An analysis of segmentation dynamics throughout embryogenesis in the centipede Strigamia maritima

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Abstract

Background: Most segmented animals add segments sequentially as the animal grows. In vertebrates, segment patterning depends on oscillations of gene expression coordinated as travelling waves in the posterior, unsegmented mesoderm. Recently, waves of segmentation gene expression have been clearly documented in insects. However, it remains unclear whether cyclic gene activity is widespread across arthropods, and possibly ancestral among segmented animals. Previous studies have suggested that a segmentation oscillator may exist in Strigamia, an arthropod only distantly related to insects, but further evidence is needed to document this.

Results: Using the genes even skipped and Delta as representative of genes involved in segment patterning in insects and in vertebrates, respectively, we have carried out a detailed analysis of the spatio-temporal dynamics of gene expression throughout the process of segment patterning in Strigamia. We show that a segmentation clock is involved in segment formation: most segments are generated by cycles of dynamic gene activity that generate a pattern of double segment periodicity, which is only later resolved to the definitive single segment pattern. However, not all segments are generated by this process. The most posterior segments are added individually from a localized sub-terminal area of the embryo, without prior pair-rule patterning.

Conclusions: Our data suggest that dynamic patterning of gene expression may be widespread among the arthropods, but that a single network of segmentation genes can generate either oscillatory behavior at pair-rule periodicity or direct single segment patterning, at different stages of embryogenesis.

Keywords: Segmentation clock, Oscillation, Pair-rule genes, Single segment patterning, Arthropod, Evolution

Background

Most arthropods, like most other segmented animals, couple the processes of segmentation and growth, adding segments progressively to the posterior of the body. This process is termed short germ or sequential segmentation, to distinguish it from the long germ mode of segmentation seen in Drosophila, where all segments are generated simultaneously by a cascade of genetic interactions taking place in the blastoderm [1].

Despite considerable effort in recent years, we still understand little of the mechanisms that drive segmentation in sequentially segmenting arthropods. Particular interest has focused on the nature of the so-called “growth zone”, or “segment addition zone”, a region of apparently undifferentiated tissue at the posterior of the embryo from which segments emerge. In all of the short germ arthropods so far examined, homologues of the Drosophila pair-rule segmentation genes are expressed in this region, often in rather broad domains around the site of invagination of the proctodeum [2]. Patterned stripes of gene expression emerge from this posterior domain, through the apparent repression of expression in the interstripe regions. One major concern of this paper is the nature of the patterning process in this posterior region, and in particular, whether dynamic oscillations of gene expression precede segment patterning. Such cyclic gene expression has recently been documented unambiguously in one insect [3,4], suggesting parallels between segment patterning in short germ insects, and somite patterning in vertebrates [5].
In some arthropods (for example, the myriapods Lithobius and Glomeris, the spider Cupiennius), this initial pattern appears to have a single segment periodicity, as defined by the subsequent expression of engrailed and other segment polarity genes in register with these primary stripes, and by the appearance of definitive segment morphology shortly afterwards [6-8]. In other cases, including several well documented cases in short germ insects (Tribolium, Schistocerca), the earliest stripes to resolve are at a double segment or “pair-rule” periodicity, just as they are in Drosophila [9,10]. The single segment repeat is then generated either by the subsequent splitting of these stripes, or by the intercalation of secondary stripes between them, before the appearance of definitive segment morphology. We have previously documented this pattern for homologues of the pair-rule genes, including even-skipped, hairy, runt and oddskipped, and also for genes of the Notch signaling pathway - most strikingly, the gene encoding the Notch ligand Delta [12,13].

In all of these cases, the early “pair-rule” patterning of the genes extends as complete rings around a focal point: initially this focus is marked by an area of reduced cell density that we interpret as being the blastopore, based on morphology [14] and on gene expression data (Jack Green, unpublished data). Later, the proctodeum will invaginate close to this point, and the pattern becomes centered on the proctodeum. Within the area closest to the center of this pattern, gene expression appears highly variable, even in embryos of similar age. We have proposed [11,12] that gene expression may be dynamic in this “peri-proctodeal” area, with cells near the proctodeum showing oscillating gene activity, and a pattern emerging through the fixation of a travelling wave of gene expression to give rise to a static pattern of cell states in the transition zone, where the germ band emerges, before a further round of gene interactions defines the single segment pattern.

However, interpretation of the observed gene expression patterns is complicated by the accompanying movement of cells, as the posterior disc converges to form the germ band, and as anterior parts of the germ band condense towards the head [14]. It has not previously been clear to what extent patterns of transcript accumulation in the posterior disc and emerging germ band reflect dynamic aspects of gene expression, and to what extent they reflect these cell movements.

In this paper we document more robustly the evidence for dynamic patterns, using embryos that are staged by independent criteria to different phases of a single cycle of gene activity. We define more precisely the limits of dynamic gene expression, concluding that this is limited to the region within, and including, the first resolved “pair-rule” stripe, but that the movement of the remainder of the pattern is largely the result of cell movement.

We extend our observations to earlier stages of patterning, when the head segments are being defined. Using the even-skipped genes as representative of the pair-rule gene network, and the Delta gene as representative of the Notch signaling pathway, we show that both of these pathways are apparently involved in the patterning of all segments from the intercalary back, and that the earliest aspects of segment patterning appear to involve dynamic gene activity qualitatively similar to the patterning of the trunk.

We also show that the relative timing of double and single segment patterning shifts as more trunk segments are added, until at about the time that the 39th segment is patterned, the oscillation that generates double segment periodicity appears to shut off; the last 10 or so segments resolve singly from a domain of ubiquitous and continuous posterior even-skipped expression.

Results

Segmentation in Strigamia proceeds from anterior to posterior [14-16]. Five segments of the head appear first during stage 3 and then, after a short pause, leg-bearing segments (LBS) appear in sequence, initially at a uniform rate of 1 segment every 3.2 hours until about 39 LBS are visible at the end of stage 4. Thereafter, segments are added much more slowly, with the process pausing completely during the movements of germ band flexure in stage 6. The final leg-bearing segment is not demarcated until shortly before hatching.

