

# Aquaculture

## THE FEASIBILITY OF USING GAS MIXTURE TO STUN SEABREAM (*Sparus aurata*) BEFORE SLAUGHTERING IN AQUACULTURE PRODUCTION

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Abstract:	<p>Current European Union regulation explicitly states that farmed fish should be spared any avoidable pain, distress or suffering at the time of slaughter. It has been shown that fish suffer when they are killed in an ice slurry, the most common method of killing farmed fish in the Mediterranean. Thus, it is necessary to find a method of slaughtering Mediterranean fish that is, (1) efficient in inducing unconsciousness with minimal pain and distress, (2) practical to be applied to a large group of animals at the same time, and (3) feasible to be used at sea. The present study assesses the welfare of Gilthead seabream (<i>Sparus aurata</i>) stunned by two different gas mixtures authorised for stunning other farmed species.</p> <p>To achieve this objective, commercial sized seabream were stunned and /or sacrificed under different protocols: a) killed directly in ice slurry, b) exposed to a mixture of 30% CO<sub>2</sub> + 70% N<sub>2</sub>, and then moved to ice slurry and c) exposed to a mixture of 40% CO<sub>2</sub> + 30% N<sub>2</sub> + 30% O<sub>2</sub> and then moved to ice slurry. Electroencephalograms (EEG) were recorded to evaluate the state of consciousness of seabream during stunning, while blood and brains were sampled to obtain acute stress indicators and relative gene expression, respectively. Additionally, dead fish were kept for in situ meat quality evaluation.</p> <p>When exposed to the gas mixtures, fish lost balance at 1min 23s ± 31s with CO<sub>2</sub> + N<sub>2</sub> and 1min 12s ± 32s, with CO<sub>2</sub> + N<sub>2</sub> + O<sub>2</sub>, respectively. Cortisol, lactate and glucose levels were significantly lower in all fish exposed to gas prior to ice slurry than in fish slaughtered directly in ice slurry (<math>p &lt; 0.05</math>). Electroencephalogram records indicated that fish started to lose consciousness when they lost balance and sank to the bottom of the tank. No differences were found in the meat quality (pH and rigor mortis) among the three treatments.</p> <p>Altogether, the study concludes that the use of carbon dioxide together with nitrogen prior to immersion in ice slurry is more humane than ice slurry alone.</p>
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# THE FEASIBILITY OF USING GAS MIXTURE TO STUN SEABREAM (*Sparus aurata*) BEFORE SLAUGHTERING IN AQUACULTURE PRODUCTION

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## ABSTRACT

Current European Union regulation explicitly states that farmed fish should be spared any avoidable pain, distress or suffering at the time of slaughter. It has been shown that fish suffer when they are killed in an ice slurry, the most common method of killing farmed fish in the Mediterranean. Thus, it is necessary to find a method of slaughtering Mediterranean fish that is, (1) efficient in inducing unconsciousness with minimal pain and distress, (2) practical to be applied to a large group of animals at the same time, and (3) feasible to be used at sea. The present study assesses the welfare of Gilthead seabream (*Sparus aurata*) stunned by two different gas mixtures authorised for stunning other farmed species.

To achieve this objective, commercial sized seabream were stunned and /or sacrificed under different protocols: a) killed directly in ice slurry, b) exposed to a mixture of 30% CO<sub>2</sub> + 70% N<sub>2</sub>, and then moved to ice slurry and c) exposed to a mixture of 40% CO<sub>2</sub> + 30% N<sub>2</sub> + 30% O<sub>2</sub> and then moved to ice slurry. Electroencephalograms (EEG) were recorded to evaluate the state of consciousness of seabream during stunning, while blood and brains were sampled to obtain acute stress indicators and relative gene expression, respectively. Additionally, dead fish were kept for *in situ* meat quality evaluation.

When exposed to the gas mixtures, fish lost balance at 1min 23s ± 31s with CO<sub>2</sub> + N<sub>2</sub> and 1min 12s ± 32s, with CO<sub>2</sub> + N<sub>2</sub> + O<sub>2</sub>, respectively. Cortisol, lactate and glucose levels were significantly lower in all fish exposed to gas prior to ice slurry than in fish slaughtered directly in ice slurry ( $p < 0.05$ ). Electroencephalogram records indicated

40 that fish started to lose consciousness when they lost balance and sank to the bottom of  
41 the tank. No differences were found in the meat quality (pH and *rigor mortis*) among  
42 the three treatments.

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44 Altogether, the study concludes that the use of carbon dioxide together with nitrogen  
45 prior to immersion in ice slurry is more humane than ice slurry alone.

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47 Keywords: Stunning, stress indicators, electroencephalogram, unconsciousness, *Sparus*  
48 *aurata*

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## 1. INTRODUCTION

Aquaculture currently provides almost 50% of all aquatic products consumed by the world's population (FAO 2018), and this proportion is rising due to an increase in the demand for fishery products when the catches obtained from extractive fishing stagnant or decline. The Mediterranean species, gilthead seabream (*Sparus aurata*) is one of the five most cultured species in Europe and the total aquaculture production of sea bream in 2018 was 89,523 tonnes (FAO 2005-2020). This production is mainly from floating sea cages.

At present, there is no legislation in the European Union (EU) to specifically protect the welfare of farmed fish at slaughter. However, the European Regulation on the protection of animal welfare during slaughter and killing (Council Regulation EC 1099/2009) states that animals, including fish, should be spared any avoidable pain, or suffering during stunning and slaughter. An effective stunning leads to a brain state that is incompatible with this capacity and persistence of consciousness (EFSA, 2004). If insensibility is gradually induced, then it should be insured that fish do not the above mentioned negative states during the induction phase. To date and in accordance with the scientific literature and EFSA report (2009), there are two alternative methods that induce immediate loss of consciousness and meet the requirements of the Regulation 1099/2009: (1) stunning by mechanical percussion method followed by bleeding (Van der Vis et al. 2003) and (2) electrical stunning followed by killing (Van der Vis et al. 2003, Lambooij et al. 2008). The main problem for automated percussive stunning is variation in the size of fish within the population, which can cause a mis- stun in fish, especially those weighing less than 1 kg (EFSA, 2009). Electrical stunning is the method commonly used in trout farms throughout the United Kingdom (HSA, 2018)

and, which has been evaluated as safe for workers on these land-based trout farms (Morzel et al. 2003, Knowles et al. 2007).

Mediterranean fish farmers, working on the deck of a boat where the harvest is collected favour a method that requires reduced space and simplicity to safely perform on thousands of individuals at a time with minimum handling. EFSA (2009) mentioned that the most common practice of slaughtering sea bream is in ice water ("ice slurry") and indicated that this method is associated with a long period (minutes) during which the animal is conscious before unconsciousness and death are achieved. During this period until unconsciousness, the fish suffers from suffocation, inferred through physiological and behavioural responses (Kestin et al. 2002, Robb and Kestin 2002, Van der Vis et al. 2003, Acerete et al. 2009). An alternative stunning method with potential to be used on boats would be the exposure to water saturated with gas mixtures such as carbon dioxide (CO<sub>2</sub>) or nitrogen (N<sub>2</sub>). Gas mixtures containing CO<sub>2</sub> induce hypercapnic hypoxia and inhibit neurones through acidosis. However, CO<sub>2</sub> narcosis is aversive to fish, which react with violently to high concentrations of with quick accelerated swimming, thrashing and attempts to escape (Marx et al. 1997, Robb and Kestin 2002, van de Vis et al. 2003, Sanderson and Hubert 2007). Immobility is reached within 2-4 minutes, however, fish would experience pain and distress even if unable to demonstrate it behaviourally (Kiessling et al. 2004). Sea bass (*Dicentrarchus labrax*) exposed to CO<sub>2</sub> remain conscious for 7-10 min and after this period, unconsciousness was demonstrated by complete cessation of rhythmic opercular respiratory movements and heartbeat, absence of VOR (vestibulo-ocular reflex) and pin-prick response (EFSA, 2009). No information was found for seabream exposed to CO<sub>2</sub>, nevertheless an adverse reaction would be expected and has been observed (personal observation by the

