The real Nemo movie: Description of embryonic development in *Amphiprion ocellaris* from first division to hatching

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**Abstract**

**Background:** *Amphiprion ocellaris* is one of the rare reef fish species that can be reared in aquaria. It is increasingly used as a model species for Eco-Evo-Devo. Therefore, it is important to have an embryonic development table based on high quality images that will allow for standardized sampling by the scientific community.

**Results:** Here we provide high-resolution time-lapse videos to accompany a detailed description of embryonic development in *A. ocellaris*. We describe a series of developmental stages and we define six broad periods of embryogenesis: zygote, cleavage, blastula, gastrula, segmentation, and organogenesis that we further subdivide into 32 stages. These periods highlight the changing spectrum of major developmental processes that occur during embryonic development.

**Conclusions:** We provide an easy system for the determination of embryonic stages, enabling the development of *A. ocellaris* as a coral reef fish model species. This work will facilitate evolutionary development studies, in particular studies of the relationship between climate change and developmental trajectories in the context of coral reefs. Thanks to its lifestyle, complex behavior, and ecology, *A. ocellaris* will undoubtedly become a very attractive model in a wide range of biological fields.

**KEYWORDS**
coral reef fish, developmental stages, embryogenesis, pigmentation, Pomacentridae, time-lapse video

**Introduction**

Developmental biology is built around model organisms such as the mouse *Mus musculus*, the chicken *Gallus gallus*, the frog *Xenopus laevis*, the zebrafish *Danio rerio*, the...
fruit fly *Drosophila melanogaster*, and the round worm *Caenorhabditis elegans*. However, such model organisms, sometimes called “supermodels,” only represent a very small and biased fraction of living biodiversity and can therefore limit our understanding of the diversity of biological processes. There is a widely recognized need for additional model species that better represent the diversity of the animal kingdom. Recently, the development of tools such as massive sequencing and CRISPR/Cas9 have rendered this possible.

Diversity is particularly lacking in the teleost fishes, for which zebrafish, and to a lesser extent medaka (*Oryzias latipes*), have emerged as invaluable model species, complete with elaborate technological tools and a vivid scientific community. However, these two species do not represent the diversity of the ca. 30,000 species of teleost fishes. Several additional models have emerged in recent years (1990-2010) such as the stickleback, *Gasterosteus aculeatus*, the cichlids (eg, the tilapia, *Oreochromis niloticus*), the poecilid fishes (eg, guppies and platys), the annual killifishes, and the *Astyanax mexicanus* cavefish. These models, whose embryonic development has been well described, have shed new light on a number of relevant biological questions. But they suffer from a limit: they are mostly freshwater fishes and capture only a fraction of teleost fishes diversity. The majority of teleosts are marine, but, mostly because of their biphasic life cycle (with extremely numerous small eggs and tiny pelagic larvae that will metamorphose into juveniles), they are notoriously difficult to raise in captivity at the laboratory scale. To date, only a few studies have examined embryogenesis in marine fishes, including flatfish, sea bream, sea bass, mahi-mahi, and grouper (see eg, 15, 16), limiting our ability to address a wide range of biological questions in this group.

In our lab, we are working to develop the false clownfish *Amphiprion ocellaris* as a marine teleost fishes model system. Living in symbiotic association with giant sea anemones, this species is native to the Indo-West Pacific region from the Ryukyu Islands in Japan to northwestern Australia. Like most coral reef fishes, *A. ocellaris* have a biphasic life cycle with a dispersive oceanic phase and a more sedentary reef phase (Figure 1). In contrast to many coral reef fishes that spawn in the open ocean, false clownfishes are benthic spawners.

*A. ocellaris* are an ideal model for studies of embryogenesis because breeding pairs spawn regularly, laying 100 to 300 eggs every 2 weeks, and because the chorion is transparent until hatching, allowing for visualization of ongoing embryonic development. Classical techniques of developmental biology, including in situ hybridization, have been successfully performed on false clownfish embryos. This fish is therefore a new model species with a high potential for further study of...
evolutionary and developmental processes in a unique ecological framework. Other anemone fish species, and in particular the sister species *Amphiprion percula*, have also been used as model species although to our knowledge *A. ocellaris* is the most widely used at present.

