diffusion of salt from the brine to the muscle.

380

### 381 ***3.2 Water state and distribution***

382 **The** water activity ( $a_w$ ) of untreated sea bass samples was **0.990 ± 0.002**. As shown in **Fig. 2**, fish  
383 tissue brining resulted in a significant decrease of water activity, explained by the bonding of  
384 residual fluid from the fish muscle by salt through ionic interactions. These interactions reduce the  
385 amount of free water contained in the fish muscle, thus lowering water activity of the product  
386 (Lupín, Boeri, & Moscidar, 1981). Statistical analysis showed that only the NaCl concentration in  
387 the brine had a significant ( $p < 0.05$ ) influence on water activity of sea bass samples during salting,  
388 leading to values in the range of 0.966 to 0.972 and 0.941 to 0.949, during the salting period for  
389 the 5 and 10% concentration respectively. **Neither** PEF intensity **(0.3 and 0.6 kV/cm)** **nor** duration  
390 of brine salting did affect water activity of the fish samples.

391 According to different authors (da Silva Carneiro et al., 2016; Mudalal, Petracci, Tappi, Rocculi,  
392 & Cavani, 2014), there are three different water populations in muscle tissues, the first one (below

393 5%) exists as true hydration water that is strictly bound to proteins by macromolecular of  
394 multimolecular adsorption, the second is water located inside organized protein structures (intra-  
395 myofibrillar), and the third one, which is the major one (>70%), is the extra-myofibrillar water,  
396 easily mobilizable. The first one is not free; it has an ice-like structure (liquid crystal), it is  
397 unfreezable, unaffected by charges on the muscle protein (pH), and it is unavailable to participate  
398 in reactions. From a calorimetric point of view, freezable water (FW) is usually associated to the  
399 second two fractions, representing the water affected during processing. FW assessment by DSC  
400 has been used to determine the gross phase changes of water in polymeric networks (Capitani et  
401 al., 2003) and in food systems, such as meat (Venturi et al., 2007; Petracci et al., 2012; Mudalal,  
402 Petracci, Tappi, Rocculi, & Cavani, 2014).

403 **Fig. 3A** reports, as an example, the obtained thermograms of sample C10 at different brining times  
404 (zero to eight days). As it is possible to observe, the FW peak was actually composed by two  
405 superimposed peaks, melting at slightly different temperatures. While in the fresh sample, this  
406 difference was small, with the first melting at around  $-3^{\circ}\text{C}$  and the second melting at around  $0^{\circ}\text{C}$   
407 being almost indistinguishable, as the brining time increased, the first peak appeared at lower  
408 temperatures, until reaching  $-6^{\circ}\text{C}$  after 8 days. In order to better understand the phenomena, the  
409 total melting enthalpy of FW were calculated and the relative amount of the two peaks were  
410 plotted, as shown in **Fig. 3B** (example of raw thermogram) and **3C** (example of deconvoluted  
411 thermogram) respectively.

412 **Fig. 4** shows the total  $\text{FW}^{\text{w}}$  content, (**4A**), the fraction of peak 1 (**4B**) and the melting temperature  
413 of the first peak (**4C**). In the fresh sample, total  $\text{FW}^{\text{w}}$  content was 0.69 g/g water. In control samples  
414 immersed in the 5% NaCl brine, this value increased slightly after two days. However, the increase  
415 of salt concentration led to a decrease of the  $\text{FW}^{\text{w}}$  to the initial values. The first raise was probably  
416 due to a fast water uptake that increased the general mobility of the water. However, the  
417 simultaneous increase of salt concentration probably counterbalanced this effect. **However,**  
418 **differences were not significant.** In PEF treated samples, no differences were observed compared  
419 to initial value at all brining days.

420 For samples in the 10% NaCl brine, the total  $\text{FW}^{\text{w}}$  water content showed a slight decrease that was  
421 maintained during all brining time, but without significant differences among the samples. The  
422 water uptake, as shown in **Fig. 1A** was similar for the two salt concentrations (**Fig. 1B**). However,

423 samples in the 10% solution showed, as expected, a higher salt diffusion during brining (**Fig. 1C**),  
424 this is the reason for the lowering of FW<sup>w</sup>.  
425 Hence, it is possible to observe that the total FW<sup>w</sup> was fairly constant in all samples; however, if  
426 we take into account the two different peaks, it is possible to be observed that, while initially the  
427 majority of the water was melting at 0°C (about 80%), as brining proceeded, the fraction (peak 1)  
428 melting at lower temperature increased progressively. In samples in the 10% solution, the increase  
429 occurred after the first two days and then values remained similar (between 0.88 and 0.95), while  
430 for the 5% samples, the transition was more progressive. The decrease in FW<sup>w</sup> and melting  
431 temperature depends on the balance between the water uptake and the salt concentration in the  
432 tissue. Although at the end of the eighth day values were similar for all samples, control samples  
433 (C5) showed higher values for peak 1 after two and five days, showing a slower decrease of the  
434 melting temperature transition. As shown by **Fig. 1C**, in PEF treated samples, salt concentration  
435 increased more compared to the control, corroborating the hypothesis of the observed differences.  
436 Moreover, in **Fig. 4C** the melting temperature related to peak 1 was evaluated for all samples  
437 during brining. In the 5% samples the temperature did not change, while for the 10% samples a  
438 significant decrease was observed already after two days. Hence, DSC data were able to  
439 discriminate samples according to the concentration of salt in the brine showing a proportional  
440 reduction of freezable water and a decrease of the melting temperature due to the increasing salt  
441 content. However, few significant differences were observed among samples. **This was not**  
442 **expected since a higher amount of salt found in PEF treated samples compared to control at**  
443 **different brining times for both 5% and 10% samples.** Moreover, the effect of ‘salting out’  
444 observed in the control samples, was not reflected in the FW measurements. **This might be due to**  
445 **a different distribution of salt in the tissue as hypothesized earlier. Indeed, sampling procedure is**  
446 **pivotal for DSC analysis, since the small sample size (about 15 mg). Hence, although we took**  
447 **extra care in collecting representative samples, this could be one of the reasons for the observed**  
448 **unexpected behavior. However, considering that,** to our knowledge, there are no reports of FW<sup>w</sup>  
449 measure by DSC in fish samples during brining, so it is not possible to compare results giving a  
450 more exhaustive explanation of the obtained results.  
451 Low-resolution NMR has been successfully used in many previous studies to investigate water  
452 mobility and distribution in fish and meat samples subjected to salting (da Silva Carneiro et al.,  
453 2016; Gudjónsdóttir, Arason, & Rustad, 2011; Aursand, Gallart-Jornet, Erikson, Axelson, &