100 authors). The gases argon (Ar), oxygen (O<sub>2</sub>) and nitrogen (N<sub>2</sub>) have been experimentally  
101 used in mixtures with CO<sub>2</sub> in animals almost always terrestrial, such as pigs, broilers or  
102 rats, in an attempt to reduce the stress caused by hypercapnia (Gerritzen et al. 2000,  
103 McKeegan et al. 2007, Kirkden et al. 2008, Coenen et al. 2009, Dalmau et al. 2010, Xu  
104 et al. 2011). These studies concluded that these gas mixtures could be used as stunning  
105 methods which induced fewer signs of aversion and breathlessness than only CO<sub>2</sub> where  
106 gas mixtures are already accepted for poultry and pigs (Council Regulation  
107 EC1099/2009). In land animals, it is known that stunning with CO<sub>2</sub> –based gas mixture  
108 has some advantages: meat quality is better than with using electrical stunning (Dich-  
109 Jørgensen et al. 2016), it is cheaper and readily available and it is compatible with the  
110 speed of operation in large slaughterhouses as animals are stunned in groups  
111 (Europgroup for animals, 2019). Nevertheless, it is necessary to evaluate and validate  
112 whether the use of these gas mixtures represents an alternative and more humane  
113 method for fish.

114  
115 Currently, there is very little information on the assessment of welfare and stress during  
116 the slaughter of Mediterranean species and its impact on meat quality (Van der Vis et al.  
117 2003, Knowles et al. 2007, Acerete et al. 2009, Matos et al. 2010), as there are no  
118 feasible and scientifically validated measures. Conscious animals have the capacity to  
119 receive, process and respond to information from internal and external environments  
120 (EFSA, 2004). Therefore, in general, consciousness is associated with the awake state  
121 and the ability to perceive, interact and communicate with the environment and others  
122 (Zeman, 2001). The opposite state, that is, unconsciousness, is defined as: “a state of  
123 unawareness (loss of consciousness) in which there is temporary or permanent  
124 disruption to brain function”. As a consequence of this disruption, the unconscious



animal is unable to respond to normal stimuli (EFSA, 2006). Disruption of brain function can occur as a result of brain concussion, administration of anaesthetics, anoxia or an electroconvulsive shock (Lopes da Silva, 1982). To establish whether the application of gas mixtures can be considered humane, a range of behavioural indicators (e.g. coordinated swimming and escape behaviours, ability to maintain equilibrium, “eye roll” reflex, and ventilatory reflexes) can be implemented to evaluate the degree of consciousness/sensibility in fish (Kestin 2002). However, it has become increasingly clear that behavioural measures alone are not sufficient to assess insensibility, as some commercially used methods may induce sedation and/or paralysis without analgesia or anaesthesia prior to insensibility. Therefore, it is necessary to obtain neurophysiological or neurochemical evidence of insensibility to ascertain the impact of various commercial slaughter procedures. One of the most reliable methods of assessing the state of consciousness is monitoring the brain activity by recording of the electroencephalogram or EEG (Raj et al. 1997, Rodriguez et al. 2008, Bowman et 2019, 2020, Brijs et al. 2021).

Measurement of indices of stress can indicate the welfare status of fish (Pickering, 1992). A typical stress response includes plasma glucose and lactate increase (Lowe-Linde and Niimi 1984, Rotllant and Tort 1997). High levels of cortisol have often been associated with increases in glycemia and plasma lactate, therefore, blood glucose and lactate are considered reliable markers of stress in fishes (Pickering et al. 1982, Simontacchi et al. 2008, Roque et al. 2010). Cortisol is the most informative and accessible marker of stress in fish (Reddy and Leatherland 1998). Elevated cortisol levels are thought to have knock-on effects on blood cells and plasma glucose and lactate; therefore, these variables are also considered representative of the stress status

of fish (Rottlant and Tort 1997). Plasma electrolytes are the most commonly measured indicators of the secondary phase stress response in fish and may provide indirect measurement of altered cortisol (Reddy and Leatherland, 1998).

In addition to optimising fish welfare, it is also necessary to evaluate the impacts of the stunning/slaughter methods on meat quality. From the time of slaughter, the fish carcass starts a process of deterioration that will condition its commercial possibilities. Considering that the loss of quality related to the perception of freshness attributes will be inevitable, efforts should be aimed at delaying the process as much as possible. Minimizing peri-mortem stress will reduce the degradation of ATP-related products (Erikson et al. 1997) and delay the time of occurrence of *rigor mortis* (Erikson 2001) to improve the characteristics of fillets (Robb et al. 2000) and texture (Roth et al. 2002). Therefore, a stunning method that induces loss of consciousness quickly and minimizes adverse reactions by fish will be favourable not only from the point of view of fish welfare, but also on the quality of the final product (Marx et al. 1997). The effects on the quality of the fish according to the method of stunning and slaughter have been studied in several species, mainly salmonids (Skjervold et al. 1999, 2001, Bahuaud et al. 2010), although there are also studies on gilthead seabream (Panebianco et al. 2006, Giuffrida et al. 2007, Campus et al. 2010, Matos et al. 2010).

The present study assesses the welfare of seabream stunned with gas mixes that have been used in other species (chickens, pigs and trout) for the slaughtering of animals in group. It responds to the legislative requirements as well as a demand from a productive sector. The effects of exposing seabream to CO<sub>2</sub>+O<sub>2</sub>+N<sub>2</sub> and to CO<sub>2</sub>+N<sub>2</sub> were evaluated by recording behaviour (loss of equilibrium) and EEG (to assess consciousness) and by

measuring acute stress indicators (cortisol, glucose and lactate). A final evaluation was made on the meat quality to validate the slaughter protocol verifying if the fillet quality was maintained or improved.

## 2. MATERIAL AND METHODS

### 2.1. Ethics statement

The housing, husbandry and use of animals for the procedures described in this manuscript were carried out according to Spanish and European legislation. The project, including this experimental procedure, was approved by IRTA's (Institute of Agrifood Research and Technology, Caldes de Montbui, Spain) Ethics Committee and the Catalan government (approval number: 6722).

### 2.2. Experimental fish:

Seabream came to IRTA from a commercial facility at nursery size (2-5 g wet weight) and were grown for 18 to 24 months in a recirculation aquaculture system (RAS) (IRTAMar®) at 20-21 °C with 100% saturation of dissolved oxygen and full-strength seawater. Fish were fed daily with a Skretting diet for their size and species. A total of 72 fish were used for the different experiments which weighed a minimum of 250 g wet weight and the average size was  $303 \pm 58$  g.

### 2.3. Experimental procedure:

#### 2.3.1. Baseline study (Control fish)

Ten seabream were directly chilled in ice slurry to have a baseline control, mimicking commercial conditions. Time to unconsciousness was not monitored as we considered that this procedure did not adequately stun and kill the fish. Blood was collected from

the caudal vein five minutes after the cessation of breathing, loss of body movements and absence of reaction during handling. Brains from eight fish were extracted immediately after blood sampling and kept in -80 °C for further molecular analysis. After sampling, dead fish were kept inside a 4 °C chamber in ice in perforated recipients to drain water and used for the *in situ* meat quality analysis. EEG was also performed on three fish (see below).

### 2.3.2. Experimental procedure 1: exposure to CO<sub>2</sub>+N<sub>2</sub>+O<sub>2</sub>

The experiment was performed with 15 fish exposed to a gas mixture of 40% CO<sub>2</sub> + 30% N<sub>2</sub> + 30% O<sub>2</sub> (Freshline 3 Mix 50/20, Carbueros Metalicos, Spain). Gas mixtures were selected in these proportions because the mixture was commercially available and had previously been used in land animals (Llonch et al. 2013). In order to define the concentrations of gas to be used, we measured the level of CO<sub>2</sub> in the water when the O<sub>2</sub> was <2 mg /L when using only CO<sub>2</sub>+N<sub>2</sub>. Then the concentration of gas to be used with the cylinder of CO<sub>2</sub>+N<sub>2</sub>+O<sub>2</sub> was defined by using the same level of CO<sub>2</sub>.