*A. ocellaris* belongs to a monophyletic group of 30 species, the anemonefishes (subfamily Amphiprioninae) within the family Pomacentridae (Teleostei; Perciformes; Pomacentridae). These fish provide an excellent example of rapid adaptation, fast speciation, and parallel evolution. The symbiotic association with sea anemones (considered to be an obligate mutualistic relationship) is thought to have been the key innovation that allowed anemonefishes to radiate rapidly in untapped ecological niches. As expected under the ecological theory of adaptive radiation, this increased diversification, rates of morphological evolution, and rapid and convergent morphological changes correlated with the different ecological niches offered by different host anemones. Thus, anemonefishes allow us to integrate studies of ecology, genomics, and development (eco-evo-devo) to understand adaptive radiation and phenotypic divergence.

Establishing a precise and well-illustrated embryonic developmental table for this species would therefore be of particular interest. Developmental staging is essential because heterogeneity among individuals means that age does not perfectly correlate with development. Staging tables are available for most model organisms (eg, zebrafish). Here, we provide high resolution pictures and time lapse videos of false clownfish embryonic development and describe with high accuracy developmental stages that can be used by the scientific community. We group the 32 developmental stages into six larger time-blocks that we call periods: zygote, cleavage, blastula, gastrula, segmentation, and organogenesis (Figure 2). Table 1 summarizes brief descriptions of the main morphological changes occurring at each stage. Figure 2 shows overall development and highlights the six periods. It also shows the corresponding available time-lapse movies.

### 2.2 Zygotic period

In our facility, the female lays eggs on a terracotta pot. Once the female has spawned, she leaves the nest, and the male fertilizes the eggs. Newly fertilized eggs of *A. ocellaris* have an ovoid shape, with the longitudinal axis (1.707 ± 0.037 mm) longer than the transverse axis (0.872 ± 0.019 mm) (Figure 3A). The egg is surrounded by a translucent envelope called the chorion. The fertilized cell is located at the end of the egg closest to the adhesive edge of the chorion (Figures 3A and A'). False clownfish eggs are telolecithal with the yolk concentrated at the vegetal pole. The yolk is composed of large dark-yellow yolk globules/platelets, giving it a grainy appearance. After spawning, the chorion sticks to the terracotta pot through a mucous secretion (Figure 3A, A').

**One-cell stage (0 hpf):** Fertilization induces an increase in the volume of the blastodisc which replaces the yolk (Figure 3A, A', B, B'). The blastodisc gradually segregates from the yolk and forms a more prominent cell (Figure 3B, B', Movie 1).

### 2.3 Cleavage period

We define the cleavage period as consisting of six stages: 2-, 4-, 8-, 16-, 32-, and 64-cell stages (Figures 2 and 3). Cleavages are synchronous and occur every 30 min at 26°C. As in most teleost fishes, they are meroblastic
and discoidal (i.e., that they only occur in the blastodisc which keep a connection with the yolk). Early cleavage divisions follow a highly reproducible pattern of meridional and equatorial planes (Figure 3I).

**Two-cell stage (0.5 hpf):** The first cleavage furrow, ending the first zygotic cell cycle, is vertically oriented (Figure 3I) dividing the blastodisc into two cells (blastomeres) of equal size. This pattern is seen in other teleost species until the 32-cell stage.11-14,32 Both cells stay connected to the underlying yolk (meroblastic cleavage) (Figure 3C and C’, Movie 1).

**Four-cell stage (1 hpf):** In the second division, the cleavage plane is oriented perpendicular to the first one, resulting in four blastomeres arranged in a 2 X 2 array when viewed from the animal pole (Figure 3D and D’, I, Movie 1).

**Eight-cell stage (1.5 hpf):** The third set of cleavages occurs in two planes parallel to the first cleavage plane, dividing the four blastomeres into eight blastomeres. They are arranged in a 2 X 4 array (Figure 3E, E’, I, Movie 1).

**16-cell stage (2 hpf):** The fourth cleavage also occurs along two planes, this time parallel to the second cleavage plane. The two rows of four blastomeres are divided into four rows of four blastomeres (4 X 4 array) (Figure 3F, F’, I, Movie 1).

**32-cell stage (2.5 hpf):** The fifth set of cleavages generates a 4 X 8 array of cells, although the pattern is less stereotypic than in previous stages. All cells are still in contact with the yolk, shaping the underlying yolk into a dome-like structure (Figure 3G, G’, Movie 1).

