

Elevated CO₂ Affects Predator-Prey Interactions through Altered Performance

Bridie J. M. Allan^{1*}, Paolo Domenici², Mark I. McCormick¹, Sue-Ann Watson¹, Philip L. Munday¹

1 ARC Centre of Excellence for Coral Reef Studies, and School of Marine and Tropical Biology, James Cook University, Townsville, Queensland, Australia, **2** CNR-IAMC, Istituto per l'Ambiente Marino Costiero, Località Sa Mardini, Torregrande, Oristano, Italy

Abstract

Recent research has shown that exposure to elevated carbon dioxide (CO₂) affects how fishes perceive their environment, affecting behavioral and cognitive processes leading to increased prey mortality. However, it is unclear if increased mortality results from changes in the dynamics of predator-prey interactions or due to prey increasing activity levels. Here we demonstrate that ocean pCO₂ projected to occur by 2100 significantly affects the interactions of a predator-prey pair of common reef fish: the planktivorous damselfish *Pomacentrus amboinensis* and the piscivorous dottyback *Pseudochromis fuscus*. Prey exposed to elevated CO₂ (880 μatm) or a present-day control (440 μatm) interacted with similarly exposed predators in a cross-factored design. Predators had the lowest capture success when exposed to elevated CO₂ and interacting with prey exposed to present-day CO₂. Prey exposed to elevated CO₂ had reduced escape distances and longer reaction distances compared to prey exposed to present-day CO₂ conditions, but this was dependent on whether the prey was paired with a CO₂ exposed predator or not. This suggests that the dynamics of predator-prey interactions under future CO₂ environments will depend on the extent to which the interacting species are affected and can adapt to the adverse effects of elevated CO₂.

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* E-mail: bridie.allan@my.jcu.edu.au

Introduction

Predation is one of the key processes structuring communities in ecological and evolutionary time [1]. Prey with well-honed antipredator responses will have high survival, while predators with low catch rates suffer slower growth and reduced reproductive output [2,3]. Any factor that influences a part of the predator-prey interaction sequence will affect the outcome for both parties. Recent research has shown that carbon dioxide (CO₂) levels projected to occur in the ocean by the end of this century (based on Representative Concentration Pathways RCPs) [4] can affect the behavior of marine organisms and significantly increase predation rates in natural habitat [5–10]. These findings emphasize the potential impact that elevated CO₂ in the ocean could have on marine population dynamics and ecosystem processes [11–13]. To date however, the dynamic mechanisms of predator-prey interactions that underlie increased predation rates in high CO₂ environments are unknown.

The concentration of CO₂ in the atmosphere has increased approximately 40% since the industrial revolution, to present-day levels >390 ppm. The atmosphere and surface-ocean are in approximate gas equilibrium; therefore the concentration of CO₂ in the ocean also increases as atmospheric CO₂ increases [14]. If the current trajectory of emissions is maintained, atmospheric CO₂ concentrations are predicted to exceed 900 ppm by the year 2100 [4]. It has recently been demonstrated that these levels of dissolved CO₂ can dramatically alter the response of fish predators to prey sensory cues [15] and of fish prey to the presence of

predators [6]. Fish exposed to elevated CO₂ exhibit impaired olfactory [6,15] and auditory responses [16], altered activity levels [7,9] and reduced behavioral lateralization (the propensity to turn left or right) [17]. The underlying mechanism for these behavioural changes appears to be impaired neurotransmitter function in fish that are permanently exposed to elevated CO₂ [18]. If elevated CO₂ alters the processing of sensory information, then it may directly influence the behavioral basis that governs the outcome of a predator-prey interaction, such as the timing of a predator strike or the distance to react to the predator by the prey. During a predator-prey interaction, fish employ a rapid acceleration called a fast-start that is used by predators to capture prey, or by prey to avoid a predatory strike [19,20,21]. Consequently, any effect of elevated CO₂ on the timing or performance of fast-starts by either the predator or prey could lead to changes in prey mortality.

This study examined the potential cause of increased mortality of prey fish that has been observed in previous studies [7,9] by asking the specific question: Does exposure to elevated CO₂ change the outcome of predator-prey encounters by altering the kinematics of the predator-prey interaction? To test this hypothesis, locomotion performance, prey reaction distance and capture success were examined in staged encounters between newly metamorphosed individuals of a prey fish, *Pomacentrus amboinensis*, and a common predator, *Pseudochromis fuscus*. Predators and prey were exposed to CO₂ levels (880 μatm) relevant to the end of the century based on the most recent representative greenhouse gas

concentration pathways [4]. Previous experiments have demonstrated that the mortality rates of high-CO₂ exposed prey when placed in the field with non-CO₂ exposed predators are 2–3 times higher than when prey are exposed to similarly treated predators in the laboratory [7,9]. This suggests that the behavior of the CO₂ exposed prey depends on whether or not the predator has also been exposed to elevated CO₂. Therefore, the responses of fish exposed to high CO₂ were compared to fish exposed to present day CO₂ levels (440 μatm) in a fully crossed design. This enabled us to tease apart the independent effects on the predator and prey as well as the interacting effects when both were exposed to elevated CO₂.