A 60 L container with 35-40 L of seawater was used. The gas mixture was bubbled in the seawater from a gas cylinder attached to a manometer with an airline and an air stone until the required concentration of gas mixture in the water was achieved. and Temperature, dissolved oxygen (WTW Oxi 3210) and CO<sub>2</sub> (Handheld OxyGuard CO<sub>2</sub> I) were continuously measured throughout the experiment. The conditions in the water were in the range of 36 - 50 ppm CO<sub>2</sub>, 3.8 - 6.1 mg / L O<sub>2</sub>. Temperature of the water in all experimental procedures was maintained at 21 - 22°C. Fish were exposed to this gas mixture individually and were left an additional 5 min after having lost balance and turned belly up. Blood was then collected. A further group of 10 fish was exposed to the

gas mixture at the same time for meat quality analysis (more details below) following the same procedure. Conditions in this case were 38 ppm CO<sub>2</sub> and 7.7 mg / L O<sub>2</sub>.

### 2.3.3. Experimental procedure 2: exposure to CO<sub>2</sub>+N<sub>2</sub>

For this experiment a combination of 30% CO<sub>2</sub> + 70% N<sub>2</sub> (Freshline 30 Alimentacion; Carburos Metalicos, Spain) was used. The same containers as in experiment 1 were used and gas was dissolved in the water as previously described. The experiment was performed in two groups of 15 and 16 fish. First group (N = 15) was used in a similar exposure as experimental procedure 1 and the second group was used to record the EEG. The conditions in the water for both groups were in the range of 41 - 57 ppm CO<sub>2</sub>, 0.6 - 1.2 mg / L O<sub>2</sub>, and 69 ppm CO<sub>2</sub>, 2.2 mg / L O<sub>2</sub> respectively for the first and second group. All the fish were left in the gas mixture 5 min after having rotated belly up and blood samples were then collected. A further group of eight fish was used *in situ* meat quality analysis. Conditions in this case were 32 ppm CO<sub>2</sub> and 1.7 mg / L O<sub>2</sub>.

All treatments, number of fish and samples taken are specified in Table 2 (see Results section).

### 2.4. Behavioural responses

For the screening experiments, behaviour was the response used to assess whether a fish was unconscious using the following criteria:

- The fish lost balance and turned belly up (onset of unconsciousness) (Raj and Gregory 1996, Dalmau et al. 2016).
- The fish did not react when strongly grabbed by the caudal fin (Schoettger and Julin 1967)

From when fish lost balance, we waited between 3 and 10 min before transferring the fish to ice slurry where it died. The exposure to anaesthetic gas was initially 10 min from the moment when the fish turned belly up. Afterwards, we observed that 5 min exposure did not change the result, i.e., no fish would react being moved from the water supplemented with gas to the ice slurry indicating they were in a non-return condition. Finally, we observed that 3 min was the minimum period of exposure after loss of equilibrium and turning belly-up to observe no return.

## 2.5. Electroencephalogram (EEG)

To ensure the behavioural responses assessed were synchronised with the electroencephalographic record, we first evaluated that fish behaved similarly when exposed to the same conditions. It was verified that the degree of variation in behaviour and time to perform these behaviours (loss of balance and duration of aversion) among individuals were not different. For this purpose, groups of three fish, which were not previously used in any experimental procedure, were exposed to a mixture of CO<sub>2</sub>+N<sub>2</sub> at the same time, and the latency to turn belly up was measured. The timing when fish turned belly up within each group was similar and within a 1-2 s period.

Once we verified the response times were not different between fish exposed to the same gas mixture, a single water mixture with CO<sub>2</sub>+N<sub>2</sub> was prepared and divided into two equal tanks for the exposure. Two fish were exposed to the gas mixture at the same time. One immobilised fish with the EEG record already started (see below) was placed in one tank at the same time the other fish was liberated in the water of the other tank. Both fish were treated similarly before being introduced into the tanks with the gas mixture, netting and time of air exposure were the same, but only the EEG fish was

attached to the EEG (see below). The fish liberated to swim freely in the tank was filmed, therefore obtaining in parallel a behaviour video and an EEG record to correlate the EEG with the screening behaviour (lose balance and turning belly up). This experimental design had previously been used and loss of posture was established as the onset of unconsciousness (Dalmau et al. 2016). The water conditions were: 17.7 °C; 69 ppm CO<sub>2</sub>, 2.2 mg / L O<sub>2</sub>. The experiment was repeated 8 times (8 fish in EEG and 8 fish free in a tank, N = 16).

The QCON Monitor® (Quantum Medical, Spain) is a cerebral consciousness monitor based on wireless technology that assesses brain activity. From the QCON Monitor® (QCON Manual version 6, Valencia et al. 2012), the Index of Consciousness (IoC) and the burst suppression index (BS%) can be estimated to assess unconsciousness during states of anaesthesia (Litvan et al. 2002). The IoC is an algorithm that analyses the raw EEG with a unitless scale from 0 (isoelectric EEG, coma) to 99 (awake) (Revuelta et al. 2008). The BS% indicates the percentage of isoelectric activity during the preceding 30 s and also ranges from 0 to 100 (Litvan et al. 2002). The QCON® monitor is currently used in human patients (Valencia et al. 2012), rabbits (Silva et al. 2011) and pigs (Llonch et al. 2011).

In order to record brain activity through EEG, fish were restrained individually by tying or strapping the fish to a division that was placed in the exposure tank. Two electrodes (Contell Asset Support, Netherlands) were placed on the animal's skull either side of the middle line at the point where the brain is located and separated 5 mm from each other for a transhemispherical electroencephalography (EEG) recording. The reference electrode was placed in the muscle 2-3 cm below the dorsal fin on the right-hand side of the fish. Subsequently, the 3 electrodes were connected to a computer by means of a

150 cm coaxial cable (QCON monitor; Quantum Medical; Barcelona, Spain) to record brain activity using EEG as described in EFSA (2013) and Llonch et al. (2015). The QCON® monitor was then fitted to the electrodes to record EEG data. The data was transferred to a Personal Computer (Acer, Aspire One) for data to be analysed. The moment when the fish became unconscious was identified by plotting the log readings of the brain suppression rate (BS%) and the index of consciousness (IoC) in the same graph and finding the exact moment where the two lines crossed, which indicated the point the fish became unconscious. Baseline EEG activity of the animals was recorded for 1 min, before the animals were placed into the tank and exposed to the gas treatment and the record was maintained 5 min after the free fish lost balance and turned belly up. The fish tied to the division were immersed in the exposure tank once the baseline EEG record was verified to be of good quality. The fish were immersed in the exposure tank leaving only the top of the head out, where the electrodes entered the skull. As previously mentioned, the other fish was released in a second exposure tank at the exact same time EEG fish was placed into an exposure tank, after having been air exposed for the same amount of time as the EEG fish. EEG is a painful and stressful method for fish that for ethical reasons should be used on as few fish as possible. Therefore, it was decided to only perform EEG for the CO<sub>2</sub>+N<sub>2</sub> group that was clearly demonstrated to induce loss of consciousness and because the results showed no significant difference between the two gas treatments (see results section). EEG fish were manipulated in the same manner and after the basal EEG was recorded, they were carefully placed under the ice slurry leaving the top of the head out.

