A hydroid (*Cordylophora caspia*)

MarLIN – Marine Life Information Network
Biology and Sensitivity Key Information Review

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**Summary**

**Description**

A tall erect colony growing up to 10 cm high, branching occasionally from alternate sides and light horn to light brown in colour. Each branch is ringed at the base and has a terminal polyp. The polyps are white to pale pink and have 12 - 16 long, colourless, extensile tentacles arranged irregularly on the surface of the polyp. The mouth is born on a conical but truncated proboscis. Each branch bears one to three pear-shaped reproductive bodies on short stalks. It produces a planula larvae but no free-living medusoid stage.

**Recorded distribution in Britain and Ireland**

The species has a sporadic distribution associated with areas of low salinity within estuaries and brackish lagoons.

**Global distribution**

Found in estuarine, lagoonal and coastal lake habitats in boreal to subtropical waters.
Habitat

This hydroid colonizes brackish waters of 2 - 12 psu but, where salinity may rise, occasionally up to 35 psu. It is found in shallow depths, often in shade, on various hard substrata, submerged vegetation, and the shells of crabs and snails.

Depth range
Low shore to ca 2m.

Identifying features

- Tall, erect colony ensheathed in perisarc.
- Polyps terminal with 12-16 tentacles.
- Polyps naked (athecate).
- Reproductive polyps (gonopores) pear-shaped.

Additional information

No text entered

Listed by

Further information sources

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Biology review

Taxonomy

- **Phylum**: Cnidaria (Sea anemones, corals, sea firs & jellyfish)
- **Class**: Hydrozoa (White weeds, sea firs, sea beard and siphonophores; hydroids)
- **Order**: Anthoathecata
- **Family**: Cordylophoridae
- **Genus**: Cordylophora
- **Authority**: (Pallas, 1771)
- **Recent Synonyms**: Cordylophora lacustris (Pallas, 1771)

Biology

- **Typical abundance**: Moderate density
- **Male size range**: Not provided
- **Male size at maturity**: Not provided
- **Female size range**: Small-medium (3-10cm)
- **Female size at maturity**: Not provided
- **Growth form**: Turf
- **Growth rate**: ca 0.05-0.1mm/hr
- **Body flexibility**: High (greater than 45 degrees)
- **Mobility**: Not provided
- **Characteristic feeding method**: Passive suspension feeder, Predator, See additional information
- **Diet/food source**: Typically feeds on Small zooplankton, small crustacea, oligochaetes, insect larvae and probably detritus.
- **Sociability**: Not provided
- **Environmental position**: Epibenthic
- **Dependency**: Independent.
- **Supports**: None
- **Is the species harmful?**: No

Biology information

**Growth form**

Growth form is highly variable in *Cordylophora caspia*. The colony of consists of a mass of stolons (hydrorhizae) growing across the surface of the substratum. Growth is apical, with side stolons arising at right angles. Upright hydrocauli bear apical polyps (hydranths), and side branches at 45° (Fulton, 1961). The degree of branching, length and spacing of hydrocauli, cell size, size and shape of polyps and the number and length of tentacles vary with environmental conditions (Fulton, 1962; Kinne, 1970, 1971; Arndt, 1989). For example, colonies have short, polyps that grow directly from the hydrorhiza at 0.5psu; more elongate polyps with longer tentacles and multiply branched uprights at 15psu, but smaller polyps and less branched uprights at 30psu than at 15psu (see Kinne...
Gaulin et al. (1986) noted that predation by the nudibranch Tenellia fuscata resulted in denser colonies.

**Growth rates**

Growth rates are variable depending on environmental or laboratory conditions. Growth in number of polyps is exponential, the colonies doubling in polyp number every 2-4 days, although growth rate can vary as much as two-fold even under standard conditions (Fulton, 1961, 1962). In addition, although old colonies could reach as much as 2000 polyps in size growth rates decreased with age (Fulton, 1962). Fulton (1961) reported that uprights grew at 0.05mm/hr while stolons extension rates vary from 0.1mm/hr (Fulton, 1961) to 2-3mm$^3$/day (Chester et al., 2000). Fulton (1962) reported that growth rates varied with temperature, salinity, ionic composition, oxygen tension and feeding rate (see sensitivity).

**Seasonal changes**

*Cordylophora caspia* dies back in late autumn and overwinters as dormant stolons and resting stages (menonts) inside the remnants of the uprights (see Roos, 1979 for figure; Arndt, 1989; Jormalainen et al., 1994). Arndt (1989) reported that colonies died back in autumn when the temperature fell to about 10°C only to germinate in spring when the temperature exceeded 5°C. Roos (1979) reported that colonies died back in October and new polyps budded again in early spring in the Netherlands. In the Baltic Sea growth was maximal in spring, uprights reaching maximal height at the peak of sexual reproduction in July, with a decline after sexual reproduction, and subsequent growth in August (Jormalainen et al., 1994). However, in one year, Jormalainen et al (1994) noted that the colonies regressed to the dormant condition after sexual reproduction then started growing again by mid August.

