

1 **A biologically relevant rapid quantification of physical and biological stress profiles**
2 **on rocky shores**

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17 **Abstract**

18 Different combinations and intensities of physical (e.g. thermal) and biological (e.g.
19 competition or predation) stress operate on organisms in different locations. Variation in
20 these stresses can occur over small to medium spatial scales (cm to 10s m) in
21 heterogeneous environments such as rocky shores, due to differences in sun and wave
22 exposure, shore topography and/or recruitment. In this study we demonstrate how simple
23 measurements can be taken that represent physical and biological stresses (stress profiles)
24 in a given location. Using a bootstrapped principal component analysis, we identified
25 significantly different stress profiles at four sites separated by only 10s to 100s of metres on
26 the Shek O peninsula in Hong Kong. We then measured response to thermal stress, as
27 determined by detachment temperature, in the limpet *Cellana grata* (which is known to be a
28 sensitive indicator species to thermal stress) from each location. Significant differences in
29 stress profile between locations were also seen in thermal stress tolerance of limpets from
30 those locations. At locations where the major stresses are likely to be more physical or less
31 biological in nature (e.g. southerly facing aspect or lower density of grazers), the mean
32 detachment temperature was higher, whereas detachment temperature was lower at sites
33 with more biological or less physical stress. This method is, therefore, able to determine
34 biologically meaningful differences in stress profiles over small to medium spatial scales, and
35 demonstrates that localised adaptation (i.e. post planktonic settlement) or acclimation of
36 species may occur in response to these different stress profiles. The technique can be
37 adapted to different environments and smaller or larger spatial scales as long as the stress
38 experienced by the study species is relevant to these scales.

39 **Key words:**

40 Rocky shore; Limpet; *Cellana*; Hong Kong; Environmental stress; Climate change; PCA

41 **1. Introduction**

42 On rocky shores few organisms live at environmentally optimal conditions, and are often
43 forced towards the edges of their fundamental niche by biological factors such as predation
44 and competition (Lubchenco, 1980; Somero, 2010; Underwood and Denley, 1984).
45 Examples of these realised niches are common in the rocky shore literature, where shore
46 height (and hence relative immersion:emersion time) provides a simple and localised
47 environmental gradient (Colman, 1933; Harley and Helmuth, 2003). For example,
48 transplants of algae to lower shore levels in the absence of grazers and other competitive
49 species typically result in much higher growth rates of algae than at the shore height they
50 normally occur (Hawkins and Hartnoll, 1985; Norton, 1985), indicating that these species
51 usually occur outside their optimal environmental conditions.

52 Typically, for hard substrate marine communities there is considerable competition for space
53 (reviewed by Paine, 1984), and as a result, many organisms will be forced towards the
54 edges of their fundamental niches (Braunisch et al., 2008; Costantini et al., 2009). To
55 contend with this, organisms on rocky shores must either tolerate less favourable
56 environmental conditions such as longer periods of desiccation or thermal stress, or have
57 the ability to be able to cope with higher levels of competition, grazing or predation (van
58 Straalen and Roelofs, 2012).

59 In the rocky intertidal zone, different physical locations (shores or sites within shores)
60 present very different sets of biological and physical challenges for a given species, and
61 these differences can change over a range of spatial scales (see examples in Harley and
62 Helmuth, 2003). For example, at headlands and peninsulas both wave exposure and direct
63 sunlight (and hence temperature and desiccation levels) can vary dramatically over 10s of
64 metres (for example, between north and south facing shores). The impact of predators,
65 competitors and facilitators can also vary over these spatial scales (Menge et al., 1994),
66 often as a result of differential physical factors (Benedetti-Cecchi, 2001). Such distances are
67 normally greater than a typical organism will displace in its adult lifetime, especially when the
68 rocky substratum is not continuous (i.e. separated by sand patches).

69 The classic theoretical models of Menge and Sutherland (1976; 1987) propose that the roles
70 of physical and biological factors such as disturbance, competition and predation, differ with
71 environmental stress. Since physical stress differs at a local scale of 10s of metres (due to
72 aspect or wave exposure), this infers that biological stresses should also vary at these
73 scales (Benedetti-Cecchi, 2001; Harley and Helmuth, 2003), as do patterns of community
74 structure (e.g. Hutchinson and Williams, 2001).

75 Adaptations to these combinations of stress (herein 'stress profiles') are likely to occur to
76 maximise survival and reproduction (and hence fitness). Such adaptations could occur
77 through natural selection, however, given that most rocky shore organisms have a planktonic
78 dispersal phase (Pechenik, 1999), localised adaptations (over distances measured in 10s or
79 100s m) are likely to be acclimation responses rather than evolutionary responses (Allcock
80 et al., 1997, Hoskin 1997; Somero 2010; but see Kruger-Hadfield et al., 2013).