**64-cell stage (3 hpf):** During the sixth set of divisions, cells start to be cleaved completely from the others, forming a second layer of cells on top of those that are still connected to the yolk (marginal cells). Unlike in previous stages, there are no regularly-patterned cleavage planes or stereotypical cell arrangements (Figure 3H, H’, Movie 1).
2.4 | Blastula period

The blastula period extends from the seventh set of cleavages until gastrulation. Cleavages occur with increasing irregularity. The blastodisc acquires a more uniform appearance, and starts to thin and spread around the yolk (epiboly). When 30% of the yolk is covered by the blastodisc, gastrulation begins (Figure 4- Movie 1).

| Stage                  | Figure | Movie | Hpf | Description                                                                 |
|------------------------|--------|-------|-----|------------------------------------------------------------------------------|
| **Zygotic period**     |        |       |     |                                                                              |
| Zygotic stage          | 3.A    | –     | 0.5 |                                                                              |
| 1-cell                 | 3.B    | 1     | 0   | The blastodisc consists of one cell                                          |
| **Cleavage period**    |        |       |     |                                                                              |
| 2-cell                 | 3.C    | 1     | 0.5 | 2 blastomeres                                                                |
| 4-cell                 | 3.D    | 1     | 1   | 2 X 2 array of blastomeres                                                   |
| 8-cell                 | 3.E    | 1     | 1.5 | 2 X 4 array of blastomeres                                                   |
| 16-cell                | 3.F    | 1     | 2   | 4 X 4 array of blastomeres                                                   |
| 32-cell                | 3.G    | 1     | 2.5 | 2 X 8 array of blastomeres                                                   |
| 64-cell                | 3.H    | 1     | 3   | Blastodisc consists of two layers of cells                                  |
| **Blastula period**    |        |       |     |                                                                              |
| seventh cleavage       | 4.A    | 1     | 3.5 | High mound of cells                                                          |
| eighth cleavage        | 4.B    | 1     | 4   | High mound of cells                                                          |
| High-cell stage        | 4.C    | 1     | 5.5 | High mound of cells                                                          |
| Sphere                 | 4.D    | 1     | 6.5 | Interface between the blastodisc and the yolk nearly flat, resulting in a hemispherical shape of the blastodisc |
| Dome                   | 4.E    | 1     | 9   | Shape of the blastula remains spherical; yolk bulging toward animal pole     |
| 30% epiboly            | 4.F    | 1     | 9.5 | Blastoderm is of uniform thickness and starts to cover the yolk              |
| **Gastrula period**    |        |       |     |                                                                              |
| Germ-ring              | 5.A    | 1     | 10.5| Germ-ring is visible                                                         |
| Shield                 | 5.B    | 1     | 12.5| Embryonic shield is visible                                                  |
| 50% epiboly            | 5.C    | 1–2   | 15  | Embryonic axis is formed                                                     |
| 75% epiboly            | 5.D    | 2     | 17.5| Head primordium, brain rudiment and tail bud are visible                     |
| 90% epiboly            | 5.E    | 2     | 19  | Formation of the yolk plug                                                  |
| 100% epiboly           | 5.F & 5.G | 2–3 | 21  | Kupffer's vesicle forms as well as 3 somite furrows                           |
| **Segmentation period**|        |       |     |                                                                              |
| 6-somite               | 6.A    | 3     | 23  | Rudiment of the brain subdivides- optic primordia is formed                  |
| 12-somite              | 6.B    | 3     | 27.5| Melanophore appear over the yolk                                             |
| 15-somite              | 6.C    | 3     | 29.5| Notochord, lens primordium, rudiment of cerebellum and ephiphysis are visible |
| 18-somite              | 6.D    | 3     | 32.5| The otic vesicle starts to form                                              |
| 22-somite              | 6.E    | 3     | 34.5| v-shaped trunk somites and twitching of the trunk muscles are observed       |
| **Organogenesis period**|    |       |     |                                                                              |
| 25%-OCV                | 7.B    | 4     | 44  | Orientation of embryo changes within the chorion. Pericardial cavity formed  |
| 40%-OCV                | 7.C    | 4     | 55  | Eyes start to show black pigmentation. The heart and medial fin fold form    |
| 80%-OCV                | 7.D    | 4     | 75  | Vascularization of the yolk, formation of the cardinal vein                  |
| 4 dpf                  | 7.E    | none  | 4 dpf| Orange xanthophores appear, pectoral fin and cloaca are visible              |
| 5 dpf                  | 7.F    | none  | 5 dpf| Heart moves anteriorly, the mouth opens, formation of opercules              |
| 6 dpf                  | 7.G    | none  | 6 dpf| Eyes are silver, lower jaw extends anteriorly and becomes vascularized       |
| 7 dpf                  | 7.H    | none  | 7 dpf| Gills are visible and the jaw becomes thicker                                |
High-cell stages (3.5 hpf to 5.5 hpf): Cleavages continue to occur; however, no clear cleavage planes can be identified. The seventh cleavage stage (3.5 hpf- Figures 4A and A'), eighth cleavage stage (4 hpf- Figures 4B and B') and high-cell stages (5.5 hpf- Figures 4C and C') result in blastomeres that become smaller, without a clear increase in the size of the blastodisc (Figures 4A-C, Movie 1).