Materials and Methods

Ethics Statement

Research was carried out under approval of the James Cook University animal ethics committee (permit: A1067) and according to the University's animal ethics guidelines. Fish collections around Lizard Island, Great Barrier Reef were carried with permission of the Great Barrier Reef Parks Authority (permit: G10/33239.1) and Queensland Government Department of Primary Industry and Fisheries (permit: 103256). Suffering was minimal as prey were consumed immediately following a successful strike.

Study Site and Species

Fishes were collected during December 2010 at Lizard Island (14° 40' S, 145° 28' E), northern Great Barrier Reef (GBR) and maintained in a flow-through seawater system at the Lizard Island Research Station (LIRS). Newly metamorphosed individuals of the common damselfish, *Pomacentrus amboinensis* (Pomacentridae) were used as the prey species. The dottyback, *Pseudochromis fuscus* (Pseudochromidae) was used as the predator. *P. fuscus* is an abundant, small, widely distributed mesopredator found throughout the Indo-Pacific. It is a gape limited, highly territorial and active predator and makes up 9.5% of the piscivorous reef fish assemblage at Lizard Island [22]. It is considered an important predator of newly settled coral reef fishes [23,24]. *P. fuscus* readily adjusts to aquarium conditions and has been observed exhibiting normal feeding and aggressive behaviors within 48 hours of collection [25]. Because of this, it has been used extensively as a model predator in predator-prey manipulation studies.

Newly metamorphosed *P. amboinensis* (range 10.3–15.1 mm, 12.6 mean standard length (SL), standard deviation (SD) 1.5) [26] were collected using light traps [27] moored ~100 m off the fringing reef of Lizard Island. On the morning of capture they were transferred to 30 L aquaria supplied with a continuous flow of either control (present-day CO₂) or elevated-CO₂ seawater (see below) for 4 days. This period of time has been found to be sufficient to elicit the full behavioural effects of high CO₂ and larval fish do not become acclimated with longer exposure [28]. Fish were fed 4 times daily *ad libitum* with newly hatched *Artemia* sp. but were starved for the 12 hours prior experimental trials to standardize for satiation.

Adult *P. fuscus* (range 64–83 mm, 72.3 mm mean SL, SD 0.6) were collected with a dilute solution of clove oil [29] from the shallow fringing reef around Lizard Island. Immediately after collection, fish were transported back to LIRS where they were housed separately in mesh breeding baskets within 30 L aquaria to avoid aggressive interactions. Fish were maintained in tanks for 4 days and were fed 2 juvenile reef fish for the first 2 days and then not fed for the last 2 days to standardize for satiation. Food deprivation in the wild is not unusual and previous work has

demonstrated a high prevalence of gut emptiness for piscivorous fish [30].

CO₂ Treatment

The pCO₂ of treatment seawater was manipulated by CO₂ dosing to a set pH_{NBS}. Seawater was pumped from the ocean into 2×60 L sumps where it was diffused with ambient air (control) or CO₂ to achieve the desired pH (CO₂ treatment). A pH of 7.89 was selected to achieve the approximate pCO₂ required, based on preliminary observations of total alkalinity, salinity and temperature of seawater at Lizard Island. A pH-controller (Tunze Aquarientechnik, Germany) was attached to the CO₂ treatment sump to maintain pH at the desired level. A solenoid injected a slow stream of CO₂ into a powerhead at the bottom of the sump whenever the pH of the seawater rose above the set point. Equilibrated seawater from each sump was then supplied at a rate of ~500 ml.min⁻¹ to eight replicate 35 L aquariums, four housing small groups of *P. amboinensis* and four housing *P. fuscus*. Temperature and pH_{NBS} of each aquarium was measured each morning and afternoon using an HQ40d pH meter (Hach, Colorado, USA) calibrated with fresh buffers. Total alkalinity (TA) of seawater was estimated by Gran titration from water samples taken twice weekly from control and treatment tanks. Alkalinity standardizations achieved accuracy within 1% of certified reference material from Dr. A. Dickson (Scripps Institution of Oceanography). Average seawater pCO₂ was calculated using measured values of pH, TA, temperature and salinity in the program CO2SYS [31] and using the constants of Mehrbach *et al.* [32] refit by Dickson and Millero [33]. Seawater parameters are shown in Table 1.

Laboratory Assays

Trials were conducted over a period of 10 days in a temperature-controlled room at LIRS, ensuring the water temperature remained between 26 and 28°C. One predatory *P. fuscus* was placed into the experimental arena (38 cm×58 cm×10 cm water height) and one *P. amboinensis* was then released into a length of PVC tube (11 cm diameter, 15 cm high) placed upright in the middle of the experimental arena. Both fish were allowed to acclimate for 30 min. The PVC tube was then carefully raised and removed from the tank using a wire connected to the top of the tube. This allowed the predator and the prey to start the interaction.

A soundproof polystyrene lid was placed on the experimental arena to minimize disturbance and to eliminate observer effect. A high-speed video camera (Casio ex-fh20; 420 fps) recorded fish behavior through a hole in the lid and trials were filmed until the prey was consumed or 10 min had elapsed. The water in the experimental arena was changed following each trial to maintain oxygen saturation. SL (defined as the length of a fish measured from the tip of the snout to the posterior end of the last vertebra) of

Table 1. Mean (±SD) seawater parameters in the experimental system.

pH _{NBS}	temp (°C)	salinity (ppt)	TA (μmol.kg ⁻¹ SW)	pCO ₂ (μatm)
8.15 (0.04)	27.66 (0.98)	35	2269.66 (15.01)	440.53 (44.46)
7.89 (0.06)	27.74 (0.99)	35	2261.23 (14.92)	879.95 (140.64)

Temperature, pH salinity, and total alkalinity (TA) were measured directly. pCO₂ was estimated from these parameters using CO2SYS.