In this paper, we have used the genes Delta, even-skipped (eve1 and eve2) and engrailed to monitor the progress of this molecular patterning. Delta is a marker for cell interaction processes that are known to be important in the coordination of vertebrate segmentation, as well as in many other developmental processes; eve is a member of the primary pair-rule gene set that generates the first periodic, double segment pattern in Drosophila. It is also expressed later during Drosophila segmentation, with single segment periodicity. Engrailed is a widely used marker for the definitive segment pattern. It is expressed in the posterior part of each segment throughout the arthropods.

In Strigamia, both Delta and eve are expressed with a primary double segment periodicity as concentric rings...
around the site of invagination of the proctodeum, but out of phase with one another [12]. As the segmentation process continues, the appearance of intercalary stripes of eve1 and Delta within the forming germ band defines the single segment repeat (Figures 1 and 2), and shortly thereafter, engrailed is activated in every segment [15]. A second even-skipped gene, eve2, is co-expressed with eve1 during the primary, double segment phase, but is never activated in segmental stripes [13]. It ceases to be expressed as segments mature, except that it is transiently expressed specifically as a stripe in the antennal segment (see below).

The dynamics of segmentation gene expression
These patterns of gene expression that presage segment formation arise near the posterior of the embryo, and then progressively move anteriorly as further segments are added behind them. This is visible, for example, in the series of embryos presented in Figure 2, where the primary Delta expression corresponding to LBS 15 arises as a patch adjacent to the proctodeum, becomes the first ring in slightly older embryos, and is then displaced anteriorly as the next ring resolves behind it.

In part, this movement results from a condensation of the whole surface epithelium of the egg towards the
anterior. This is most obvious in the germ band, as the head extends forward and the first formed segments become relatively smaller (Figure 3; see also [14]). However, this movement also affects the territory outside the germ band. This is clear from the expression of a number of markers, including *decapentaplegic (dpp)* (Figure 1), which is expressed in a double segment pattern that mirrors earlier segmentation markers. This expression persists in the extra-embryonic territory after the resolution of segments in the germ band. The whole pattern can be seen to extend anteriorly in continuity with the morphological segments of the germ band.

We believe that this tissue movement may account for most of the movement of the gene expression patterns after the resolution of the first or second well-resolved rings around the proctodeum. It is presumably driven by

![Figure 2 Dynamics of Delta across two cycles of expression during stage 3](image)

Embryos are aligned from top to bottom according to developmental age, indicated on the left by the last formed morphological segment visible in each embryo (mx2: second maxilla, mxp: maxilliped, 1LBS: first leg-bearing segment). On each row the same embryo is viewed laterally with fluorescent nuclear staining (SYBR green) to show the morphology (A1, B1, C1, D1, E1), in normal light from the same view (A2, B2, C2, D2, E2), and from a posterior orthogonal view (A3, B3, C3, D3, E3). Within this period, Delta expression around the proctodeum is dynamic, showing an oscillation of expression in each region of the peri-proctodeal area. Arrows of the same color mark a single cycle of expression, from its initiation as bilateral patches around the proctodeum (black asterisk) to its resolution as a primary pair-rule stripe. Magenta arrowhead marks the second LBS in lateral views: red asterisk marks artefactual breaks in the epithelium. Scale bar: 400 μm.
cell proliferation in the posterior of the embryo, with space for the movement being generated by the continuing condensation of the head and anterior germ band, which continues throughout the period of segment addition.

However, this condensation of the germ band and surrounding epithelium is clearly not sufficient to account for the very dynamic patterns of gene expression seen in the region immediately around the site of invagination of the proctodeum (Figures 1, 2, 3, 4). We here define this region as the “peri-proctodeal region” to distinguish it from the larger area of unsegmented tissue posterior to the emergence of segments. We have previously suggested that the appearance of the double segment pre-pattern in this peri-proctodeal region reflects a dynamic process of gene expression akin to that observed in the vertebrate pre-somitotic mesoderm [11,12].

To demonstrate the generation of this pattern more precisely, Figure 4 illustrates four closely staged embryos from the same clutch that have been put into a developmental series by the number of leg-bearing segments (LBS) morphologically visible, and by the number of intercalated eve1 stripes (panels A-D). Embryo A (six LBS, intercalated eve1 stripe 17 visible), has a relatively well resolved primary ring of eve1 expression corresponding to LBS 26, but little or no eve1 expression within this ring, closer to the proctodeum. The slightly older embryo B (also six LBS, but with intercalated stripe 19 visible) shows eve1 expression in two bilateral patches adjacent to the proctodeum. This is the beginning of primary eve1 ring 28. In embryo C, (with seven and possibly an eighth LBS visible), eve1 expression now fills most of the area within ring 26, and in embryo D (eight LBS clearly visible, but still intercalated eve1 stripe 19), the central region closest to the proctodeum is now
clearing of eve1 expression, defining the beginning of primary eve1 stripe 28.

In Figure 4, panel E, embryo A has been montaged with bisected images of embryos B, C and D. These composite images suggest that primary eve1 ring 26 resolves by the loss of expression from its internal margin, closer to the proctodeum, and by a limited expansion of the outer part of the ring. This expansion will presumably continue as subsequent primary rings appear, but any later expansion must be much slower than the resolution of the primary rings: on the timescale of this series, (and accepting the limitations of these composite embryo montages) there is no perceptible shift in the position of the older rings (for example, ring 24 marked by arrows; see also in Additional file 1: Movie 1).

