Export–import dynamics of zooplankton on a coral reef in Palau

W. M. Hamner1,*, P. L. Colin2, P. P. Hamner3

1, 3Department of Ecology and Evolutionary Biology, University of California Los Angeles, Box 951606, Los Angeles, California 90095-1606, USA
2Coral Reef Research Foundation, PO Box 1765, Koror, Palau PW 96940

ABSTRACT: Although coral reefs are widely viewed as oases in an oceanic desert, we demonstrate that the oceans around coral reefs in Palau are not deserts devoid of food, nor are coral reefs oases isolated from the surrounding sea. On flood tides, oceanic zooplankton advects onto the Palau barrier reef, where it is consumed by a wall of planktivorous fish. Similarly, on ebb tides lagoon zooplankton is consumed as it advects across the reef flat. Coral reef planktivores ingest zooplankton imported from both the sea and the lagoon, but coral reef fish also export meroplanktonic fish eggs that disperse offshore. Every day about 30 species of reef fishes, some forming spawning aggregations (parrotfish and surgeonfish), spawn at the forereef when the tide turns, releasing clouds of buoyant eggs and sperm into the ebb-tide waters that flow off the reef flat. Fish eggs disperse seaward, but remain entrained within several kilometers of the reef in a tidal, neritic water mass, separated from oceanic currents by tidal fronts. Flood tides return fish embryos and offshore zooplankton to the reef. The flux of materials between oceanic, neritic, and lagoon waters near coral reefs must be reexamined.

KEY WORDS: Coral reef zooplankton · Fish spawning · Egg dispersal · Palau

INTRODUCTION

Coral reefs exchange water, plankton, and non-living materials with the adjacent ocean and lagoon. Early literature (Odum 1953, Odum & Odum 1955, Johannes et al. 1972, Johannes & Gerber 1974) claimed coral atolls were ‘oases of life in an oceanic desert’, energetically closed ecosystems with limited exchange with the surrounding sea, retaining production, recycling nutrients, and flourishing in oligotrophic oceans. Unfortunately this paradigm has been generalized to all categories of coral reefs. Barrier reefs, however, surround high islands or occur along continental margins, where terrestrial nutrients discharge into barrier reef lagoons with longer residence times and higher productivity than exhibited by coral reef atoll lagoons (Birkeland 1982, Sammarco & Crenshaw 1984). Our data demonstrate that the barrier reef in Palau is strongly connected to the islands, lagoon, and sea, a finding supportive of more recent research (Pickard 1986, Hamner et al. 1988, Wolanski & Hamner 1988, Sorokin 1990, Kinsey 1991, Hatcher 1997). We assess import and export of zooplankton for a barrier reef around high islands in Palau, at a site where circulation is controlled by tidal currents, and we document changes in zooplankton composition as it crosses the reef. We also describe episodic spawning of reef fishes with planktonic eggs and evaluate contribution of fish eggs to the total plankton biomass exported from the barrier reef each day.

At Enewetak in the mid-Pacific Ocean (Odum & Odum 1955, Johannes & Gerber 1974, Atkinson et al. 1981) and Davies Reef in the Great Barrier Reef (Pickard 1986, Hamner et al. 1988) trade winds and wave-overturning generate an almost constant unidirectional flow from ocean to lagoon. Our study area in Palau (see Fig. 1), however, is protected from easterly waves by barrier reef topography and from southeasterly swells by an offshore, submerged barrier reef, and semidiurnal tides control flow from sea to lagoon.

*Email: whamner@ucla.edu
(flood) and lagoon to sea (ebb), allowing assessment of export and import of zooplankton across the reef.

We expected in Palau that on flood tides ocean waters would advect onto the reef, where mesozooplankton would be consumed by planktivorous fish and filter feeders (Emery 1968, 1973, Hobson & Chess 1978, Hamner et al. 1988, Sorokin 1990, Holzman & Genin 2003, Fabricius & Metzner 2004, Heidelberg et al. 2004). We also expected that breaking waves on the forereef (modest at our study site) and grazing fishes would generate modest amounts of reef detritus, which would be transported across the reef toward the lagoon (Alongi 1997, Crossman et al. 2001, Yahel et al. 2002). On ebb tides, lagoon mesozooplankton, different from that found on reefs or in the open sea (Emery 1968, Tranter & George 1972, Hobson & Chess 1978, 1979, Hamner & Carleton 1979), flows onto the Palau barrier reef, and we expected that lagoon zooplankton would also be consumed by planktivores in transit across the reef.

