Effects of disturbance timing on community recovery in an intertidal habitat of a Korean rocky shore

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Intertidal community recovery and resilience were investigated with quantitative and qualitative perspectives as a function of disturbance timing. The study was conducted in a lower intertidal rock bed of the southern coast of South Korea. Six replicates of artificial disturbance of a 50 cm × 50 cm area were made by clearing all visible organisms on the rocky substrate in four seasons. Each of the seasonally cleared plots was monitored until the percent cover data reached the control plot level. There was a significant difference among disturbance timing during the recovery process in terms of speed and community components. After disturbances occurred, Ulva pertusa selectively preoccupied empty spaces quickly (in 2-4 months) and strongly (50-90%) in all plots except for the summer plots where non-Ulva species dominated throughout the recovery period. U. pertusa acted as a very important biological variable that determined the quantitative and qualitative recovery capability of a community. The qualitative recovery of communities was rapid in summer plots where U. pertusa did not recruit and the community recovery rate was the lowest in winter plots where U. pertusa was highly recruited with a long duration of distribution. In this study, U. pertusa was a pioneer species while being a dominant species and acted as a clearly negative element in the process of qualitative recovery after disturbance. However, the negative effect of U. pertusa did not occur in summer plots, indicating that disturbance timing should be considered as a parameter in understanding intertidal community resilience in temperate regions with four distinct seasons.

Key Words: macroalgae; pulse disturbance; recruitment; resilience; Ulva pertusa

INTRODUCTION

Disturbance is an event that reduces a substantial amount of living biomass from a habitat due to various physical and biological causes. It often resets ecological equilibrium, temporarily increases or decreases the diversity of a community, and brings a habitat into a mosaic pattern (Bulleri et al. 2016). The cause of disturbance is different between land (fire, storms, pests, floods, etc.) and ocean (storms, pollution, oil spills, trampling, etc.), and it appears in various patterns depending on areal extent, magnitude, frequency, predictability, and turnover rate or rotation period (Sousa 1984b). Particularly, disturbance in the rocky intertidal zones involves unpredictable and diverse causes compared to other ecosystems. In addition, changes and the intensity of physical marine environments such as waves, water temperature, and salinity add additional variables to the composition and formation of a biological community in an intertidal shore. Especially, disturbance by human influences has changed community structure in rocky intertidal zones over a long period, and the magnitude and frequency has

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The resilience pattern of algal communities in rocky intertidal zones after disturbance is extremely dynamic. Due to the year-round availability of propagule supply and often short life span of organisms, the speed of community recovery after disturbance is faster than any other ecosystem (Sousa 1984a). Speed and pattern of a community’s recovery has been reported to be related to disturbance size, timing, and early colonists (Connell and Slatyer 1977, Sousa 1984b). Also, the interactions between disturbance size and timing, geographical characteristics and the environment were reported as important factors in community resilience (Turner 1983, Farrell 1989, Kim and DeWreede 1996, Benedetti-Cecchi et al. 2003). Benedetti-Cecchi (2000) emphasized the influence of direct and indirect interactions on the recovery patterns of a rocky intertidal community through verifying various research hypotheses. Viejo (2009) compared the recovery patterns of communities in relation to disturbance along the tide level of rocky intertidal zones and concluded that the presence of physical stress and the biological characteristics of dominant species were the determining factors. Recently, Diez et al. (2009) showed that the recovery of a community in an intertidal zone was different in species composition depending on pollution levels, indicating pollution could be an another factor affecting community recovery. Therefore, understanding the role of various biological and physical factors affecting post disturbance processes is essential, but complicated (Viejo et al. 2008).

The ecological importance of disturbance timing has been suggested by multiple researchers (Peterson and Stevenson 1992, Addessi 1994, Hutchinson and Williams 2003). Recently, species diversity was reported to increase faster in the dry season than in the rainy season on a tropical rocky shore (Mayakun et al. 2010). In a temperate climate with four distinct seasons a more precise and detailed interpretation on disturbance timing may be required. The timing of a disturbance can be important since the availability of propagules is often temporally variable (Sousa 2001, Bellgrove et al. 2004), with differences in seasonal recruitment patterns influencing community structure and patterns of succession (Underwood and Anderson 1994). However, some studies showed that effects of disturbance timing were less important depending on dispersal patterns and temporal variation of dominant species in the community (Kim and DeWreede 1996, Foster et al. 2003). More recently, research on disturbance timing tend to extend their scope with relation to climate change and physical interferences (Huff 2011, Deepananda and Macusi 2012, Bernhardt and Leslie 2013). Additional studies on disturbance timing, therefore, are still required in various habitat types and geographic areas to give a habitat-specific pattern of community resilience in coastal rocky shores.