**Feeding**

Hydroids are passive carnivores that capture prey that swim into, or are brought into contact with their tentacles by currents. Prey are then killed or stunned by the nematocysts born on the tentacles and swallowed. Diet varies but is likely to include small zooplankton (e.g. nauplii, copepods), small crustacea, chironomid larvae, detritus and oligochaetes, but may include a wide variety of other organisms such as the larvae or small adults of numerous groups (see Gili & Hughes, 1995).

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**Habitat preferences**

- **Physiographic preferences**
  - Sea loch / Sea lough, Ria / Voe, Estuary, Isolated saline water (Lagoon), Enclosed coast / Embayment

- **Biological zone preferences**
  - Lower eulittoral, Lower infralittoral, Sublittoral fringe, Upper infralittoral

- **Substratum / habitat preferences**
  - Artificial (man-made), Bedrock, Caves, Cobbles, Large to very large boulders, Overhangs, Pebbles, Small boulders, Under boulders

- **Tidal strength preferences**
  - Moderately Strong 1 to 3 knots (0.5-1.5 m/sec.), Strong 3 to 6 knots (1.5-3 m/sec.), Very Weak (negligible), Weak < 1 knot (<0.5 m/sec.)

- **Wave exposure preferences**
  - Extremely sheltered, Very sheltered

- **Salinity preferences**
  - Low (<18 psu), Reduced (18-30 psu)

- **Depth range**
  - Low shore to ca 2m
Other preferences None entered

Migration Pattern

Habitat Information

Habitat preferences
The distribution of *Cordylophora caspia* is determined by availability of suitable hard substratum, food availability, range and variability of temperature and salinity. *Cordylophora caspia* can survive between -10 °C (as resistant dormant stages, menonts) and 35 °C. Colonies tolerate 5 to 35 °C, and reproduce between 10 to 28 °C. It can also survive 0 to 35psu as resistant stages, grow between 0.2 to 30 psu, reproduce between 0.2 to 20psu and possesses the ability to ionic regulate (Kinne, 1971; reviewed by Arndt, 1986, 1989). In nature, well developed colonies are usually found in water of 2 -12psu where tidal influence is considerable or between 2 -6psu where conditions are constant (Arndt, 1989). It may also occur at full salinities, and fast flowing, well oxygenated freshwater containing Ca, Mg, Na Cl and K ions (Fulton, 1962; Arndt, 1989). Arndt (1986, 1989) suggested that respiration, growth and reproduction were optimal between 4-7psu and that food intake was high in comparison to other hydroids so that growth and reproduction rates required for the survival of the species could only occur in eutrophic or hypertrophic waters where food is plentiful. Its marine distribution is probably limited by food availability, competition from *Clava* spp. or *Laomedea* spp. and predation e.g. from the nudibranch *Tenellia adspersa* (as *Embletonia pallida*) (Arndt, 1989). *Cordylophora caspia* prefers conditions of low light (Allman, 1871-1872, Arndt, 1989), although light intensity did not affect growth (Fulton, 1963), which probably reflects the settlement preferences of the planula larvae.

Substrata
Most hydroids do not show a high specificity of substrata. *Cordylophora caspia* has been recorded from a wide variety of hard substrata including rocks, shells and artificial substrata (pilings, harbour installations, bridge supports), floating debris and occasionally from the leaves of reeds (*Phragmites*) or stalks of water lilies (MBA, 1957; Roos, 1979; Morri & Boero, 1986; Arndt, 1986, 1989; JNCC, 1999; Foster-Smith, 2000).

Non-native status
*Cordylophora caspia* was thought to have been introduced to British waters on foreign timber (Allman, 1871-1872). *Cordylophora caspia* was introduced into the Baltic Sea in ca 1803 and was reported as an alien species in the Baltic Sea and the Chesapeake Bay region, USA (Folino, 1999 (summary only); Olenin et al., 2000). Folino (1999, summary only) suggested that the distribution of *Cordylophora* spp. was expanding globally due to increased boat travel and ballast discharge.