At a low frequency in our population of eggs, we observe embryos that resemble these photographic montages. Embryo F in Figure 4, for example, is a slightly abnormal embryo, in which the development of the eve pattern (eve2 in this case) is slightly asymmetric on the two sides. We suggest that this represents an embryo in which the right side is slightly ahead of the left in the cycle of segment generation, naturally reproducing the situation seen in the first montage.
The appearance of primary rings of Delta shows this same pattern of cyclic activation initiating from bilateral patches (Figure 2). For eve and Delta, the fully resolved primary rings are out of phase (Figure 5); the expanding patterns close to the proctodeum do overlap transiently, but these too show different phases of oscillation, as illustrated in Figure 5 by two embryos about a half cycle apart (A, B) and by a bilaterally asymmetric embryo that shows a similar discrepancy between left and right sides (Figure 5C). Intercalated stripes, when they appear, are directly superposed - eve intercalated stripes within primary Delta rings, and vice-versa (Figure 5A1).

Our model for gene expression in the periproctodeal region implies that cells are cycling between expressing and non-expressing states. To examine this in more detail, we have carried out hybridizations with a probe from the Delta gene that contains a largely (>95%) intron sequence (Additional file 2: Figure S1) to monitor the distribution of nascent transcripts, and hence to reveal where transcription is most active (Figure 6). This probe reveals intense “nuclear dots” of nascent Delta transcripts in cells of the periproctodeal region, suggesting that Delta is strongly transcribed in this region. In the same embryos, the nuclear dots are much less frequent, and much less intense, in the resolved rings of Delta transcript accumulating further from the proctodeum. In these rings, the relatively weak hybridization of exon sequences in the probe to cytoplasmic transcript dominates the nuclear signal, suggesting that rates of transcription are much lower here. (Figure 6; see Additional file 3: Figure S2, Additional file 4: Figure S3 and Additional file 5: Figure S4 for similar preparations of older embryos). These observations are consistent with the proposal that expression in the periproctodeal region of the Strigamia embryo is highly dynamic, with transcripts being cleared from cells once every cycle of the pattern. In the resolved stripes, Delta transcripts may be more stable here than they are in the periproctodeal region.

In the presomitic mesoderm of zebrafish, dynamic waves of gene expression across the tissue are revealed by an offset between the distribution of nascent nuclear transcripts and the accumulation of cytoplasmic transcript for genes, such as Delta [17]. This offset is due to the delay in the appearance of cytoplasmic transcript relative to the onset of transcription, and hence depends in part on the length of the transcription unit. We have not observed convincing evidence for an offset between nascent and cytoplasmic Delta transcript, in either the peri-proctodeal region, or in the resolved stripes. The transcription unit of the Delta gene is only 8.3 kb long though, which, in the context of the relatively slow rate of Strigamia segmentation, may not generate a sufficient delay between the onset of transcription and the appearance of cytoplasmic transcript.

**Concentric proctodeal patterning and germ band formation are separable processes**

We have observed a rare but telling instance of a spontaneous abnormal embryo in which the process of concentric posterior patterning has proceeded in the absence of any germ band formation (Figure 7A). At least 10 rings of eve1 expression are present, but there is no sign of germ band elongation/convergent extension and only very limited condensation, possibly of the precursor of the head. The embryo is apparently at a late blastoderm stage, but eve1 expression is much more finely patterned than would be expected at this stage, and the embryo may in fact be much older. This appears to be a case where dorsoventral and/or axial patterning have failed, but concentric patterning has continued.

This exceptional embryo is one of a clutch where all other embryos showed defects in germ band elongation.

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**Figure 5 Oscillation within posterior area of Delta and eve.** Posterior area, proctodeum included (asterisk), of flat mounted germ bands double stained for eve1 and Delta, anterior to the top. Embryos A and B, very close in developmental age (around 24 LBS), show different phases of oscillation of eve1 and Delta internal to the 38th LBS ring of eve1 expression (arrow). Delta is broadly expressed close to the proctodeum in A, largely overlapping with eve, but is down-regulated in this region in B, and resolving to a 39th stripe. Once fully resolved, primary stripes of eve (magenta arrow) and Delta (blue arrow) are out of phase, while the following intercalated stripes are directly superposed (A1, high magnification of A). C: An abnormal asymmetric embryo showing different phases of oscillation on the two sides of the embryo. Asterisk marks the proctodeum. Scale bar: 100 μm in A, B, C, 50 μm in A1.
Figure 7C depicts an embryo from another clutch, stained for Delta, which showed a similar defect. The germ band has formed, but is shorter and extends over less of the egg surface than in normal embryos with similar numbers of segments and primary rings of eve expression (Figure 7B). In these embryos segment patterning in the posterior disc seems to be compressed, and there are fewer cells between Delta stripes than in a normal embryo: four to five cells between primary stripes just before intercalation in the short embryo of Figure 7C, compared with seven to nine cells between stripes at the same position in a normal embryo (Figure 7B).

Loss of oscillatory expression correlates with a late transition from double to single segment addition

The dynamic patterns described above are observed throughout the period of trunk segment patterning until the primary rings corresponding to about the 38th leg-bearing segment have been generated (approximately the 26 LBS stage, beginning of stage 4.3).

At the early stages of trunk segment patterning, the double segment pattern persists for about five repeats before the intercalation of segmental stripes. This implies that there is a considerable delay (about 30 hours at 13C) between the generation of the double segment pre-pattern, and the resolution of the single segment repeat. However, as segmentation progresses, the appearance of intercalating stripes “catches up” with primary ring formation. This is demonstrated in Figure 8, which plots, for a series of embryos ordered by age, the appearance of morphological segments (in blue), the activation of eve1 in segmental intercalated stripes that presage the appearance of the odd numbered LBS (in red) and the first appearance of the double segment “primary stripes” that will resolve to define the even numbered LBS (in yellow). For example, for the embryo highlighted by the inset photo in Figure 8 (embryo B-15), only one leg-bearing segment is morphologically visible; eve segmental stripes are resolved up to the 9th LBS but primary eve expression corresponding to the 22nd leg-bearing segment is already appearing. However, the yellow and red lines approach one another progressively as segmentation proceeds, until in embryos with more than 30 leg-bearing segments, the two phases of eve expression are no longer distinct. eve and Delta stripes corresponding to later segments appear individually, with no double segment pre-patterning.

