

Effect of temperature variation in *Agarophyton chilensis*: contrasting the response of natural and farmed populations

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Abstract

During the domestication process, farmers influence the reproduction and care of organisms to ensure a predictable supply of the resource of interest, causing changes in phenotypic and genotypic character frequencies. In Chile, as a result of unconscious selection and domestication process, farmed populations of the red alga *Agarophyton chilensis* have most likely undergone a reduction in genetic diversity and a modification in life-history traits compared to wild populations. In order to understand the implications that these processes may have in *A. chilensis*, we investigated how temperature variations (10 °C, 15 °C, and 20 °C) affect growth and photosynthetic responses of natural and farmed populations from three different localities along the Chilean coast. Natural population's growth decreased at low and high temperature levels while all three farmed populations respond in a very similar way to temperature variation. We propose that a possible outcome of farming, in the *A. chilensis*, vegetatively propagated crops, could have been the selection of general-purpose-genotypes able to perform adequately across the range of temperature tested in our experiment. Furthermore, our results showed that photosynthetic activity was also affected by temperature treatments (e.g., different maximum maximal electron transport rate and quantum yield values depending on the population type and temperature). In a context of climate change, *A. chilensis* farmed populations may be better able to cope with impacts of anthropogenic activities than natural populations due to the buffer effect of their general-purpose-genotypes, tolerant to a wide range of conditions.

 $\label{eq:constraint} \begin{array}{l} \mbox{Keywords} \ \mbox{Rhodophyta} \ \cdot \mbox{Abiotic factor} \ \cdot \ \mbox{Domestication} \ \cdot \ \mbox{Management} \ \cdot \ \mbox{General-purpose-genotypes} \ \cdot \ \mbox{Photosynthesis} \ \cdot \ \mbox{Origin} \ \cdot \ \mbox{Seaweed} \ \cdot \ \mbox{Selection} \end{array}$

Introduction

Domestication is considered a long and complex process during which domesticators influence the reproduction and care of domesticated species to guarantee predictable supply of

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resources presenting selected traits of interest for human use (Zeder 2015). This selection process generates changes in phenotypic and genotypic character frequencies of cultivated populations (Zohary 1984). Even if domestication of marine species is much more recent than the one of terrestrial animals and plants (Duarte et al. 2007), strong evidence for domestication has been found in a few cultivated seaweeds (Valero et al. 2017). As for terrestrial plants (Meyer et al. 2012), some domesticated seaweeds are characterized by a shift in their reproductive strategy (e.g., changes from sexual reproduction to vegetative propagation) between natural and farmed populations (Valero et al. 2017). This shift in reproductive strategy has been demonstrated for Agarophyton (referred as Gracilaria in Guillemin et al. 2008) and is probably also present in Kappaphycus (Ask and Azanza 2002). Asexual propagation enables farmers to selectively multiply superior genotypes and maintain desired phenotypes through vegetative propagation (Valero et al. 2017).

In Chile, intensive seaweed farming is limited to the domesticated red alga *Agarophyton chilensis* used mainly for agar extraction (Buschmann et al. 2017). Even if the domestication

process of this species has begun only a few decades ago, the almost complete predominance of diploid individuals in farms demonstrate that farming practices had significantly modified life-history traits as compared to wild populations (Guillemin et al. 2008). Moreover, recent investigations have demonstrated that this red alga colonized the Chilean coast from New Zealand, likely at the end of the Last Glacial Maximum (Guillemin et al. 2014). The lower genetic diversity of the Chilean populations, when compared to the ones from New Zealand, is indeed consistent with a genetic bottleneck resulting from a transpacific range extension and was probably reinforced by the overexploitation of natural Chilean populations during the 1990s (Guillemin et al. 2014). In Chile, active transport and exchange of inoculums of A. chilensis between coastal communities of fishermens for cultivation purposes have contributed to the artificial expansion of the species distribution. Nowadays, natural populations are distributed between 30°S to 45°S while farming extends further north, up to Antofagasta, 17°S (Bird et al. 1986; Guillemin et al. 2008). Human activities have also led to a loss of genotypic diversity in farmed populations that could be partially linked to involuntary selection of faster growing thalli, during the first steps of the domestication process (Guillemin et al. 2008; 2013; Valero et al. 2017).

Reduced genetic diversity can severely affect the ability of populations to resist pests and pathogens and limit the scope for future genetic improvement in domesticated crops (Robinson et al. 2013). Besides, it has been reported that a reduction in genetic diversity could lead to lower growth and resilience in highly stressful and/or variable environment (Simms 2000). However, to date, few studies have focused on the potential ecophysiological differences between natural and farmed populations of A. chilensis. Gallegos-Sánchez et al. (2018) detected a significant and negative effect of low salinity conditions on thalli sampled from both natural and farmed A. chilensis populations but with farmed population's thalli being less affected than the natural ones. Results suggested that farmed populations might be more tolerant to salt stress than wild ones in this species and the authors proposed that this difference between population types could be due to previous selective process carried out by farmers.

