Clustered brachiopod Hox genes are not expressed collinearly and are associated with lophotrochozoan novelties

Sabrina M. Schiemann¹, José M. Martín-Durán¹, Aina Børve¹, Bruno C. Vellutini¹, Yale J. Passamanecck², Andreas Hejnol¹*

¹ Sars International Centre for Marine Molecular Biology, University of Bergen, Bergen 5006, Norway
² Kewalo Marine Laboratory, Pacific Biosciences Research Center, University of Hawaii, Honolulu, HI, USA

*Corresponding author: Andreas Hejnol (andreas.hejnol@uib.no)
Abstract

Temporal collinearity is often regarded as the force preserving Hox clusters in vertebrate genomes. Studies that combine genomic and gene expression data in invertebrates would allow generalizing this observation across all animals, but are scarce, particularly within Lophotrochozoa (e.g., snails and segmented worms).

Here, we use two brachiopod species—*Terebratalia transversa*, *Novocrania anomala*—to characterize the complement, cluster and expression of their Hox genes. *T. transversa* has an ordered, split cluster with ten genes (*lab*, *pb*, *Hox3*, *dfd*, *scr*, *lox5*, *antp*, *lox4*, *post2*, *post1*), while *N. anomala* has nine (missing *post1*). Our *in situ* hybridization, qPCR and stage specific transcriptomic analyses show that brachiopod Hox genes are neither strictly temporally nor spatially collinear; only *pb* (in *T. transversa*), *Hox3* and *dfd* (in both brachiopods) show staggered mesodermal expression. The spatial expression of the Hox genes in both brachiopod species correlates with their morphology and demonstrates cooption of Hox genes in the chaetae and shell fields, two major lophotrochozoan morphological novelties.

The shared and specific expression of a subset of Hox genes, *Arx* and *Zic* orthologs in chaetae and shell-fields between brachiopods, mollusks, and annelids supports the deep conservation of the molecular basis forming these lophotrochozoan hallmarks. Our findings challenge that collinearity alone preserves lophotrochozoan Hox clusters, indicating that additional genomic traits need to be considered in understanding Hox evolution.
Introduction

Hox genes are transcription factors that bind to regulatory regions via a helix-turn-helix domain to enhance or suppress gene transcription [1, 2]. Hox genes were initially described in the fruit fly *Drosophila melanogaster* [3, 4] and later on in vertebrates [5-7] and the nematode *Caenorhabditis elegans* [8]. In all these organisms, Hox genes were shown to provide a spatial coordinate system for cells along the anterior-posterior axis [9]. Remarkably, the Hox genes of these organisms are clustered in their genomes and exhibit a staggered spatial [3] and temporal [10, 11] expression during embryogenesis that corresponds to their genomic arrangement [3, 12, 13]. These features were used to classify Hox genes in four major orthologous groups –anterior, Hox3, central and posterior Hox genes– and were proposed to be ancestral attributes to all bilaterally symmetrical animals [1, 13, 14].

However, the study of the genomic arrangements and expression patterns of Hox genes in a broader phylogenetic context has revealed multiple deviations from that evolutionary scenario. Hox genes are prone to gains [15-17] and losses [18-21], and their arrangement in a cluster can be interrupted, or even completely disintegrated [22-25]. Furthermore, the collinear character of the Hox gene expression can fade temporally [24, 26, 27] and/or spatially [28]. Hox genes have also diversified their roles during development, extending beyond providing spatial information [29]. In many bilaterian embryos, Hox genes are expressed during early development, well before the primary body axis is patterned [26, 30-32]. They are also involved in patterning different tissues [33] and have been often recruited for the evolution and development of novel morphological traits, such as vertebrate limbs [34, 35], cephalopod funnels and arms [28], and beetle horns [36].
It is thus not surprising that Hox genes show diverse arrangements regarding their genomic organization and expression profiles in the Spiralia [37], a major animal clade that includes a high disparity of developmental strategies and body organizations [38-42]. A striking example is the bdelloid rotifer Adineta vaga, which belongs to the Gnathifera, the possible sister group to all remaining Spiralia [41, 42]. As a result of their reduced tetraploidy, its Hox complement includes 24 genes, albeit it lacks posterior Hox genes and a Hox cluster [43]. The freshwater flatworms Macrostomum lignano and Schmidtea mediterranea also lack a Hox cluster [44, 45] and parasitic flatworms have undergone extensive Hox gene losses, likely associated with their particular life style [21]. Interestingly, the limpet mollusk Lottia gigantea [16] shows a well-organized Hox cluster. Other mollusks (e.g. the pacific oyster Crassostrea gigas) and the segmented annelid Capitella teleta exhibit organized split Hox clusters [46, 47]. On the other hand, the cephalopod mollusk Octopus bimaculoides has lost several Hox genes and lacks a Hox cluster [22]; and the clitellate annelids Helobdella robusta and Eisenia fetida do not show a Hox cluster and have greatly expanded some of the Hox classes [16, 17].