## 2.6. Blood analysis



At the end of the two experiments (procedures 1 and 2) and baseline study, blood was collected ( $\approx 1$  ml) from the caudal vein with 5 mL heparinised syringes with a needle 21Gx 1 1/2". Once the blood samples were collected, the haematocrit was measured and the plasma was obtained by centrifugation, and subsequently frozen at -80 °C until further analysis. The parameters analysed were cortisol (meditec kit, ELISA method), lactate (Abcam kit), glucose (Cromotest kit), magnesium (Cromotest kit) and total protein (Bradford microplate method). All the kits were used according to the manufacturer's instructions and if the reaction was to be developed in volumes higher than 300  $\mu$ L, at the end processed samples were loaded in microplates to facilitate the reading of the optical density in a plate reader (Tecan, Infinite M200 Series). Each fish was sampled 5 min after the fish turned belly up.

## 2.7. Molecular analysis

In the end of the two experiments (procedures 1 and 2) and baseline study, whole brains were extracted from the dead fish and immersed in RNA later and placed for 48h at 4 °C. Brains were then frozen at -80 °C for further analysis. The RNA was extracted from 100 mg of the preserved brains using TRI Reagent RNA Isolation Reagent (SigmaAldrich, Germany) following manufacturer's instructions. The cDNA was synthesized using 1  $\mu$ g of total RNA and oligo dT (20) in 20  $\mu$ L reactions and the SuperScript1 III First-Strand Synthesis SuperMix 50 rxn kit (Invitrogen, Life technologies, USA) following the manufacturer's protocol. Before performing the rt-qPCR, primers (Table 1) were validated by conventional PCR using a cDNA pool from all the samples.

Table 1. Primers used in this study specific for seabream species.

Gene name	Amplicon size	Primer sequence (5'→3')	Accession number	Reference
<i>18s rRNA</i>	134 bp	F: GCA TTT ATC AGA CCC AAA ACC R: AGT TGA TAG GGC AGA CAT TCG	AY993930	Perez Sanchez et al. 2011
<i>efla</i>	134 bp	F: CCC GCC TCT GTT GCC TTC G R: CAG CAG TGT GGT TCC GTT AGC	AF184170	Perez Sanchez et al. 2011
<i>gapdh</i>	111 bp	F: ATCAAGAAGGTCGTCAAGGC R: AGATGGAGGAGTGGCTGTC	DQ641630	Malandrakis et al. 2014
<i>hsp70</i>	174 bp	F: ATT GTT CTG CGC ATC ATC AA R: GCC TCC ACC AAG ATC AAA GA	EU805481	Benhamed et al. 2016
<i>COX2</i>	192 bp	F: GAG TAC TGG AAG CCG AGC AC R: GAT ATC ACT GCC GCC TGA GT	AM296029	Sepulcre et al. 2007

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350 MyTaq™ HS Mix (Bioline) was used to run the conventional PCR with the following  
351 conditions: initial activation step at 95°C for 3 min, followed by 40 cycles: denaturation  
352 at 95 °C for 5 s, annealing at T<sub>m</sub> (58–60°C) 95 °C for 15 s and extension at 60°C for 1  
353 min and 95 °C 15 s, hold 50 °C 10 min. Primer efficiency was evaluated by serial  
354 dilutions to ensure that it was close to 100% performing real time PCR. Target  
355 transcripts (*gapdh*, *efla* and *hsp70*) were analysed by real-time quantitative PCR (rt-  
356 qPCR) (see primers in Table 1). The qPCR was run using a Biometra Optical  
357 Thermocycler (Analytik Jena, Goettingen, Germany) in 96-well plates in duplicate 20  
358 µL reaction volumes containing 10 µL of Luminaris Color HiGreen qPCR Master Mix

(Thermo Scientific), 1 µL of the primer corresponding to the analysed gene (10 pmol), 3 µL of RNA/DNA water free and 5 µL of cDNA in its corresponding dilution. Furthermore, amplifications were carried out with a systematic negative control (NTC; no template control) containing no cDNA. Standard amplification conditions contained an UDG pre-treatment at 50°C for 2 min, an initial activation step at 95°C for 10 min, followed by 35 cycles: 15 s at 95°C, 30 s at the annealing T<sub>m</sub> and 30 s at 72°C. Elongation 1 min 95 °C, 30 s 55 °C and 30s 95 °C. Results were normalised using the housekeeping gene *18S*. The mRNA abundance for each gene was determined using the Pfaffl method (Pfaffl, 2001) on relative quantification.

## 2.8. In situ meat quality analysis

*In situ* meat quality analysis data was collected from 8 fish per treatment where two parameters were assessed:

a) pH. Measurements were made with a pH meter (pHmeter Crison pH25+) attached to a probe which was inserted in a cut made in the muscle with a scalpel. The side of the fish where the cut was made was changed at each measurement. Measurements were taken at 0, 2, 6, 10, 24, 48 and 72h *postmortem*.

b) *Rigor mortis* (RM). Measurements were made at the same time as pH measurements. Rigor development was monitored by carefully placing the fish on a plane surface with two thirds of its length beyond the edge of the surface, i.e. without support provided by the surface. The sag of the tail from the horizontal plane was recorded after five seconds and the rigor index calculated:  
$$\text{Rigor index (\%)} = 100 (\text{current height} - \text{height before entering rigor}) / \text{height}$$

before entering rigor. The fish was then carefully replaced back inside their box until the next measurement.

For the *in situ* meat analysis fish were kept in polystyrene boxes and placed inside a 4°C camera, but the actual measurements were performed at room temperature

## 2.9. Statistical analysis

The data obtained from the QCON monitor (IoC and BS%) were analysed using linear general models, with Proc Mixed proceeding for repeated measures of SAS (SAS 9.4).

In both cases, the variables were submitted to symmetrical composition covariance structure (CS). When the variance analysis showed significant differences ( $p < 0.05$ ), the comparison of least square mean values (LSMEANS) was adjusted to Tukey multiple comparison test (Rodriguez et al. 2008, Dalmau et al. 2016).

One-way ANOVA on ranks was used to detect differences among treatments for the blood parameters analysed. Multiple comparisons were made with the Dunn's method.

Although time to loss of balance could be assessed as an independent variable for the gas treatments, this was not possible for the control group thus a *t-test* was made to see if the time take to lose balance was significantly different between the two gas treatments. Another *t-test* was also performed for the moment when the values of the IoC and the BS% in the EEG crossed versus the moment when the free fish lost balance.

Alle these variables fulfilled the requisites for the use of a parametric test.

A repeated measures (RM) two-way ANOVA was performed to verify the meat quality indexes did not change with the stunning with gases. The two factors were treatment and time. Meat quality values were not normally distributed and the homoscedasticity test was not passed for either parameter, we still preferred to this test instead of ranked

one way ANOVA for each point in time. All ANOVAs and *t-tests* were performed using Sigmaplot (version 12.0).

Level of significance in all statistical tests was considered lower than 0.05 (*P-value* < 0.05).

### 3. RESULTS

Fish were successfully stunned and killed by both gas treatments. Average times per treatment for fish to lose balance and turn belly up varied from 1 to 3 minutes with gases to 52 minutes in ice slurry (Table 2). The longest and shortest periods of time passed **between exposure to treatment and loss of posture** are shown in table 2. This table also lists the type of samples taken from each group of fish. In the case of ice slurry, it was difficult to evaluate when fish were unconscious and therefore, we decided to present the results of when fish were dead. In the case of exposure to gas mixtures, fish started to swim calmly around the tank and between 30 to 80 seconds later, all fish became aware of their situation displaying signs of aversion for periods of around 10 to 12 seconds just immediately before losing balance and turning upside down. Signs of aversion were a very strong acceleration of swimming and raising the head out of the water, but not jumping. Once fish had turned belly up and did not move, tail grabbing was applied without any reaction from the fish. Time elapsed from entering the tank until loss of posture and balance was  $01:12 \pm 00:32$  for fish exposed to  $\text{CO}_2+\text{N}_2+\text{O}_2$  and  $01:23 \pm 0:31$  for fish exposed to  $\text{CO}_2+\text{N}_2$ . A t-test indicated no differences ( $P > 0.1$ ) between the time elapsed until loss of balance in both gas treatments.