Life history

Adult characteristics

| Reproductive type | Vegetative |
| Reproductive frequency | Annual episodic |
| Fecundity (number of eggs) | See additional information |
| Generation time | <1 year |
| Age at maturity | Less than 1 month |
Season
Spring - Autumn

Life span
See additional information

Larval characteristics

Larval/propagule type
- 

Larval/juvenile development
Direct development

Duration of larval stage
< 1 day

Larval dispersal potential
< 10 m

Larval settlement period
See additional information

Life history information

Most hydroids (including Cordylophora caspia) are dioecious. The reproductive organs are carried in gonophores. Sperm are released into the sea and eggs are fertilized within the female gonophores where the embryos develop into planulae. Sperm swim towards female gonophores, however, sperm probably have a limited lifespan and hence a limited range for fertilization of only a few metres. Hence the growth of hydroids in clumps may enhance fertilization rates, albeit at the cost of intraspecific competition. Temperature is critical for stimulating or preventing reproduction in hydroids (see distribution; Gili & Hughes, 1995).

Sexual reproduction
Early seasonal growth from winter dormancy in early spring is rapidly followed by formation of gonophores and sexual reproduction in midsummer followed by active growth in late summer. However, sexual reproductive effort may retard growth (see general biology). Jormalainen et al. (1994) reported that reproduction began in early June, peaked in July (80% uprights with gonophores) and rapidly reduced by August (30% uprights with gonophores). Similar reproductive periods have been reported by other authors (Allman, 1871-1872; MBA, 1957; Roos, 1979; Foster-Smith, 2000). Roos (1979) and Jormalainen et al. (1994) reported that the sex ratio was biased in favour of females.

Each upright branch may bear between 1-3 gonophores each with between 10 - 6 eggs, the number decreasing in autumn (Hincks, 1868; Jormalainen et al., 1994). Therefore, fecundity is dependant on the number of branches and hence the number of gonophores, and in large colonies of 70-2000 polyps (Fulton, 1962), may be high. The larvae are released as planulae and no medusoid stage occurs. However, in some cases the larvae may develop directly into juvenile polyps in the gonophore before release (Bouillon, 1963).

Asexual reproduction
Hydroids may reproduce asexually by budding to from another colony. A common form of asexual reproduction in hydroids is the formation of vertical stolons, which then adhere to adjacent substratum, detach and form another colony (Gili & Hughes, 1995). Hydroids exhibit remarkable powers of regeneration and Cordylophora caspia can be cloned in culture from detached uprights or excised tissue (Moore, 1952; Fulton, 1961, 1962). Asexual reproduction by fission or mechanical fragmentation of the colony may be an important factor in dispersal (Gili & Hughes, 1995).

Longevity
While uprights have a short, finite lifespan from about early spring to autumn, no information concerning the lifespan of the dormant stages (menonts) was found. Unless destroyed by predators or physical damage, the colony may have a long lifespan (perhaps very long (Gili &
Hughes, 1995). The ability to reproduce asexually and regenerate from damaged sections means that although any individual colony may have a finite lifespan the genetic individual (genet) may be considerably longer lived (Gili & Hughes, 1995).

Dispersal
Rapid growth, budding and the formation of stolons allows hydroids to colonize space rapidly. Hydroids are often the first organisms to colonize available space in settlement experiments (Gili & Hughes, 1995). Planula larvae swim or crawl for short periods (e.g. <24hrs) so that dispersal away from the parent colony is probably very limited (Gili & Hughes, 1995). Fragmentation may also provide another route for short distance dispersal. However, it has been suggested that rafting on floating debris (or hitch hiking on ships hulls or in ship ballast water) as dormant stages or reproductive adults, together with their potentially long lifespan, may have allowed hydroids to disperse over a wide area in the long-term and explain the near cosmopolitan distributions of many hydroid species, including Cordylophora caspia (Gili & Hughes, 1995; Folino, 1999).
Sensitivity review

This MarLIN sensitivity assessment has been superseded by the MarESA approach to sensitivity assessment. MarLIN assessments used an approach that has now been modified to reflect the most recent conservation imperatives and terminology and are due to be updated by 2016/17.

Physical Pressures

<table>
<thead>
<tr>
<th>Substratum Loss</th>
<th>Intolerance</th>
<th>Recoverability</th>
<th>Sensitivity</th>
<th>Confidence</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>High</td>
<td>High</td>
<td>Moderate</td>
<td>Moderate</td>
</tr>
</tbody>
</table>

Removal of the substratum would result in the loss of colonies, their hydrorhizae and any resting stages. Therefore, an intolerance of high has been recorded. Recovery will depend on whether nearby colonies or colonies within the same water body have survived. If so, recoverability will be rapid, although the original population abundance may take several growing seasons to achieve, and an intolerance of high has been recorded. If the population has been completely destroyed recoverability will be low or not at all (see additional information below).