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the predator and prey and the water temperature were recorded for each trial.

In order to partition the relative effects of CO₂ on predators, prey and the full interaction, four combinations of CO₂-treated and control predators and prey were undertaken in a crossed design: control predator vs control prey (n = 21); treated predator vs treated prey (n = 21); control predator vs treated prey (n = 16); treated predator vs control prey (n = 16). Predator and prey fish were used only once in each trial. This ensured both were naïve to the experimental procedure. All combinations of treatments were undertaken daily to control for any potential daily variation. Video-analysis could not be performed on all interaction trials, due to fish leaving the screen at the time of the response and/or technical problems with the video. As a result, the sample size varied slightly among the performance traits measured. Trials were only used when the predator was at the opposite end of the tank to the prey at the start of the interaction. This was done to standardize for predator position. Maximum predator attack speed and maximum prey escape speed (UMAX_{pred} and UMAX_{prey}) were measured based on the center of mass (COM) of the fish when stretched straight based on Webb [34]. COM was assumed to be at 35% of the body length from the tip of the snout as it is the case for generalist fish [31]. Speed was smoothed using a 5-point differentiation-based moving polynomial regression [35]. Stage 1 and 2 were defined based upon directional changes of the anterior part of the body of the fish, based on Domenici and Blake [19].

Prey escape variables were measured only when prey performed a C-start. Predator attacks were measured only when a predator showed a fast-directed burst towards the prey (>3 body lengths s⁻¹). All variables with the exception of number of prey caught were measured using only the first attack that occurred within a trial. This was done to control for any anaerobic stress either the predator or prey may have experienced due to prolonged attacks.

Kinematic Analysis

Video recordings were analyzed using WinAnalyze motion-analysis software (v. 1.9 2D; Mikromak Service Brinkmann, Berlin, Germany). In each frame the snout and the COM (center of mass) were located on each fish. These points were chosen to standardise each frame.

The following performance variables were measured:

Predator

- 1) Capture success: percentage of trials in which the predator ingested the prey within the 10 min filming period, out of the total number of trials for each treatment.
- 2) Attack rate: number of attacks per unit time, measured for each interaction.
- 3) Predation rate: capture success divided by the number of attacks per unit time.
- 4) Predator attack distance (D_{pred}; m): the straight-line distance between the predator centre of mass (COM) at the time the attack commenced and the end of the attack. The end of the attack is defined as when the predator came to a halt.
- 5) Maximum predator attack speed (UMAX_{pred}; m s⁻¹): the top speed achieved at any point in time during the attack, based on the predator COM (see electronic supplemental material).

Prey

- 6) Prey reaction distance (R_D; m): the distance between the prey COM and the tip of the predator's snout at the onset of the escape response to a predator attack.
- 7) Apparent Looming Threshold (ALT) is defined as the apparent looming threshold for prey avoidance responses to a predatory strike and is a measure of the magnitude of the prey's response to the perceived threat of predation. The higher the perceived threat, the higher the angle, (ALT; radians s⁻¹): measured at the onset of the escape response and measured as the rate of change of the angle (α) subtended by the predator's frontal profile as seen by the prey. Previous work has shown that fish tend to react to an approaching stimulus (a predator) when a given threshold of dα/dt (i.e. ALT) is reached. ALT is calculated as (4US)/(4D²+S²), based on Dill [36] and Webb [37]. where U = predator speed, calculated as the speed of the predator in the frame prior to the prey's response; S = (Max. Depth+Max. Width)/2, where Max. Depth (DMAX) was estimated to be positioned at one-quarter of the body length of the predator (L_{pred}) (pers. obs.) and maximum width (WMAX) at 0.25L_{pred}; D = R_D +0.25L_{pred}.
- 8) Prey escape distance (D_{prey}; m): the straight-line distance between the prey COM at the onset of the escape response and at the end of the escape response (i.e. when the prey comes to a halt).
- 9) Maximum prey escape speed (UMAX_{prey}; m s⁻¹): the top speed achieved at any point in time during the escape response, measured using the prey COM.
- 10) Mean prey escape distance during stage 1 & 2 (Ds1s2; m): the distance between the COM of the fish at frame 0 and 24 ms later. This fixed duration was based on the average duration (22.8±3.2 ms), of the first two tail flips of the tail (the first two axial bends, i.e. stages 1 and 2 defined based on [19], which is the period considered crucial for avoiding ambush predator attacks [19,34]).

Statistical Analyses

To test the null hypothesis that feeding success is independent of predator and prey CO₂ exposure, predator success was compared by 2×4 contingency table analysis. The effects of CO₂ elevation on predator-prey interactions were examined using a 2-factor MANOVA with the fixed factors: Prey treatment (Control and CO₂ elevated) and Predator treatment (Control and CO₂ elevated). Univariate 2-factor ANOVAs with Tukey's HSD post-hoc tests were performed to determine the nature of any differences found by MANOVA. Predation rate data was arcsine transformed to meet the assumption of homogeneity of variance. Residual analysis indicated that data met the assumptions of normality and homogeneity of variance.

