

# Positional information, in bits

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**Cells in a developing embryo have no direct way of “measuring” their physical position. Through a variety of processes, however, the expression levels of multiple genes come to be correlated with position, and these expression levels thus form a code for “positional information.” We show how to measure this information, in bits, using the gap genes in the *Drosophila* embryo as an example. Individual genes carry nearly two bits of information, twice as much as would be expected if the expression patterns consisted only of on/off domains separated by sharp boundaries. Taken together, four gap genes carry enough information to define a cell’s location with an error bar of  $\sim 1\%$  along the anterior/posterior axis of the embryo. This precision is nearly enough for each cell to have a unique identity, which is the maximum information the system can use, and is nearly constant along the length of the embryo. We argue that this constancy is a signature of optimality in the transmission of information from primary morphogen inputs to the output of the gap gene network.**

gene regulatory networks | embryonic development | optimization

**B**uilding a complex, differentiated body requires that individual cells in the embryo make decisions, and ultimately adopt fates, that are appropriate to their position. There are wildly diverging models for how cells acquire this “positional information” (1), but there is general consensus that they encode positional information in the expression levels of various key genes. A classic example is provided by anterior/posterior patterning in the fruit fly, *Drosophila melanogaster*, where a small set of gap genes and then a larger set of pair rule and segment polarity genes are involved in the specification of the body plan (2). These genes have expression levels that vary systematically along the body axis, forming a blueprint for the segmented body of the developed larva that we can “read” within hours after the start of development (3).

Although there is consensus that particular genes carry positional information, less is known quantitatively about how much information is being represented by the expression levels in individual cells. Do the broad, smooth expression profiles of the gap genes, for example, provide enough information to specify the exact pattern of development, cell by cell, along the anterior/posterior axis? How much information does the whole embryo use in making this pattern? Answering these questions is important, in part, because we know that crucial molecules involved in the regulation of gene expression are present at low concentrations and even low absolute copy numbers, so that expression is noisy (4–10), and this noise must limit the transmission of information (11–14). Is it possible, as suggested theoretically (15–18), that the information transmitted through these regulatory networks is close to the physical limits set by the irreducible randomness of counting individual molecular events? To answer this and other questions, we need to measure positional information quantitatively, in bits. We do this here using the gap genes in *Drosophila* as an example.

There are many ways in which positional information could be represented during the process of development. Cells could make decisions based on the integration of signals over time or by comparing their internal states with those of their neighbors.

Eventually, the internal state of each individual cell must carry enough information to specify that cell’s fate, but it is not clear at what point in development this happens. Thus, when we look at the gap genes during the 14th nuclear cycle after fertilization, there is no guarantee that their expression levels will carry all the information that cells eventually will acquire, either from maternal inputs or via communication with their neighbors. Because our experimental methods give us access to snapshots of gene expression levels, however, we will start by asking how much positional information is carried by local measurements in individual cells at a moment in time. These expression levels themselves reflect an integration of many inputs over space and time (9, 19), but these molecular mechanisms do not influence the definition or measurement of the information that the expression levels carry.

## Quantifying Information

In the early stages of development, different cells have essentially the same morphology, at least in the bulk of the embryo, away from the poles. Thus, if we do not look at the expression levels of the relevant genes, we have no information about the position of the cell; it could be anywhere along the anterior/posterior axis of the embryo. Mathematically, this is equivalent to saying that, a priori, the position  $x$  of the cell is drawn from a distribution of possibilities,  $P_x(x)$ . Once we observe the expression level  $g$ , we still do not know the precise position  $x$  of the cell, but our uncertainty is greatly reduced. In Fig. 1, we illustrate this idea using the gap gene *hunchback* (*hb*). Expression levels of *hb* vary systematically along the anterior/posterior axis of the *Drosophila* embryo, but these expression levels also are variable across cells in the same position, both within a single embryo and across multiple embryos. Thus, if we make a “slice” through the expression profile at some particular level  $g$ , we cannot point

## Significance

**In a developing embryo, individual cells need to “know” where they are to do the right thing. How much do they know, and where is this knowledge written down? Here, we show that these questions can be made mathematically precise. In the fruit fly embryo, information about position is thought to be encoded by the concentration of particular protein molecules, and we measure this information, in bits. Just four different kinds of molecules are almost enough to specify the identity of every cell along the long axis of the embryo, and we argue that the way in which this information is distributed reflects an optimization principle, maximizing the information available from a limited number of molecules.**

Author contributions: This work is a close collaboration between theorists (G.T. and W.B.) and experimentalists (J.O.D., E.F.W., and T.G.). All authors contributed to all aspects of the work.