Considering the possible consequences of unconscious selection (as defined in Zohary 2004) and domestication on *A. chilensis*, we propose that natural populations will be less sensitive to temperature variations than farmed populations. Genotypic diversity has been shown to be higher in natural than farmed *A. chilensis* Chilean populations, and we propose that the farms will show less effective mechanisms of acclimation (e.g., enhancing of photosynthetic performance) than the one observed in natural populations when confronted with temperature considered as high in their natural environment (i.e., water of the southern coast of Chile do not generally reach 20 °C; Westermeier et al. 1993). The aim of the present study was to assess the effect of temperature on growth and photosynthetic responses of both natural and farmed stands of *A. chilensis* and tetrasporophyte thalli sampled from three sites along the Chilean coast that were followed up during 1 month in controlled laboratory conditions at 10 °C, 15 °C, and 20 °C.

Materials and methods

Study sites and life cycle phase determination of the sampled thalli

A total of 600 individuals were sampled from 3 natural and 3 farmed populations (i.e., 100 individuals in each population) during a spring season. One farmed and one natural population were sampled from three sites: Concepción, Maullín, and Ancud (Fig.1). The distinction between natural and farmed populations was first based on whether Agarophyton chilensis thalli were actively planted or not by farmers. As previously reported for the species (Guillemin et al. 2008), thalli were attached to small rocks and pebbles by a holdfast in the three natural A. chilensis beds while unattached thalli were found embedded in the sandy bottom in the three farmed stands. In each site, natural and farmed populations were separated by 1 km approximately. In natural beds, individuals sampled correspond to distinct holdfasts (i.e., distinct genotypes produced by sexual reproduction and spore settlement). In farms, to avoid sampling fragments of the same asexually propagated genotype, sampled thalli were separated by at least by 2 m. All the collected thalli were transported in isolated boxes to the CEACIMA hatchery (Centro de Investigación de Acuicultura y Ciencias del Mar, Universidad de Los Lagos) located in the Metri Bay (41°36'S, 72°43'W). Once in the hatchery, each thallus was cleaned with fresh water and all epiphytes were removed by hand. Thalli were individually marked with a numbered tag and maintained in 400 L tanks at 12 °C with constant aeration, 12 L:12D photoperiod, 20 µmol photons $m^{-2} s^{-1}$ photon flux and weekly filtered seawater exchange.

Collected thalli were observed under stereoscope microscope (Stemi DV4, Zeiss, Jena, Germany) to determine phase (i.e., diploid tetrasporophytes or haploid gametophytes) of mature individuals. For vegetative individuals, a 3-cm fragment of tissue was excised from each thallus, placed into plastic bags with silica gel for rapid dehydration. The sex markers available for *A. chilensis* were amplified following Guillemin et al. (2012) and the amplification products were visualized in 1.5% agarose gel (w/v) after adding 2 μ L of GelRed (Biotium, USA). Results were used to determine sex and phase of vegetative individuals. In order to prevent experimental bias due to ecophysiological variability between life cycle phases (Guillemin et al. 2013), only diploid tetrasporophytes were selected for our experiments.



Fig.1 Location of the three sites sampled along the Chilean coast. Photographs of farmed and natural populations studied are given for each site. All photographs by S. Usandizaga

Experimental design

The experimental design consisted of 90 2-L Erlenmeyer flasks (i.e., 6 populations of origin of thalli × 3 temperature treatments × 5 replicates per population per temperature treatment) arranged in 15 60-L plastic water tanks fitted with temperature control systems. Three temperature treatments were used: 10, 15, and 20 °C. These temperatures were chosen since they roughly represent the temperature range encountered in the field by the study species. Temperature conditions varied widely within sites were A. chilensis is found and between regions populated by the species along the Chilean coasts with values between 9 and 16 °C recorded in Maullín (Westermeier et al. 1993) and between 10 and 20 °C in areas further north such as Concepción, Coquimbo, and Antofagasta (Santelices and Ugarte 1990). In each 60-L plastic water tank, one 50 W automatic heater (Whale VK-1000, Regent) and one stainless steel thermometer (Hagen, Phelan) were used to maintain constant temperature. For each population under study, five replicates (i.e., five 2-L flasks) were followed per temperature treatment and eight thalli (5 cm length each), selected randomly from distinct tetrasporophytes, were placed in each 2-L flask. Thalli were selected without replacement from a pool of 40 tetrasporophytes, available for each population of origin (see above). Once a week, thalli were transferred to clean 2-L flasks with fresh Provasoli culture medium (McLachlan 1973). After 1 week of acclimation at 12 °C, the laboratory experiment was run for 30 days. All Erlenmeyer flasks were under constant conditions of aeration, photoperiod (12 h light:12 h dark) and photon flux (20 μ mol photons m⁻² s⁻¹).

