First evidence for backcrossing of F₁ hybrids in Acropora corals under sperm competition

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Acropora is a species-rich genus of reef-building corals with highly diverse morphologies. Hybridization among intercrossing species potentially influences species diversity within Acropora. However, the mechanisms that allow hybridization/backcrossing remain unknown. Although we tested a limited number of species, we hypothesized that Acropora gametes in the Indo-Pacific may preferentially fertilize conspecific gametes despite their compatibility with heterospecific gametes, leading to infrequent hybridization between potentially intercrossing species. In this study, F₁ hybrids of Acropora florida and A. intermedia showed specific fertilization trends. For example, sperm had the ability to backcross with the parental species even in the presence of sperm from the parental species. Also, eggs of the hybrids produced from A. florida eggs and A. intermedia sperm ("FLOint") exhibited self-fertilization. Since a low ratio of hybridization between A. florida and A. intermedia is predicted, the population size of hybrids should be small. Therefore, self-fertilization would facilitate reproduction of the hybrid in nature, while remaining sperm could outcompete parental species sperm to backcross with eggs. Although we succeeded in breeding two colonies of hybrids, it is reasonable to speculate that hybrids show a high tendency to choose the most efficient sexual reproduction tactics.

Hybridization is considered a mechanism for evolutionary innovation. Introgressive hybridization is caused by the repeated backcrossing of hybrids to the parental species¹. Introgressive hybridization results in new gene combinations, leading to transgressive phenotypes². Extensive hybridization is associated with the rapid diversification of species³. Moreover, hybrids can potentially occupy new habitats, differentiating them from the parental species⁴. Implications of hybridization for adaptation and hybrid fitness have been suggested⁵–⁷, but the ways in which introgression occurs in nature are still unknown.

In the Indo-Pacific, the reef-building coral Acropora spp. is species rich (> 110 species)⁸, and there is a potential relationship between hybridization and high species diversity⁹. For example, tabular species such as A. hyacinthus can form species complexes⁹,¹⁰, and gene flow among such species complexes occurs in a complex manner¹¹. In addition, intermediate morphologies among intercrossing species imply that admixture events are associated with morphological diversity and similarity¹²,¹³. Morphological similarity is associated with hybridization/introgression in Caribbean¹⁴–¹⁷ and Indo-Pacific Acropora¹⁸,¹². In both the Indo-Pacific and Caribbean, co-occurrence of spawning times/dates and gamete compatibility is related to introgression¹⁸–²⁰. For repeated hybridization events among intercrossing species, the backcrossing of F₁ hybrids to the parental species must occur, but the reproduction of F₁ hybrids has not yet been fully investigated in the Southern Japanese Indo-Pacific, which is a hybrid hotspot area with high species richness of the coral Acropora¹⁹.

For introgression between two species, F₁ hybrids from the two species must backcross with the parental species. However, such reproductive strategies, including the fertilization mechanisms of F₁ hybrids in the Indo-Pacific, have not been fully elucidated, because Indo-Pacific hybrids other than A. florida and A. intermedia¹¹ have not been successfully raised to their spawning age of approximately 7 years²¹. In nature, repeated hybridization between parental species arising from the backcrossing of F₁ hybrids has been demonstrated¹⁴ in A. prolifera, the hybrid of two species inhabiting the Caribbean¹⁴. Therefore, although the importance of hybridization in the species-rich Indo-Pacific reef-building coral Acropora has been posited, there is no evidence showing how hybrids reproduce at the gametic level, or how they backcross and/or mate with each other.

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Although introgression in the coral Acropora has been shown7,11,22, how F1 hybrids backcross with the parental species remains unknown. Our previous study showed that sperm of the F1 hybrids named “FLOint” and “INTflo”, bred from A. florida and A. intermedia, are compatible with eggs of the parental species21. In addition, eggs of the hybrid FLOint, raised from A. floridensis and A. intermedia sperm, showed high selfing21. Ecologically, mating opportunities within F1 hybrids are more limited than backcrossing opportunities with the parental species due to smaller numbers of F1 hybrid colonies. For hybrids to backcross with the parental species, the eggs of the parental species must accept hybrid sperm but tentatively prefer to mate with conspecific sperm in the presence of both conspecific and heterospecific sperm22. To clarify how gametes of the F1 hybrid mate, we examined whether hybrid sperm can outcompete parental species sperm, and whether hybrid FLOint eggs always show selfing. Using the results of these analyses, we show fertilization trends of the gametes of F1 hybrids (A. florida and A. intermedia) in the Indo-Pacific that lead to backcrossed and F2 generations of hybrids.

