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**HOKKAIDO UNIVERSITY**
Inducible defenses in prey intensify predator cannibalism

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Abstract. Trophic cascades are often a potent force in ecological communities, but abiotic and biotic heterogeneity can diffuse their influence. For example, inducible defenses in many species create variation in prey edibility, and size-structured interactions, such as cannibalism, can shift predator diets away from heterospecific prey. Although both factors diffuse cascade strength by adding heterogeneity to trophic interactions, the consequences of their interaction remain poorly understood. We show that inducible defenses in tadpole prey greatly intensify cannibalism in predatory larval salamanders. The likelihood of cannibalism was also strongly influenced by asymmetries in salamander size that appear to be most important in the presence of defended prey. Hence, variation in prey edibility and the size structure of the predator may synergistically affect predator–prey population dynamics by reducing prey mortality and increasing predator mortality via cannibalism. We also suggest that the indirect effects of prey defenses may shape the evolution of predator traits that determine diet breadth and how trophic dynamics unfold in natural systems.

Key words: behavior; cannibalism; diet; food webs; Hynobius retardatus (Dunn); inducible defense; predation risk; Rana pirica (Matsui); salamander; trait-mediated indirect effects; trophic cascade.

INTRODUCTION

Classical views of trophic dynamics used simple food chains to argue that carnivores strongly control herbivore biomass, which in turn, influences plant biomass (Hairston et al. 1960, Rosenzweig 1971, 1973, Oksanen et al. 1981, Carpenter et al. 1987). Such “green world” or “exploitation” hypotheses therefore emphasize the importance of top-down control, and some systems are clearly driven this way (Estes and Palmisano 1974, Silliman and Bertness 2002). However, it is also clear that such striking trophic cascades do not operate in many systems because the reticulate nature of food webs, as well as other factors, can buffer their influence (Strong 1992, Polis and Strong 1996). For example, intraspecific heterogeneity or differentiation (Strong 1992), such as variation in plant edibility or quality, can strongly determine how effectively one species harvests the biomass of another in the food web and, thus, the trophic dynamics of the system (also see Leibold 1989, Persson 1999, Vos et al. 2004b).

Heterogeneity within a species can arise in a number of ways, but two factors seem to be emerging in terms of their importance to population and community dynamics. First, there is often considerable variation in the edibility of species (Leibold 1989, Strong 1992), and much of this variation may arise because of inducible defenses. The ecological significance of inducible defenses is well established (Agrawal 2001), and theory suggests that such plasticity can promote species coexistence and stabilize population dynamics (Matsuda et al. 1993, Bolker et al. 2003) as well as provide insight into the paradox of enrichment (Vos et al. 2004a, b) and the complexity–stability debate (Kondoh 2007). Moreover, because inducible defenses occur in numerous plant and animal species (Tolllian and Harvell 1999), the heterogeneity they create is likely to produce effects that operate throughout natural food webs.

Second, many species are composed of different size cohorts (Polis 1984) and the size-structured interactions that emerge in these situations can also serve to increase trophic heterogeneity within the food web. Cannibalism is one form of size-structured interaction that is prevalent in many food webs (Polis 1981, Persson 1999), and it can lead to interactions and dynamics that are not considered or predicted by unstructured models (Persson et al. 2003, Rudolf 2007a, b, 2008). For example, in amphibian systems these different dynamics can emerge because indirect interactions between just two species are possible when size-structured interactions like cannibalism are operating (Rudolf 2006). Moreover, recent models (Rudolf 2007b) suggest that the indirect interactions resulting from cannibalism can modify trophic cascades as well as the effects of enrichment on population dynamics.

Although the ecological importance of inducible defenses and cannibalism is increasingly recognized, to our knowledge, no empirical study has examined how their combined effects may influence the outcome of species interactions in natural populations. In this paper,
we show that induced morphological defenses in tadpole prey greatly intensify cannibalism among predatory salamanders. Moreover, asymmetries in the size of cannibalistic salamanders and conspecific prey also exerted a stronger influence on the probability of cannibalism in the presence of tadpole defenses. Hence, variation in the edibility of tadpole prey can exert strong trait-mediated indirect effects in this system, which may shape salamander trait evolution by increasing the likelihood of cannibalism.