In Palau a suite of 30 to 40 species of reef fishes, principally surgeonfishes, parrotfishes, and wrasses, spawn for a few hours after high tide along the forereef throughout the year (P. L. Colin pers. obs.). At our study site, the parrotfish *Chlorurus sordidus* and the 2 surgeonfishes *Acanthurus nigrofuscus* and *Ctenochaetus striatus* dominate spawning aggregations at intervals of about 50 m along the reef face (Colin & Bell 1991, Domeier & Colin 1997, P. L. Colin pers. obs.). The eggs of the scarids *Chlorurus* and *Scarus* are spindle-shaped, making them easy to distinguish, while those of most other reef fishes (acanthurids, labrids, some scarids and other families) are spherical. Both types of eggs are transparent, buoyant, and relatively large (*C. sordidus*: 1.38 × 0.47 mm; *Acanthurus* sp.: 0.58 mm diameter), and are collected readily with neuston nets (Zaitsev 1971, Omori & Ikeda 1984). We hypothesized that on falling tides a substantial biomass of these fish eggs would be exported from the forereef into the open sea, because the spawning fish aggregations are invariably seaward of most planktivorous fish along the forereef.

We quantified import and export of mesozooplankton across the Palau barrier reef and documented changes in zooplankton assemblages as they advected across the reef. In addition, we documented episodic spawning of reef fishes with planktonic eggs, and quantified the contribution of meroplanktonic fish eggs exported from, and to a lesser degree, imported onto the reef each day. Although our sampling effort emphasized collection of fish eggs exported offshore during ebb tide, we also collected mesozooplankton from the lagoon and from the sea during flood and ebb tides, permitting us to address the issue of tidal export–import dynamics.

**MATERIALS AND METHODS**

We sampled for fish eggs and mesozooplankton between the open sea, the eastern barrier reef of Palau (Fig. 1), and the lagoon during flood and ebb tides during the day for 35 d between 10 July and 14 August 2000, collecting 132 samples. We used moored, metered, 0.305 mm neuston nets, with 0.305 mm mesh plankton sock cod ends, to fish at the surface in cross reef tidal currents 1 km south of Malakal Lighthouse, at the entrance to Malakal Harbor (Fig. 2). Paired nets were moored 10 m apart, seaward of the forereef and in the lagoon. Permanent moorings marked by a surface float were used to station the nets. Forereef nets were placed 80 m seaward of the reef crest over a 15 m bottom near deep water (Fig. 3). Lagoon nets were in 1 m of water at low tide above a sandy bottom 5 m from the backreef. Floats held the top of the 65 × 70 cm (0.455 m²) nets slightly above the surface, with the lower edge 65 cm below the surface. General Oceanic flow meters were suspended in each net 30 cm below the surface. Tidal currents fished the nets. The tide in Palau is semidiurnal, with a maximum amplitude of 1.0 to 2.0 m at neap and spring tides. As a temporary tide gauge at our study site matched tide recordings in Malakal Harbor 5 km away, we used the Malakal Harbor records in our study.

We sampled fish eggs and mesozooplankton during the first 2 h of ebb tide for 35 consecutive days, over an...
entire lunar period, from 10 July to 13 August 2000 (see Fig. 4). On 14 occasions we fished the first 2 h of ebb tide (when most fish spawned), then removed the nets and recorded flow meter data, washed net plankton into plankton socks, preserved the socks and zooplankton in 2% formalin, attached clean socks, and fished the nets for a further 4 h. Handling time was 5 min. We did not fish nets for precisely 2, 4, or 6 h, so all flow meter data and wet weights were adjusted to 2, 4, or 6 h per linear meter of reef or presented as mg m\(^{-3}\) (see Table 1). We collected no plankton on 14 July because the fish shifted from spawning at dusk on 13 July (16:40 to 18:40 h) to dawn on 15 July (06:00 to 08:10 h). Data for July 14 (see Fig. 4) were interpolated.

Flow through all nets was measured using General Oceanic flow meters, with low-speed rotors calibrated over a known distance by hanging the flow meters 2 m below a boat at speeds similar to flow over the reef. Although initial calibration was acceptable, meters in nets were often fouled by flotsam. Less often, wave chop caused rotor spinning and/or introduced air into water-filled flow meters, producing meter error. We calculated percentage difference between paired meters in adjacent nets, normalized for a 2 h period on all days when both meters had readings above or below 5000 revolutions in 2 h. When both meters rotated rapidly (>5000 rev. 2 h\(^{-1}\)), their difference was 11.3% (n = 14); when rotating slowly (<5000 rev. 2 h\(^{-1}\)) the difference was 52.6% (n = 11), 5 times more variable. Although we used large, slow-flow rotors, these meters perform erratically at very slow speeds. Consequently, for paired nets, we always used the flow rate and zooplankton data (wet weight m\(^{-3}\)) from the net with the higher meter reading. Although we use flow meter data and wet weight m\(^{-3}\) throughout our analysis and discussion, we also emphasize the quantity of zooplankton collected per net per unit time (i.e. during the first 2 h or last 4 h of ebb tide, or in a 6 h tidal cycle).