81 In this study, we characterise small sections of shore separated by short distances (10s –
82 100s m) in relation to a number of physical and biological parameters, to determine the
83 combinations of physical and biological factors that influence the stress profile of organisms.

84 Using a bootstrapped PCA approach (modified from Catlin-Groves et al., 2009 and Stafford
85 et al., 2012), we test for significant differences in types of stress acting on limpets (*Cellana*
86 *grata*) on four sections of shores (each 10 m long, but all separated by < 500 m) on the Shek
87 O peninsula on Hong Kong Island. We collected limpets from these sections and tested their
88 response to thermal stress by investigating detachment temperatures. We hypothesise that,
89 where stress profiles differ between sections of shore, those sections where physical
90 parameters are more important (i.e. a lower density of grazers, or a southerly aspect), will
91 have limpets with higher detachment temperatures (i.e. animals are more resistant to
92 temperature extremes). Equally, where stress profiles are more associated with biological
93 stress and less with physical stresses (e.g. higher grazer density meaning more competition,
94 northerly aspect), limpets will have lower detachment temperatures. Support for these
95 hypotheses would indicate that the calculation of stress profiles is biologically meaningful
96 and a useful tool to investigate differences in stress profiles over small spatial scales.

97 **2. Methods**

98 Four sites (continuous sections of rocky shore of 10 m in length) were selected on the Shek
99 O peninsula on Hong Kong Island (Figure 1) in late May 2013. Sites were selected based on
100 initial inspection of community structure, and knowledge of the direction they faced as well
101 as predominant wave conditions to ensure a variety of potential physical and biological
102 stresses were captured. A 10-m line transect was placed at the height on the section of
103 shore with the highest density of *Cellana grata* (this shore height differed between sites, but
104 was recorded by measuring the vertical distance from the sea using a cross staff, see Baker
105 and Crothers, 1987 for details). The GPS position (accuracy $\pm 3\text{m}$) was noted at each site,
106 and a number of parameters measured. Grazer density (for *Cellana grata* and combined for
107 all other grazers) was obtained from five haphazardly placed 50 x 50 cm double strung
108 quadrats along each 10 m transect. Crevice and rockpool percentage cover and barnacle
109 and mussel percentage cover were also recorded from these five quadrats using the point
110 intercept technique (as described in Bohnsack, 1979). Five recordings of shore slope were
111 taken, using a clinometer placed in the centre of each quadrat (one recording per quadrat).
112 This sampling covered 50% of the area being considered, which has been shown to be
113 sufficient to capture details of even the most patchily distributed organisms or shore
114 characteristics (Stafford, 2002). Photographs of the site, in direct sunlight with a white sheet
115 of paper in the field of view were also taken to determine rock colour (see below).

116 Simple computer-based analysis followed the field-based data collection. Sites were scored
117 on an ordinal scale for exposure to sun (north facing = 1, east = 2, west = 3, south = 4) and
118 wave exposure (west = 1, north = 2, south = 3, east = 4). Wave exposure was based on
119 predominant swell direction for Shek O, obtained from seasonal data available from
120 MetCentral Ltd and from analysis of fetch from the sites (i.e. shelter from headlands or near
121 shore islands reduced the exposure of north facing beaches). Photographs of the site were
122 analysed to determine the rock colour using the GIMP image analysis package
123 (www.gimp.org), by converting to greyscale, transforming the image so the white paper had
124 a pixel colour value of 255, then calculating the mean pixel colour value of 10 rock sample
125 points (average of 4 x 4 pixels) to determine the darkness of the rock (converted so that
126 darker values have higher values by subtracting the mean value obtained from 255).

127 Raw data for each site were transformed into a percentage weighting for each measured
128 parameter. This presented data from different variables on a uniform scale, but also was

129 fundamental to the assumption that all sites are stressed, just by different factors or
130 magnitudes of those factors. For example, if a site faced south, yet had few other grazers,
131 the importance of the direction in terms of thermal stress would be emphasised and the
132 contribution of biological stress would be reduced. To achieve this percentage weighting; 1)
133 all measured variables at a given site were standardised by dividing by the mean value for
134 that variable across all sites. 2) The percentage value of each factor at each site was then
135 calculated by dividing the standardised values, obtained in step 1, of each factor at each site
136 by the sum of all factors for each site and then multiplying by 100 (see supplementary
137 material for details of calculation). This way, the standardised percentage weighting of each
138 factor for each site was determined (Table 1).

139 Data were analysed using a bootstrapped principal component analysis (see Stafford et al.,
140 2012 for full details). Briefly, sites were classified by 100 data points, where x of those points
141 were equal to the percentage of the considered component of the physical and biological
142 factors at each site (i.e. the first factor is density of *Cellana grata*, the second the density of
143 other grazers and so on, following the order of parameters given in Table 1).

