The developmental hourglass model and recapitulation: An attempt to integrate the two models

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Abstract
Recapitulation is a hypothetical concept that assumes embryogenesis of an animal parallels its own phylogenetic history, sequentially developing from more ancestral features to more derived ones. This concept predicts that the earliest developmental stage of various animals should represent the most evolutionarily conserved patterns. Recent transcriptome-based studies, on the other hand, have reported that mid-embryonic, organogenetic periods show the highest level of conservation (the developmental hourglass model). This, however, does not rule out the possibility that recapitulation would still be detected after the mid-embryonic period. In accordance with this, recapitulation-like morphological features are enriched in late developmental stages. Moreover, our recent chromatin accessibility-based study provided molecular evidence for recapitulation in the mid-to-late embryogenesis of vertebrates, as newly evolved gene regulatory elements tended to be activated at late embryonic stages. In this review, we revisit the recapitulation hypothesis, together with recent molecular-based studies that support the developmental hourglass model. We contend that the recapitulation hypothesis does not entirely contradict the developmental hourglass model and that these two may even coexist in later embryonic stages of vertebrates. Finally, we review possible mechanisms underlying the recapitulation pattern of chromatin accessibility together with the hourglass-like evolutionary conservation in vertebrate embryogenesis.

KEYWORDS
chromatin accessibility, developmental hourglass model, evo-devo, gene expression regulation, parallelism, recapitulation

1 | INTRODUCTION

Diverse multicellular animals with different body plans originally evolved from a simple unicellular organism. Similarly, animal development starts with a single zygote that develops into a multicellular adult with complex morphological patterns. This tendency of increasing complexity along a time course has inspired scientists to propose parallel relationships between development and evolution (or taxonomic hierarchy) (Gould, 1977; Hall, 1992; Oppenheimer, 1959; Richardson & Keuck, 2002). One of the most
famous of these theories is Haeckel’s recapitulation theory (Haeckel, 1866), which predicted that development is a rapid replay of the evolutionary trajectory.

Although the theory of recapitulation is no longer accepted in its original form, the essence of the theory led to the early conservation model, which predicts that the earliest stage of development is expected to show the most ancestral, or most evolutionarily conserved features. This early conservation model appears to be reasonable from a developmental perspective, at least intuitively, because changes in earlier developmental processes are thought to have greater and usually harmful impacts on subsequent developmental processes (Arthur, 1997; Garstang, 1922; Riedl, 1978; Wimsatt, 1986), possibly leading to the conservation of earlier stages (Figure 1a). On the contrary, recent transcriptome-based studies have demonstrated that it is the mid-embryonic phase, and not the earliest one, that is the most conserved across species of a given phylum (Domazet-Lošo & Tautz, 2010; Drost et al., 2015; Gildor et al., 2019; Hu et al., 2017; Irie & Kuratani, 2011; Israel et al., 2016; Kalinka et al., 2010; Levin et al., 2012; Malik et al., 2017; Marlé et al., 2018; Wang et al., 2013; Yanai et al., 2011). Notably, this tendency was reported in several phyla, indicating that “the developmental hourglass model” (Duboule, 1994) would be one of the most plausible hypotheses that clarifies the apparent relationship between development and evolution in each animal phylum (Figure 1b). While the hourglass-like conservation of animal embryogenesis appears to rule out the recapitulation theory, there remains a possibility that “recapitulation pattern” may hold true, at least partially, in the diversifying phase of embryogenesis. Furthermore, several morphological features appear to favor the recapitulation theory (see below). In this review, we propose a plausible concept involving the integration of the hourglass model with recapitulation and review possible evolutionary mechanisms underlying the tendency toward embryonic evolution.

**FIGURE 1** Schematic diagrams that depict evolutionary divergence along embryogenesis. (a) The early conservation model, which assumes that the earliest stage is the most highly conserved. (b) The developmental hourglass model (Duboule, 1994), which assumes the highest conservation across species of the same phylum at the mid-embryonic period or the phylotypic period. (c) The nested hourglasses model with shifting bottlenecks. The most conserved mid-embryonic phase shifts to later stages when comparisons were made in smaller phylogenetic groups, which may be regarded as the recapitulation pattern.
heart, they do not show features that resemble those of adult ancestral fish. Moreover, different features that appear at different stages of development may often evolve independently (Garstang, 1922; Richardson & Keuck, 2002), thereby contradicting predictions of the original recapitulation theory. Considered together, it may be more appropriate to regard the recapitulation as a tendency that evolutionarily more derived features tend to appear sequentially at later developmental stages, in the same order of their evolutionary origins. The current review aims to follow this definition.