Table 2: Time elapsed for fish to turn belly up according to the experimental procedure.

<b>Treatment</b>	<b>Ice slurry (N=10)</b>	<b>CO<sub>2</sub>+N<sub>2</sub>+O<sub>2</sub> (N= 15)</b>	<b>CO<sub>2</sub>+N<sub>2</sub>+O<sub>2</sub> (N=8) (a)</b>	<b>CO<sub>2</sub>+N<sub>2</sub> (N=15)</b>	<b>CO<sub>2</sub>+N<sub>2</sub> (N=8) (a)</b>	<b>CO<sub>2</sub>+N<sub>2</sub> (N=16)</b>
<b>Concentration (ppm)</b>	—	36 - 50 CO <sub>2</sub> , 3.8 - 6.1 O <sub>2</sub>	38 CO <sub>2</sub> , 7.7 O <sub>2</sub>	41 – 57 CO <sub>2</sub> , 0.6 - 1.2 O <sub>2</sub>	32 CO <sub>2</sub> , 1.7 O <sub>2</sub>	69 CO <sub>2</sub> , 2.2 O <sub>2</sub>
<b>Mean± SD (mm:ss)</b>	52:00 ± 10:00*	01:12 ± 00:32	01:23	01:23 ± 0:31	01:29	03:03± 0:38
<b>Max (hh:mm:ss)</b>	01:13:00	02:15	01:23	02:45	01:29	04:12
<b>min (mm:ss)</b>	36:00	00:25	00:59	00:40	00:50	02:15
<b>Other samples</b>	Meat, Blood, RNA	Blood and RNA	Meat	Blood and RNA	Meat	EEG

(a) This group was exposed at the same time and this value corresponds to the last fish that moved.

\* Time until death.

Max = the longest period recorded in a fish to turn belly up in a particular treatment.

min = the shortest period recorded in a fish to turn belly up in a particular treatment.

SD = standard deviation.

The IoC-view® recordings were successful in 7 animals assessed out of 8 exposed to CO<sub>2</sub>+N<sub>2</sub>. One pair of fish from the exposure to CO<sub>2</sub>+N<sub>2</sub> was discarded due to bad reading. The mean ( $\pm$  SD) basal IoC was 90.2 ( $\pm$  11). The IoC started to decrease significantly ( $P < 0.05$ ) at 63 ( $\pm$  20.2) s after placing the fish in the water saturated with the gas mixture (IoC = 89 [ $\pm$  3.7]). It continued decreasing and reached its lowest value on average (IoC = 2) at 343 ( $\pm$  203.99) s after the start of the exposure to the saturated water (Table 3).

The mean ( $\pm$  SD) basal BS% was 0. The BS% started to increase significantly ( $P < 0.05$ ) at 63 ( $\pm$  20.2) s after placing the fish in the water saturated with the gas mixture. It continued increasing and reached its highest value on average (BS% = 94) at 379 ( $\pm$  182) s after the start of the exposure to the saturated water.

All basal EEG were significantly different from the final readings for both IoC and BS% ( $p < 0.05$ ).

Time of unconsciousness by EEG was 3:07  $\pm$  1:17 and to loss of balance (turning belly up) was 3:03  $\pm$  0:38 which was statistically the same with  $P = 0.89$ .

Table 3 Time elapsed to loss of consciousness (EEG) of seabream exposed to CO<sub>2</sub>+N<sub>2</sub>.

Fish	IoC value at start	1) IoC decrease (mm:ss)	2) Loss of consciousness (mm:ss)	3) Loss of balance (mm:ss)
1	67	02:57	05:00	04:12
2	95	01:10	02:45	02:15



3	88	00:01	02:30	03:04
4	91	00:11	03:00	03:04
5	99	00:20	02:38	02:52
6	98	00:50	01:20	02:30
7	97	01:57	04:42	03:25
8	99	00:50	07:50	NA
9	91	00:10	02:50	NA
10	99	01:01	08:11	NA

348

349 The moment when fish lost consciousness was estimated by plotting the log of the  
350 readings as shown in Figure 1. Values of BS% would raise sharply from 0 to close more  
351 than 80 in seconds as the IoC started decreasing (see Figure 1).

352 1) Moment when IoC starts to decrease significantly (mm:ss)

353 2) Moment when fish lost consciousness according to EEG signal (mm:ss)

354 3) Moment when the free fish lost balance and turned belly- up (mm:ss)

355 NA- Fish were under a layer of ice slurry and could not be observed.

356 For the 7 pairs of fish exposed in parallel, a *t-test* indicated the moment where the  
357 values of IoC crossed the BS% values was not significantly different from the moment  
358 when fish lost posture and balance ( $p > 0.1$ ).

359

360 Blood parameters mean values exhibited variation between treatments. Haematocrit  
361 varied from 71.4 % (CO<sub>2</sub>+O<sub>2</sub>+N<sub>2</sub>) to 40.3 % (ice slurry); Glucose from 14.09 (CO<sub>2</sub>+N<sub>2</sub>)  
362 to 247.3 g/dL (Ice slurry); Cortisol from 2.48 (CO<sub>2</sub>+N<sub>2</sub>) to 474.1 nmol/uL (Ice slurry);  
363 Lactate from 0.464 (CO<sub>2</sub>+O<sub>2</sub>+N<sub>2</sub>) to 12.44 nmol/mL (ice slurry); Protein from 5.29  
364 (CO<sub>2</sub>+N<sub>2</sub>) to 18.26 mg/mL (CO<sub>2</sub>+O<sub>2</sub>+N<sub>2</sub>); Magnesium from non-detected (ND, CO<sub>2</sub>+N<sub>2</sub>)

and CO<sub>2</sub>+O<sub>2</sub>+N<sub>2</sub>) to 9.86 mg/dL (CO<sub>2</sub>+O<sub>2</sub>+N<sub>2</sub>). Statistical analysis showed that in general treatment with ice slurry was significantly different from exposure to gas mixtures for the following parameters: cortisol, glucose, lactate and magnesium (P < 0.05, Table 4).

Table 4 Biochemical plasmatic parameters measured in fish from the different treatments and control.

Treatment	Haematocrit %	Glucose g/dL	Cortisol nmol/uL	Lactate nmol/mL	Protein mg/mL	Magnesium mg/dL
Ice slurry	51.0±1.7	145.8±16.5	109.5±38.9	521±28.6	14.6±0.31	2.7±0.7
CO <sub>2</sub> +O <sub>2</sub> +N <sub>2</sub>	52.4±6.8	135.3±41.9	41.9±41.1*	60.6±34.1*	15.5±1.67	4.36±2.11*
CO <sub>2</sub> +N <sub>2</sub>	50.8±4.9	79.1±14.1*	35.7±32.2*	132.5±31.5*	13.7±3.2	5.02±1.76*

\* indicates the treatment is significantly different from the direct exposure to ice slurry (control).

Relative gene expressions were estimated using *18S* gene as the housekeeping gene. Relative gene expression had a very high variation both intra and inter groups and no differences were obtained between treatments. The gene expression results have been included as supplementary material.

The pH started descending as soon as the fish were dead although in the fish killed with ice slurry this decrease was slower. Initial values of pH were 7.21 ± 0.14, 6.83 ± 0.16 and 6.74 ± 0.15, and they decreased until 72h where pH values were 6.41 ± 0.07, 6.38 ± 0.12, 6.35 ± 0.05 for slaughtering in ice slurry, exposure to CO<sub>2</sub>+N<sub>2</sub>+O<sub>2</sub> and exposure to CO<sub>2</sub>+N<sub>2</sub>, respectively. Meat will be of better quality when the pH is lower (Love, 1980),

after 72h there were no differences among treatments (Figure 2A). Again from 8 hours onwards all treatments showed a parallel pH progress.