Although Hox gene expression is known for a handful of spiralian species [26, 44, 46, 48-58], the relationship between genomic organization and expression domains is known for only three of them, namely the annelids C. teleta and H. robusta, and the planarian S. mediterranea. Consistent with the lack of a Hox cluster, H. robusta and S. mediterranea show neither temporal nor spatial collinearity [44, 54-56]. Conversely, C. teleta, which has an organized, broken cluster, does exhibit these features [46]. In general, these observations suggest that the presence of collinearity –
in particular, temporal collinearity– could be associated with the retention of a more or less intact spiralian Hox cluster, as it seems the case for the vertebrate cluster [14, 23, 59, 60]. However, more studies combining genomic and expression information, and including the vast spiralian morphological diversity, are essential to draw robust conclusions about Hox gene evolution and regulation in Spiralia and Metazoa [61]. These studies would also allow to test if hypotheses about the correlation between collinearity and cluster organization as observed in deuterostomes [23] stand true for protostomes.

Here, we present a comprehensive study of the genomic arrangement and expression of Hox genes in Brachiopoda, a lineage of the Spiralia whose origins date back to the Lower Cambrian [62]. Brachiopods are marine, sessile, filter-feeding animals. They are protected by two dorsoventral mineralized shells and reproduce by external fertilization, often developing through an intermediate, free-living larval stage [63]. In this study, we use two brachiopod species –the ‘articulate’ Terebratalia transversa and the ‘inarticulate’ Novocrania anomala– that respectively belong to the two major brachiopod lineages, thus allowing the reconstruction of putative ancestral characters for Brachiopoda as a whole (Figure 1A). By transcriptomic and genomic sequencing we demonstrate that the Hox complement consists of ten Hox genes in T. transversa and nine in N. anomala. In addition, the ten Hox genes of T. transversa are ordered in a split Hox cluster that differs from the genomic arrangement reported for the brachiopod Lingula anatina [personal communication, Luo and 64]. We show that Hox gene expression is restricted to the ‘trunk’ region of the larva, and is overall neither temporally nor spatially collinear. However, the genes pb (only in T. transversa), Hox3 and dfd show spatially collinear expression in the mesoderm of
both brachiopod species. Additionally, the Hox genes \textit{lab}, \textit{scr}, \textit{anp} and \textit{post1} appear to be associated with the development of two brachiopod features: the chaetae and the shell-forming epithelium. Altogether, our findings demonstrate that the presence of a split Hox cluster in the Brachiopoda is likely not associated with a temporally collinear expression of Hox genes, which differs from the hypothesized correlation between temporal collinearity and the retention of the vertebrate Hox cluster [14, 23, 59, 60] and suggests that alternative/additional genomic forces might shape Hox clusters during spiralian evolution, such as low genomic rearrangement frequency.

\section*{Results}

\textbf{The Hox gene complement of \textit{T. transversa} and \textit{N. anomala}}

Transcriptomic and genomic searches resulted in the identification of ten Hox genes in \textit{T. transversa}. In the brachiopod \textit{N. anomala}, we identified seven Hox genes in the transcriptome and two additional fragments corresponding to a Hox homeodomain in the draft genome assembly. Attempts to amplify and extend these two genomic sequences in the embryonic and larval transcriptome of \textit{N. anomala} failed, suggesting that these two Hox genes might be expressed only during metamorphosis and/or in the adult brachiopod. Maximum likelihood orthology analyses resolved the identity of the retrieved Hox genes (Figure supplementary 1). The ten Hox genes of \textit{T. transversa} were orthologous to \textit{labial (lab)}, \textit{proboscipedia (pb)}, \textit{Hox3}, \textit{deformed (dfd)}, \textit{sex combs reduced (scr)}, \textit{lox5}, \textit{antennapedia (anp)}, \textit{lox4}, \textit{post2} and \textit{post1}. The nine Hox genes identified in \textit{N. anomala} corresponded to \textit{lab}, \textit{pb}, \textit{Hox3}, \textit{dfd}, \textit{scr}, \textit{lox5}, \textit{anp}, \textit{lox4}, and \textit{post2}.

\textbf{Genomic organization of Hox genes in \textit{T. transversa} and \textit{N. anomala}}
We used the draft assemblies of *T. transversa* and *N. anomala* genomes to investigate the genomic arrangement of their Hox genes. In *T. transversa*, we identified three scaffolds containing Hox genes (Figure 1B). Scaffold A spanned 81.7 kb and contained *lab* and *pb* in a genomic region of 15.4 kb, flanked by other genes with no known linkage to the Hox cluster in other animals. Scaffold B was the longest (284.8 kb) and included *Hox3, dfd, scr, lox5, antp, lox4* and *post2*, in this order (Figure 1B) including the micro RNA *mir-10* between *dfd* and *scr*. As in scaffold A, other genes flanked the Hox genes, which occupied a genomic region of 76.2 kb. Finally, *post1* aligned to various short scaffolds. We could not recover any genomic linkage between the identified Hox genes in *N. anomala* due to the low contiguity (N50 of 3.5 kb) of the draft genome assembly. Altogether, these data demonstrate that *T. transversa* has a split Hox cluster broken into three sub-clusters, each of them with an organized arrangement. Importantly, the potential genomic disposition of these three sub-clusters is similar to that observed in other spiralians, such as *C. teleta* and *L. gigantea* (Figure 1C), which suggests that the lineage leading to the brachiopod *L. anatina* experienced genomic rearrangements that modified the ordered and linkage of the Hox genes.

**Hox gene expression in T. transversa**

To investigate the presence of temporal and/or spatial collinearity in the expression of the clustered Hox genes in *T. transversa*, we first performed whole-mount *in situ* hybridizations in embryos from blastula to late, competent larval stages (Figure 2).