2.2 The developmental hourglass model

According to the re-defined recapitulation mentioned above, earlier embryonic features may be expected to display more ancestral features than later embryonic stages. This early conservation model (Figure 1a) has been upheld by several theoretical studies (Arthur, 1997; Garstang, 1922; Riedl, 1978; Wimsatt, 1986). However, no consensus has been reached as to whether the earliest stage is evolutionarily the most conserved. For example, morphologically diverse features observed during early developmental stages, such as variable cell cleavage patterns, gastrulation, and axis formation, are often referred to as typical examples of evidence that counter the early conservation model (Kalinka & Tomancak, 2012; Raff, 1996; Schierenberg, 2006; Slack et al., 1993). Moreover, the situation has become far more complicated as several counter models have been proposed (Bininda-Emonds et al., 2003; Duboule, 1994; Poe & Wake, 2004; Raff, 1996; Richardson, 1999; Slack et al., 1993). The developmental hourglass model, for instance, assumes the highest conservation within each phylum at the mid-embryonic period (Duboule, 1994; Raff, 1996). This originates from morphological similarities that were apparent at vertebrate organogenesis stages, which have been pointed out so far by many biologists (von Baer, 1828; Ballard, 1981; Duboule, 1994; Haecikel, 1874; Raff, 1996; Slack et al., 1993). For example, the vertebrate embryos around pharyngula period share a variety of primordial organs, such as notochord, somite, dorsal nerve cord, the pharyngeal arch, rhombomere, and optic anlagen. Despite these apparent similarities, another morphological study with detailed quantitative analyses showed that the highest conservation did not occur at the mid-embryonic phase, but rather proposed that the mid-embryonic phase was the most divergent stage (Bininda-Emonds et al., 2003). Furthermore, others proposed the ontogenetic adjacency model, which predicts that there is no global tendency in the evolutionary conservation of embryonic stages, except for small changes (Poe & Wake, 2004).

While comparative morphological approaches were unable to resolve the controversy (Bininda-Emonds et al., 2003; Poe, 2006; Poe & Wake, 2004; Richardson et al., 1997), the situation changed with the advent of molecular-based studies focusing on gene expression profiles during embryogenesis (Domazet-Lošo & Tautz, 2010; Hazkani Covo et al., 2005; Irie & Kuratani, 2011; Irie & Sehara-Fujisawa, 2007; Kalinka et al., 2010; Roux & Robinson-Rechavi, 2008). Specifically, transcriptomic technologies, such as microarray and RNA sequencing, provided quantitative approaches that allowed conserved developmental stages between different species to be evaluated. The findings of most studies, including ours, have supported mid-embryonic conservation rather than early conservation in various animal groups, including chordates (Domazet-Lošo & Tautz, 2010; Hu et al., 2017; Irie & Kuratani, 2011; Irie & Sehara-Fujisawa, 2007; Marlétaz et al., 2018; Wang et al., 2013), echinoderms (Gildor et al., 2019; Li et al., 2020; Malik et al., 2017), arthropods (Domazet-Lošo & Tautz, 2010; Kalinka et al., 2010), nematodes (Levin et al., 2012; Schiffer et al., 2018), and mollusks (Xu et al., 2016). Thus, accumulating evidence suggests that the mid-embryonic period reflects the most highly conserved expression profiles in a phylum or less inclusive groups, thereby contradicting the early conservation model. However, this does not necessarily refute the possibility that the recapitulation pattern square with the developmental hourglass model at later developmental stages with increasing diversity (Abzhanov, 2013; Kuratani, 2004). Specifically, if the most conserved stage of the lower taxa shifts sequentially to later periods, these two ideas may coexist with each other at the latter part of embryogenesis (Figure 1c; the nested hourglasses model with shifting bottlenecks).

