Biology of the conifer needle scale, *Nuculaspis abietis* (Hemiptera: Diaspididae), in northern Iran and parasitism by *Aspidiotiphagus citrinus* (Hymenoptera: Aphelinidae)

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**Key words.** *Nuculaspis abietis*, Diaspididae, Hemiptera, *Aspidiotiphagus citrinus*, Aphelinidae, Hymenoptera, *Picea abies*, Norway spruce, development, host range, parasitism, Iran

**Abstract.** The conifer needle scale, *Nuculaspis abietis* (Schrank) emerged as an important pest of conifers in the Kelardasht region of Mazandaran province, Iran, in the late 1990’s. This pest feeds on conifer needles and twigs causing needle drop and branch desiccation. Its discovery in Kelardasht in Mazandaran Province necessitated a local quarantine of conifers in the genera *Picea* and *Pinus*. We studied the life history and ecology of this scale on Norway spruce, *Picea abies*, under laboratory and natural field conditions. The complete life cycle of females required 206.4 ± 5.7 days in the laboratory (25 ± 1°C, 65–75% RH, 14L : 10D) and 315.7 ± 40.0 days under field conditions. The sex ratio ranged from 59% female for second instar nymphs to 71% female for adults. *Aspidiotiphagus citrinus* (Crawford) (Hymenoptera: Aphelinidae) was found naturally parasitizing the scale and overwintering in the larval stage on second instar nymphs. First generation adult wasps emerged in spring from overwintered second instar nymphs to parasitize 64.75% of first instar scales. Second generation wasps emerged from early September to mid-October and parasitized 19.75% of second instar scales, for a cumulative parasitism rate of 84.5%.

**INTRODUCTION**

The conifer needle scale, *Nuculaspis abietis* (Schrank) (= *Coccus abietis*, *Aspidiotus abietis*) (Danzig, 1967) is known to attack conifers from the genera *Abies*, *Juniperus*, *Picea*, *Pinus*, and *Pseudotsuga* in Europe, Russia, and North America (Danzig, 1993). Borchsenius (1939) described the second instar nymph and Graf & Weber (1996) described the damage done by this pest to *Pinus sylvestris* L. In southern Germany, *N. abietis* has one generation per year and hibernates as a second instar nymph (Kostarab & Kozar, 1988), but little else is known about the biology of this insect. In 1995, *N. abietis* was reported in the Kelardasht region on blue and green Norway spruce (*Picea pungens* and *P. abies*), its first record in Iran (Abai, 1995). Given the pest’s absence from other regions of Mazandaran province, a regional quarantine of conifers in the genera *Picea*, *Abies*, and *Pinus* in Kelardasht.

The Norway spruce, *Picea abies* (L.), is a large tree that can reach 40 m in height and >100 cm dbh (diameter at breast height). The natural range of this tree encompasses most of northern, central, and southern Europe but it is largely restricted to montane habitat (elevations above 400 m) south of 47°N latitude. An ecologically flexible species (Platonova, 2004), it prefers a cool climate and abundant rainfall (Banan & Fazilat, 1984). Norway spruce was planted for the first time in Kelardasht, Lagim and Sangedeh (Mazandaran Province, Iran) on 75–80 ha of land in the 1970’s (Mirbadin & Sagheb, 1991). Since that time, it has been planted extensively in government-sponsored reforestation projects in northern Iran and has become popular as an ornamental tree in urban settings.

The primary objective of this investigation was to characterize the biology and seasonal population dynamics of the conifer needle scale on Norway spruce in Mazandaran Province. Secondary objectives were to survey patterns of host use in the area and identify any parasitoids of *N. abietis* that might be present.

**MATERIAL AND METHODS**

**Study location**

Kelardasht is located in the province of Mazandaran in northern Iran, 55 km south of the Caspian Sea at 1200 m a.s.l. Yearly rainfall averages 496 mm, and the climate is cool and humid (Mirbadin & Sagheb, 1991).

**Morphological description of life stages**

A series of Norway spruce terminals bearing *N. abietis* were collected from naturally infested trees in Kelardasht. A total of 40 individuals of each representative life stage were examined, measured and photographed in order to characterize distinguishing features.

