An epibiont mediated increase in the susceptibility of *Mytilus edulis* to predation by *Nucella lapillus*

Jess Johansson*

*This study was conducted under the supervision of Professor Todd Gillis, Department of Integrative Biology, University of Guelph, Guelph, Ontario, Canada*

In marine organisms, the relationship that arises between two individuals where one organism biofouls the surface of a typically larger host organism is defined as epibiosis. Colonization of an organism by an epibiont can severely limit the feeding and reproductive abilities of the host and reduce overall fitness. Therefore, it is important to consider the benefits, if any, that the basibiont gains from this epibiotic relationship. In previous experiments, it was observed that barnacle epibionts protected *Mytilus edulis* from predation by *Carcinus maenas*, an intertidal predator. This study uses a choice experiment to deduce whether *Semibalanus balanoides* can mediate predation by another intertidal predator, the dogwhelk *Nucella lapillus*. *N. lapillus* preferred the substrate of barnacle-covered mussels to smooth-surface mussels, however specific preference to predate on these mussels was inconclusive from the results. The presence of *S. balanoides* on the shell surface of *M. edulis* decreased defensive behaviors, suggesting that the basibiont may be more vulnerable to predation because of this epibiotic relationship. If so, the relationship may border on parasitism if *M. edulis* fitness is significantly reduced.

Subject to extreme hydrodynamic forces, the body surfaces of intertidal organisms have evolved unique designs in response to the constant physical stresses in their environment. External modifications on the body surfaces of these organisms can alter the profile of the individual in the water column thereby increasing drag, which changes how environmental factors such as wave exposure influence the organism [1]. Such effects can have significant consequences on an animal’s fitness and reproductive success.

Epibiosis describes the unique ecological interaction between two marine organisms whereby one individual – the epibiont – colonizes the body surface of a larger host organism known as the basibiont. By definition, the respective fitness of both the epibiont and basibiont are not impaired by this non-symbiotic relationship [2]. However, an epibiont modifies the host’s morphology and subsequently alters the host’s interactions with the ambient environment [3]. In an extreme and fluctuating environment such as the intertidal zone where physical constraints are more pronounced, the consequences of morphological alterations have the potential to be more severe for the basibiont.

Basibionts provide novel habitat and refuge to their associated organisms and in certain contexts are considered ecosystem engineers. Ecosystem engineers represent an ecological classification of taxa whose widespread distribution and abundance have a significant impact on the structure of their ambient community [4]. In the intertidal zone the highly competitive blue mussel *Mytilus edulis* reduces available resources such as space for competitors, although its hard shell surfaces provide substrate for colonization by epibionts such as the barnacle *Semibalanus balanoides* [5]. However, colonization of their outer shell surface by *S. balanoides* significantly alters the morphology of the individual by increasing its surface area, which in turn increases drag on the organism [4]. This poses a significant threat to the survival of *M. edulis*, whose life history is greatly influenced by hydrodynamic forces due to their sessile benthic adult stage.

As adults, *M. edulis* attach to its substrate through a specialized organ in their foot called the byssus that secretes mechanically strong protinaceous filaments. These filaments, called byssal threads, are produced at a rate dependant on the hydrodynamic properties of their environment and adhere to the surrounding substrate in both rocky and muddy shores [6]. As constraining environmental forces increase, individuals will require greater byssal thread production in order to remain attached to the substrate, which is met by a greater energy demand for the organism. Most often, *M. edulis* form large aggregates referred to as ‘mussel beds,’ which reduce the hydrodynamic forces acting upon a solitary individual. In these instances, secreted byssal threads attach to both the substrate as well as neighboring mussels to create a high density cluster of individuals. This also provides simultaneous defense against predation by drilling organisms, such as the dogwhelk *Nucella lapillus* by...
reducing the total exposed surface area of each individual [7]. Dislodgement of *M. edulis* from their substrate represents a significant threat to individual survival [8]. In this scenario, the organism may be washed into areas of low flow which reduces the resources available to the individual and decreases fitness.