Smothering

<table>
<thead>
<tr>
<th>Smothering</th>
<th>Intolerance</th>
<th>Recoverability</th>
<th>Sensitivity</th>
<th>Confidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>Immediate</td>
<td>Not sensitive</td>
<td>Low</td>
<td></td>
</tr>
</tbody>
</table>

Hydroids usually colonize overhanging, vertical or steeply sloping surfaces presumably to avoid the possibility of siltation and smothering and competition from macroalgae. Smothering by 5cm of sediment (see benchmark) is likely to cover a large proportion of the colony, preventing feeding and hence reducing growth and reproduction. In addition, local hypoxic conditions are also likely to inhibit growth (Fulton, 1961, 1963). However, the colony is likely to become dormant, or otherwise survive for a period of at least a month, and recover rapidly once the sediment is removed. Therefore, an intolerance of low has been recorded to represent the affects of smothering on growth. Recovery is likely to be immediate (see additional information below).

Increase in suspended sediment

<table>
<thead>
<tr>
<th>Increase in suspended sediment</th>
<th>Intolerance</th>
<th>Recoverability</th>
<th>Sensitivity</th>
<th>Confidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tolerant</td>
<td>Not relevant</td>
<td>Not sensitive</td>
<td>Low</td>
<td></td>
</tr>
</tbody>
</table>

*Cordylophora caspia* is found in estuarine and sheltered lagoonal habitats, which are characterized by high suspended sediment loads. *Cordylophora caspia* was also reported in saltmarsh pools (JNCC, 1999) and saltmarshes are a depositional environment characterized by siltation. Therefore, *Cordylophora caspia* is probably tolerant of increases in suspended sediment loads at the benchmark level.

Decrease in suspended sediment

<table>
<thead>
<tr>
<th>Decrease in suspended sediment</th>
<th>Intolerance</th>
<th>Recoverability</th>
<th>Sensitivity</th>
<th>Confidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>Immediate</td>
<td>Not sensitive</td>
<td>Low</td>
<td></td>
</tr>
</tbody>
</table>

A reduction in suspended sediment is unlikely to directly affect *Cordylophora caspia*. A decrease in suspended sediment may also reduce the availability of organic particulates and hence reduce food availability. Arndt (1986, 1989) suggested that *Cordylophora caspia* had a high food requirement for growth and reproduction. It is therefore, likely to be intolerant of any reduction in food availability and an intolerance of low has been recorded. Recovery is likely to be immediate.

Dessication

<table>
<thead>
<tr>
<th>Dessication</th>
<th>Intolerance</th>
<th>Recoverability</th>
<th>Sensitivity</th>
<th>Confidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>High</td>
<td>Moderate</td>
<td>Low</td>
<td></td>
</tr>
</tbody>
</table>

Intertidal populations of *Cordylophora caspia* are restricted to damp habitats such as underboulders and overhangs. The branched growth form of this species is likely to retain water on emersion (see image). However, an increase in desiccation at the benchmark level is likely to result in drying and death of the uprights. Increased dessication may result in the
formation of resistant, dormant stages, however, no information on their desiccation
tolerance was found. Therefore, an intolerance of high has been recorded.
If hydrorhizae or dormant stages survive recovery is likely to be very rapid and colonies may
appear rapidly once conditions return to their prior state. If, resting stages are destroyed then
recovery will depend on recruitment from nearby subtidal colonies, and is likely to be rapid,
although the original population abundance may take several growing seasons to achieve (see
additional information below).

Increase in emergence regime

<table>
<thead>
<tr>
<th></th>
<th>Intermediate</th>
<th>Very high</th>
<th>Low</th>
<th>Very Low</th>
</tr>
</thead>
</table>

An increase in emergence is likely to adversely affect colonies. While *Cordylophora caspia*
would probably survive the extremes of temperature resulting form increased emergence (see
below), colonies are likely to succumb to increased desiccation (see above) if increased
emergence exposes them to mid shore (or higher) conditions. Therefore, the upper shore
proportion of the population is likely to be lost and an intolerance of intermediate has been
recorded. Recoverability is likely to be very high (see additional information below).

Decrease in emergence regime

<table>
<thead>
<tr>
<th></th>
<th>Tolerant*</th>
<th>Not relevant</th>
<th>Not sensitive*</th>
<th>Very high</th>
</tr>
</thead>
</table>

A decrease in emergence is likely to increase the availability of suitable habitats and may allow
the population to extend its range. Therefore, tolerant* has been recorded.