**Development under laboratory and field conditions**

Two-year-old potted Norway spruce saplings (*n* = 10) were obtained from the Kelardasht Tree Nursery and held in a growth chamber at 25 ± 1°C, 65–75% RH, and a 14L : 10D photoperiod under fluorescent lighting. Infestation was accomplished by...
placing a branch with crawlers of *N. abietis* onto each sapling for a period of two weeks. After the crawlers had settled, 20 nymphs on each sapling were marked by attaching numbered tags to needles bearing one or two nymphs each. Nymphs were sexed in second nymphal instar, the first stage at which gender can be distinguished. All tagged nymphs were examined under a hand lens daily during the first instar, and weekly thereafter, to monitor their development. All sampling was non-destructive. The first molt is clearly discernable as an abrupt color change to orange as the shed exuvia adheres to the scale cover. Thereafter, the scale does not adhere to the body of the insect and was carefully raised from one edge with a dissecting needle to detect ovi-position by females and pupation by males. Scale covers were then replaced with care so as not to harm the insects beneath.

We determined the developmental period under field conditions following the method of McClure (1985). Four infested Norway spruce trees in Kelardasht were sampled weekly from January to December, 2000, by removing one infected branch from each tree. Branches were taken to the laboratory and ten insects were dissected from each branch under a stereomicroscope to determine their growth stages. Mean numbers of insects in each growth stage were then calculated from the average counts of each stage on all four branches.

**Reproductive biology**

In order to estimate female egg load, we collected infested branches from Norway spruce trees in the Kelardasht tree-nursery in early June, 2000, and transported them to the laboratory. There, we dissected out the ovaries of gravid females (*n* = 100) in a drop of saline solution on a microscope slide under a
TABLE 1. Mean (± SEM) duration (in days) of immature life stages of *N. abietis* raised (n = 200) on Norway spruce saplings under controlled laboratory conditions of 25 ± 1°C, 65–75% RH, and 14L : 10D photoperiod and those observed on established Norway spruce trees under field conditions in Kelardasht, Iran, in 2000.

| Life stage                      | Females  | Males  |
|---------------------------------|----------|--------|
| **Laboratory**                  |          |        |
| Egg                             | 4.1 ± 0.2|        |
| First instar nymph              | 21.6 ± 0.7|       |
| Second instar nymph             | 153.8 ± 3.5| 174.6 ± 2.2|
| Third instar nymph / prepupa    | 27.3 ± 2.5| 6.1 ± 0.2|
| Pupa                            | 10.0 ± 1.1|        |
| Total                           | 206.4 ± 5.7| 215.2 ± 4.9|
| **Field**                       |          |        |
| Egg                             | 7.1 ± 0.5 |        |
| First instar nymph              | 29.2 ± 1.4|        |
| Second instar nymph             | 279.5 ± 4.2| 273.7 ± 4.1|
| Third instar nymph / prepupa    | 57.9 ± 3.2| 7.5 ± 0.8|
| Pupa                            | 14.2 ± 1.4|        |
| Total                           | 315.7 ± 9.3| 329.4 ± 8.7|

stereomicroscope as described by McClure (1983). To determine fertility, gravid females (n = 30) were each isolated on an excised twig of *P. abies* in a Petri dish (11 cm diam. × 1 cm ht.) held in a growth chamber under the same conditions as described above. To preserve plant condition, the ends of the twigs were encased in a balls of moistened cotton that were refreshed every second day. To prevent the escape of crawlers, the edges of each Petri dish was glued closed. The number of eclosing crawlers were counted and removed daily and the dishes re-sealed after each observation.

Realized female fecundity was assessed in a separate trial. Gravid females (n = 30) were each placed on a twig of *P. abies* in a glass tube (2.4 cm diam. × 10 cm ht.). The ends of the twigs were embedded in balls of moist cotton. Each day, the female’s scale was partially raised and all the eggs produced were removed and counted until the female died.

The sex ratio of the scale population was estimated by collecting branches from the aforementioned infested trees, selected to bear suitable numbers of either second instar nymphs (n = 10) or adults (n = 10). Both life stages were then sexed by observing 100 individuals from each branch under the stereomicroscope.

**Host range and local distribution**

The distribution of *N. abietis* in Mazandaran was determined by inspecting potential coniferous host plants in various regions of the province from early April to early June, 2000. Three sites at least one kilometer apart were sampled in each region and a minimum of 10 trees of each species were inspected at each site. To estimate the relative susceptibility of the primary host species, one branch was collected from each of three infested trees of *Picea abies*, *Picea pungens*, *Picea nordmanniana*, *Abies alba* and *Abies nordmanniana* in the Kelardasht tree nursery in early September, 2000. These were transported to the laboratory where all second instar nymphs were counted on 100 needles of each branch under a stereomicroscope.

**Seasonal population dynamics**

From December 1999 to December 2000, one branch was sampled weekly from each of four infested Norway spruce trees near Kelardasht. Following transportation to the laboratory, all scale life stages were counted on 100 needles of each branch under a stereomicroscope as described by Kostarab & Kozar (1988). These data were used to estimate seasonal scale population dynamics.