Results

Capture success was significantly associated with the CO₂ exposure treatment experienced by the predator and prey (χ² = 8.95, df = 3, p = 0.03; Fig. 1a). When both the predator and prey had been exposed to elevated CO₂ the capture rate (52%) was similar to that of the control group where both predators and prey were exposed to present-day CO₂ (51%). In contrast, the capture rate of these two treatment groups was markedly higher than the CO₂ predator-Control prey group. When predators exposed to elevated CO₂ were given prey exposed

to present-day CO₂, capture success of the predator was 33% less than the treatment groups where both predators and prey had been exposed to the same levels of CO₂ (either elevated or present-day CO₂). Capture success was also 14% less for predators exposed to present-day CO₂ with elevated CO₂ exposed prey compared with treatment groups where both predators and prey had been exposed to the same levels of CO₂.

The 2-factor MANOVA revealed a significant interaction between the effects of elevated CO₂ on the performance characteristics of predators and prey (Pillai's trace $(3,52) = 7.50$, $p < 0.0001$). ANOVA detected a significant interaction in the CO₂ levels of predators and prey for six of the tested behavioral

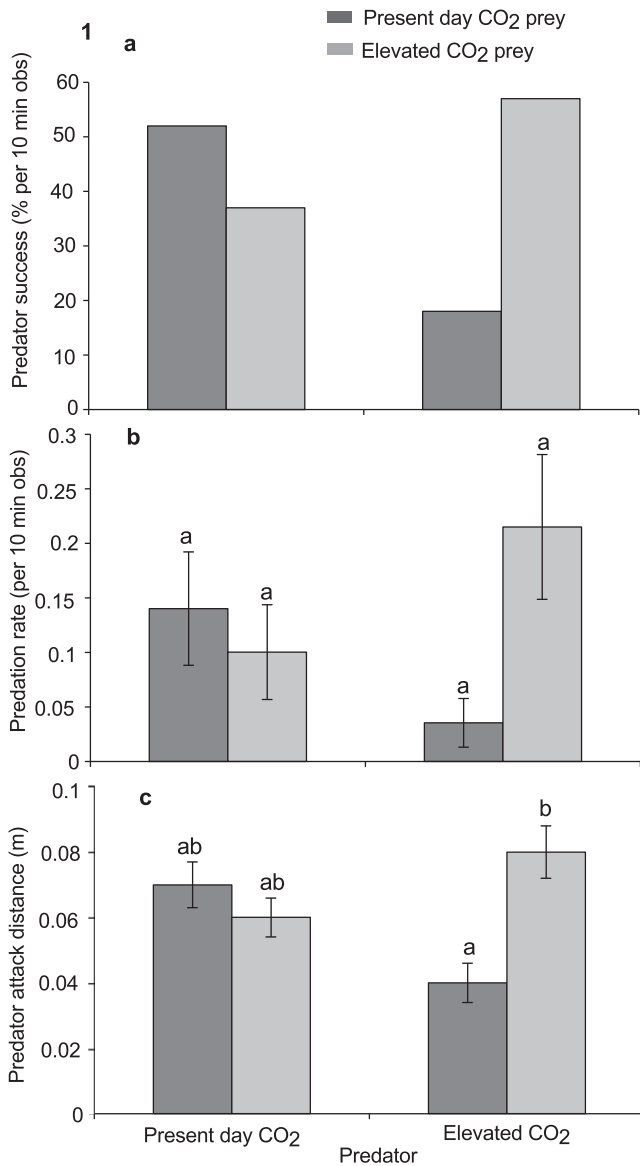


Figure 1. Effects of elevated CO₂ on predator attack performance. Comparison of the effects of elevated CO₂ (440, 880 μ atm) on interactions between a predator (*Pseudochromis fuscus*) and prey (*Pomacentrus amboinensis*) on 3 performance variables of the predator: (a) predator success (b) predation rate and (c) predator attack distance. (a) N = 21, 16, 16, 21, (b) N = 21, 16, 16, 21 and (c) N = 17, 15, 12, 16 (left to right). Errors are standard errors. Letters above bars represent Tukey's HSD groupings of means.

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Table 2. Comparison of the effects of elevated CO₂ (440, 880 μ atm) on interactions between a predator (*Pseudochromis fuscus*) and prey (*Pomacentrus amboinensis*) on 6 performance variables: (a) attack rate (b) predation rate (c) predator attack distance (d) prey reaction distance (e) ALT and (f) prey escape distance.

behavior	source of variation	df	MS	F	P
(a) attack rate	predator	1	0.0042	0.143	0.705
	prey	1	0.0070	0.238	0.626
	predator*prey	1	0.1427	4.845	0.030
	error	70	0.0294		
(b) predation rate	predator	1	0.0049	0.464	0.829
	prey	1	0.1459	1.358	0.247
	predator*prey	1	0.4387	4.083	0.020
	error	67	0.1074		
(c) predator attack speed	predator	1	2017	0.213	0.646
	prey	1	38512	4.070	0.048
	predator*prey	1	61256	6.474	0.013
	error	56	9462		
(d) prey reaction distance	predator	1	685.5	0.422	0.519
	prey	1	8898.2	5.471	0.022
	predator*prey	1	16555.2	10.179	0.002
	error	56	1626.4		
(e) ALT	predator	1	228	0.011	0.916
	prey	1	12548	0.616	0.435
	predator*prey	1	313294	15.396	<0.001
	error	54	20349		
(f) prey escape distance	predator	1	57241	6.365	0.014
	prey	1	40388	4.491	0.038
	predator*prey	1	69008	7.674	0.007
	error	56	0.4445		

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attributes: attack rate, predation rate, predator attack distance, reaction distance, apparent looming threshold and prey escape distance (see Table 2).