Materials and methods

Coral collection. In 2016, Acropora florida fragments from six colonies were collected from Majanohama, Akajima (Aka Island), Japan (26° 120 N, 127° 170 E), and two F1 hybrid colonies (INTflo: A. intermedia × A. florida sperm; FLOint: A. florida × A. intermedia sperm) were kept at Aka Island port. In 2016, we detected no A. intermedia with mature eggs, and thus A. intermedia spawning was supposed to have occurred during the previous full moon. In 2017, the hybrids (FLOint and INTflo) were used in experiments12,21, and two F1 hybrids, measuring approximately 30–50 cm, were transferred to Sesoko Station from Aka Island and maintained in an aquarium tank. In 2017, fragments from seven colonies of A. florida and 12 colonies of A. intermedia were collected from Sesoko Island (26° 37 N, 127° 51 E). All colonies and fragments were kept in a running seawater tank at the Akajima Marine Science Laboratory in 2016 or Sesoko Station at the University of the Ryukyus in 2017 until 1–5 days before their predicted spawning date.

Spawning observation and gamete collection. Corals were observed at 20:30 from 5 days before their predicted spawning date on a full moon. When bundles were observed at the mouth of each polyp, the corals were transferred to a tank filled with seawater, and the time of spawning was recorded. The bundles were collected from the colony using plastic pipettes (Table 1), and gametes were separated into sperm and eggs using 100-µm plankton mesh, following Morita, et al.24. The sperm concentration of the isolated spermatozoa was determined using a hemocytometer. The final sperm concentrations were adjusted to approximately 10^4, 10^5, or 10^6 sperm/mL for use in the subsequent crossing experiments.

Crossing experiments and paternity tests. In the crossing experiments, approximately 200 eggs were transferred to 5 mL filtered seawater and 5 mL sperm suspension according to previously described methods25,26. The experiments were performed using A. florida and FLOint hybrid gametes on 19 June 2016 at Aka Island, and A. intermedia and INTflo hybrid gametes on 6 June 2017 at Sesoko Island. The spawning dates of A. florida and the two hybrids (INTflo and FLOint) did not coincide in 2017; INTflo spawned on 6 and 16 June and FLOint on 16 June, but A. florida spawned on 7, 8, and 11 July. Therefore, crossing experiments using FLOint and A. florida were not conducted.

For the crossing experiments, sperm concentrations were 10^4, 10^5, and 10^6 sperm/mL for the sperm non-choice and sperm choice choice tests (fertilization trials in the presence of both parental species and hybrid sperm). In the crossing experiments, only colonies that spawned on the same day were used. A. florida and the FLOint hybrid were used in 2016, and INTflo and A. intermedia were used in 2017 (Table 1, Supplementary Table 1). The fertilization ratio was recorded based on whether or not eggs showed cell division within 2–4 h after mating.
at a temperature of 29 °C. Larvae from 3 to 4 days after fertilization were preserved in 99.5% ethanol and used for paternity tests. Paternity tests were performed to confirm which sperm fertilized the egg in the sperm choice test. DNA was extracted from the larvae, and microsatellite analysis was performed using the extracted DNA as a template with the markers 11745m3 and 11401m4, according to Kitanobo et al. For each marker, fewer than two alleles were detected in the hybrids, and the same alleles were consistently detected from sperm and tissues, suggesting that the hybrids were not chimera (Supplementary data 1). Some analyses were conducted using acrylamide gel electrophoresis and others using fragment analysis with the ABI 3130xl or 3730xl DNA Sequencer (Applied Biosystems, Waltham, MA, USA). Microsatellite Analysis v1.0 (Applied Biosystems) software (https://www.thermofisher.com/order/catalog/product/4381867) was used to score the sizes.

**Statistical analyses.** We conducted Tukey's honestly significant difference (HSD) tests to evaluate differences in multiple comparisons. Welch's two-sample t-tests were used to confirm differences in the fertilization ratio when using heterospecific sperm. All statistical analyses were performed using R version 4.0.1.

**Ethical approval.** All applicable international, national, and/or institutional guidelines for sampling, care, and experimental use of organisms for the study have been followed, and all necessary approvals have been obtained (No. 31-30).

**Results**

**F1 hybrid spawning times and dates.** Hybrids (FLOint; one colony) and *A. florida* (five colonies) released gametes in June 19–21, 2016 on Aka Island. *A. intermedia* did not release gametes synchronously with the hybrids. *A. intermedia* with mature eggs were not found, and it is likely that most *A. intermedia* had already spawned around the full moon at the end of May (Table 1). In 2017, spawning of the hybrid INTflo and *A. intermedia* coincided, but *A. intermedia*, *A. florida*, and the hybrid FLOint did not spawn on the same date (Table 1). Most colonies of *A. florida* spawned in July rather than June.