**Materials and Methods**

*Study system and background*

Tadpoles of Japanese brown frogs (*Rana pirica* [Matsui]) and predatory larval salamanders (*Hynobius retardatus* [Dunn]) provide an excellent predator–prey system for studying the evolutionary and ecological significance of adaptive morphological plasticity (Kishida and Nishimura 2005, 2006). Both amphibian larvae exhibit antagonistic morphological plasticity in response to one another. Prey tadpoles typically develop bulgier bodies (Fig. 1a, left, Fig. 1b; see also Plate 1) by thickening their epithelium tissues in response to larval salamander risk cues (Kishida and Nishimura 2004). Because salamanders are gape-limited predators that swallow their prey whole, the bulgy morph is highly effective in reducing tadpole vulnerability to salamander predation because it is more difficult to swallow than non-induced tadpole morphs. In contrast, salamander larvae can produce a predaceous morph (Fig. 1a, right) having an enlarged gape that allows them to swallow larger prey (Michimae and Wakahara 2002). Although salamanders primarily consume tadpoles when they are available, they also can cannibalize conspecifics, and this diet shift is likely influenced by the prevalence of bulgy tadpoles, whose bodies are significantly wider than those of salamander conspecifics (Fig. 1b). Hence, we predicted that when salamanders cohabit with bulgier tadpoles, their difficulty in consuming this defended morph would indirectly intensify cannibalism among salamanders. We tested this and other hypotheses with field and laboratory experiments that compared tadpole and salamander survivorship and morphology when salamanders are present with either defended or undefended tadpoles.

*Field experiment in a natural pond*

Our field experiment was conducted in a mountain pond (area = 55 m²) located in Hekirichi, Hokuto, Hokkaido, Japan (41°53’ N, 140°34’ E). This pond had a limited tree canopy, a maximum depth of 0.5 m, and a soil bottom. Adult salamanders (*Hynobius retardatus*) and frogs (*Rana pirica*) typically begin spawning in April, and the larvae of both amphibians coexist and interact from spring to summer. Invertebrate predators such as dragonfly larvae were not very abundant; those that were present were restricted to patches of aquatic vegetation along the periphery of the pond.
After spawning, frog and salamander eggs began to hatch from early to mid May. Soon after hatching, interactions between both amphibian larvae led to induced morphological changes as they grew. By early June, tadpoles had fully expressed their defensive bulgy morphs whereas the salamander population was composed of large individuals having a large gape (predaceous morph) and small individuals having a small gape (non-predaceous morph; Fig. 1a). Before initiating the field experiment, we first estimated the density of both amphibian larvae by randomly placing 10 quadrats (80 x 80 cm) throughout the pond. Because it was difficult to reliably identify the species of each larva, this approach was only able to provide an estimate of overall amphibian larvae density (89.8 ± 21.1, mean ± SD). Hence, we also used dip nets to collect many larvae and returned these samples to the laboratory to determine the proportion of tadpole and salamander larvae in the population and their size distributions. Defended tadpoles and salamanders collected from this pond were classified (in 1-mm categories) according to body length (snout–vent length), which is an appropriate measurement of size for the study of morphological plasticity in *R. pircia* and *H. retardatus* (Kishida and Nishimura 2005, Kishida et al. 2009). After size measurements, larvae were maintained individually in containers having 200 mL of aged tap water.

We also collected sufficient numbers of undefended tadpoles (basic morph) from several nearby ponds that contained no salamanders. These defended tadpoles were conditioned to salamander predation risk for two days before initiating the experiment. We did so by randomly placing fifty undefended tadpoles into 13-L tanks (N = 15) having 4 L of aged tap water and three, freely roaming salamander larvae to serve as inducers of predation risk. During this conditioning period, the undefended tadpoles became acclimated to predation risk as evidenced by their reduced activity levels, which is a common response to predation risk in many amphibian larvae species (Skelly 1994, Relyea 2001, Van Buskirk 2002, Kishida et al. 2009). Although tadpoles clearly responded behaviorally to salamander predation risk, they continued to maintain the non-induced morph because our conditioning period was too short to allow development of the bulgy morph. After this conditioning period, undefended tadpoles were also classified in 1-mm increments of body length (snout–vent length).