Sphere-stage (6.5 hpf): After the high-cell stage, the interface between the lower part of the blastodisc and
the upper part of the yolk is flat or nearly flat, resulting in a hemispherical shape of the blastodisc (Figures 4D and D'). We can distinguish the sphere-stage from the high-cell stage by the limit between the upper part of the yolk and the blastodisc that is becoming conspicuous (white arrowhead, Figure 4D'-Movie 1), which seems to result from a uniform increase in pressure against the yolk.

*Dome-stage (9 hpf):* The flattening of the blastodisc continues (Figures 4E and E'), starting to cover the top of the yolk, which bulges towards the animal pole in a dome-like shape (black arrowhead, Figure 4E').

*30% epiboly stage (9.5 hpf):* The blastodisc, which gradually transforms into a uniformly thick layer, starts to cover the yolk and is now called the blastoderm. This stage can be measured by the percentage of epiboly, specifically the ratio of the distance from the animal pole to the blastoderm margin and the distance from the animal to the vegetal pole of the embryo (Figures 4F and F').

**2.5 Gastrula period**

Gastrulation is defined as the period during which cellular movements (involution, convergence and extension) of the blastoderm tend to form the embryonic axis and the organization of the germ cell layers: the epiblast that will give rise to a part of the ectoderm, the hypoblast that corresponds to the future mesoderm and endoderm. 32
When 30% epiboly is reached, the germ-ring forms and the embryonic shield is visible at the dorsal side of the blastoderm margin. Gastrulation starts at 30% epiboly in *A. ocellaris* and epiboly continues until the blastoderm completely covers the yolk, with a speed of epiboly of about 9.5% per hour. The gastrula period is differentially divided into 6 stages: Germ-ring stage, shield stage, 50%-epiboly, 75%-epiboly, 90%-epiboly, and 100%-epiboly (Figure 5-Movies 1 and 2).

**Germ-ring stage (10.5 hpf):** The percent-epiboly is almost the same as at the 30%-epiboly stage. The germ-ring can be observed as a stripe at the most vegetal pole-side of the blastoderm margin along the equatorial plane (Figures 5 A, A’- Movie 1).

**Shield stage (12.5 hpf):** A thickening appears at one position of the blastoderm margin (now defined as the dorsal side) (Figures 5B, B’-Movie 1). This thickening is referred to as the “embryonic shield” and is the result of cellular movements. Gastrulation and cell involution take place in this part of the blastoderm.

**50%-epiboly stage (15 hpf):** Epiboly moves the blastoderm margin to 50% of the distance between the animal and vegetal pole (Figures 5 C, C’-Movies 1 and 2). At this stage, the dorsal side of the blastoderm thickens further and the future embryonic axis becomes visible, with the anterior end in the direction of the animal pole.

**75%-Epiboly (17.5 hpf):** Epiboly displaces the blastoderm margin to 75% between the ends of the animal and vegetal poles (Figures 5 D, D’- Movie 2). At this stage, the head primordium develops anteriorly and the brain rudiment is visible. The posterior end of the embryonic axis develops a distinct swelling, the tail bud.

**90%-Epiboly (19 hpf):** The remnant of uncovered yolk protruding from the neighborhood of the vegetal pole

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**FIGURE 5** Gastrula period. (A-F and the corresponding magnification A’-F’) Lateral and (G-G’) dorsal stereomicroscope images of embryos at each stage of gastrula period: (A-A’) germ-ring stage, (B-B’) shield stage, (C-C’) 50%-epiboly stage, (D-D’) 75%-epiboly stage, (E-E’) 90%-epiboly stage, (F-F’ and G-G’) 100%-epiboly stage. Black arrowheads in F’ and G’ show somite furrow. White arrowheads (A to E) point to the edge of the blastoderm to better visualize epiboly’s progression. ES, embryonic shield; GR, germ ring; HP, head primordium; Kv, Kupffer’s vesicle; TB, tail bud; YP, yolk plug. Scale bar = 200 μm
may be now considered a yolk plug, the section of the yolk at the vegetal pole that has not been covered by the blastoderm (Figures 5 E, E'-Movie 2).