This study is the first report for Korean rocky shores of the influence of the timing of disturbance on macroalgal community recovery. We describe the patterns of community resilience in quantitative (e.g., speed of recovery) and qualitative (e.g., diversity or similarity) perspectives after seasonal experimental disturbances. Different compositions of residence species with season in the Korean habitat may provide valuable implications to disturbance ecology.

MATERIALS AND METHODS

Study site

The study site is located on a rocky shore of Jeongdo-ri, Wando-gun, Korea (34°17′45″ N, 126°42′50″ E), within the Dadohae Marine National Park, designated in 1981 (Fig. 1). Wando is one of the main islands in the Korean South Sea, surrounded by many aquaculture farms producing Sargassum fusiforme, Pyropia tenera, and Haliotis discus. Nevertheless, the rocky shores of Jeongdo-ri represent a typical intertidal habitat for the southern coast of Korea. The shore of Jeongdo-ri consists mainly of a long pebble beach and gently sloping bedrock. The experiment was carried out on flat bedrock, which emerged only at low tides (<60 cm below mean sea level) (National Oceanographic Research Institute 2007). The range of tidal height in the study area was between 8 and 379 cm (as of March 2009). Choi and Kim (2004) reported Gelidium sp., articulate coralline algae, Grateloupia elliptica, Chondracanthus tenellus, S. fusiforme, and Chondrus ocellatus as the dominant species in the lower rocky intertidal zone of the study site. From our field observations, the six most common species were Ulva pertusa, Chondracanthus intermedius, articulate coralline algae, Gelidium sp., Chondrus ocellatus, and Chondracanthus tenellus, indicating that not much change had occurred in the community structure for about 6–7 years except for the appearance of U. pertusa as the most dominant species. Recently, U. pertusa was again reported as the most dominant species (Park et al. 2009). Invertebrate species were not at a significant amount at the tidal level of the experiment conducted, but limpets (Cellana toreuma and Acmaea palli-
Experimental design and data collection

Artificial disturbance was achieved by clearing all organisms on the rock surface using a chisel and metal brush scraper. All visible organisms in 50 cm × 50 cm plots were removed and rechecked carefully after the subsequent high tide, although small pieces of crustose corallines were difficult to remove. We did not use any chemicals or a fire torch to avoid possible damage to adjacent seaweeds and invertebrates. A permanent transect line (25 m in length) marked in every 0.5 m was placed horizontally on the granite bedrock and the line position was fixed using stainless steel anchor bolts at both ends. Therefore, we could place the 50 cm × 50 cm quadrat on the exact positions every sampling time. Six replicate plots (50 cm × 50 cm) were randomly assigned and cleared each season, and an additional six control plots marked at the beginning of experiment and monitored throughout the study period (6 replicate plots × 4 seasons, plus 6 replicate controls = 30 plots in total). Seasonal pulse disturbances were made in July 2007 (summer), April 2008 (spring), November 2008 (autumn), and January 2009 (winter). Field observations and data collections were mostly done at 1-2 month intervals with a total 24 surveys from July 2007 to July 2010 when the communities of the treatment plots and control plots became similar.

Data were obtained by non-destructive methods in the field, and some species that were difficult to identify were collected and fixed with 5% formalin in the field and identified in the lab. Algal percent covers were estimated from subplot-based percentage counts using 5 cm × 5 cm subplots made up with plastic lines in the quadrat as described in Kim et al. (2004a). In the case of relatively larger canopy algae like *Sargassum thunbergii* and *S. fusiforme*, data were counted only when the holdfast was located inside the quadrat. Digital photography, together with data estimates in the field, was sometimes used to supplement field data to obtain unbiased percent cover values.