144 Samples were then taken (with replacement) of these 100 data points for each site (100
145 points were sampled, but because this was with replacement, many would be sampled more
146 than once, and many not at all, following recommended bootstrapping procedures in Efron,
147 1979). From this sample, the first three principal components were calculated. This process
148 of determining the first three principal components was conducted 10,000 times (with
149 appropriate transformations to allow for the arbitrary nature of principal component axes, and
150 'reflection' of the axes – full details of this are given in Stafford et al., 2012). The samples
151 that were the furthest 5% from the mean PC value for the site were then excluded to provide
152 95% confidence spheres on a three dimensional plot of the first three principal components.
153 An overlap of these spheres indicates no significant difference between the stress regimes
154 of the corresponding sites (see Stafford et al., 2013, for discussion of overlapping of
155 bootstrapped-derived 95% confidence limits for determining significance). The technique
156 was developed to determine differences between limited cases, and hence the variable to
157 case ratio can easily exceed 1 (Stafford et al., 2012). The limitation of the technique is that
158 the variables need to be appropriately compressed so that the first three principal
159 components explain > 90 % of the variability in the data, otherwise the positions of the
160 spheres may not be reliable enough to determine significant differences (Stafford et al.,
161 2012). The code for running the bootstrapped PCA is provided at
162 <http://rickstafford.com/software/software.html>. To explore the critical differentiating factors
163 between the sites, a biplot from non-bootstrapped PCA was produced. However, this plot
164 should be treated with some caution, as the case to variable ratio for PCA is exceeded
165 (Tabachnick and Fidell 1989).

166 Two days after taking shore measurements, *Cellana grata* were collected from each site (all
167 *C. grata* from all sites collected within 90 min on the same day). Where possible 10
168 individuals in the size range 20 - 28 mm were collected. Where this was not possible, due to
169 restricted numbers of limpets in this size range, as many as could be obtained were
170 collected (minimum $n = 7$). Limpets were stored at ambient temperature for no more than 90
171 min before laboratory experiments. Initially all limpets were placed in plastic pots (diameter
172 40 mm) in air and then refreshed under a filtered and aerated seawater spray for 5 min on
173 return to the laboratory. All limpets attached firmly to the bottom of the plastic pots. The pots
174 were then drained of water, sealed, and placed upside down in a water bath, initially at 30

175 °C, and maintained at 30 °C for 2 minutes, before the temperature was raised by 1 °C every
176 2 minutes. A measure of thermal tolerance was obtained by observing when limpets lost
177 adhesion and fell from the top of the pots (the detachment temperature, as described by
178 Wolcott, 1973, after which he considered limpets 'ecologically dead'). Detachment
179 temperatures were analysed by multiple regression, with size of limpet as a linear
180 continuous variable, and site from which the limpet was collected a factorial variable in the
181 analysis. Ideally we would have collected limpets in a narrower size range, then excluded
182 'size' as a variable in the model, and just tested for differences in mean detachment
183 temperature between sites. However, due to the restricted number of limpets at each site,
184 and differing, but overlapping sizes between sites, we included size as a variable in case this
185 directly influenced detachment temperature.

186 **3. Results**

187 The bootstrapped PCA process indicated that two sites were statistically similar in terms of
188 their physical and biological characteristics (i.e. their spheres overlapped), and two sites
189 were significantly different from each other and from the two similar sites (i.e. spheres did
190 not overlap; Figure 2). The first three principal components explained 97.7% of the variability
191 in the data. An examination of the biplot showed that sites 1 and 4 were discriminated from
192 the other sites by increased physical stress, especially insolation, and also showing lighter
193 rock colouration (Figure 3). The importance of biological stress was reduced in these sites
194 whereas site 2 had increased number of limpets and site 3 had increased numbers of other
195 grazers (Figure 3).

196 During the detachment temperature experiment, ~ 10 % of limpets crawled down the side of
197 the pots, and as such, detachment temperature could not be obtained for these individuals.
198 Two limpets did not detach from the pots due to adhesion with mucus, but were found to be
199 dead after termination of the experiment at 60 °C, and these limpets were excluded from the
200 analysis. All remaining limpets detached before 40 °C. Overall, 8 limpets provided data from
201 site 1, 10 from site 2, 5 from site 3 and 6 from site 4. The regression for detachment
202 temperature including the factor 'site' and the continuous variable of limpet size was
203 significant with an adjusted $r^2 = 0.205$ (Table 2a). There was no effect of limpet size on
204 detachment temperature, but there was a difference between the sites the limpets were
205 collected from and their detachment temperatures (Table 2; Figure 4). Sites 1 and 4 did not
206 differ significantly from each other, but sites 2 and 3 were significantly different from site 1
207 (Table 2a; Figure 4). In a separate analysis, sites 1 and 4 were combined (giving 14 limpet
208 detachment temperatures), because these sites were not significantly different according to
209 the bootstrapped principal component analysis. Again, the regression was significant, but
210 explained slightly less variability ($r^2 = 0.181$), in this case, detachment temperatures of
211 limpets from sites 2 and 3 were significantly different from the combined site, but not from
212 each other (Table 2b; Figure 4).