**Parasitism**

A single branch was cut weekly from each of four Norway spruce trees and taken to the laboratory where infested plant parts were excised and held in glass tubes (as described above). The containers were kept in a growth chamber, under the conditions described above, until the emergence of parasitoids.

We determined percentage parasitism using the method of McClure (1983). One branch was cut weekly from each of the four infested *P. abies* trees and a sample of 100 nymphal scales from each branch was selected at random. These were examined under a stereomicroscope to determine the presence of parasitism. Percent parasitism was calculated as the sum of parasitized scales plus those with emergence holes, divided by the total number of scales, and multiplied by 100. This approach is expected to slightly underestimate actual rates of parasitism because eggs are not directly observable. However, it was possible to distinguish emerged mummies of the first and second generations.

**RESULTS**

**Morphological description of life stages**

Eggs of *N. abietis* averaged 0.23 ± 0.02 mm (mean ± SEM) in length by 0.13 ± 0.01 mm in width. As they are
laid, the oval eggs are yellowish orange, soft and attached together in a string. As the eggs dry, they separate and become paler in color. Ocular points become visible on the eggs one or two days prior to eclosion. The emergence of first instar nymphs, or crawlers, requires 70 ± 20 min (n = 30) (Fig. 1A). Crawlers average 0.24 ± 0.02 mm in length by 0.14 ± 0.01 mm in width. They are yellow, oval in shape, with antennae and three pairs of legs. Shortly after settling, crawlers secrete a white cap, the first scale (mean = 0.48 ± 0.01 mm in length by 0.33 ± 0.01 mm in width; Fig. 1B) that gradually changes to orange in the nipple stage. The gender of crawlers cannot be distinguished by visual inspection.

The female second instar nymph is oval in shape and pale yellow in color with indistinct abdominal segmentation. The scale is brown or pale brown and averages 1.66 ± 0.04 mm in length by 0.62 ± 0.02 mm in width. In comparison, the male second instar nymph is spindle-shaped, dark yellow in color, has distinct abdominal segmentation, and ocular points clearly visible on the head (Fig. 1D). The scale is dark brown or black and averages 1.65 ± 0.03 mm in length by 0.61 ± 0.01 mm in width. The male insect secretes a silk wax underneath the scale, whereas no such silk is secreted by females.

The female third instar nymph, or virgin adult, averaged 0.79 ± 0.02 mm in length by 0.50 ± 0.02 mm in width. These are oval in shape, yellow in color, have indistinct abdominal segmentation, and secrete a third white scale that male insects do not. In contrast, male insects pass through preupal and pupal phases. The pre-pupa averages 0.82 ± 0.02 mm in length by 0.48 ± 0.02 mm in width, is spindle-shaped and dark yellow with two pairs of ocular points on the head, one pair on the frons and one pair on the ventral surface of the head (Fig. 1E). The pupa is spindle-shaped, dark yellow, and averages 1.02 ± 0.04 mm in length by 0.36 ± 0.01 mm in width.

In the mature adult female (Fig. 1C), both the body and the scale are larger than in virgin females, (body mean = 1.72 ± 0.01 mm in length by 0.55 ± 0.01 mm in width; scale mean = 2.70 ± 0.03 mm in length by 0.79 ± 0.01 mm in width). Once gravid, the female ceases growth and honeydew secretion. The free-living adult male lacks functional mouthparts, is spindle-shaped (mean = 1.08 ± 0.03 mm in length), yellow with a wide black band on thorax and possesses a single pair of clear wings, one pair of ocelli, antennae with 10 segments, and a penis (0.24 ± 0.01 mm in length). Since neither the prepupa nor the pupae of males secrete a scale, these stages remain under the scale of the second instar nymph until emergence of the mature adult.

**Development under laboratory and field conditions**

The mean durations of immature life stages and total developmental times for female and male insects were considerably shorter under laboratory conditions than under the field conditions prevailing in Kelardasht in 1999 (Table 1). Males took nine and 14 days longer to complete development than did females in the two studies, respectively. The longevity of adult females under natural conditions was 31.6 ± 5.7 days.