Data analyses

In this study, community resilience was analyzed both quantitatively (% cover recovery) and qualitatively (species diversity, similarity). One-way ANOVAs were used to compare seasonal effects with total algal cover and selected dominant species. A separate test was done at 2, 6, 11 months after disturbance, because we assumed that these three time frames reflect the early, middle and later stages of community recovery in this study. Kruskal-Wallis tests were often conducted where a log transformation did not meet the parametric assumptions.

PRIMER 5 (for windows ver. 5.2.9) was used for statistical and qualitative analyses. We also used Bray-Curtis similarity analysis (Clarke and Warwick 2001) for the comparison of recovery patterns. Shannon-Wiener’s diversity index was used to qualitatively evaluate community resilience (Wolda 1981).
RESULTS

A total of 16 taxa of macroalgae were recorded during the study period including 10 Rhodophyta, 1 Chlorophyta, and 5 Phaeophyta. The major taxa of >5% average percent cover from the all plots of treatment and control during the 3 years of study period were *Ulva pertusa* (13.3%), *Gelidium* sp. (10.1%), articulate coralline algae (8.3%), *Chondracanthus intermedius* (7.5%), *Chondrus ocellatus* (5.7%), and *Chondracanthus tenellus* (5.3%) (Table 1). The relative frequency of all 16 taxa during the observational period was 0.67 on average and three taxa, *Gelidium* sp., articulate coralline algae, and *Chondrus ocellatus*, were found at all sampling times (1.0), followed by *Grateloupia elliptica* (0.96), *Acrosorium uncinatum* (0.92), *Chondracanthus tenellus* (0.88), and *U. pertusa* (0.83) as the high frequency species.

At each sampling time, an average of 10.7 taxa was observed, the highest being 15 taxa in April 2009 and the lowest 8 in July 2009. Seasonal abundance in percent cover of the major groups changed distinctively. *U. pertusa*, *Gelidium* sp., *Chondrus ocellatus* levels increased from spring and dropped after summer. On the other hand, *Chondracanthus intermedius*, *Chondracanthus tenellus* were relatively more abundant in winter than the other seasons. Articulate coralline algae did not show any seasonal patterns, but the abundance of *U. pertusa* changed the most drastically over time among all the macroalgae (Fig. 2).

Effects of seasonal disturbance on community structure during the recovery process are shown in Fig. 3. In the plots with spring disturbance (April 2008), *U. pertusa* was recruited at a very fast pace and marked 74.83% cover within two months, comparable to 38.17% cover in the control plots (p < 0.05). This pattern persisted over summer and autumn, and then levels were gradually reduced over the winter (24.83% in the treatment plots and 0.5% in the control plots in January 2009). In the following spring (May 2009), the difference between the treatment and control disappeared (55.67% for treatment and 47.3% for control, p > 0.05). The plots with summer disturbance (July 2007) showed a unique pattern from other seasonal plots. *U. pertusa* was not included in the top 5 abundant species, and instead, *Chondria crassicaulis* occupied the plots distinctively throughout the experimental period. Particularly, as *Chondria crassicaulis* started to be recruited in September 2007, they covered 13.67% in the treatment plots, compared with 1.33% in the control plots (Fig. 3). This pattern persisted until January 2008. Together with *Chondria crassicaulis*, *Gelidium* sp., and *Chondracanthus tenellus* were relatively abundant in the summer plots, compared to the other seasonal plots. In the autumn plots (October 2008), *U. pertusa* was selectively recruited to the treatment plots, comprising 49.5% in the treatment plots and 0.5% in the control plots in January 2009. With the seasonal bloom of *U. pertusa* in May 2009, the pattern of the difference between treatment and control was continuously maintained. It is worth noting that although the percent cover of *U. pertusa* dropped substantially in the control plots (18.17%) in December 2009, the percent cover was still high (63.67%) in the treatment plots, disturbed 1 year ago; this pattern persisted until May 2010 (p < 0.05). Recruitment and coverage of *U. pertusa* in the treatment plots of winter disturbance (January 2009) showed an extreme case. Until March 2009, plots were bare for 2 months after disturbance, however recruitment occurred explosively in May 2009 and covered 93.83% of the treatment plots, which was about twice of the coverage in control plots (47.33%). Although the coverage was reduced to 18.17% in the control plots in December 2009, the coverage in the treatment plots was maintained at high levels (>70%) until May 2010 (p < 0.05), 16 months after disturbance. The coverage of *U. pertusa* then reduced rapidly in July 2010 to result in no difference between treatment and control (19.33 and 17.67%, respectively, p > 0.05).