100%-Epiboly stage (21 hpf): Epiboly ends as the blastoderm completely covers the yolk plug, defining 100%-epiboly. The Kupffer's vesicle forms (Figure 5F') and the first three somites are visible. (Figures 5 F, F', G, G'-Movies 2 and 3).

2.6 | Segmentation period

The long axis of the embryo starts to extend even before epiboly is complete, and continues to do so during the segmentation period. Structures including the somites, tail, eyes and auditory vesicles begin to form. Additionally, the brain subdivisions start to become visible, and pigmentation appears, first on the yolk sac and later on

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**Figure 6** Segmentation period. (A-E and the corresponding magnification A’-E’)
Steremicroscope images of embryos at all stages of segmentation period: (A-A’) 6-somite stage, (B-B’) 12-somite stage, (C-C’) 15-somite stage, (D-D’) 18-somite stage and (E-E’) 22-somite stage. (F) Cartoon showing the development of the brain between 6 and 22-somite stages. Ce, cerebellum; Di, Diencephalon; Ep, epiphysis; EyP, eye primordium; Me, metencephalon; Mel, melanophore; No, notochord; Lp, lens primordium; OtV, otic vesicle; Te, telencephalon; V-S, V-shaped somites. Scale bar = 200 μm
the body axis (Figure 6-Movie 3). From the 1- to 18-somite stage, the rate of somite appearance is approximately one somite every 35 min.

Using somite number, we categorize the segmentation period of false clownfish embryos. Because false clownfish embryos at the late gastrula stage already possess three somites, the gastrula stage and segmentation period overlap slightly.

6-somite stage (23 hpf): The epiboly is complete (100%-epiboly) (Figures 6 A, A’-Movie 3). Six somites have formed and the tail bud is prominent at the posterior end of the body axis. At the anterior end of the embryo, the rudiment of the brain begins to subdivide and we can distinguish the telencephalon, the diencephalon and the mesencephalon (Figure 6F). The optic primordium (or eye primordium) is formed.

12-somite stage (28.5 hpf): At the 12-somite stage, the first melanophores are visible on the yolk (Figures 6B, B’-Movie 3).

15-somite stage (30.5 hpf): The notochord is clearly visible (Figures 6C, C’- Movie 3). At the posterior end, the tail bud begins to separate from the body as it becomes more elongated (Figure 6C, C’-Movie 3). At the anterior end, the lens primordium starts to be visible as well as the rudiment of the cerebellum and epiphysis (Figure 6F). The optic primordium (or eye primordium) is formed.

18-somite stage (32.5 hpf): The otic vesicle starts to form (Figure 6D, D’- Movie 3).

22-somite stage (34.5 hpf): This stage is characterized by the formation of V-shaped trunk somites (Figure 6E, E’). The twitching of the trunk muscles can also be observed (Movie 3).

2.7 Organogenesis

From the late segmentation stage onward, the embryo starts to turn itself within the chorion (Movie 4). Organogenesis starts when the embryo has a well-developed notochord, the brain is highly sculptured, and the lens primordia and otic vesicles are visible. The following changes then occur: the tail (post-cloacal region) begins to elongate and the heart, the median fin fold (ventral and dorsal) begins to elongate posteriorly (Figure 6E’-Movie 3). At the anterior end, the lens primordium starts to be visible as well as the rudiment of the cerebellum and epiphysis (Figure 6F). The optic primordium (or eye primordium) is formed.

4 dpf: At 4 dpf (Figure 7E to E’’’), a few pale orange xanthophores form on the ventral side of the trunk and posterior to the eye (Figure 7E’’- orange arrowheads). The common cardinal vein can be recognized as a pink tube in the lateral view (Figure 7E’’- and E’’’-white arrowheads). From this stage onward, the pectoral fin is elongated posteriorly (Figure 7EE’’’). The posterior part of the median fin fold (ventral and dorsal) is enlarged (Figure 7E’’’). The cloaca is formed and the mouth opens antero-ventrally (Figure 7E’’’).

5 dpf: The posterior part of the median fin (ventral and dorsal) begins to expand in a dorsoventral direction. The cardinal vein extends to the posterior region (Figure 7F’’). At this stage (Figures 7F-F’’’), the pectoral fin elongates posteriorly. The lens' black pigmentation increases. The position of the heart is visibly different between previous stages and this stage; the heart moves to the anterior, and is located at the anterior aspect of the yolk at this stage. Mouth openings are more anterior than at 4 dpf. The opercula can be observed in the ventral cranial region (Figure 7F’’’).

6 dpf: The embryo further increases in size (Figure 7G to G’’’). The eyes become silver. The lower jaw extends anteriorly, elongating the head in a more anterior direction. The jaw becomes vascularized.

7 dpf: Melanophores form two horizontal stripes on the tail and concentrate ventrally just dorsal to the yolk (Figure 7H to H’’’). Gills are clearly visible.
FIGURE 7  Organogenesis period. (A) Cartoon representing the “Otic vesicle closure” index (OVC). OVC corresponds to the ratio between the otic vesicle length (Lo) and the distance between the otic vesicle and the eye border (De-o + Lo). Graph represents the %OVC from 40 hpf until 5 dpf. Stereomicroscope images of embryos during organogenesis period: (B) 25%-OVC, (C) At 40%-OVC, (D) 80%-OVC, (E, E’, E”, E’’) 4 dpf; (F, F’, F”, F’’) 5 dpf, (G, G’, G”, G’’) 6 dpf, and (H, H’, H”, H’’) 7 dpf. Formation of blood circulation is indicated with white stars and arrowheads. Orange arrowheads point to orange xanthophores. Cl, cloacal; Cv, cardinal vein; Gi, gills; He, heart; Ir, iridophores; Lj, Lawer jaw; Mff, median fin fold; Mo, mouth; Op, opercula; PC, Pericardial cavity; Pf, pectoral fin. Scale bar = 200 μm
2.8 | Hatching

False clownfish embryos hatch generally once the lights are off 7 days after fertilization at 26°C (in our conditions). However, in some clutches we observed a one-day delay between one part of the clutch and the other.

The embryo hatches by breaking the egg capsule with its active wriggling, and the hatchlings emerge tail first (Figure 8A to A”). Because hatching was first observed in eggs that were scraped off of the terracotta pot, we wondered how hatching occurs when eggs are still fixed to their substrate. We succeeded in directly observing this event and the hatching sequence was the same (Figure 8B). Larvae in eggs which have been detached from their substrate may have difficulty extracting themselves from the chorion, but this is of course not the case with attached eggs. Once hatching occurs, young larvae still have some yolk (Figure 8C) and are immediately capable of swimming (data not shown).

2.9 | The development of larval pigmentation pattern

As the development of pigmentation is prominent during the embryonic development of *A. ocellaris*, particularly in the later stages, we thought it is important to summarize the different stages of the emergence of the larval pattern. Here, we describe development of two types of chromatophores, melanophores and xanthophores. We do not describe iridophore development because they are absent or not visible under the stereomicroscope, except those in the eye.

2.9.1 | Melanophores

Pigmentation by the melanophores is first observed from stage 12-somites (28 hpf) over the yolk; the melanophores migrate from the dorsal side of the yolk to reach the ventral side (29,5 hpf) (Figure 9A–D). Melanophores appear in the head region (Figure 9E) at 30 hpf, first localizing close around and behind the eyes (Figure 9E–H). On the trunk, pigmented melanophores appear in the posterior trunk between the embryo and the yolk (Figure 9G and H). Between 40 hpf and 7 dpf, the overall melanophore pattern does not change. At 7 dpf, tail melanophores localize along the myosepta to form a stripe, and ventral melanophores concentrate between the yolk and the ventral side of the embryo (Figure 9K).

2.9.2 | Xanthophores

At 4 dpf, a few pale orange xanthophores form on the ventral side of the trunk and posterior to the eye (Figure 9I-crop). At 6 dpf, the dorsal side of the larvae is pigmented by xanthophores (Figure 9J). At 7 dpf, xanthophores form a stripe on the head that runs through the eye (Figure 9K).

3 | DISCUSSION

In this study, we provide a staging series organized similarly to other models such as zebrafish, goldfish, sea bass and Atlantic salmon allowing for comparative studies. We also document the emergence of the larval
Figure 9  Pigmentation throughout larval development. (A, B, C, D) Time-lapse pictures of a ventral view of a single embryo from 28 hpf to 29.5 hpf. Black circles follow single melanophores migration over the yolk. (E, F, G, H) Time-lapse pictures of dorsal view of an embryo. (I, J, K) Pictures of dechorionated embryos at 4 dpf (I and its associated focus), 6 dpf (J) and 7 dpf (K). Black and orange arrows point respectively to melanophores and xanthophores.
pigmentation pattern that we have recently provided, linking embryonic development with the conspicuous adult pigmentation pattern of this species. These data provide a general framework that can be used as a tool to standardize studies and experimental procedures throughout false clownfish development.

3.1 False clownfish embryonic development in the teleost fishes context

Similar to previous studies on teleost development, we describe six periods of embryogenesis during embryonic development of *A. ocellaris*, namely the zygote, cleavage, blastula, gastrula, segmentation, and organogenesis periods, as well as 32 stages (Table 1 and Figure 2) reminiscent of those described in other models such as zebrafish, medaka, cichlids, and goldfish.11,14,32,36 These well-defined stages provide a framework for testing hypotheses of homologies within the various processes that occur during the embryonic development of teleost fishes. We observed interesting morphological differences between the embryos of false clownfish and other teleost species, whether they are aquaculture species (the sea bass *Dicentrarchus labrax*,34 the turbot *Scophthalmus maximus*,37 the batfish *Platax teira*38), other coral reef fish (the damselfish *Neopomacentrus cyanomus*,19 the dottybacks *Pseudochromis dilecticus*,17 the mandarinfish *Synchiropus splendidus*,39 the goby *Elacatinus puncticulatus*,40 several angelfishes of the genus Centropyge18,41) or other fish models (the stickleback *G. aculeatus*, the salmonid *Coregonus clupeaformis*,19 the gadid *Trisopterus luscus*,42 the wrasse *Labrus bergylta*43). These include the shape of the yolk/chorion, different early chromatophore patterns, and variation in developmental timing. Because of its specific life history traits (fixed eggs from a benthic marine fish, relatively long embryonic development with an immediately active larval phase), the false clownfish differs from most other coral reef fishes, whether they are aquaculture species (the sea bass *Dicentrarchus labrax*,34 the turbot *Scophthalmus maximus*,37 the batfish *Platax teira*38), other coral reef fish (the damselfish *Neopomacentrus cyanomus*,19 the dottybacks *Pseudochromis dilecticus*,17 the mandarinfish *Synchiropus splendidus*,39 the goby *Elacatinus puncticulatus*,40 several angelfishes of the genus *Centropyge*18,41) or other fish models (the stickleback *G. aculeatus*, the salmonid *Coregonus clupeaformis*,19 the gadid *Trisopterus luscus*,42 the wrasse *Labrus bergylta*43).

By comparing the growth and development of the larvae of four damselfish including an anemonefish (*Premnas biaculeatus*), Kavanagh and Alford, 2003 found a highly significant negative correlation between the duration of embryonic development and PLD that was also observed when more species were taken into account.44,45 This is not surprising, as damselfish life history strategies vary widely,45 and a shift in one life history trait can trigger the coordinated evolution of many other traits (eg, imprinting on sea anemone odor at hatching might require earlier development of the olfactory system46). Changing the respective duration of embryonic development and settlement time or altering with growth and developmental rates constitutes the range of levels on which evolution can operate to ensure the best possible survival in a vast series of ecological possibilities. This flexibility undoubtedly explains the great success of damselfishes and, this has yet to be demonstrated in other teleost fishes clades. The link between egg size, duration of embryonic development and short PLD, three features that differ between most other coral reef fishes, will be very interesting to study in more detail.44,47

3.2 The emergence of the larval pigmentation pattern

False clownfishes are well known for their iconic pigmentation patterns with three white bars formed by iridophores, lined with black melanophores on an orange background of xanthophores.48,49 This pattern emerges during metamorphosis, differs from the simpler larval pattern, and appears to be formed differently than the pattern of horizontal stripes well-studied in zebrafish.50,51