An additional flow meter was moored next to a large aggregation of spawning fish every day, adjusted by a diver at 15 min intervals so that it was always 1 m below the surface for the first 2 h of ebb tide. This diver also collected information every day for 35 d on species and abundance of fishes in the spawning aggregation (P. L. Colin unpubl.). The reef-top flow meter was moored taut between the reef and a large subsurface float, it rotated smoothly, independent of winds or waves, and it regularly recorded ebb flows 2 to 5 times faster than did flow meters in offshore nets (see Fig. 4A).

Vertical distribution of fish eggs on ebb tide was determined from 2 sets of horizontally stratified plankton samples collected 1 h after high tide by swimming a plankton net at the surface and at 3 m depth intervals over a 20 m transect, across and back between the

Fig. 2. Location of study transect, (A–B) on Lighthouse Reef, Palau. *: locations where nets were moored

Fig. 3. Diagram of forereef with location of moored zooplankton nets during flood and ebb tides (solid line; upper and lower panel, respectively). Note planktivorous fishes (‘wall-of-mouths’) present only on flood tides
2 seaward mooring lines. Fish eggs are buoyant, so the deepest samples (at 10 m) were collected first to reduce net contamination. After each sample had been collected, we shook the net vigorously from front to back to move plankton into the sock, placed the sock and zooplankton into a labeled jar, attached a clean plankton sock to the sleeve at the cod end, and swam the net at the next shallower depth. Most fish eggs (90%) were in the top 4 m; 60% of these were in the top meter. Thus, our moored, floating plankton nets undersampled fish eggs (and probably other mesozooplankton also) by about 40%. Stratified samples were collected on a calm day without waves and minimal vertical mixing of the water column. On windy days with surface waves, fish eggs and zooplankton flowing off the reef would mix deeper, with greater undersampling.

Zooplankton samples and collection socks were preserved in 2% formalin in the field. plankton socks were washed later, and non-zooplankton (Sargassum, eel grass, large algal fragments, coral sand, and lagoon debris) was removed prior to weighing. Wet samples were weighed to the nearest 0.1 g after draining for 2 min. Sample weights were converted to weight per cubic meter (moored nets sampled 0.455 m²). Samples were mixed with 500 ml of water and stirred for 2 min with a magnetic stirrer, then zooplankton in a 1 ml sub-sample were counted. We used data from 15 consecutive days (16 to 30 July) to evaluate counting errors for eggs, which were 3.0% for spherical and 2.2% for spindle-shaped eggs.

For statistical analysis of mesozooplankton assemblages, we evaluated samples for 5 combinations of site location and tidal stage. These were designated as Ocean Ebb-2 (n = 31), collected during the first 2 h of ebb tide from the ocean side of the forereef, and Ocean Ebb-4 (n = 15), for samples from the same location but during the last 4 h of ebb tide. We analyzed zooplankton for the entire 6 h of flood tide from the ocean site (Ocean Flood, n = 9), for the 6 h of ebb tide from the lagoon site (Lagoon Ebb, n = 9), and also from the lagoon during the entire flood tide (Lagoon Flood, n = 6). Zooplankton were subsampled, categorized initially into 25 taxonomic groups, with approximately 200 individuals of the most common taxa counted, although only 13 of the 25 taxa were sufficiently abundant for further statistical consideration (see Table 2). We elected to evaluate statistical properties via routine SPSS 11.0 procedures rather than more sophisticated approaches (Carleton et al. 2001, Legendre & Gallagher 2001) because we collected our samples according to a balanced, replicated sampling design and because we intentionally eliminated rare taxa from statistical analysis (Legendre & Gallagher 2001). Eliminating rare spe-

RESULTS AND DISCUSSION

In Palau, water moves between the lagoon and the ocean both as channel-flow through reef passes such as Malakal Channel (Fig. 2) and as sheet-flow across barrier reefs at all but lowest tides; our reef transect was emergent during spring low tide. We sampled zooplankton with 2 pairs of neuston nets that fished on rising and falling tides on both outer and inner margins of the barrier reef, documenting zooplankton that arrived at the reef, that survived cross-reef transport, and that left the reef (Fig. 3). On rising tides, seaward nets captured zooplankton coming from ocean to reef, seaward of planktivorous fishes, while lagoon nets captured materials that flowed over the reef from ocean to lagoon. On falling tides, lagoon nets captured zooplankton from the lagoon, while seaward nets captured plankton that survived the passage across the reef and fish eggs that were spawned at the forereef. There is a ‘wall of mouths’ of planktivorous fishes along the ocean side of the barrier reef in Palau on flood tides, but not on ebb tides (Fig. 3). Planktivorous fishes never occurred along the back reef.
Import of zooplankton from the sea on flood tides

During flood tide, ocean water moved slowly toward the reef (1.4 ± 1.1 cm s⁻¹), with 299.5 ± 263.9 mg m⁻³ mesozooplankton (Ocean Flood; Table 1, Fig. 4B). During a 6 h flood period, 71.7 ± 63.0 g zooplankton were imported onto each linear meter of forereef. Ocean Flood samples (Tables 1 & 2) had large copepods (29.9%), appendicularians (29.0%), chaetognaths (26.5%), and developing fish eggs (both round and spindle-shaped; 6.6%), typical of tropical oceanic water seaward of diurnal planktivorous fishes (Emery 1968, 1973, Hobson & Chess 1978, Hamner et al. 1988). There was no reef debris (filamentous algae, fish feces, sand) in water reaching the reef from the sea.

