How is developmental stability sustained in the face of genetic variation?

JAMES R. S. WHITTLE*

School of Biological Sciences, University of Sussex, Brighton, United Kingdom

ABSTRACT The number and arrangement of scutellar bristles on the thorax of Drosophila melanogaster is largely invariant in wild-type stocks. This character therefore appears to be buffered against changes in phenotype, and has previously been described as a canalized character. Mutations that do alter this phenotype increase the variability in bristle number and can reveal otherwise cryptic genetic differences at other loci. This phenomenon is examined and possible mechanisms contributing to stability of this developmental event are discussed, but the notion that the character is canalized is found not to be heuristic.

KEY WORDS: developmental stability, bristle pattern, canalization

Introduction

Like begets like. Fascination with the phenomenon of reproduction drives us to understand the mechanisms underpinning the process of development, including an explanation of the regularity with which a particular animal form develops from each fertilized egg of that same species. I want to look at a particular developmental process, the spatial patterning of scutellar bristles, which are sense organs on the Drosophila melanogaster adult. We know that the properties (and thus the behavior) of cells and groups of cells that compose tissues reflect the abundance and types of different proteins that they contain. Each of those proteins is encoded by a gene and the protein structure derives directly from the translation or ‘expression’ of the gene in that particular cell. When geneticists sample a natural population and examine the DNA sequence of a gene or the amino acid sequence of its product, they find considerable variability or heterogeneity within and between individuals. A population contains a large reservoir of genetic variants, yet most of them do not cause detectable phenotypic changes. I want to see whether progress in our understanding of the ‘developmental machinery’ offers any hints regarding the basis of developmental stability. The first point is to explain how geneticists can untangle the complexities of development.

Geneticists have devised successful and effective methods for ‘dissecting’ a developmental process

Geneticists use perturbation analysis, often calling it ‘mutational dissection’, to identify genetic components in a biological process. I will explain how this method brings us to two conclusions: first that all these genes must normally contribute to the reliability or stability of the process, and second, that if many genes are involved in the process this does not seem to make it more ‘fragile’ than if fewer genes were involved. The ‘logic’ of mutational dissection runs as follows. If we mutagenize animals and then breed from them to produce homozygotes in the F2 generation, then any mutation that has altered the integrity, abundance or stability of a gene product involved in development will be recognizable in the F2 segregation because of the upset it creates. If the mutation is recessive in its effect, then it can be described and catalogued, and maintained as a heterozygous stock displaying a wild-type phenotype. Crosses between different recessive mutations make clear whether any two mutations found are in the same gene or are changes in two quite different genes (the complementation test). Each mutation will have identified a gene that must be essential for the developmental event: otherwise why would any change in it have shown up? Locating the site of each mutation in the chromosomes provides a useful step towards obtaining the DNA sequence for that gene. When a gene has been sequenced, its protein product can be deduced by ‘decoding’ it or can actually be synthesized in vitro. The ‘expression’ of that gene can be documented by searching tissues for the messenger RNA or the protein. Combining the descriptive knowledge of what goes wrong when the mutant version of a gene replaces the wild-type one with the molecular information about the protein product usually provides a reasonable account of how

Abbreviations used in this paper: Km, Michaelis constant; RFLP, Restriction Fragment Length Polymorphism; SOP, sense organ precursor; Vmax, maximum velocity.

*Address for reprints: School of Biological Sciences, University of Sussex, Brighton BN1 9QG, United Kingdom. FAX: 0044-1273-678433. e-mail: j.r.s.whittle@sussex.ac.uk

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the activity of that gene contributes to the process. It is self-evident that normality depends upon every single one of the genes found by ‘mutational dissection’ being operational.

