Polyp dropout in a solitary cold-water coral

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Abstract Scleractinian corals feature both sessile and mobile stages and diverse modes of development. In some cases, development can be reversed. Examples include polyp detachment in response to environmental stress (bail-out or polyp expulsion) and reverse metamorphosis, where juveniles detach from the primary skeleton and revert to the mobile stage. Here, we provide aquaria and field evidence of a new form of reverse development: polyp dropout in the solitary cold-water coral Caryophyllia huinayensis. It features tissue retraction and detachment of an entire adult polyp from the skeleton in the putative absence of a stressor. The dropout polyp remains viable and continues to live for many weeks, albeit in a rather collapsed state lacking a well-developed hydroskeleton. We carried out a long-term (37 months) rearing experiment under constant aquaria conditions and found polyp dropout in four out of 83 individuals. Detachment was accompanied by the extrusion of mesenterial filaments through perforations in the body wall. We believe this resulted in the loss of the hydroskeleton, which prevented the dropouts to subsequently resettle or form a new skeleton. As opposed to other known forms of reverse development, the new form is not accompanied by reversible metamorphosis, abandonment of the colonial way of life, nor is it a survival or asexual reproduction strategy. We found field indications of polyp dropout in Patagonian field populations of C. huinayensis, where 1.4 ± 0.8% (mean ± SD, N = 9322) of the polyps of the natural population showed partial detachment indicative of imminent dropout in the putative absence of external impact. Polyp dropout is the first record of polyp detachment in a solitary CWC with possible repercussions for adult coral mobility, evolution and Stanley’s (2003) ‘naked coral’ hypothesis.

Keywords Cold-water scleractinia · Caryophyllia huinayensis · Polyp detachment · Polyp dropout · Chilean Fjord Region · Asexual dispersal

Introduction

Corals are sessile animals with diverse modes of development, generally involving a mobile stage of dispersal, the planula larva, which later settles, undergoes metamorphosis and develops into the primary polyp (e.g. Richmond and Hunter 1990). Under environmental stress, development may become reverse. For example, a settled, calcified polyp may build back its radially compartmented body plan, reverse metamorphosis to become a mobile secondary larva, disperse with the currents and reattach elsewhere (Richmond 1985). Reverse development may also affect the colony: through ontogenetic reversal, genetic programs specific to earlier stages are reactivated, leading to back-
transformation to the previous morph (called ‘rejuvenation’) resulting in resting developmental stages with inert metabolic functions (Piraino et al. 2004). Polyp detachment, including polyp expulsion and polyp bail-out, is another form of reverse development, where a sessile polyp abandons its initial structure and becomes mobile again, maintaining its biological organization. Both polyp expulsion and polyp bail-out have been reported from shallow-water colonial scleractinians. Polyp expulsion is observed in physiologically healthy corals inhabiting chronically physically disturbed environments. Here, whole polyps, including their calices, leave the colony to settle elsewhere (asexual reproduction). This process seems to be regulated by the colony (Kramarski-Winter et al. 1997). Polyp bail-out, by contrast, refers to the escape of polyps without calices from a parent colony in response to acute environmental stress (e.g. Sammarco 1982; Kružić 2007; Capel et al. 2014; Serrano et al. 2018) and was also observed for octocoral species (Rakka et al. 2019; Wells and Tonra 2020). Thus, polyp bail-out provides a route of escape to new locations for possible resettlement. Additionally, fission (intratentacular budding), extratentacular budding (both with different modes), transverse division and asexual planula production have been reported for a number of tropical and temperate coral species as asexual reproduction modes (e.g. Cairns 1988; Tokuda et al. 2017), in some cases allowing the species to recover after external physical disturbance (Wilson 1979; Coppari et al. 2019). So far, the fragmentation of colony parts is the only reported asexual mode of dispersal in scleractinian cold-water corals (CWCs) (Wilson 1979; Le Goff-Vitry et al. 2004; Roberts et al. 2009; Dahl et al. 2012) and polyp detachment processes have not yet been described for solitary scleractinians.