In most specimens of our population, the last resolved primary ring of eve that generates two segments is that
corresponding to the 38th or 40th segment. For Delta, the last primary pair-rule ring is that corresponding to LBS 39 or 41. Some variability here probably reflects variation in the final segment number of the adult animals, which in our population ranges from 43 to 53, with the great majority of the male individuals having 47 segments, and the females 49 segments.

At this time, the pattern of gene expression in the periproctodeal area changes suggesting that this segmenting tissue has entered a new regulatory regime. From about the stage of embryo G-03 in the plot of Figure 8 (that is, 32 LBS visible), eve1 is uniformly expressed in the periproctodeal area, with no apparent oscillation (Figure 9G, K, P U); the eve2 gene, which only ever shows double segment periodicity, is progressively repressed (Figure 9F, O, T), with the expression fading most quickly from the germ band tissues (Figure 9 F2). At the anterior margin of the periproctodeal area, segmental eve1 stripes resolve from the uniform domain of expression. These are limited to the germ band - they do not encircle the proctodeum. Delta is co-expressed with eve in these stripes (Figure 9H), in sharp contrast to the alternating expression that characterizes the earlier pair-rule activation of Delta and eve (for example, Figure 9C). By the time segment addition has paused at late stage 5, around the 43 LBS stage, Delta expression has become restricted to a single apparently stable band just posterior to the last formed segment (Figure 9Q, R) and only one or two eve stripes persist after resolution (Figure 9P, Q). By this time, the activation of engrailed expression occurs close to the Delta stripe, appearing just one segment’s width anterior to it (Figure 9S, Additional file 6: Figure S5). Eventually, segmental Delta expression fades completely just before the onset of germ band spreading in stage 6, although stable Delta expression persists in the proctodeal ring itself (Figure 9V, W). The lack of resolving eve stripes at stage 6 (Figure 9U) correlates with the pause in segment addition at this stage.

These changes also affect some genes that are not cyclically expressed - for example, knirps. knirps is expressed throughout the posterior of the egg at very early stages (C. Brena, unpublished), but during stage 4 it is repressed in all territory posterior to the second leg-bearing segment, except for the proctodeal ring itself (Figure 9E, J, N, Y). It is re-expressed specifically in the unsegmented territory anterior to the proctodeum around the 34 LBS stage, at about the time that oscillation stops (Figure 9J), and expression is maintained throughout stages 5 and 6 (Figure 9N, Y).

The rate of appearance of morphologically defined segments, which has been constant throughout stage 4 [14], continues unchanged for a period after the cessation of oscillation, while the remaining primary bands resolve from double to single segment periodicity (Figure 8, Additional file 6: Figure S5). However, the segments that
are actually patterned from this time on appear much more slowly than those generated from the double segment prepattern: segmentation slows to a rate of about 1.6 segments per day, compared with 7.5 segments per day during stage 4 [14].

The early onset of periodic patterning
Expression of the segmentation genes that we are considering here initiates during the blastoderm stage, while the head segments are being patterned. Our understanding of these earliest stages of segment patterning is less clear, in part because these early embryos are extremely difficult to preserve during the process of in situ hybridization. However, the data that we have show significant parallels with the later phase of trunk segment addition, though with some unique features.

A key question to address for these early stages is whether anterior segments are patterned by a process similar to that which occurs in the trunk, which seems to involve dynamic patterning of the pair-rule genes as the primary process, or whether they are patterned by a process more akin to that in Drosophila, where subdivision of the embryo occurs with respect to localized transcription factors of the gap gene type, which generate a non-periodic spatial pre-pattern that instructs the pair-rule pattern.

Below, we present the data that are available for eve and Delta gene expression during these early stages, before addressing this question explicitly in the Discussion.

even-skipped
At the uniform blastoderm stage (Stage 2.1, Figure 10), the even-skipped genes are expressed in a broad posterior domain that occupies the whole of the posterior hemisphere of the egg. The few embryos that we have at early blastoderm stages suggest that this expression initiates at the posterior pole of the egg and spreads anteriorly as a growing cap (Figure 10B; eve2 staining).
As the head primordium begins to condense ventrally, and to move anteriorly, the uniform expression of eve becomes modulated by the first signs of segment patterning. For eve1, a single broad anterior stripe resolves at the anterior of the expression domain, near the equator of the egg (Figure 10A, D). Within the territory of the head condensation, this quickly resolves into two segmental stripes that define the mandibular and first maxillary segments (Figure 10G; Note that this figure presents only a subset of the embryos used to assign segment identities to bands; for eve1, the complete series of available embryos is shown in Additional file 7: Figure S6.). Outside of the germ band, eve1 expression fades completely. A second broad stripe resolves behind the first, initially at 25 to 30% egg length from the posterior pole (Figure 10D), but moving forwards until it too lies near the equator, close behind the now resolved mandibular and first maxillary stripes (Figure 10G). Further stripes resolve more posteriorly, defining the double segment periodicity that will characterize...
eve expression throughout later development (Figure 10G, K, O, S).

eve2 shows a similar pattern of primary stripe formation, but expression fades completely without ever resolving to single segment periodicity. The transcript disappears slightly faster, if anything, from the germ band than from the surrounding territory. The most anterior stripe is barely resolved before fading (Figure 10E); the second stripe persists slightly longer, becoming well resolved as it moves forward towards the middle of the egg (Figure 10I), before fading as it reaches about 60% egg length (Figure 10M).