Elevated CO₂ significantly affected predator attack rates, however, this was influenced by prey CO₂ exposure (Table 2). There was no difference in attack rate when predators and prey were exposed to the same CO₂ treatments. However, when predators were exposed to elevated CO₂, they displayed a significantly decreased attack rate against prey that had not been exposed to elevated CO₂. Similarly, predators exposed to present-day CO₂ levels had a reduced attack rate when paired with prey that had been exposed to elevated CO₂ levels.

Exposure to elevated CO₂ significantly affected predation rates (Table 2; Fig. 1b). For predators, exposure to elevated CO₂ resulted in a significant reduction in the predation rates of prey that had not been exposed to elevated CO₂. Post-hoc tests yielded a border line p-value ($p = 0.05$) for the comparison between the elevated CO₂ exposed predators that interacted with present-day CO₂ exposed prey and the elevated CO₂ exposed predators that interacted with similarly exposed prey. Furthermore, when

predators exposed to elevated CO₂ were paired with similarly exposed prey, there was a significant increase in predation rates.

Elevated CO₂ significantly affected predator attack distance, but the nature of the effect differed according to whether prey had been exposed to elevated CO₂ or not (Table 2; Fig. 1c). There was no effect of CO₂ on attack distance when prey encountered control predators, however, when predators had been exposed to elevated CO₂ they displayed significantly increased attack distances against prey that had also been exposed to elevated CO₂ compared to control prey. The reaction distance of the prey to the predator at the onset of the first attack was affected by the CO₂ treatment of the prey and predator (Table 2; Fig. 2a). Prey exposed to elevated CO₂ allowed CO₂ exposed predators to get closer to them before undertaking an escape response. Max predator speed was found not to differ between treatments.

Apparent looming threshold (ALT) was significantly affected following exposure to elevated CO₂. When prey that had been exposed to elevated CO₂ were paired with similarly exposed predators, ALT increased significantly. In contrast, when control prey were paired with CO₂ exposed predators, ALT decreased substantially. There was no significant difference in ALT between the crossed trials (Table 2; Fig. 2b).

Prey escape distance was also found to be significantly affected following exposure to elevated CO₂ (Table 2; Fig. 2c) with prey exposed to present-day CO₂ having the highest escape distance compared to the other three treatment combinations.

Prey escape speed during stage 1 and 2 and maximum prey escape speed were not influenced by the exposure of either prey or predator to elevated CO₂ and no interactions between predator and prey treatments were found.

Discussion

Although recent studies have demonstrated that exposure to elevated CO₂ significantly increases prey mortality rates [9,28,38], the effects of elevated CO₂ on the kinematics at the basis of predator-prey interactions has not been investigated. Here, we demonstrated that CO₂ levels that may occur in the surface oceans by the end of the century impact on both the kinematics and the timing of predator-prey interactions. More specifically, prey treated with elevated CO₂ showed changes in escape performance, such as shorter reaction distances, reduced escape distances and changes in apparent looming threshold (ALT). Because CO₂ exposed prey were closer to the predator at the onset of their escape reaction and they swam shorter distances, elevated CO₂ had a clear negative effect on the reactivity and locomotion performance of the prey. The fast kinematics of the escape responses are likely to be under the control of Mauthner cells (although control by other reticulospinal cells cannot be ruled out [39]), which are triggered as a reaction to the fast approach of a predator. It is therefore possible that increased CO₂ levels may have an intrinsic effect on the sensory performance and neural control by the Mauthner cells or other reticulospinal neurons at the basis of the escape response, leading to increased prey vulnerability and ultimately increased mortality. Because escape distance is affected by CO₂, it is possible that CO₂ affects mainly the motivational component of the motor response, which acts upon the duration of the burst. Nevertheless, the effects of CO₂ on the onset of the response (i.e. reaction distance), suggest that the sensori-motor performance and the timing of the Mauthner cell's firing are also affected. Previous work has demonstrated that the regulation of plasma and cellular HCO₃⁻ and Cl⁻ following exposure to high CO₂ may lead to the excitation of GABA-A receptors [18]. Given that GABA-A receptors are found through-