We placed eight replicate, rectangular (80 x 80 x 80 cm) enclosures having PVC framing and nylon mesh (1-mm openings) on all sides into the pond. We added 5 L of pond silt and sand on the mesh bottom of each enclosure to serve as natural substratum. Enclosures were stocked with natural density and size distributions of both amphibian larvae based on the sampling just described. We randomly applied two treatments ("undefended" and "defended") to the enclosures, and each treatment was replicated four times. In the undefended enclosures, we added 16 salamanders and 74 tadpoles having the undefended morph. In the defend enclosures, we added 16 salamanders and 74 tadpoles having the defended morph. All animals used in the experiment were measured for body length and width (tadpoles) or body length and gape and head width (salamanders) before placing them in the enclosures. Consistent with our experimental objectives, initial tadpole morphology (Fig. 2a) differed between defended and undefended treatments (MANOVA, F_{1,6} = 706.20, P < 0.0001) but a significant trait x treatment interaction (MANOVA, F_{1,6} = 3764.81, P < 0.0001) indicated that only body width (F_{1,6} = 2944.79, P < 0.0001) and not body length (F_{1,6} = 1.27, P = 0.3025) was different. There were no differences in the body length and gape and head width...
(MANOVA, $F_{1,6} = 0.55$, $P = 0.4845$; Fig. 2b) of salamanders placed in the defended and undefended enclosures. This experiment was conducted for 3 days, which minimizes growth and the potential for morphological induction.

At the end of the experiment, all surviving tadpoles and salamanders were returned to the laboratory for measurement of body length and body width (tadpoles) and body length and gape and head width (salamanders). All measurements were made with digital calipers and the total number of surviving larvae was recorded.

Statistical analyses: field experiment

The proportion of salamanders and tadpoles surviving to the end of the experiment was analyzed with a one-way ANOVA that considered tadpole phenotype (defended, undefended) as a fixed effect. To analyze amphibian morphology at the end of the experiment, we performed MANOVAs on the enclosure means of final body length and body width (tadpoles) and final body length and gape and head width of salamanders. When necessary (e.g., a significant trait $\times$ treatment interaction), MANOVAs were followed up with one-way ANOVAs on each trait for each amphibian.

Laboratory experiment

To obtain a better understanding of the mechanisms underlying the patterns observed in our field experiment, we conducted a laboratory experiment using animals collected from the same ponds. We randomly applied two treatments (“defended” and “undefended”) to experimental units (polypropylene tanks, $24.5 \times 37 \times 13$ cm, $L \times W \times D$) filled with 10 L of aged tap water. Because we were not able to collect as many undefended tadpoles, this treatment was replicated 15 times, whereas the defended treatment was replicated 23 times. We placed two salamanders and 10 undefended tadpoles into each undefended replicate, and two salamanders and 10 defended tadpoles into each defended replicate. The number of amphibians placed in each tank was thus scaled down appropriately to match the densities of animals used in our field experiment. We were also careful to use defended (body length, $11.42 \pm 0.30$ mm; body width, $9.07 \pm 0.69$ mm; mean $\pm$ SE) and undefended (body length, $11.43 \pm 0.28$ mm; body width, $6.21 \pm 0.60$ mm; mean $\pm$ SE) tadpoles that were of similar size to those used in the field experiment. Salamanders were also categorized based on body size (large vs. small) and photographed ventrally and laterally in a glass chamber before placing one individual of each size in each replicate. Resulting images were projected onto a computer monitor to measure salamander body length and gape and head width.

At 24 hours after the start of the experiment, we recorded tadpole activity (moving or stationary) to determine how morphological status (defended or
undefended) influenced their behavior. After three days, the experiment was terminated and we counted the number of surviving tadpoles and salamanders. We also took ventral and lateral photographs of all surviving salamanders to allow measurement of final morphological traits. In addition, comparison of these photos with our original photos allowed us to identify those individuals (large and/or small) that had survived to the end of the experiment. Individual identification was based on shape and small scars in body and tail parts evident in initial and final photos.

Statistical analyses: laboratory experiment

We used one-way ANOVAs to examine how tadpole phenotype (defended, undefended) influenced tadpole activity levels after 24 hours and tadpole survivorship at the end of the 3-day experiment. Multiple logistic regression on binomial survivorship data of salamanders in the defended treatment to determine if tadpole activity levels, in addition to salamander morphology, were important to the probability of cannibalism. Finally, because we found that the gape to head-width ratio was important, we examined whether these variables were influenced by differential gape and head-width growth among our experimental treatments.