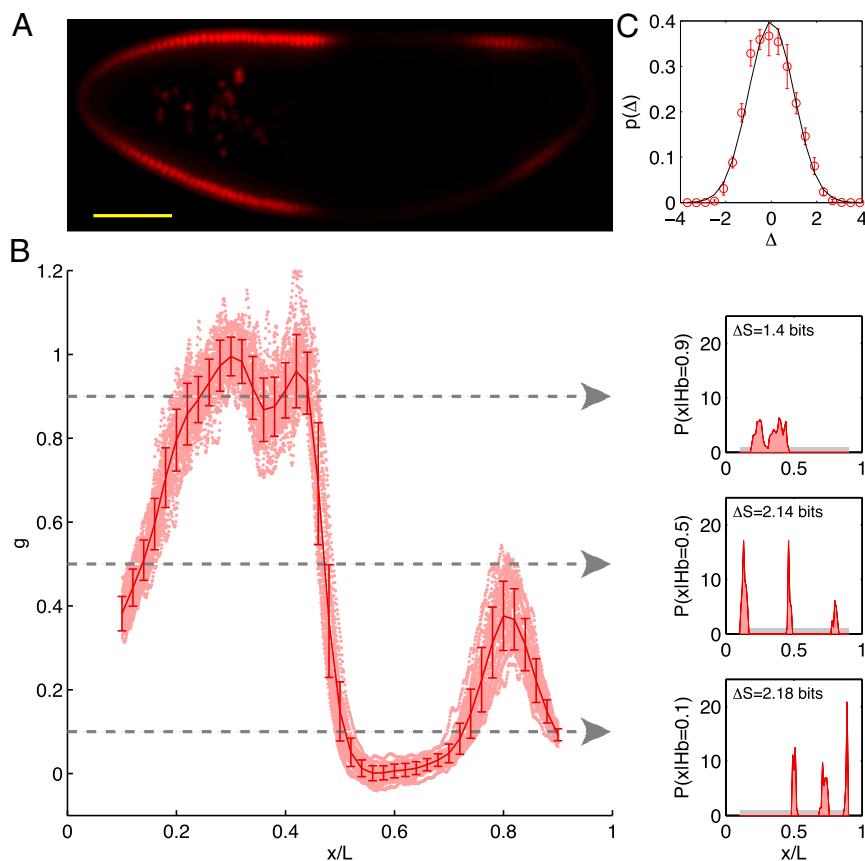
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**Fig. 1.** Positional information carried by the expression of Hb. (A) Optical section through the midsagittal plane of a *Drosophila* embryo with immunofluorescence staining against Hb protein. (Scale bar = 100  $\mu\text{m}$ .) (B) Normalized dorsal profiles of fluorescence intensity, which we identify as Hb expression level  $g$ , from 24 embryos (light red dots) selected in a 38- to 48-min time interval after the beginning of nuclear cycle 14. Position  $x$  along the anterior/posterior axis is normalized by the length  $L$  of the embryo;  $x/L=0$  corresponds to the anterior end of the embryo, and  $x/L=1$  corresponds to the posterior end. Means  $\bar{g}(x)$  and SDs  $\sigma_g(x)$  are plotted in darker red. Considering all points with  $g=0.1, 0.5$ , or  $0.9$  (Left), yields the conditional distributions with probability densities  $P(x|g)$  (Right). Note that these distributions are much more sharply concentrated than the uniform distribution  $P_x(x)$  shown in light gray; correspondingly, the entropies  $S[P(x|g)]$  are very much smaller than the entropy  $S[P_x(x)]$ . For each  $g$ , we note the reduction of uncertainty in  $x$  by reading out  $g$ ,  $\Delta S = S[P_x(x)] - S[P(x|g)]$ . (C) Variations in expression level around the mean at each position, estimated by the distribution of normalized relative expression, given by  $\Delta = [g - \bar{g}(x)]/\sigma_g(x)$  (red circles with SEMs). The solid line is a zero mean/unit variance Gaussian.

uniquely to the position  $x$  of the nucleus in which the Hb protein has that exact concentration. Instead, there is a range of positions that are consistent with the value of  $g$ , and we can summarize this range of possibilities by the conditional probability distribution,  $P(x|g)$ , that a cell with expression level  $g$  will be found at position  $x$ . For all values of  $g$  that occur in the embryo, we see that this conditional distribution is narrower or more concentrated than the nearly uniform distribution  $P_x(x)$ .

The probability distributions  $P_x(x)$  and  $P(x|g)$  provide the ingredients we need to make a mathematically precise version of the qualitative statement that “the expression level  $g$  of a gene provides information about the position  $x$  of the cell.” Crucially, the foundational result of information theory is that there is only one way of doing this that is consistent with simple and plausible requirements, for example, that independent signals give additive information (20–22).

For any probability distribution, we can define an entropy  $S$ , which is the same quantity that appears in statistical mechanics and thermodynamics; for the two distributions here,

$$S[P_x(x)] = - \int dx P_x(x) \log_2 [P_x(x)], \quad [1]$$

$$S[P(x|g)] = - \int dx P(x|g) \log_2 [P(x|g)]. \quad [2]$$

For example, if we measure  $x$  from 0 to  $L$  along the length of the embryo, then a uniform distribution of cells corresponds to  $P_x(x)=1/L$ , and this has the maximum possible entropy  $S[P_x(x)]=\log_2(L)$ . The intuition that the conditional distribution  $P(x|g)$  is narrower or more concentrated than  $P_x(x)$  is quantified by the fact that the entropy  $S[P(x|g)]$  is smaller than  $S[P_x(x)]$ , and this reduction in entropy is exactly the information that observing

$g$  provides about  $x$ , measured here in bits. As an example, if observing the expression level  $g$  tells us, with complete certainty, that the cell is located in a small region of size  $\Delta x$ , then the gain in information is  $I(g) \equiv S[P_x(x)] - S[P(x|g)] = \log_2(L/\Delta x)$  bits. Notice that entropies of continuous variables, such as position, depend on our choice of units, while the information, being the difference of entropies, is independent of this choice (22).