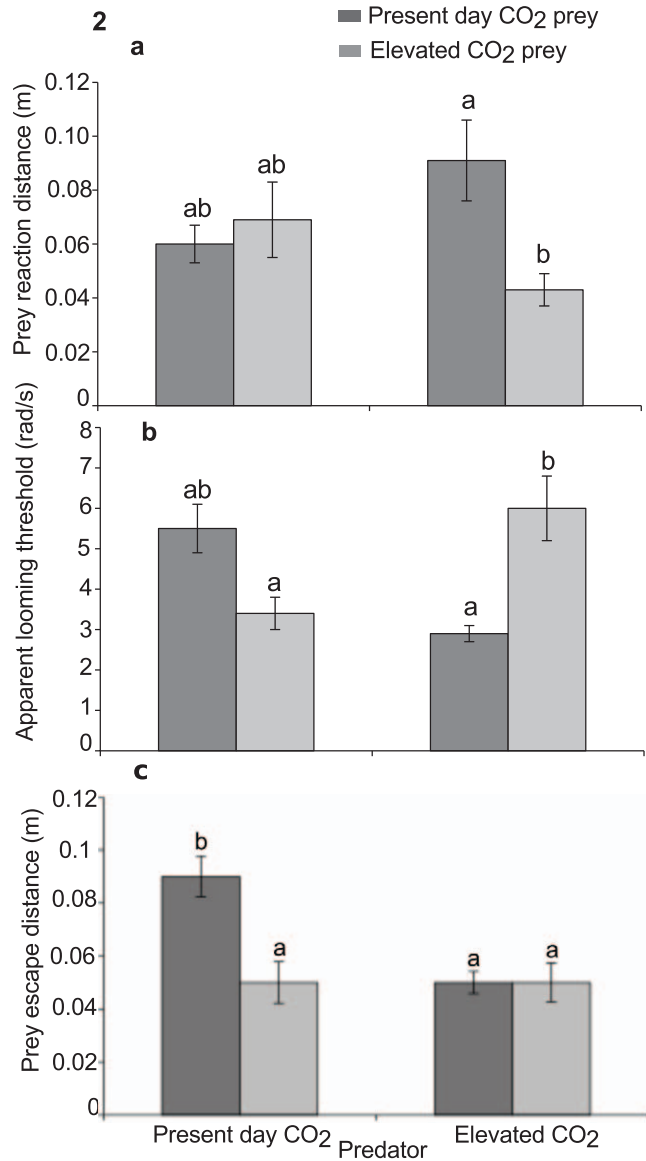


Figure 2. Effects of elevated CO₂ on prey escape performance. Comparison of the effects of elevated CO₂ (440, 880 μatm) on interactions between a predator (*Pseudochromis fuscus*) and prey (*Pomacentrus amboinensis*) on 3 performance variables of the prey: (a) prey reaction distance (b) apparent looming threshold (ALT) and (c) prey escape distance. (a) n = 19, 14, 11, 19 (b) N = 17, 14, 11, 16 and (c) N = 17, 14, 11, 18 (left to right). Errors are standard errors. Letters above bars represent Tukey's HSD groupings of means. doi:10.1371/journal.pone.0058520.g002

out the Mauthner neuron [40], these results suggest that high CO₂ interferes with GABA-A receptor function resulting in the misfiring of action potentials. Therefore, it is possible that this interference with brain neurotransmitters may be responsible for the changes observed. These results are consistent with the increased mortality rates observed in CO₂-exposed prey [9,28] and suggest that higher vulnerability to predation may be caused by a combination of changes in escape performance and other behavioural traits, such as increased activity levels and distance from shelter.

The results for the predators are complex to interpret, because the overall predation rate is the result of both predator

performance during the attack and its motivation to attack. Attack rates and attack distances were affected by elevated CO₂, with the lowest attack rates and distances occurring when predators and prey had experienced different CO₂ histories. This demonstrates that high CO₂ has an effect on predator performance during a predator-prey encounter, however the extent of this effect appears to be dependent upon the extent to which prey are affected by high CO₂. Predatory attacks require not only high speed but also precision to aim at the prey. The neural mechanisms that control predatory attacks are poorly understood, although it has been suggested that in some cases they may also be controlled by Mauthner cells [41]. It is possible that future levels of high CO₂ will impact on the complex circuitry needed to carry out attack processes. This may explain why we see behavioural changes that result in either decreased or increased capture success when predators and prey are exposed to different CO₂ histories, but when both are treated similarly, these changes appear to 'level out' resulting in no change in overall capture success compared with controls. Similar results have been suggested in estuarine systems involving crustaceans and molluscs, where for individual species there was a negative effect following CO₂ exposure, but these effects were not manifest at the community level (i.e. within the predator-prey interaction) [42]. This suggests that elevated CO₂ will have a marked impact on the dynamics and outcome of predator-prey encounters, but the population and community-level effects are likely to be dependent on the sensitivity of species to elevated CO₂.

This is the first study to examine performance-based attributes of both predators and prey under a high CO₂ environment within the context of a predator-prey interaction. While our findings are specific to the species used, the fact that the behavior of both predator and prey were affected is strongly suggestive that elevated CO₂ will affect the behavior of other predator and prey fish. Whether in other cases the increase in CO₂ will provide an

advantage for the prey, the predator or neither, will depend on the extent to which the specific predators and prey individuals are tolerant to CO₂ changes. Recent studies have shown that effects of elevated CO₂ can differ markedly between species [9]. Moreover, the effects of elevated CO₂ are not necessarily restricted to single fish-fish interactions, but might be manifest through other trophic pathways.