**Anterior Hox genes**

The anterior Hox gene *lab* was first detected in the mid gastrula stage in two faint
bilaterally symmetrical dorsal ectodermal domains (Figure 2Ad, Ae). In late gastrulae, 
lab expression consisted of four dorsal ectodermal clusters that corresponded to the 
position where the chaetae sacs form (Figure 2Af, Ag). In early larva, the expression 
was strong and broad in the mantle lobe (Figure 2Ah, Ai), and in late larvae it became 
restricted to a few mantle cells adjacent to the chaetae sacs (Figure 2Ij, Ik). These 
cells do not co-localize with tropomyosin, which labels the muscular mesoderm of the 
larva (Figure 3A). This suggests that lab expressing cells are likely ectodermal, 
although we cannot exclude localization in non-muscular mesodermal derivates.

The Hox gene pb was first detected asymmetrically on one lateral of the ectoderm of 
the early gastrula (Figure 2Bb, Bc). In the mid gastrula, the ectodermal domain 
located dorsally and extended as a transversal stripe (Figure 2Bd, Be). Remarkably, 
this domain disappeared in late gastrula embryos, where pb was detected in the 
anterior mantle mesoderm (Figure 2Bf, Bg). This expression was kept in the early and 
late larva (Figure 2Bh–Bk; Figure 3B)

Hox3

The gene Hox3 was detected already in blastula embryos in a circle of asymmetric 
intensity around the gastral plate (Figure 2Ca). In early gastrulae, Hox3 is restricted to 
one half of the vegetal one, which is the prospective posterior side (Figure 2Cb, Cc). 
With axial elongation, Hox3 becomes expressed in the anterior mantle mesoderm and 
in the ventral ectoderm limiting the apical and mantle lobe (Figure 2Cd, Ce). This 
expression is maintained in late gastrula stages and in the early larva (Figure 2Cf–Ci). 
In the late larva, Hox3 is detected in part of ventral, internal mantle ectoderm and in 
the most anterior part of the pedicle mesoderm (Figure 2Cj, Ck; Figure 3C)
Central Hox genes

The Hox gene *dfd* was asymmetrically expressed on one side of the vegetal pole of the early gastrula of *T. transversa* (Figure 2Db, Dc). This expression was maintained in the mid gastrula, and corresponded to the most posterior region of the embryo (Figure 2Dd, De). In the late gastrula, *dfd* becomes strongly expressed in the posterior mesoderm (Figure 2Df, Dg). In the early larva, the expression remained in the pedicle mesoderm, but new domains in the posterior ectoderm and in the anterior, ventral pedicle ectoderm appear (Figure 2Dh, Di). These expression domains are also observed in the late larva (Figure 2Dj, Dk; Figure 3D).

The central Hox gene *scr* was first expressed in the medial dorsal ectoderm of the mid gastrula (Figure 2Ed, Ee). In late gastrula stages, the expression expanded towards the ventral side, forming a ring (Figure 2Ef, Eg). In the early larva, *scr* was detected in a ring encircling the most anterior ectoderm of the pedicle lobe and extending anteriorly on its dorsal side (Figure 2Eh, Ei). With the outgrowth of the mantle lobe in the late larva, the expression became restricted to the periostracum, the internal ectoderm of the mantle lobe that forms the shell (Figure 2Ej, Ek; Figure 3E).

The Hox gene *Lox5* is expressed on one side of the early gastrula (Figure 2Fb, Fc). During axial elongation, the expression became restricted to the most posterior ectoderm of the embryo (Figure 2Fd–Fg). This domain remained constant in larval stages, where it was expressed in the whole posterior ectoderm of the pedicle lobe (Figure 2Fh–Fk).
The *antp* gene is weakly detected at the mid gastrula stage, in one posterior ectodermal domain and one dorsal ectodermal patch (Figure 2Gd, Ge). In the late gastrula, the posterior expression is maintained and the dorsal domain extends ventrally, encircling the embryo (Figure 2Gf, Gg). These two domains remained in the larvae: the ectodermal anterior-most, ring-like domain localized to the periostracum, and the posterior domain limited to the most posterior tip of the larva (Figure 2Gh–Gk).

The Hox gene *Lox4* is first detected in the dorsal, posterior end of the late gastrula and early larva (Figure 2Hf–Hi). In the late larva, *Lox4* is expressed dorsally and posteriorly, although it is absent from the most posterior end (Figure 2Hj, Hk).

**Posterior Hox genes**

The posterior Hox gene *post2* was first detected in mid gastrula stages at the posterior tip of the embryo (Figure 2Id, Ie). This expression was maintained in late gastrulae (Figure 2If, Ig). In early larva, *post2* expression extended anteriorly and occupied the dorso-posterior midline of the pedicle lobe (Figure 2Ih, Ii). In late, competent larvae, *post2* was detected in a T-domain in the dorsal side of the pedicle ectoderm (Figure 2Ij, Ik).

The Hox gene *post1* was transiently detected in late gastrula stages in the four mesodermal chaetae sacs (Figure 2Jf, Jg).

We verified the absence of temporal collinearity in the expression of the Hox genes in *T. transversa* by quantitative real-time PCR and comparative stage-specific RNA-seq
data (Figure supplementary 2).

**Hox gene expression in N. anomala**

In order to infer potential ancestral Hox expression domains for the Brachiopoda, we investigated the expression of the nine Hox genes of *N. anomala* during embryogenesis and larval stages (Figure 4).