454 Rustad, 2008; Wu et al., 2006). As in previous studies, in the present research it was possible to  
455 reveal the presence of 3 water populations (displayed in **Fig. 5**), characterized by short, medium  
456 and long proton relaxation times.  $W_B$  ( $T_2=1-3$  ms) relates to water bound by secondary bonds to  
457 the proteins,  $W_1$  ( $T_2=40-80$  ms) describes capillary water found in the myofibrillar network, while  
458  $W_2$  ( $T_2=100-190$  ms) is mechanically immobilized water or extra-myofibrillar which can be further  
459 released as drip loss. Table 2 reports the relative intensities expressed as arbitrary units (AU) and  
460 the  $T_2$  of the three water populations for all the analyzed seabass samples. According to Aursand,  
461 Gallart-Jornet, Erikson, Axelson, & Rustad (2008) populations  $W_1$  and  $W_2$  represent more than  
462 90% of the total water in the muscle.

463 In the present study, an evident migration of water from pools  $W_B$  and  $W_1$  towards pool  $W_2$ , with  
464 longer relaxation times was observed from the untreated raw sample to all brined samples. This  
465 indicates a migration of water from the myofibrillar network towards extra-myofibrillar pools.  
466 Indeed, NaCl not only has a preservation effect, but it also acts as a structures-breaker, allowing  
467 the muscle fibers to expand and entrap water. This occurs due to electrostatic repulsion within the  
468 myofibrils, exposing protein sidechains to water binding (Strasburg, Xiong & Chiang, 017).  
469 Similar results were found in the study of Aursand, Gallart-Jornet, Erikson, Axelson, & Rustad  
470 (2008) investigating water distribution and behavior in brine salted cod and salmon by low-field  
471 NMR technique. However, in the present research, apart from a few exceptions, no significant  
472 differences were observed among samples, neither according to NaCl concentration, nor according  
473 to the treatment. **The only variable that showed consistently a significant effect on water**  
474 **distribution parameters was brining time ( $p<0.001$ ).**

475 With regard to relaxation times (**Table 2**), Wu et al (2006) found a decrease for the bound water  
476 ( $T_{2B}$ ) and an increase related to  $T_{21}$  and  $T_{22}$  populations during salting of pork meat. In the present  
477 research  $T_{2B}$  showed a decrease but the difference was not significant. Instead, salting in 5% and  
478 10% NaCl brine, led to a shift toward longer relaxation times for the other two water populations.  
479  $T_{21}$  (intra-myofibrillar water) shifted from about 45 ms to 65-85 ms, while  $T_{22}$  (extra-myofibrillar  
480 water) from about 106 ms to 130-190 ms, directly reflecting the increased amount of water, which  
481 was also observed in other studies conducted on brine salting of fish (Aursand, Gallart-Jornet,  
482 Erikson, Axelson, & Rustad, 2008). However, also for this parameter, few significant differences  
483 were observed. Specifically, while in  $T_{22}$  a significant increase was found during brining time, no

484 differences were observed among samples according to the PEF treatment (0.3 and 0.6 kV/cm). A  
485 significant effect was found only for brining time and for NaCl concentration for T<sub>21</sub> and T<sub>22</sub>.

486

### 487 *3.3. Protein functionality*

488 The pH values of sea bass samples after PEF-treatment and salting performed for 2, 5 and 8 days  
489 are shown in **Table 3**. Untreated sample showed an initial value of 6.7 that decreased progressively  
490 during brining, but the only significant differences was observed for C10 after 8 days (pH= 6.18).

491 The results of PEF treated samples (0.3 and 0.6 kV/cm) have shown significantly lower pH values  
492 compared to control samples on day 2 and 5 of brining. This could be due to a release of ions from  
493 PEF-disrupted cells or structural changes of proteins allowing release of acidic groups (Zhao, Sun,  
494 & Tiwari, 2019). Values, however, did not change during brining but apart from the initial  
495 decrease, remained stable. Nevertheless, result of multifactorial ANOVA showed that this  
496 parameters is influences significantly by all considered variables and their interaction.