Increase in water flow rate

<table>
<thead>
<tr>
<th></th>
<th>Low</th>
<th>Very high</th>
<th>Very Low</th>
<th>Very low</th>
</tr>
</thead>
</table>

Water movement is essential for hydroids to supply adequate food, remove metabolic waste
products, prevent accumulation of sediment and disperse larvae or medusae. Hydroids are
expected to be abundant where water movement is sufficient to supply adequate food but not
cause damage (Hiscock, 1983; Gili & Hughes, 1995). Flexibility of the otherwise rigid perisarc
of hydroids is provided by annulations at the base of branches in *Cordylophora caspia*. In
addition, in athecates, the neck of the polyp is flexible enough to allow the polyp adopt an
efficient 'lee-side' feeding posture in water flow. However, most hydroids have a narrow range
of water flow rates for effective feeding. For example in the athecate *Tubularia indivisa*, food
capture rate increased up to 20cm/s, but decreased as water flow rates increased (Hiscock,
1983). In *Cordylophora inkermania* food capture rates were higher in fluctuating flows than in
unidirectional flows (Gili & Hughes, 1995), presumably because more polyps were brought
into play in fluctuating flow, than in unidirectional flow where upstream branches 'shaded'
down stream branches. Loomis (in Fulton, 1961) noted that *Cordylophora caspia* did not grow
in still water cultures presumably because of the build up of CO$_2$ from respiration.
*Cordylophora caspia* was reported to dominate steep rock surfaces in strong tidal flows in the
Tamar (JNCC, 1999). Although *Cordylophora caspia* tolerates strong water flow, a further
increase in water flow to very strong is likely to reduce feeding efficiency and hence growth
and reproduction and may even remove or damage colonies. Damaged colonies may survive as
resting stages until water flow rates return to prior condition. Therefore, intolerance has been
assessed as low and recoverability as very high (see additional information below).

Decrease in water flow rate

<table>
<thead>
<tr>
<th></th>
<th>Low</th>
<th>Immediate</th>
<th>Not sensitive</th>
<th>Very high</th>
</tr>
</thead>
</table>

Water movement is essential for hydroids to supply adequate food, remove metabolic waste
products, prevent accumulation of sediment and disperse larvae or medusae (see above).
*Cordylophora caspia* has been recorded from areas of negligible or weak water flow, e.g. saline
lagoons. A further decrease in water flow is unlikely, although Loomis (in Fulton, 1961) noted
that colonies did not grow in still water cultures presumably because of the build up of CO$_2$
from respiration. Therefore, an intolerance of low has been recorded. Recovery is likely to be
very high (see additional information below).
**Increase in temperature**

*Cordylophora caspia* can survive as resistant dormant stages (menonts) at -10 °C and at 35 °C. Colonies tolerate 5 to 35 °C, and reproduce between 10 to 28 °C. Arndt (1989) reported that colonies died back in autumn when the temperature fell to about 10 °C only to germinate in spring when the temperature exceeded 5 °C. Arndt (1989) concluded that *Cordylophora caspia* was thermophilic but that low temperature had an important influence on growth and reproduction. In addition, the distribution of *Cordylophora caspia* extends into subtropical habitats (Arndt, 1986, 1989). Therefore, this species is unlikely to be adversely affected by chronic or acute temperature change at the benchmark level in British waters.

**Decrease in temperature**

*Cordylophora caspia* can survive as resistant dormant stages (menonts) at -10 °C and at 35 °C. Colonies tolerate 5 to 35 °C, and reproduce between 10 to 28 °C. Arndt (1989) reported that colonies died back in autumn when the temperature fell to about 10 °C only to germinate in spring when the temperature exceeded 5 °C. Arndt (1989) concluded that *Cordylophora caspia* was thermophilic but that low temperature had an important influence on growth and reproduction. Therefore, while low temperatures may trigger premature die back or regression, colonies are likely to survive changes in temperature at the benchmark level and an intolerance of low has been recorded.

**Increase in turbidity**

*Cordylophora caspia* is unlikely to be directly influenced by light intensity (Fulton, 1962). *Cordylophora caspia* and other hydroids tend to shun well lit conditions, planulae becoming negatively phototactic prior to settlement, presumably to avoid competition with macroalgae (Gili & Hughes, 1995). Therefore, a decrease in light penetration may decrease competition for space with macroalgae. However, decrease light penetration may also decrease phytoplankton and hence zooplankton productivity and potentially decrease food availability, and an intolerance of low has been recorded.

**Decrease in turbidity**

Increased light penetration may increase phytoplankton and hence zooplankton productivity and potentially increase food availability. Therefore, tolerant* has been recorded.

**Increase in wave exposure**

The oscillatory water flow caused by wave action is potentially more damaging to delicate marine organisms than unidirectional flow. Although, the annuli at the base of branches gives the branched colony of *Cordylophora caspia* some flexibility it has only been recorded from very or extremely wave sheltered habitats (JNCC, 1999). Therefore, it is likely than an increase in wave exposure at the benchmark level (e.g. from 'very sheltered' to 'moderately exposed') is likely to result in loss of the colonies. Populations occupying small rocks, cobbles or pebbles are likely to be more intolerant, and the resultant movement of the substratum and sediment scour may also remove attached hydorhizae and even resting stages. Therefore, an intolerance of high has been recorded.