Caryophylliidae are a diverse family of scleractinians with 269 species, colonizing water depths below 50 m (Roberts et al. 2009). *Caryophyllia huinayensis* Cairns et al. 2005 is a solitary, azooxanthellate member found in South Chile (36°S to 51°S) at water depths within the depth range 11–800 m (Fig. 1; Cairns et al. 2005; Häussermann and Försterra 2009; Sellanes et al. 2008). In spite of its small size (< 18.7 mm height and ≤ 8.7 mm calyx diameter), it is an important epibenthic component forming dense aggregations of up to 2211 ± 180 ind. m⁻² on steep walls of fjords and channels between 16 and 265 m (Häussermann and Försterra 2007; Wurz 2014). In Comau Fjord, terrestrial runoff leads to a permanent halocline subjecting the corals at depth to hypoxic and hypercapnic conditions (Silva 2008; Fillinger and Richter 2013). As with most CWCs, very little is known about its biology and development. Here, the first field and aquarium observations of reverse development in a solitary, scleractinian CWC are reported: polyp dropout in *C. huinayensis*, where an apparently unstressed, adult, solitary polyp shows tissue retraction, autonomously detaches as a unit from its skeleton and continues to live without calcium carbonate skeleton, with no visible fundamental modifications in its external body plan.

### Material and methods

In February 2014, a total of 30 *C. huinayensis* specimens were collected by SCUBA divers from Comau Fjord (Patagonia, Chile). In March and May 2015, a second and third batch of 15 and 24 individuals were sampled, respectively. Specimens were chiselled off the substrate (~ 21 m water depth, photic zone) and shipped in oxygen-replete seawater to Germany. After arrival (< 35 h from dispatch), corals were visually checked for polyp activity, tentacle appearance and colour (0% mortality) and maintained in two separate closed circuits (A and B; A: 30 individuals from 2014, B: 39 individuals from 2015) each filled with 285 L of artificial seawater mimicking the Comau Fjord environment at the sampling site of the corals (Online supplementary material 1). Corals were reared in the dark to avoid aquaria contamination with algae/cyanobacteria and thus repetitive cleaning which may cause stress. As populations of this species also occur in the aphotic zone (Sellanes et al. 2008), it is assumed that photoperiod is not of high importance for this species. The maintained corals were fed three times a week with live *Artemia franciscana*-nauplii and juvenile krill [*Euphausia pacifica*(ind. × week)] (Zierfischfutterhandel Norbert Erdmann e.K., Ritterhude, Germany). Each week, nutrient concentrations (nitrate, nitrite, phosphate and ammonium) were determined photometrically (Spectroquant® test, Merck KGaA, Darmstadt, Germany) just
before a third of the water in each circuit was exchanged. Deviations between the water changes were < 5% of the respective target value.

During the study, adults of circuit A reproduced and 14 larvae settled on the aquarium glass in July 2015. The recruits (Calyx $\varnothing$ 2.3 ± 0.3 mm) were removed with razor blades without any visual damage, glued (Super Flex Glue Gel, UHU GmbH and Co KG, Bühl, Germany) on glass slides and maintained in circuit A. After this transfer, they grew at a similar rate (3.4 ± 1.9 mm a$^{-1}$) as individuals in situ (Wurz 2014). This indicates good rearing conditions in the aquaria system.

Polyp appearance and activity were visually monitored on a daily basis. Based on that, polyps were classified to one of the following three stages: (1) Attached polyp—extended tissue in full contact with skeleton; (2) partially detached polyp—recession of tissue from lower to upper calyx and between septae and partial loss of contact of tissue with skeleton, with individual strands of tissue remaining; and (3) fully detached polyp—complete loss of contact with skeleton, tissue sinking to the bottom. The detached polyps were kept in Petri dishes (Ø 10 cm) submerged in the respective circuit and continued to be fed as specified above. Two of the detached polyps were inadvertently lost when cleaning circuit A five weeks after detachment, but one specimen (#3, circuit B) was monitored for more than seven months. Specimens were examined under a stereomicroscope biweekly to check for reattachment, skeleton precipitation and polyp activity. In order to calculate potential in situ dispersal by currents, the sinking rate of this specimen was determined in five runs, five weeks after the detachment by transferring it to a gridded 350 mm cylinder and estimating the time it took the specimen to sink to the bottom.