As the second stripe fades, posterior expression of eve2 becomes up-regulated immediately around the blastopore, which is now sharply defined as an area free of eve expression (Figure 10E). A series of closely staged eggs suggest that this expression spreads until it uniformly fills a posterior cap that extends to about 30% egg length. A third eve stripe then resolves from the margin of this cap. The process repeats, as trunk segmentation becomes

Figure 10 Expression of even-skipped and Delta genes at the onset of periodic patterning. Embryos are aligned as in Figure 1, from blastoderm stage 2.1, to the appearance of the first morphological segment, the mandible, at early stage 3.1. Panels with the same letter present different views of the same embryo (either lateral, or a view from the posterior pole orthogonal to these lateral views, with the blastopore/protodermum at the center (black or white arrow). Ventral is to the bottom in all images. Fluorescent nuclear staining (SYBR green) for the embryos hybridized with eve1 (lateral view) and eve2 (posterior view) represents the morphology for the whole row. The temporal alignment of embryos within rows is based on double in situ for eve1 and eve2 (H, L, P, T) and for eve1 and Delta (data not shown). Note that embryo B is younger than embryos A and C. The entire pattern is dynamic at this stage, condensing anteriorly as the pattern resolves. Based on a larger series of embryos (see Additional file 7: Figure S6), we have inferred continuity of expression between stages, as for example in the comparison of K2 and O2 in this plate. Expression domains are shown in this figure with arrows corresponding to the identity of the morphological segments eventually associated with each stripe, color-coded as indicated. A lighter color has been used for stages 2.1 and 2.2, where this identification is uncertain. (ant/int - antennal/intercalary segment (apparently deriving from same initial stripe); mn, mandible; mx1, first maxilla; mx2, second maxilla; mp, maxilliped; 4LBS, fourth leg-bearing segment). Note that in N1, R1, V1, the green arrow marks Delta expression associated with a single stripe that splits to give rise to both the maxilliped and the first LBS. Scale bar: 400 μm.
established, with each new stripe fading shortly after it is formed (Figure 10M, Q, U). (Later, a single transient stripe of eve2 expression arises in the antennal segment, but the relation of this to early eve2 expression is unclear; Figure 10P-Q, T-U).

**Delta**

During the uniform blastoderm stage, while the eve genes are expressed as a posterior cap, Delta is expressed as a single broad stripe around the ventral two-thirds of the egg’s circumference, at 50% egg length. There is also a faint patch of Delta expression at the posterior pole (Figure 10C). The ventral stripe appears to move forward until it lies at about 60% egg length, when a second stripe appears behind it (Figure 10F). It is not clear whether this second stripe is ever preceded by broad expression of Delta in the posterior hemisphere.

At around the onset of head condensation, these two anterior Delta stripes resolve into a pair of major stripes coincident with the engrailed stripes of the intercalary and first maxillary segments, and a minor stripe of the intervening mandibular segment (Figure 10J). At the same time, posterior expression of Delta appears, first as a uniform posterior cap, and then resolving to a series of primary bands. Like evenskipped, Delta expression is quickly excluded from the region of the blastopore itself, but each new primary band arises with strong expression around the blastopore (Figure 10J, N, R, V). The primary bands of Delta arising from the posterior cap mark the first, third, fifth and successive leg-bearing segments. Exceptionally, the first of these primary Delta bands appears to split to give rise to both the maxilipedal and the first leg-bearing segment (Figure 10R, V). Thereafter, secondary stripes intercalate to mark the second, fourth and successive even-numbered LBS.

**Discussion**

There are currently two models for the generation of segments in arthropods. In the Drosophila paradigm, initially graded signals lead to the localized activation of the segment “gap genes”, encoding transcription factors that provide spatial cues unique to each position along the anteroposterior axis. These cues instruct periodic activation of segmentation genes using distinct transcriptional circuitry for different segments. We will refer to this as a “gap gene” based model.

The other model involves a cyclic, dynamic process of gene expression analogous to that which has been analyzed in some detail for vertebrate somitogenesis [5,18]. Such a model has been proposed as the mechanism for segment formation in short germ arthropods, and is supported by experimental data in a number of species [19-22], although these data are open to alternative interpretations [23,24]. However, two recent papers provide clear evidence for cyclic waves of gene expression preceding segmentation in the short germ beetle Tribolium [3,4]. We will refer to this second class of model as a clock-based process.

**A clock-like mechanism in Strigamia**

The evidence supporting a clock-like oscillation of eve and Delta expression in the peri-proctodeal area of Strigamia can be summarized as follows:

- the coherent transformation of the observed expression patterns in embryos placed into a closely staged developmental series by other criteria (Figure 4).
- the rapid dynamics of change in the peri-proctodeal area, contrasting with the relative stability of the surrounding resolved pattern (Figures 4E and 5A, B).
- the observation of asymmetric patterns in single embryos that closely resemble different stages of the inferred cycle (Figures 4F and 5C).
- the extent of the egg surface that shows dynamic expression in the earliest stages of patterning (Figure 10), which appears to rule out evolution of these patterns by wholesale cell movement.

This evidence falls short of that which could be provided by embryo culture experiments coupled with cell marking, but such techniques have not yet proved possible in Strigamia. However, the available data make a strong argument that the patterns of expression observed for even-skipped and Delta reflect repeated cycles of transcription within this whole cell population. It is likely that the similar patterns shown by homologues of the pair rule genes hairy, runt and oddskipped reflect similar dynamic patterns [11-13] (C. Brena, data not shown).

This oscillatory activity is apparent during the main phase of trunk segmentation. For eve2 and Delta (at least), each cycle initiates at paired lateral sites adjacent to the proctodeum, once in each double segment cycle. This activation spreads, somewhat unevenly, throughout the majority of the peri-proctodeal area. Transcript is then cleared from the inner part of this area, leaving an ill-defined ring that sharpens and becomes more regular as it expands somewhat further.

The dynamic nature of this pattern is particularly clear during early stages of the segmentation process, when the primary rings that will give rise to the mouthparts and the most anterior leg-bearing segments are generated. During these earliest cycles, transcription of eve is activated throughout almost the entire posterior hemisphere of the egg excepting the area of the blastopore itself, and then cleared from all of this territory except an anterior marginal ring. Given that the cell density remains broadly uniform across the posterior hemisphere of the egg (which
at this stage is the multilayered blastoderm [14], it is not feasible that the loss of expression from the posterior territory in each cycle is due to movement of all of these cells into the marginal ring.