RESULTS

Field experiment

Tadpole survivorship was significantly ($F_{1,6} = 85.71, P < 0.0001$; Fig. 3a) higher in the defended treatment (85%) than in the undefended treatment (58%). In contrast, salamander survivorship was significantly ($F_{1,6} = 8.73, P = 0.026$) lower in the defended treatment (77%) than in the undefended treatment (89%; Fig. 3b).

At the end of the experiment, tadpole morphology continued to differ among the defended and undefended treatments (MANOVA, $F_{1,6} = 107.29, P < 0.0001$; Fig. 2a), but each trait responded differently (MANOVA, trait × treatment interaction, $F_{1,6} = 330.51, P < 0.0001$). Tadpoles in the defended treatment still had wider bodies than those in the undefended treatment ($F_{1,6} = 406.54, P < 0.0001$), but tadpoles in the undefended treatment were longer than those in the defended treatment ($F_{1,6} = 25.00, P = 0.0025$).

Salamanders also responded strongly to our experimental treatments with those in the defended treatment having longer bodies, wider heads, and wider gapes than those in the undefended treatment (MANOVA, $F_{1,6} = 16.05, P = 0.0071$; Fig. 2b). The absence of a significant trait × treatment interaction ($F_{2,5} = 0.88, P = 0.47$) revealed that all three traits responded similarly to the presence of defended tadpoles.

Laboratory experiment

Tadpole survivorship was significantly (ANOVA, $F_{1,36} = 21.88, P < 0.0001$) higher in the defended treatment (88.3%) than in the undefended treatment (70.1%; Fig. 4a) and defended tadpoles were more active (ANOVA, $F_{1,36} = 4.59, P = 0.039$) than undefended tadpoles (Fig. 4b). Multiple logistic regression examined the importance of the ratio of initial body length of the large vs. small salamander and the ratio of the initial gape width of the large salamander to the initial head width of the small salamander. This full model revealed no significant effect of length ratio or any interaction involving length ratio (likelihood ratio tests, all $\chi^2 < 2$)
Hence, we sequentially removed terms involving length ratio from the model (\(P\) values of removed terms were always \(P \geq 0.58\)). This approach yields a final model comprised of tadpole phenotype, the gape-width to head-width ratio, and their interaction. A nonsignificant lack-of-fit statistic (\(\chi^2_{30} = 28.45, P = 0.5462\)) indicated that the final model would not benefit from additional terms. We found that salamander survivorship was significantly (likelihood ratio test, \(\chi^2_1 = 9.71, P = 0.0018\)) lower in the defended treatment than in the undefended treatment (Fig. 4c). In 13 out of 23 replicates of the defended treatment, and three out 15 replicates of the undefended treatment, the small salamander was cannibalized. Furthermore, as differences in the gape-width to head-width ratio increased, the probability of cannibalism increased (\(\chi^2_1 = 15.35, P = 0.0001\)) and there was a strong trend (\(\chi^2_1 = 3.32, P = 0.0685\)) for this effect to be significantly stronger in the defended vs. the undefended treatment (Fig. 4c).

Our analysis exploring the importance of salamander gape-width to head-width ratio and defended tadpole activity levels to salamander survivorship confirmed that the probability of cannibalism increased as differences in the gape-width to head-width ratio increased (likelihood ratio test, \(\chi^2_1 = 9.95, P = 0.0016\)). However, we were unable to detect any significant effect of defended tadpole activity level (\(\chi^2_1 = 0.43, P = 0.5114\)) or its interaction with gape-width to head-width ratio (\(\chi^2_1 = 0.003, P = 0.9575\)).

We detected no significant differences in the gape-width and head-width growth of surviving salamanders among our experimental treatments (MANOVA, \(F_{1,58} = 1.63, P = 0.2062\)). Hence, interactions between large and small salamanders were not driven by differential growth during the experiment.