497 WHC of sea bass samples (**Table 3**) showed very small variations remaining in the range of 97.7  
498 to 98.99%. In some samples, a slight but not always significant increase of WHC appeared. This  
499 may have been due to the increased salt concentration as observed by Thorarinsdottir, Arason,  
500 Bogason, & Kristbergsson (2004) and Aursand et al (2008). However, no significant effect of PEF  
501 pre-treatment (0.3 and 0.6 kV/cm) or of salt concentration on WHC during salting period was  
502 observed in the present study. The only variable affecting WHC was indeed brining time and its  
503 interaction with other variables.

504 The solubility of sarcoplasmic and myofibrillar proteins in sea bass samples during brine salting  
505 is reported in **Fig. 6 A and B**.

506 Solubility of water soluble (sarcoplasmic) protein was strongly and significantly reduced during  
507 brining in all samples. In seabass brined in the 10% NaCl solution, PEF treated samples showed  
508 always significantly lower values compared to the control, but with no differences according to  
509 the intensity of the electric field applied, 0.3 or 0.6 kV/cm. For samples in the 5% brine solution,  
510 differences were not always significant.

511 Solubility of salt-soluble (myofibrillar) proteins showed a very different behavior. In control  
512 samples, it did not change compared to the initial untreated sample for all brining times. Instead,  
513 PEF treated samples reported a remarkable decrease already after 2 days for both 0.3 and 0.6

514 **kV/cm treated samples**. However, there were no differences in the values found between salt  
515 concentration and during brining.

516

### 517 **3.4 Correlation results**

518 In order to get a better understanding on the observed phenomena and of their relation, correlations  
519 among the parameters of mass transfer, water mobility and distribution, and protein functionality  
520 measured in the sea bass samples were evaluated through the Pearson's correlation. Results are  
521 shown in **Table 4**.

522  $\Delta M^o_t$  is positively correlated to both  $\Delta M^w_t$  and  $\Delta M^{NaCl}_t$ , as they showed similar behavior during  
523 brining, but it was also negatively correlated to  $W_B$  and to the solubility of both water- and salt-  
524 soluble proteins. No significant correlation was observed with any of the other parameters, that, as  
525 observed before, did not reflect the effect of salting out.

526 Water activity and total FW were positively correlated (0.64), however, the evolution of peak 1 of  
527 FW (water fraction freezing at a lower temperature) was actually correlated to all the other water  
528 state and mobility parameters, measured by LF-NMR and solubility of water-soluble proteins.

529 Specifically, the solubility of myofibrillar proteins positively correlated with  $W_B$ -water pool  
530 expressing water bound by secondary bonds to the proteins in PEF-treated samples, while the  
531 solubility of sarcoplasmic proteins negatively correlated with  $W_2$ -water pool representing  
532 mechanically immobilized water. This suggests that the water pool  $W_B$  diffused to the extra-  
533 myofibrillar spaces of the fish muscle ( $W_2$ -water pool) as a result of the PEF-induced increased  
534 solvation. Supported by previous investigations (Nguyen, Thorarinsdottir, Gudmundsdottir,  
535 Thorkelsson, & Arason, 2010), this could be caused by the reduced hydration due to the increased  
536 solvation capacity of salt ions that reduced the hydrodynamic radius of proteins, increasing  
537 substantially protein-protein interactions compared to protein-water interactions. The weaker  
538 associations between the water molecules bound to proteins resulted in their increased mobility  
539 and penetration into extra-myofibrillar spaces of the muscle. At the same time, polar and  
540 hydrophobic interactions between proteins became stronger, contributing to their increased  
541 hydrophobicity and aggregation (Stefansson & Hultin, 1994; Lin & Park, 1998).

542

## 543 **4. Conclusions**

544 The results of this study have shown that PEF treatment at 0.3-0.6 kV/cm allowed to significantly  
545 increase the salt uptake during sea bass brining, that may be due to a more homogeneous  
546 distribution of salt in the fish muscle. The study of water state and distribution however did not  
547 show many differences among samples that were generally discriminated according to the  
548 concentration of salt in the brining solution but not to the PEF treatment applied. On the other side,  
549 a remarkable reduction of myofibrillar protein solubility was observed, as a consequence of the  
550 application of the electric field.

551 To sum up, the obtained results suggest that PEF pre-treatment allowed to obtain a significant  
552 reduction of the duration of salt brining (more than 50%) or an increase of salt uptake (up to 77%)  
553 compared to conventional brining process. However, aspects related to the effect on protein  
554 structure and functionality should be further clarified, and different parameters of this innovative  
555 processing deeply investigated.

556

## 557 **5. Acknowledgments**

558 Janna Crobotova gratefully acknowledges the financial support provided by the *International*  
559 *Research Mobility Support* offered as part of NTNU Postdoc Action Pilot Programme to conduct  
560 the displayed study at University of Bologna.

561 Pietro Rocculi and Silvia Tappi acknowledge the financial support of EU project FuturEUAqua  
562 H2020-BG-2018-2020 (Blue Growth).

563

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703

704 **Figure captions**

705

706 **Figure 1.** Total weight change ( $\Delta M^0_t$ ) (A), water uptake ( $\Delta M^w_t$ ) (B) and NaCl uptake ( $\Delta M^{NaCl}_t$ )  
707 (C) of control and PEF treated sea bass samples during the brining process at 5% and 10% salt  
708 concentrations. Results are expressed as means  $\pm$  standard deviations (error bars) of n=5. Values  
709 with different letters in the auxiliary tables differ significantly (p<0.05).

710

711 **Figure 2.** Water activity of control and PEF treated sea bass samples during the brining process at  
712 5% and 10% salt concentrations. Results are expressed as means  $\pm$  standard deviations (error bars)  
713 of n=4. Values with different letters in the auxiliary table differ significantly (p<0.05).