With regard to this viewpoint, the nested hourglasses model, which was tested at the transcriptomic-level using several vertebrate species, demonstrated that the target of the mid-embryonic conservation remained mostly the same, irrespective of the phylogenetic scales analyzed (Hu et al., 2017; Wang et al., 2013). For example, the highly conserved mid-embryonic stage identified in chicken-turtle comparison (around stage HH16 in chicken) largely overlapped those identified in other tetrapod comparisons (Wang et al., 2013). Similarly, Hu et al. (2017) examined eight chordates covering different taxa and showed that conserved developmental stages in different phylogenetic groups (i.e., Osteichthyes, Tetrapoda, Aminota, Xenopus, and Archelosauria) largely overlapped each other (Figure 2a). Furthermore, concerning the higher level of taxa, mid-embryonic conservation was observed up to Vertebrata, but was only weakly supported in Chordata and Olfactores (Figure 1a; Hu et al., 2017). This does not contradict the study by Marlétaz et al. (2018) which suggested that the most conserved stages are not comparable between vertebrates and chordates. Considered together, these comparative transcriptomic studies imply that the mid-embryonic period may have been the target of conservation persistently through the vertebrate evolution (persistent conservation; Irie & Kuratani, 2014), and the recapitulation pattern does not seem to coexist even partially during later developmental stages.
The persistently conserved mid-embryonic period at the transcriptomic level and recapitulation pattern observed in chromatin accessibility in the late embryogenesis. In both panels, the results of chicken development are shown. (a) Conserved developmental stages in different taxonomic groups are estimated by ortholog-group-based comparative transcriptome (Hu et al., 2017). The percentages of chicken developmental stages included in the most similar combinations of staged embryos from five different taxonomic groups are shown as Ptop in different colored bar graphs (for detail, see Methods in Hu et al., 2017). Higher values in Ptop represent higher levels of whole-embryonic transcriptomic conservation for the species compared. Note that stages that show the body plan for vertebrates (potential phylotypic period, highlighted in gray) largely overlap each other, except for the bottom graph showing conservation for chordate-conserved stages in chicken. Meanwhile, mid-embryonic conservations up to Chordata were observed in the analysis with strictly conserved 1-to-1 orthologs (see Supplementary Information in Hu et al., 2017). Essentially the same results were observed for the other vertebrate species (see Hu et al., 2017). (b) Evolutionarily categorized chromatin accessibility during chicken development (Uesaka et al., 2019). For each developmental stage, the percentage on the vertical axis represents the relative chromatin accessibilities of the genomic regions categorized by the evolutionary ages of their sequences (see color and phylogenetic tree in the right for the category). In each graph after the conserved mid-embryonic period, the developmental stages with the highest signal (highlighted in the corresponding colors) shift sequentially to later stages by their evolutionary category. While earlier stages before the conserved stages show high signals of ancestral genomic regions (e.g., those conserved in vertebrates and olfactores), newly acquired genomic regions (e.g., those acquired since the chicken-alligator split) also show higher chromatin accessibilities. Essentially the same patterns were observed for the other species (see Uesaka et al., 2019), and these indicate that embryogenesis before the conserved stages does not show the recapitulation pattern with the unknown evolutionary mechanism. The figures are adapted and modified from Hu et al. (2017) and Uesaka et al. (2019)

3 | POSSIBLE COEXISTENCE OF THE DEVELOPMENTAL HOURGLASS MODEL AND THE RECAPITULATION PATTERN IN VERTEBRATE EMBRYOGENESIS

3.1 | Morphological examples showing the recapitulation pattern

In addition to the denial of the recapitulation pattern at transcriptomic level in later developmental stages, previous studies further supported the idea that the body plan of each animal phylum is defined by the most conserved mid-embryonic stage (phylotype hypothesis; Duboule, 1994; Hu et al., 2017; Irie et al., 2018; Raff, 1996; Wang et al., 2013). This implies that embryos develop the body plan of their phylum first and then develop lineage-specific features afterward, indicating that no recapitulation pattern emerges after the conserved mid-embryonic phase. However, similarities between whole-embryonic gene expression profiles do not necessarily indicate morphological conservation (Irie & Kuratani, 2014), although these presumably reflect the composition of homologous cells. It is
thus worth overviewing studies based on comparative morphological approach.