If we look at one cell and observe expression level  $g$ , then we gain information

$$I(g) = S[P_x(x)] - S[P(x|g)]. \quad [3]$$

However, when we choose a cell at random, we will see an expression level  $g$  drawn from the distribution  $P_g(g)$ . The average information that this expression level provides about position is then

$$I_{g \rightarrow x} = \int dg P_g(g) (S[P_x(x)] - S[P(x|g)]), \quad [4]$$

$$= \int dg \int dx P(g, x) \log_2 \left[ \frac{P(g, x)}{P_g(g) P_x(x)} \right], \quad [5]$$

where  $P(g, x)$  is the joint probability of observing a cell at  $x$  with expression level  $g$ , and we have rearranged the terms to emphasize the symmetry: Information that the expression level provides about the position of the cell is, on average, the same as the information that the position of the cell provides about the expression level,  $I_{g \rightarrow x} = I_{x \rightarrow g}$ . This average information is called the mutual information between  $g$  and  $x$ . Again, we emphasize that this measure of information is not one among many equally good possibilities; rather, it is unique (20).

Because information is mutual, we can also write  $I_{g \rightarrow x}$  in terms of the distribution of expression levels  $g$  that we find in cells at a particular position,  $P(g|x)$ ,

$$I_{g \rightarrow x} = \int dx P_x(x) (S[P_g(g)] - S[P(g|x)]). \quad [6]$$

This emphasizes that the amount of information that can be conveyed is limited both by the overall dynamic range of expression levels, which determines  $S[P_g(g)]$ , and by the variability or noise in expression levels at a fixed position, which is measured by  $S[P(g|x)]$ . It will be useful that the distribution of expression levels at each point,  $P(g|x)$ , is approximately Gaussian, as shown in Fig. 1C.

In what follows, we will use Eq. 6 to make a “direct” measurement of information, whereas Eq. 4 invites us to try and “decode” the information carried by the expression levels to recover estimates of the position  $x$  of each cell. Each approach has a natural generalization to the case where information is conveyed not by the expression level of one gene but by the combined expression levels of multiple genes  $\{g_i\}$ , and we will explore this as well. It is important to emphasize that the number of bits of information carried by the gene expression levels has meaning independent of the mechanisms by which this coding is established. Thus, at one extreme, it could be that each cell sets its expression levels independently in response to some primary morphogen [e.g., Bicoid in the *Drosophila* embryo (23–25)] whereas at the other extreme, the spatial patterns of expression could arise entirely from communication between neighboring cells, in a Turing-like mechanism (26, 27). In these different extremes, the precise value of the positional information places different quantitative constraints on the underlying mechanisms; however, in all cases, the number of available bits tells us about the reliability and complexity of the pattern that can be constructed from the local expression levels alone.

### Information Carried by Single Gap Genes

Estimating the mutual information that one gene expression level provides about position requires, from Eq. 6, that we obtain a good estimate of the conditional distribution  $P(g|x)$ . Using immunofluorescent staining, we can measure  $g$  vs.  $x$  along the anterior/posterior axis of single *Drosophila* embryos, and by making such measurements on multiple embryos, as shown in Fig. 1, we obtain many samples of the expression level at corresponding positions; from these samples, we can then build up an estimate of the distribution  $P(g|x)$ . Armed with this estimate, we can use Eq. 6 to compute the positional information. To be sure that the answer is meaningful, we have to address a number of technical issues (28).

First, as explained at the outset, we would like to measure the information carried by a snapshot of the expression levels, so we need to make measurements on embryos at a well-defined time, and we use the length of the cellularization membrane as a precisely calibrated proxy for time (29–32). We choose this time to be the window from 38 to 48 min after the start of nuclear cycle 14, because we have seen that gap gene expression levels are at a plateau in this window. We also confine our attention to the central 80% of the anterior/posterior axis, because quantitative imaging at the poles is more difficult and because there are additional genes associated specifically with terminal patterning, and we make measurements along the dorsal edge of the midsagittal plane.

Second, Fig. 1 shows that the SD of expression levels typically is less than 10% of the maximum expression level. To draw convincing quantitative conclusions, then, we must be sure that our measurements have accuracy much better than this, lest we confuse experimental error for real noise and variability in the system. As discussed by Dubuis et al. (28), the intensity of immunostaining is linear in protein concentration over the relevant dynamic range (also ref. 9), and errors can be minimized by careful attention to the orientation and age of the embryos. By comparing large numbers of embryos stained in a single batch, we find that there is little or no sign of errors due to variations in

the efficiency of staining, which means we can avoid previously troubling issues surrounding the normalization of profiles across embryos (details are provided in *Materials and Methods*). When the dust settles, our experimental or measurement errors are below  $\sim 3\%$  of the maximal expression level, and hence well below the observed noise levels (28). Note that measurement errors will always reduce the information, and so our estimate defines lower bounds on the information carried by the real biological signals.

Finally, as has been addressed in other contexts (*Materials and Methods*), care is required to be sure that the finite number of samples we collect is sufficient to get a reliable estimate of  $I_{g \rightarrow x}$ ; however, once we have control over the potential systematic errors, the statistical errors in our measurements are very small. Analysis of the data in Fig. 1 shows that the expression level of Hb provides  $I_{g_{Hb} \rightarrow x} = 2.26 \pm 0.04$  bits of information about the position of a cell along the middle 80% of the anterior/posterior axis. We can repeat this analysis for the gap genes *krüppel* (*Kr*), *giant* (*Gt*), and *knirps* (*Kni*), in addition to *Hb*, and we find  $I_{g_{Kr} \rightarrow x} = 1.95 \pm 0.07$  bits,  $I_{g_{Gt} \rightarrow x} = 1.84 \pm 0.05$  bits, and  $I_{g_{Kni} \rightarrow x} = 1.75 \pm 0.05$  bits.