**Anterior Hox genes**

The Hox gene *lab* was first detected at the mid gastrula stage in three bilaterally symmetrical ectodermal cell clusters that appear to correlate with the presumptive site of chaetae sac formation (Figure 4Ad, Ae). The expression in the most posterior pair was stronger than in the two most anterior ones. This expression was maintained in the late gastrula (Figure 4Af, Ag). In larval stages, *lab* was detected in the two most anterior chaetae sacs of the mantle lobe (Figure 4Ah, Ai), expression that fainted in late larvae (Figure 4Aj, Ak).

The Hox gene *pb* was asymmetrically expressed already at blastula stages, in the region that putatively will rise to the most posterior body regions (Figure 4Ba). With the onset of gastrulation, the expression of *pb* extended around the vegetal pole, almost encircling the whole blastoporal rim (Figure 4Bb, Bc). During axial elongation, *pb* was first broadly expressed in the region that forms the mantle lobe (Figure 4Bd, Be) and later on the ventral mantle ectoderm of the late gastrula (Figure 4Bf, Bg). In early larvae, *pb* was detected in the anterior ventral mantle ectoderm (Figure 4Bh, Bi). This domain was not detected in late, competent larvae (Figure 4Bj, Bk).
**Hox3**

The Hox gene *Hox3* was asymmetrically detected around half of the vegetal pole of the early gastrulae (Figure 4Cb, Cc). In mid gastrulae, the expression almost encircled the whole posterior area and the blastoporal rim (Figure 4Cd). In addition, a domain in the mid-posterior mesoderm became evident (Figure 4Ce). By the end of the axial elongation, *Hox3* was strongly expressed in the posterior mesoderm and weakly in the ventral posterior mantle ectoderm (Figure 4 Cf, Cg). Noticeably, the posterior most ectoderm did not show expression of *Hox3*. This expression pattern was maintained in early and late larval stages (Figure 4Ch–Ck).

**Central Hox genes**

The central Hox gene *dfd* was first detected in the posterior ectodermal tip of mid gastrulae (Figure 4Dd, De). In late gastrula stages, *dfd* was expressed in the posterior ectodermal end (Figure 4Df) and in the posterior mesoderm (Figure 4Dg). Early larvae showed expression of *dfd* in the posterior mesoderm and posterior mantle ectoderm (Figure 4 Dh, Di). This expression remained in late larvae, although the most posterior ectodermal end was devoid of expression (Figure 4Dj, Dk).

The Hox gene *scr* was only detected in late larval stages, in a strong dorsal ectodermal domain (Figure 4Ej, Ek).

The gene *Lox5* was detected asymmetrically around half of the blastoporal rim in early gastrula stages (Figure 4Fb, Fc). During axial elongation, the expression progressively expanded around the blastoporal rim (Figure 4Fd, Fe) and limited to the
ventral midline (Figure 4Ff, Fg). In the larvae, *Lox5* was expressed in the ventral, posterior-most midline (Figure 4Fh–Fk).

The Hox gene *antp* was first expressed asymmetrically in one lateral side of the early gastrula (Figure 4Gj, Gk). In the mid gastrula, *antp* was detected in the dorsal ectodermal mantle in a cross configuration: dorsal midline and the mantle cells closer to the apical-mantle lobe boundary (Figure 4Gd, Ge). In late gastrulae, *antp* was only expressed in a mid-dorsal ectodermal region (Figure 4Gf, Gg). This expression pattern was also observed in early larval stages, although the size of the domain reduced (Figure 4Gh, Gi). In late larvae, antp was detected in a small mid-dorsal patch and a weak ventro-posterior ectodermal domain (Figure 4Gj, Gk).

We could neither identify nor amplify *Lox4* in a transcriptome and cDNA obtained from mixed embryonic and larval stages, suggesting that either it is very transiently and weakly expressed during embryogenesis or it is only expressed in later stages (metamorphosis and adulthood).

*Posterior Hox genes*

The only posterior Hox gene present in *N. anomala*, *post2*, could not be amplified in cDNA obtained from mixed embryonic and larval stages, suggesting that it is not expressed—or at least expressed at really low levels—during these stages of the life cycle. The absence of larval expression of *Lox4* and *post2* could be related to the lack of the pedicle lobe of craniiform brachiopod larva, which is a characteristics of the lineage [65, 66].
Discussion

The brachiopod Hox complement and the evolution of Hox genes in Spiralia

Our findings on T. transversa and N. anomala reveal an ancestral brachiopod Hox gene complement consistent with what has been hypothesized to be ancestral for Spiralia and Lophotrochozoa on the basis of degenerate PCR surveys [15, 67-69]. This ancient complement comprises eight Hox genes – lab, pb, Hox3, Dfd, Scr, Lox5, Lox4 and Post2 – and has been confirmed by genomic sequencing of representative annelids and mollusks [16, 22, 47], rotifers and platyhelminthes [21, 43-45] and the linguliform brachiopod L. anatina [64]. While T. transversa and L. anatina (N. Satoh and Y.-J. Luo, personal communication) have retained this ancestral Hox complement, N. anomala has apparently lost Post1 (Figure 1).