Growth

Fresh weight of each thallus was assessed weekly on an analytical balance (Sartorius TE 313 DS, Germany) and the specific growth rate (SGR) was calculated as the percentage of wet weight gain per day according to the formula: SGR = $[\ln (W_f - W_i) / (t_f - t_i)] \times 100$; where W_i = initial fresh weight, W_f = final fresh weight, and t = time (days). Initial fresh weight in the 2-L flasks was of 0.12 ± 0.06 g, some variation in fresh weight exist between each 2-L flask at the beginning of the experiment since calibration of algal material was based on thallus length (see above).

Physiological variables

Thallus pieces were collected at the end of the experiment to measure rapid light curves (RLC). As an indicator of quantum efficiency and photoinhibition, we used, F_v/F_m , which was determined after incubation of 20 min of the thalli in darkness (Schreiber et al. 1995) with a Junior PAM (Walz GmbH, Germany). The electron transport rate (ETR, µmol electrons m⁻² s⁻¹) was determined after 20 s exposure in 12 increasing intensities of PAR (up to 1500 µmol photons m⁻² s⁻¹) provided by a blue light of the Junior PAM device. ETR was calculated according to Schreiber et al. (1995) as follows:

$$\text{ETR} = \Delta F / F'_{\text{m}} \cdot E \cdot A \cdot F_{\text{II}} (\mu \text{mol electrons m}^{-2} \cdot \text{s}^{-1}) \quad (1)$$

Where $\Delta F/F'_{m}$ is the effective quantum yield, E is the incident PAR (photosynthetically active radiation) irradiance expressed in μ mol photons m⁻² s⁻¹, A is the thallus absorptance and F_{II} is the fraction of chlorophyll related to photosystem II (PSII, 400-700 m), being 0.15 in red seaweed (Grzymski et al. 1997). As an estimator of photosynthetic efficiency, the initial slope of ETR (α_{ETR}) and maximum ETR (ETR_{max}) were obtained from the tangential function reported by Eilers and Peeters (1988) and the irradiance saturation (Ek_{ETR}) was calculated from the intercept between these two parameters. Representing potential thalli photoprotective mechanism, non-photochemical quenching (NPO) was measured according to Schreiber et al. (1995). The maximal nonphotochemical quenching (NPQmax) was calculated from the tangential function of NPQ versus irradiance function according to Eilers and Peeters (1988).

Statistical treatment

All analyses were performed in *R* (3.2.4 version) (Cayuela 2011). Assumptions of homogeneity of variances and normal distribution were tested using Levene's test and Shapiro-Wilk, respectively. When non-normal residuals and heteroscedasticity were detected, the data were transformed using logarithm (for *Ek* and SGR) or Box-cox (for F_v/F_m). The experimental design fitted a three-way ANOVA with treatment (temperature: 10 °C, 15 °C, and 20 °C), site of origin (Concepción, Maullín, and Ancud) and population type (natural or farmed) considered as fixed factors. Statistical differences between groups were analyzed using comparisons of means (Tukey's HSD). Significances were set at *p* < 0.05.

Results

Significant differences were detected in Agarophyton chilensis specific growth rate (SGR) after 30 days of experimentation between temperature treatments ($F_{(2, 72)} = 15.46$; P < 0.0001), between population types($F_{(1, 72)} = 4.93$; P =0.03) and a significant interaction was also detected between population type and site of origin ($F_{(2, 72)} = 5.00; P = 0.009$). In Maullín and Ancud, no significant differences between temperature treatments were detected for farmed thalli (Fig. 2C and E), while farmed thalli from Concepción showed a slightly but significantly lower SGR at 20 °C than at 15 °C $(7.26 \pm 1.48 \text{ and } 9.45 \pm 1.92 \text{ g} \cdot \text{day}^{-1}$, respectively; Fig.2A). In addition, differences among Concepción farmed and natural populations were observed (Tukey test; P < 0.05, see Fig. 2A and B). In contrast, SGR was significantly higher at 15 °C for thalli sampled in natural A. chilensis stands whatever the site under study (SGR at 15 °C; 7.49 ± 2.07 g·day⁻¹in Concepción; $8.81 \pm 2.60 \text{ g} \cdot \text{day}^{-1}$ in Maullín; and $8.67 \pm 0.84 \text{ g} \cdot \text{day}^{-1}$ in Ancud; Fig. 2B, D and F). Finally, the growth rate did not decrease significantly (Tukey test; P < 0.05) at the lower (10 °C) and the higher (20 °C) temperature tested for Maullin and Ancud populations (Fig. 2C, D, E and F).