Diversity levels after disturbance were reduced and

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Table 1. Mean percent covers of top 5 species in the four seasonal plots

<table>
<thead>
<tr>
<th>Season</th>
<th>Species</th>
<th>Cover (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spring</td>
<td><em>Ulva pertusa</em></td>
<td>43.4</td>
</tr>
<tr>
<td></td>
<td><em>Chondrus ocellatus</em></td>
<td>7.3</td>
</tr>
<tr>
<td></td>
<td><em>Chondracanthus intermedius</em></td>
<td>5.2</td>
</tr>
<tr>
<td></td>
<td><em>Gelidium</em> sp.</td>
<td>4.6</td>
</tr>
<tr>
<td></td>
<td><em>Chondracanthus tenellus</em></td>
<td>2</td>
</tr>
<tr>
<td>Summer</td>
<td><em>Chondria crassicaulis</em></td>
<td>8.8</td>
</tr>
<tr>
<td></td>
<td><em>Chondracanthus intermedius</em></td>
<td>8.5</td>
</tr>
<tr>
<td></td>
<td>Articulate coralline algae</td>
<td>4.8</td>
</tr>
<tr>
<td></td>
<td><em>Chondracanthus tenellus</em></td>
<td>4</td>
</tr>
<tr>
<td></td>
<td><em>Gelidium</em> sp.</td>
<td>3.3</td>
</tr>
<tr>
<td>Autumn</td>
<td><em>Ulva pertusa</em></td>
<td>53</td>
</tr>
<tr>
<td></td>
<td><em>Chondrus ocellatus</em></td>
<td>9.7</td>
</tr>
<tr>
<td></td>
<td><em>Chondracanthus intermedius</em></td>
<td>4.8</td>
</tr>
<tr>
<td></td>
<td><em>Gelidium</em> sp.</td>
<td>3.9</td>
</tr>
<tr>
<td></td>
<td>Articulate coralline algae</td>
<td>3.9</td>
</tr>
<tr>
<td>Winter</td>
<td><em>Ulva pertusa</em></td>
<td>65.6</td>
</tr>
<tr>
<td></td>
<td><em>Chondrus ocellatus</em></td>
<td>4.9</td>
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<tr>
<td></td>
<td><em>Gelidium</em> sp.</td>
<td>3.4</td>
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<td></td>
<td>Articulate coralline algae</td>
<td>3.2</td>
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<tr>
<td></td>
<td><em>Chondracanthus intermedius</em></td>
<td>1.6</td>
</tr>
</tbody>
</table>
then recovered in all seasonal plots, but the recovery patterns were different depending on disturbance timing (Fig. 4). For the spring plots (April 2008), diversity dropped greatly (1.79 to 0.28) and remained low until January 2009, when rapid recovery occurred to similar levels of the control plots. In January 2009, the abundance of *U. pertusa* (24.83%), which had been recruited immediately after disturbance, started to decrease as other species such as *Chondrus ocellatus* and *Chondracanthus intermedius* increased. After January 2009, diversity levels in the spring plots showed no difference between treatment and control plots. For the summer plots (July 2007) where no dramatic appearance of a particular species like *Ulva* sp., diversity fluctuation was relatively small compared to other seasonal plots. However, diversity levels jumped up (1.46 to 1.92) to the pre-disturbance level in

Fig. 2. Changes in percent cover for six major species during the study period.
Fig. 3. Colonization patterns of major species after each seasonal disturbance.
Fig. 4. Changes in species diversity level during the recovery process.