Our genomic information shows that the Hox cluster of T. transversa is split in three parts, with lab and pb separate from the major cluster and Post1 also on a separate scaffold (Figure 1B). Overall, the cluster extends over 100 kb, which is significantly shorter than those of other lophotrochozoans, such as C. teleta (~345kb) [46] and L. gigantea (~455 kb) [16]. Its compact size is related to short intergenic regions and introns, comparable to the situation observed in vertebrate Hox clusters [23]. The order and orientation of the Hox genes in T. transversa is preserved and more organized than in the Hox cluster reported for the brachiopod L. anatina, which exhibits a genomic rearrangement that placed a portion of the cluster upstream lab and in reverse orientation [64]. Indeed, the split Hox clusters reported so far in lophotrochozoan taxa exhibit all different conformations, indicating that lineage-specific genomic events have shaped Hox gene clusters in Spiralia.
Non-collinearity of Hox expression in T. transversa despite the presence of a split cluster

The analysis of Hox clustering in different animal species together with the temporal and spatial expression patterns of their Hox genes grounded the hypotheses that the regulatory elements required for their collinearity—mostly temporal—maintain the clustered organization of the vertebrate Hox genes and possibly other animals [13, 23, 59-61, 70, 71] (Figure supplementary 3). Although there are cases in which spatial collinearity is displayed in the absence of a cluster, as in the appendicularian chordate O. dioica [24], all investigated clustered Hox genes show at least one type of collinearity that could account for their genomic organization [23, 61] (Figure 6).

Since there are exceptions to the spatial collinearity in vertebrates, for instance Hoxa2 and Hoxb2 are expressed more anteriorly than Hoxl genes in the vertebrate hindbrain [72], temporal collinearity is seen as a manifestation of Hox clustering. But whether temporal collinearity is the agent keeping the cluster together, e.g. through enhancer sharing [73], is still subject of debate.

Within Spiralia, this evolutionary scenario appears to be supported by the staggered temporal and spatial expression of the Hox genes in the split cluster of the annelid C. teleta [46]. In the other investigated spiralian species, there is only either genomic information (e.g. the mollusks L. gigantea and C. gigas) or expression analysis (e.g. the mollusks G. varia, Haliotis asinina) [16, 47, 52, 58]. Most of these gene expression studies have demonstrated coordinated spatial or temporal expression of Hox genes along the anteroposterior axis of the animal [48, 49, 74] or in organ systems, such the nervous system [52, 58]. However, the absence of studies that can reveal a correlation between the expression of Hox genes and their genomic
organization in these animals hampers the reconstruction of the putative mechanisms that preserve Hox clusters in Lophotrochozoa, and thus prevent generalizations about possible scenarios of Hox cluster evolution across all animals.

Our findings robustly demonstrate that the Hox genes of the split Hox cluster of T. transversa overall show neither strictly spatial nor temporal collinearity (Figures 2, 3), and lack quantitative collinearity [61], as it has been shown for example in mouse [75]. These observations are also supported by the absence of a coordinated spatial and temporal expression of the Hox genes in N. anomalala (Figure 4). Although a general trend of spatial collinearity is present (e.g. the posterior Hox genes are expressed in posterior tissues), the early expression of Hox3 breaks temporal collinearity in T. transversa, while it is pb that becomes first expressed in N. anomalala. In both species, the gene Lox5 is also expressed before Scr, as it is also the case in the annelid N. virens [74]. Ectodermal spatial collinearity is absent in the two brachiopods even when considering the future location of the larval tissues after metamorphosis [76, 77]. The most anterior class gene lab is exclusively expressed in the chaetae of T. transversa and N. anomalala, and thus is not affiliated with anterior neural or foregut tissues as in other lophotrochozoans, such as annelids [46, 78]. Similarly, the most posterior Hox gene, Post1, is very transiently expressed in the chaetae sacs, which occupy a mid-position in the larval body. We only detected a strict spatial collinearity in the staggered expression of the Hox genes pb, Hox3 and Dfd along the anterior-posterior axis of the developing larval mesoderm in both T. transversa and N. anomalala (Figure 5).

Altogether, the absence of a strict temporal and spatial collinearity in the brachiopod
*T. transversa*, albeit the presence of a split Hox cluster, indicates that temporal collinearity is likely not the underlying factor keeping spiralian Hox genes clustered, as it seems to be the case in vertebrates [14, 23, 59-61]. Therefore, alternative mechanisms might need to be considered. In this regard, why do Hox clusters split in different positions between related species, as seen for instance in brachiopods (this study) and drosophilids [79], but still display similar expression profiles? This might indicate that the control of expression in large split Hox clusters relies more on gene-specific short-range transcriptional control than on a global, coordinated regulation, as seen in the small Hox vertebrate clusters [23, 75, 80]. The conservation of Hox clusters in many animals could then be a consequence of the general conservation of syntenic relationships in their genomes (Figure 6). Our findings thus highlight the necessity of further detailed structure-function analyses of spiralian Hox clusters to better understand the intricate evolution of the genomic organization and regulation of Hox genes in metazoans.