In accordance with the transcriptomic studies, clear counter examples against the recapitulation pattern in the mid-to-late embryogenesis has also been reported for morphological features (Bininda-Emonds et al., 2003; Koyabu & Son, 2014; Poe, 2006; Prum, 1999; Prum & Brush, 2015; Richardson et al., 2009; Weisbecker et al., 2008; Ziermann & Diogo, 2014). Nonetheless, a reasonable number of morphological traits display recapitulation patterns during evolution, especially in vertebrates (Gould, 1977; Richardson & Keuck, 2002). One example is the developmental pattern of ribs and scapulae during turtle embryogenesis (Nagashima et al., 2009). Unlike other amniotes, adult turtles are exceptional in that they have scapulae (shoulder blades) beneath the ribs, or the carapace. Despite the unique anatomy of adult turtles, they retain a typical amniote pattern early in their embryogenesis as follows: primordia of scapulae emerge at the outside of rib primordia and then tilt to locate at the ventral side of the ribs, which in intermediate stage resemble that of an ancestral fossil turtle, Odontochelys (Li et al., 2008), seemingly recapitulating the ancestral state. In addition, there are more examples that seem to display the recapitulation-like pattern, such as the caudal morphology of teleost fishes (Metscher & Ahlberg, 2001; see also Sallan, 2016), the elongated jaw of some needlefishes (Lovejoy et al., 2004; Lovejoy, 2000), the hind limb buds that become regressed during embryogenesis in snakes and cetaceans (Bejder & Hall, 2002; Cohn & Tickle, 1999; Thewissen et al., 2006), the opposable hallux of birds (Botelho et al., 2015), the position of the hallux in the foot of birds (Botelho et al., 2017), the position of the external naris of cetaceans (Thewissen, 2018), and the three-digit hind limb of jerboas and camels (Cooper et al., 2014). Diogo et al. (2015) also argued that the majority of head muscles of zebrafish, salamanders, and turtles tend to develop sequentially in the order of their evolutionary origins. Importantly, all these examples that seem to show the recapitulation pattern were observed in later embryogenesis following the organogenesis phase.

3.2 Testing the recapitulation pattern in the context of chromatin accessibility

Considering that previous comparative studies did not support coexistence between the hourglass model and recapitulation at the level of the whole embryonic transcriptome (Hu et al., 2017, Wang et al., 2013), how could we reconcile the recapitulation pattern observed for a significant number of morphological traits with the hourglass model (Bejder & Hall, 2002; Botelho et al., 2015, 2017; Cohn & Tickle, 1999; Cooper et al., 2014; Diogo et al., 2015; Gould, 1977; Lovejoy et al., 2004; Metscher & Ahlberg, 2001; Nagashima et al., 2009; Richardson & Keuck, 2002; Thewissen et al., 2006). One possibility would be that, although some traits display recapitulation patterns during development both at morphologic and transcriptomic levels, the overall recapitulation pattern in developing whole embryos disappears when all traits of the embryos are taken into account. Briefly, if only a minority of traits follow recapitulation, then the recapitulation pattern would not be detected at whole embryo level. In this case, no reconciliation is needed. Another possibility is that the transcriptomic approach lacks sufficient resolution to detect the recapitulation pattern. This concern arises from the fact that a cross-species comparative transcriptomic approach mainly focuses on the expression of orthologous coding genes, whereas many protein-coding genes are known to be repeatedly recruited at different developmental stages (Hu et al., 2017). In addition, most genomic mutations that accumulated during the evolution of major vertebrate groups have been identified in noncoding regions rather than in protein-coding regions (Mikkelsen et al., 2007; Seki et al., 2017). Thus, it would be helpful to measure the activities of noncoding regulatory regions to test recapitulation patterns during development. One of a proxy of the regulatory region activity at various developmental stages is chromatin accessibility in embryos. Accessible chromatin marks active regulatory regions, including enhancers, promoters, and silencers (Gross & Garrard, 1988; Klemm et al., 2019), because accessible chromatin regions become physically accessible to transcription factors for the execution of their functions. ATAC-seq (an assay for transposase-accessible chromatin using sequencing; Buenrostro et al., 2013) is widely utilized to profile chromatin accessibility on a genome-wide scale. Thus, with this method, we measured chromatin accessibilities of developing embryos in a recent study of ours (Uesaka et al., 2019). The chromatin accessibilities of whole embryos are expected to reflect the number of cells in which the chromatin state of the examined region is accessible, which would be consistent with the recently confirmed similarities between ATAC-seq profiles of bulk samples and aggregate profiles of single-cell ATAC-seq (Lareau et al., 2019; Satpathy et al., 2019). Along with this postulate, we quantitatively examined chromatin accessibilities of genomic regions during development, and found that these showed a recapitulation pattern (Uesaka et al., 2019). In brief, we estimated the activities of gene regulatory elements through embryogenesis of three vertebrates (chicken, mouse, and medaka) using ATAC-seq, and analyzed whether sequence conservation of these accessible chromatin regions showed any evolutionary tendencies along with the development of each species. The results indicated that genomic regions conserved at the sequence level in lower taxa tended to show the highest chromatin accessibilities at later developmental stages (Figure 2b for chicken; for mice and medaka, see Uesaka et al., 2019). In other words, evolutionarily newer genomic regions tend to become accessible at later developmental stages, suggesting a parallel relationship between the evolutionary age of genomic regions, and the timing of their accessibility along the development. Unlike the embryogenesis from the conserved mid-embryonic period of vertebrates onwards, early-to-mid embryogenesis did not show a recapitulation pattern in chromatin accessibility (Figure 2b), indicating that a chromatin-level recapitulation pattern can only be observed after the conserved mid-embryonic period. The recapitulation pattern observed in whole embryonic chromatin
accessibility (Uesaka et al., 2019) coincides well with morphological traits that show recapitulation patterns, at least in terms of timing in development, as both take place mainly after the conserved mid-embryonic phase of the hourglass model.