The experimental gas mixture treatments used induced a faster instauration of *rigor mortis* (RM) when compared to killing directly in ice slurry. The later the instauration of RM the better, so that there is time to process fish (Figure 2B). At 2 hours, RM was the following  $37.94 \pm 20.12$ ,  $79.22 \pm 20.41$ ,  $81.87 \pm 5.66$  for killing in ice slurry, exposure to  $\text{CO}_2+\text{N}_2+\text{O}_2$  and exposure to  $\text{CO}_2+\text{N}_2$ , respectively. This meant that, after 2h the group of fish placed directly in ice slurry was the only group that had not entered RM phase, however from 8h onwards all groups showed a parallel RM evolution. At 72h, RM values were  $68.03 \pm 8.91$ ,  $67.67 \pm 17.74$ ,  $71.63 \pm 7.22$  for slaughtering in ice slurry, exposure to  $\text{CO}_2+\text{N}_2+\text{O}_2$  and exposure to  $\text{CO}_2+\text{N}_2$ , respectively.

#### 4. DISCUSSION

The aim of this study was to evaluate the effectiveness of two gas mixtures to be used as a method to stun Mediterranean fish from aquaculture production using seabream as a model species. Most fish took less than 1 minute and 30 seconds to initiate loss of equilibrium, irrelevant of the gas mixture. In the exposure to  $\text{CO}_2+\text{N}_2$ , an EEG demonstrated that fish start losing consciousness at the point they lose balance and turn upside down. Thus, fish losing balance and turning upside down might be defined as the moment when fish start losing consciousness. In practice, these behavioural responses can be used an operational indicator for stunning and killing fish. Nevertheless, fish can only be considered properly stunned 3 minutes after having lost balance, since it is important to ensure they will not recover when moved to the ice slurry, in order to ensure the welfare of all fish. However, this suggestion must be taken with caution due

410 to the reduced number of fish used in the experiment. At this point, soon (5 minutes)  
411 after losing balance and consciousness some fish were blood sampled for primary  
412 indicators of stress (glucose, cortisol and lactate) and significant differences were found  
413 between fish killed directly in ice slurry *versus* fish exposed to the gas mixture. In  
414 addition, *in situ* meat analysis was not different among treatments leading us to  
415 conclude that flesh quality is not affected by introducing this stunning method. Both  
416 mixtures seemed to induce similar reactions and no differences between treatments were  
417 perceived.

418  
419 To our knowledge, there is no data available on stunning seabream with gas mixtures.  
420 However, Zampacavallo and collaborators (2003, 2015) stun-killed seabass in ice water  
421 saturated with 60% CO<sub>2</sub> + 40% N<sub>2</sub> and with 30% CO<sub>2</sub> + 70% N<sub>2</sub>, respectively. In their  
422 studies, the authors confirmed a significant reduction in the time take to achieve death  
423 from 20 minutes to 6 and 10 minutes respectively.

424  
425 In the present study, no treatment rendered the fish unconscious in an immediate  
426 manner. Nevertheless, fish only displayed aversion to their situation for 10-12 seconds  
427 immediately before turning belly up and showing signs of losing consciousness. This  
428 aversion moment has been observed in other species where gas mixtures were used for  
429 stunning (Llonch et al. 2013, Dalmau et al. 2016, Verhoeven et al. 2016). The time  
430 which fish were, most likely, in a situation that impaired their welfare were those 10  
431 seconds before the fish started to lose consciousness. In the beginning of the exposure,  
432 fish swam calmly around the tank. Exposure to CO<sub>2</sub> alone is problematic since fish  
433 display several signs of aversion (Van der Vis et al. 2003, Erikson 2011, Roque  
434 personal observation), however, adding N<sub>2</sub> and / or O<sub>2</sub> has been suggested to mitigate

this aversion (Gerritzen et al. 2000, McKeegan et al. 2007, Kirkden et al. 2008, Coenen et al. 2009, Dalmau et al. 2010, Xu et al. 2011). All these studies showed that mixing gases with N<sub>2</sub> worked better than CO<sub>2</sub> alone for the species concerned (EFSA, 2009). The addition of oxygen (O<sub>2</sub>) for Arctic char did not increase time to loss of balance showing that O<sub>2</sub> does not antagonise the anaesthetic capacity of CO<sub>2</sub> (Sandblom et al. 2013).

Rodríguez et al. (2008) and Llonch et al. (2011) concluded that a significant decrease in the electrical activity of the brain is considered a sign of the onset of unconsciousness in pigs. This is also the case for rabbits (Dalmau et al. 2016). Moreover, EFSA's review (2013) clearly states that changes in EEG power are considered a good indicator of brain activity in studies where animals were stunned with gas. In the present study, an IoC significantly lower than basal values occurred from a few seconds to nearly 3 minutes after the immersion in a bath containing a mixture of CO<sub>2</sub> and N<sub>2</sub>. The experiment showed that fish lose consciousness, with the IoC decreasing as the BS% increased. According to the manufacturer's manual a IoC between 0 and 40 corresponds to deep anaesthesia in humans, but as the device was not developed for fish species, we never finished the record with a IoC < 5 to ensure the fish was at a point of no return. Van der Vis et al. (2003) observed a difference of approximately 5 minutes between a salmon exposed to CO<sub>2</sub> losing balance and being declared unconscious by losing the VER. In the present study, the mean time difference between loss of balance and IoC value being lower than BS% was 4.87 s. Nevertheless, we still used loss of balance as the operational indicator of unconsciousness as it is very easy to appreciate even when observing a group of fish instead of individual fish, where other indicators such as VER would be difficult to appreciate. Still, as stated in material and methods section, fish

were left in the exposure tanks for a minimum period after having lost balance, in order to ensure they could not react or recover. Timings in this experiment were longer than those when just observing fish and this is most likely explained by a longer handling procedures. Both fish (free swimming and EEG fish) were outside the water for more than one minute (setting up of the EEG and one-minute record) and this increases the level of stress in the fish making it more difficult for them to anaesthetise (Zahl et al. 2013).

Acute stress parameters (glucose, cortisol and lactate) were significantly higher in fish sacrificed directly in ice slurry implying that this treatment was more stressful for fish.

A typical stress response includes plasma glucose and lactate increase (Lowe-Linde and Niimi, 1984; Rotllant and Tort 1997). High levels of cortisol have often been associated with increases in glycemia and plasma lactate; therefore, blood glucose and lactate are considered reliable markers of stress in fishes (Pickering et al., 1982; Simontacchi et al. 2008; Roque et al. 2010). Cortisol response peaks after 2.5 to 60 minutes (Pankhurst 2011) and present experimental design follows previous literature demonstrating significant differences among treatments (Zampacavallo et al. 2003, 2015, Daskalova et al. 2016, Gräns et al. 2016). In Arctic char exposed to a mixture of CO<sub>2</sub> + O<sub>2</sub> (50-50), cortisol increased significantly from basal levels only 30 minutes post exposure (Sandblom et al. 2013), which would not be a problem in the present study since fish would be dead by then. This delayed cortisol response could be related to long deep anaesthesia (Sandblom et al. 2013) which makes these mixes fit for the purpose of this study. In the present study, no recovery investigation was made since the purpose was an irreversible stunning method and personal observations established that seabream weighing between 250-500 g (commercial size) do not recover if exposed to the gas

mixture for at least 3 minutes after they lost the balance. In the present study, stress parameters were measured 5 minutes after fish had lost balance and consciousness and demonstrated that fish in ice slurry had higher levels of stress at this point. However, the time period to this point was different in different treatments and show to be much longer in ice slurry ( $52:00 \pm 10:00$  m). It cannot be discounted that the stress response was similar across treatments but had more time to develop in the ice slurry treatment. This also seems to be the case with seabass where fish stunned in ice water saturated with  $\text{CO}_2+\text{N}_2$  and sampled 30 to 60 minutes later, presented much higher values than in present case (Zampavallo et al. 2003, 2015). However, clearly at the point of loss of consciousness ice slurry fish had higher levels that can indicate higher stress and these ice slurry fish had a considerably longer period ( $52:00 \pm 10:00$  m) in this stressful state before losing consciousness compared to the  $< 4$  minutes registered in gas treatments. Magnesium was significantly higher in the fish exposed to gases, and this is most likely due to acidification of water and blood by the  $\text{CO}_2$  (Shrivastava et al. 2019) and fish must compensate for the blood acidosis. The significant alteration in plasma ions as magnesium in fish exposed to gas mixture might represent disturbances in acid-base balance, oxygen and carbon dioxide transport (Roque et al. 2010). This result was in accordance with the findings of Tort et al. (2003).