714

715 **Figure 3.** Example of (A) the obtained thermograms for sample C10 at different brining times (0  
716 to 8 days), (B) of a raw thermogram and (C) of a deconvoluted thermogram related to freezable  
717 water (FW<sup>w</sup>).

718

719 **Figure 4.** DSC data of (A) freezable water (FW<sup>w</sup>) content, (B) fraction of the first peak composing  
720 FW and (C) melting temperature of water of control and PEF treated sea bass samples during the  
721 brining process at 5% and 10% salt concentrations. Results are expressed as means  $\pm$  standard  
722 deviations (error bars) of n=8. Values with different letters in the auxiliary tables differ  
723 significantly (p<0.05).

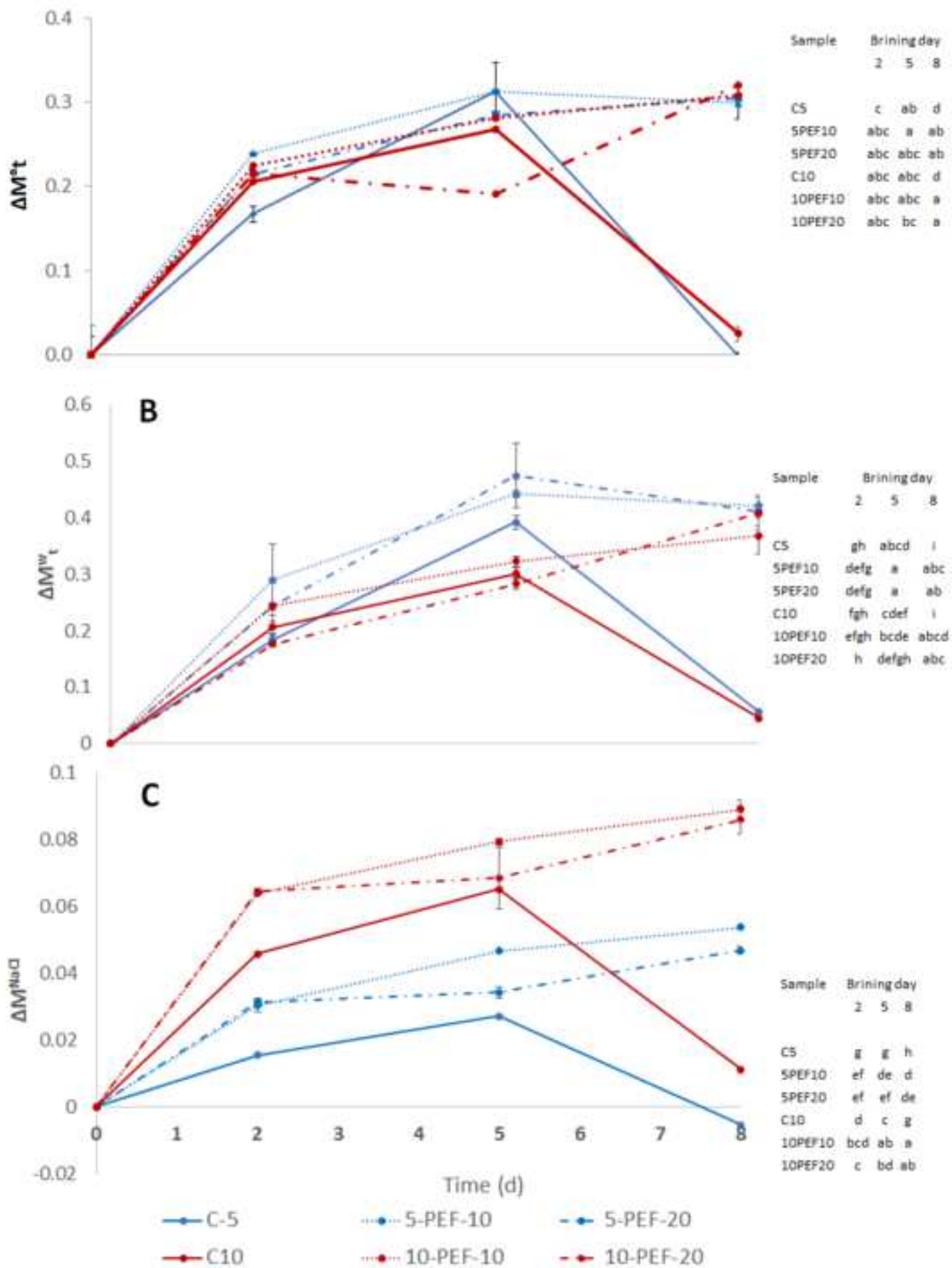
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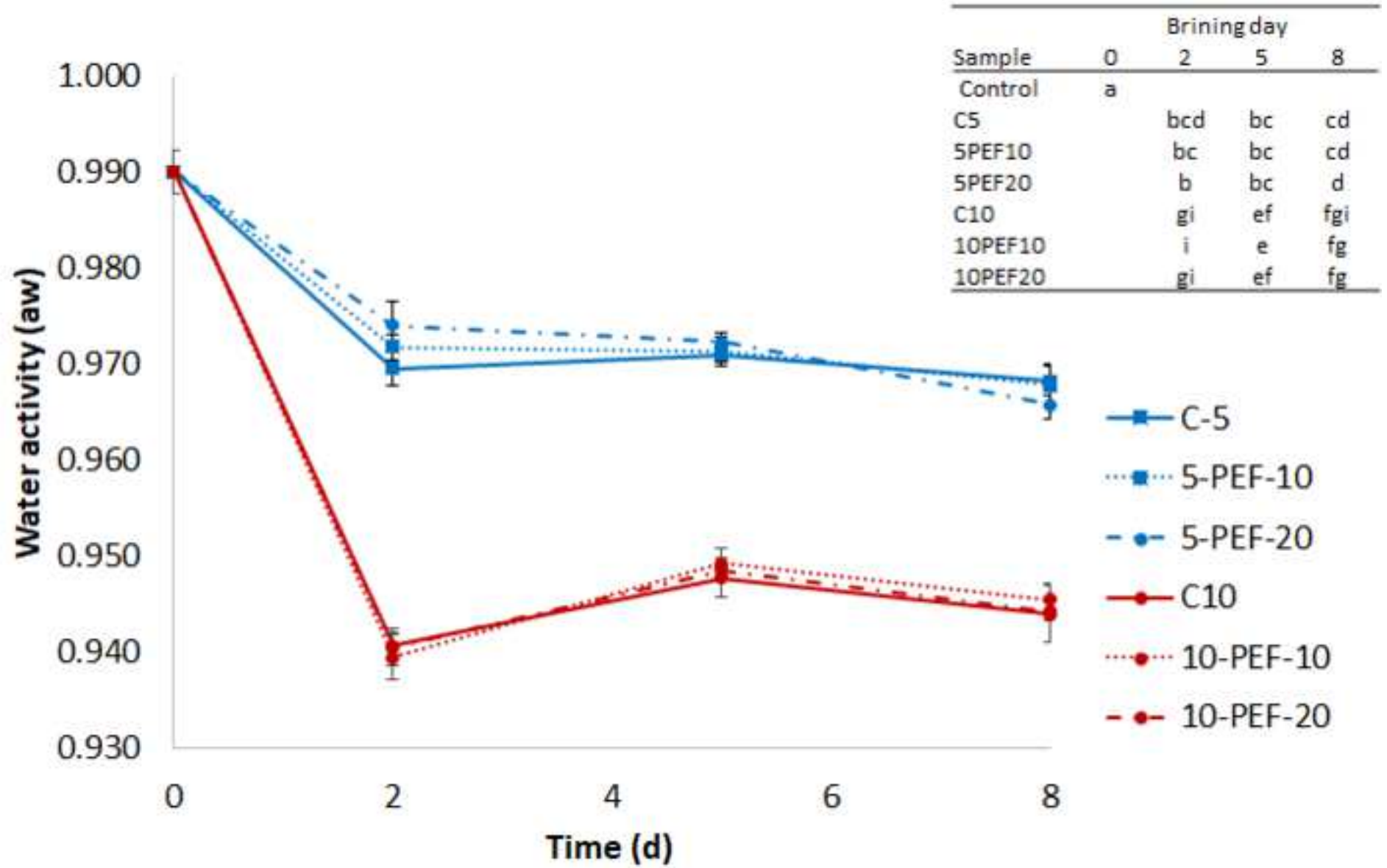
725 **Figure 5.** Three typical transverse relaxation time relaxograms ( $T_2$ ) obtained on a control sample  
726 at day 0 (dashed black line) and at day 8 (solid black line) and on sample salted in 10% brine and  
727 treated at 10 A (solid gray line). To allow for a direct comparison among them, the intensities are  
728 scaled so that the total area equals one arbitrary unit.