213 **4. Discussion**

214 The results support the hypotheses presented in the introduction; namely that shores with
215 relatively higher levels of physical stress such having southerly aspects would have higher
216 limpet detachment temperatures than sites with relatively higher levels of biological stress.
217 Sites 1 and 4 had lower proportions of intra- and inter-specific competition, as measured by
218 grazer density, than sites 2 and 3, indicating lower relative levels of biological stresses such

219 as competition. These sites were also likely to have higher levels of physical stress, being
220 darker in colour and higher on the shore than sites 2 and 3. While the proportion of insolation
221 was lower at site 1 than at sites 2 and 4, both sites 1 and 4 were south facing, whereas site
222 2 was north facing and site 3 west facing.

223 Using a principal component analysis technique to determine differences between shores
224 does demonstrate the inter-relatedness of some variables. For example, sites 1 and 4
225 showed higher levels of insolation, yet by examining the biplot, this was closely related to
226 shore height, exposure and colour of the rock. In some cases, these relationships between
227 factors might be slightly counterintuitive. For example, wave exposure might mean reduced
228 desiccation or temperature stress, if the rock surface were wet by waves during emersion.
229 However, *Cellana grata* suffers high levels of mortality due to acute physical stress, often
230 occurring over one or a few consecutive tidal cycles in mid-summer (Ngan 2006; Williams
231 and Morritt, 1995). At these times, insolation and temperature would be very high, and
232 normally related to high pressures and calm seas.

233 Colour was measured as the whiteness of the rock, and as such, it may be related to lack of
234 biofilm. Because of this, the variable space largely divides into physical and biological factors
235 with the top left corner of the biplot being dominated by physical factors. Differences
236 between sites 2 and 3 largely occur because of differences in water retaining features and
237 complexity (rockpools and crevices vs mussel beds and barnacles) and the type of
238 competition occurring (intra- vs. inter-specific). It is unclear if these differences could be
239 important in determining stress acting on the shore, and further details of adaptation to
240 biological stress may need to be measured to determine this – in this study shores with
241 higher physical stress do show differences in responses of their limpets to detachment
242 temperature.

243 This study demonstrates that the stress profile of a limited section of rocky shore can be
244 described by a small number of simple field parameters augmented by follow-up image
245 analysis of photographs. Using open source statistical techniques, it is possible to determine
246 significantly different stress profiles on spatially proximate sections of shores. These
247 calculated stress profiles demonstrate real biological relevance, because the heat tolerance
248 of limpets from these different stress profiles also differed significantly and in a predictable
249 manner based on the measured stress profiles.

250 Predicting different stress profiles at small spatial scales is important. Traditional broad scale
251 (10s – 100s km) climate envelope and ‘niche’ models, which largely concentrate on air
252 temperatures and rainfall (e.g. Peterson, 2001) are proving ineffective at mapping fine scale
253 species distributions, leading to the development of smaller microclimate models (Gillingham
254 et al., 2012a, b). However, biological factors such as competition, predation, parasitism and
255 facilitation are known to be important in determining the ecological niche of most organisms
256 (Liebold, 1995). Despite the recognized importance of biological factors, determining the
257 strength of such interactions is difficult and time consuming without intensive manipulative
258 experiments or modern modelling approaches (reviewed by Underwood, 2000; Wootton and
259 Emmerson, 2005).

260 The technique presented here provides a rapid, largely *in-situ* method to determine stress
261 regimes, indicative of the biological and physical factors acting on organisms. Localised
262 acclimation by limpets also appears to result from these different stress regimes (possibly as

263 a result in changes in heat shock proteins Haplin et al., 2002; Harley et al., 2009).
264 Understanding that in limpets different types of acclimation may occur on different sections
265 of shores where the factors structuring communities and acting on individuals differ
266 significantly (for example, a greater proportion of competition vs. a greater proportion of
267 physical stress through direct insolation at different sections of the shore), and may not only
268 be related to large scale variables such as latitude, is important in understanding species
269 responses to disturbances such as climate change (see discussion in Helmuth, 2009).