**Recruitment of Hox genes for patterning lophotrochozoan chaetae and shell fields**

The bristle-like chaetae (or setae) of annelids and brachiopods, and shell valves in mollusks and brachiopods are the most prominent hard tissues found in lophotrochozoan spiralian [63] and provide fossilized hallmarks of the Cambrian explosion [81]. It has been already recognized that the ultrastructural morphology of the brachiopod and annelid chaetae is nearly identical [82-84] and with the placement of brachiopods as close relatives of annelids and mollusks [85], the homology of these structures appeared more likely [86]. In this context, the anterior Hox gene *lab* is expressed in the chaetae of *Chaetopterus* sp. [26] and *Post1* is expressed in the chaetae of *C. teleta, P. dumerilii* and *N. virens* [46, 74]. Our results show that
similarly, \textit{lab} and \textit{Post1} are expressed specifically in the chaetal sacs of the brachiopods \textit{T. transversa} and \textit{N. anomala} (Figures 2, 4) and follow the different arrangement of the chaetae in both species. Further evidence of a common, and probably homologous, molecular profile comes from the expression of the homeodomain gene \textit{Aristaless-like (Arx)} and the zinc finger \textit{Zic}. These genes are expressed at each chaetae sac territory in the \textit{Platynereis} larva [87], in \textit{Capitella teleta} [88], and also in the region of the forming chaetae sac territories in \textit{T. transversa} (Figure supplementary 4). Therefore, the expression of the Hox genes \textit{lab} and \textit{Post1} and the homeodomain gene \textit{Arx} indicate that similar molecular signature underlays the development of chaetae in annelids and brachiopods. This, together with the evident and striking morphological similarities shared by brachiopod and annelid chaetae, support considering these two structures homologous, and thus, common lophotrochozoan innovations. This would be consistent with placing the iconic Cambrian fossil \textit{Wiwaxia}, which contains chaetae, as a stem group lophotrochozoan [89].

The protective shell is a mineralized tissue present in brachiopods and mollusks. In the gastropod mollusk \textit{G. varia}, the Hox genes \textit{lab}, \textit{Post1} and \textit{Post2} are first expressed in the shell field, and later is \textit{Dfd} [57]. In \textit{H. asinina} also \textit{lab} and \textit{Post2} are related to shell formation [52]. In brachiopods, \textit{Dfd} is associated to the adult shell in \textit{L. anatina} [64]. During embryogenesis of \textit{T. transversa} and \textit{N. anomala}, however, only \textit{Scr} and \textit{Antp} are expressed in the shell fields, but not \textit{lab} or \textit{Post1}, which are expressed in the chaetae sacs. This could support the homology of the chitin-network that is formed at the onset of brachiopod and mollusk shell fields. However, the different deployment of Hox genes in the shell fields of brachiopods and mollusks
might indicate that these genes do not have an ancient role in the specification of the shell-forming epithelium. However, their consistent deployment during shell development might reflect a more general, conserved role in shaping the shell fields according to their position along the anterior posterior axis.

**Conclusions**

In this study, we characterize the Hox gene complement of the brachiopods *T. transversa* and *N. anomala*, and demonstrate the last common ancestor to all brachiopods likely had ten Hox genes (*lab, pb, Hox3, dfd, scr, Lox5, antp, Lox4, post2, post1*). Noticeably, brachiopod Hox genes do not show global temporal and spatial collinearity, albeit *T. transversa* exhibits an ordered, split Hox cluster. Only the genes *pb* (in *T. transversa*), *Hox3* and *dfd* (in both brachiopods) show spatial collinearity in the ‘trunk’ mesoderm. In addition, the Hox genes *lab* and *post1*, as well as the homeobox *Arx*, are expressed in the developing chaetae, as also described for other annelid species [46, 53, 74]. These molecular similarities, together with evident morphological resemblances [83], support considering brachiopod and annelid chaetae homologous structures and reinforce considering the fossil *Wiwaxia* as a stem group lophotrochozoan [89]. Altogether, our findings challenge a scenario in which temporal collinearity is the major force preserving Hox clusters [12, 14, 23, 60, 61], and indicate that alternative/additional genomic mechanisms might account for the great diversity of Hox gene arrangements observed in extant animals.

**Material and Methods**

*Animal cultures*

Gravid adults of *Terebratalia transversa* (Sowerby, 1846) were collected around San
Juan Island, Washington, USA and *Novocrania anomala* (Müller, 1776) around Bergen, Norway. Animal husbandry, fertilization and larval culture were conducted following previously published protocols [90-92].

**Hox cluster reconstruction in *T. transversa* and *N. anomala***

Male gonads of *T. transversa* and *N. anomala* were preserved in RNAlater (Life Technologies) for further genomic DNA (gDNA) isolation. Paired end and mate pair libraries of 2 kb and 5 kb insert sizes of *T. transversa* gDNA were sequenced using an Illumina HiSeq2000 platform. First we trimmed Illumina adapters with Cutadapt 1.4.2 [93]. Then, we assembled the paired end reads into contigs, scaffolded the assembly with the mate pair reads, and closed the gaps using Platanus 1.21 [94]. The genomic scaffolds of *T. transversa* including Hox genes are published on GenBank with the accession numbers KX372775 and KX372776. Paired end libraries of *N. anomala* gDNA were sequenced using an Illumina HiSeq2000 platform. We removed Illumina adapters as above and assembled the paired end reads with MaSuRCA 2.2.1 [95].

**Gene isolation**

Pooled samples of *T. transversa* and *N. anomala* embryos at different developmental stages (cleavage, blastula, gastrula, mid gastrula, late gastrula, early larva, and late/competent larva) were used for RNA isolation and Illumina sequencing (NCBI SRA; *T. transversa* accession SRX1307070, *N. anomala* accession SRX1343816). We trimmed adapters and low quality reads from the raw data with Trimmomatic 0.32 [96] and assembled the reads with Trinity 2.0.6 [97]. Hox genes were identified by BLAST searches on these transcriptomes and their respective draft genomes (see above). First-strand cDNA template (SuperScript™, Life Technologies) of mixed
embryonic stages was used for gene-specific PCR. RACE cDNA of mixed embryonic stages was constructed with SMARTer RACE cDNA Amplification Kit (Clontech) and used to amplify gene ends when necessary. All fragments were cloned into the pGEM-T-Easy vector (Promega) and sequenced at the University of Bergen sequencing facility. T. transversa and N. anomala Hox gene sequences were uploaded to GenBank (accession numbers KX372756–KX372774).