3.3 The relationship between the recapitulation pattern of chromatin accessibility and the hourglass-like embryonic divergence

Both transcriptome and chromatin accessibility-based studies indicated that the mid-embryonic phase retained the most ancestral features, followed by derived features that developed afterward. According to this perspective, the recapitulation pattern of chromatin accessibility and transcriptomic hourglass-like divergence may coexist during mid-to-late embryogenesis (Figure 3; coexistence model). Although there is less empirical evidence, several assumptions can be made for possible mechanisms underlying this model. One is that evolutionarily newer gene regulatory elements could be the mechanism behind diversified gene expression profiles, as well as the morphological traits showing recapitulation patterns that appear in the later embryonic stages. Another assumption is that evolutionarily newer genomic regions activated at earlier stages lead to divergence in early embryogenesis at both the transcriptomic level (Hu et al., 2017; Irie & Kuratani, 2011; Wang et al., 2013) and the morphological level, such as the size of eggs, forms of blastula, and developing gastrula (Kalinka & Tomancak, 2012; Raff, 1996; Schierenberg, 2006; Slack et al., 1993). Meanwhile, higher chromatin accessibility of conserved genomic regions at early stages, such as the genomic regions conserved in vertebrates and in Olfactores, could be associated with deeply conserved embryonic patterns, such as egg cleavage, gastrulation, and neurulation (Figure 2b).

The above-stated assumptions require detailed investigation, mainly because the association between developmental patterns in chromatin accessibility and hourglass-like gene expression divergence as well as morphological traits that show recapitulation patterns remain unclear. Even the relationship between the recapitulation pattern of chromatin accessibility and hourglass-like transcriptomic divergence requires clarification on several points. For example, genomic regions with conserved sequences do not necessarily show a conserved pattern in chromatin accessibility (Stergachis et al., 2014; Vierstra et al., 2014), implying that such conserved genomic regions may not be linked with evolutionarily conserved gene expression. Another issue is raised by the fact that not all accessible chromatin regions contribute to the regulation of gene expression (The ENCODE Project Consortium, 2012). Therefore, identification of regulatory sequences, such as enhancers and repressors that promote hourglass-like divergence of gene expression is needed. More detailed analyses of single-cell-based genome-

![FIGURE 3](image_url)
wide chromatin accessibility and transcriptomes in the same cell may help elucidate the relationship between the recapitulation pattern of chromatin accessibility and hourglass-like gene expression divergence (Cao et al., 2018; Chen et al., 2019; Zhu et al., 2019). In addition, the previous study conducted by us (Uesaka et al., 2019) did not include a cross-species comparison of developmental chromatin accessibility to confirm whether evolutionarily newer chromatin accessibility displays a higher diversity to match with the latter part of the hourglass model.