During austral winter 2019, spring 2019 and summer 2019/2020, a field study was conducted by SCUBA divers at Comau Fjord (Cross-Huinay North, 42°23’12.8” S, 72°27’46.3” W and Lliguapi 42°9’43” S, 72°35’55” W) (3 seasons × 2 sites = 6 transects) to determine the proportion of C. huinayensis undergoing polyp detachment in situ. Photographs were taken with a digital camera (Nikon D7000 DSLR, 24 mm lens, 19.5 cm × 29.6 cm frame) between 24 and 27 m water depth along a horizontal 35-m line transect in 1 m intervals. C. huinayensis individuals were counted, and the polyps were classified into three stages of polyp detachment, described from the aquaria experiment: polyp attached; partially detached; and fully detached.

**Results and discussion**

This is the first report of reverse development in a scleractinian CWC. Four out of 83 aquarium specimens (4.8%), two in each circuit, dropped out and exposed the white skeletons of the coralites (Fig. 2d, Online supplementary material 2). Microscopic examination of the vacated skeletons indicated that no tissue remained on them. The two dropouts of circuit A were former recruits (see Materials and Methods, now adults), and the two dropouts in circuit B were from wild corals. In one of the latter, the tissue began to recede twelve weeks before polyp dropout (Fig. 2, Online supplementary material 2, 3). An additional adult individual (circuit B) showed tissue recession, however, remained partially attached for ten weeks (Fig. 2b, Online supplementary material 2, 3). The other corals (A: $n = 42$, B: $n = 36$) remained attached. No unusual behaviour was observed neither before nor after the event in any of the attached fellow corals, so that contamination or disease in the rearing system seems unlikely, as it would have affected more individuals.

The observed polyp dropout in the solitary C. huinayensis is considered to be an autonomous detachment of a previously unstressed polyp. It is preceded by tissue retraction and leads to detachment of the entire adult polyp which subsequently lives on without a calcium carbonate skeleton in an unattached state. No fundamental changes in the external body plan are apparent, and the oral disc with the mouth and the tentacles as well as the mesenteries remain (Fig. 2a–c).

Polyp dropout differs from other reported forms of reverse development confined to stressed shallow-water colonial scleractinians in tropical, subtropical and temperate environments in a number of ways (Table 1). In contrast to (i) reversible metamorphosis (Richmond 1985) and (ii) ontogeny reversal (Piraino et al. 1996, 2004), polyp dropout is not accompanied by fundamental changes in the body plan. Polyp dropout also differs from (iii) polyp expulsion (Kramarski-Winter et al. 1997) and (iv) polyp bail-out (Sammarco 1982; Kružić 2007; Capel et al. 2014; Serrano et al. 2018). These are escape mechanisms of polyps from chronically physically disturbed (polyp expulsion) or acutely stressed disintegrating (polyp bail-out) colonies, resulting in dispersal and, after successful resettlement, asexual reproduction. The latter requires an adult and a reproductive product, but in the case of polyp dropout, the whole adult moves away from its skeleton and there is no reproductive product. In contrast to the polyp detachment processes described so far, polyp dropout occurs in the absence of abiotic or biotic stressors and does not appear to be related to reproduction (Table 1).
The dropout polyps had a constricted appearance with mesenterial filaments protruded apically through the mouth, but also laterally through the body wall (Fig. 2c). While the apical exposure is not uncommon, the later feature is absent in attached specimens not least because of the barrier built by the calcareous skeleton. The behaviour of the mesenterial filaments (e.g. whether they can be retracted and the openings in the body wall can be closed again) cannot be described because a more detailed examination would have caused a disturbance for the polyp. Similar observations of protruded mesenterial filaments were reported for detached polyps of the tropical scleractinian *Mussa angulosa* (Goreau et al. 1971) and planula larvae of *Pocillopora damicornis*, when the formation of an organic film was prevented in an attempt to keep the larvae from settling (Richmond 1985). Also, some
Table 1  Reported stressors causing polyp detachment in scleractinian corals and likelihood of their involvement in the observed detachment of *Caryophyllia huinayensis*