Recovery will depend in part on recruitment from other areas. However, any resting stages and fragments of colonies remaining may contribute to the recovery, although the original population abundance may take several growing seasons to achieve. Therefore, a recoverability of high has been suggested.

**Decrease in wave exposure**

*Cordylophora caspia* has only been recorded from very or extremely wave sheltered habitats.
A decrease in wave exposure may allow the species to colonize additional habitats and increase its extent. Therefore, tolerant* has been recorded.

### Noise

| Tolerant | Not relevant | Not sensitive | High |

Hydroids are unlikely to be sensitive to noise or vibration at the benchmark level.

### Visual Presence

| Tolerant | Not relevant | Not sensitive | High |

Hydroid polyps may retract when shaded by potential predators, however hydroids are unlikely to be affected by visual presence.

### Abrasion & physical disturbance

| Intermediate | Very high | Low | Low |

Abrasion by an anchor or fishing gear is likely to remove relatively delicate upright parts of the colony. However, the surface covering of hydrorhizae may remain largely intact, from which new uprights are likely to grow. In addition, the resultant fragments of colonies may be able to develop into new colonies (see displacement). Populations on small hard substrata (e.g. cobbles, pebbles or stones) may be removed by fishing gear, constituting substratum loss (see above). Overall, a proportion of the colonies is likely to be destroyed and an intolerance of intermediate has been recorded. However, recovery from surviving hydrorhizae and occasional fragments is likely to be rapid (see additional information below).

### Displacement

| Intermediate | Very high | Low | Low |

*Cordylophora caspia* colonies have been cultured by securing cut uprights to slides to which they subsequently attach. Fragmentation is thought to be a possible mode of asexual reproduction in hydroids (Gili & Hughes, 1995). Therefore, it is possible that a proportion of displaced colonies (or fragments thereof) may attach to new substrata and an intolerance of intermediate has been recorded. Recovery is likely to be rapid (see additional information below).

### Chemical Pressures

| Intolerance | Recoverability | Sensitivity | Confidence |

#### Synthetic compound contamination

The species richness of hydroid communities decreases with increasing pollution but hydroid species adapted to a wide variation in environmental factors and with cosmopolitan distributions tend to be more tolerant of polluted waters (Boero, 1984; Gili & Hughes, 1995). Stebbing (1981) reported that Cu, Cd, and tributyl tin fluoride affected growth regulators in *Laomedea* (as *Campanularia* flexuosa) resulting in increased growth. Bryan & Gibbs (1991) reported that virtually no hydroids were present on hard bottom communities in TBT contaminated sites and suggested that some hydroids were sensitive to TBT levels between 100 and 500 ng/l.

However, Calder (1976) suggested that hydroids found in the low salinity areas of south Carolina, such as *Cordylophora caspia*, were also present in relatively polluted waters, such as Charleston Harbour. *Cordylophora caspia* was also a dominant species on settlement plates placed on a floating shipyard dock in Warnock river (Sandrock et al., 1991). Floating docks are likely to result in local contamination with heavy metals and antifouling agents from ship paints, as well as oils and other chemicals used in ship maintenance. As a member of fouling communities, *Cordylophora caspia* is probably less intolerant of antifouling measures than other hydroids. Therefore, an intolerance of low has been suggested albeit at very low confidence.

#### Heavy metal contamination

Low | Immediate | Not sensitive | Moderate |
Stebbing (1981) reported that Cu, Cd, and tributyl tin fluoride affected growth regulators in *Laomedea* (as *Campanularia*) *flexuosa* resulting in increased growth. Ringelband (2001), however, reported that 1.74-7.96 mg/l vanadium inhibited growth of *Cordylophora caspia* at low salinities. Various heavy metals have also been shown to have sublethal effects on growth in the few hydroids studied experimentally (Bryan, 1984). Therefore, an intolerance of low has been recorded.

**Hydrocarbon contamination**  
Low  Immediate  Not sensitive  Very low  

Little information of the effects of hydrocarbons on hydroids was found. Hydroid species adapted to a wide variation in environmental factors and with cosmopolitan distributions tend to be more tolerant of polluted waters (Boero, 1984; Gili & Hughes, 1995). Calder (1976) suggested that hydroids found in the low salinity areas of south Carolina, such as *Cordylophora caspia*, were also present in relatively polluted waters, such as Charleston Harbour. *Cordylophora caspia* was also a dominant species on settlement plates placed on a floating shipyard dock in Warnock river (Sandrock *et al.*, 1991). Floating docks are likely to result in local contamination with heavy metals and antifouling agents from ship paints, as well as oils and other chemicals used in ship maintenance. However, the above evidence is primarily anecdotal.