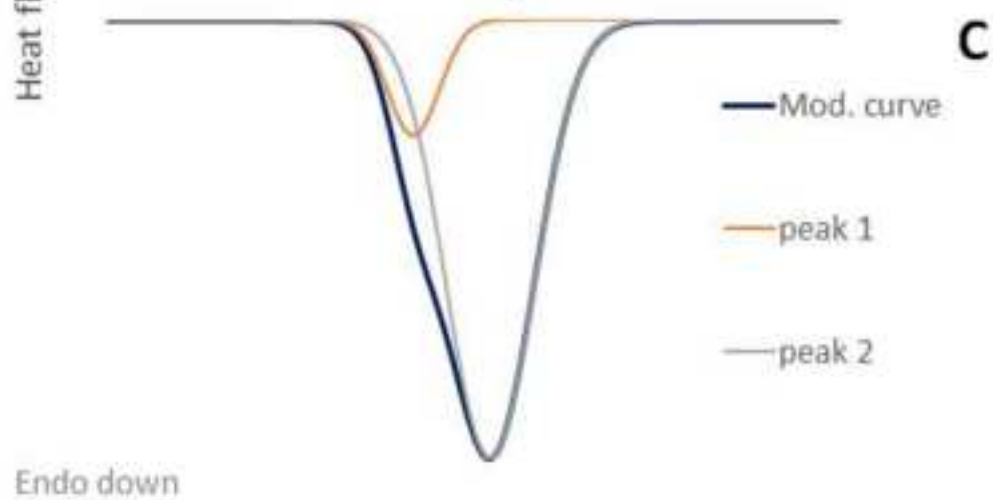
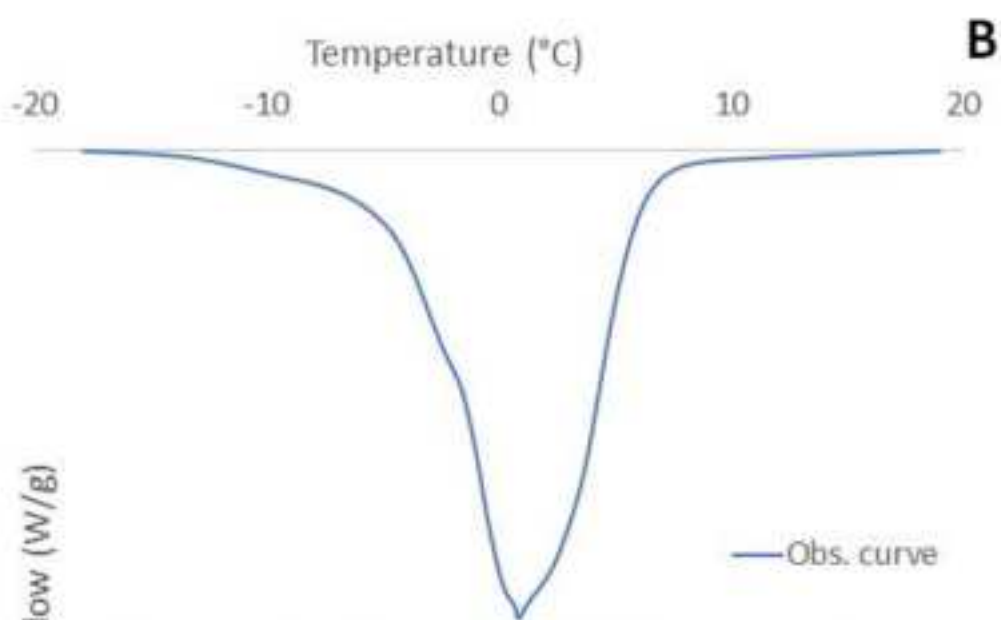
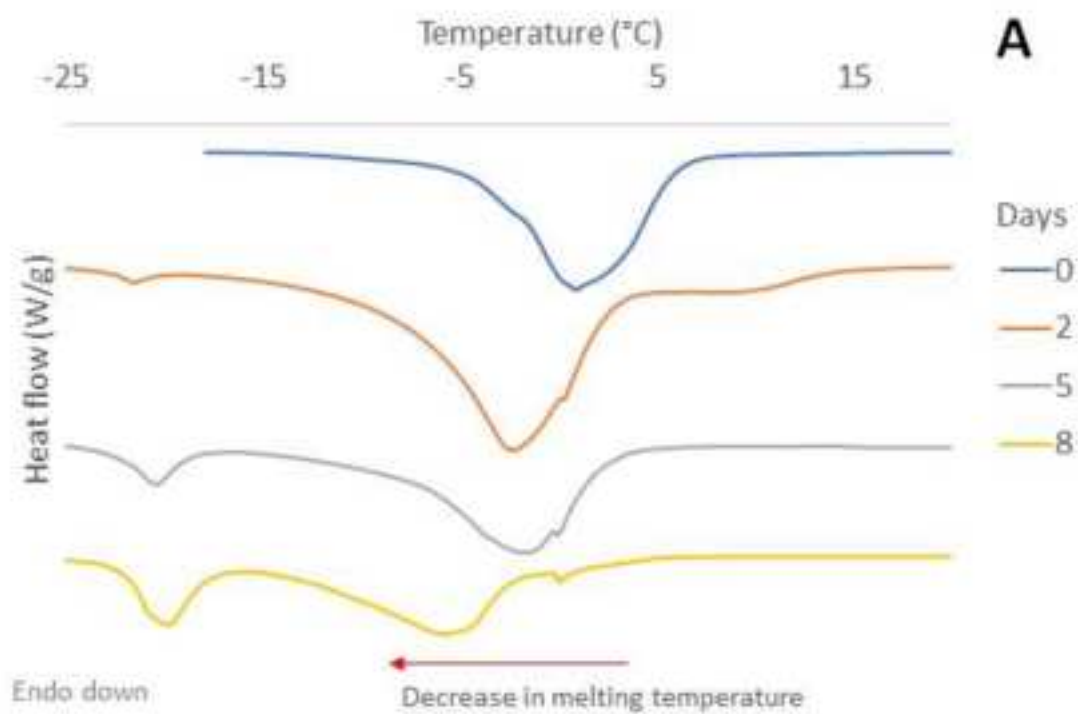
729