With the barnacle epibiont *S. balanoides*, the compensation for loss of fitness by the basibiont is returned in the form of defense against predation by a top predator, the starfish, who display an inherent preference for the smooth shelled mussels of the sublittoral [9]. However, this benefit could be negated in the northern Atlantic Ocean where the common intertidal predatory dogwhelk *N. lapillus* readily consumes both *M. edulis* and the epibiont *S. balanoides* [10, 11]. In defense to these attacks, species of mussels developed the capacity to use the byssal threads secreted from the byssus of their foot to trap and immobilize predatory dogwhelks [12, 13, 14]. Predation of thick-shelled organisms by *N. lapillus* is attributable to a modification to the ancestral radula, enabling the gastropod to drill through the hardened exterior of its chosen prey. Therefore, the addition of an epibiont to the outer shell of a prey organism does not necessarily hinder the feeding methods of *N. lapillus* compared to the methods engineered by other predators. Although the presence of an epibiont would provide a thicker surface through which the dogwhelk would be required to drill, the epibiont is typically unable to colonize the entire shell surface and portions of the original surface remain exposed to predation.

Although *M. edulis* is preferred over *S. balanoides* as a food source by *N. lapillus*, the increased prey density could further attract the dogwhelk to prey upon these individuals [10]. Therefore, if the presence of the epibiont *S. balanoides* on the shell of *M. edulis* increases the mussel’s susceptibility to predation by *N. lapillus*, then: (1) the distribution of *N. lapillus* will follow the distribution of barnacle-covered *M. edulis*, (2) barnacle-covered *M. edulis* individuals will experience greater predation rates by *N. lapillus* and (3) *N. lapillus* individuals foraging on barnacle-covered mussels will experience fewer defensive byssal thread attacks by *M. edulis*. As the presence of an epibiont increases drag on the basibiont, *M. edulis* would have to produce a greater number of byssal threads to attach to the substrate to prevent dislodgement. The actual production of byssal threads does not have the ability to increase due to the design of the byssus organ and only a finite number of byssal threads can be made. Barnacle-covered mussels will likely allocate more energy to producing byssal threads to further cement them to the substrate. This will reduce the risk of dislodgement.

**Materials and Methods**

The outlined experimental methods were primarily derived from Casey and Chattopadhyay, 2008 [7].

**Specimen Collection Methods**

A total of 72 *N. lapillus* individuals were collected from Green Point, St. Andrews, New Brunswick on August 11th 2009. Individuals were starved for four days to increase the probability that they would feed during the experimental period. The shell heights of all specimens were measured and individuals from each stage structure were separated into nine groups with similar size distributions to limit the influence of predator size on prey choice.

All *M. edulis* groups were chosen based on number of individuals within the bed and the presence of *S. balanoides* on their outer shell. All groups contained five *M. edulis* individuals; nine groups were colonized by *S. balanoides* and nine groups possessed smooth surfaces. Unwanted organisms and debris such as chitons, gastropods and rocks were manually removed from the mussel groups prior to the start of the experiment using a scrub brush.

**Experimental Procedure**

Three sea tables with dimensions of 50 x 35 x 11 cm were divided into three equal chambers by mesh lining to create a total of nine experimental replicates of the uniform treatment. Attached to each sea table was a constant source of flow to ensure that the properties of the water such as temperature, pH and salinity remained constant throughout the experiment.

Within each replicate were two mussel groups placed on opposite sides of the partitioned area – one group which was colonized by *S. balanoides* and one group with smooth shell surfaces. Prior to the start of the experiment, eight *N. lapillus* individuals were placed in the middle of the partition area after which no more manipulations occurred to the experimental design.

Throughout the first two days of the experimental period, qualitative observations were made of the behaviour and displacement of *N. lapillus*. From these observations, an appropriate observation interval of two hours over the course of a 12 hour period was initiated for the final three days of the experiment. Each observational account recorded the distribution of *N. lapillus* with regards to substrate; whether the dogwhelks were located on the tank bottom, the mussel group with barnacles or the smooth mussel group. For the mussel group colonized by *S. balanoides* it was also noted whether the dogwhelk was located on the shell of the individual mussel or on the portion covered by barnacles.