Regarding the photosynthetic responses, significant effect of temperature treatments were detected for all parameters measured: the optimal quantum yield offluorescence (F_v/F_m) , the maximal electron transport rate (ETR_{max}), the saturation irradiance (E_k) , and the maximal non-photochemical quenching (NPQ_{max}) (see Table 1). Interactions between temperature treatment and population type were also observed for ETR_{max} and $E_{\rm k}$ (Table 1). Values of ETR_{max} and $E_{\rm k}$ were significantly higher for thalli sampled in the farmed population of Concepción and grown at 20 °C than for thalli sampled in the farmed population of Ancud and grown at 10 °C (18.63 \pm 12.76 and 1.44 \pm 0.37 μ mol electrons·m⁻²·s⁻¹, 442.97 ± 306.60 and 31.86 ± 4.78 μ mol photons $m^{-2} \cdot s^{-1}$, for ETR_{max} and E_k , respectively; Tukey's HSD tests; P < 0.05; Table 2). Significant interactions between temperature treatment, population type, and site of origin were observed for F_v/F_m and NPQ_{max} (Table 1). In Maullín at 15 °C, values of NPQ_{max} were significantly higher for thalli sampled in the farmed bed than the natural population (Tukey's HSD tests; P < 0.05; Table 2).

Discussion

The present study confirms that natural and farmed *Agarophyton chilensis* populations respond differentially to temperature variations. Indeed, contrarily to our expectations, our results indicate that farmed populations are less sensitive to temperature variations than natural ones (i.e., thalli growth rate are mostly similar at 10, 15, and 20 °C for farms while thalli from natural populations have a higher growth rate at 15 °C). In

Fig.2 Mean (±SE) of the specific growth rate (SGR) of Agarophyton chilensis thalli sampled from farmed and natural populations in Concepción (A and B), Maullín (C and D), and Ancud (E and F). Thalli were submitted to three temperature treatments (black bars, $T = 10 \,^{\circ}\text{C}$; light gray bars, T = 15 °C; and dark gray bars, $T = 20 \,^{\circ}\text{C}$). Values are given after 30 days of experiment. Different letters denote significant differences between temperature treatments (Tukey's hsd posthoc tests; P < 0.05; results given independently within each sampling site and temperature treatment)



Chile, farmed population has been under unconscious human selection pressure for, at least, three decades and we propose that a possible outcome could have been the selection of generalpurpose-genotypes (as in Baker 1974) in these vegetatively propagated crops. Our results also showed that photosynthetic activity was affected by temperature treatments (e.g., different maximum maximal electron transport rate and quantum yield values depending on the population type and temperature). Regardless of the origin of the natural population under study, our experiment shows that thalli specific growth rate (SGR) was higher at 15 °C than at lower (i.e., 10 °C) or higher (i.e., 20 °C) temperatures. Even if SGR tended to be much more homogeneous for farmed thalli at all temperatures, one slight difference was observed for the farmed thalli from Concepción that show significantly less growth at 20 °C. Supporting our results, a study realized by Santelices and

Tempera	uture trea	atmer	nt (1.1;	10 °C, 15	C, and	120 %	U), site	of ongin (S	; Conce	spciór	ı, Maı	ullin and	Ancuc	l), and _l	population ty	pe (P; ni	atural	or farmec	l) were c	onside	red as	tixed	factors			
	Tempe	ratur	e treatr	nent (T.T)	Popul	lation	type (P	(Site of (origin	1 (S)	Inte	eraction	ר × T) ר	(T.)	Interac	tion (S	(T.T)	Intera	ction ($(\mathbf{P} \times \mathbf{S})$		Interact	ion (T	T × P	\times S)
	SumSç	q df	ы	b	SumS	df df	Ц	d	SumSq	df 1		sur Sur	nSq d	ΓF	d	SumSq	df]	d	Sum	df df	Ь	d	SumSq	df I	Г ц	
$F_{\rm v}/F_{\rm m}$	0.00	7	13.20	< 0.01***	¢ 0.00	-	1.89	0.18	0.00	2	1.32 (.28 0.00	0	10.8	3 < 0.01***	0.00	4	2.87 0.04	;* 0.00	5	0.88	0.42	0.00	4	2.86	0.04*
ETR _{max}	326.2	0	6.17	$< 0.01^{**}$	52.9	-	2.00	0.17	1.61	5	0.03 (0.97 506	5 2	9.57	$< 0.01^{***}$	270	4	2.55 0.06	79.3	7	1.50	0.24	109.9	4	1.04	0.40
$E_{\rm k}$	6.34	0	6.97	$< 0.01^{**}$	0.16	-	0.35	0.56	1.86	0	2.04 ().14 4.80	6 2	5.34	< 0.01**	1.46	4	0.80 0.53	0.57	7	0.63	0.54	1.01	4	0.55	0.70
NPQmax	0.45	7	6.02	$< 0.01^{**}$	0.40	1	10.68	< 0.01** (0.01	2	0.16 (0.85 0.1	1 2	1.48	0.24	0.26	4	0.1€	0.23	2	3.10	0.06	0.49	4	3.31	0.02*

Results of ANOVA analyses on of the specific growth rate (SGR) and photosynthetic parameters for Agarophyton chilensis thalli submitted to three temperature treatments during 1 month.