| Stressor                      | Species                  | Climatic zone | Reference                               | Probability for the present event | Justification                                                                 |
|-------------------------------|--------------------------|---------------|-----------------------------------------|-----------------------------------|--------------------------------------------------------------------------------|
| Low oxygen                    | *Seriatopora hystrix* gr. | Tropics       | Sammarco (1982)                         | Very unlikely                     | Minimal concentration of oxygen 8.33 mg/l, water circulated constantly, (Comau Fjord: 4-12 mg/l; 2 h hypoxia did not cause detachment (A. Castrillon, pers. comm.) |
|                               | *Pocillopora damicornis* |               | Richmond (1985)                         |                                   |                                                                                  |
| High/low temperature          | *Acropora tenuis*        | Tropics       | Yuyama et al. (2012)                    | Unlikely, unless one assumes that reduced variability induces change | Temperature was monitored constantly in a climatized cool room; measured values ranged between 12.0 and 13.3 °C, without seasonal variation (Comau Fjord: 9.5–16 °C) |
|                               | *Pocillopora damicornis* | Subtropics    | Fordyce et al. (2017)                   |                                   |                                                                                  |
|                               | *Favia favus*            |               | Kramarsky-Winter et al. (1997)          |                                   |                                                                                  |
|                               | *Cladocora caespitosa*   | Temperate region | Kružić (2007)                          |                                   |                                                                                  |
|                               | *Oculina patagonica*     | Temperate region | Kramarsky-Winter et al. (1997)          |                                   |                                                                                  |
| High/low salinity             | *Pocillopora damicornis* | Tropics       | Shapiro et al. (2016)                   | Very unlikely                     | Salinity of new water introduced to the system was measured and adjusted; values of the water in the system ranged between 30.4 and 33.2 reflecting the natural salinity range in the coral’s habitat (see Table Online supplementary material 1). There were no abrupt changes in salinity in the aquarium system |
|                               |                         |               | Liu et al. (2020)                       |                                   |                                                                                  |
| High/low pH                    | *Pocillopora damicornis* | Tropics       | Kvitt et al. (2015)                     | Very unlikely                     | The pH of the maintenance water and that of new water introduced to the system during water exchange are controlled and ranged between 7.8 and 8.0 reflecting the natural pH range in the coral’s habitat (see Table Online supplementary material 1). There were no abrupt changes in pH in the aquarium system |
| Sedimentation                 | *Favia favus*            | Subtropics    | Kramarsky-Winter et al. (1997)          | Very unlikely                     | Water was constantly, automatically treated, and no sediment was added to the system; sediment stress experiments (sediment: water = 1: 1000) did not cause detachment (unbub. data.) |
|                               | *Oculina patagonica*     | Temperate region | Kramarsky-Winter et al. (1997)          |                                   |                                                                                  |
| High nutrient concentrations  | *Acropora tenuis*        | Tropics       | Yuyama et al. (2012)                    | Unlikely                          | Nitrate concentration (0.1–5.9 mg/l) exceeded natural values (Comau Fjord: 0.18–1.1 mg/l) 95% of the time, however, remained stable (no sudden increases) over the entire observation period and was within the acceptable range for coral husbandry (Borneman 2008). As the other coral individuals remained unaffected, an effect of nitrate is unlikely. In addition, warm water scleractinians are optimally maintained at 5-10 mg nitrate/l. Phosphate concentration (0.06–0.28 mg/l) was in the natural range (Comau Fjord: 0.12–0.36 mg/l) |
anemones expel filament ends (acontia) through body wall openings (cinclides) (Manuel 1988). The perforations of the coral body wall likely affected the hydroskeleton supporting the body and tentacles of attached polyps, causing the rather retracted state (Fig. 2c). It is also likely that the impaired hydroskeleton prevented the reattachment of the polyps. Accordingly, stereomicroscopical observations confirmed no signs of skeleton precipitation under the detached polyps within the four-week (circuit A) and seven-month (circuit B) observation period, respectively, similarly to the Mediterranean scleractinian Astroides calycularis (*12-week observation; Serrano et al. 2018). By contrast, a small portion of bailed-out polyps of the tropical coral Seriatopora hystrix was able to initiate skeletogenesis as early as nine days after detachment (Sammarco 1982).

Some of the C. huinayensis dropouts attached to the glass with their tentacles. However, we have no evidence for resettlement on the glass, although C. huinayensis larvae settle on it successfully. The swimming anemone Stomphia coccinea is able to routinely detach and reattach to substrate (Robson 1961) and may serve as a model for polyp detachment/reattachment. Before adhesion, its pedal disc becomes sticky due to the discharge of nematocysts (Ellis et al. 1969). The latter may be lacking in the aboral calicoblastic ectoderm of dropped out C. huinayensis. Indeed, after settlement of scleractinian planula larvae, the aboral ectoderm undergoes ultrastructural changes including the loss of secretory cells and nematocysts (Vandermeulen 1975). While it is uncertain whether the detached C. huinayensis are able to restore this capacity beyond our observation period (seven months), we cannot rule out this possibility, given that expelled polyps of the tropical scleractinian coral Tabastreaea coccinea have been shown to resettle after seven months (Capel et al. 2014).