Changes in seawater μCO_2 has been shown to impact fishes at all stages of development [10,43], with significant impacts on individual fitness occurring at the vulnerable juvenile stage. Alterations to the dynamics of regulating processes at this early life stage could have significant effects on the replenishment and sustainability of marine populations [28]. In addition to affecting the physiology and behavior of given species, elevated CO₂ is likely to influence interactions between species, including predator-prey encounters. This may have important ecological effects, such as changing the balance of an interaction in favor of the predator or the prey. This study highlights the importance of considering species interactions when making predictions concerning the response of communities to climate change [15,44].

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Author Contributions

Conceived and designed the experiments: BJMA PD MIM PLM. Performed the experiments: BJMA. Analyzed the data: BJMA PD. Contributed reagents/materials/analysis tools: BJMA PD MIM SAW PLM. Wrote the paper: BJMA PD MIM PLM.

References

- Lima SL, Dill LM (1990) Behavioral decisions made under the risk of predation: a review and prospectus. *Canadian Journal of Zoology* 68: 619–640.
- Bengtson S (2002) Origins and early evolution of predation. *Paleontological Society Papers* 8: 289–318.
- Planes S, Romans P (2004) Evidence of genetic selection for growth in new recruits of a marine fish. *Molecular Ecology* 13: 2049–2060.
- Meinshausen M, Smith SJ, Calvin K, Daniel JS, Kainuma MLT, et al. (2011) The RCP greenhouse gas concentrations and their extensions from 1765 to 2300. *Climatic Change* 109: 213–241. doi:10.1007/s10584-011-0156-z.
- Bibby R, Cleall-Harding P, Rundle S, Widdicombe S, Spicer J (2007) Ocean acidification disrupts induced defences in the intertidal gastropod *Littorina littorea*. *Biology Letters* 3: 699–701. doi:10.1098/rsbl.2007.0457.
- Dixson DL, Munday PL, Jones GP (2010) Ocean acidification disrupts the innate ability of fish to detect predator olfactory cues. *Ecology Letters* 13: 68–75.
- Munday PL, Dixson DL, McCormick MI, Meekan M, Ferrari MCO, et al. (2010) Replenishment of fish populations is threatened by ocean acidification. *Proceedings of the National Academy of Sciences* 107: 12930–12934. doi:10.1073/pnas.1004519107.
- de la Haye KL, Spicer JI, Widdicombe S, Briffa M (2011) Reduced sea water pH disrupts resource assessment and decision making in the hermit crab *Pagurus bernhardus*. *Animal Behaviour* 82: 495–501.
- Ferrari MCO, Dixson DL, Munday PL, McCormick MI, Meekan MG, et al. (2011) Intrageneric variation in antipredator responses of coral reef fishes affected by ocean acidification: implications for climate change projections on marine communities. *Global Change Biology* 17: 2980–2986. doi:10.1111/j.1365-2486.2011.02439.x.
- Briffa M, de la Haye K, Munday PL (2012) High CO₂ and marine animal behaviour: Potential mechanisms and ecological consequences. *Marine Pollution Bulletin* 64: 1519–1528. doi:10.1016/j.marpolbul.2012.05.032.
- Harley CDG, Randall Hughes A, Hultgren KM, Miner BG, Sorte CJB, et al. (2006) The impacts of climate change in coastal marine systems. *Ecology Letters* 9: 228–241. doi:10.1111/j.1461-0248.2005.00871.x.
- Wisenden BD (2012) Cognitive dysfunction and risk assessment by prey: predictable changes in global climate have unpredictable effects. *Functional Ecology* doi:10.1111/j.1365-2435.2011.01956.x FE SPOTLIGHT.
- Fabry VJ, Seibel BA, Feely RA, Orr JC (2008) Impacts of ocean acidification on marine fauna and ecosystem processes. *ICES Journal of Marine Science: Journal du Conseil* 65: 414–432.
- Doherty PJ, Dufour V, Galzin R, Hixon MA, Meekan MG, et al. (2004) High mortality during settlement is a population bottleneck for a tropical surgeonfish. *Ecology* 85: 2422–2428.
- Cripps IL, Munday PL, McCormick MI (2011) Ocean Acidification Affects Prey Detection by a Predatory Reef Fish. *PLoS One* 6: e22736.
- Simpson SD, Munday PL, Wittenrich ML, Manassa R, Dixson DL, et al. (2011) Ocean acidification erodes crucial auditory behaviour in a marine fish. *Biology Letters* 7: 917–920. doi:10.1098/rsbl.2011.0293.
- Domenici P, Allan B, McCormick MI, Munday PL (2011) Elevated carbon dioxide affects behavioural lateralization in a coral reef fish. *Biology Letters* 8: 78–81. (doi:10.1098/rsbl.2011.0591).
- Nilsson GE, Dixson DL, Domenici P, McCormick MI, Sørensen C, et al. (2012) Near-future carbon dioxide levels alter fish behaviour by interfering with neurotransmitter function. *Nature Climate Change* 2: 201–204. doi:10.1038/nclimate1352.
- Domenici P, Blake RW (1997) The kinematics and performance of fish fast-start swimming. *Journal of Experimental Biology* 200: 1165–1178.
- Walker JA, Ghalambor CK, Griscti OL, McKenny D, Reznick DN (2005) Do faster starts increase the probability of evading predators? *Functional Ecology* 19: 808–815. doi:10.1111/j.1365-2435.2005.01033.x.
- Langerhans RB, Reznick DN (2009) Ecology and evolution of swimming performance in fishes: predicting evolution with biomechanics. *Fish locomotion: an etho-ecological perspective* Enfield: Science Publishers: 200–248.
- Stewart BD, Jones GP (2001) Associations between the abundance of piscivorous fishes and their prey on coral reefs: implications for prey-fish mortality. *Marine Biology* 138: 383–397.
- Munday PL, Eyre PJ, Jones GP (2003) Ecological mechanisms for coexistence of colour polymorphism in a coral-reef fish: an experimental evaluation. *Oecologia* 137: 519–526. doi:10.1007/s00442-003-1356-7.
- Holmes T, McCormick M (2010) Size-selectivity of predatory reef fish on juvenile prey. *Marine Ecology Progress Series* 399: 273–283. doi:10.3354/meps08337.