Furthermore, the relationship between the chromatin accessibility and morphological features that display recapitulation patterns also remains unclear. Based on the reports indicating that the evolution of lineage-specific morphological traits in vertebrates is often associated with evolutionary acquisitions of gene regulatory elements (Carroll, 2005; Nishihara et al., 2016; Seki et al., 2017; Shim et al., 2012; Tashiro et al., 2011), one might expect that morphological development showing a recapitulation pattern could be attributed to the sequential activation of gene regulatory elements in a recapitulation manner. However, it is not clear whether this relationship holds true for the overall tendency of morphological traits. In certain cases, newly acquired gene regulatory elements may replace ancestral regulatory elements during evolution, while still maintaining their function (Odom et al., 2007; Weirauch & Hughes, 2010; Wilson & Odom, 2009). Finally, the recapitulation pattern of developmental chromatin accessibility has only been observed in a limited number of vertebrates, and thus it is unclear how widely this pattern can be observed in the animal kingdom. Therefore, further studies involving both comparative transcriptomic analyses as well as whole-embryonic chromatin accessibility analyses would be needed to test the generality of the possible coexistence.

### 4 | POTENTIAL EVOLUTIONARY MECHANISMS UNDERLYING THE RECAPITULATION PATTERN OBSERVED IN VERTEBRATE EMBRYOGENESIS

Based on the findings that developmental chromatin accessibility at the whole embryo level shows the recapitulation pattern only after the conserved mid-embryonic phase of the vertebrates and that many morphological features that show the recapitulation pattern appear during the later embryonic stages, we have proposed the potential coexistence of the recapitulation and the hourglass model. Meanwhile, the evolutionary mechanisms underlying this relationship remain largely unknown. In this last section, we discuss the possible mechanisms that may underlie the evolutionary tendency of vertebrate embryogenesis.

#### 4.1 | Possible mechanisms underlying the evolutionary tendency of chromatin accessibility

Why do the chromatin accessibilities of more evolutionarily derived genomic regions tend to become higher at later stages to show a recapitulation pattern? One attractive concept to explain this would be “developmental burden” (Riedl, 1978). According to this concept, since preceding developmental processes serve as a prerequisite for later processes, earlier developmental processes are less prone to change, leading to the evolutionary conservation of earlier processes. Similar mechanisms, such as the stepping stone model (Garstang, 1922), somatic program (Mayr, 1994), epigenetic characters (Levtrup, 1978), and generative entrenchment (Wimsatt, 1986) have been proposed by other scientists.

Developmental burden has often been used to explain the conservation of certain morphological traits, such as the pharyngeal arches and notochord of vertebrate embryos. For example, while adult terrestrial vertebrates do not have gills, they still develop pharyngeal arches during embryogenesis, and this could be attributed to its responsibility for the development of other organs, such as the thymus. Similarly, a possible scenario based on the concept of the developmental burden would be that gene regulatory activities at earlier stages exert a greater impact on the following developmental processes when mutated, and thus are more difficult to change during evolution. This could have led to the conservation of these gene regulatory elements in both nucleotide sequences and their activities during early stages, which furthered the recapitulation pattern.

However, our recent report measured only sequence-level conservation of potential regulatory regions, and whether or not the activities of these regions show cross-species conservation remains unknown. Critically, the recapitulation pattern of chromatin accessibility was observed only up to the conserved mid-embryonic phase, indicating that simple application of developmental burden throughout embryogenesis may not be justified. This problem will be discussed further in the next section.

#### 4.2 | Why is the recapitulation pattern restricted to the later part of embryogenesis?

Our recent findings indicated that the most evolutionarily new genomic regions showed higher chromatin accessibilities not only in the latest developmental stages, but also during the stages before the transcriptomically conserved mid-embryonic period (Figure 2b). Considered together with transcriptomic divergence observed during the earlier stages, these appear to debunk the assumption that developmental burden acts on all stages and proceeds throughout embryogenesis.

In interpreting the developmental burden concept using these observations, one assumption would be that the developmental burden acts upon all stages of embryogenesis, but its effect is masked by stronger diversifying effects from other evolutionary mechanisms such as natural selection. For example, some studies contend that strong positive selective pressure is exerted on early embryonic stages during diversification and adaptation to a variety of environments (Kalinka & Tomancak, 2012; Slack et al., 1993). This could have led to high chromatin accessibilities in recently acquired genomic regions during early stages, as reported for transcriptomic divergence (Hu et al., 2017; Irie & Kuratani, 2013; Wang et al., 2013).
and some lineage-specific morphological traits (Kalinka & Tomancak, 2012; Raff, 1996; Schierenberg, 2006; Slack et al., 1993). In addition, selective pressures on maternal reproductive strategies could also exert similar effects at the beginning of embryogenesis, as developmental features that emerge during initial stages, such as egg size and yolk amount, are established and regulated by maternal phenotypes (Abzhanov, 2013; Kalinka & Tomancak, 2012). To further expand this burden-oriented perspective, the activation of highly ancestral chromatin accessibility during earlier stages also requires further explanation (Figure 2b), as these may be the remnants of a strong conservation effect due to developmental burden. Unfortunately, the above-mentioned burden-oriented assumptions have no experimental basis, and thus far, no study has succeeded in measuring the force of developmental burden to substantiate its existence. Thus, potential evolutionary mechanisms that underlie the later-stage recapitulation patterns of chromatin accessibility remain largely unknown.