Table 1 continued

| Stressor                      | Species              | Climatic zone | Reference                                                                 | Probability for the present event | Justification                                                                 |
|-------------------------------|----------------------|----------------|---------------------------------------------------------------------------|-----------------------------------|--------------------------------------------------------------------------------|
| Other allelopathic effects    | *Tabastreaea coccinea* | Tropics        | Capel et al. (2014); Algae: pers. comm. K. Capel Sin et al. (2012)          | Very unlikely                    | Autotrophs cannot grow in the system as it is permanently dark                  |
| Presence synthetic toxins     | *Pocillopora damicornis* | Tropics        | Wecker et al. (2018)                                                      | Very unlikely                    | These substances are not used in or near the aquarium unit                      |
| High/low food                 | *Mussa angulosa*      | Tropics        | Goreau et al. (1971); Richmond (1985); Goreau and Goreau (1959)           | Very unlikely                    | Fed three times a week with live Artemia salina nauplii, with the spacing of the corals greater than their diameter with expanded tentacles, additionally each coral specimen was handled once a week with a juvenile krill. Furthermore, it is likely that the water contained other food resources such as dissolved and particulate organic matter and micro-organisms. As the other coral individuals remained unaffected, an effect of starvation/ lack of nutritional diversity is unlikely |
|                               | *Pocillopora damicornis* | Not defined    |                                                                           |                                   |                                                                                 |
|                               | *Astroides calycularis* | Temperate region |                                                                           |                                   |                                                                                 |

If we assume that reattachment beyond our observation period might be possible, and because coral tissue is only slightly denser than seawater, we can suppose that in Comau Fjord dropouts may potentially drift with the currents, reach new habitats, resettle and reproduce. Sinking rate was 2.6 ± 0.3 cm s⁻¹. Tidally corrected current velocities of 5 cm s⁻¹ (maximum 15 cm s⁻¹) were recorded in the vicinity of the coral banks (unpubl. data). Considering the steep walls and the species’ distribution to at least 265 m (Häussermann and Försterra 2007), polyps dropped out at 20 m water depth may potentially drift lateral up to 3.7 km until they reach 265 m.

No direct evidence for polyp dropout was found in situ, but the partially detached stage, which precedes dropout over a longer period of time (twelve weeks for specimen #5 in vitro), provided indirect evidence for dropout in natural field populations of C. huinayensis. In Comau Fjord, we documented partial polyp dropout in 1.4 ± 0.8% of the polyps (six phototransects, total number of polyps analysed: 9322). Partially detached polyps were detected at both sites (1.3 ± 1.1% of the polyps at Cross-Huinay...
North, 1.5 ± 0.5% of the polyps at Liliguapi) and did not follow a common pattern in terms of frequency over the three seasons analysed. These field observations provide a strong indication that polyp dropout is a process which occurs naturally, however, under normal environmental conditions in only a few individuals. Thus, sexual dispersal apparently dominates over the asexual one (here drift of dropouts), similar to the tropical scleractinian coral *Pocillopora meandrina* (Magalon et al. 2005).

Although the respective five aquarium specimens did not show any external signs of physical stress or disease during the > 20 months maintenance period and also shortly before the polyp dropout, the possibility that some unmonitored external agent may have accounted for the observed changes cannot be excluded. It is unlikely that the removal process of the former recruits and the collection of adults had an impact 20 months after the transfer, especially as the corals grew as expected and partial detachment, indicative of imminent dropout, was also observed in situ. Furthermore, the lack of light during the maintenance period hardly seems to be a trigger, since only 6% of the aquarium corals showed symptoms of dropout. Moreover, partially detached corals were observed in situ, exposed to a light–dark cycle even under summer conditions. Weekly monitoring of water parameters did not reveal changes in the physico-chemical environment at the time or twelve weeks prior to the event (ANOVA $p > 0.05$).

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Declarations

Conflict of interest On behalf of all authors, the corresponding author states that there is no conflict of interest.

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