**DISCUSSION**

Our field experiment found that tadpole survivorship was higher in the defended than in the undefended
treatment (Fig. 3a). The body length and width of defended tadpoles did not change during the experiment, suggesting that their improved survivorship was directly due to their defended status (bulgy body) at the beginning of the experiment. In contrast, the mortality of undefended tadpoles was 27% higher, indicating that salamander foraging success is strongly influenced by tadpole morphology. These results agree with those of our laboratory experiment where tadpole mortality was 18.2% higher in the undefended than in the defended treatments (Fig. 4a).

By the end of the field experiment, undefended tadpoles still had narrower bodies than defended tadpoles (Fig. 2a) and were thus more vulnerable to salamander predation. The lack of change in the body width of undefended tadpoles also indicates that our experimental duration was insufficient for morphological induction to occur. Indeed, the time required for expression of the bulgy morph under similar experimental conditions is ~1 week (Kishida et al. 2006, 2007). Moreover, at the end of the experiment, surviving tadpoles from the undefended treatment did not have the thickened epithelial tissue that is a diagnostic feature of the defended morphology (Kishida and Nishimura 2004). In contrast, surviving tadpoles in the undefended treatment were significantly longer than those in the defended treatment, suggesting that shorter individuals are also more vulnerable to salamander predation.

Although the increased survival of defended vs. undefended tadpoles in the field likely reflects differences in their respective morphologies, other factors, such as behavior, also may have been operating. We attempted to homogenize potential behavioral effects by conditioning undefended tadpoles to predation risk before initiating our experiment because tadpole experience or naivete can obviously influence survivorship under predation risk. Previous research has shown that other frog tadpole species reduce activity levels to reduce their likelihood of being detected by predators (Skelly 1994, Relyea 2001, Van Buskirk 2002). In our field experiment, the defended tadpoles had experienced salamander predation risk in the pond before we collected them so one might expect them to be less active than naive, undefended tadpoles. Reduced activity levels are expected to reduce both detection by and encounters with salamanders, thereby promoting increased tadpole survivorship. Our laboratory results do not support this hypothesis because defended tadpoles were actually more active than undefended tadpoles (Fig. 4b). Hence, the protection conferred by the bulgy morph may allow tadpoles to forage more actively in the presence of predation risk, whereas undefended tadpoles must compensate for their lack of this defense with lower activity levels that likely reduce their conspicuousness to predators. Such trait compensation, especially between morphological and behavioral traits, is a strategy that can provide fitness advantages in environments where predation risk varies temporally and spatially (DeWitt et al. 1999, Rundle and Brönmark 2001).

Salamander survivorship in the field was reduced by 12% when defended vs. undefended tadpoles were available as prey (Fig. 3b). This reduction in survivorship likely occurred because the bulgy bodies of defended tadpoles, which are wider than those of salamander conspecifics (Fig. 1b) and thus more difficult to swallow, intensified cannibalism among salamanders. This interpretation is supported by our laboratory results showing that the likelihood of cannibalism increased considerably as the ratio of gape width to head width of large vs. small salamanders increased (Fig. 4c). Although this pattern was evident in both the defended and undefended tadpole treatments, our data suggest that the influence of changes in this ratio on cannibalism was more important in the defended treatment. For example, an 80% probability of cannibalism occurred at a gape-width to head-width ratio of ~1.3 in the defended treatment, whereas a gape-width to head-width ratio of ~2.0 was required before such high cannibalism rates occurred in the undefended treatment. Although undefended tadpoles are also significantly wider than salamander conspecifics (Fig. 1b), overall cannibalism in this treatment was very low (only three out 15 small salamanders were cannibalized). In fact, two of the three cannibalism events we observed in the undefended treatment did not occur until the asymmetry in the gape width and head width of large and small salamanders, respectively, was quite high (a ratio of ~1.4 or more). Hence, these salamanders appear capable of discriminating between heterospecific and conspecific prey (see Pfennig and Collins 1993 for an example involving kin recognition by cannibalistic salamanders) and such discrimination may be strongly influenced by this asymmetry.

Our analysis highlights the importance of salamander size asymmetries to cannibalistic interactions in the presence of defended tadpoles, but the higher activity of defended tadpoles may have increased salamander activity that, in turn, increased encounters among salamanders and the likelihood of cannibalism. Although this intuitive mechanism may have been operating, further analysis of our laboratory results provided no evidence for it. Indeed, the probability of cannibalism in the presence of defended tadpoles was again strongly influenced by salamander gape-width to head-width ratio, whereas tadpole activity levels had no significant effect. Hence, while we acknowledge that tadpole activity levels have the potential to influence this predator–prey interaction, their effect was surprisingly weak in our laboratory experiment where we were able to measure both variables carefully.