In all cases, the expression of a single gene carries much more than one bit of information; indeed, it carries more nearly two bits. The conventional view of the gap genes is that they are characterized by domains of expression, with boundaries, and the sharpness of the boundary often is taken as a measure of precision. However, if the patterns of expression were perfect on/off domains with infinitely sharp boundaries, then the expression level could provide at most one bit of information about position. Our result that gap genes provide nearly two bits of information about position demonstrates that intermediate expression levels are sufficiently reproducible from embryo to embryo that they carry significant amounts of positional information, and that the view of domains and boundaries misses almost half of this information.

### How Much Information Does the Embryo Use?

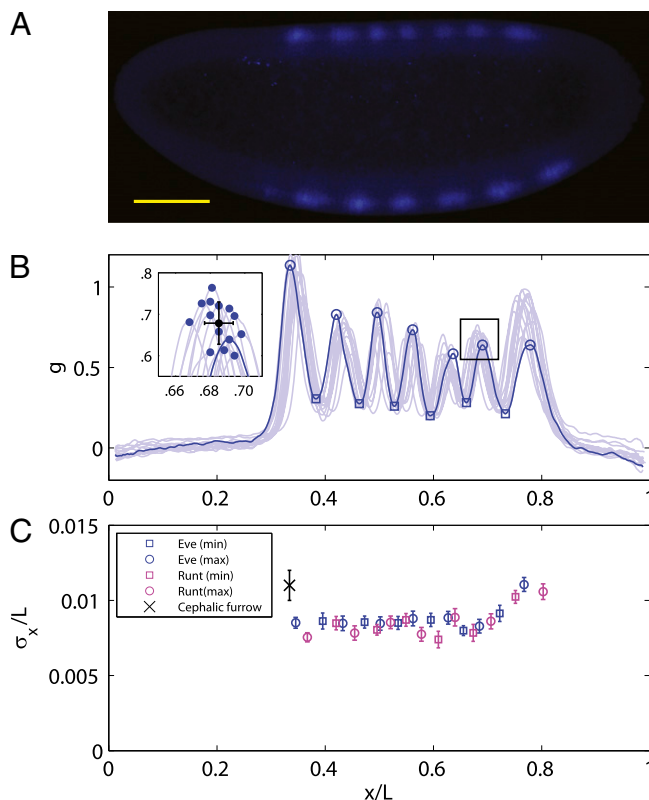
At best, every nucleus could be labeled with a unique identity, so that with  $N$  nuclei, the embryo could make use of  $\log_2 N$  bits (21). Along the anterior/posterior axis, we can count nuclei in a single midsagittal slice through the embryo, and in the middle 80% of the embryo, where the images are clearest, we have  $N = 58 \pm 4$  along the dorsal side and  $N = 59 \pm 4$  along the ventral side, where the error bars represent SDs across a population of 57 embryos in nuclear cycle 14; this corresponds to  $5.9 \pm 0.1$  bits of information. However, do individual cells, in fact, “know” their identity? More precisely, are the elements of the anterior/posterior pattern specified with single-cell resolution?

Several experiments suggest that elements of the body plan in the larval fly that emerges from the embryo can be traced to identifiable rows of cells along the anterior/posterior axis (33), which is consistent with the idea that at least some single rows of cells have a reproducible identity. Quantitatively, we can ask about the reproducibility of various pattern elements in early development, elements that appear not long after the expression patterns of the gap genes are established. A classic case is the cephalic furrow, which can be observed in live embryos and is known to have a position along the anterior/posterior axis that is reproducible with an accuracy of  $\sim 1\%$  of the embryo length (34).

Is the cephalic furrow special, or can the embryo more generally position pattern elements with  $\sim 1\%$  accuracy? The striped patterns of pair rule gene expression allow us to ask about the position of multiple pattern elements, seven peaks and six troughs of expression along the anterior/posterior axis. As shown in Fig. 2, all these elements have positions that are reproducible to within 1% of the embryo length. This strongly suggests that all cells know their position along the anterior/posterior axis with a precision  $\sigma_x/L \sim 1\%$ .

The distance between neighboring nuclei is  $\delta x/L = 0.8/N = 0.014 \pm 0.001$  of the embryo's length. If cells know their position





**Fig. 2.** Reproducibility of multiple pattern elements along the anterior/posterior axis. (A) Optical section through the midsagittal plane of a *Drosophila* embryo with immunofluorescence staining against *even-skipped* (Eve) protein. (Scale bar = 100 μm.) (B) Normalized dorsal profiles of fluorescence intensity from 12 embryos selected in a 45- to 55-min time window after the beginning of nuclear cycle 14 (light blue lines); the dorsal profile of the top panel embryo is shown in darker blue. (Inset) Zooming in on a single peak, we can measure the SD of both the expression level and position of this element in the pattern. (C) Summary of results from such measurements on Eve (blue) and Runt (magenta), plotting the SD of the position  $\sigma_x$  as a function of the mean position  $\bar{x}$ , together with a similar measurement on the reproducibility of the cephalic furrow (33). Note that all the elements are positioned with 1% accuracy or better.

with 1% accuracy, this error is smaller than the internuclear spacing, suggesting that every cell indeed has a specified position. However, this is not quite right, because errors are probabilistic and probability distributions have tails. Specifically, if the best we (or the cells) can do is to specify positions with an error that has an SD of  $\sigma_x/L = 0.01$ , and the errors come from a Gaussian distribution, then there is a probability  $P \sim 0.08$  that we will be off by  $\delta x/L = 0.014$  or more in one direction. This confusion means that the reproducibility of pattern elements in Fig. 2 provides evidence for individual nuclei having access to  $I = \log_2(0.8L/\sigma_x\sqrt{2\pi e}) = 4.27$  bits of information (22), although more may be available, as discussed below.