Flood tide waters passed through a ‘wall of mouths’ of planktivorous fishes at the forereef (Emery 1973, Hamner et al. 1988), then flowed over the reef crest, across the broad reef flat, and into the lagoon. Biomass m⁻³ of zooplankton flowing into the lagoon was 7 times lower than biomass seaward of the ‘wall of mouths’ (40.7 ± 21.5 mg m⁻³; Lagoon Flood, Table 1). Flood tide current in the lagoon after crossing the reef was faster than offshore (6.2 ± 3.7 cm s⁻¹ vs. 1.4 ± 1.1 cm s⁻¹). Lagoon nets collected 43.1 ± 19.9 g zooplankton per linear meter of reef during 6 h of flood tide, half the amount imported onto the forereef front in the same time (Table 1). A large percentage of the biomass entering the lagoon was reef debris (algae and sediment; Alongi 1997, Crossman et al. 2001, Yahel et al. 2002). Further, the composition of living zooplankton biomass entering the lagoon was different from that captured seaward of the reef, consisting of small copepods (66.4%), small chaetognaths (11.2%), small appendicularians (9.4%), zoea (3.2%), fish eggs (2.3%), and foraminifera (1.8%) (Table 2). Copepods captured in Lagoon Flood nets were significantly smaller than copepods captured in Ocean Flood nets (Table 1), presumably because of selective feeding on large copepods by planktivorous fishes at the forereef (Emery 1968, 1973, Hobson & Chess 1978, Hamner et al. 1988, Heidelberg et al. 2004).

Table 1. Zooplankton imported onto and exported from reef in 6 h on flood and ebb tides (Ocean Ebb-2: first 2 h, ebb tide; Ocean Ebb-4: last 4 h, ebb tide; n: number of samples). Wet weight in g: mean wet weight per linear meter of reef; wet weight in mg m⁻³: wet weight per m³ of water. Values are means ± SD. Dominant plankton collected at each location in **bold type**

<table>
<thead>
<tr>
<th></th>
<th>n</th>
<th>Wet wt (g m⁻³ reef)</th>
<th>Water speed (cm s⁻¹)</th>
<th>Wet wt (mg m⁻³)</th>
<th>Plankton collected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ocean Ebb-2</td>
<td>31</td>
<td>75.0 ± 44.8</td>
<td>4.9 ± 2.7</td>
<td>277.3 ± 177.4</td>
<td><strong>Fish eggs</strong>, zoea</td>
</tr>
<tr>
<td>Ocean Ebb-4</td>
<td>15</td>
<td>20.8 ± 5.7</td>
<td>3.8 ± 1.8</td>
<td>36.3 ± 16.4</td>
<td><strong>Fish eggs</strong>, foraminifera, copepods, zoea</td>
</tr>
<tr>
<td>Ocean Flood</td>
<td>9</td>
<td>71.7 ± 63.0</td>
<td>1.4 ± 1.1</td>
<td>299.5 ± 263.9</td>
<td><strong>Large copepods</strong>, <strong>appendicularians</strong>, chaetognaths, fish eggs</td>
</tr>
<tr>
<td>Lagoon Flood</td>
<td>6</td>
<td>43.1 ± 19.9</td>
<td>6.2 ± 3.7</td>
<td>40.7 ± 21.5</td>
<td><strong>Small copepods</strong>, small chaetognaths, larvaceans, zoea</td>
</tr>
<tr>
<td>Lagoon Ebb</td>
<td>9</td>
<td>158.4 ± 85.6</td>
<td>14.2 ± 6.3</td>
<td>51.9 ± 23.5</td>
<td><strong>Small copepods</strong>, zoea, crustacean larvae, chaetognaths</td>
</tr>
</tbody>
</table>

aBody lengths were 1.13 ± 0.19 mm (n = 9)
bBody lengths were 0.52 ± 0.19 mm (n = 6)
During ebb tide, lagoon water flowed onto and across the reef (14.2 ± 6.3 cm s⁻¹), transporting 51.9 ± 23.5 mg zooplankton m⁻³. During the 6 h ebb period, the biomass of zooplankton transported from lagoon to barrier reef was 158.4 ± 85.6 g m⁻¹ (Lagoon Ebb, Table 1). Lagoon water (Lagoon Ebb, Tables 1 & 2) contained small copepods (24.1%), zoea (21.4%), crustacean larvae (20.1%), chaetognaths (15.5%), fish eggs (10.1%), and appendicularians (6.3%), typical of coral reef lagoons (Emery 1968, Tranter & George 1972, Hobson & Chess 1978, 1979). Most fish eggs from the lagoon had developed embryos, were unfertilized (opaque), or were decomposing (fungal growths). The wet weights of lagoon mesozooplankton were highest at new and full moon, due to zoea and larval crustaceans.