Fig. 5. Changes in similarity index during the recovery processes.
January 2008 as the percent cover of *Chondria crassicaulis*, which had been recruited at substantial amounts directly after disturbance, decreased. In the autumn plots (October 2008), the decreased diversity (1.51 to 0.68) after disturbance continued for an amount of time until December 2009 and fluctuated with a similar pattern to the control plots. Autumn plots began to recover 16 months after disturbance (i.e., February 2010), which timing was matched to the reduction (63.67 to 43.17%) of *U. pertusa* coverage. Post-disturbance diversity in the autumn plots remained at a similar level as control plots, which period overlapped with low *U. pertusa* coverage. A similar pattern was observed in the winter plots (January 2009), but slightly more time (i.e., 18 months) was required for recovery (1.75 for control and 1.54 for treatment in July 2010).

Community similarity of pre-disturbance conditions of each seasonal plot was approximately 70% (72.11% for spring plots, 71.5% for summer plots, 69.42% for autumn plots, and 66.24% for winter plots). The similarity index between treatment and control plots showed, as expected, a reduction and gradual recovery, but the pattern was different depending on the season of disturbance. The spring plots formed the most heterogeneous community (35.59%) at 4 months after disturbance (August 2008), and after that the similarity was gradually increased reaching 74.5% in May 2009 and maintained at around 70% afterwards. Similarity in the summer plots showed the least fluctuation, reflecting the least diversity changes, which dropped to 57.26% after disturbance (August 2008) with gradual recovery. For the autumn plots, similarity at the pre-disturbance stage was 69.42% which dropped to 33.99% after disturbance. With gradual recovery, similarity reached 67.18% in May 2009 and remained at around 60% afterwards. In the winter plots, similarity of the post-disturbance stage started at 29.95% followed by partial recovery maintained until February 2010. After that, similarity reached 63.24% in May 2010 and 75.7% in July 2010, which was assumed to be complete recovery (Fig. 5).

The seasonal differences in *U. pertusa*, which is an important variable in the process of recovery of a community after disturbance, were compared between periods divided into the early period (2 months after disturbance), middle period (6 months), and later period (11 months) after disturbance. With regard to seasonal differences in the distribution of *U. pertusa* according to the periods of time after disturbance, clear seasonal differences were shown in all of the early, middle, and later periods. In particular, seasonal differences were larger in the early and later periods than in the middle period (p < 0.05).

Two months after the occurrence of disturbance, *U. pertusa* was recruited rapidly and extremely in spring plots (74.83%) and the degree was higher than in autumn plots (49.5%). However, in the case of winter plots, virtually no recruitment of *U. pertusa* occurred in the early period after disturbance (p < 0.05). In the case of the middle period, *U. pertusa* was in the process of gradually decrease in the spring plots (57.83%), maintained high coverage values (72.17%) along with spontaneous generation in the autumn plots, and was extremely recruited to the winter plots (82.67%) (p > 0.05). In the case of the later period, *U. pertusa* coverage values decreased in the spring plots (22.83%) and was still maintained at high levels in the autumn (74.5%) and winter (93%) plots (p < 0.05) (Fig. 6).

**DISCUSSION**

This study aimed to examine the influence of the seasonal timing of pulse disturbances on the patterns of rocky intertidal macroalgal community recovery. After disturbances occurred, *U. pertusa* preoccupied open spaces quickly and exclusively in all plots except for the summer plots where *Chondria crassicaulis* temporarily dominated. The patterns of community recovery thereafter seem to be directly affected by whether *U. pertusa* were recruited or not and the extent of the occupancy. That is, after disturbances, *U. pertusa* acts as a very important biological variable that determines the recovery capability of a community quantitatively and qualitatively. The rapid occupation by *U. pertusa* of even small spaces temporarily made by storms has been previously reported (Turner 1983), with rapid recruitment of *Ulva* sp. suppressing the settlement and recruitment of oth-
er algae (Sousa 1979, Viejo et al. 2008). In this study, we also verified that the coverage of other species such as *Chondrus ocellatus* and *Chondracanthus intermedius* increased from the time when *U. pertusa* coverage began to decrease, indicating the strong space-preoccupying ability of *U. pertusa*. In particular, the preoccupation of bare spaces by the morphologically broad blades of *U. pertusa* acts as a negative influence of the recruitment of secondary species.