### Decoding the Information Carried by Multiple Genes

Do the four gap genes, taken together, carry enough information to specify position with  $\sim 1\%$  accuracy? To answer this question, we need to know not just the distribution of expression levels for single genes at each point  $x$  along the anterior/posterior axis but the joint distribution of all the expression levels. The major difficulty in such an experiment is to avoid spectral cross-talk among the different fluorescence signals, but for the experiments shown in Fig. 3, we have shown that cross-talk is  $\sim 1\%$  or less (28, 30), and, as noted in *Materials and Methods*, modest amounts of cross-talk actually do not change our estimate of  $\sigma_x$  or the

information. Given that we can sample the joint distribution of expression levels, how do we estimate the information that these expression levels carry?

We observe the expression levels  $g_i$ , with  $i = 1(\text{Hb}), 2(\text{Kr}), 3(\text{Gt}), 4(\text{Kni})$ . At each point  $x$ , there are average values of these expression levels  $\bar{g}_i(x)$ , and across an ensemble of embryos, there are fluctuations  $\delta g_i$ . Let us assume that these fluctuations have a Gaussian distribution. If we look just at one gene, this means that the statistics of the fluctuations are described completely by the mean and the variance  $\sigma_i^2(x)$ , so that if we look at the same position  $x$  in many embryos, we will see a distribution of expression levels

$$P(g_i|x) = \frac{1}{\sqrt{2\pi\sigma_i^2}} \exp\left[-\frac{(g_i - \bar{g}_i(x))^2}{2\sigma_i^2(x)}\right], \quad [7]$$

and this is in reasonable agreement with the data, as shown for the case of *Hb* in Fig. 1C (results for other genes are similar). If we look at many genes simultaneously, we have not just the variances of each gene but the correlations or covariances among the genes, which define a matrix  $C_{ij}(x)$ . The joint distribution of expression levels at one point is then

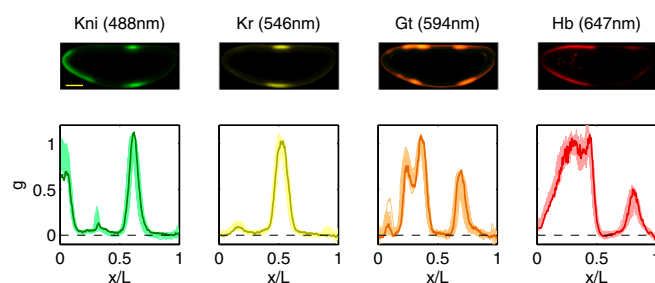
$$P(\{g_i\}|x) = \frac{1}{\sqrt{(2\pi)^4 \det C}} \exp[-F(\{g_i\})] \quad [8]$$

$$F(\{g_i\}) = \frac{1}{2} \sum_{i,j=1}^4 (g_i - \bar{g}_i(x))(C^{-1})_{ij}(g_j - \bar{g}_j(x)),$$

where  $C^{-1}$  denotes the inverse of the matrix  $C$  and  $\det C$  denotes its determinant. We can estimate all the elements of the covariance matrix, at every position  $x$ , in the usual way, averaging over samples taken from multiple embryos.

As an aside, we note that most of the significant off-diagonal elements of the covariance matrix are negative. For example, if the expression level of *Hb* happens to be a bit above average at one point in a single embryo, then the expression of *Kr* will be a bit below average at that same point. Presumably, this reflects the mutually repressive interactions among the gap genes (35–37).

The distribution  $P(\{g_i\}|x)$  characterizes the measurements that we can make as an outside observer of the embryo. However, a single nucleus does not have access to the position  $x$ ; rather, the whole idea of positional information is that this position is encoded in the expression levels. To assess the quality of this code, we can try to read it, asking for the distribution



**Fig. 3.** Simultaneous measurements on four gap genes reproduced, in part, from the work of Dubuis et al. (28). (Upper) Optical sections through the midsagittal plane of a single *Drosophila* embryo with immunofluorescence staining against Kni (green), Kr (yellow), Gt (orange), and Hb (red), with fluorescence excitation wavelengths in parentheses. (Scale bar = 100 μm.) (Lower) Normalized expression levels along the dorsal edge for 24 embryos in a 38- to 48-min time interval after the start of nuclear cycle 14 (light colors); the sample embryo is highlighted (dark colors).

of positions that are consistent with a particular set of expression levels that we might observe. By Bayes' rule, this can be written as

$$P(x|\{g_i\}) = \frac{P(\{g_i\}|x)P_x(x)}{P_g(\{g_i\})}, \quad [9]$$

where  $P_x(x)$  is, as before, the (nearly uniform) distribution of cell positions and  $P_g(\{g_i\})$  is the (joint) distribution of expression levels averaged over all cells in the embryo.

If the noise levels are small, then  $P(x|\{g_i\})$  will be sharply peaked at some  $x_*(\{g_i\})$ , which is the best estimate of the position, given the expression levels. Expanding around this estimate, the distribution is approximately Gaussian,

$$P(x|\{g_i\}) \approx \frac{1}{\sqrt{2\pi\sigma_x^2}} \exp\left[-\frac{(x-x_*(\{g_i\}))^2}{2\sigma_x^2}\right], \quad [10]$$

where the error in our position estimate is defined by

$$\frac{1}{\sigma_x^2} = \sum_{i,j=1}^4 \left[ \frac{d\bar{g}_i(x)}{dx} (C^{-1})_{ij} \frac{d\bar{g}_j(x)}{dx} \right] \bigg|_{x=x_*(\{g_k\})}. \quad [11]$$

All the terms in Eq. 11 are experimentally accessible.

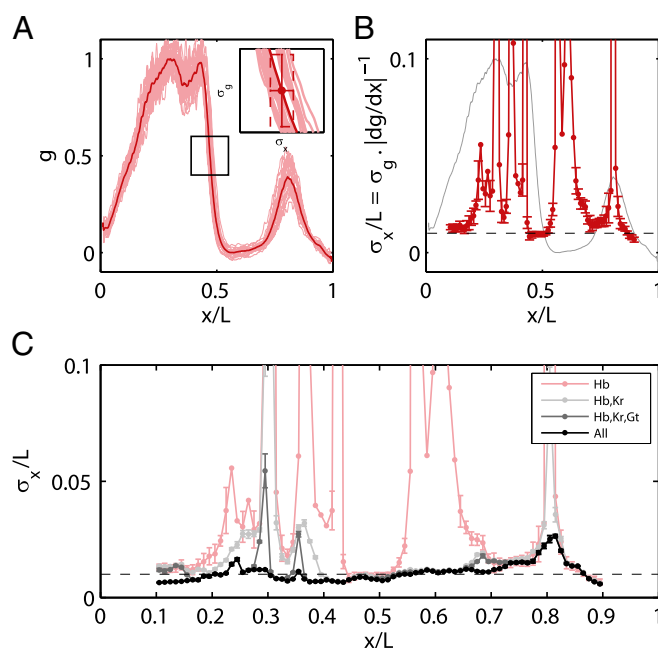
Eq. 11 tells us the precision with which expression levels encode position: Observing the expression levels  $\{g_i\}$  allows us (or the cell) to specify position, at best, with an "error bar"  $\sigma_x$ ; this error could be different at different points in the embryo, so we really should write  $\sigma_x(x)$ . Checking our intuition, we see that this error bar is smaller when the variability in expression is smaller (smaller  $C$ ), when the mean slopes of the expression levels are larger (larger  $d\bar{g}_i/dx$ ), or when we can sum over more genes. We can define a similar quantity based on measurements of a single gene,

$$\frac{1}{\sigma_x(x)} = \left| \frac{d\bar{g}_i(x)}{dx} \right| \frac{1}{\sigma_i(x)}, \quad [12]$$

and this construction is shown schematically in Fig. 4A and B in the case of *Hb*. Note that when  $\sigma_x$  is small, we can justify our approximation that  $P(x|\{g_i\})$  is sharply peaked, but when  $\sigma_x$  becomes large, it is more rigorous simply to say that we do not have much information about  $x$  rather than trying to give a more quantitative interpretation.

Analyzing the spatial profiles and variability of gene expression as suggested by Eq. 11, we obtain the estimates of  $\sigma_x$  shown in Fig. 4C. Remarkably, the reliability of position estimates based on the four gap genes is  $\sigma_x/L \sim 1\%$  (compare with dashed line), almost precisely equal to the observed reproducibility with which pattern elements are positioned along the anterior/posterior axis. This is strong evidence that the gap genes, taken together, carry the information needed to specify the full pattern. Further, this positional accuracy is almost constant along the length of the embryo, which again is consistent with what we see in Fig. 2. This constancy emerges in a nontrivial way from the expression profiles, the noise levels, and the correlation structure of the noise. If we try to make estimates based on one gene, we can reach  $\sim 1\%$  accuracy only in a very limited region of the embryo, but the detailed structure of the spatial profiles ensures that these signals can be combined to give nearly constant accuracy.

If the errors in estimating position really are Gaussian, as in Eq. 10, then we can substitute into Eq. 4 to show that  $I = \langle \log_2[0.8L/(\sigma_x\sqrt{2\pi e})] \rangle$ , where  $L$  is the length of the embryo, and  $\langle \dots \rangle$  denotes an average over position. Computing this average, we have  $I = 4.14 \pm 0.05$  bits. Alternatively, we can use the distribution of expression levels at each position, Eq. 8, and we find



**Fig. 4.** Positional error as a function of position. (A) Geometrical interpretation of the positional error for a single gene (*Hb*) at a given position. From Eq. 12,  $\sigma_x(x)$  is proportional to the reproducibility of the profiles and is inversely proportional to the derivative of the mean profile. (B) Positional error based on the expression of *Hb* alone (red; mean  $\pm$  SEM from bootstrapping) compared with the mean profile (gray). (C) Positional error based on combinations of gap genes, from Eq. 11. Note that once we combine information from all the gap genes, the net positional error is nearly constant and equal to 1% along the entire anterior/posterior axis.

$I = 4.1 \pm 0.23$  bits. The agreement between these estimates supports our approximations and gives us confidence that the measurement of  $\sigma_x$  in Fig. 4 really does characterize the encoding of positional information by the gap genes.