Bold significant at *P < 0.05, **P < 0.01, ***P < 0.00

Table 1

Ugarte (1990) on A. chilensis (as Gracilaria chilensis) natural populations from Maullín, also revealed better growth rates at 15 °C than at 10 °C or 20 °C. Most temperate species of the genus Agarophyton has been shown to grow faster in temperature ranging between 15 and 20 °C (McLachlan and Bird 1984). A study performed in Gracilaria gracilis (as G. verrucosa) reported a slow growth rate at 10 °C and high mortality after a 14-day heat-wave in Saldanha Bay, South Africa (Engledow and Bolton 1992). Agarophyton chilensis is a temperate-water species with a distribution limited to southern part of the Pacific (Bird et al. 1986; Guillemin et al. 2008) and it is possible that the species present a metabolism with limited temperature tolerance.

Although farming has begun only a few decades ago (i.e., during the 1980s) in Chile, the predominant mechanism for stock propagation by cuttings used in A. chilensis farms has already significantly decreased their genotypic diversity when compared with wild populations (Guillemin et al. 2008). There is an overall agreement that genetically more variable populations may be associated with higher resilience, increased productivity and population growth rate as compared with less variable populations (Forsman 2014). However, contrasting with our expectations, farmed thalli in our experiment clearly show a strong ability to grow in contrasting environmental conditions, including the quite "extreme" temperature of 20 °C tested. In the same way, Gallegos-Sánchez et al. (2018) concluded that A. chilensis farmed populations may be less sensitive to salt stress and able to grow in a greater range of salinity than natural populations. One possible explanation for these results is that farmed populations of A. chilensis are composed mostly of general-purpose genotypes, able to grow in highly stressful and/or variable environments. General-purpose-genotypes are sometime also referred as "Jack-of-all-trades, master of none" since they are described as versatile genotypes that are able to perform adequately across a range of environments but are not superior in any of them. These general-purpose-genotypes can confer a species or population a broad tolerance to environmental changes and are often associated with species invasion (Baker 1974; Richards et al. 2006). In Chile, farms have been developed using material growing embedded in muddy estuaries and sandy bays. These habitats are typically highly heterogeneous and present strong seasonal variations in temperature and salinity (Westermeier et al. 1993; Buschmann et al. 1995). It have been demonstrated that intraspecific competition for resources utilization in clonal individuals living in habitats characterized by fluctuating environmental conditions could lead to positive selection of generalpurpose-genotypes (Arnaud-Haond et al. 2012).

In benthic algae, temperature variations affect photosynthetic metabolism (Davison 1991) determining, for example, seasonal distribution (De Nicola 1996). However, habitats characterized by strong spatial and temporal variations of
 Table 2
 Photosynthetic parameters measured in Agarophyton chilensis

 natural and farmed thalli submitted to three temperature treatments. Thalli
 from one natural and one farmed populations were sampled in three sites

(i.e., Concepción, Maullín, and Ancud). Data are given as mean \pm S.E. (n = 5). Distinct uppercase letters denote significant differences after Tukey test