**Reproductive biology**

As is generally true for most diaspидid species, the embryonic development of eggs is staggered in accordance with their sequence of oviposition and is temperature-dependent, the eclosion of crawlers requiring anywhere from a number of hours to eight days (AR, personal observation). Consequently, both egg and first instar nymphs are often seen together under the same adult female’s scale. Newly eclosed nymphs leave the scales and settle on needles within 6–12 h, although a few sometimes settle underneath their mother’s scale. Under field conditions in Kelardasht, the oviposition period ranged from 16 to 32 days (mean = 18.7 ± 1.4 d, n = 30). Dissections of gravid females collected from infested branches revealed egg loads ranging from 41 to 74 eggs (mean = 64.2 ± 5.2). In the fecundity trial, females (n = 30) laid an average of 6.5 ± 1.7 eggs per day (range = 3–14) for an average of 12.3 ± 2.1 days, yielding an average lifetime fecundity of 57.3 ± 5.1 eggs (range = 32–64). Collection of dispersing crawlers from Petri dishes yielded a mean of 54.0 ± 4.4 nymphs per female (n
s = 30), suggesting that egg fertility was very high. The sex ratio was female-biased in both second instar nymphs (0.59) and adults (0.71), but neither of these values represented significant departures from 0.5 (Chi square, Goodness-of-fit, α = 0.05).

Host range and local distribution

The survey revealed that N. abietis was absent from all sampled regions of Mazandaran province except for Kelardasht, where it occurred primarily on species of Picea and Abies (Table 2). The nursery samples revealed significant variation among tree species (F4,10 = 276.82, P < 0.001) in numbers of scale per 100 needles, P. abies and P. pungens bearing significantly more (Tukey HSD, Dα = 0.05) than the other species that were not different from each other (Mean ± SE, P. abies: 123 ± 4.3, P. pungens: 118 ± 5.2, P. nordmanniana: 23 ± 3.0, P. alba: 15 ± 1.7, and A. nordmanniana: 11 ± 1.5).

Seasonal abundance

Nuculaspis abietis was observed to complete one generation per year in Kelardasht, hibernating as a second instar nymph. Adult females first appeared in early April, peaked in abundance in late April, and had disappeared by late July (Fig. 2). Male second instars became prepupa in late April and pupation was observed in early May. Adult males peaked in abundance in the middle of June and had largely disappeared by the beginning of July.

Oviposition began in late May and the number of eggs and neonate crawlers found under scales peaked in early June at 777.3 ± 62.4 per 100 needles. First instar nymphs began settling on needles at the beginning of June and reached peak abundance in the middle of July, whereas second instar nymphs were first observed in late June and peaked in abundance in early September (Fig. 3). The mean oviposition period under field conditions was 18.7 ± 1.4 days (minimum = 16, maximum = 23, n = 30).

Parasitism

The only parasitoid species emerging from samples was Aspidiotiphagus citrinus (Crawford) (Hymenoptera: Aphelinidae) (Fig. 4). The wasp larva develops as a solitary ectoparasitoid, although superparasitism can occur and in one case we observed two larvae in a single scale host. In the Kelardasht region, A. citrinus has two generations per year. Overwintered wasps emerged as adults from the end of April until late June and began attacking first instar nymphs of N. abietis in May (Fig. 5). Wasps of the second generation emerged over a period from early September to mid October and attacked second instar N. abietis beginning in early September, their progeny overwintering as larvae on their hosts.

Under field conditions in Kelardasht, cumulative parasitism of N. abietis by first (overwintered) generation A. citrinus reached 64.75% by the middle of June when all second generation wasps had emerged from their hosts. Wasps of the second generation parasitized 19.75% of
scales from early September until late October, for a cumulative parasitism rate of 84.5%.

**DISCUSSION**

Our results indicate that *N. abietis* has one generation per year in northern Iran, primarily feeding on species of *Picea* (*P. abies, P. alba, P. pungens, P. nordmanniana* and *P. nitidiformis*) and *Abies* (*A. sibrica, A. alba* and *A. nordmanniana*). Although *Pinus sylvestris* may also be attacked, the specimens of *Juniperus, Thuja* and *Taxus* spp. that we examined in the infested region of Kelardasht were not. Scale outbreaks are often confined to one host species even when other host species occur in the area (Edmunds, 1973; McClure, 1986). This is thought to occur because nymphal survival is typically highest on the parent tree, lower on other trees of the same species, and very low on host trees of other species. Based on the samples obtained from the infested nursery, both *P. abies* and *P. pungens* appear to be preferred in northern Iran and should be priority targets for phytosanitary inspectors in adjacent localities.