The pacemaker for these oscillations seems to lie at or adjacent to the blastopore throughout the segmentation process. Initially, new cycles of gene expression appear concentrically around the extended blastopore, but as the morphology of the proctodeum develops, the initiation becomes bilateral, with new cycles of expression initiating at foci that appear as two patches lying antero-laterally just outside the proctodeal ring; tissues of the proctodeum itself do not show oscillating gene expression. This posterior territory is already multilayered [14], and clearly divided into a number of populations expressing distinct cell markers (data not shown). Oscillatory expression of the pair rule genes is restricted to the outer, ectodermal cell layer [13] at least by the 15 to 20 LBS stage. The observation that this oscillation can be maintained in an embryo that lacks any obvious sign of germ band formation (Figure 7A) suggests that axial patterning is distinct from the generation of periodicity that underlies segmentation.

It appears that gene expression becomes stable in individual cells at or shortly after the time that each ring becomes well resolved. The further slow expansion of this primary pattern reflects tissue movement driven by continuing proliferation of the posterior disc, and convergent extension of the forming segments as the whole egg epithelium condenses anteriorly and cells move into the germ band. However, we cannot at present say precisely where the transition from oscillatory to stable gene expression occurs.

The prevalence of clock-like segmentation processes in arthropods

The oscillation of eve and Delta expression that we infer in Strigamia is similar to that recently reported for odd-skipped and eve in Tribolium, by Sarrazin et al. [3] and El-Sherif et al. [4]. These authors have shown by culturing bisected embryos, by comparing series of closely staged embryos and by tracking cell movements in the posterior of the germ band that the early stripes of odd-skipped and eve expression are indeed waves of oscillatory gene expression moving across cells. These waves are of double segment periodicity, and move in register but out of phase.

In no other arthropod has oscillatory gene expression been documented unambiguously, but the phenomenology of gene expression suggests that it exists quite widely. It was first suggested for opisthosomal segment patterning in spiders [19,20], and has been proposed for abdominal segment addition in at least one hemimetabolous short germ insect [21,25]. The concentric expression of segmentation genes around the proctodeum in the millipede Glomeris appears strikingly similar to what we observe in Strigamia [26]. Concentric patterns have been observed, at least for even-skipped, in another centipede, Lithobius [7]. However, in all of these cases the patterns of gene expression define a single segment periodicity, rather than the double segment periodicity observed in Strigamia and in Tribolium. This raises some question as to the homology of the oscillatory mechanisms that may exist across the arthropods. However, it may be that the processes which generate primary single segmental stripes in Glomeris, and perhaps other arthropods, may be homologous to the mechanisms that generate primary pair-rule patterning in Strigamia.

Dynamic oscillation of gene expression also occurs in the pre-somitic mesoderm of vertebrates, where the oscillatory mechanism has been analyzed in some detail [5]. The precise mechanism of the oscillator appears to vary between species [27], but in all vertebrates examined, the oscillation is of single somite periodicity, and is limited to the mesoderm, whereas in arthropods primary segment patterning involves the ectoderm. In zebrafish (at least) the oscillation seems to be a cell autonomous process, and the role of Notch signaling is to co-ordinate the phase of oscillation between cells [28]. The extent to which similar mechanisms underlie oscillatory gene expression in vertebrates and arthropods remains to be documented: modeling suggests that it may be rather easy to generate either (or both) spatial and temporal oscillation from gene networks that generate pattern [29]. Intriguingly, a cell autonomous circuit of pair-rule gene interaction has been described in Tribolium [30] but this might serve either to generate dynamic behavior, or to ensure that cell states are mutually exclusive once pair-rule periodic expression is established.

Anterior segment patterning - clock-like or gap-gene based?

In even the most extreme of short germ insects, the segments of the mouthparts and anterior thorax appear to be patterned from cells generated during cleavage. Most authors have assumed that these segments in all insects are patterned by a gap-like mechanism dependent on localized factors upstream of the pair-rule genes [31] and we thought it likely that the same would be true in Strigamia. We therefore expected to see that the dynamic patterning of Delta in association with eve would initiate posterior to the first formed segments, approximately at the position of the first leg-bearing segment.

Our data confirm that segments of the trunk, from the second leg-bearing segment backwards, do appear to be generated by a simple repeat of the same cycle of gene expression states. However, to our surprise, both Delta and eve show patterns of dynamic expression that initiate
at the very onset of segment patterning, and relate to the generation of pattern from the intercalary segment back.

For *eve*, this early expression takes the form of an initial activation throughout the entire posterior hemisphere of the egg, and then the resolution of stripes within this domain. This is reminiscent of early *eve* expression in insects, the only other arthropods that have been studied at these early stages. In *Drosophila*, the resolution of stripes within this early domain depends on the regulation of *eve* by gap genes.

It is possible that for *Strigamia* too, the early resolution of the *eve* pattern depends on upstream “gap like” factors. Gap gene domains themselves may shift dynamically due to gene interactions, and the expression of dependent pair-rule genes then follows them, as has been documented in the dipteran insect *Clognia albipunctata* [32]. However, the observation that the dynamics of Delta parallel those of *even-skipped*, even for these very first stripes, and that the whole early pattern appears to move so dynamically across the blastoderm as it resolves, raises the question as to whether the patterning of even the most anterior segments, the intercalary and mouthpart segments, might depend on signaling interactions similar to those inferred for the more posterior trunk segments.

**The origins and significance of pair-rule patterning**

Pair-rule patterning during segmentation was first described in *Drosophila*, where the mechanism is relatively well understood. In most other insects studied, at least some homologues of the *Drosophila* pair-rule genes are expressed with a double segment periodicity, although the function of some genes appears to differ between species. No pair-rule expression has been reported for segmentation genes in some short germ insects [33,34], in the few crustaceans studied [35,36], or for most segments in spiders [37], though in the latter the most anterior stripes of gene expression are involved in a splitting phase [38,39]. There is one report of pair-rule patterning for a *paired* (Pax3/7) homologue in the prosoma of a mite [40].