Concepció	n			Maullín			Ancud		
	Т	Р	Mean±S.E.	Т	Р	Mean±S.E.	Т	Р	Mean±S.E.
$F_{\rm v}/F_{\rm m}$	10 °C	Natural	0.60 ± 0.01^{ab}	10 °C	Natural	$0.64\pm0.03^{\rm ac}$	10 °C	Natural	0.66 ± 0.01^{ac}
	15 °C		0.64 ± 0.02^{abc}	15 °C		0.37 ± 0.36^{b}	15 °C		0.64 ± 0.02^{abc}
	20 °C		0.66 ± 0.02^{ac}	20 °C		0.66 ± 0.01^{ac}	20 °C		0.66 ± 0.01^{abc}
	10 °C	Farmed	$0.68\pm0.02^{\rm c}$	10 °C	Farmed	$0.69\pm0.00^{\rm c}$	10 °C	Farmed	0.66 ± 0.01^{ac}
	15 °C		0.61 ± 0.02^{ab}	15 °C		0.61 ± 0.02^{ab}	15 °C		0.64 ± 0.02^{abc}
	20 °C		0.64 ± 0.03^{abc}	20 °C		0.63 ± 0.01^{abc}	20 °C		0.65 ± 0.02^{abc}
ETR _{max}	10 °C	Natural	5.33 ± 0.39^{ab}	10 °C	Natural	5.94 ± 3.28^{ab}	10 °C	Natural	3.06 ± 1.19^{ab}
	15 °C		9.52 ± 5.54^{ab}	15 °C		2.98 ± 2.98^{ab}	15 °C		17.48 ± 5.61^{ab}
	20 °C		4.46 ± 3.02^{ab}	20 °C		4.29 ± 1.64^{ab}	20 °C		4.16 ± 2.06^{ab}
	10 °C	Farmed	4.51 ± 3.97^{ab}	10 °C	Farmed	3.73 ± 1.83^{ab}	10 °C	Farmed	1.44 ± 0.37^b
	15 °C		3.97 ± 0.68^{ab}	15 °C		8.25 ± 3.32^{ab}	15 °C		9.21 ± 5.67^{ab}
	20 °C		18.63 ± 12.76^{a}	20 °C		16.11 ± 8.19^{ab}	20 °C		10.03 ± 8.65^{ab}
$E_{\mathbf{k}}$	10 °C	Natural	143.97 ± 53.30^{ab}	10 °C	Natural	129.40 ± 88.27^{ab}	10 °C	Natural	73.94 ± 45.65^{ab}
	15 °C		185.94 ± 70.68^{ab}	15 °C		111.63 ± 58.52^{ab}	15 °C		229.16 ± 92.04^{ab}
	20 °C		200.16 ± 196.5^{ab}	20 °C		111.39 ± 48.50^{ab}	20 °C		124.41 ± 91.02^{ab}
	10 °C	Farmed	94.85 ± 87.88^{ab}	10 °C	Farmed	66.76 ± 32.14^{ab}	10 °C	Farmed	31.86 ± 4.78^{b}
	15 °C		93.56 ± 21.17^{ab}	15 °C		173.11 ± 32.20^{ab}	15 °C		110.38 ± 70.53^{ab}
	20 °C		442.97 ± 306.60^{a}	20 °C		233.80 ± 108.15^{ab}	20 °C		184.44 ± 134.12^{ab}
NPQ _{max}	10 °C	Natural	0.74 ± 0.03^{ab}	10 °C	Natural	0.68 ± 0.14^{ab}	10 °C	Natural	0.71 ± 0.09^{ab}
	15 °C		0.94 ± 0.28^{ab}	15 °C		0.42 ± 0.50^a	15 °C		1.03 ± 0.27^{ab}
	20 °C		0.64 ± 0.14^{ab}	20 °C		0.58 ± 0.10^{ab}	20 °C		0.55 ± 0.20^a
	10 °C	Farmed	1.56 ± 0.92^{ab}	10 °C	Farmed	0.66 ± 0.13^{ab}	10 °C	Farmed	0.66 ± 0.04^{ab}
	15 °C		0.93 ± 0.12^{ab}	15 °C		1.16 ± 0.23^{b}	15 °C		0.99 ± 0.30^{ab}
	20 °C		0.69 ± 0.09^{ab}	20 °C		1.04 ± 0.23^{ab}	20 °C		0.86 ± 0.10^{ab}

abiotic factors, request constant adjustment of photosynthetic processes in species populating them (Ensminger et al. 2001). In our study, the highest values of ETR_{max} and E_k were observed in the A. chilensis farmed population from Concepción grown at 20 °C. These results suggest that the effect of high temperatures on photosynthetic metabolism of this farmed population could be mild. Driven by seasonal changes in river discharge, precipitation, and coastal upwelling, high variability in abiotic conditions (e.g., temperature, salinity, and turbidity; Saldías et al. 2016) has been observed in the river mouths were A. chilensis thalli are planted in Concepción. These characteristics could be associated to distinctive heat susceptibility of the photosynthetic metabolism of the A. chilensis thalli growing in Concepción farms. In our experiment, the response of the maximum quantum yield (F_v/F_m) to temperature was quite variable. The lowest value of F_v/F_m (indicating photoinhibition) was observed at intermediate temperature (15 °C) in the farm of Maullín. In plants, it is well known that limitation of electron transport that reduces the ability of plants to use light result in an excess light energy that may cause photoinhibition due to damage to the PSII apparatus (Moll and Steinback 1986). However, short-term response of photosynthesis to temperature cannot easily be used to infer the long-term response of algal growth (Wienke and Dieck 1989). Indeed, despite the possible signal of photoinhibition detected at 15 °C, no limited growth could be observed in these *A. chilensis* thalli.