270 Many studies are recognising the need for incorporation of biological and physiological
271 processes along with environmental variables. The notion of the 'physiological niche' has
272 become popular, with many authors realising it may be an interplay of physical and biological
273 processes that determine species distributions, growth rates and ultimately evolutionary
274 fitness (e.g. Helaouët and Beugrand, 2009; Helmuth, 2009; Kearney and Porter, 2009; Sarà
275 et al., 2011). However, the approach detailed in this paper of classifying micro- or meso-
276 habitats as significantly different in the makeup of physical and biological factors can be
277 used to rapidly determine differences in realised niches of organisms in the field. Significant
278 differences detected between sites can also validate the use of 'natural experiments', where
279 different sites are compared, rather than true manipulations being carried out (Diamond,
280 1983; Sagarin and Pauchard 2010). Using this field technique along with measures to
281 predict and measure physiological limits of study organisms may allow for validation of some
282 of the more predictive approaches that apply the physiological niche concept. Combining
283 these approaches may be vital for understanding biological responses to environmental
284 change, which may embody changes to many physical and biological variables (Sagarin and
285 Pauchard 2010), and can play an important role in informing predictive models of
286 environmental change.

287 The limited movement of many adult forms of rocky shore species makes small scale
288 acclimation (i.e. over distances of 10s to 100s m) more important than in many other marine
289 and terrestrial environments. Indeed, there is considerable variability in stress on rocky
290 shores at the spatial scale of centimetres, with crevices or shade providing refuges from heat
291 and desiccation stress (Chappon et al., 2013; Denny et al., 2011; Garrity 1984; Stafford
292 2002). Furthermore, at distances of metres (rather than 10s m), there can be considerable
293 changes in vertical distribution patterns and the stresses associated with these patterns. For
294 most grazing molluscs, these small scale patterns may be important over short temporal
295 scales (such as tidal cycles), but less important for long-term adaptation than the 10s of
296 metres scale examined here. *Cellana grata*, for example, is not a homing limpet, and unless
297 crevices are plentiful, is frequently found on flat rock (Williams and Morritt, 1995; Williams et
298 al., 2005). It can displace by several metres each day, but will be confined to short sections
299 of shore separated by sand or water for its adult life. As such, examining a 10 m stretch of
300 shore for a biological variable related to long-term adaptation is logical. However, to assess
301 a shorter term acute response to stress as may occur during the emersion period of a tidal
302 cycle, examining shore characteristics on the scale of centimetres would be more
303 appropriate. For more mobile rocky shore organisms, such as crabs, larger spatial scales,
304 including much wider vertical ranges would need to be considered. On some shores,
305 predation risk also needs careful consideration, as it greatly affects community structure and
306 likely acts as a strong selective pressure. Molluscan predators such as dogwhelks can be
307 quantified in a similar manner to grazers, using quadrat counts; however, these were not
308 seen in the quadrat counts in this study. Fish as predators and grazers can be important on

309 many rocky shores, especially in tropical regions (although less so in Hong Kong; Williams et
310 al., 2000), and if this is an important consideration, then high tide surveys would also need to
311 be conducted.

312 To examine evolutionary pressures it should be noted that intergenerational dispersal may
313 be higher for many rocky shore (and other marine) species than many terrestrial species
314 such as plants or insects. Therefore, even in terrestrial systems adaptation (through long-
315 term acclimation or evolution) may be localised for many species (e.g. Ayre, 1995 and
316 references within). As such, the ability to calculate stress regimes at local, biologically
317 relevant, scales has considerable implications for successfully predicting organisms'
318 responses to climate change. Rapid in-field measurements, such as those presented here,
319 allow for quantification of differences in stress regime of organisms, and could be used to
320 rapidly assess large numbers of micro- to meso-scale habitats.

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Table 1. Physical and biological measurements of each site. Raw data collected *in situ* or from examination of site aspect or photographs and the percentage contribution of this parameter to the overall stress profile of the site (see methods for details). The final column indicates the rationale for choosing this measurement.

Variable	Raw data Site 1	Raw data Site 2	Raw data Site 3	Raw data Site 4	Percentage contribution Site 1	Percentage contribution Site 2	Percentage contribution Site 3	Percentage contribution Site 4	Rationale
Limpet (mean number per quadrat)	13.6	38.6	23.2	4.6	8	15	10	2	Intraspecific competition
Other grazers (mean number per quadrat)	4.4	3.0	23.6	3.8	6	3	24	4	Interspecific competition
Barnacle/Mussel (mean percentage cover)	1.0	45.6	0.0	9.8	1	28	0	7	Facilitation and provision of biotic microhabitats
Crevice/Rockpool (mean percentage cover)	6.2	5.2	18.4	15.4	9	5	20	17	Provision of abiotic microhabitats
Angle (degrees from horizontal)	28.4	31.8	63.4	24.2	10	7	16	6	Related to potential shade and water run off
Shore Height (m + C.D)	2.0	1.85	1.2	2.2	17	10	7	15	Time of emersion
Colour (mean pixel value)	135	99	111	144	17	8	10	14	Heat absorption
Insolation Index (1-4)	4.0	3	1.5	4.0	20	10	6	16	Exposure to direct sunlight and associated temperature
Exposure Index (1-4)	2.0	3.5	1.6	4.0	12	14	7	19	Exposure to wave action and possible modification of effective shore height