**Percentage of eggs fertilized in crosses among conspecífics.** Crossing experiments using hybrids and parental species were conducted using *A. florida* and FLOint hybrid gametes on 19 June 2016, and *A. intermedia* and INTflo hybrid gametes on 6 June 2017. Fertilization ratios between the parental species were low in *A. florida* (<11%) (Table S1). In contrast, fertilization ratios among *A. intermedia* were high under both low and adequate sperm concentrations, but the colony (13_12) had very low fertility ratios (Table S1).

**Inherent selfing and backcrossing of F1 hybrid eggs.** Self-fertilization of the INTflo and FLOint gametes was examined. INTflo gametes did not show self-fertilization in the presence of low (10⁴ sperm/mL) to high (10⁶ sperm/mL) sperm concentrations (Table S1). By contrast, as in our previous study, the eggs of FLOint showed high selfing ratios of 76%, 89%, and 76% under low, moderate and high sperm concentrations, respectively (10⁴, 10⁵, and 10⁶ sperm/mL) (Table S1).

To investigate the inherent ability of hybrid eggs to backcross with the sperm of the mother species, *A. florida* or *A. intermedia*, gametic compatibility was examined. Hybrid eggs showed a high ratio of fertilization to the sperm of the mother species; INTflo crossed with the sperm of *A. intermedia*, and FLOint showed a high ratio of fertilization to *A. florida* sperm (Fig. S1). Moreover, there was no significant difference in fertilization ratio among the different sperm concentrations (10⁴, 10⁵, and 10⁶ sperm/mL) for each hybrid (FLOint eggs × *A. florida* sperm: Tukey HSD P > 0.05, INTflo eggs × *A. intermedia* sperm: Tukey HSD P > 0.05).

Since high self-fertilization was observed for the eggs of FLOint in this and our previous study, we performed paternity testing to determine whether FLOint eggs can be fertilized by *A. florida* sperm in sperm choice experiments (fertilization trials in the presence of both parental species and hybrid sperm). The results showed that most eggs were self-fertilized in the presence of both *A. florida* sperm and FLOint sperm (Fig. 1). On the other hand, most INTflo hybrid eggs backcrossed with *A. intermedia* sperm in the presence of INTflo sperm (Fig. 1).

**High compatibility of F1 hybrid sperm and eggs of the parental species.** INTflo and FLOint sperm both backcrossed with eggs of the maternal parent of each hybrid (Fig. S2). For *A. florida* eggs, the ratio of fertilization to conspecific sperm was much lower (Table S1) than in previous trials, such as that conducted in 2015, although the reason for this is unclear. For *A. intermedia* eggs, the fertilization ratio did not change with sperm concentration (*A. intermedia* eggs × INTflo sperm: Tukey HSD P > 0.05). As in *A. intermedia*, the fertilization ratio of *A. florida* eggs also did not significantly differ with sperm concentration (*A. florida* eggs × FLOint sperm: Tukey HSD P > 0.05).

**Can F1 hybrid sperm compete with conspecific sperm to backcross with eggs of the maternal species?** To examine backcrossing and self-fertilization ratios in the presence of parental species and hybrid sperm (*A. intermedia* and INTflo sperm or *A. florida* and FLOint sperm, respectively), paternity tests were performed. Fertilization ratios in the sperm choice experiments were not significantly different among combinations (Fig. S3; INTflo eggs × INTflo sperm and *A. intermedia* sperm: Tukey HSD P > 0.05, *A. intermedia* eggs × INTflo sperm and *A. intermedia* sperm: Tukey HSD P > 0.05).

Eggs of the parental species (*A. florida* and *A. intermedia*) were fertilized by hybrid sperm in the presence of conspecific sperm. In case of INTflo sperm backcrossing, the hybrid sperm fertilized *A. intermedia* eggs independent of the sperm concentration of the parent species (Fig. 2, Tukey HSD P > 0.05) or hybrid sperm in...
concentration (Tukey HSD P > 0.05). FLOint sperm also backcrossed with parent species A. florida eggs independent of hybrid sperm concentration (Fig. 2, A. florida egg × A. florida sperm and FLOint sperm; T-test, mt = 0.1, df = 1.78, P > 0.05).