Populations of *A. chilensis* from different sites in Chile have been reported to present ecological differences, potentially linked to local adaptation in response to specific abiotic and biotic environmental characteristics (Santelices and Ugarte 1990; Usandizaga et al. 2018). The success of *Agarophyton* farming depends in part on the origin of the initial inoculum since differences in thallus morphology, agar yield and gel strength and susceptibility to epiphytes exist among regions and populations. Indeed, it have been suggested that random transplantation between distinct habitats could lead to cultivation failure (Santelices and Ugarte 1990). However, in our experiment, even if we included sampling sites located more than 700 km apart, no major effect of the site of origin were detected on growth or photosynthesis. A possible explanation for this discrepancy is that continuous transplantations and exchanges during the last decades have leaded to the homogenization of the genetic diversity among the whole Chilean coast. However, this hypothesis is not in accordance with population genetic studies showing the presence of clear genetic divergence between regions in Chile (Guillemin et al. 2008; 2014). Studies focused on the effect of other stressors (e.g., nutrient supply, salinity, irradiance) and the cumulative effects of various of these stressors on the physiological responses of distinct genotypes is now needed in order to better explore the resistance of *A. chilensis* populations to stress.

Conclusion

Agarophyton chilensis tolerance of a wide range of abiotic conditions has been proposed as one of the main reasons of the species successful expansion in the Pacific and establishment in a wide array of habitats (Santelices and Ugarte 1990; Chow et al. 2001). These successful extension waves have probably also been facilitated by intrinsic characteristics of the species, such as its capacity to shift between sexual and asexual reproduction (Guillemin et al. 2014). We suggest that the possible selection for general-purpose-genotypes in the asexually reproducing farmed populations may help modulating the impact of environmental variation on population dynamics (Reed et al. 2010) and Chilean Agarophyton crop to cope better with impacts of climate change and direct anthropogenic activities. Nevertheless, implementation of breeding strategy and cultivar selection for mariculture systems improvement has not yet begun in Chile and development of long-term management plans for the sustainable exploitation of A. chilensis populations is dearly needed.

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Compliance with ethical standards

Conflict of interest The authors declare no conflict of interest.

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References

- Arnaud-Haond S, Duarte C, Diaz-Almela E, Marbà N, Sintes T, Serrão EA (2012) Implications of extreme life span in clonal organisms: millenary clones in meadows of the threatened seagrass *Posidonia oceanica*. PLoS One 7:e30454:7
- Ask EI, Azanza RV (2002) Advances in cultivation technology of commercial eucheumatoid species: a review with suggestions for future research. Aquaculture 206:257–277
- Baker HG (1974) The evolution of weeds. Annu Rev Ecol Syst 5:1-24
- Bird CJ, McLachan J, Oliveira EC (1986) Gracilaria chilensis sp. nov; (Rhodophyta, Gigartinales), from Pacific South America. Can J Bot 64:2928–2934
- Buschmann AH, Westermeier R, Retamales C (1995) Cultivation of Gracilaria in the sea-bottom in southern Chile: a review. J Appl Phycol 7:291–301
- Buschmann AH, Camus C, Infante J, Neori A, Israel A, Hernández-González MC, Pereda SV, Gomez-Pinchetti JL, Golberg A, Tadmor-Shalex N, Critchley AT (2017) Seaweed production: overview of the global state of exploitation, farming and emerging research activity. Eur J Phycol 52:391–406
- Cayuela L (2011) Modelos lineales: Regresión, ANOVA y ANCOVA. Universidad Rey Juan Carlos, Madrid
- Chow FY, Macchiavello J, Cruz SS, Fonck E, Olivares J (2001) Utilization of *Gracilaria chilensis* (Rhodophyta: Gracilariaceae) as a biofilter in the depuration of effluents from tank cultures of fish, oysters, and sea urchins. J World Aquacult Soc 32:215–220
- Davison IR (1991) Environmental effects on algal photosynthesis: temperature. J Phycol 27:2–8
- Duarte CM, Marbá N, Holmer M (2007) Rapid domestication of marine species. Science 316:382–383
- Eilers PHC, Peeters JCH (1988) A model for the relationship between light intensity and the rate of photosynthesis in phytoplankton. Ecol Model 42:199–215
- Engledow HR, Bolton JJ (1992) Environmental tolerances in culture and agar content of *Gracilaria verrucosa* (Hudson) Papenfuss (Rhodophyta, Gigartinales) from Saldanha Bay. S Afr J Bot 58:263–267
- Ensminger I, Xyländer M, Hagen C, Braune W (2001) Strategies providing success in a variable habitat: III. Dynamic control of photosynthesis in *Cladophora glomerata*. Plant Cell Environ 24:769–779
- Forsman A (2014) Rethinking phenotypic plasticity and its consequences for individuals, populations and species. Heredity 115:1–9
- Gallegos-Sánchez CF, Beltrán J, Flores V, González AV, Santelices B (2018) Testing the effects of heterozygosity on growth rate plasticity in the seaweed *Gracilaria chilensis* (Rhodophyta). Ecol Evol 8: 5741–5751
- Grzymski J, Johnsen G, Sakshug E (1997) The significance of intracellular self-shading on the bio-optical properties of brown, red and green macroalgae. J Phycol 33:408–414
- Guillemin M-L, Faugeron S, Destombe C, Viard F, Correa JA, Valero M (2008) Genetic variation in wild and cultivated population the haploid-diploid red alga *Gracilaria chilensis*: how farming practices favour asexual reproduction and heterozygosity. Evolution 62: 1500–1519
- Guillemin M-L, Huanel OR, Martínez EA (2012) Characterization of genetic markers linked to sex determination in the haploid-diploid red alga *Gracilaria chilensis*. J Phycol 48:365–372
- Guillemin M-L, Sepúlveda RD, Correa JA, Destombe C (2013) Differential ecological responses to environmental stress in the life history phases of the isomorphic red alga *Gracilaria chilensis* (Rhodophyta). J Appl Phycol 25:215–224
- Guillemin M-L, Valenzuela P, Gaitán-Espitia JD, Destombe C (2014) Evidence of reproductive cost in the triphasic life history of the red alga *Gracilaria chilensis* (Gracilariales, Rhodophyta). J Appl Phycol 26:569–575