Table 2. Results of multiple regression analysis for detachment temperature of limpets against size of limpet and the site it was collected from (a) for all four sites. Results for different sites demonstrate significant differences compared to site 1. (b) for sites 1 and 4 combined (as these sites were shown not to differ significantly in stress profile). Results for different sites demonstrate significant differences compared to the combined sites 1 and 4.

a)

	Estimate	s.e.	t	p
(Intercept)	30.2021	6.4871	4.656	< 0.0001
Size	0.3560	0.2888	1.232	0.2297
Site 2	-3.8428	1.3549	-2.836	0.0091
Site 3	-4.2801	1.6687	-2.565	0.0170
Site 4	-2.1689	1.6335	-1.328	0.1968

b)

	Estimate	s.e.	t	p
(Intercept)	32.2021	6.4053	5.027	< 0.0001
Size	0.2285	0.2765	0.826	0.4164
Site 2	-2.9862	1.2095	-2.469	0.0207
Site 3	-3.2885	1.5148	-2.171	0.0396

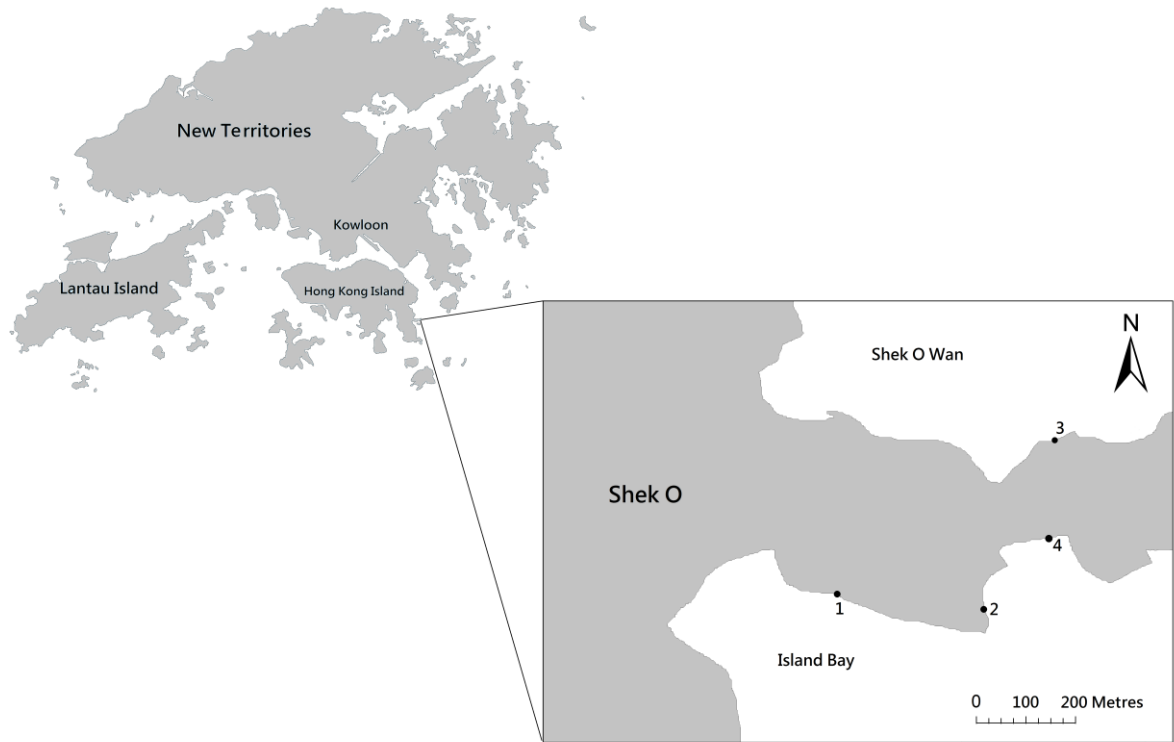


Figure 1. Locations of the four sites on the Shek O peninsula on Hong Kong Island.