Longer awareness of the fish killed in ice slurry is probably the explanation of the increase in glucose, since this results as a response to the release of stress-induced hormones in the blood circulation, which trigger muscle or liver glycogenolysis, releasing glucose for the increased energy requirement during stress (Eslamloo et al. 2014). This increased energy demand also leads to the increment of blood lactate, caused by the anaerobic activity of muscles (Wang and Richards 2011, Zampacavallo et

al. 2003) and hypoxic stress (Eslamloo et al. 2014) in the case where the gas mix did not contain oxygen.

The gene expression (supplementary data) in relation to the treatments was very variable and no conclusions could be drawn. On reflexion, we realise the very short-term exposure to the experimental treatments (stunning lasting less than 2 minutes) most likely did not induce a marked response on the synthesis of mRNA and thus we would be measuring more than anything the pre-stunning *antemortem* period which was common to all the fish. The differences would then be explained by the individuality of the fish (Jolles et al. 2020) where even though the fish had been exposed to the same circumstances, the individual fish varied either the response or at least the abundance of the response. The mRNA had high quality when measured by spectrophotometry and visualised in a gel, however, it must be pointed that some mRNA are very short lived, 5 to 10 minutes (Guaniyogi and Brewer 2001), and the fish were kept in the water five minutes before sampling. Another aspect to potentially contribute to this situation, was that the fish were stunned in a very hypoxic environment which leads to an increase of ATP and consequently to cellular degradation, including the mRNA. Many studies use sacrifice in anaesthesia as the negative control, a decision was taken that for the present study such control would not be used because there is a vast amount of literature demonstrating the use of anaesthesia is not without stress for the fish (Toni et al. 2015, Bodur et al. 2018, Freitas Souza et al. 2019, Teles et al. 2019). Slaughtering the fish by hitting them on the head would be a solution to this, unfortunately this is not feasible when you need to sample the brain and cutting of the spinal cord leaves the heart and the brain in most cases on the same half of the fish which is not a good situation from a consciousness perspective.



535

536 In the present study, the treatment did not seem to affect the meat quality. This was  
537 slightly surprising because the fish in the ice slurry treatment were more stressed than  
538 the ones submitted to gas exposure according to the blood parameters. In fact, at the  
539 beginning (2 hours), both meat quality parameters had better values for the fish killed  
540 directly in ice slurry (pH: 7.01 versus 6.75 and 6.79, *rigor mortis*: 37.94 versus 79.22  
541 and 81.87 for ice slurry, CO<sub>2</sub>+N<sub>2</sub>+O<sub>2</sub> and CO<sub>2</sub>+N<sub>2</sub>, respectively). For both parameters,  
542 these values are similar to those previously reported in the literature for seabream  
543 (Tejada and Huidobro 2002) and similar to seabass (Zampacavallo et al. 2003, 2015).  
544 When fish are stressed during crowding, they deplete their energy reserves prior to  
545 slaughter and *rigor mortis* occurs sooner than when the fish have been crowded  
546 carefully. A delayed *rigor mortis* allows processing to take place before rigor occurs  
547 (Sigholt et al. 1997). With early *rigor mortis* the flesh can be difficult to process,  
548 reducing both the yield and flesh quality and resulting in a shorter shelf-life (Morzel et  
549 al. 2003). *Rigor mortis* is characterised by a progressive rigidity of the body due to a  
550 reduction of ATP levels in the muscle. Thus, an intense stress *antemortem* will increase  
551 the anaerobic metabolism which will consume energy reserves and will accelerate both  
552 the start and the resolution of the *rigor mortis* (Nakayama et al. 1992, Eriksson et al.  
553 1997, Sigholt et al. 1997, Robb 2001). Under stressful conditions, there will be an  
554 accumulation of lactic acid in the muscle that will induce a reduction of the pH during  
555 sacrifice. This pH reduction will contribute to a faster *postmortem* drop of the muscle  
556 pH in the stressed fish. However, this was not the case in the present study. Even  
557 though, the muscle pH in ice slurry treatment was higher for the first 2 hours  
558 *postmortem*, from 8 hours onwards there were no differences among treatments  
559 indicating that if stunning was not improving the quality of the meat, it did not

significantly deteriorate it, which was also in accordance with studies in other fish species, such as Atlantic salmon (Sigholt et al. 1997) and eels (Morzel and Van der Vis, 2003). Flesh pH is considered a good indicator of the muscle texture (Love, 1980) and of the shelf-life of the fish (Foegeding et al. 1996, Zampacavallo et al. 2003, 2015). *Postmortem* muscle pH is around 7 and then it decreases to 6.5 or less due to lactic acid accumulation.

Altogether, the results obtained in the present study suggested that both gas mixture treatments, CO<sub>2</sub>+N<sub>2</sub>+O<sub>2</sub> and CO<sub>2</sub>+N<sub>2</sub>, have a high feasibility to be used as stunning method by the aquaculture industry to preserve the welfare of a Mediterranean fish species like gilthead seabream.

## 5. CONCLUSIONS

Although stunning with gas mixtures is not an immediate stunning method for seabream, the exposure to either 30% CO<sub>2</sub> and 70% N<sub>2</sub>, or 40% CO<sub>2</sub>, 30% N<sub>2</sub> and 30% O<sub>2</sub> induce less suffering than ice slurry treatment alone (EFSA 2009) which is a clear advantage to the seabream production.

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## List of figures

**Figure 1A** Evolution of an electroencephalogram (EEG) record along time in seabream.

The Index of consciousness (IoC) represented in continuous line can be seen decreasing whereas the brain suppression rate (BSR), in dash line, can be seen increasing. Fish were immersed in the water with gas approximately 1 minute after starting the record. Green arrow indicates when lines crossed and the moment when the fish is defined to become unconscious.

**Figure 1B** Example of the general EEG of a seabream. Time 0 line indicates when the exposure to the gases started.

**Figure 2** Evolution of the pH (A) and *rigor mortis* (B) for 72h at 4°C in seabream.

Measurements were made at room temperature, but samples were kept refrigerated outside the brief moments of measurement. Error bars represent the standard deviation of the mean of the fish at that point in time. Graph lines represent the mean of 8 fish per treatment.