730 **Figure 6.** Content of (A) water- and (B) salt-soluble proteins (% net weight) of control and PEF  
731 treated sea bass samples during the brining process at 5% and 10% salt concentrations. Results are  
732 expressed as means  $\pm$  standard deviations (error bars) of n=3. Values with different letters in the  
733 auxiliary tables differ significantly (p<0.05).

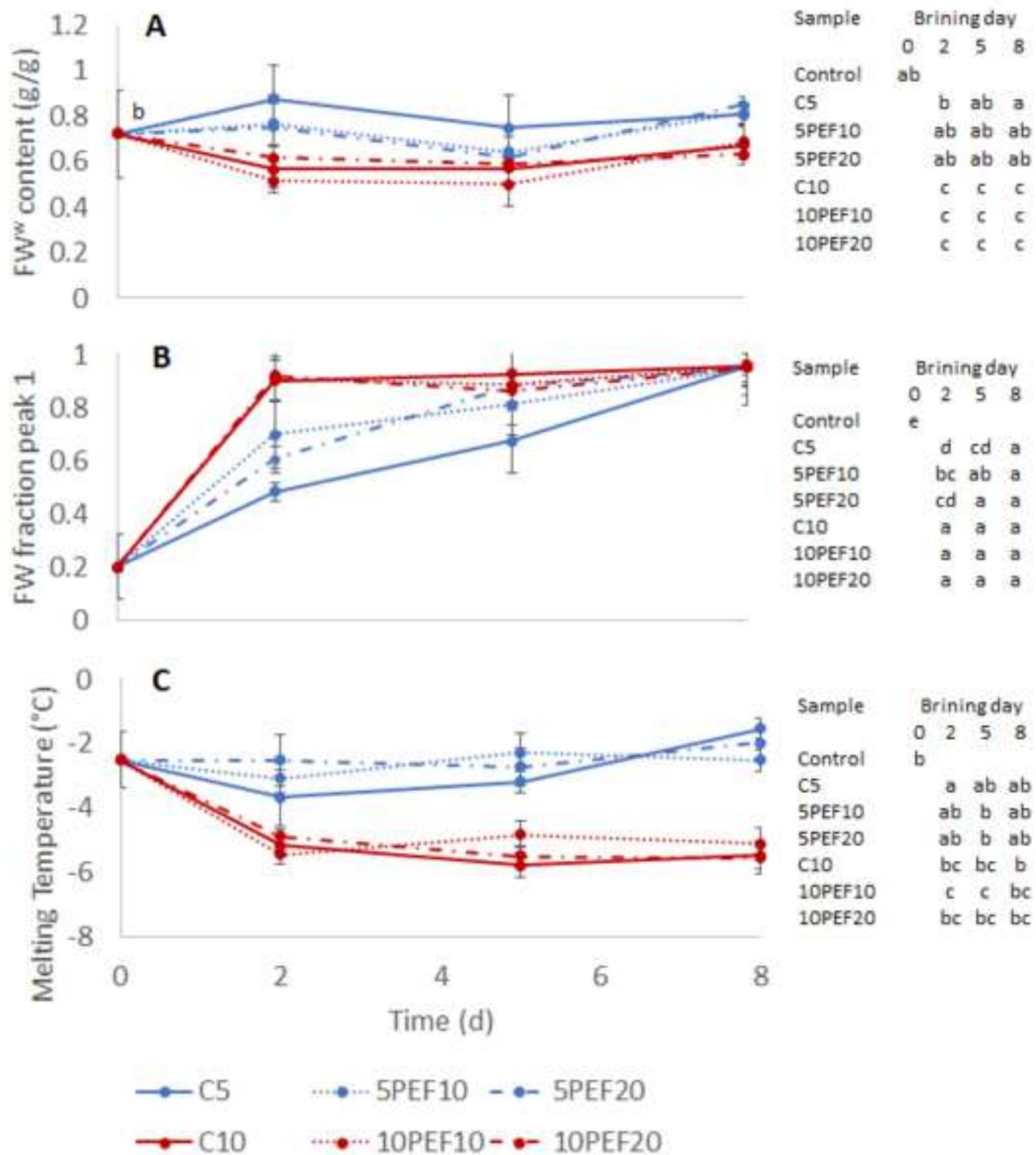
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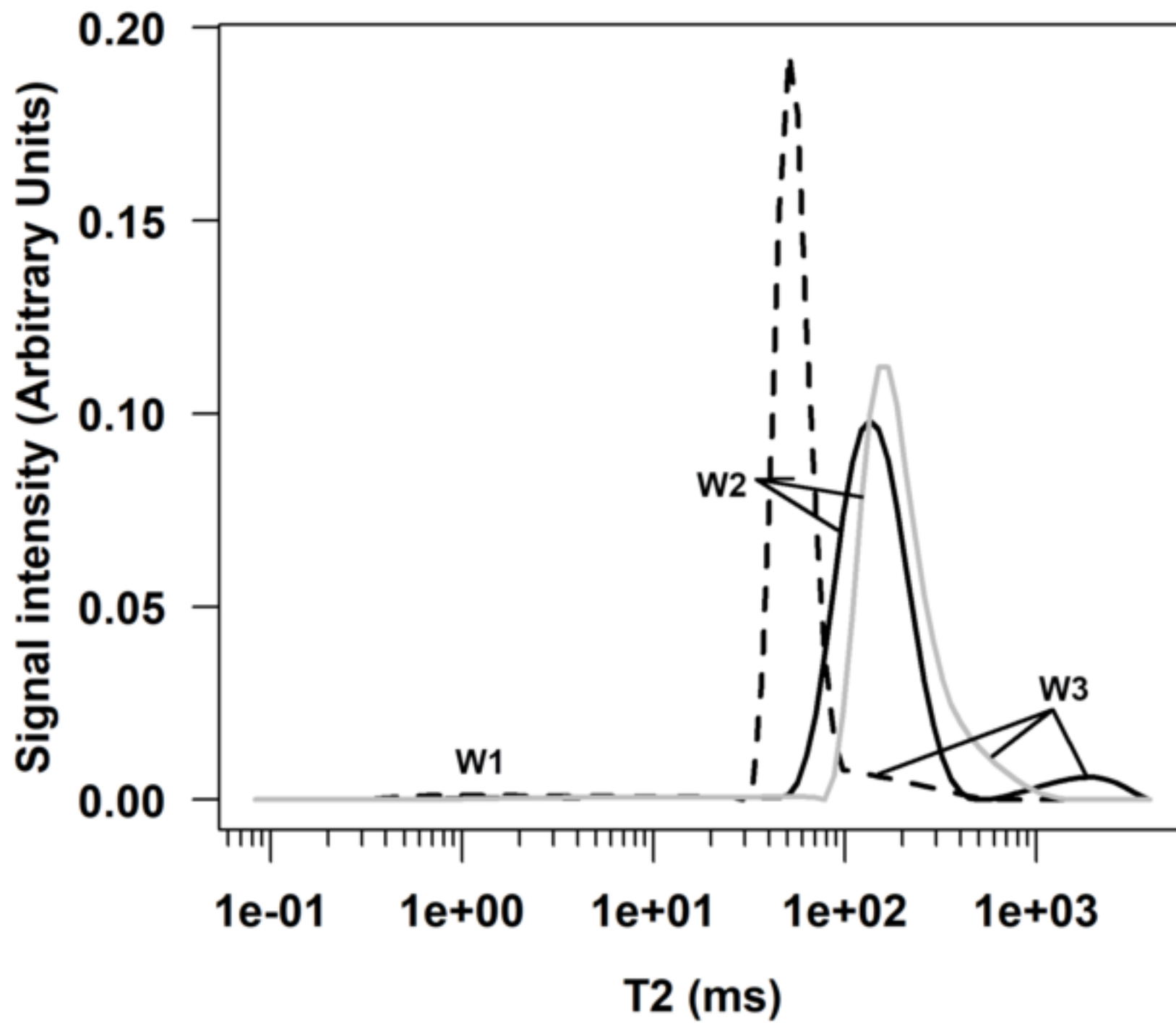