The pair-rule nature of primary segment patterning in *Strigamia* is very clear. However, no such pair-rule patterning is apparent in the lithobiomorph centipede *Lithobius atkinsoni*, where *even-skipped* expression has been studied in relation to segment polarity marker genes [7]. There is also no trace of pair-rule patterning during trunk segment patterning in the millipede, *Glomeris marginata* [26], although stripe splitting is observed in the head [41]. Thus, among the myriapods, pair-rule patterning of trunk segments has been observed uniquely in *Strigamia*. It may, therefore, be a feature that has evolved within centipedes, and may prove to be unique to geophilomorphs, or to the single derived lineage leading to geophilomorphs and scolopendromorphs [42,43], which is the clade of centipedes that shows the highest segment numbers. *Lithobius*, showing single segment addition, lies basal to the origin of this clade.

**The final phase of segment addition - from pair-rule to single segment periodicity**

The transition from double to direct single segment patterning appears to be accompanied by a broad change in the regulatory regime occurring in the unsegmented posterior tissue: *eve1* expression becomes stable, and *eve2* and *Delta* expression are switched off in the posterior tissue at the time of this transition. At least one other transcription factor not previously active during the main phase of segmentation, *knirps*, is switched on. In contrast to the earlier phase of patterning, segmental stripes resolve only in the region of the germ band, and not around the entire periphery of the peri-proctodeal area. There is no indication that the whole tissue is undergoing dynamic cycles of gene expression.

This final phase of single segment addition undermines the argument we have made previously [11], that pair rule patterning might explain an observed constraint on the number of segments in centipedes. All centipedes have an odd number of pairs of leg-bearing segments as adults and, hence, an even number of trunk segments including the poison claw segment. This constraint is particularly striking for the geophilomorph centipedes, which show great inter- and intra-specific variability in the number of leg-bearing segments, which ranges from 27 to 191 [43].

This observation is readily explained as a developmental constraint if all trunk segments are generated in pairs. However, if the last segments are generated singly, we can only preserve this explanation by proposing that there is no variation in the number of singly patterned segments, and that the variation available within populations is restricted to the phase of double segment patterning. In this context, it is relevant that *Lithobius*, which appears to make its trunk segments singly [7], belongs to a clade that indeed shows no variation in segment number: all lithobiomorph centipedes have 15 pairs of leg-bearing segments. Invariant segment number is in fact frequent in all major arthropod lineages, particularly among relatively short bodied forms (for example, of no more than 20 segments). Thus it is not unreasonable to propose that, in the long bodied geophilomorphs, the variable generation of segments by pair rule patterning has been imposed on an invariant underlying body plan.

**Conclusions**

The dynamic appearance of gene expression during segment patterning in *Strigamia* reflects two processes - an
intrinsically dynamic process of gene expression in cells around the proctodeum, and the overall movement of cells in the germ band more anteriorly. A pair-rule segmentation oscillator underlies the generation of all anterior trunk segments, and may also be involved in patterning of head segments from the intercalary back. However, there appears to be a transition in the dynamics of segment addition at stage 4.3, such that oscillatory gene expression is suppressed, and the last 10 trunk segments are patterned individually, without a pair-rule pre-pattern. These alternative modes of behavior for the segmentation gene network in one species suggest that the different modes of segmentation observed in different arthropods may evolve rather readily from a largely conserved underlying gene network.

Methods

*Strigamia maritima* eggs were collected, cultured, fixed, stained and photographed as described in [14].

For the preparation of *in situ* hybridization probes, previously cloned fragments of the *S. maritima* genes eve1, eve2 [12] *knirps* (NCBI Accession: EF175909.1 to EF175912.1) and *dpp* (A. Chipman, unpublished data) were extended with gene-specific primers through 3’ and 5’ rapid amplification of cDNA ends (RACE) (BD SMART™ RACE cDNA Amplification Kit, Clontech, Mountain View, CA, USA). For *en* and *Delta* published clones were used to make the probes [12,15]. Probes for exon sequences were prepared using the longest available cDNA clone of each gene (2.2 kb for *dpp*, 1.3 kb for *knirps*, 0.7 kb for *en* for Delta, eve1 and eve2, see Additional file 2: Figure S1). The intron probes for Delta and eve1 were designed as reported in Additional file 2: Figure S1, after mapping cDNA data against genomic scaffolds from the Smar_1.0 genome assembly [44]. The Delta ‘intron’ probe contains >95% intron sequence, but does span two short (60 bp) exons in the 2.8 kb sequence, and so detects cytoplasmic transcript weakly, as well as strong nuclear signal from nascent transcript. The eve1 intron probe is derived purely from intron; this probe yielded no signal following *in situ* hybridization (see Additional file 2: Figure S1).

*In situ* hybridization was carried out essentially as described by [15,45], but in general with the anti-DIG antibody (Roche, Mannheim, Germany) at 1:3,000 dilution. In the case of intron probes, hybridization and staining reaction times have been increased to up to 20 h and 8 h, respectively, (in the case of the unsuccessful eve1 intron probes to up to 40 h and 13 h, respectively).

Stages of embryos were estimated as described in [14]. During stages 3 to 5, embryos were further staged by recording the number of the last leg-bearing segment morphologically visible, that is, defined by both anterior and posterior segmental grooves.

**Additional files**

**Additional file 1:** Movie 1. A single oscillation of eve1 expression. Possible representation of a single oscillation of eve1 expression, obtained by overlapping ordered embryos A-D of Figure 4 (younger embryo on top), with increasing transparency of the younger embryo in each sequential frame. The ring corresponding to the 24th LBS (marked by red arrows), external to the dynamic peri-proctodeal area, shows no detectable movement over this time window.

**Additional file 2:** Figure S1. Gene structure for the genes Delta, eve1 and eve2. Gene structure and extension of *in situ* probes for the genes Delta, eve1 and eve2, mapped against genomic scaffolds from the Smar_1.0 genome assembly (http://www.ncbi.nlm.nih.gov/assembly/322118/), and gene models derived from the current genome annotation (pending submission to Ensemble Metazoom - web link to be provided at proof stage), which incorporates RNAseq data as well as the cDNA information. The genome map is represented on a light blue background. Bars represent exons, the blue-filled parts being the coding regions, and the open parts the UTRs; 5’ on the left, red lines mark the stop codons. Hybridization in *situ* probes are represented on a black background, with exon probes in yellow (derived from cDNA clones) and intron probes in light blue, derived from genomic DNA. Note that in the case of Delta the two intron probes each include a small exon (60 nucleotides). The two exclusively intronic probes of eve1 have not yielded any detectable signal on *in situ* hybridization. (Data from Strigamia maritima genome assembly Smar_1.0. Note that the three gene maps are not to the same scale.

**Additional file 3:** Figure S2. Distribution of Delta nascent transcripts in a late stage 3 embryo. A1 Flat mount of an embryo at the stage with one leg-bearing segment morphologically visible (1LBS stage), hybridized with a Delta probe comprising approximately 95% intron and 5% exon sequence (see Additional file 2: Figure S1). This probe detects unprocessed nascent transcripts much more strongly than spliced cytoplasmic transcripts. At this early stage there are prominent nuclear dots of nascent transcript (magenta arrowheads) in many cells of the peri-proctodeal region (enlarged in panel A3); double nuclear dots are often visible, showing transcription associated with both copies of the gene. In more mature Delta stripes, transcript is still present at high level in the cytoplasm, but nuclear dots are less frequent and less prominent (enlarged in A2). Note that embryos at this stage extend around much of the egg, and are therefore distorted and split at the lateral margins when flattened. The head is to the left. Black arrowhead: mandibular segment; an asterisk marks the proctodeum. Scale bar in A1: 200 μm, in A2-3: 20 μm.

**Additional file 4:** Figure S3. Distribution of Delta nascent transcripts in an early stage 4 embryo. A1 Flat mount of an embryo at the 15 LBS stage, hybridized with a Delta probe that detects unprocessed nascent transcripts much more strongly than spliced cytoplasmic transcripts (see Additional file 2: Figure S1). At this stage Delta transcripts are no longer detectable in *in situ* hybridization (see Additional file 2: Figure S1). At this stage nuclear dots of nascent transcript (magenta arrowheads) are still prominent in many cells of the peri-proctodeal region (enlarged in A4, A5), although the signal intensity may be somewhat lower than in earlier stages. In more mature Delta stripes, transcript is still present at high level in the cytoplasm, but nuclear dots are less frequent (insert A3). Nuclear dots are almost undetectable in the oldest, more anterior stripes (insert A2). The head is to the left. Black arrowhead: mandibular segment; an asterisk marks the proctodeum. Scale bar in A1: 200 μm, in A2-5: 20 μm.

**Additional file 5:** Figure S4. Distribution of Delta nascent transcripts in an early stage 5 embryo. A1 Flat mount of an embryo at the 41 LBS stage, hybridized with a Delta probe that detects unprocessed nascent transcripts much more strongly than spliced cytoplasmic transcripts (see Additional file 2: Figure S1). At this stage Delta transcripts are no longer detectable in the peri-proctodeal region. Two posterior stripes of Delta expression remain, defined by an accumulation of cytoplasmic transcript, but nuclear dots of nascent transcript are no longer detectable in these stripes (insert A4). Nuclear dots show that transcription of Delta is activated at high level in more anterior medial cells (magenta arrowheads in insert A3). These cells are neuronal precursors. In more mature segments they will form clusters where Delta is present at high level both in the nuclei and the cytoplasm (insert A2). The head is to the left. Black arrowhead: mandibular segment; an asterisk marks the proctodeum. Scale bar in A1: 200 μm, in A2-4: 20 μm.
Additional file 6: Figure S5. Last phase of segmentation from double (primary Delta) to single-segment periodicity (enlarged secondary Delta). In a 34 LBS embryo (A) Delta still appears at double segment periodicity (large blue arrowheads); intercalated Delta stripes defining the single segment periodicity appear only slightly later, that is, more anteriorly, as the segment pattern matures (small blue arrowheads). Engrailed expression (magenta arrowheads) appears yet slightly later than intercalated Delta, overlapping with the segmental stripes of Delta as these fade. Engrailed is then expressed persistently in the whole maturing germ band anterior of the segment addition zone. In a 36 LBS embryo (B), the single segment intercalation of Delta appears just after the earliest resolved Delta band; engrailed expression appears closer to this first Delta stripe. By the 43 LBS stage (C), only a single stripe of Delta transcript persists. The transcription of engrailed initiates at a position just one segment anterior to this Delta stripe, but Delta expression has already faded from this anterior region. Scale bar in A1, B1, C1: 100 μm, in A2-C2: 50 μm.

Additional file 7: Figure S6. The complete set of 28 embryos used to deduce the maturing pattern of eve expression as presented in main text Figure 10. These embryos have been put into a developmental series using both morphological staging markers and the transitions in the eve pattern itself. Embryos A, D, G, K, O and S are the embryos indicated with the same letters in Figure 10, selected from this full series. The correspondence of bands at the different stages was inferred with reference to the full series, allowing for example the conclusion that the Mx2 band (blue arrow in Figure 10, red arrow in Additional file 7: Figure S6) moves anteriorly during stages 2 and 3. Similar, though less extensive, series were used to infer the correspondence of Delta and engrailed expression patterns between stages. Each embryo is presented in a single row: Columns 1 and 2 - lateral views; columns 2 and 3 - ventral views; columns 4 and 5 posterior views; paired nuclear fluorescence and bright field images.

Abbreviations
Dgk, Digoxygenin; dppl, decapentaplegic; en, engrailed; eve, even-skipped; LBS, leg bearing segment.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
CB carried out all of the experimental work, recognized the transition from double to single segment patterning, and developed the first interpretation of the data. MA participated in the interpretation of the data, and both authors wrote the manuscript. Both authors read and approved the final manuscript.

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