A long-established empirical finding is that if one induces new mutations in a genotype that already has a mutant gene affecting a particular phenotype, then it generally becomes easier to detect new mutations in other genes affecting the same phenotype. This has been of great benefit in mutational dissection, and there are two stratagems. The first is to start with a stock showing a mutant phenotype and to recover mutations in other genes that act as suppressors or enhancers of that phenotype. There are situations in which a newly induced mutation is only detectable in a genotype that is already mutant for another gene, and many instances where mutations of a subtle kind would have been overlooked except in the ‘mutant background’. For example, we have successfully isolated mutations in other genes which when heterozygous (therefore they are ‘dominant’ mutations) modify the phenotype of homozygotes for the mutation spade in the wingless gene (Buratovich et al., 1997). A second example also features the wingless gene which is necessary for the appearance of the scutellar bristles (Phillips and Whittle, 1993). Flies that are heterozygous for a mutation in wingless, encoding a secreted ‘signal’ protein, and that are also simultaneously heterozygous for a mutation in arrow, which has a role in the response of cells to this ‘signal’, have a low frequency of leg defects. Heterozygous mutations at a third locus can enhance the frequency of the leg defect. This enhancement can be used to identify related genes in the same process (Kim A. Caldwell, S. Dougan, E. Matunis, J. Tran, and S. DiNardo, personal communication).

The second stratagem has been to use a heterozygote for a recessive mutation (it therefore has a wild-type phenotype) as a background in which to search for phenotypic change in this heterozygote, either when one of its parents has been subjected to mutagenesis, or when the heterozygote also carries a known deletion or additional copy of genetic material elsewhere in the genome. Antonio García-Bellido pioneered this latter approach in Drosophila, calling it “gene-dose titration analysis” (Botas et al., 1982). A way to interpret the genetic situation underlying this approach is to say that the attenuation or loss of activity of one gene product makes the developmental process more sensitive to small changes in the activity of other gene products. Using mutations in different genes to subtract successively more proteins that are part of the same ‘developmental machine’ makes the system more labile or error-prone, often in a multiplicative way (each successive mutation having a larger de-stabilising effect than the previous one). This is a familiar experience in relation to failures in components of designed machinery, like those of a car. This analysis also reminds us that whether the effect of a mutation is recessive or dominant can depend upon the composition of the rest of the genotype and should not be thought of as some absolute molecular property of that DNA sequence.

A case study: the pattern of scutellar bristles

I have chosen to consider the spatial patterning of large bristles (sense organs of the peripheral nervous system) on a part of the dorsal thorax called the scutellum. In wild-type flies, the pattern and number of these bristles (four) is almost invariant (Fig. 1a). The scutellum comes from the fusion during pupation of two separate (and mirror-imaged) groups of cells (called the wing imaginal discs) found within the larva on the left and right sides. I will recount the history of the scutellar bristles in the reverse order to that which occurs during development (see also the paper by Modolell and Campuzano, this issue). Each functioning bristle, including its neuron connecting it to the central nervous system, is the product of the co-ordinated differentiation of a clone of four cells from a ‘sense organ precursor’ cell (called an SOP). The SOPs appear within a ‘proneural cluster’ of cells, a group related by position rather than by cell lineage, and all of which express the gene scute (of the achaete-scute cluster of genes, García-Bellido and Santamaría, 1978; see also the paper by Campos-Ortega in this issue) which is inactive outside each cluster. The borders of the cluster and the scutellar SOPs themselves appear at fixed sites in the disc epithelium so the pattern and number of scutellar bristles must depend upon the precision of the expression of scute.