At the seaward side of the reef, the biomass of zooplankton flowing off the reef into the ocean over an entire 6 h ebb tide was 95.8 ± 35.4 g m⁻¹. During the first 2 h of ebb tide (Ocean Ebb-2), zooplankton in seaward nets included 60% copepods, zoea, and crustacean larvae, very few of these were captured in seaward nets (except zoea, twice); lagoon zooplankton were either captured by reef planktivores or settled in transit.

Export of zooplankton from the lagoon and barrier reef on ebb tides

During ebb tide, lagoon water flowed onto and across the reef (14.2 ± 6.3 cm s⁻¹), transporting 51.9 ± 23.5 mg zooplankton m⁻³. During the 6 h ebb period, the biomass of zooplankton transported from lagoon to barrier reef was 158.4 ± 85.6 g m⁻¹ (Lagoon Ebb, Table 1). Lagoon water (Lagoon Ebb, Tables 1 & 2) contained small copepods (24.1%), zoea (21.4%), crustacean larvae (20.1%), chaetognaths (15.5%), fish eggs (10.1%), and appendicularians (6.3%), typical of coral reef lagoons (Emery 1968, Tranter & George 1972, Hobson & Chess 1978, 1979). Most fish eggs from the lagoon had developed embryos, were unfertilized (opaque), or were decomposing (fungal growths). The wet weights of lagoon mesozooplankton were highest at new and full moon, due to zoea and larval crustaceans.

At the seaward side of the reef, the biomass of zooplankton flowing off the reef into the ocean over an entire 6 h ebb tide was 95.8 ± 35.4 g m⁻¹. During the first 2 h of ebb tide (Ocean Ebb-2), zooplankton from reef to sea was primarily composed of eggs spawned by herbivorous fish at the forereef (75.0 ± 44.8 g m⁻¹; 89% eggs; Table 2). During the last 4 h of ebb tide (Ocean Ebb-4), less zooplankton was captured (20.8 ± 5.7 g m⁻¹), with fewer fish eggs (Tables 1 & 2), and, although flow rates did not decrease significantly, biomass per cubic meter was much lower. Most plankton, therefore, was exported seaward during the first 2 h of ebb tide in the form of meroplanktonic fish eggs. Although lagoon waters that flowed onto the back reef during ebb tide contained 60% copepods, zoea, and crustacean larvae, very few of these were captured in seaward nets (except zoea, twice); lagoon zooplankton were either captured by reef planktivores or settled in transit.

Export of zooplankton from the lagoon and barrier reef on ebb tides

During ebb tide, lagoon water flowed onto and across the reef (14.2 ± 6.3 cm s⁻¹), transporting 51.9 ± 23.5 mg zooplankton m⁻³. During the 6 h ebb period, the biomass of zooplankton transported from lagoon to barrier reef was 158.4 ± 85.6 g m⁻¹ (Lagoon Ebb, Table 1). Lagoon water (Lagoon Ebb, Tables 1 & 2) contained small copepods (24.1%), zoea (21.4%), crustacean larvae (20.1%), chaetognaths (15.5%), fish eggs (10.1%), and appendicularians (6.3%), typical of coral reef lagoons (Emery 1968, Tranter & George 1972, Hobson & Chess 1978, 1979). Most fish eggs from the lagoon had developed embryos, were unfertilized (opaque), or were decomposing (fungal growths). The wet weights of lagoon mesozooplankton were highest at new and full moon, due to zoea and larval crustaceans.

At the seaward side of the reef, the biomass of zooplankton flowing off the reef into the ocean over an entire 6 h ebb tide was 95.8 ± 35.4 g m⁻¹. During the first 2 h of ebb tide (Ocean Ebb-2), zooplankton from reef to sea was primarily composed of eggs spawned by herbivorous fish at the forereef (75.0 ± 44.8 g m⁻¹; 89% eggs; Table 2). During the last 4 h of ebb tide (Ocean Ebb-4), less zooplankton was captured (20.8 ± 5.7 g m⁻¹), with fewer fish eggs (Tables 1 & 2), and, although flow rates did not decrease significantly, biomass per cubic meter was much lower. Most plankton, therefore, was exported seaward during the first 2 h of ebb tide in the form of meroplanktonic fish eggs. Although lagoon waters that flowed onto the back reef during ebb tide contained 60% copepods, zoea, and crustacean larvae, very few of these were captured in seaward nets (except zoea, twice); lagoon zooplankton were either captured by reef planktivores or settled in transit.

Export of zooplankton from the lagoon and barrier reef on ebb tides

During ebb tide, lagoon water flowed onto and across the reef (14.2 ± 6.3 cm s⁻¹), transporting 51.9 ± 23.5 mg zooplankton m⁻³. During the 6 h ebb period, the biomass of zooplankton transported from lagoon to barrier reef was 158.4 ± 85.6 g m⁻¹ (Lagoon Ebb, Table 1). Lagoon water (Lagoon Ebb, Tables 1 & 2) contained small copepods (24.1%), zoea (21.4%), crustacean larvae (20.1%), chaetognaths (15.5%), fish eggs (10.1%), and appendicularians (6.3%), typical of coral reef lagoons (Emery 1968, Tranter & George 1972, Hobson & Chess 1978, 1979). Most fish eggs from the lagoon had developed embryos, were unfertilized (opaque), or were decomposing (fungal growths). The wet weights of lagoon mesozooplankton were highest at new and full moon, due to zoea and larval crustaceans.

At the seaward side of the reef, the biomass of zooplankton flowing off the reef into the ocean over an entire 6 h ebb tide was 95.8 ± 35.4 g m⁻¹. During the first 2 h of ebb tide (Ocean Ebb-2), zooplankton from reef to sea was primarily composed of eggs spawned by herbivorous fish at the forereef (75.0 ± 44.8 g m⁻¹; 89% eggs; Table 2). During the last 4 h of ebb tide (Ocean Ebb-4), less zooplankton was captured (20.8 ± 5.7 g m⁻¹), with fewer fish eggs (Tables 1 & 2), and, although flow rates did not decrease significantly, biomass per cubic meter was much lower. Most plankton, therefore, was exported seaward during the first 2 h of ebb tide in the form of meroplanktonic fish eggs. Although lagoon waters that flowed onto the back reef during ebb tide contained 60% copepods, zoea, and crustacean larvae, very few of these were captured in seaward nets (except zoea, twice); lagoon zooplankton were either captured by reef planktivores or settled in transit.
Lagoon plankton, therefore, were imported into the reef economy, whereas eggs spawned at the forereef by herbivorous fish during the first 2 h of ebb tide were exported from the reef to the sea.

Our largest net sample of zooplankton was collected from 1 net offshore during the first 2 h of ebb tide on 28 July. It collected >300,000 round surgeonfish eggs (236.0 g m⁻¹ in 2 h⁻¹; 600.5 mg m⁻³), but not a single spindle-shaped parrotfish egg or any other live zooplankton. On the first days of full and new moon, crab zoea contributed a substantial percentage to the ebb tide catch (e.g. Ocean Ebb-4, 15 July, zoea 43.7%).

The flux of water across the barrier reef for the first 2 h of ebb tides was plotted using the faster of the current meters in the paired plankton nets, with greatest flow after full moon and least during first quarter. We also measured water flow across the reef crest adjacent to a large, recurrent aggregation of spawning reef fish. Flow recorded atop the reef crest was faster than in the seaward nets, but both exhibited lunar periodicities in water velocity (Fig. 4A). Increased flow rates across the reef during spring tides suggested that more zooplankton would be transported across the reef during spring tides, so we plotted biomass of exported zooplankton during the first 2 h of ebb tide for the entire lunar cycle, with wet weights normalized to 120 min (Fig. 4B). We did not find a statistically significant lunar periodicity for zooplankton biomass because of biomass variability of up to 5-fold between successive days (12 to 13 July; 28 to 29 July). Ebb tide abundance of surgeonfish and parrotfish eggs was plotted separately (Fig. 4C); both groups of fishes exhibited highly significant lunar spawning rhythms. Surgeonfish spawned near high-water spring tides in late afternoon, with peak spawning on 12 July at 17:32 h, on 28 July at 17:39 h, and on 10 August at 17:00 h, 4 or 5 d prior to new and full moon. Parrotfish spawned in the morning after high tide, but also throughout the day, primarily during neap tides (Colin & Bell 1991, Domeier & Colin 1997).

During the last 4 h of ebb tide, wet weight of zooplankton exported from the reef was much lower than during the first 2 h (Table 1). The taxonomic composition of the catch during the last 4 h of ebb tide was still dominated by fish eggs (58.5%), although some eggs collected during the last 4 h of ebb tide had advanced embryos or were coated with fungus. These eggs were not recently spawned, but apparently had advected across the reef flat from the lagoon. When we compared the first 2 h and last 4 h of ebb tide, we found that 98.5% of the spindle-shaped parrotfish eggs and 77.9% of the round surgeonfish eggs were captured during the first 2 h of ebb tide. During the last 4 h of ebb tide, benthic foraminifera were the second most important category collected, followed by zoea and copepods (Table 2). Crab zoea and larval crustaceans were important components of late ebb tide samples,
but were abundant only following full and new moon. As the tide ebbs, waves resuspend more reef material because the waves are closer to the reef’s surface. All late ebb tide samples contained reef debris, crustacean exoskeletons, copepod fecal pellets, and algae.

Dendrograms of samples and station relationships (Fig. 5) show that zooplankton collected during ebb tide on the ocean side of the barrier reef (Ocean Ebb-2, Ocean Ebb-4) were similar to each other; this is not surprising since these 2 groups were temporal subdivisions of the same location and tidal phase. A multidimensional scaling plot (Fig. 6) also shows these similarities, with all Ocean Ebb samples clustered together. However, samples in the first 2 h differed significantly from those in the last 4 h, because the latter contained fewer fish eggs and because lagoon waters had crossed the reef by the last 4 h of ebb tide, carrying some lagoon plankton, zoea, and foraminifera. The flood tide samples are somewhat intermixed in the sample dendrogram (Ocean Flood, Lagoon Flood; Fig. 5B) because both sample locations were dominated by copepods, appendicularians, and chaetognaths on flood tides. However, cluster analysis (Fig. 6) shows these 2 sets of samples to be distinctive. Ocean Flood samples collected seaward of the wall of planktivorous fishes contained large copepods, appendicularians, chaetognaths, doliolids, and siphonophores typical of oceanic water (Emery 1968, Hobson & Chess 1978, Hamner et al. 1988, Sorokin 1990). Importantly, there were also substantial numbers of eggs (6.5%) identical in size and shape to those spawned on ebb tides by the reef fishes at our study site. These eggs, however, were not recently spawned, but had visible embryos, spawned roughly 12 h earlier, apparently retained in coastal boundary waters and returned to the reef. Lagoon Ebb samples (LE; Fig. 5) were taxonomically diverse and evenly represented, with small copepods (30%), crab zoea (24%), chaetognaths (18%), shrimp larvae (14%), and fish eggs (10%). Crab zoea and larvae were only abundant in Lagoon Ebb waters, so they must have been produced by lagoon crabs. Appendicularians were abundant (6 to 11%) in 5 of 9 Lagoon Ebb samples.

Drifters and returning eggs

We released drogues amid the clouds of fish eggs at the forereef (Fig. 7), tracking drogues as proxies of fish eggs, because it is difficult to track recently spawned eggs and/or larvae at sea (but see Willis & Oliver 1990). Drogues drifted seaward only 2 to 3 km during ebb tide, and all but 1 drogue reversed or changed direction when the tide began to flood. Oscillatory movements of drogues defined a tidal, neritic, water mass that remained near the barrier reef. This water mass was separated from the equatorial currents around the archipelago by fronts and small-scale and meso-scale eddies visible from small boats and from airplanes on calm days. The nature of entrained, tidal waters has not been well investigated (Kingsford et al. 1991, Wolanski & Spagnol 2000, Zeidberg & Hamner 2002), but fronts between tidal waters and the surrounding oceanic currents create hydrographic barriers around oceanic islands that permit neritic waters to act as nurseries for fish and invertebrates that have pelagic developmental stages in their early life history (Leis 1982, Willis & Oliver 1990, Kingsford et al. 1991).

Many of our surface drogues released at the reef front during ebb tide returned to the vicinity of the reef and/or reentered the lagoon 12 h later on the flood tide (Fig. 7). Some fish eggs spawned on the previous high tide also returned (6.6%; Ocean Flood, Table 2). It is important to clarify that these returning eggs were indeed eggs of the fish that had spawned some 12 h earlier at or close to our study site. All plankton was identified and counted using a microscope with sufficient resolution to discern quite subtle differences in egg shape, size, color, and location of oil droplets and embryos for different species of fish. Most of these imported eggs would be eaten during the day at the forereef (see fish gut data in Hamner et al. 1988), but some eggs crossed the reef and entered the lagoon (eggs 2.3%; Lagoon Flood, Table 2). Eggs spawned during the day would usually return to the reef 10 to 12 h later at night, when eggs would be hard to see. Most returning eggs were healthy, with early-stage developing embryos. The return of drogues and recently spawned fish eggs to their natal reef supports the hypothesis that neritic waters, probably entrained by topographic fronts and eddies, enhance self-recruitment (Willis & Oliver 1990, Kingsford et al. 1991, Jones et al. 1999, Swearer et al. 1999).

Cross-reef flow and zooplankton dynamics

Flow across coral reefs is controlled by reef height and morphology, wind, wave-overtopping, tides, or a combination of these (Atkinson et al. 1981, Pickard 1986, Hamner & Wolanski 1988, Wolanski & Hamner 1988). On the ocean side of the barrier reef, at or near low tide, the water column has no density gradient to depths of at least 15 to 20 m. As rising tide drives flow onto the barrier reef, the entire water column (10 m or more in depth) feeds the shallow reef flat. As water moves onto the reef front, current speed increases compared to speed of currents in deeper water offshore (Roberts & Suhayda 1983, Hamner et al. 1988). Flood tide flow offshore of Lighthouse Reef was only...
1.4 cm s\(^{-1}\), considerably less than where the reef was shallower and flow averaged 6.2 cm s\(^{-1}\).

Ebb tide flow dynamics differ because lagoon water is warmer and/or less saline than ocean water. When lagoon water flowed onto the reef, it was vertically compressed and moved rapidly (14.2 cm s\(^{-1}\)). At the forereef the less dense lagoon water flowed out to sea in a 2 to 3 m surface layer. The ebb tide speed of this shallow layer seaward of the forereef was faster than the current at the same spot during rising tides (Ocean Ebb-2, 4.9 cm s\(^{-1}\); Ocean Ebb-4, 3.8 cm s\(^{-1}\); Ocean Flood, 1.4 cm s\(^{-1}\); Table 1).

It is also clear that the assemblage of plankton in a given body of water changes qualitatively and quantitatively, gradually or abruptly, as it flows across the reef. When oceanic or neritic water flows onto the forereef during the day, schools of planktivorous fish abruptly remove larger zooplankton (Emery 1968, 1973, Hamner et al. 1988). Thereafter, the water flows over the reef crest, where breaking waves break loose filamentous algae and resuspend sediments from the reef surface, abruptly adding new particulate materials to the reefwater column. This water then flows across the reef flat, where grazing fishes gradually resuspend even more materials (Alongi 1997, Crossman et al. 2001, Yahel et al. 2002). This water finally flows into the lagoon, where it quickly mixes with lagoon water. During ebb tide, mesozooplankton composition in lagoon water similarly changes when it flows across the reef, gradually depleted of prey by planktivores and filter feeders, then abruptly enriched at the forereef during the day by a massive infusion of fish eggs. Lastly, the water flowing off the reef into the sea is buoyant (warmer or less saline) and the original parcel of lagoon water, now enriched by fish eggs and zoea, advects offshore as a floating lens rich in neuston that becomes concentrated into buoyant, convergent fronts and slicks formed by tides and topographic eddies (Wolanski & Hamner 1988, Willis & Oliver 1990, Kingsford et al. 1991, Wolanski & Spagnol 2000, Parr 2001, Wolanski et al. 2003).

The dynamics of cross-reef flow therefore markedly affect the quantity of food potentially available to coral reef organisms, and it is clear that there is far more plankton available to coral reefs than implied by the older literature (Odum & Odum 1955, Johannes et al. 1972, Johannes & Gerber 1974). For example, our offshore zooplankton biomass measurements in Palau were approximately 300 mg m\(^{-3}\), similar to published data on standing stock near coral reefs (Tranter & George 1972, Sammarco & Crenshaw 1984, Hamner et al. 1988, Sorokin 1990). These relatively robust estimates for offshore mesozooplankton biomass, however, must be multiplied by volume flow across the reef, which is often enormous. For example, at Enewetak, Johannes & Gerber (1974) estimated that 18 100 to 33 000 m\(^{3}\) of water (devoid of zooplankton by the time it reached their nets atop the reef) flowed across each linear meter of forereef at their transect location every 12 h. Pickard (1986) calculated that 6000 m\(^{3}\) flowed across each meter of forereef at Davies Reef in 12 h, producing a flux of mesozooplankton of 400 g 12 h\(^{-1}\) m\(^{-2}\) (Hamner et al. 1988). In Palau, flood tides transported 72 g 6 h\(^{-1}\) m\(^{-2}\) and ebb tides 158 g 6 h\(^{-1}\) m\(^{-2}\), a total of 230 g 12 h\(^{-1}\) m\(^{-2}\). Lighthouse Reef and Davies Reef are enriched by similar quantities of imported zooplankton despite hydrographic differences, but Lighthouse Reef also exported 95.8 g 6 h\(^{-1}\) m\(^{-2}\) zooplankton, primarily fish eggs, on ebb tide into the sea.

Coral reefs were originally considered conservative recyclers of materials (Odum 1953, Odum & Odum 1955), but vast amounts of fish eggs are exported from Palau reefs every day, all year long (P. L. Colin pers. obs.). Although a seemingly flagrant waste of materials valuable to reef organisms, exporting buoyant eggs from reefs full of egg predators is advantageous to the fish (Colin & Bell 1991, Domeier & Colin 1997, Craig 1998). Neritic waters apparently retain the eggs long enough for larvae to become behaviorally competent and remain near natal reefs prior to settlement (Leis 1982, Kingsford et al. 1991, Hearn et al. 2000, Wolanski & Spagnol 2000, Parr 2001, Zeidberg & Hamner 2002). Thereafter, other factors limit settlement of fish larvae on the reef (Doherty et al. 2004). Clearly, export of materials and energy from the forereef and residence times of larval fish in neritic waters must be reexamined (Hatcher 1997).

Acknowledgements. We thank John Carleton for his advice on statistical procedures, Amatzia Genin for reading a very rough early draft, and 9 anonymous reviewers for helpful suggestions and almost never conflicting advice.

LITERATURE CITED