Thus, the gap genes carry enough information for each nucleus to know its position with an error bar  $\sim 1\%$  of the embryo's length, and this is equal to the variability in localization of features that emerge in later stages of development. On the other hand, as noted above, this is not quite enough to specify the position of every nucleus uniquely. Is it possible that more information is "hiding" in the expression profiles? In particular, if the noise in neighboring cells is correlated, the errors in specifying relative positions (e.g., that one cell is more posterior than another) could be much smaller than the errors in specifying absolute positions. As a first step, we can ask how much information the expression levels of the gap genes provide about position measured from a "center of mass" that we compute from the whole spatial profile, rather than position in the fixed coordinate system that starts with  $x=0$  at the anterior end of the embryo. This relative positional information is 0.7 bits larger than the absolute positional information; although the data are very preliminary, we see hints of a similar gain of information about relative position for the peaks of *Eve* expression in Fig. 2. These results indicate that, through spatial comparisons, there may be enough information available to specify each cell's identity.

### More Than One Bit per Gene?

The positional information carried by single gap genes is more nearly two bits than one, as described above, suggesting that spatial variations in gene expression define much more than on/off expression domains. However, when we combine information from different genes, redundancy among the spatial profiles of the different genes limits the information gain, with the result





fraction of this capacity is achieved by the embryo? The result, from ref. 15, is that

$$I_{\max} = \log_2 \left[ \frac{1}{\sqrt{2\pi e}} \int_0^L \frac{dx}{\sigma_x(x)} \right]. \quad [17]$$

The observed information transmission is  $I/I_{\max} = 0.984 \pm 0.003$ , within a few percent of the optimum.

## Discussion

The final result of embryonic development appears precise and reproducible. Less is known about the degree of this precision, and about the time at which precision first becomes apparent. Our central result is that in the early *Drosophila* embryo, the patterns of gap gene expression provide enough information to specify the positions of individual cells with a precision of  $\sim 1\%$  along the anterior/posterior axis. This is the same precision with which subsequent pattern elements are specified, from the pair rule expression stripes through the cephalic furrow, so that all the required information is available from a local, instantaneous readout of the gap genes.

The precise value of the information that we observe is also interesting. It corresponds to being able to locate any nucleus with an error bar that is smaller than the distance to its neighbor, but the total number of bits is not quite large enough to specify the position of every cell uniquely. The difference is that when we make an estimate with error bars, the estimate comes from a distribution with tails, and the (small) overlap of the tails of these distributions means that one cannot quite identify every cell. It is possible that cells, in fact, do not quite have unique identities or that these identities emerge only later in development. Alternatively, although the gap genes encode position with an error bar, the difference between positions coded by expression levels in neighboring cells could have a much smaller error bar, and we have preliminary evidence for this idea. Although further experiments are required to settle this issue, we find it remarkable that the gap gene expression levels carry so much information, such that an enormously precise pattern is available very early in development.

The fact that precision is available early does not mean that there is no enhancement of precision by subsequent processes. In particular, because the joint distribution of expression levels does not fill the full space of possibilities, it would be possible for the embryo to recognize a large error, and perhaps to correct it, with no additional inputs. The question of whether the embryo achieves such an error-correcting code (21) for positional information is completely open.

The information that gene expression levels can carry about position is limited by noise. In particular, both because the concentrations of transcription factors are low and because the absolute copy numbers of the output proteins are small, there are physical sources of noise that cannot be reduced without the embryo investing more resources in making these molecules. Given these limits, it still is possible to transmit more information through the gap gene network by “matching” the distribution of input signals to the noise characteristics of the network. Although this matching condition is generally complicated, in the limit that the noise is small, it can be expressed very simply: The density of cells along the anterior/posterior axis should be inversely proportional to the precision with which we can infer position by decoding the signals carried in the gap gene expression levels. Because cells are almost uniformly distributed at this stage of development, this predicts that an optimal network would have a uniform precision, and this is what we find. This uniformity emerges despite the complex spatial dependence of all the ingredients, and thus seems likely to be a signature of selection for optimal information transmission.

## Materials and Methods

**Experiments.** To allow simultaneous imaging of proteins encoded by all four gap genes, polyclonal antibodies were generated (Panigen, Inc., Blanchardville,

WI) in mice, rats, and guinea pigs against His-Trx-tagged full length Hb, Kni, and Gt fusion proteins (42); procedures were under the approval of Princeton University’s Institutional Animal Care and Use Committee, Protocol No. 1798A to E.F.W. To image Kr protein, we use a rabbit anti-Kr antibody generated by Chris Rushlow (New York University). Fixation and staining were done as described by Dubuis et al. (28); details of the imaging, profile extraction, and staging (embryo age determination during nuclear cycle 14) are described by Dubuis et al. (28). We draw attention to the discussion of experimental errors in the study of Dubuis et al. (28), because this issue is especially important for our analysis.