This is a specific instance of the puzzle about spatial patterning that has tantalized developmental biologists for many years. The presence of active Scute product depends upon the integrity of at least three other genes, wingless, pannier and extramacrochaetae. Of these, the last two must ‘repress’ scute, because loss-of-activity mutations in these genes prompt more bristles to form, while the first one must ‘activate’ scute, for its mutations result in a disappearance of bristles. The Wingless protein is secreted, and therefore has one of the properties of a signal molecule, but originates very locally in the...
tissue. There is evidence for the implication of two other secreted ‘signalling’ molecules participating in the process, the products of the genes *decapentaplegic* and *hedgehog*. It is known in other systems that the binding of the protein Hedgehog to its receptor protein Patched removes a negative regulatory effect upon the signal transduction in the recipient cell. Because cells homozygous for mutations in the *patched* gene make more bristles than expected on the scutellum (Roberts, Phillips, Warner and Whittle, unpublished), this suggests that the Hedgehog/Patched signaling system contributes to the determination of the scutellar bristles. In contrast, the inactivation or removal of a kinase enzyme by mutations in the gene *zeste-white 3 shaggy* causes extra cells to behave as if they have received the Wingless signal and to become SOPs, resulting in the formation of extra bristles. Interestingly, clones of cells defective for the kinase Shaggy respond as if the signal from their wild-type neighbors is propagated through the tissue from its posterior edge (Warner, Phillips and Whittle, unpublished).

**The concept of developmental canalization**

Waddington (1942) coined the term canalization to describe the ability of some developmental processes to persist in progressing to their normal end-point in the face of genetic or environmental disturbances. In a series of papers from 1959 onwards, J. M. Rendel and A. S. Fraser and their colleagues explored canalization with reference to scutellar bristle number (Rendel 1959,1967; Fraser, 1970). Rendel introduced a recessive X-linked mutation sc 

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from the gene *scute* into a wild population so that it was segregating. He tracked the mutation in his experiments from one generation to the next by the closely linked recessive mutation *white* (eyes). White-eyed flies had the *scute* mutation whilst red-eyed flies had the wild-type copy of *scute*. Mutant homozygotes had fewer scutellar bristles but there was more variability in number. In this population, the mean scutellar bristle number of *sc 

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males was 0.92, comprising flies with 0,1,2 or 3 bristles, whereas the wild-type males all had four scutellar bristles. Selection for increased scutellar bristle number on the *scute* sibs in this population caused the mean scutellar bristle number to rise towards four (to 2.81 in males, comprising flies with 1,2,3 or 4 bristles) showing that at least some of the variability must have been genetic. In the wild-type sibs, sharing all their genotype in common with these flies except for the region close to *scute* and *white*, the bristle number only rose to 4.06 (188 males with four bristles and 12 males with five bristles).

As the mean scutellar bristle number of the *scute* individuals in the selected population approached four, the *variability* in bristle number in this genotype fell again. This outcome was described by saying that selection had moved the process back into a ‘buffered zone’ at four bristles, despite the presence now in the genotype of the *scute* mutation. He concluded that the ‘buffering’ or ‘canalization’ could not be a function of the *scute* gene (because that was mutant), but that the introduction of the *scute* mutation had upset the process sufficiently so that previously cryptic genetic variation was now revealed and could then be seized upon in the selection response. Rendel went on to postulate that there was a normally distributed underlying propensity to form these bristles (which he called ‘make’), and he used a statistical transformation to estimate the ‘distances’ between the 2-3, the 3-4 and the 4-5 bristle thresholds for this hypothetical underlying variable. His analysis showed that the distance between the 3-4 and the 4-5 thresholds (the ‘4’ class) was larger than that of the other intervals. The canalization concept in relation to scutellar bristle pattern has been defended (Waddington, 1972), and critically reviewed by Scharloo (1991).

Thirty eight years later, and having the benefit of hindsight, I do not consider that Rendel’s proposal of a underlying hypothetical variable with normal distribution, ‘make’, as part of an explanation for scutellar bristle variation, is any longer an heuristic notion, for the following reasons. SOPs for the left and right scutellar bristles are detectable in the two wing discs whilst they are quite separate structures, rather than part of a common ‘unit’ within which the control of SOP number could be exerted. The hypothetical ‘make’ distribution therefore relates poorly to any biologically meaningful single entity. Second, there was debate in the literature at the time as to whether the statistical consideration ought to have been the individual bristle site and the bristle/no bristle threshold (Robertson, 1956) rather than the sum total of scutellar bristles (see also Scharloo, 1991). Simultaneous losses at one site and increases at the other site in the same fly have been seen. The anterior and posterior bristles are determined at different developmental ages (Poody, 1975). Many authors have shown that flies with five scutellar bristles are most likely to have their extra bristle at an anterior site, which suggests that the developmental events preceding the appearance of the anterior and posterior scutellars cannot be identical. Counting together bristles at anterior and posterior sites therefore sacrifices important spatial information about this developmental event. These objections, together with the accumulating knowledge of molecular events in the history of scutellar bristle formation, seem to leave the concept of canalization without utility in this example.

**How might attributes of genes and gene products confer stability upon a developmental process?**

First, stability might arise from the ‘logic’ of the system, in other words, from the molecular or cellular ‘circuitry’ or connectivity that has evolved. The formal representation of control circuitry, including sequences of routines and ‘decisions points’, for which the ‘output’ usually remains constant in the face of fluctuations in different ‘inputs’, is familiar to us outside the biological sphere. Negative feedback is well known from engineer-designed self-regulating machines, and molecular counterparts have been found in biological systems, particularly in the area of biosynthetic pathways. Positive feedback systems, able to amplify a ‘firm’ and fixed response from a very small initial ‘bias’ or signal, are characteristic of, for example, electronic detector systems for photon counting.

The idea being put forward here is that the developmental machinery may have the property of robustness. A recent paper by Barkai and Leibler (1997) proposes that there may be mechanisms for robust adaptation in simple signal transduction networks, although they consider a prokaryotic example. My suggestion is that the behavior of a developmental ‘routine’ or response might be derived from the way the elements of the network are connected rather than from the precise kinetic properties of each molecular component. The prediction would be that if such connectivity conferred stability, then a mutation that severed a feed-forward or feed-back loop altogether would severely upset the behavior while a mutation that changed the abundance or the specific activity of
an individual protein involved might not. The system would therefore be insensitive to (some) variation in the genes encoding the individual components.

A second alternative explanation is that the stability of a developmental process may be an outcome of the kinetic properties of the total assembly of component gene products. The work of H. Kacser and his colleagues (Kacser and Burns, 1973) has clearly shown that ‘through-put’ or flux in a metabolic pathway comprising several coupled enzymes is relatively insensitive to small changes in the catalytic properties of each individual enzyme. Gene products that have catalytic properties (rather than structural roles) might give a system a flexibility or tolerance to changes precisely because of their particular properties (for instance, the enzyme parameters Km and Vmax in a cascade of phosphorylation events of the type that mediate signal transduction from some receptors). Such an imagined structure should be relatively insensitive to protein changes affecting these parameters stemming from alleles segregating in the population.

A third explanation for stability is redundancy. Geneticists have long recognized that if a genome carries duplicate copies of genes (as an autopolyplid does), then the species would be relatively refractory to phenotypic change following mutation in one of the several copies of the same coding unit. In common parlance this would be described as ‘safety in numbers’ but it only predicts stability of phenotype in the face of mutation and offers no explanation of any dynamic buffering of the developmental process following fluctuations in other ‘inputs’ or parameters. Wilkins (1997), has suggested that canalization may in part reflect the contributions of paralogous genes to a given process. Paralogs are gene products closely related structurally (meaning that their DNA sequences are similar). The effect of removing one paralog by genetic ‘knock-out’ or by disruption may be masked by the contribution of paralogs encoding similar proteins, if they are present in the same cells at the same time. In the absence of one particular gene product, the activity of a second and related protein present becomes relevant. The missing protein might have the higher affinity or efficiency in the process but in its absence, even a ‘poor fit’ alternative might protect the developmental process from the consequence of a total loss of a crucial component or molecular activity. Wilkins (1997) cites a number of examples where it is clear that phenotypic change is more severe when both (of two) members of a paralogous family of genes are mutant than when either is singly mutant.

A fourth explanation for developmental stability is provided by homeostasis at the level of cell behavior. The ‘machinery’ driving cell division within an imaginal disc is an excellent example self-regulating process of a higher level than the previous examples. García-Bellido and his colleagues (García-Bellido et al., 1973) were pioneers in recognizing that, within a group of cells that share the same developmental fate (he called it compartment), the cell lineage history was indeterminate rather than fixed. They showed this by creating genetic mosaics with cell-limited differing growth parameters and observing that heterogeneity in cell cycling time, a central cell process, was without any effect upon the size or the fidelity of the developmental patterns (for instance, the scutellum and its bristles) formed by that compartment of cells. This property confers on the compartment of cells a stability in the face of stochastic cell loss or local growth impairment. More recently, they have documented the fact that despite the indeterminate lineage within a compartment, there is a regular dynamic pattern of cell division with respect to position within the disc. This implies a high-level and dynamic interaction between the cells of a compartment within a disc so that in successive intervals of developmental time, ‘permission to divide’ is reassigned according to a spatial pattern irrespective of the previous (and possibly unequal) division contributions of those cells (Milan et al., 1996a,b).

Conclusions

Scutellar bristle pattern and number in Drosophila show little variability, even in the face of segregating genetic differences. We conclude that the developmental events involved seem to be ‘buffered’ or dampened against changes. Yet there is a low level of genetic variation affecting scutellar bristles in natural populations. Many lines with increased scutellar bristle number have been created by selection in samples from wild-type populations (e.g., Fig. 1b, taken from Payne 1918; Fraser, 1970). Quantitative trait loci affecting bristles in wild populations have been mapped using restriction fragment length polymorphisms (RFLPs), and have identified genetic variants at or in the immediate vicinity of all the following loci known to developmental biologists analyzing bristle formation: achaete-scute, Notch, bobbed, daughterless, scabrous, extramacrochaetae, hairy, Delta and Enhancer of split (Falco and Mackay, 1996). Many gene products indeed participate during the specification of bristles (see Simpson 1997, for a recent review).

Work towards a complete genetic inventory and detailed descriptions of the properties of the gene products is yielding secrets about individual steps in the ‘developmental mechanics’, and this reductionist approach to analysis is still in the ascendancy. We are approaching the point at which there could be a complete description of the gene expression events presaging the appearance of each SOP. At that point, it will be valuable to model the process to see if we indeed have sufficient understanding to predict and then test experimentally the stability in behavior of the process, as has been achieved for the behavior of gap genes in segmentation in Drosophila by H. Kitano (Kitano et al., 1998). Provided that the model can subsume all the known information and makes verifiable new predictions, then it would suggest that there are unlikely to be other important attributes contributing to stability that are not rather obvious emergent properties following from the integrated function of the individual genetic components.

In other words, developmental stability of this character is unlikely to be an emergent property beyond the reach of the molecular biology and genetics of contemporary developmental biologists, and is more obvious and less mysterious than the concept of canalization might have suggested.

A personal perspective

I am proud to have known Antonio García-Bellido, to have worked in his laboratory as a visitor where, in 1973, I heard about compartments, and the power of genetic mosaics in understanding the bithorax complex. I acknowledge the continuing influence of his inspiration and vision. In 1967 my doctoral work was on the mapping of gene differences affecting scutellar bristle number found in selected lines from wild populations (Whittle, 1967), after which I vowed never to count another bristle and moved to Paramecium genetics in Bloomington, Indiana where Fernandus Payne, then an Emeritus faculty member, presented me with a
copy of his pioneering study on genetic analysis of scutellar bristles (Payne, 1918 and Fig. 1). In 1997 I find myself looking at scutellar bristle formation again with molecular tools. One hopes this is humility in the presence of an intriguing biological problem rather than the recurrence of narrow obsessional behavior flying in the face of reason!

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