The sex ratio of *N. abietis* was slightly female-biased in the second instar, the earliest stage in which sex determination is possible, and tended toward greater female-bias in the adult stage. Although the deviation from a 50% sex ratio was not significant, the trend is suggestive of differential male mortality during the course of development. Changes in the sex ratio of black pine leaf scale, *Nuculaspis californica*, have been inferred to reflect differential mortality according to gender (Alstad et al., 1980; Alstad & Edmunds, 1983, 1987). In the latter studies, hatching ratios were significantly female-biased, and subsequent differential mortality further increased this bias prior to emergence of adult males. In addition, Edmunds & Alstad (1985) showed that haploid males of *N. californica* were more sensitive to malathion than diploid females, so insecticide applications may have the potential to further skew sex ratios in these insects.

Reported parasitoids of *N. abietis* include *Aphytis abnormis* (Howard), *A. mytilaspis* (Le Baron), *Prospaltella aspidioticola* Mercet, *P. aurantii* (Howard), *P. gigas* Chumakova, *P. leucaspis* Mercet, *Coccophagus similis* (Masi) (Kostarab & Kozar, 1988) and *Encarsia aspidioticola* Mercet (Bataglia & Viggiani, 1985). *Aspidiotiphagus citrinus* is known to attack a range of diaspidid scale species (Tao & Chang, 1963; Walker & Deitz, 1979), including other species of *Nuculaspis* (McClure, 2002), but this is the first report of it attacking *N. abietis*. Although this species regularly parasitizes more than 90% of * Fiorinia externa* Ferris and *Nuculaspis tsugae* (Marlatt) in Japan, it has been less effective in the eastern United States (McClure, 1986). In Kyoto, Japan, *A. citrinus* and its scale hosts both have two generations per year and the emergence of adult parasitoids is well synchronized with the peak abundance of vulnerable scale stages (McClure, 1986). However, in Connecticut, USA where *F. externa* completes only one generation a year (McClure, 1978) the peak emergence of bivoltine *A. citrinus* occurs almost eight weeks after the peak abundance of *F. externa* second instar nymphs. This results in highly variable parasitism rates from year to year and equilibrium scale densities much higher than observed in Japan. However, because *N. tsugae* completes two generations per year in Connecticut, its life cycle is more synchronous with that of *A. citrinus* and parasitism rates around 55% are sustained (McClure, 1986). Despite the fact that *N. abietis* has one generation in Kelardasht while *A. citrinus* completes two, the phenology of the scale population is such that adult parasitoids of both generations tend to encounter vulnerable stages of *N. abietis*, at least in the year of this study. Although it is conceivable that aberrant weather conditions could desynchronize the parasitoid and scale populations in some years, parasitoids that complete two or more generations in the period required for their host to complete one are considered to possess a favorable “generation time ratio” (Kindlmann & Dixon, 1999), a situation generally viewed as auspicious for biological control outcomes. For example, excellent biological control of the pseudococcid *Maconellicoccus hirsutus* Green has been obtained with *Anagrus kamali* Moursi and *Gyranusoida indica* Shafee, Alam & Agarwal (Hymenoptera: Encyrtidae) and this success is often attributed to the parasitoids’ ability to complete two generations in the time it takes their host to complete one (Michaud, 2003).

During the course of this study, it was apparent that *N. abietis* achieved much higher densities on young trees in the Kelardasht nursery than it did on established trees in the local forests. A number of factors likely contribute to this difference. Young trees in a nursery are frequently fertilized and grow rapidly, which could make them more nutritious hosts than older, established trees. Nurseries are managed environments that receive periodic applications of herbicides and insecticides that may kill or repel parasitoids. Similarly, Yanin (1975) concluded that *N. abietis* and other coccids were more likely to become pests of woody ornamentals in artificial than in natural settings. Insecticide applications targeting *N. abietis* in the Kelardasht tree nursery are normally timed with the appearance of crawlers (typically in early June) and unfortunately this coincides with the peak emergence of adult parasitoids.

The study region was revisited in August, 2010 to assess the state of scale infestation some ten years after the original study. Densities of *N. abietis* on established trees were noticably reduced in comparison with the study period and the infestation still appeared to be restricted to the Kelardasht region. The scale remained detectable on established trees of *Picea abies, P. pungens, P. orientalis, P. nitidiformis, Abies alba* and *A. nordmanniana*, but not at levels that would be considered economic. Scale management in the tree nursery has shifted from periodic broadcast applications of insecticides to spot-treatment of only severely affected saplings on an as-needed basis, an approach which has been successful in reducing the frequency and intensity of pesticide applications and favored the conservation of the parasitoid on untreated trees, improving the overall level of biological control. Thus, *A. citrinus* appears to make an important
contribution to the control of *N. abietis* in Kelardasht, even though it may be necessary to supplement its impact with selective spot-treatments of insecticides in tree nurseries where young, vigorously growing trees under intensive fertilization and irrigation schedules may be particularly vulnerable.

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