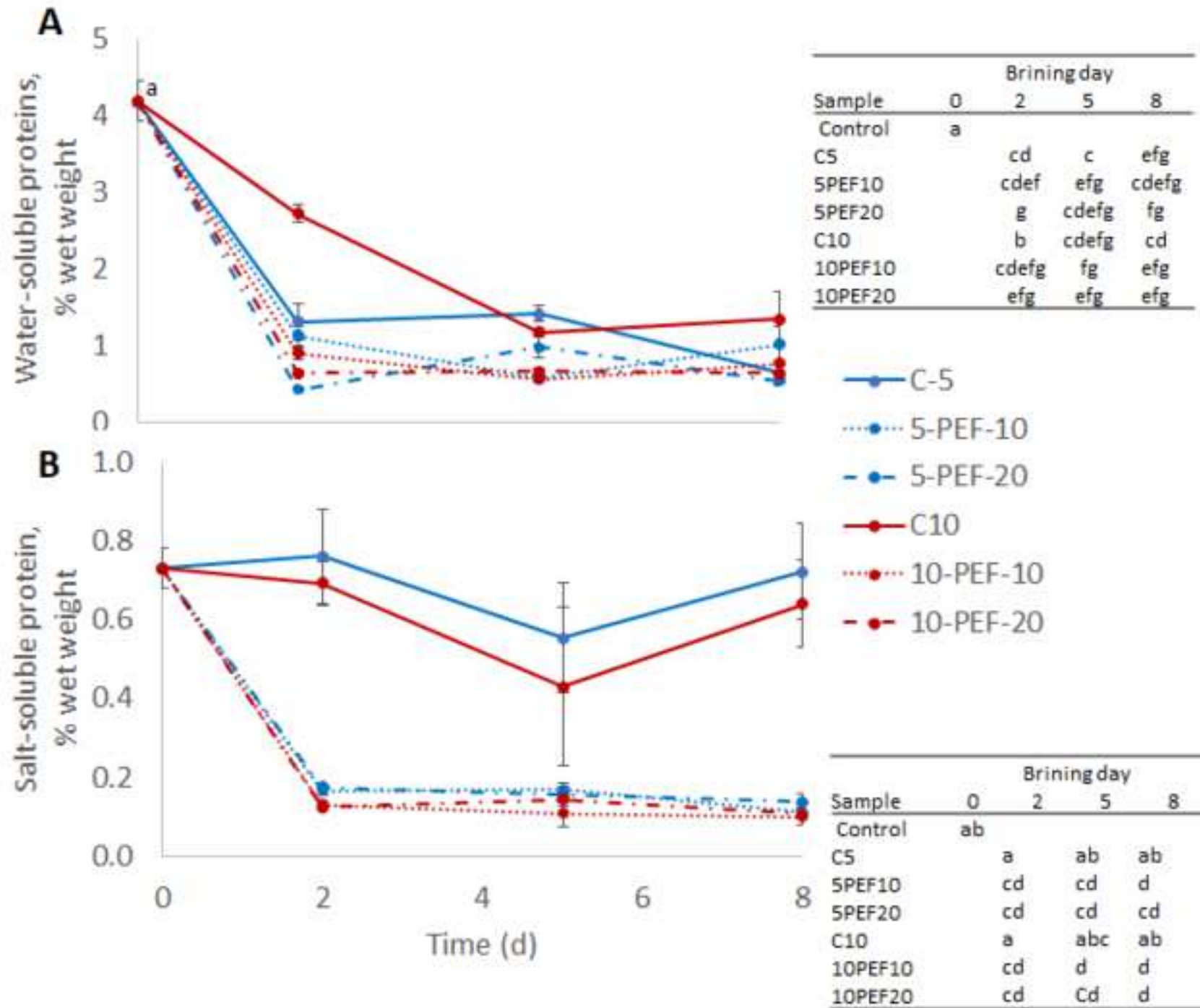














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**Conflict of interest**

The authors declare no conflict of interest.



## **CRediT Author statement**

**Janna Cropotova:** Conceptualization, Formal analysis, Investigation, Writing - Original Draft;  
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