The water soluble fractions of Monterey crude oil and drilling muds were reported to cause polyp shedding and other sublethal effects in the athecate *Tubularia crocea* in laboratory tests (Michel & Case, 1984; Michel *et al.*, 1986; Holt *et al.*, 1995). The athecate *Cordylophora caspia* may show similar sublethal effects assuming similar physiology. Therefore, an intolerance of low has been recorded albeit with a very low confidence.

**Radionuclide contamination**  
Not relevant  Not relevant  

**Changes in nutrient levels**  
Tolerant*  Not relevant  Not sensitive*  Moderate  

*Cordylophora caspia* became one of the dominant species to colonize settlement plates placed beneath a floating dock in the Warnock river (Sandock *et al.*, 1991). This station was characterized by low salinities, and higher organic and mineral nutrient loads (ca 20-100 µmol NO$_3$/l) than their other experimental station. Arndt (1986, 1989) suggested that food intake in *Cordylophora caspia* was high in comparison to other hydroids so that growth and reproduction rates required for the survival of the species could only occur in eutrophic or hypertrophic waters where food is plentiful. Therefore, *Cordylophora caspia* is likely to tolerate relatively high nutrient levels, and may benefit from moderate increases in nutrients at the benchmark level. Hence, tolerant* has been recorded.

**Increase in salinity**  
Intermediate  Very high  Low  Moderate  

*Cordylophora caspia* can survive 0 to 35psu as resistant stages, grow between 0.2-30 psu, reproduce between 0.2-20psu and possesses the ability to ionic regulate (Kinne, 1971; reviewed by Arndt, 1986, 1989). In nature, well developed colonies are usually found in water of 2-12psu where tidal influence is considerable or between 2-6psu where conditions are constant (Arndt, 1989) but it may also occur at full salinities. Kinne (1971) noted that high salinities (24 or 30psu) occasionally resulting in developmental abnormalities in older colonies in the laboratory. Arndt (1989) suggested that its marine distribution was probably limited by food availability, competition from *Clava* spp. or *Laomedea* spp. and predation e.g. from the nudibranch *Tenellia adspersa* (as *Embletonia pallida*).

Therefore a short term increase in salinity (at the benchmark level) is likely to affect growth
and reproduction but otherwise not adversely affect colonies. However, a change in salinity from reduced to variable in the long term (see benchmark) may result in loss of a proportion of the population, and an intolerance of intermediate has been recorded. Survival of resting stages is likely to result in rapid recovery (see additional information below).

**Decrease in salinity**

*Cordylophora caspia* can survive 0 to 35psu as resistant stages, grow between 0.2-30 psu, reproduce between 0.2-20psu and possesses the ability to ionic regulate (Kinne, 1971; reviewed by Arndt, 1986, 1989). In nature, well developed colonies are usually found in water of 2 -12psu where tidal influence is considerable or between 2 -6psu where conditions are constant (Arndt, 1989). It may also occur at full salinities, and fast flowing, well oxygenated freshwater containing Ca, Mg, Na, Cl and K ions (Fulton, 1962; Arndt, 1989). It has been reported from estuaries that receive significant seasonal freshwater input, and tolerates variable salinities (Arndt, 1986; 1989). Therefore, it is probably relatively tolerant of a change in salinity at the benchmark level. A reduction from full to reduced salinity may be beneficial and allow *Cordylophora caspia* to colonize new habitats. Therefore, not sensitive* has been recorded.

**Changes in oxygenation**

Fulton (1962) found that some polyps of *Cordylophora caspia* fell off or were reabsorbed after 7 days in the complete absence of oxygen, however, the remaining polyps began feeding shortly after the re-introduction of oxygen. Fulton (1962) concluded that *Cordylophora caspia* had a low oxygen requirement for growth and was able to grow at oxygen levels of >2mg/l (ca 1.4ml/l). Arndt (1986) reported that an increase in temperature from 10 to 20 °C resulted in a marked increase in metabolic rate and hence oxygen consumption. Similarly, metabolic rates increased at supra- or subnormal salinities (Arndt, 1986) and *Cordylophora caspia* may be more intolerant of low oxygen concentrations at high temperatures and extreme salinities. However, the isolated saline waters, the upper reaches of estuaries and saltmarsh pools, in which *Cordylophora caspia* occurs, are likely to experience high summer temperatures and hence low oxygen levels. Therefore, *Cordylophora caspia* is likely to survive exposure to low oxygen concentrations at the benchmark level, although growth is likely to be reduced, and an intolerance of low has been recorded.