**Orthology analyses**

Hox gene sequences of a representative selection of bilaterian lineages (Supplementary Table S1) were aligned with MAFFT v.7 [98]. The multiple sequence alignment, which is available upon request, was trimmed to include the 60 amino acids of the homeodomain. ProtTest v.3 [99] was used to determine the best fitting evolutionary model (LG+G+I). Orthology analyses were conducted with RAxML v.8.2.6 [100] using the autoMRE option. The resulting trees were edited with FigTree and Illustrator CS6 (Adobe).

**Gene expression analyses**

T. transversa and N. anomala embryos at different embryonic and larval stages were fixed in 4% paraformaldehyde in sea water for 1 h at room temperature. All larval stages were relaxed in 7.4% magnesium chloride for 10 min before fixation. Fixed samples were washed several times in phosphate buffer saline (PBS) with 0.1% tween-20 before dehydration through a graded methanol series and storage in 100% methanol at -20 °C. Single colorimetric whole mount in situ hybridization were carried out following an established protocol (detailed protocol available in Protocol Exchange: doi:10.1038/nprot.2008.201) [101, 102]. Double fluorescent in situ
hybridizations were conducted as described elsewhere [103]. Representative stained specimens were imaged with bright field Nomarski optics using an Axiocam HRc connected to an Axioscope Ax10 (Zeiss). Fluorescently labeled embryos were mounted in Murray’s clearing reagent (benzyl alcohol: benzyl benzoate, 1:2) and imaged under a SP5 confocal laser-scanning microscope (Leica). Images and confocal z-stacks were processed with Fiji and Photoshop CS6 (Adobe) and figure panels assembled with Illustrator CS6 (Adobe). Contrast and brightness were always adjusted to the whole image, and not to parts of it.

Quantitative Hox gene expression in T. transversa

Thousands of synchronous T. transversa embryos collected at 14 specific stages (oocytes, 8h mid blastula, 19h late blastula, 24h moving late blastula, 26h early gastrula, 37h asymmetric gastrula, 51h bilateral gastrula, 59h bilobed, 68h trilobed, 82h early larva (first chaetae visible), 98h late larva (long chaetae, eye spots), 131h competent larva, 1d juvenile, 2d juvenile) were pooled together and preserved in RNAlater (Life Technologies). Total RNA was isolated with Trizol Reagent (Life Technologies). For quantitative real time PCR, total RNA was DNase treated and preserved at -80 ºC. Gene specific primers bordering an intron splice-site and defining an amplicon of 80-150 bp sizes were designed for each gene (Supplementary Table S2). Expression levels of two technical replicates performed in two biological replicates were calculated based on absolute quantification units. For comparative stage-specific transcriptomic analyses, total RNA was used for constructing Illumina single end libraries and sequenced in four lanes of a HiSeq 2000 platform. Samples were randomized between the lanes. To estimate the abundance of transcripts per stage, we mapped the single end reads to the transcriptome of T. transversa with
Bowtie, calculated expression levels with RSEM, and generated a matrix with TMM normalization across samples by running Trinity’s utility scripts. Expression levels obtained after quantitative real-time PCR and comparative stage-specific transcriptomics were plotted with R.

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Author contributions

A.H. designed the study. A.H., S.M.S. and J.M.M.D. conducted the gene isolation and *in situ* hybridization studies. J.M.M.D. performed the gene orthology analyses. A.H., J.M.M.D., Y.P. and B.V. collected the stage-specific samples of *T. transversa* embryos. A.B. and J.M.M.D. isolated the genomic DNA of *T. transversa* and *N. anomala*. J.M.M.D. and B.V. did the draft genome assemblies and S.M.S. analyzed the Hox genomic organization. J.M.M.D. performed the stage-specific RNA isolations; A.B. did the quantitative real time PCR experiments, and B.V. conducted
the analysis of the stage-specific transcriptomes. A.H. and J.M.M.D. wrote the manuscript. All authors discussed the data and edited the text.