**Discussion**

In our study, the F₁ hybrid FLOint showed adequate fertilization patterns for reproduction, with high rates of backcrossing to the parental species and most of the eggs showing selfing. Our results showed that hybridization may arise when colony numbers decline due to heavy bleaching events. For example, A. florida eggs could hybridize with A. intermedia sperm at low sperm concentrations (FLOint)23. Conversely, A. intermedia eggs were preferentially fertilized by conspecific sperm when exposed to both A. intermedia and A. florida sperm23. Since the recovery of reefs after heavy bleaching often takes more than 10 years, and reef species composition changes frequently28,29, the post-bleaching F₂ generation is predicted that they need to reproduce within lower number of colonies. In contrast to the Caribbean hybrid A. prolifera (< 25% selfing)23, FLOint showed more than 95% selfing (Fig. 1). Six to fourteen eggs and 10⁵ sperm are packed into Acropora gamete bundles30,31; thus sperm not involved in the self-fertilization of bundled eggs can mate with other unfertilized eggs. However, for backcrossing, spawning synchronicities of the hybrids FLOint and A. intermedia and INTflo and A. florida were
Sperm from the hybrids FLOint and INTflo potentially mate with parental species eggs when hybrid and parental species sperm compete. Sperm choice experiments in this study showed that the competencies of FLOint and INTflo sperm were high enough to outcompete parental sperm even when the number of hybrid sperm was far lower than that of the parental species sperm (Fig. 2). To support this result, repeated backcrossing is suggested in Caribbean Acropora. In addition, there were fewer than two alleles of microsatellites, indicating polyploidy of the hybrids involving two chromosomes, unlike that reported in a previous study. Therefore, fusion of the parental species is predicted to occur, but as the two parental species A. florida and A. intermedia are morphologically distinct, lineage fusion does not occur extensively at present.

From the present study, interspecific hybridization and introgression are suggested, but molecular based analyses for examining admixture events between the parental species are needed. From our preliminary SNP-based analyses, gene flow occurred among intercrossing species showing spawning synchronicity and high gamete compatibility (Kitanobo et al., unpublished data). However, the detection of hybridization is influenced by methodological differences, and hybrid lineages are rare (five species). Moreover, integrative approaches from breeding trials and morphological and molecular based analyses indicate that morphologically distinguishable species can be reproductively isolated and can evolve independently. Contrary to a previous study, tabular species do not cross with other morphospecies, but A. florida and A. intermedia gametes show high rates of intercrossing. In addition, gametes showed specific fertilization patterns according to sperm concentration, and the patterns of the F₁ hybrids also indicate that backcrossing is highly probable. However, it seems difficult to

**Figure 2.** Backcrossing of hybrid sperm with parental species. Sperm choice experiments using eggs of the parental species are shown. Light grey bars indicate the fertilization ratio of parental species sperm, and white bars indicate the sperm of the hybrid species. Microsatellites were checked using electrophoresis or Microsatellite Analysis v1.0 (Applied Biosystems) software (https://www.thermofisher.com/order/catalog/product/4381867).
distinguish between hybridization and incomplete lineage sorting if an admixture event occurred in the early speciated *Acropora* (< 6 Ma)\(^3\). From our study, a slight admixture event between *A. florida* and *A. intermedia* may be ongoing, but detailed comprehensive studies involving other intercrossing species are needed. In this study, we focused on only two intercrossing species, *A. florida* and *A. intermedia*, but *A. intermedia* shows high rates of crossing with other sympatric and synchronous spawning species\(^3\). As discussed above, the morphologically distinct species of *A. florida* and *A. intermedia* show slight differences in spawning times and dates (Table 1); thus, these two species are tentatively at lower risk of hybridization. *A. intermedia* and other intercrossing species such as *A. gemmifera* are more likely to hybridize due to their overlapping spawning times\(^3\).

Our study also shows that delimitation of species is suspected in the morphologically distinct intercrossing species *A. florida* and *A. intermedia*. Although our study used limited numbers of hybrids, the sperm competency of F\(_1\) hybrids was sufficient to provide opportunities for mating with parental species (backcrossing) and selfing in FLoint. This would be beneficial to the production of the F\(_2\) generation in cases of solitary spawning due to reduced spawning synchronicity or a reduced number of colonies of the parental species (Fig. 3). These features are congruent with high rates of introgression events under past climate changes\(^2\),\(^6\), and hybrid hotspots are located at biogeographic borders including Southern Japan\(^8\). Although the unique fertilization patterns of F\(_1\) hybrids are potentially not involved in ongoing hybridization, they may be a footprint of the past hybridization of ancestral species, and these results can be used to understand the complex history of the coral *Acropora*.

**Figure 3.** Schema of F\(_1\) hybrid reproduction. Flow from *A. intermedia* and *A. florida*, F\(_1\) hybrids, to reproduction of F\(_1\) hybrids. According to our studies, hybridization can occur when the sperm concentration is low, and an F\(_1\) hybrid (FLoint) can backcross or reproduce asexually. Bold lines indicate probable pathways, which are supported by this and our previous studies; faint grey lines indicate pathways that may not occur because of the low probability of hybridization.

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Author contributions
S.K. conducted all of the experiments, and wrote a paper. K.I., H.F., N.I. supported fertilization experiments and edited a paper. M.M. conceived experiments, support microsatellite analyses, and wrote a paper.

Competing interests
The authors declare no competing interests.

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