At the end of the five-day experimental period, several measurements were made. For all replicates and within each group, the mortality of *M. edulis* and *S. balanoides*, the number of byssal threads attached to the substrate and the number of byssal threads attached to each dogwhelk were recorded for further analysis. Images of defensive byssal thread attacks by *M. edulis* on *N. lapillus* were recorded on a camera and pieced together to create a time lapse video of
RESULTS

The distribution pattern of *N. lapillus* over the experimental period across the three substrate types – barnacle-covered mussels, smooth surface mussels and the aquarium substrate - was analyzed using ANOVA and found to be statistically significant ($F_{2,24} = 14.875$, $p=6.28 \cdot 10^{-5}$). This distribution pattern is depicted in Figure 1, whereby it should be noted that the surface of barnacle-covered mussels were significantly preferred to the other substrate types (Table 1).

The predation rate of *N. lapillus* on *M. edulis* was inferred from the number of drill holes present on the shell of each individual mussel. Comparisons between the number of holes drilled in the shells of barnacle-covered mussels versus smooth-surface mussels was not found to be significant (Student’s $t$-test, $p=0.1028$).

The number of defensive byssal threads attached to *N. lapillus* individuals distributed on *M. edulis* was used to assess the defensive capabilities of the differing mussel types. Statistically significant correlations were found between the number of defensive byssal threads produced and the number of *N. lapillus* individuals situated on an individual mussel for both barnacle-covered and smooth-surface mussel types (Pearson’s product moment correlation, $p=0.002756$ and $0.001786$ respectively). It was observed that smooth-surface mussels produced significantly more defensive byssal threads when predated by *N. lapillus* (Student’s $t$-test, $p=0.02572$) (Figure 2). However, no significant differences were observed between the differing mussel types with respect to the number of byssal threads attached to the aquarium substrate (Student’s $t$-test, $p=-0.232076$).

DISCUSSION

This lab study suggests that the presence of barnacle epibionts on the surface of mussels significantly alters the ecological distribution of their predator *N. lapillus*. However, no significant differences were observed with respect to the...
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predation rate of *N. lapillus* on either mussel type. It was seen that barnacle-covered mussels inherently produced fewer defensive byssal threads than smooth-surface barnacles when under predation by *N. lapillus* [13]. For the purposes of scientific discourse all data will be discussed regardless of statistical significance and the inability of data to meet this criterion will be further explained by the sample size limitations of the proposed experimental design.

When selecting a habitat by choice, the bias of an individual will opt to occupy the space that increases its personal fitness along a gradient [15]. Previous choice experiments by Davenport et al. 1996 have demonstrated that *N. lapillus* prefers both the substrate of barnacle encrusted rock and bare rock to that of mussel carpets, attributing the choice to the availability of a low-risk food source [13]. However, in the Northern Atlantic intertidal zone the epibiotic relationship between *S. balanoides* and *M. edulis* is widespread and has important implications for basibiont predation risk as well as the distribution of *N. lapillus*. Of the available substrata, *N. lapillus* preferred the direct proximity of a food source provided by either smooth-surfaced or barnacle-covered mussels (Figure 1). However, the observed significant preference of barnacle-covered mussels by *N. lapillus* must provide a selective advantage to the predator. Consequently, this preference creates a disadvantage for the basibiont *M. edulis* who now experiences greater predation pressures as a product of increased *N. lapillus* density. Predation risk is a considerable stressor for *M. edulis* as the presence of both dogwhelk effluent and individuals significantly increases the heart rate of *M. edulis* individuals [16]. The positive relationship between predator density and basibiont stress will ultimately result in an increase in predation susceptibility for the basibiont. Similarly, this susceptibility can also be argued as a decrease in basibiont fitness due to the additional energy expenditure required to maintain the stressed state of the individual. The regulation of predation pressure against *M. edulis* by barnacle epibionts has been previously observed in the green crab *Carcinus maenas*, where epibiont colonization of mussel beds results in increased predation [1]. By adversely affecting the fitness of the basibiont in terms of biotic stress, the neutrally defined epibiotic relationship between *M. edulis* and *S. balanoides* should fall under questioning.