25. Messmer V, van Herwerden L, Munday PL, Jones GP (2005) Phylogeography of colour polymorphism in the coral reef fish *Pseudochromis fuscus*, from Papua New Guinea and the Great Barrier Reef. *Coral Reefs* 24: 392–402. doi:10.1007/s00338-005-0001-9.
26. Kerrigan BA (1996) Temporal patterns in size and condition at settlement in two tropical reef fishes (Pomacentridae: *Pomacentrus amboinensis* and *P. nagasakiensis*). *Marine ecology progress series* 135: 27–41.
27. Meekan M.G., Wilson S.G., Halford, A. & Retzel, A. (2001) A comparison of catches of fishes and invertebrates by two light trap designs, in tropical NW Australia. *Marine Biology* 139: 373–381. doi:10.1007/s002270100577.
28. Munday PL, Dixon DL, McCormick MI, Meekan M, Ferrari MCO, et al. (2010) Replenishment of fish populations is threatened by ocean acidification. *Proceedings of the National Academy of Sciences* 107: 12930–12934. doi:10.1073/pnas.1004519107.
29. Munday PL, Wilson SK (1997) Comparative efficacy of clove oil and other chemicals in anaesthetization of *Pomacentrus amboinensis*, a coral reef fish. *Journal of Fish Biology* 51: 931–938.
30. Sweatman HPA (1984) A field study of the predatory behavior and feeding rate of a piscivorous coral reef fish, the lizardfish *Synodus leuognathus*. *Copeia*: 187–194.
31. Pierrot D, Lewis E, Wallace DWR (2006) MS Excel Program Developed for CO₂ System Calculations. ORNL/CDIAC-105a. Oak Ridge, Tennessee: Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory, U.S. Department of Energy.
32. Mehrbach C, Culbertson CH, Hawley JE, Pytkowicz RM (1973) Measurement of the apparent dissociation constants of carbonic acid in seawater at atmospheric pressure. *Limnology and Oceanography*: 897–907.
33. Dickson AG, Millero FJ (1987) A comparison of the equilibrium constants for the dissociation of carbonic acid in seawater media. *Deep Sea Research Part A Oceanographic Research Papers* 34: 1733–1743. doi:10.1016/0198-0149(87)90021-5.
34. Webb PW (1976) The effect of size on the fast-start performance of rainbow trout *Salmo gairdneri*, and a consideration of piscivorous predator-prey interactions. *Journal of Experimental Biology* 65: 157–177.
35. Lancelotti C (1956) *Applied analysis*, Prentice Hall, Eaglewood Cliffs, New Jersey.
36. Dill LM (1974) The escape response of the zebra danio (*Brachydanio rerio*) I. The stimulus for escape. *Animal Behaviour*, 22, 711–722.
37. Webb PW (1982) Avoidance responses of fathead minnow to strikes by four teleost predators. *Journal of Comparative Physiology A: Neuroethology, Sensory, Neural, and Behavioral Physiology* 147: 371–378.
38. Ferrari MCO, McCormick MI, Munday PL, Meekan MG, Dixon DL, et al. (2011) Putting prey and predator into the CO₂ equation—qualitative and quantitative effects of ocean acidification on predator–prey interactions. *Ecology letters*. 14: 1143–1148.
39. Kohashi T, Oda Y (2008) Initiation of Mauthner-or non-Mauthner-mediated fast escape evoked by different modes of sensory input. *The Journal of Neuroscience* 28: 10641–10653.
40. Diamond J, Huxley AF (1968) The activation and distribution of GABA and L-glutamate receptors on goldfish Mauthner neurones: an analysis of dendritic remote inhibition. *The Journal of Physiology* 194: 669–723.
41. Wöhl S, Schuster S (2007) The predictive start of hunting archer fish: a flexible and precise motor pattern performed with the kinematics of an escape C-start. *Journal of Experimental Biology* 210: 311–324.
42. Landes A, Zimmer M (2012) Acidification and warming affect both a calcifying predator and prey, but not their interaction. *Marine Ecology Progress Series* 450: 1–10. doi:10.3354/meps09666.
43. Munday P.L., McCormick M.I., Nilsson G.E. (2012) Impact of global warming and rising CO₂ on coral reef fishes: what hope for the future? *Journal of Experimental Biology*. In press.
44. Gilman SE, Urban MC, Tewksbury J, Gilchrist GW, Holt RD (2010) A framework for community interactions under climate change. *Trends in Ecology & Evolution* 25: 325–331. doi:10.1016/j.tree.2010.03.002.