**Biological Pressures**

<table>
<thead>
<tr>
<th>Biological Pressure</th>
<th>Intolerance</th>
<th>Recoverability</th>
<th>Sensitivity</th>
<th>Confidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Introduction of microbial pathogens/parasites</td>
<td>Not relevant</td>
<td></td>
<td></td>
<td>Not relevant</td>
</tr>
<tr>
<td>No information found.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Introduction of non-native species</td>
<td>Not relevant</td>
<td></td>
<td></td>
<td>Not relevant</td>
</tr>
<tr>
<td><em>Cordylophora caspia</em> is a non-native species (Allman, 1871-1872). But has not been reported to compete with other species, including other non-native species, in British or Irish waters.</td>
<td></td>
<td></td>
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<tr>
<td>Extraction of this species</td>
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<td>Not relevant</td>
<td>Not relevant</td>
<td>Not relevant</td>
</tr>
<tr>
<td>Hydroids are not known to be subject to specific extraction.</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Extraction of other species</td>
<td>Not relevant</td>
<td>Not relevant</td>
<td>Not relevant</td>
<td>Not relevant</td>
</tr>
<tr>
<td><em>Cordylophora caspia</em> is not known to be associated with species or habitats subject to extraction.</td>
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</tbody>
</table>
Additional information

intolerance assessment

*Cordylophora caspia* and other hydroids have the ability to produce dormant resting stages (menonts) that are far more resistant to environmental change than the colony itself. Therefore, although colonies may be removed or destroyed, the resting stages may survive in remnants of the hydrorhizae attached to the substratum. For the sake of assessment, the intolerance of the branched colonies themselves (the clearly visible component) has been recorded. The resting stages provide a mechanism for rapid recovery.

Recoverability

Hydroids are often initial colonizing organisms in settlement experiments and fouling communities (Jensen *et al.*, 1994; Hatcher, 1998). In settlement experiments in the Warnow estuary, *Cordylophora caspia* was found to colonize artificial substrata within ca one month of deployment, its abundance increasing from June to the end of September with a peak in July (Sandrock *et al.*, 1991). Long term panels at the low salinity station became dominated by *Cordylophora caspia*, %*Balanus improvisus*% and %*Nais elinguis*%. Similarly, Roos (1979) reported that *Cordylophora caspia* recruited to and grew luxuriantly on water lily stalks in summer after early reproduction of nearby colonies in early spring. Therefore, it is likely that *Cordylophora caspia* will recruit to available space rapidly in its growing season, in the vicinity of other populations. Once colonized the hydroids ability to grow rapidly and reproduce asexually is likely to allow it to occupy space and sexually reproduce quickly, possibly recruiting to additional space before dying back in winter. Therefore, where colonies or dormant resting stages are present in the habitat, or within isolated habitats (e.g. lagoons), recovery is likely to be rapid and occur within less than a year.

Long distance dispersal is probably limited in hydroids, including *Cordylophora caspia*. Long distance dispersal is probably dependant on passive dispersal by currents on floating debris or shipping. Although, *Cordylophora caspia* has probably been introduced to the coasts of several countries (e.g. Chesapeake Bay USA and the Baltic Sea), passive transportation is a sporadic and un-predictable event. Therefore, if the whole population is destroyed, and resting stages removed, recolonization rates will depend on distance from nearby colonies, and may take many years. In isolated habitats such as lagoons and coastal lakes, recruitment from other areas may take a many decades or not occur at all.
Importance review

Policy/legislation
- no data -

Status

<table>
<thead>
<tr>
<th>National (GB) importance</th>
<th>Global red list (IUCN) category</th>
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Non-native

<table>
<thead>
<tr>
<th>Native</th>
<th>Non-native</th>
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<td></td>
<td>East Europe, Soviet Middle Asia</td>
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</table>

Date Arrived 1868

Importance information

Hydroids are preyed on by chitons, gastropods (especially nudibranchs), polychaetes and pycnogonids (Gili & Hughes, 1995). For example, the pycnogonid Anoplodactylus sp. was reported to feed on Cordylophora caspia in Australasia (Staples & Watson, 1987) and may do so in the British Isles.

The life cycle of nudibranchs is often closely linked with the seasonal availability of their preferred prey. Arndt (1989) noted that the marine distribution of Cordylophora caspia was limited in part by predation by the nudibranch Tenellia adspersa (as Embletonia pallida). Gaulin et al. (1986) noted that a healthy colony would survive limited nudibranch predation by Tenellia fuscata but be removed by large numbers of predators. For example, Chester et al. (2000) suggested that Tenellia adspersa showed rapid population growth, removing the hydroid and preventing its reestablishment.

Nudibranch predation is probably an important factor limiting the presence of hydroids in early community succession, removing the hydroids and allowing colonization of other species.
Bibliography


Atmosphériques.


Datasets