**Analysis.** Measurements on the expression profiles of a single gene in multiple embryos provide many samples of the joint distribution  $P(g, x)$ . To compute the mutual information between  $g$  and  $x$ , we discretize the two continuous axes into a number of bins; along the  $g$  axis, we use these bins adaptively so that the histogram of  $g$  in these bins is nearly flat. We then take the (normalized) counts in each bin as an estimate of the probability, compute the information, and examine the dependence on the number of bins and the number of samples. Following refs. 43 and 44, we search for the expected systematic dependencies and extrapolate to the limit where the number of bins and samples both become large. We can obtain an upper bound on the information by assuming that the conditional distribution  $P(g|x)$  is Gaussian, and we can obtain an approximation to the information by taking this Gaussian approximation through to the construction of  $P_g(g)$ ; all these estimates agree within error bars. With simultaneous measurements of expression levels for multiple genes, we can estimate the information that they carry jointly. The difficulty is that the space of expression levels is now much larger but our number of samples is not. Having calibrated the Gaussian approximation against more direct calculations for single genes (above), we can use this approximation in the case of multiple genes, using Eq. 8 directly in the multidimensional generalization of Eq. 5; we use a Monte Carlo method to evaluate these integrals numerically and estimate errors by a bootstrap method. Means and covariance matrices are calculated from our multiple samples of joint expression levels in the usual way. Importantly, if the signals that we observe are invertible linear combinations of the true signals, as might happen, for example, because of a small amount of cross-talk among the different imaging channels, then the invariance of the information to coordinate transformations tells us that this will not change our estimate. The other path to the analysis of multiple genes is through the computation of  $\sigma_x$ , as described in the discussion leading to Eq. 11. Here, too, we have to be careful about the dependence of our estimates on the number of samples that we include in our analysis, and quoted results are extrapolated as by Strong et al. (43) and Slonim et al. (44). In the discussion leading to Fig. 5, we set thresholds to quantize the expression levels and then estimate the mutual information between the four-bit words and the position  $x$ ; the results we show are for the settings of the four thresholds that maximize the information.

**Derivation of Optimality Condition.** To derive Eq. 13, consider the case where information flows from a single input transcription factor (e.g., Bicoid) to a set of  $K$  output genes (the gap genes). The concentration of the input is  $c$ , and the output genes have expression levels  $g_1, g_2, \dots, g_K$  (16–18). Different cells in the embryo experience different values of  $c$ , depending on their position, and if we choose a cell at random, it sees a concentration drawn from the distribution  $P_c(c)$ . The network responds to this input, generating expression levels that are drawn from the distribution  $P(\{g_i\}|c)$ ; it will also be useful to define the (joint) distribution of output expression levels,

$$P_g(\{g_i\}) = \int dc P_c(c) P(\{g_i\}|c). \quad [18]$$

The information that flows from input to output can be written, as in Eq. 4, as

$$I(\{g_i\}; c) = - \int dc P_c(c) \log_2 P_c(c) - \int d^K g P_g(\{g_i\}) S[P(c|\{g_i\})], \quad [19]$$

where, from Bayes’ rule, we have

$$P(c|\{g_i\}) = \frac{P(\{g_i\}|c) P_c(c)}{P_g(\{g_i\})}. \quad [20]$$

The transmitted information  $I(\{g_i\}; c)$  depends both on the characteristics of the gene network, expressed as  $P(\{g_i\}|c)$ , and on the distribution of input signals,  $P_c(c)$ . In particular, noise associated with the finite number of available molecules is encoded by the details of  $P(\{g_i\}|c)$ . Given these constraints, it still is possible to maximize information transmission by the proper choice of the input distribution (20, 21). In general, this optimization

is a hard problem, but we can make progress if we assume that the noise is small, and we will argue that this is a good approximation.

In Eq. 19, we need to take an average over the full distribution of output expression levels,  $P_g(\{g_i\})$ . This distribution is broadened by two effects. First, the inputs  $c$  are varying, and the outputs vary in response. Second, even when the input  $c$  is fixed, the outputs  $\{g_i\}$  vary because of noise. We assume that noise is small in the sense that the first effect is much larger than the second, so that we can average over outputs by assuming that the output is always equal to its average value,  $g_i = \bar{g}_i(c)$ , and then average over the input  $c$ . In this approximation,

$$I = - \int dc P_c(c) \log_2 P_c(c) - \int dc P_c(c) S_{\text{cond}}^{(c)}(\{g_i = \bar{g}_i(c)\}), \quad [21]$$

where  $S_{\text{cond}}^{(c)}(\{g_i\}) = S(P(c|\{g_i\}))$ . To find the distribution of inputs that maximizes the information, we introduce, as usual, a Lagrange multiplier to fix the normalization of  $P_c(c)$  and solve

$$\frac{\delta}{\delta P_c(c)} \left[ I - \lambda \int dc P_c(c) \right] = 0. \quad [22]$$

The result is

$$P_c(c) = \frac{1}{Z} \exp \left[ -(\ln 2) S_{\text{cond}}^{(c)}(\{g_i = \bar{g}_i(c)\}) \right], \quad [23]$$

where  $Z$  is chosen to normalize the distribution. If the noise is also approximately Gaussian—given knowledge of the gene expression levels  $\{g_i\}$ , we know the input concentration to within some error bar  $\sigma_c^{\text{eff}}(c)$ , which itself depends on the actual value of the input—then  $S_{\text{cond}}^{(c)} = \log_2[\sqrt{2\pi}\sigma_c^{\text{eff}}(c)]$  and

$$P_c(c) = \frac{1}{Z} \cdot \frac{1}{\sigma_c^{\text{eff}}(c)}, \quad [24]$$

corresponding to Eq. 13. The system can optimize information transmission by using the symbols  $c$  in proportion to their reliability (15).

The noise in the system can be summarized by  $\sigma_x$  itself, which is smaller than the distances over which the output of any single gap gene varies significantly. Thus, in retrospect, the effective noise really is small, as assumed above, which justifies the approximation leading to Eq. 23. This derivation can be generalized to cases where there are multiple independent morphogen inputs, each varying along  $x$ .

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