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Figures

Figure 1. Genomic organization of Hox genes in Brachiopoda. (A) Images of an adult *T. transversa* and *N. anomala*, and phylogenetic position of these species within Brachiopoda and Lophotrochozoa. (B) The ten Hox genes of *T. transversa* are ordered along three genomic scaffolds and are flanked by external genes (vertical lines; gene orthology is based on best blast hit). Thus, *T. transversa* has a split Hox cluster composed of three sub-clusters. No predicted ORFs were identified between the Hox genes in scaffold A and B. A colored box represents each Hox gene, and below each box there is the direction of transcription and the exon-intron composition. The genomic regions containing Hox genes are represented in scale. (C) The genomic organization of brachiopod Hox genes in a phylogenetic context (adapted from [22]).
The genomic order of Hox genes in *T. transversa* is similar to that observed in other spiralians (e.g. *Capitella teleta* and *Lottia gigantea*), which suggests that the translocation of the posterior Hox cluster upstream to lab is a lineage-specific feature of *L. anatina* (in *T. transversa* and *L. anatina* the arrows below the genes show the direction of transcription; conformation of the Hox cluster in *L. anatina* was kindly provided by N. Satoh and Y.-J. Luo). The low contiguity of the draft genome assembly of *N. anomala* hampered recovering genomic linkages between the identified Hox genes. Each ortholog group is represented by a particular color.
Figure 2. Expression of Hox genes in *T. transversa*. (A–J) Whole mount *in situ* hybridization of each Hox gene during embryonic and larval stages in *T. transversa*. The Hox genes *lab* and *post1* are expressed during chaetae formation. The genes *pb*, *Hox3* and *dfd* are collinearly expressed along the mantle and pedicle mesoderm. The Hox genes *scr* and *antp* are expressed in the periostracum, the shell-forming epithelium. *Lox5*, *Lox4* and *post2* are expressed in the posterior ectoderm of the pedicle lobe. See main text for a detailed description of each expression pattern. Black arrowheads indicate expression in the chaetae sacs. Orange arrowheads highlight mesodermal expression. Green arrowheads indicate expression in the periostracum.
The genomic organization of the Hox genes is shown on the left. On top, schematic representations of each analyzed developmental stage on its respective perspective. In these schemes, the blue area represents the mesoderm. Drawings are not to scale. The red line indicates the onset of expression of each Hox gene based on in situ hybridization data. The blastula stage is a lateral view (inset is a vegetal view). The other stages are in a lateral view (left column) and dorsoventral view (right column). The asterisk demarcates the animal/anterior pole. al, apical lobe; bp, blastopore; ch, chaetae; em, endomesoderm; gp, gastral plate; gu, gut; me, mesoderm; ml, mantle lobe; mo, mouth; pl, pedicle lobe.
Figure 3. Hox expression in mesoderm and periostracum of *T. transversa*. (A–E)

Double fluorescent *in situ* hybridization of *lab, pb, Hox3, dfd* and *scr* with tropomyosin (Tropo, in green) in late larval stages of *T. transversa*. (A) The gene *lab* is expressed in relation to the chaetae sacs, but does not overlap with the tropomyosin-expressing mesoderm. (B–D) The Hox genes *pb, Hox3* and *Dfd* show spatial collinearity along the mantle and pedicle mesoderm. (E) The gene *scr* is expressed in the periostracum, which is the epithelium that forms the shell.
Figure 4. Expression of Hox genes in *N. anomalae*. (A–G) Whole mount *in situ* hybridization of the Hox genes during embryonic and larval stages in *N. anomalae*. The gene *lab* is expressed in the chaetae. The Hox genes *Hox3* and *dfd* are collinearly expressed in the mantle mesoderm. The genes *scr* and *antp* are expressed in the prospective shell-forming epithelium. The genes *pb* and *Lox5* are detected in the ectoderm of the mantle lobe. The genes *Lox4* and *post2* were not detected in transcriptomes and cDNA during embryonic stages. See main text for a detailed description of each expression pattern. Black arrowheads indicate expression in the chaetae sacs. Orange arrowheads highlight mesodermal expression. Green arrowheads indicate expression in the periostracum. On top, schematic representations of each analyzed developmental stage on its respective perspective. In these schemes, the blue area represents the mesoderm. Drawings are not to scale. The red line indicates the onset of expression of each Hox gene based on *in situ* hybridization data. The blastula
stage is a lateral view (inset is a vegetal view). The other stages are in a lateral view (left column) and dorsoventral view (right column). The asterisk demarcates the animal/anterior pole. al, apical lobe; bp, blastopore; ch, chaetae; em, endomesoderm; gp, gastral plate; gu, gut; me, mesoderm; ml, mantle lobe; mo, mouth.
Figure 5. Summary of Hox gene expression in *T. transversa* and *N. anomala*. (A, B) Schematic drawings of late larvae of *T. transversa* and *N. anomala* depicting the expression of each Hox gene. The Hox genes *pb* (not in *N. anomala*), *Hox3* and *dfd* show staggered expression, at least in one of their domains, associated with the mesoderm (light blue box). In both brachiopods, the genes *scr* and *antp* are expressed in the periostracum, or the shell-forming epithelium (red boxes) and lab and post1 are associated to the developing chaetae (green boxes; asterisk in *post1*: *post1* is expressed in the chaetae only during late embryonic stages, not in the mature larva, and only in *T. transversa*). The expression of *Lox4* and *post2* in *N. anomala* could not be determined in this study. The gene *post1* is missing in *N. anomala*. Drawings are not to scale.
Figure 6. Hox gene cluster evolution. The absence of temporal collinearity in the
ordered, split Hox cluster of the brachiopod *T. transversa* weakens the scenario that
temporal collinearity is a major force keeping Hox genes together. Alternatively, we
propose that the ancestral Hox cluster was organized and regulated by short-range
regulators, which elaborated a collinear expression, as observed in amphioxus
(temporal and spatial collinearity) and hemichordates (only spatial collinearity).
Direct selective pressures could have promoted genomic changes that led to the
evolution of compact, tightly regulated, collinear Hox clusters (e.g. in vertebrates). In
other animal lineages, the original organized Hox cluster would become broken,
rearranged or atomized indirectly, as a result of general genome-wide events. In these
cases, the conservation/loss of the ancestral regulatory elements, together with the
evolution of new ones associated with novel morphologies, would have influence the
collinear expression of Hox genes. Under this scenario, collinearity is only one of
multiple direct or indirect factors affecting Hox clustering.