*U. pertusa* is a representative marine alga on the coasts of South Korea which is present all year round, but since unpredictable blooms occur frequently, it is not easy to evaluate the role of *U. pertusa* in intertidal community structures (Lee et al. 2000, Oh et al. 2002, Choi and Kim 2004, Kim et al. 2011) and its growth strategies to future climate conditions (Kang and Kim 2016, Kim et al. 2016). Interestingly, in this study, *U. pertusa* did not recruit to the summer plots and this is likely attributable to the natural life cycle of *U. pertusa*. Kim et al. (2004b) showed that *U. pertusa* distribution reached its peak in May and quickly decreased between July and September when temperature rose. Similarly, the coverage of *Ulva* sp. was reported to be the lowest in summer in Hong Kong which has a subtropical climate (Kennish et al. 1996). However, since *Chondria crassicaulis* was temporarily dominant after disturbances although not to a level as high as *U. pertusa*, the argument that the recruitment of certain species (i.e., *C. crassicaulis* in this study) after disturbances affects community recovery is considered valid.

Although we expected that the effects of disturbances would be offset as time passed after disturbance occurrence, the actual patterns of *U. pertusa* distribution showed clear differences among times (seasons). In particular, differences between plots are large in the early period and later period after a disturbance and these differences are attributable to the times of recruitment and periods of distribution of *U. pertusa* after the disturbance. In cases where a disturbance occurred in spring, *U. pertusa* relatively quickly recruited and continuously decreased thereafter. However, in the case of winter plots where a disturbance occurred in December, the initial recruitment of *U. pertusa* occurred later compared to spring plots and the coverage sharply increased at 4 months after the disturbance occurrence and is maintained for more than one year, so that the space preoccupation effect of *U. pertusa* is still maintained in the later period after disturbance.

The seasonal differences in the process of community recovery are noteworthy in terms of quality as well. When a disturbance occurs in the spring, the diversity of a community decreases rapidly and the recovery rate is high. However, when a disturbance occurs in autumn, the decrease in diversity is relatively smaller compared to spring and the recovery rate is very low. This is attributable to the fact that in the case of spring plots, the recruitment of other species such as *Chondrus ocellatus* and *Gelidium* sp. progressed together with that of *U. pertusa* in the process of community recovery, while in the case of autumn plots, *U. pertusa* recruited extremely and the period of persistence was relatively long. An interesting fact is that the pattern of similarity recovery of spring plots and autumn plots is completely different from the pattern of diversity recovery. The similarity recovery of autumn plots is approximately six months faster compared to spring plots and this is likely attributable to the community structure of the control plots. In other words, in spring plots, *U. pertusa* intensively recruited in plots where the disturbance occurred compared to control plots, while in autumn plots, high recruitment of *U. pertusa* occurred simultaneously in both treatment plots and control plots. This means that *U. pertusa* formed clearer independent communities in treatment different from control in the spring plots, and this is likely attributable to whether *U. pertusa* recruitment occurred selectively in plots where disturbances happened or occurred in the entire intertidal zone in the form of a bloom regardless of disturbance. As such, the values of diversity and similarity may be relatively different. In particular, since the results were obtained through comparison with control plots, the significance of the results may be distorted in terms of the resilience of the community. In other words, the community cannot be assumed to have recovered because the community structure is similar to control plots. In fact, similarity increased rapidly between March and May 2009 but diversity decreased. Lande (1996) and Jost et al. (2010) emphasized the necessity to take caution in analysis with ecological indexes, because of their statistical weakness. These points emphasize that interpretation of data regarding the process of community recovery should take into account of both qualitative and quantitative perspectives.

In summary, transitional communities are formed in treatment plots in the early period after a disturbance, but the communities recover to their original state (i.e., more similar to the control plots at the time) over time. As expected, the size of the decrements of diversity and similarity according to disturbance timing and recovery rates are different from plot-to-plot. The qualitative recovery of communities was rapid in summer plots where *U. pertusa* did not recruit and the community recovery rate was
the lowest in winter plots where *U. pertusa* was extremely recruited and the duration of distribution was long. This phenomenon can be explained by the space preemption of early colonists and their sustainability. Sousa (1980) reported that species diversity increased at the beginning of the successional process before a dominant species appeared. However, in this study, that trend was not shown at all. This is likely attributable to the time difference until the dominant species is recruited. That is, in all plots except for summer plots, the recruitment of the pioneer species (*U. pertusa*) occurred very rapidly, before any other species had an opportunity for recruitment. Viejo (2009) stated that the resilience of community was determined according to the characteristics of the dominant species, and their growth and recruitment rates. Thus our results on resilience perspectives provide supportive evidence with the *U. pertusa* space preoccupancy capability appearing in this study.