- McLachlan J (1973) Growth media-marine. In: Stein JR (ed) Handbook of phycological methods. Cambridge University Press, Cambridge, pp 25–52
- McLachlan J, Bird CJ (1984) Geographical and experimental assessment of the distribution of species of *Gracilaria* in relation to temperature. Helgoländer Meeresun 38:319–334
- Meyer RS, Duval AE, Jensen HR (2012) Patterns and processes in crop domestication: an historical review and quantitative analysis of 203 global food crops. New Phytol 196:29–48
- Moll BA, Steinback KE (1986) Chilling sensitivity in Oryza sativa: the role of protein phosphorylation in protection against photoinhibition. Plant Physiol 80:420–423
- De Nicola DM (1996) Periphyton responses to temperature at different ecological levels. In: Stevenson RJ, Bothwell ML, Lowe RL (eds) Algal ecology: freshwater benthic ecosystems. Academic Press, San Diego, pp 149–181
- Reed TE, Waples RS, Schindler DE, Hard JJ, Kinnison MT (2010) Phenotypic plasticity and population viability: the importance of environmental predictability. Proc R Soc B 277:3391–3400
- Richards CL, Bossdorf O, Muth NZ, Gurevitch J, Pigliucci M (2006) Jack of all trades, master of some? On the role of phenotypic plasticity in plant invasions. Ecol Lett 9:981–993
- Robinson N, Winberg P, Kirkendale L (2013) Genetic improvement of macroalgae: status to date and needs for the future. J Appl Phycol 25:703–716
- Saldías GS, Largier JL, Mendes R, Pérez-Santos I, Vargas CA, Sobarzo M (2016) Satellite-measured interannual variability of turbid river plumes off central-southern Chile: spatial patterns and the influence of climate variability. Prog Oceanogr 146:212–222

- Santelices B, Ugarte R (1990) Ecological differences among Chilean population of commercial *Gracilaria*. J Appl Phycol 2:17–26
- Schreiber U, Endo T, Mi H, Asada K (1995) Quenching analysis of chlorophyll fluorescence by the saturation pulse method: particular aspects relating to the study of eukaryotic algae and cyanobacteria. Plant Cell Physiol 36:873–882
- Simms EL (2000) Defining tolerance as a norm of reaction. Evol Ecol 14: 563–570
- Usandizaga S, Camus C, Kappes JL, Guillemin M-L, Buschmann AH (2018) Nutrients, but not genetic diversity, affect *Gracilaria chilensis* (Rhodophyta) farming productivity and physiological responses. J Phycol 54:860–869
- Valero M, Guillemin M-L, Destombe C, Jacquemin B, Gachon C, Badis Y, Buschmann AH, Camus C, Faugeron S (2017) Perspectives on domestication research for sustainable seaweed aquaculture. Perspect Phycol 4:33–46
- Westermeier R, Gómez I, Rivera P (1993) Suspended farming of *Gracilaria chilensis* (Rhodophyta, Gigartinales) at Cariquilda river, Maullín, Chile. Aquaculture 113:215–229
- Wienke C, Dieck I (1989) Temperature requirements for growth and temperature tolerance of macroalgae endemic to the Antarctic region. Mar Ecol Prog Ser 54:189–197
- Zeder MA (2015) Core questions in domestication research. Proc Natl Acad Sci 112:3191–3198
- Zohary D (1984) Modes of evolution in plants under domestication. In: Grant WF (ed) Plant biosystematics. Academic Press Canada, Montreal, pp 579–586
- Zohary D (2004) Unconscious selection and the evolution of domesticated plants. Econ Bot 58:5–10