It is important to consider that gastropod densities and distributions do not always generate accurate measures of foraging activity and therefore may provide poor inferences of predation rates [17]. Despite the consistency of the observed distribution pattern, where the majority of *N. lapillus* was distributed on the barnacle-covered mussels, the predation rates against the two differing mussel types were

![Figure 2: Barnacle covered mussels produce less byssal threads when attacked by *N. lapillus*. Statistically significant correlations were observed between the number of defensive byssal threads produces and the number of *N. lapillus* individuals predating on a given mussel for both barnacle-covered mussels and smooth-surface mussels. Only data collected from *M. edulis* mussels that possessed *N. lapillus* individuals on their shell surface was used (n=25 and 14 respectively). It predation rate of *N. lapillus* on either mussel type. It was seen that barnacle-covered mussels inherently produced fewer defensive byssal threads than smooth-surface barnacles when under predation by *N. lapillus* [13]. For the purposes of scientific discourse all data will be discussed regardless of statistical significance and the inability of data to meet this criterion will be further explained by the sample size limitations of the proposed experimental design.

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Table 1: Statistical Analysis of Collected Data.
Statistical analysis of the collected data for each of the three experimental predictions is listed. The data sets compared, statistical test utilized and subsequent p-values are noted. Statistical significance was inferred at an alpha value of 0.05.

| Distribution of N. lapillus over Experimental Period (Day 3 to 5) |
|---------------------------------------------------------------|
| **Comparison** | **Test** | **Test Statistic** | **Df** | **P value** |
| Barnacle-covered Mussels, Smooth-surface Mussels, Substrate | ANOVA | 14.87510152 | 2,24 | 6.28e^{-5} |
| Barnacle-covered Mussels, Smooth-surfaced Mussels | Student's T-Test | // | 16 | 3.47e^{-24} |
| Barnacle-covered Mussels, Substrate | Student's T-Test | // | 16 | 1.52e^{-25} |
| Smooth-surface Mussels, Substrate | Student's T-Test | // | 16 | 2.62e^{-6} |

**Predation by N. lapillus on M. edulis**

| Comparison | Test | Test Statistic | Df | P value |
|------------|------|----------------|----|---------|
| Number of holes drilled in Barnacle-covered Mussels, Smooth-surface Mussels | Student's T-Test | // | 10 | 0.103 |

**Byssal Threads Attached to N. lapillus**

| Comparison | Test | Slope (b) | Correlation Coefficient | P value |
|------------|------|-----------|-------------------------|---------|
| Byssal threads attached to N. lapillus on Barnacle-Covered Mussels | Linear Regression | 2.05 | 3.375637 | 0.00276 |
| Byssal threads attached to N. lapillus on Smooth-Surface Mussels | Linear Regression | 3.876712 | 3.61117 | 0.00179 |

| Comparison | Test | Test Statistic | Df | P value |
|------------|------|----------------|----|---------|
| Defensive byssal threads produced by Barnacle-covered Mussels, Smooth-surface mussels | Student's T-Test | // | 17 | 0.0257 |

not found to be statistically significant. As N. lapillus was observed actively feeding upon S. balanoides, it can be extrapolated that the higher density of dogwhelks on the mussel surface forced some individuals to consume the alternative prey. In field experiments, observations of N. lapillus have described a lower prey-handling time for S. balanoides than M. edulis [18]. This is attributable to both the smaller size of S. balanoides and the risk associated with its consumption. However, as S. balanoides sits erect on the outer surface of M. edulis shells, there is a considerable lack of refuge for the predator when foraging on its prey. Due to the smaller size of S. balanoides, it would be necessary to consume a greater number of individuals to gain the nutritional equivalent to one M. edulis. This increased foraging time is directly associated with an increase in predation risk by N. lapillus [18]. Although it may be more adaptive to prefer to consume barnacle-covered mussels, it is not feasible for all N. lapillus individuals distributed on these mussels to predate upon the basibiont simultaneously. There may be strong competitive forces amongst N. lapillus which
forces less dominant individuals to predate upon *S. balanoides* as opposed to the desired *M. edulis* [9].