Figure 1A

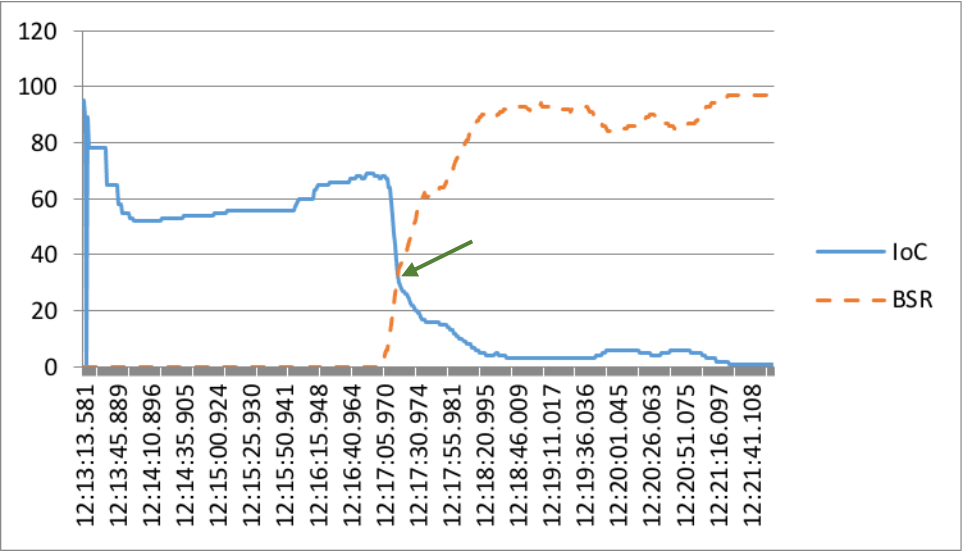


Figure 1B

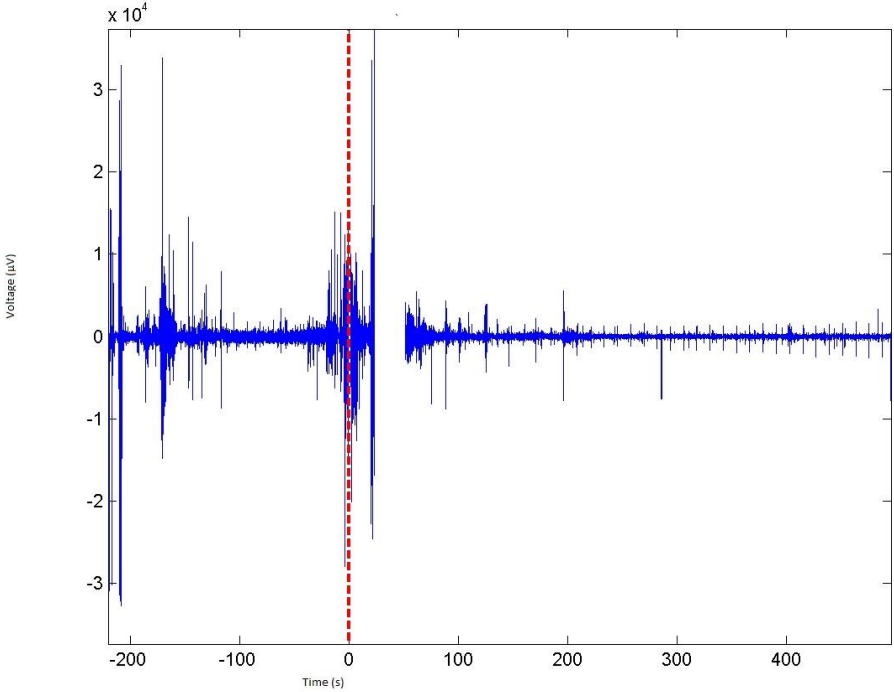


Figure 2 A

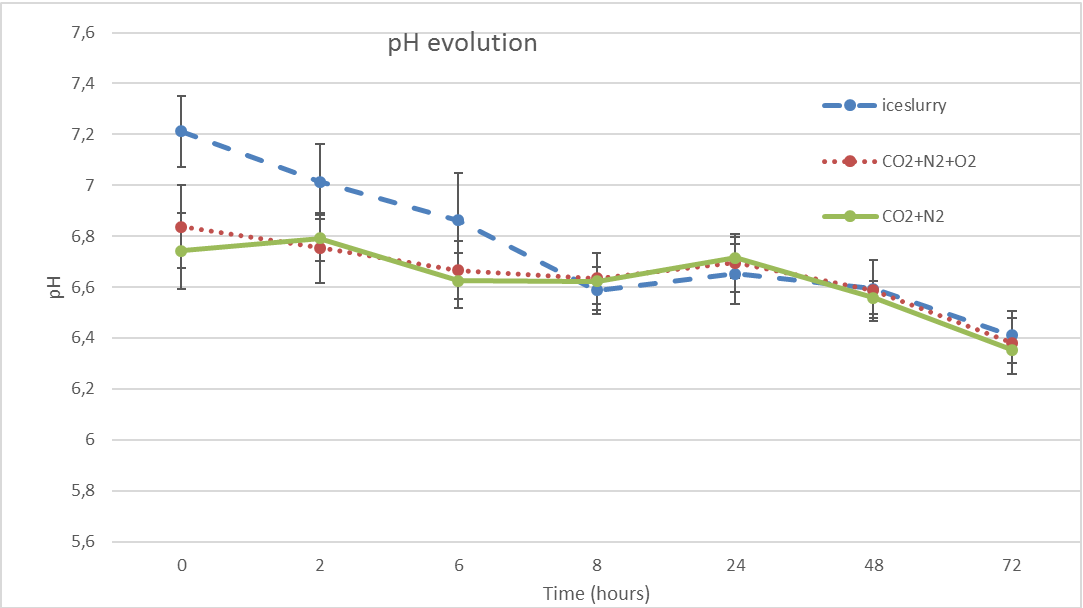
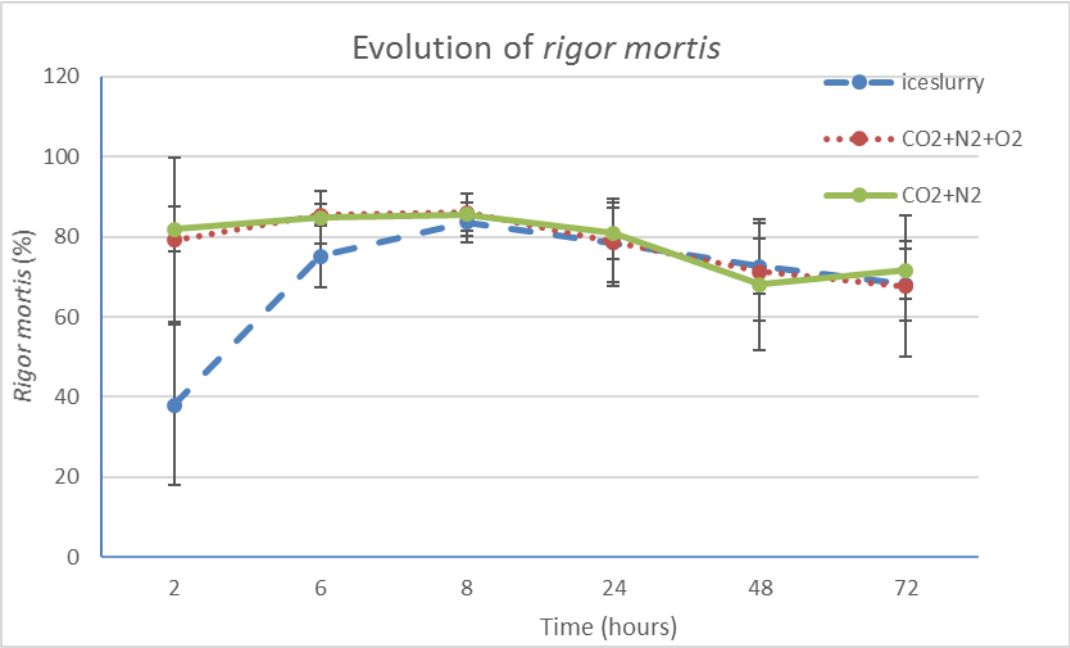


Figure 2B



Supplementary material

Relative gene expressions were estimated using *18S* gene as the house keeping gene. A one way ranked ANOVA to search for differences among treatments showed that the expression was different for *gapdh*, *hsp70*, *cox2* and *ef2α*, where:

Fish exposed to CO<sub>2</sub>+O<sub>2</sub>+N<sub>2</sub> differentially expressed *gapdh* (P = 0.014) from fish exposed directly to ice slurry; fish exposed to CO<sub>2</sub>+N<sub>2</sub> differentially express *hsp70* (P = 0.002) and *ef2α* (P = 0.005) from fish exposed directly to ice slurry (Figure S1).

Figure S1:

