Split hand/foot malformation genetics supports the chromosome 7 copy segregation mechanism for human limb development

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Genetic aberrations of several unlinked loci cause human congenital split hand/foot malformation (SHFM) development. Mutations of the DLX5 (distal-less) transcription factor-encoding gene in chromosome 7 cause SHFM through haploinsufficiency, but the vast majority of cases result from heterozygous chromosomal aberrations of the region without mutating the DLX5 gene. To resolve this paradox, we invoke a chromosomal epigenetic mechanism for limb development. It is composed of a monochromatid gene expression phenomenon that we discovered in two fission yeasts with the selective chromosome copy segregation phenomenon that we discovered in mouse cells. Accordingly, one daughter cell inherits both expressed DLX5 copies while the other daughter inherits both epigenetically silenced ones from a single deterministic cell of the developing limb. Thus, differentiated daughter cells after further proliferation will correspondingly produce proximal/distal-limb tissues. Published results of a Chr. 7 translocation with a centromere-proximal breakpoint situated over 41 million bases away from the DLX locus, centromeric and DLX5-region inversions have satisfied key genetic and developmental biology predictions of the mechanism. Further genetic tests of the mechanism are proposed. We propose that the DNA double helical structure itself causes the development of sister cells’ gene regulation asymmetry. We also argue against the conventionally invoked morphogen model of development.

This article is part of the themed issue ‘Provocative questions in left–right asymmetry’.

1. Background

Biological research is in crisis—Technology gives us the tools to analyze organisms at all scales, but we are drowning in a sea of data and thirsting for some theoretical framework with which to understand it [1].

Vertebrate limb development has been an active area of research in the field of developmental biology for many decades (reviewed in [2]). The morphogen gradient model [3] has been the primary paradigm followed for guiding research on limbs, as well as for the development of other organs during embryogenesis. However, despite decades of extensive research on development in all sorts of organisms, it is not understood precisely how developmental control genes are regulated, expressed or silenced at the correct position and with their exact timing in the course of development, nor how the development of different cells types and tissues is coordinated during embryogenesis. In the case of limb development, naturally arising limb developmental anomalies have provided a rich source of genetic material for discovering developmental mechanisms operating in humans. The congenital split hand/foot disorder (SHFM; online Mendelian inheritance in man (OMIM) 225300), also referred to as ectrodactyly (figure 1), is such a congenital defect in limb digit formation. SHFM consists of a spectrum of distal portion malformations of the hand/foot owing to a deep median cleft, missing digits and bones and missing
digits and hypoplasia of the central rays [4]. The SHFM occurs as an isolated limb defect (non-syndromic and sporadic) or as part of a syndrome associated with defects of other organs, such as mental retardation, hearing loss and clift lip and palate [5]. Also, the extent of the SHFM malformation phenotype is highly variable, and inexplicably, zero to four limbs can be malformed in the same genetically predisposed individual in familial cases.

As the relationship between SHFM genotype and phenotype is not understood, here, we present a new developmental biology framework for explaining the baffling genetics of chromosome (Chr.) 7 SHFM1 locus aberrations affecting limb development without invoking mutations of the relevant DLX5 gene as the cause of malformation. The DLX5 gene encodes a transcription factor related to the Drosophila distal-less homeobox domain-containing protein, hence named DLX gene in humans. We exploit the genetics of Chr. 7 aberrations to dissect the molecular aetiology of the SHFM1 locus and of the biology of limb development. Non-syndromic cases occur in about one in 18 000 births, and most of them are sporadic in nature. The aetiology of sporadic cases also remains unknown [6]; we propose the same chromosomal basis for their origin too. Our analysis aims to dis- cover unknown aspects of the limb developmental mechanism. Here, we argue that the complementarity of DNA strands itself can be converted into a/symmetric gene expression of daughter/sister cells, a phenomenon likely required as a key feature of eukaryotic development.

2. DLX5 gene defines the SHFM1 locus

Many of the genes and pathways for limb development have been remarkably conserved from Drosophila to humans [7]. SHFM is a heterogeneous condition caused by abnormalities of several loci in humans. At least seven loci have been identified for the familial inherited cases of SHFM through linkage or cytogenetic analyses [4,5,8–11]. These loci represent several chromosomal regions: SHFM1 maps on Chr. 7q21 (MIM 183600); SHFM2 on Chr. Xq26 (OMIM 313350); SHFM3 on Chr. 10q24 (MIM 600095); SHFM4 on Chr. 3q27 (MIM 605289) and SHFM5 on Chr. 2q31. Homozygous cathedrin-3 gene mutations cause SHFM, macular dystrophy syndrome and ectodermal dysplasia [12]. A p63 transcription factor, encoded by the p63 gene comprising the SHFM3 locus, binds to enhancers of the DLX5/6 genes of the SHFM1 locus to regulate gene expression during limb development [13]. p63 functions genetically upstream of the DLX5 expression, and thus, the DLX–p63 pathway defines the role of corresponding SHFM1 and SHFM4 loci in limb development [14]. These genes have been implicated in the development of the limb, craniofacial structures, the inner ear and the brain [15]. Importantly, the DLX gene functions in the Wnt signalling pathway required for limbs’ skeletal development [10].

Relatively recently described intragenic DLX5 gene mutations very clearly cause SHFM when in the heterozygous condition [11,16], but surprisingly, most cases are associated with heterozygous chromosomal aberrations of this 7q21 locus where the DLX gene itself is not interrupted or mutated [5,17–19]. These aberrations include deletions, translocations and inversions of the locus. Thus, a well-appreciated paradox persists in the literature whereby a gene mutation causes genetic disorder but paradoxically disorder is caused in most patients by chromosomal aberrations without the relevant gene having been mutated. Nearly all studies have hypothesized a position-effect control to explain this paradox such that the genomic rearrangements disrupt the normal expression pattern of the SHFM1 locus DLX5 gene by separating it from the required long-range-acting, cis-regulatory elements, resulting in decreased, increased or ectopic gene expression. Long-range gene regulators that function in developmental processes are well known. Surprisingly, such regulatory elements are dispersed in regions spread over hundreds of kilobases (Kb) upstream or downstream of the gene itself [20]. It is also well known that developmental control genes undergo tissue-specific and temporally express in time during development. However, the mechanisms for controlling the expression of DLX gene by position effects, tissue-specific expression and the biological basis of disorder-causing SHFM1 locus chromosomal aberrations are not fully understood [20]. For the SHFM1 locus-associated cases, an autosomal-dominant, incomplete penetrance and haploinsufficiency model for DLX gene regulation has been proposed in numerous studies [5,10]. Although this model helps to describe the preponderance of SHFM1 cases very well, it is not clear how to experimentally scrutinize its validity. Here in this hypothesis paper we propose a chromosomal epigenetic mechanism, a mechanism first discovered in studies of fission yeasts [21,22] and mouse cells [23,24], for differential regulation of the DLX gene of a specific pair of sister cells produced during limb development. In short, we aim to explain the enigmatic genetics of chromosomal aberrations that cause SHFM1 although their DLX5 gene is not mutated.

3. The DNA SSIS produces sister cells of different cell types in two evolutionarily diverse fission yeasts

Pertinent to defining the developmental biology mechanisms at large, it is crucial to understand how developmentally
equivalent or non-equivalent daughter cells are produced at specific stages of embryogenesis and during tissue homeostasis. Production of non-equivalent daughter cells is generally thought to be accomplished by regulated distribution of differentiation-specifying cellular factors and/or by differential exposure of daughter cells to cell-extrinsic factors [25]. Because different mechanisms might have evolved in different organisms, research with different organisms on this topic has continued for many decades. As a case in point, the mating-type switching phenomenon of fission yeasts has provided a powerful model system for understanding the biological mechanism of asymmetric cell division. In the highly diverged Schizosaccharomyces pombe [21,22] and S. japonicus [26] yeasts (reviewed in [27]), a DNA strand-specific, non-canonical imprint is installed at the mating-type locus (mat1) during its replication in mitotic cells. The imprint causes sister chromatids to differ by epigenetic means for their mat1 gene activity. By this mechanism, only one of the two sister cells produces one daughter of switched mating/cell type, such that strictly one of four granddaughters of a cell switches in over 80% of cellular pedigrees. This unique mechanism of cellular differentiation is based fundamentally on the inherent DNA sequence differences in the mat1 DNA strands. A key genetically predicted result that initially established the DNA strands asymmetry model was that yeast cells genetically engineered to carry an inverted mat1 gene duplication produced equivalent sister cells because both sister cells became capable of producing switched progeny of their own [22]. That is, the usually non-equivalent sister cells became developmentally equivalent once the mother cell carries the inverted mat1 duplication construct. The two chromosomal DNA strands carry DNA sequences that are complementary to one another, but they have opposite chemical polarities [28]. Furthermore, each strand serves as a template for synthesizing the complementary strand during chromosome duplication through the semi-conservative replication mechanism [29]. Thereby, each chromosome replication process produces paired daughter chromosome copies, which, in the G2 phase of the cell cycle, are called sister chromatids. Moreover, one chromatid always contains the arbitrarily designated Watson (W) DNA replication template strand and the newly synthesized Crick (C) strand, and the sister chromatid contains the template C strand and the newly synthesized W strand. Yeast sister chromatids are additionally differentiated by the inherent leading- versus lagging-strand mode of replication at the mat1 locus [30,31]. To accomplish such precision, mat1 is replicated strictly in a single chromosomal direction conducive for the imprinting process [32]. Altogether, the developmental asymmetry of daughter cells in fission yeasts is simply owing to the production and inheritance of the epigenetically differentiated mat1 gene residing on sister chromatids. Mechanistically, installation of the epigenetic mark is based on the specific W versus C strand, older versus newly synthesized, and the leading- versus lagging-strand mode of DNA replication at the mat1 locus. In short, the daughter cell's developmental asymmetry results from the parental cell's mat1 replication history [21,22], and that is strictly based on the double-helix structure of DNA. The understanding of the DNA strand's chirality mechanism of cellular differentiation in fission yeasts has motivated us to propose the somatic strand-specific imprinting and selective sister chromatid segregation (SSIS) model as a mechanism for explaining the origin of vertebrates' visceral organs' left–right laterality [33,34] and of the human brain’s hemispheric laterality development [35,36]. Here we advocate the same mechanism to explain both the precise regulation of DLX gene expression at a critical cell division during limb development and the aforementioned SHFM1 locus chromosomal aberrations affecting limb development in humans.

4. The SSIS mechanism proposed for the DLX5 gene regulation of a deterministic cell dividing in the limb bud anlagen

As stated above, correct spatial and temporal control of gene expression is essential for embryonic development, including limb development. Regulation of developmental control genes can be influenced by regulatory elements located some distance from the promoter regions, both in upstream and downstream regions of the gene [37], but their precise mechanism of regulation is not yet understood. Their transcriptional regulation is thought to be under chromosomal 'position-effect' controls. Many of these genes encode transcription factors with tissue- and stage-specific expression patterns. It is not known precisely how embryonic and stem cells establish a unique programme of gene expression that determines what kinds of daughter cells they will produce. Notably, the DLX gene encodes an evolutionarily conserved group of homeodomain transcription factors related to the Drosophila distal-less (dll) genes [14]. The DLX gene family consists of six gene clusters existing in vertebrates, which are generally organized into bi-gene clusters. In four-limbed (i.e. tetrapods) animals, the DLX5 gene is expressed in the apical ectodermal ridge of the developing limb bud and it is clearly required for limb development because altered function of DLX5 factor causes the human split hand/foot malformation 1 (SHFM1) development. The molecular signalling pathways involved in limb digit patterning and limb bud growth are best described by studies of the mouse and chicken model organisms [14]. In comparison, the mechanisms of how the DLX5 gene for limb development is precisely expressed or silenced in different human limb tissues remain undefined.

The adjoining DLX5 and DLX6 genes are regulated in a strict spatial pattern during embryogenesis. As stated above, these genes are supposedly regulated by the action of both long- and close-range cis-acting regulatory elements of the SHFM1 locus. Interestingly, regulatory elements, such as enhancers, that control DLX gene expression in vertebrates are spread in a region of several hundred Kb located both in the 5′- and 3′-flanking regions of the locus. Notably, disrupting the action of these regulatory elements causes limb defects phenotypically similar to those arising from conventional DLX5 gene coding region mutations [11,16,17,19]. Taken together, ectodactyly results from the 7q21.1–q22.1 region aberrations, where the dysregulation and/or haploinsufficiency model of DLX5 is well supported by numerous studies [4,14,18]. Haploinsufficiency is the term used when two copies of the gene are essential for the normal phenotype development. The SSIS mechanism (figure 2) is advanced here to help express DLX5 from both Chr. 7 homologues in one daughter cell and also to keep both alleles epigenetically silenced in the other daughter cell. This is hereafter named the monochromatid gene expression mechanism proposed to operate in the deterministic progenitor cell [39]. The SSIS
SSIS mechanism for chromosome 7 copies segregation of distal/proximal-fate-specifying determinant cell

Figure 2. Monochromatid gene expression and selective chromatid segregation components of the hypothetical SSIS mechanism proposed for limb development. The mechanism is advanced for the developmental regulation of the DLX5 (i.e. SHFM1) locus of Chr. 7 in the distal and proximal tissue fates—producing deterministic cell and its daughter cells produced during limb development. The mechanism is based on DNA bases and chemical polarity differences of the arbitrarily designated ‘Watson’ (W = 5’ top) and ‘Crick’ (C = 3’ top) strands of DNA and their inherently asymmetric leading- versus lagging-strand mode of replication for the distal/proximal tissue-specifying developmental DLX5 gene. The blue lines represent the DNA replication template W strands, the red lines represent the template C strands, and the grey lines represent the strand synthesized in the present replication cycle. For the sake of simplicity, DNA strands are deliberately drawn as straight lines and not as the normally existing double-helix configuration. By this mechanism, designed to achieve monochromatid gene expression, transcriptionally active (ON) and silenced (OFF) entities of the DLX5 gene are generated for both Chr. 7 copies in the limb determining cell in a DNA strand-specific manner, as drawn here. Accordingly, epigenetically unequal daughter chromosome copies (called chromatids in the G2 phase of the cell) are produced for the DLX gene of both Chr. 7 homologue replicates in an ‘ON/OFF’ manner owing to a non-canonical strand-specific imprinting process operating in mitotic cells. By our hypothesis, an unknown segregation factor operates at Chr. 7 centromeres to cause selective chromatid segregation by the designated W,W::C,C biased segregation mode drawn in the diagram. The deterministic cell undergoes stem-cell-like asymmetric cell division precisely with respect to adjoining cells that specify tissues of the anterior/posterior (formally equivalent to the left–right body axis) and dorsal/ventral axes. To coordinate the development of these three axes, perhaps precise cell-to-cell molecular signalling interactions occur between adjoining cells by maintaining planar coordinate the development of these three axes, perhaps precise cell-to-cell molecular signalling interactions occur between adjoining cells by maintaining planar

mechanism also employs the selective chromatid segregation process such that both template W strand-containing chromatids are segregated to a specific daughter cell whereas both C strand-containing chromatids are segregated to the other daughter cell; this was named the W,W::C,C chromatid segregation phenomenon (figure 2) by referring specifically to those DNA (older) strands that were used as templates for DNA replication in the parental cell [23,36].

After discovering the monochromatid gene expression phenomenon in yeast cells [21,22], we searched for the existence of the selective chromatid segregation phenomenon in biology with the goal of exploiting it for understanding multicellular organisms development. The chromosome-specific selective chromatid segregation phenomenon was first discovered in studies of mouse cells. It concerns a Chr. 7 segregation pattern that changes with cell type [23], and the left–right dynein molecular motor was proposed to act at the centromere to execute the selective chromatid segregation function [24]. It was subsequently discovered that Drosophila autosomes usually follow the biased W,W::C,C segregation mode during male germline asymmetric stem-cell division, whereby 50% of stem-cell daughters inherit both template W strands and 50% inherit both template C strands [40]. Moreover, Drosophila chromatids carrying the old histones are delivered to the stem-cell daughter and those carrying new histones are delivered to the other daughter cell differentiating into a different cell type [41]. Thus, chromatids of autosomal chromosomes are selectively segregated both by the nature of their W versus C template strands and independently by recognizing the old versus new histones residing on them for this germline asymmetric cell division. Two independent controls therefore operate for the segregation of Drosophila autosomes during germline cell mitosis. It is worth noting that precedence exists for the discovery of co-segregation of chromosomes containing ‘immortal’ DNA strands undergoing asymmetric stem-cell division in mouse cells [42,43]. Also, a recent study has unambiguously demonstrated that asymmetric inheritance of template strands in the mouse and human embryonic stem cells occurs at a high frequency when stem cells are induced to differentiate into the three primary germ layers in the embryoid bodies [44]. It may seem unusual, but evidence is accumulating to show that the selective chromatid segregation process in some cases might involve the segregation of single DNA strands to sister nuclei during stem-cell division at critical stages during embryogenesis and cancer development, so named the amitotic mitakaryotic cell division [45]. However, in comparison with these examples describing biased segregation of the entire or most of the genome, SSIS concerns biased segregation of one or a set of specific chromosomes to function as an epigenetic mechanism for cellular differentiation. Moreover, different chromosomes might be involved in differentiating cells of different cell types.

The SSIS mechanism is composed of two unrelated phenomena—monochromatid gene expression and of selective chromatid segregation—that function together in mitosis of a specific cell. Although seemingly independent phenomena, the advent of the monochromatid gene expression process had probably provided biological material for the evolution of the selective chromatid segregation phenomenon in diploid organisms. Here, we propose that SSIS operates during mitosis of a limb deterministc progenitor cell to produce one daughter cell that inherits both chromosomally borne transcriptionally expressed DLX5 copies, whereas the other daughter cell inherits both chromosomally borne, epigenetically silenced DLX ‘epialleles’ (figure 2). Such differentiated daughter cells are proposed
to activate or repress different sets of target genes in their progeny cells to help develop features specific to each anatomical region of the limb. Currently, it cannot be determined whether the cell with expressed or the one with silenced DLX genes activates the distal versus the proximal limb tissue developmental programme. Either case equally supports the SSIS mechanism for precisely controlling the expression versus silencing of the DLX5 gene at the single-cell level in the limb deterministic cell. Here, we searched the literature to find SHFM1 cases associated with chromosomal aberrations that can be employed to scrutinize validity of the SSIS mechanism proposed here for limb development (figure 2).

5. The SHFM translocation t(7q11.21;9p12) aetiology supports the SSIS mechanism for limb development

Although the relevant gene's mutation had not been identified, for many years it appeared that only Chr. 7q21 aberrations cause SHFM1 [5], and only recently have recent reports established that conventional mutations in DLX5 cause autosomal-dominant SHFM1 [11,16]. Thus, intragenic DLX5 mutations do cause limb malformations, therefore, the model of DLX5 gene haploinsufficiency for the SHFM1 development is very well established. Inexplicably, however, most of the syndromic SHFM1 cases involve chromosomal aberrations, including translocations of the SHFM1 locus itself or of regions around it. Such translocations within the Chr. 7q21–q22 region exhibit breakpoints that are located hundreds of Kb away from the DLX locus, and inexplicably, breakpoints reside both in the centromere-proximal as well as the telomeric side of the locus. As these translocations do not harbour DLX5 mutations, they were explained by hypothesizing the dissociation of long-range, cis-acting regulatory gene controls, such as enhancer sequences, to cause dysregulation of DLX5 to cause the SHFM1 disorder [4,14,18]. Partially supporting this notion, several SHFM1 locus proximal sequences were shown to bind to the p63 transcription factor at the SHFM1 locus. Moreover, haploinsufficiency of the p63 gene of the SHFM3 locus likewise causes SHFM [13,14]. Clearly, the precise level of expression of both p63 and DLX5 genes is required for healthy limb development.

As stated before, the model of autosomal-dominant, incomplete penetrance with variable expression of DLX5 has been proposed in numerous SHFM1 locus studies, although the authors state the conundrum that the genetic aetiology remains elusive in a substantial proportion of affected individuals because the majority of cases are associated with chromosomal aberrations of the region where subjects harboured the wild-type DLX5 gene [11,46]. In this situation, interpretation of the existing data has not yet unravelled the mechanism of limb development. We advance here the alternative SSIS mechanism to understand the mechanism of DLX5 gene regulation likely required for limb development to help to explain the chromosome aberrations aetiologi of SHFM1 molecularly (figure 2). To scrutinize validity of the SSIS mechanism, here, we chose specifically to focus on the aetiology of t(7q11.21;9p12) translocation associated with SHFM in which the breakpoint in the centromere-proximal side is located over 41 million bases away from the DLX5 gene (figure 3) [47,48]. Understanding its aetiology has remained highly enigmatic and we believe its analysis provides a unique window into the mechanism of limb development. In this family, all three family members, a daughter, her father and a grandfather, carry the translocation-developed malformed limbs. Collectively, five limbs of these three members exhibited SHFM and seven limbs did not, while no other family member without translocation-developed malformation. As the disorder occurs rarely (1 in 18 000 births) in the general public, SHFM of all three members in this family is proposed by authors to have been clearly caused by the translocation [47,48]. In addition, it was pointed out that the authors could not exclude the possibility that Chr. 7q11.21 and/or Chr. 9p12 breakpoints might contribute to the SHFM phenotype. As no known SHFM locus has been mapped to these regions, such a possibility was suggested by previous workers to be unlikely. Here we advance the alternative SSIS mechanism to explain the biological basis of at least this translocation’s aetiology (figure 3).

We speculate here that the translocation interferes with the selective chromatin segregation process of the SSIS mechanism in one-half of limbs because of random segregation operating there (figure 3). As a result, SHFM is predicted to develop when by random chance both developmentally equivalent daughter cells are produced in the limb bud. This anomaly results presumably because the DLX gene product level is insufficient owing to the DLX5ON/DLX5OFF epiallelic constitution and/or both daughter cells have inherited the novel equivalent potential for development; either way, proper development is disrupted. Stated another way, essentially functional haploinsufficiency of DLX5 is generated in one-half of limbs in a

![Figure 3. A hypothetical random Chr. 7;9 translocation chromosome's segregation mechanism.](image-url)
novel way without DLX5 having been mutated there. Equally likely, when the normally found asymmetric cell division occurs in a limb bud owing to the random mode of chromatid segregation, a healthy limb will develop. This predicted outcome was observed (figure 3). Other translocations with breakpoints lying closer to the DLX5 gene (but still located hundreds of kb away from the locus) have been explained by the position-effect dysregulation hypothesis, but the q11 breakpoint of the t(7q11.21;p12) translocation located over 41 million bases away is unlikely to have caused dysregulation of DLX5 by the usually invoked cis-effects hypothesis. It is this feature of the location of the breakpoint far away from the DLX5 locus that has led us to specifically feature this translocation in this communication. Thus, in a novel way SSIS explains all the unexplained features of the autosomal-dominant, incomplete penetrance and haploinsufficiency model previously proposed to explain aetiology of chromosomal aberrations. We surmise that the observation of limbs developing SHFM by random chance while the translocation breakpoint is situated far away from the DLX5 locus provides experimental support for the SSIS mechanism. We propose that the location of the precise translocation breakpoint with respect to the DLX5 gene is irrelevant to the aetiology and that the observation of random chance of developing SHFM in a limb supports the key feature of the SSIS mechanism positing a single deterministic cell in the limb bud where this mechanism is hypothesized to operate.

In comparison with this translocation causing SHFM in nearly one-half of limbs, three different chain-terminating nonsense mutations in heterozygous condition caused SHFM in 25% (seven out of 24) limbs owing to reduced penetrance and/or unequal sex distribution [11]. These differences in penetrance by translocation and nonsense mutations are not statistically significantly different from each other because of small sample size. Alternatively, this difference may indicate alternative ways in which the disorder develops: one way by making developmentally equivalent sister cells in the case of the translocation and another way by haploinsufficiency in the case of nonsense mutations (figure 3). With the limited data available, it is not yet possible to distinguish between these alternatives.

6. The pericentric Chr. 7 inversion aetiology supports the SSIS mechanism for limb development

Familial rearrangements involving Chr. 7 have been reported very infrequently [49]. SSIS proposes that the orientation of the centromere with respect to the DLX5 locus in the chromosome is a key component of the mechanism and that the mechanism operates simultaneously on the two Chr. 7 homologues (figure 2). Therefore, an inversion of the Chr. 7 centromere with respect to the DLX5 locus, when existing in the heterozygous constitution, is predicted to produce symmetric cell divisions in all four limbs deterministic cells. And, as a consequence, SHFM should develop in all limbs of the subject (figure 4). We searched the literature and indeed found such a patient carrying a de novo originating Inv(7;p22;q21.3) pericentric inversion associated with all limbs with SHFM [50]. We surmise that the results of this Chr. 7 inversion with one of the inversion breakpoints located in the centromere-proximal side over 700 Kb away from the DLX5 gene [10] have satisfied a second prediction of the SSIS mechanism; the

![Figure 4](image-url)

Figure 4. A hypothetical SSIS mechanism proposed for the pericentric-inverted Chr. 7 heterozygous individual. The Chr. 7 should follow the usual selective segregation mode as described in figure 2, but chromatids of the centromeric inverted chromosome should segregate with an opposite orientation to generate the indicated W,C::W,C segregation mode for the DLX5 epialleles during mitosis of the deterministic cell. As a result, only developmentally equivalent daughter cells, both containing ON/Off epiallelic constitution, will be produced in each limb: therefore, all limbs are predicted to develop SHFM. Results satisfying the prediction have been described [50]. All other details are the same as those defined in figure 2. The chromosomal line length is not drawn to scale.

7. A DLX5-region inversion supports the SSIS mechanism for limb development

A third genetic test of the SSIS mechanism predicts that an inversion of the DLX5 gene-containing region in the heterozygous constitution should produce symmetric cell divisions in all limb deterministic cells owing to the W,C::W,C selective segregation mode occurring specifically for the DLX5 locus. And consequently, SHFM should develop in all limbs of the inversion carrier (figure 5). Following a literature search we indeed found a subject containing inv(7)(q21.1;q36.3) paracentric inversion associated with all malformed limbs [51].

8. Results of a larger inversion encompassing both centromere and DLX5 regions of Chr. 7 are consistent with the SSIS mechanism for limb development

A fourth genetic test of the SSIS mechanism predicts that a pericentric inversion that includes the centromere and the DLX5 locus together in the same inverted segment should not cause SHFM because the normal DLX epialleles biased segregation mode will not be altered. Indeed, such a karyotype with Inv(7)p22;q22 pericentric inversion, found in three
members of a family, did not cause SHFM [49]; note that DLX5 is located at q21.3. It is worth pointing out that both of these inversions discussed here have one of their breakpoints located at p22 but only one of them caused SHFM. Although the precise locations of the two p22 breakpoints have not been molecularly defined, observations with two pericentric inversions add weight to the earlier suggestion [10] that a presumptive mutation in the p22 region is unlikely to have caused the disorder by the Inv(7)p22q21.3 karyotype because no such disorder-causing mutation mapping in this region had been reported previously. Another t(7q22;1p22) translocation did not cause SHFM in four family members [52]; we interpret this observation to mean that the SHFM locus is not located centromere-distal to the Chr. 7q22 region.

Altogether, results of a translocation described in figure 3, two pericentromeric inversions (one described in figure 4) and DLX5-region inversion (figure 5) have strongly satisfied unique genetics and developmental biology predictions of the SSIS mechanism. The orientation of the DLX5 gene relative to the Chr. 7 centromere appears to be essential for proper differentiation of distal-limb structures in humans. We conclude that Chr. 7 centromeric orientation plays a critical role in the selective chromatid segregation hypothesis of the SSIS mechanism, that Chr. 9 centromere supports a random segregation pattern in the limb deterministic cell and that the SHFM locus resides within the Chr. 7q21.3–q22 region: indeed, that is where the DLX5 locus is situated. Encouraged by four independent corroborations of the SSIS mechanism presented here, we propose that SSIS should be considered as the primary mechanism to define the impact of Chr. 7 aberrations on limb development in future studies, and, perhaps by extension, for understanding the regulatory mechanisms of other developmental control genes in general.

9. SHFM caused by the extra copy of the DLX5/6 region inserted in Chr. 3p21 is consistent with the SSIS mechanism

In humans, malformation-causing mutations act in dominant fashion both through gain or loss of gene function. The SSIS mechanism proposes regulation of the DLX5 gene such that a precise level of gene product is produced in a specific daughter cell but it is silenced in the other daughter cell (figure 2). Accordingly, the observation of Chr. 7q22–q34 region’s insertion into Chr. 3p21 and its association with the three-limb SHFM developmental phenotype of a person [53] are consistent with the SSIS mechanism. Similarly, results of a de novo duplication of the 719 Kb region, harbouring only the DLX5 and DLX6 genes, found in a patient with SHFM [54] are consistent with the SSIS mechanism. We imagine that such rearrangements affecting gene dosage level might also interfere with the production of the precise level of DLX5 gene product in the cell.

10. Further tests of the SSIS mechanism for limb development

Here we have advanced the SSIS mechanism to explain existing data and to guide future research on limb development. Of necessity, we are obliged to focus on only the final stage of the limb malformation phenotype and its association with specific genetic loci. With this limited knowledge we can only speculate on the details of the SSIS mechanism for limb development. It is not yet possible to identify the hypothesized deterministic limb bud cell, let alone to directly demonstrate the hypothesized monochromatid gene expression and the selective chromosome segregation phenomena operating there (figure 2). Despite these technical limitations, we are pleased with the power of the SSIS mechanism to explain the association of genotype with the phenotype. Additional tests of the SSIS mechanism with predicted developmental outcome should be entertained, some are proposed below, and they should be considered in future research.

As noted above, most cases of SHFM are sporadic in nature and of unknown aetiology [6]. According to the SSIS mechanism proposal, possibly one cause of sporadic cases may be the result of spontaneous, rare somatic recombination events occurring between non-sister chromatids in the genetic interval between the centromere and the DLX5 locus in the G2 phase of the cell cycle in the deterministic cell (figure 2). A molecular test of this hypothesis predicts the loss of heterozygosity of single-nucleotide polymorphic molecular markers located near the DLX gene in tissues of the malformed limb. Also, overall much reduced recombination frequency occurring in mitosis in comparison with that of meiotic recombination frequency may have evolved lest recombination interfere with the operation of the SSIS mechanism of cellular differentiation and development (figure 2).

The mouse equivalent Dlx5 and Dlx6 genes mutations cause SHFM, although both copies of both genes must be deleted to generate limb malformation [15,55]. Thus, unlike the situation in humans, haploinsufficiency of these genes does not cause malformation in the mouse. On the other hand, genetic stocks of heterozygous translocations of the Dlx5/6 loci located on mouse Chr. 6 might provide research material for conducting interesting genetic tests of the SSIS mechanism. For example, such a translocation in the heterozygous condition is predicted to produce developmentally equivalent daughter cells owing to random segregation mode, and SHFM should result in 50% of limbs. This way, one can test whether the precise location of the breakpoint with respect to the DLX locus matters and whether only the DLX region’s orientation with respect to centromeric orientation is relevant. Moreover, translocation homozygous animals are predicted to produce healthy limbs.
or limbs with unknown phenotype (flipped limbs?). The existing mouse cell lines harbouring site-specific loxP recombination sites in the genome [56] could be used to produce the required translocations. Generating mouse stocks with genetically engineered inversions of Chr. 6 centromere or of Dlx5/6 genes can provide material for additional genetic tests, equivalent to the analysis we presented above for the human Chr. 7 pericentric inversion.

11. Should the SSIS mechanism be considered as a general mechanism for development?

Embryonic development requires precise regulation of many developmental genes, likely located in different chromosomes, which are expressed and/or silenced simultaneously and in a developmentally programmed manner. Inappropriate expression of developmental genes in tissues is a major cause of developmental anomalies. Analogous to the analysis presented here for understanding the Chr. 7 aberrations’ aetiology, translocations that cause SHFM have provided evidence for the SSIS mechanism operating on Chr. 2 [39]. Notably, three independent Chr. 2q14.1–q14.2 region translocations with breakpoints separated by a relatively large distance of 2.5 million bases, these translocations involving three other chromosomes, were found to be associated with SHFM. Also, the transcriptional regulation of a hypothetical gene located distal to the 2q14.2 breakpoint was speculated to conform to the SSIS mechanism [39]. Supporting this proposal, the SHFM5 locus located far away from the translocation breakpoints at centromere-distal 2q31.1 region contains the HOXD-genes cluster and mutations in this cluster indeed cause SHFM [57]. HOX genes, also known as homeotic genes, comprise a functionally related cluster of genes that famously control the embryonic body plan development along the anterior–posterior axis of multicellular organisms. There are four such gene clusters located in different chromosomes in humans. Their encoded homeodomain-containing proteins function by regulating the timing and extent of local growth rates of tissues required for patterning during limb and external genitalia development [57]. HOX genes function by coordinating the expression of sonic hedgehog, fibroblast growth and other signalling-cascade factors [2]. Interestingly, haploinsufficiency of the HOXD-gene cluster [58] causes SHFM in humans, just as we propose here for the DLX5 gene’s functional haploinsufficiency condition generated instead by the Chr. 7 translocation (figure 3) and by a specific centromeric inversion (figure 4). Thus, the diploid dose of human HOXD and of the DLX5 genes is deemed essential for the patterning of limb tissues along the anterior–posterior and/or the distal–proximal axes in humans. Moreover, the haploinsufficiency feature suggests that both alleles of the developmental gene must be expressed simultaneously in a specific cell, just as we propose here for the regulation of the DLX5 gene though the SSIS mechanism (figure 2).

It was speculated that evolutionary changes in the number of HOX-gene clusters and their expression may have caused body-pattern evolution [59]. The order of specific genes in the HOX clusters has been very well conserved during evolution from worms to vertebrates, and moreover, homeotic transformation of axial structures results when expression of these genes is altered. Curiously, the order of HOX genes in the chromosome is the same as the order of their expression in the anterior–posterior axis of the developing embryo. Reasons for conserving the genes’ structural and functional colinearity relationship in evolution are under debate; we propose here that one reason could be for preserving the SSIS mechanism to operate on clustered genes during embryogenesis.

Interestingly and paradoxically, the HOXD locus is subject to epigenetic regulation through both gene-repressing Polycomb and gene-activating Trithorax factors. These factors possess histone modifying enzymatic activities to promote active versus inactive chromatin states on gene targets. In mouse ES cells, ‘bivalent domain’ promoters carrying both active and repressive chromatin signatures mark the HOXD locus [60]. Note that SSIS requires monochromatid gene expression such that the promoter of the HOXD locus is transcriptionally active in one chromatid and epigenetically repressed in its sister chromatid. Thus, the finding of bivalent domain promoters of the HOXD locus is in accord to what is predicted by the SSIS mechanism. Moreover, chromatid-specific mat1 imprinting occurs at a specific sequence of bases on a specific strand and only when it is synthesized by the lagging-strand replication complex in yeast (reviewed in [27]). Therefore, by SSIS, the chromosomal aberrations spanning both the 5’- and the 3’-regions of the SHFM1 locus might interfere with the function of DLX gene regulatory controls byaltering the direction of replication of these elements. Thus, the SSIS mechanism provides an alternative explanation to the long-range, cis-acting gene regulation controls previously hypothesized to function on both sides of the HOXD [61,62] and the DLX 5 (SHFM1) loci [18]. We predict that both Chr. 2 and Chr. 7 undergo the SSIS mechanism in the specific limb cells and that at least one locus that resides centromere-distal to the breakpoints discussed above for either chromosome is required for limb development.

We point out that mechanistic aspects of some of the features of the generic SSIS model are only known from studies with other systems. It is not known which cell is the deterministic cell conforming to the SSIS mechanism in the developing limb (figure 2). We know of only one case, Caenorhabditis elegans, where an equivalent deterministic cell is discovered to be the one-celled embryo itself that dictates bilateral left–right neuronal asymmetry to develop many cell divisions later in the adult worm by arguably following the SSIS mechanism [63,64]. For accomplishing development, regulated generation of equivalent and non-equivalent daughter cells at specific stages in development is likely required. By the SSIS mechanism, the process of segregation of W,W::C,C chromatids is proposed to produce non-equivalent daughter cells (figure 2), while the hypothetical W,C::W,C segregation mode (similar to that drawn in figure 4) will produce equivalent daughter cells.

One can imagine that mutations in factors required for the specific mode of chromatid segregation will uncover a default segregation mode, which should cause congenital developmental anomalies in mutation carriers. By applying this logic, the SSIS mechanism has been advanced to explain several congenital anomalies that have developed in diverse organisms. These include mouse visceral organs’ laterality development, such as situs inversus and 50% embryonic lethality of the left–right dynavin molecular motor protein mutants [33,34,64], a factor implicated in the Chr. 7 selective chromatid segregation phenomenon that operates during mouse cells mitosis [24]; the generation of bilaterally symmetrical neurons in the C. elegans worm by injecting mutant tubulin message into the one-celled embryo [63,64]; human mirror hand movement disorder development owing to rad51/RAD51 heterozygosity [65]; and the two-coloured wing-spots pigmentation...
development of the Brachus beetle’s Piebald gene mutant [66]. Additionally, the mechanism predicts that chromosomal aberrations consisting of translocations or inversions could hinder the distribution of epialleles of developmental control genes to cause congenital developmental anomalies. Similar to the SHFM malformation resulting from Chr. 7 aberrations discussed in this paper and of Chr. 2 aberrations presented above [39], the human psychoses aetiology associated with various translocations with breakpoints covering over 40% of the Chr. 11q arm have been explained by the SSIS mechanism [36,67].

In all these examples, the authors of the studies that first described those disorders stated that the genotype–phenotype relationships have not been understood. We surmise that the SSIS mechanism has explained the molecular basis of their aetiologies very well. Collectively, different features of the SSIS mechanism are evidenced in studies varying from fission yeasts to invertebrates and vertebrates.

Concerning an unrelated issue in humans, approximately 3% to 5% of autosomal genes undergo monoallelic gene expression, and furthermore, they are chosen randomly from maternal and paternal chromosomes by an unknown mechanism [68]. We hypothesize here that the SSIS process, by employing the W,C::W,C biased segregation mode, may provide the mechanism of mammalian monoallelic expression by thus assuring random choice of maternal and paternal autosomal alleles and of genes subject to X-inactivation.

As stated at the beginning of this communication, the morphogen gradient model has been the main paradigm followed for guiding research on biological pattern formation. This model proposes that the fate of a cell is controlled by its position in the developing embryo and by its response to concentration gradients of morphogens with a consistent directional bias laid across the embryo or the developing organ [3]. Morphogen-like mechanisms do operate in limited cases, such as for the morphogenesis of Dictyostelium slime mould consisting of cells of only two (prespore and prestalk) cell types [69]. It should be noted, however, that the morphogen model does not readily explain the genetic behaviour of a Chr. 7 translocation discussed here, in which no gene mutation has been implicated for SHFM development and also where SHFM develops in some but not in other limbs of the same patient. The morphogen model of visceral organ laterality development in mice, usually explained by the fluid/morphogen flow of rotating cilia found on cells of embryonic node tissue, has been recently questioned [64] because surprisingly as few as only two cilia located at any place on the entire node are sufficient for laterality development [70]. Moreover, to explain the generation of asymmetric cell division, such as of stem cells, a model proposing the long-range-acting morphogens is unlikely to be the mechanism of cell-fate determination differentially operating on adjoining cells with the precision required. We suspect that the molecules demonstrated to function in cell-to-cell and local, close-range signalling processes might have been misinterpreted as providing convincing support for the morphogen model. Note for example, that the famous Wnt (wingless)/planar cell polarity signalling factor is thought to act as a diffusible morphogen by eliciting long-range-acting cellular controls; stunningly, however, Drosophila embryos develop normally after Wnt diffusion is prohibited by tethering the genetically engineered Wnt protein to the membrane of the cell producing it [71].

By realizing these developments, further studies should be designed to test whether an epigenetic SSIS-like mechanism operates in diverse organisms. Concerning the SHFM1 disorder, there is no shortage of molecular and mechanistic studies published in prominent journals for decades, but so far its molecular aetiology has remained undefined. While considering this point, workers in the field would benefit by heeding the advice from Dr James A. Birchler [74], who pointed out that the perception of good research emphasizing only descriptive, mechanistic studies is hindering science because new ideas remain unappreciated. Lacking mechanistic details at the outset, new ideas are considered too risky, unfit both for funding and for publication in prominent journals. The monochromatid differentiation concept, based on DNA strands’ chirality for achieving asymmetric gene expression of sister cells, is one such a new idea. Because it has been found to function only in two fission yeasts [27], and due to technical difficulties, it has not been researched whether it operates in any other organism. This might create doubt of SSIS’s existence in biology at large because assumptions/details of the SSIS mechanism remain unknown. Concerning limb development, SSIS provides a conceptual framework with which to understand vast amounts of data, a challenge posed by Dr Brenner quoted at the start of this communication [1]. Our thesis should help guide future research on the elusive mechanism of limb development. In sum, evidence summarized here supports our thesis that strand chirality of the double-helix structure of DNA provides the physical basis for mysteries of development during cell proliferation [21,22,27,36,39,65–67], a function that is performed in addition to providing the basis of heredity [28] by following the Mendelian principles of genetics primarily operating in meiosis.

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References

1. Brenner S. 2012 Tuning centenary: life’s code script. Nature 482, 461. (doi:10.1038/482461a)
2. Zeller R, Lopez-Rios J, Zuniga A. 2009 Vertebrate limb bud development: moving towards integrative analysis of organogenesis. Nat. Rev. Genet. 10, 845 – 858. (doi:10.1038/nrg2681)
3. Wolpert L. 1969 Positional information and the spatial pattern of cellular differentiation. J. Theor. Biol. 25, 1–47. (doi:10.1016/S0022-5193(69)80016-0)

4. Bernardini L, Palka C, Cecarini C, Capalbo A, Bottillo I, Mingarelli R, Novelli A, Dallapiccola B. 2008 Complex rearrangement of chromosomes 7q21.13-q22.1 confirms the ectodactyly-deafness locus and suggests new candidate genes. Am. J. Med. Genet. A 146A, 238 – 244. (doi:10.1002/ajmg.a.32093)

5. Scherer SW et al. 1994 Fine mapping of the autosomal-dominant split hand split foot locus on chromosome-7, band q21.3-q22.1. Am. J. Hum. Genet. 55, 12 – 20.

6. Czezel AE, Vitez M, Kodaj I, Lenz W. 1993 An epidemiological study of isolated split hand/foot in Hungary. J. Med. Genet. 30, 593 – 596. (doi:10.1136/jmg.30.7.593)

7. Shubin N, Tabin C, Carroll S. 1997 Fossils, genes and the evolution of animal limbs. Nature 388, 639 – 648. (doi:10.1038/41770)

8. Elliott AM, Evans JA. 2006 Genotype–phenotype correlations in mapped split hand foot malformation (SHFM) patients. Am. J. Med. Genet. A 140, 1419 – 1427. (doi:10.1002/ajmg.a.31244)

9. Babbs C, Heller R, Everman DB, Crocker M, Twigg SR, Schwartz CE, Giele H, Wilkie AOM. 2007 A new locus for split hand/foot malformation with long bone deficiency (SHFD1) at 2q14.1 identified from a chromosome translocation. Hum. Genet. 122, 191 – 199. (doi:10.1007/s00439-007-0390-7)

10. van Silfhout AT et al. 2009 Split hand/foot malformation due to chromosome 7q aberrations (SHFM1): additional support for functional haploinsufficiency as the causative mechanism. Eur. J. Hum. Genet. 17, 1432 – 1438. (doi:10.1038/ejhg.2009.72)

11. Sowinska-Seidler A, Socha M, Jamsheer A. 2014 Split hand/foot malformation with autosomal-dominant split hand/foot malformation of a Chinese family with a heterozygous DLX5 mutation in a Chinese family. Hum. Mol. Genet. 27, 479 – 481. (doi:10.1136/jmg.27.8.479)

12. Crackower MA et al. 1996 Characterization of the split hand/split foot malformation locus SHFM1 at 7q21.3-q22.1 and analysis of a candidate gene for its expression during limb development. Hum. Mol. Genet. 5, 571 – 579. (doi:10.1093/hmg/5.5.571)

13. Shamseldin HE, Faden MA, Alshamr W, Alkuraya FS. 2012 Identification of a novel DLX5 mutation in a family with autosomal recessive split hand and foot malformation. J. Med. Genet. 49, 16 – 20. (doi:10.1136/jmedgenet-2011-100556)

14. Spitz F, Herkener C, Morris MA, Dubrule D. 2005 Inversion disruption of the Hand cluster leads to the partition of regulatory landscapes. Nat. Genet. 37, 889 – 893. (doi:10.1038/ng1597)

15. Klar AJS. 1987 Differentiated parental DNA strands confer developmental asymmetry on daughter cells in fission yeast. Nature 326, 466 – 470. (doi:10.1038/326466a0)

16. Klar AJS. 1990 The developmental fate of fission yeast cells is determined by the pattern of inheritance of parental and grandparental DNA strands. EMBO J. 9, 1407 – 1415.

17. Armakolas A, Klar AJS. 2006 Cell type regulates the epigenetic mechanism of asymmetric cell division. EMBO J. 25, 6791 – 6795. (doi:10.1038/emboj.20130710)

18. Watson JD, Crick FH. 1953 Molecular structure of nucleic acids; a structure for deoxyribose nucleic acid. Nature 171, 737 – 738. (doi:10.1038/171737a0)

19. Meiselson M, Stahl FW. 1958 The replication of DNA of Schizosaccharomyces pombe. Proc. Natl Acad. Sci. USA 5, 681 – 682. (doi:10.1073/pnas.54.8.671)

20. Watson JD, Crick FH. 1953 Molecular structure of nucleic acids; a structure for deoxyribonucleic acid. Nature 171, 737 – 738. (doi:10.1038/171737a0)

21. Roberts SH, Hughes HE, Davies SJ, Meredith AL. 1991 Bilateral split hand and split foot malformation in a boy with a de novo interstitial deletion of 7q21.3. J. Med. Genet. 27, 479 – 481. (doi:10.1136/jmg.27.8.479)

22. Klar AJS. 1990 The developmental fate of fission yeast cells is determined by the pattern of inheritance of parental and grandparental DNA strands. EMBO J. 9, 1407 – 1415.

23. Armakolas A, Klar AJS. 2006 Cell type regulates the epigenetic mechanism of asymmetric cell division. EMBO J. 25, 6791 – 6795. (doi:10.1038/emboj.20130710)

24. Kimble J, Wood WD. 1975 A device for the activation of yeast cells. Proc. Natl Acad. Sci. USA 72, 3962 – 3966. (doi:10.1073/pnas.72.8.3962)

25. Klar AJS. 1994 A model for specification of the left–right axis in vertebrates. Trends Genet. 10, 392 – 396. (doi:10.1016/0168-9525(94)00055-B)

26. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Genetics 160, 1745 – 1747.

27. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

28. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

29. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

30. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

31. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

32. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

33. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

34. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

35. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

36. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

37. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

38. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

39. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.

40. Klar AJS. 2004 A genetic mechanism implicates chromosome 11 in schizophrenia and bipolar affective disorders. Nature 410, 1745 – 1747.
46. Lango Allen H, Caswell R, Xie W, Xu X, Wragg C, Timpenny PD, Turner GL, Weedon MN, Ellard S. 2014 Next generation sequencing of chromosomal rearrangements in patients with split-hand/split-foot malformation provides evidence for DNMT1/ exon enhancers of DLX5/6 expression in humans. J. Med. Genet. 51, 264 – 267. (doi:10.1136/jmedgenet-2013-102142)

47. Hasegawa T. 1991 EEC syndrome (ectodactyly, ectodermal dysplasia and cleft lip/palate) with a balanced reciprocal translocation between 7q11.21 and 9p12 (or 7p11.2 and 9q12) in three generations. Clin. Genet. 40, 202 – 206. (doi:10.1038/clingen.1991.1)

48. Fukushima Y, Ohashi H, Hasegawa T. 1993 The EEC syndrome (ectrodactyly, split hand/foot malformation and inv(7)(p22q21.3). Zwierstra RP, Leegte B, Castedo S. 1995 Bilateral sampling.

49. Gordon CT, Krasnewich D, White B, Lenane M, Naritomi K, Izumikawa Y, Tohma T, Hirayama K. 1991 EEC syndrome (ectrodactyly, split hand/foot malformation and inv(7)(p22q21.3). EEC syndrome (ectrodactyly, split hand/foot malformation and inv(7)(p22q21.3). J. Med. Genet. 46, 492 – 493. (doi:10.1002/ajmg.1320460005)

50. Loboledo RF, Rajan LX. 2002 L. Luftkin T. The Dlx5 and Dlx6 homeobox genes are essential for craniofacial, axial, and appendicular skeletal development. Genes Dev. 16, 1089 – 1101. (doi:10.1101/gad.988402)

51. Yu Y, Bradley A. 2001 Engineering chromosomal rearrangements in mice. Nat. Rev. Genet. 2, 780 – 790. (doi:10.1038/sj.nrg3593564)

52. Duboule D. 1992 The vertebrate limb: a model system to study the Hox/HOM gene network during development and evolution. BioEssays 14, 375 – 384. (doi:10.1002/bies.950140606)

53. Del Campo M, Jones MC, Venaka AN, Curry CI, Jones RL, Mascarello JF, Ali-Kahn-Catts Z, Drumheller T, McGinns W. 1999 Monodactylyal limbs and abnormal genitalia are associated with hemizygosity for the human 2q31 region that includes the HOXD cluster. Am. J. Hum. Genet. 65, 104 – 1206. (doi:10.1086/302467)

54. Hughes CL, Kaufman TC. 2002 Hox genes and the evolution of the arthropod body plan. Science 41, 1089 – 1101. (doi:10.1126/science.1171488)

55. Soshnikova N, Duboule D. 2009 Epigenetic temporal control of mouse Hox genes in vivo. Science 324, 1320 – 1323. (doi:10.1126/science.1171488)

56. Duboule D. 2009 Epigenetic temporal control of mouse Hox genes in vivo. Science 324, 1320 – 1323. (doi:10.1126/science.1171488)

57. Loomis WF. 2015 Genetic control of morphogenesis in Drosophila. Dev. Biol. 402, 146 – 161. (doi:10.1016/j.ydbio.2015.03.016)

58. Shimohara K et al. 2012 Two rotating cilia in the node cavity are sufficient to break left–right symmetry in the mouse embryo. Nat. Commun. 3, 1 – 8. (doi:10.1038/ncomms1624)

59. Alexandre C, Baena-Lopez A, Vincent JP. 2014 Patterning and growth control by membrane-tethered Wingless. Nature 505, 180 – 185. (doi:10.1038/nature12879)

60. Vandenberg LN, Levin M. 2013 A unified model for left–right asymmetry? Comparison and synthesis of molecular models of embryonic laterality. Dev. Biol. 379, 1 – 15. (doi:10.1016/j.ydbio.2013.03.021)

61. Richardson MK. 2009 Diffusible gradients are out—molecular models of embryonic laterality. Dev. Biol. 379, 1 – 15. (doi:10.1016/j.ydbio.2013.03.021)

62. Riddle R 2010 Control of vertebrate Hox clusters by remote and global cis-acting regulatory sequences. Adv. Exp. Med. Biol. 689, 63 – 78. (doi:10.1007/978-1-4419-6637-5_4)

63. Lobbink M, Wang G, Xu G, Hsieh YY, Chuang CF, Lemire JM, Levin M. 2012 Early, nonciliary role for pericentric hybridization.

64. Sauer S, Klar AJS. 2012 Left–right symmetry breaking in mice by left–right dynein may occur via a biased chromatid segregation mechanism, without directly involving the Nodal gene. Front. Oncol. 2, 1 – 10. (doi:10.3389/fonc.2012.00166)

65. Klar AJS. 2014 Selective chromatid segregation mechanism invoked for the human congenital mirror hand movement disorder development by RAD51 mutations: a hypothesis. Int. J. Biol. Sci. 10, 1018 – 1023. (doi:10.7150/ijbs.9886)

66. Klar AJS. 2015 Selective chromatid segregation mechanism for Brachus wing spots piebald color. Front. Biosci. 7, 322 – 333. (doi:10.2741/c734)

67. Klar AJS. 2014 Selective chromatid segregation mechanism explains the etiology of chromosome 11 translocation-associated psychiatric disorders: a review. J. Neurol. Disord. 2, 1 – 6.

68. Gimelbrant A, Hutchinson JN, Thompson BR, Chess A. 2007 Widespread monoallelic expression on human autosomes. Science 318, 1136 – 1140. (doi:10.1126/science.1148910)

69. Birchler JA. 2015 Mendel, mechanism, models, marketing, and more. Cell 163, 9 – 11. (doi:10.1016/j.cell.2015.09.008)