The addition of *S. balanoides* to the shell surface of *M. edulis* also severely alters the exterior morphology of the organism. As intertidal organisms, mussels have evolved a specific shape to withstand the hydrodynamic forces of their ambient environment; therefore changes to this morphology pose a significant risk to dislodgement. Since *M. edulis* remain attached to the substrate via byssal threads, it would be expected that the barnacle-covered mussels would utilize a higher proportion of their energy for producing byssal threads for better adhesion. As such, it would be expected that these *M. edulis* individuals would have less available free energy to allocate to the production of defensive byssal threads to attack predators. However, only the latter proposed phenomenon was reflected in the results of this experiment (Figure 2). Although both mussel types produced a similar amount of byssal threads that adhered to the substrate of the aquarium, the barnacle-covered mussels produced significantly fewer defensive byssal threads. This is especially intriguing as these individuals experienced higher densities of *N. lapillus* on their surfaces, yet this increase in predation risk was met with fewer defensive behaviors. It is therefore possible that the presence of *S. balanoides* is causing stress to the basibiont, which results in weakened defensive abilities. This inability to cope with stress and subsequent poor allocation of resources could increase the susceptibility of *M. edulis* to predation by *N. lapillus*.

It is important to note that although possible explanations and preliminary theories have been mentioned in this discussion, many of the comparisons on which they were based are not statistically significant. This is ultimately due to limitations of experimental design. Constraints of time and space only allowed a limited number of repetitions to run simultaneously. This resulted in a small number of experimental replicates, which in turn reduces the significance of the trends observed. The aquaria used for the replicates were small which provided a limit for sample size of each replicate. This resulted in a low number of individuals within each repetition, which in turn reduced the significance of the trends observed.

Within the experimental design, a serious effort was made to exclude confounding variables. The height of all *N. lapillus* individuals was measured and size-structured to create a normally distributed predator population within each replicate. All dogwhelks were collected from the same shoreline to avoid the effects of wave-exposure from impounding the results. This was especially important as wave-exposed *N. lapillus* have greater foraging rates than wave-sheltered individuals [18].

The distribution pattern of *N. lapillus* atop of barnacle-covered mussels supports the hypothesis that the addition of a barnacle epibiont increases the mussel’s susceptibility to predation by dogwhelks. By attracting additional predatory individuals, the barnacle epibionts indirectly increase the stress experienced by their basibiont host – ultimately impairing their abilities to properly defend against attacks. However, despite this distribution pattern the predation rates on both barnacle-covered and smooth-surface mussels were not significantly different. The similar predation rates upon mussel individuals may have been affected by intraspecific competition among dogwhelks whereby weaker individuals are forced to consume the less desired prey, *S. balanoides*. An inclusive look at the overall predation rate (both barnacle and mussel prey types) should be analyzed in order to compare the environmental predation rate of both mussel types. If the overall predation rate of barnacle-covered mussels is greater than smooth-surface mussels, the original hypothesis may be supported. Future experiments should include fewer dogwhelk individuals per replicate in order to reduce intraspecific competition among the predators. The significant reduction in defensive byssal thread production by barnacle-covered mussels supports the hypothesis that *N. lapillus* increases the basibiont’s susceptibility to predation. However, the mechanisms have not been inferred and are beyond the scope of this paper.

The addition of the barnacle epibiont, *S. balanoides*, to the surface of *M. edulis* increases the basibiont’s susceptibility to predation by *N. lapillus*. From the collected data, it is impossible to detect which aspects of *M. edulis*’ life cycle, survival and fitness would be most affected by this epibiotic relationship. This is an emerging area for future research.

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