4.3 | Gene regulatory background for the mid-embryonic conservation of gene expression

The question remains as to why developmental stages only after organogenesis exhibit the recapitulation pattern in chromatin accessibility. As observed so far, even the most attractive hypothesis, such as developmental burden, fails to fully explain these findings. It is tempting to speculate that the conserved mid-embryonic phase acts as a singular point, or boundary that inhibits the recapitulation pattern before this phase. With this perspective, a classic argument is that the mid-embryonic phase is the target of the strongest negative selection due to embryonic lethality (Raff, 1996). This concept has been referred to as the conservation mechanism for the bottleneck part of the hourglass model; however, it was not supported by our recent study (Uchida et al., 2018). Namely, embryonic lethality of vertebrate embryos under normal and perturbation conditions suggested that it was the early embryonic phase, rather than the mid-embryonic phase, that was targeted by negative selection pressure owing to lethality (Uchida et al., 2018). Another feature unique to the conserved mid-embryonic phase is the enrichment of pleiotropically expressed genes (Hu et al., 2017; Li et al., 2020; Liu & Robinson-Rechavi, 2018). Based on this, Hu et al. suggested that pleiotropic constraint (Duboule & Wilkins, 1998; Galis, 1999) at the gene regulation level could be a contributing factor for mid-embryonic conservation. Although the exact molecular mechanism of pleiotropic constraint at the gene regulation level remains to be clarified, it would be of interest to investigate the relevance of regulatory elements, such as pleiotropic enhancers (Sabarís et al., 2019) or chromatin accessibility, in this process. These factors, however, do not provide any reasonable mechanisms for the boundary of the recapitulation pattern at the mid-embryonic phase. Furthermore, it is also problematic that the recapitulation pattern has been observed only for chromatin accessibility, and not for the transcriptome (Hu et al., 2017; Irie & Kuratani, 2011; Uesaka et al., 2019; Wang et al., 2013; see Figure 1c; the nested hourglasses model with multiple bottlenecks). Notably, a theoretical study suggested that gene activations showing the recapitulation pattern arise with a computer-based simulation that evolved linearly aligned artificial cells with simple gene regulatory networks (Kohskakabe & Kaneko, 2016). However, the discrepancies observed between this and the experimental studies would be due to the different conditions of development and evolution between artificial and real organisms. Further analyses utilizing the epigenome and transcriptome data from multiple developing tissues, including public databases, such as the ENCODE database (Davis et al., 2018), would provide a basis for understanding the relationship between the gene regulatory elements and hourglass-like transcriptomic divergency.

5 | CONCLUSIONS AND FUTURE ISSUES

While the recapitulation pattern was not observed for whole-embryonic transcriptomes, it was observed for developmental chromatin accessibility during vertebrate embryogenesis after the most conserved mid-embryonic period. This discrepancy remains to be clarified. However, it is possible that the recapitulation and the developmental hourglass model coincide with each other during later vertebrate embryogenesis, as both imply that more derived features appear after the most conserved mid-embryonic period. The evolutionary mechanism underlying such coexistence at later stages remains largely unknown. However, it may be useful to clarify whether the recapitulation pattern of chromatin accessibility underlies the diversifying gene expression profiles and the recapitulation pattern in the morphological development of some traits. It also has to be noted that the coexistence model has not been validated enough even in vertebrates, and thus the generality of this model remains unclear. Thus, further studies focusing on not only various vertebrates but also species from different phyla are awaited. Clarifying these remaining questions will further advance our understanding not only of the laws on the evolution of animal embryogenesis but also of how animal embryogenesis has evolved and would evolve.

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CONFLICT OF INTERESTS

The authors declare that there are no conflict of interests.

DATA AVAILABILITY STATEMENT

Original data are available from the authors upon request.
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