Recent work demonstrates that trait-mediated indirect interactions are important in cannibalistic systems (Rudolf 2006, 2007a, b, 2008), and our study uniquely shows that inducible defenses in prey may be ultimately responsible for their emergence. The existence of trait-
mediated indirect effects in three-species food chains is well established (Werner and Peacor 2003, Schmitz et al. 2004), as is their influence on trophic cascades (Schmitz et al. 1997, 2004, Trussell et al. 2002, 2006a) and ecosystem function (Trussell et al. 2006b, 2008, Schmitz et al. 2008). Much of this work has focused on the cascading effects of prey habitat or diet shifts in response to predation risk (Schmitz et al. 1997, 2004, Werner and Peacor 2003, Trussell et al. 2006a), but little effort has explored how such indirect effects are transmitted by predator diet shifts in response to induced morphological defenses in prey (but see Raimondi et al. 2000). Our study demonstrates that the heterogeneity in prey edibility created by tadpole inducible defenses causes strong trait-mediated indirect effects in this amphibian system by increasing the intensity of cannibalistic behavior among salamanders and relaxing predation on tadpole prey. Although it is clear that prey heterogeneity and cannibalism can independently affect trophic cascades (Strong 1992, Persson et al. 2003, Rudolf 2007a, b), we suggest that the synergistic effects of tadpole defenses and cannibalism, both of which are ubiquitous in natural systems, may have particularly strong effects on cascade strength.

It is also likely that emergence of cannibalism as a trait-mediated indirect effect of inducible defense can strongly influence amphibian population dynamics. Considerable theory has explored how inducible defenses may regulate population dynamics. In some cases, inducible defenses create negative feedbacks between predator density and prey mortality that stabilize predator–prey dynamics (Ruxton and Lima 1997, Vos et al. 2004b), whereas others have shown that they can destabilize these dynamics (Luttbeg and Schmitz 2000, Kopp and Gabriel 2006). Our results suggest that tadpole defenses and the emergence of cannibalism may ultimately stabilize tadpole–salamander dynamics, as recent models suggest (Rudolf 2007a, b), by relaxing the intensity of salamander predation on tadpoles and increasing salamander mortality rates. However, longer term experiments are needed to fully understand the population and community consequences of cannibalism in this and other systems.

Our study also suggests that tadpole defenses can strongly influence phenotypic evolution in salamanders by intensifying phenotypic selection on salamanders through their enhancement of cannibalistic behavior. In the field, surviving salamanders that were maintained with defended tadpoles were longer, and had wider gaps and wider heads than those maintained with undefended tadpoles (Fig. 2b), suggesting that selection on salamander morphology was intense. Although previous work has shown that predaceous salamander morphs having a large gape can be induced by high tadpole density (Michaemae and Wakahara 2002), such induction, as in tadpole defenses, typically requires more time (7–10 days, Kishida et al. 2009) to develop than was possible in our experiment. Moreover, in our laboratory experiment, where we were able to measure individual salamander growth, we did not detect significant differences in the gape and head-width growth of surviving salamanders among our treatments. Hence, because larval salamanders are gape-limited predators, defended tadpoles may directly and indirectly, via their enhancement of cannibalism, drive selection for larger gape and head width in salamanders.

The prevalence of intraguild predation, including cannibalism (Polis 1981, Persson 1999, Rudolf 2006a, 2007a, b, 2008, Rosenheim 2007), in natural systems may be strongly connected to the expression of inducible defenses in numerous taxa (Tollrian and Harvell 1999). We found that cannibalism was intensified by the expression of inducible changes in prey morphology, but predator-induced changes in prey behavior also may cause predators to shift to less vigilant prey, including conspecifics. Regardless of the nature of the defense, we suggest trait-mediated indirect effects, which emerge because of predator diet shifts in response to prey defenses, play a key role in the evolution of predator traits and in maintaining predator diet breadth and cannibalism that, in turn, determine how trophic dynamics unfold in natural systems.

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LITERATURE CITED