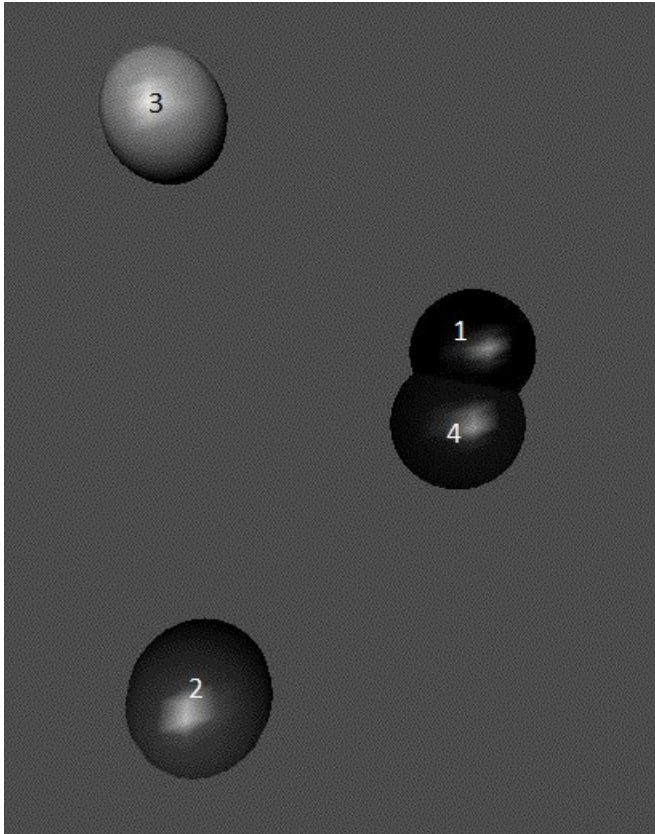


Figure 2. Bootstrapped PCA (with 95% confidence spheres) of the measured physical and biological variables at the four sites (1-4). Overlap indicates that no significant difference occurs between the sites.

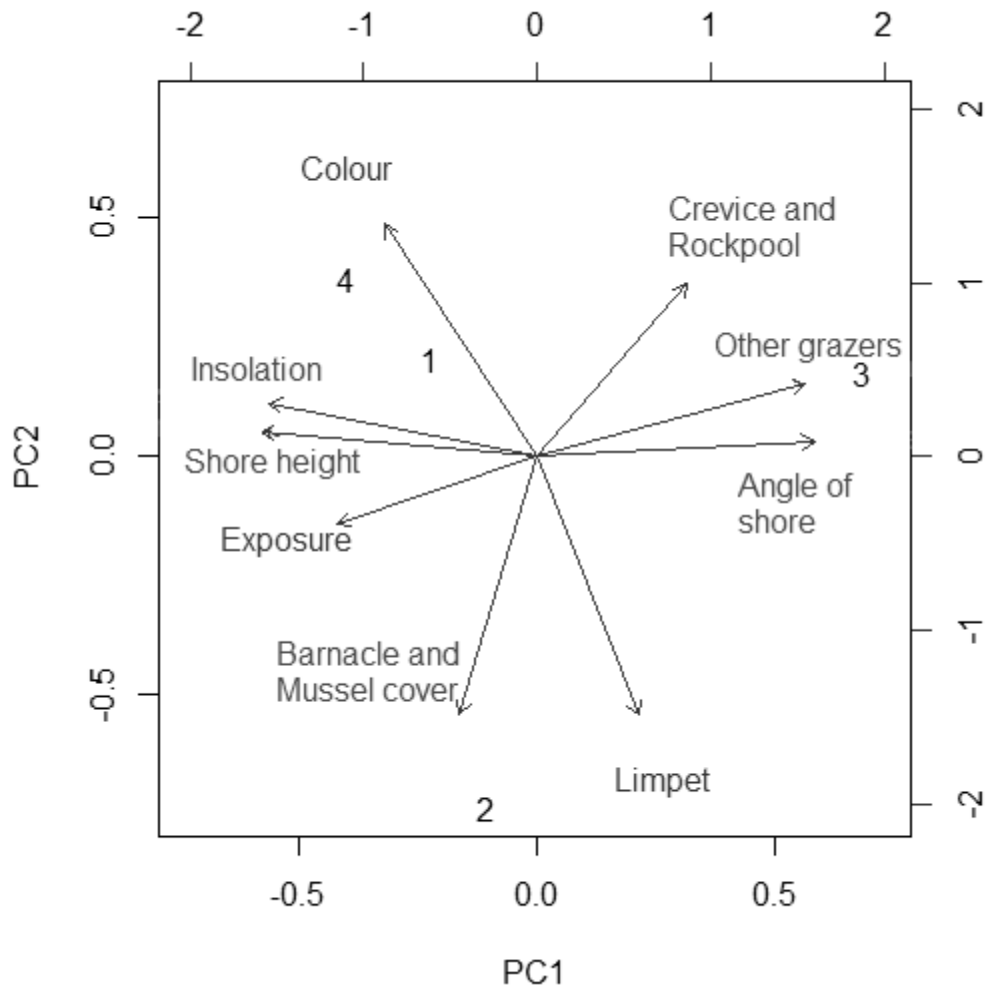


Figure 3. Biplot of non-bootstrapped PCA based on percentage data in table 1. Arrows indicate key discriminatory factors between sites (e.g. site 3 is discriminated from other sites by a greater number of other grazers and a steeper angle of shore). Note, variable to case ratio is exceeded for PCA (see methods).