The emergence of pigmentation during embryonic development occurs in three main steps (i) it starts by the appearance of melanophores over the yolk at the 12-somites stages (28 hpf). These cells then migrate toward and invade the entire yolk at 30 hpf (Figure 9 A-D), (ii) At 30 hpf, melanophores appear in the vicinity of the eyes (Figure 9E) and later on the tail and in the posterior trunk at the border with the yolk (Figure 9F-H). Around 4dpf pale orange xanthophores are visible on the ventral side of the trunk and posterior to the eye (Figure 9D) (iii) Lastly, at 7 dpf, some melanophores localize along the myosepta to form a stripe and ventrally concentrate at the border between the yolk and the ventral side of the embryo (Figure 9K). At that stage xanthophores form a stripe that runs through the eye (Figure 9K). Interestingly, this pattern seems to be very similar between all anemonefishes, including *P. biaculeatus* and *Amphiprion perideraion*, in which lateral black marks of melanophores over a yellow pale body are clearly visible at hatching.52,53

The embryonic melanophore pattern that we observe in *A. ocellaris* is very different from those described in other teleost models such as cichlid fishes, medaka and zebrafish. Zebrafish, at hatching, have two dorsal and lateral melanophore stripes that originate from waves of migrating neural crest cells.51 In *A. ocellaris* these larval stripes are not present, suggesting a different migration pattern of neural crest cells,51 which are thought to generate all but the yolk melanophores in teleosts.51,54 In cichlids a single prominent ventral stripe of melanophores is visible.11 Further analysis using neural crest
markers could clarify the difference in ontogenetic processes that causes the emergence of such different melanophore patterns in teleosts. The genetic dissection of the diversity of this early and relatively simple pattern can be a relevant model to study the molecular and developmental basis of trait diversity.

4 | CONCLUSIONS

In this study, we provide a detailed illustrated and filmed embryonic developmental table for a coral reef fish: the false clownfish *A. ocellaris*. It is our hope that this staging will facilitate work with *A. ocellaris* as a model species, which can be used to better understand the genesis of coral reef fish developmental trajectories and how they are impacted by environmental perturbations. This emerging model will also allow investigation of a wide variety of biological questions such as the control of behavior, the diversification of pigmentation patterns and the link between embryonic development and life history strategies.

5 | EXPERIMENTAL PROCEDURES

5.1 | *A. ocellaris* rearing conditions

*A. ocellaris* used for imaging were obtained from different clutches laid by a single *A. ocellaris* breeding pair at the ICOB Marine Research Station in Taiwan, but the timing of events presented here aligns with that of the embryos of other pairs raised in Observatoire de Banyuls-Sur-Mer, France. The breeding pair was held in a 120-L tank of natural sea water at a temperature of 26°C, and a 12:12 h light: dark photoperiod. Adult fish were fed twice daily with live food including shrimp, squid, and the Japanese scad *Decapterus maruadsi*. Egg clutches were laid on a terra-cotta pot in the breeding tank.

5.2 | Stereomicroscope acquisition of high-resolution images and videos of *A. ocellaris* embryonic development

To follow the developmental changes occurring during embryonic development, we performed time-lapse using brightfield transmitted illumination of at least three embryos. Larvae were placed in a drop of filtered sea water on a Petri dish under the stereomicroscope. Live images were acquired on stereomicroscope (OLYMPUS SZX16, 1XPF) using a camera (DP80 using software CellSens DP80) with a picture resolution of 1360 x 1024 pixels and a time exposure of 142.9 ms (time-frame of 1 image/min). Measurements were performed using ImageJ software (V1.51k). For time-lapse of hatching, because embryos will not hatch under bright light, we placed a red plastic bag under the petri dish to decrease brightfield transmitted illumination.

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CONFLICT OF INTEREST

The authors declare that they have no competing interests.

ETHICS APPROVAL

These experiments were approved by the C2EA-36 Ethics Committee for Animal Experiment Languedoc-Roussillon (CEEA-LR), number A6601601. We have an approval number (A6601601) of premises for animal testing issued by the Regional Directorate of Food, Agriculture and Forestry of Occitania, and the Departmental Directorate of Protection of Populations of the Pyrenees Orientales. The animals were raised in our lab from breeding stock. Experimental protocols were based on the regulations in force in France (Articles R214-87 to R214-137 of the Rural Code), updated by Decree 2013-118 and by five decrees dated February 1, 2013, and published February 7, 2013, pursuant to Directive 2010/63/EU. This regulation is under the responsibility of the Ministry of Agriculture.

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SUPPORTING INFORMATION

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