Although this study took note of the time of disturbance occurrence, many researchers emphasized the distribution and effects of herbivores on the formation of intertidal communities (Cubit 1984, Tuchman and Stevenson 1991, Navarrete 1996, Underwood 1998, Wootton 2002, Mayakun et al. 2010, Tamburello et al. 2014). However, in this study, the effect of herbivores was quite limited. *Littorina* sp., which is a representative herbivore, was partially distributed in the upper part of the intertidal zone, and was barely distributed in the middle and lower parts, where the experimental plots were located. Only filter feeders such as *Serpula vermicularis* and *Serpulobis imbricatus* were present in extremely low densities. Therefore, after the pioneer species settled, there would have been very few biological factors that could possibly reduce the species. Viejo (2009) indicated that the upper part of the intertidal zone showed faster community recovery compared to other tidal levels and evaluated the reason as relatively less external stress such as herbivores. However, in this study, such results were judged applicable to the lower part of the intertidal zone as well. This is set forth as “speed-herbivory-*Ulva’s inhibition*” mechanism in addition to the “speed-herbivory” relationship of Viejo (2009).

In the process of community recovery, the kinds and duration of distribution of initially present species such as *U. pertusa* are very important biological elements (Allison 2004). Benedetti-Cecchi (2000) reported that the space preoccupancy of filamentous algae interrupted the recruitment of later settlement species through a hypothesis testing experiment regarding intertidal bedrock succession. A recent study on the recovery patterns of a coral community in the subtidal zone also revealed that when the percent cover of the turf algae that preoccupied the space decreased, the resettlement of coral clearly increased (Emslie et al. 2008). Other studies also reported that the coralline turf algae that had been recruited after removing the dominant species had a negative influence on the recruitment of *Hormosira* (Bellgrove et al. 2010, Schiel and Lilley 2011). All of these studies show that competition between species occurs in the process of the resilience of a community.

In this study, *U. pertusa* was a pioneer species in the study area while being a dominant species and acted as a clearly negative element in the process of qualitative recovery after disturbance. This seems to support the typical inhibition model (Sousa 1979). However, this model cannot be applied to summer plots, and therefore, to concretely identify the effects of *U. pertusa*, comparison and tests through selective removal of target species should be conducted. Dayton (1975) experimentally verified the competition effects between species by selectively removing the canopies of macroalgae distributed in the same area. In addition, in changes in communities involve diverse temporal and spatial variables, studies and analyses from more composite viewpoints considering differences between sea levels, seasonal and yearly variations are also necessary.

Excessive dominance by a certain species could have an equivalent effect as the occurrence of disturbances. This is also the reason why the role of *U. pertusa* is emphasized in this study. Here, it is noteworthy that most of the times when the diversity of communities largely recovered include winter. This indicates that diversity is recovered when *U. pertusa* recruitment is limited. Therefore, in the case of mid-latitudinal areas where four seasons are distinctive as with South Korea, in addition to the timing of disturbance, the season that follows a disturbance is also considered to be an important environmental factor. Since physical environmental changes depending on seasons lead to the occurrence, recruitment, dominance, and settlement periods of particular species, the meaning of seasons is not at all trivial.

Intertidal zones are ecological spaces exposed to diverse physical environments, where the structures of benthic communities are changed easily by diverse events. Recently, artificial or natural disturbances to rocky intertidal zones occur more frequently and at more diverse intensities compared to the past. The effects of human leisure activities, large-scale development, and stronger storms due to climate change make the prediction of the results of disturbance more difficult. In par-

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ticular, external disturbances can cause direct damage to communities as well as opportunities for securing spaces for intertidal living dwellers with short life cycles. Therefore, understanding their spatial patterns and elucidating the mechanisms are meaningful in terms of ensuring biodiversity, and the conservation and restoration of living things.

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