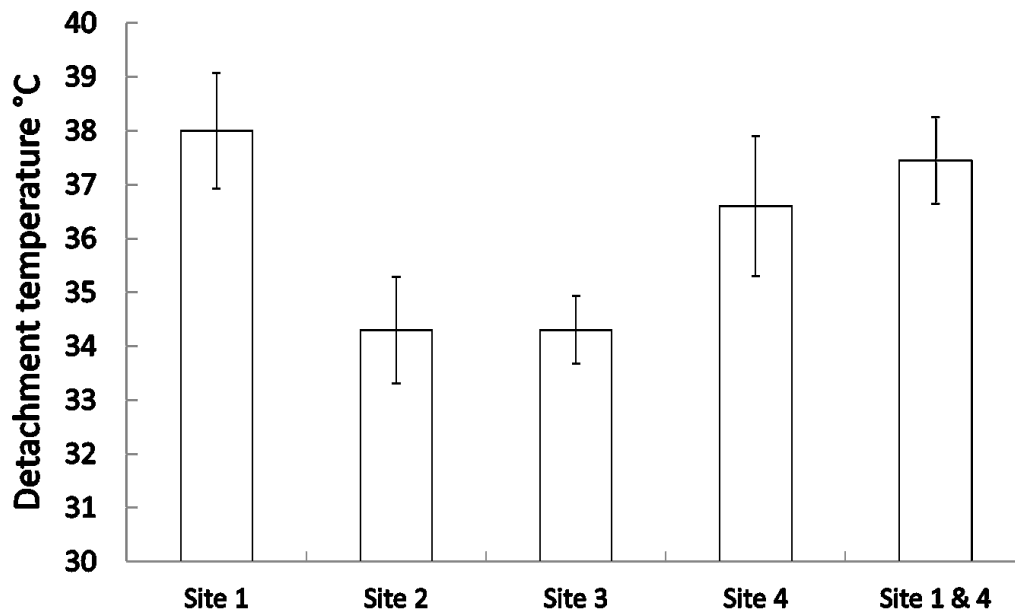


Figure 4. Mean (\pm S.E. n = 8, 10, 5, 6 for sites 1-4, respectively) detachment temperature of limpets from each site. The final bar indicates the combination of sites 1 and 4 (n = 14) as these shores did not demonstrate significantly different stress profiles.

Supplementary material 1. Worked example of calculating percentage contribution of different types of stress. Excel workbook with macros.

	Limpet (mean % cover)	Other grazers (mean % cover)	Barnacle/Mussel (mean % cover)	Crevice/Rockpool (mean % cover)	Angle (deg)	Shore Height (m)	Colour (m)	Insolation (h)	Exposure Index (1-4)	
site1	13.6	4.4	1	6.2	28.4	2	135	4	2	
site2	38.6	3	45.6	5.2	31.8	1.85	99	3	3.5	
site3	23.2	23.6	0	18.4	63.4	1.2	111	1.5	1.6	
site4	4.6	3.8	9.8	15.4	24.2	2.2	144	4	4	
	25.13333333	10.33333333	15.53333333	9.93333333	41.2	1.683333	115	2.833333	2.366667	Mean Stage 1 - Ca
Stage 2 - Then divide each value in the column by the mean of that column										
	0.541114058	0.425806452	0.064377682	0.624161074	0.68932	1.188119	1.173913	1.411765	0.84507	Sum of Row Stage 3 - ca
	1.535809019	0.290322581	2.935622318	0.523489933	0.771845	1.09901	0.86087	1.058824	1.478873	6.963647
	0.923076923	2.283870968	0	1.852348993	1.538835	0.712871	0.965217	0.529412	0.676056	10.55466
	0.183023873	0.367741935	0.630901288	1.55033557	0.587379	1.306931	1.252174	1.411765	1.690141	9.481689
										8.980391
Stage 4 - divide each cell in each row by the sum of the row, and then multiply by 100 (see cell formula)										
	7.770555952	6.114705035	0.924482326	8.963135355	9.898842	17.06173	16.85773	20.27335	12.13546	Step 5 - check it adds to
	14.55099765	2.750656583	27.81350605	4.959796882	7.31283	10.41255	8.156295	10.03181	14.01156	100
	9.735364241	24.0871754	0	19.5360665	16.22955	7.5184	10.1798	5.583518	7.130126	100
	2.038038914	4.094943266	7.025320556	17.26356336	6.54068	14.55316	13.94342	15.72053	18.82035	100
Step 6 - now round to whole numbers - but then check it still adds to 100 Keep the rounded values for the input file										
	8	6	1	9	10	17	17	20	12	100
	15	3	28	5	7	10	8	10	14	100
	10	24	0	20	16	7	10	6	7	100
	2	4	7	17	6	15	14	16	19	100