vestigate longer-term trends in $[CO_3^{2-}]$ and PCO_2 . The observations made here also provide potential to refine the estimates of deepocean $[CO_3^{2-}]$ based on the assumption that foraminiferal shell weight is an index of dissolution (12). Such estimates clearly require prior knowledge of initial weight, which might be attained from a nearby shallow-core calibration or possibly by using an atmospheric PCO_2 record. It is interesting to note that the shell weights of undissolved foraminiferal calcite appear to trace surface-ocean $[CO_3^{2-}]$, whereas those of dissolved foraminiferal calcite appear to trace $[CO_3^{2-}]$ of the deep ocean.

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- 21. The CO₂ record used here is plotted on the time scale calculated by tuning Vostok δ^{18} O to the June insolation curve at 65°N (29). Using this time scale allows for better correlation between ice-cores and marine records.
- Empirical values of K_{0,1,2} in terms of T and S were taken from Millero (30). Carbonate alkalinity, CA,

 $([HCO_3^{-}]+2[CO_3^{2-}])$ is considered equal to 96% of TA, allowing for the contributions from borate and water (plus minor others) alkalinity. The value of 96% is taken from Zeebe and Wolf-Gladrow (31) as a typical seawater example (TA \approx PA = 2300 μ mol kg⁻¹, T = 25°C, S = 35).

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Supporting Online Material

www.sciencemag.org/cgi/content/full/297/5582/833/

DC1 Methods Fig. S1

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12 April 2002; accepted 1 July 2002

Single-Cell Gene Expression Profiling

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A key goal of biology is to relate the expression of specific genes to a particular cellular phenotype. However, current assays for gene expression destroy the structural context. By combining advances in computational fluorescence microscopy with multiplex probe design, we devised technology in which the expression of many genes can be visualized simultaneously inside single cells with high spatial and temporal resolution. Analysis of 11 genes in serum-stimulated cultured cells revealed unique patterns of gene expression within individual cells. Using the nucleus as the substrate for parallel gene analysis, we provide a platform for the fusion of genomics and cell biology: "cellular genomics."

The first step in the translation of genomic sequence into physiology or pathophysiology is transcription. Transcriptional regulation has been studied almost exclusively on nucleic acids extracted from cultured cells or tissues by Northern blot (1), differential display (2), serial analysis of gene expression (SAGE) (3), or forms of microarray (4, 5). Here, we describe a complementary approach that monitors mRNA synthesis by visualizing specific sites of transcription (6, 7). The use of transcription sites for expression profiling allows analysis of coordinated transcription events and organization of gene expression. To achieve sensitive and specific detection of RNAs by fluorescence in situ hybridization (FISH), we used oligomer DNA probes that were each tagged with a single fluorophore at multiple sites (8). To detect many mR-NAs simultaneously, we used combinations of these probes labeled with spectrally distinct colors (Fig. 1A) (9). A combinatorial approach to labeling probes (Fig. 1B) provided a large number of virtual "colors" for distinguishing many transcripts (supporting online text and table S1). Spectral "barcodes" with a minimum of two distinct fluorophores were used to increase specificity. Schemes based on color combinations have been applied to detection of entire chromosomes for cytogenetics (10, 11) and for analysis of subchromosomal regions of DNA (12). However, the targets in these genomic assays are orders of magnitude larger than transcripts and do not contain functional information.

Cell culture and preparation, fluorescence microscopy, and image acquisition were performed and images were subjected to computational analysis to detect sites of transcription (9). Transcription sites have a

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discernible three-dimensional volume and shape, as well as the highest mRNA signal due to the presence of multiple nascent transcripts at the location of the transcription unit in chromatin (Fig. 1B). The color combination, or spectral barcode, defined transcript identity (Fig. 1C). Colocalization of multiple colors in the image represented independent hybridization events that signified the presence of a transcription site. Signals were analyzed for individual cells on the basis of contiguity of the 4',6'diamidino-2-phenylindole (DAPI) counterstain (9). We identified 10 sites of transcription simultaneously in starved and serum-stimulated human colon adenocarcinoma (DLD-1) cells (13) (Fig. 2, A to C). Even accessibility to probe was assured by presence of transcription sites in more than 97% of cells (176/181). Sites of the most prevalent gene, y-actin, occurred in 80% of nuclei (144/181). Before the mix of probes for 10 genes was made, each gene was first hybridized individually to ensure that measured expression was independent of multiplexing (14).

Assaying expression by transcription site visualization preserves population heterogeneity. For the DLD-1 cell line, the number of transcription sites per cell was not uniform throughout the population. There was a strong correlation with nuclear cross-sectional area and total DNA signal, measured as fluorescence intensity of DAPI counterstain (Fig. 3A) (r^2 value, 0.56). One additional transcription site was observed for each 11.9 μ m² of nuclear area [95% confidence interval (CI), 10.5 to 13.7 µm²]. In contrast, starved and stimulated normal fibroblasts did not show a correlation of sites and size. Analysis of 162 fibroblast nuclei fit a linear correlative model quite poorly (r^2 value, 0.0025). Starvation of the cancer cells did not completely abolish increased numbers of transcription sites associated with increased DNA content, as it did in normal fibroblasts. Although levels of transcription were far reduced in starved DLD-1 and cell cycle state was grossly synchronized, differences among cells were detectable in situ. The ability to assay this heterogeneity is a clear advantage in the evaluation of mixed tissue samples, such as is common in clinical pathology.

Population profiles were determined for cells that were previously subjected to microarray and other molecular analyses (15, 16). This provided a direct comparison of data based on extracted RNA with single-cell expression levels, a process we termed "FISH & Chips." Thirty min after serum stimulation, the average DLD-1 nucleus had detectable expression of five individual genes and roughly six to seven transcription sites, including both alleles of some genes (Fig. 3B). Genes with frequently occurring sites often had two active alleles, with an approximate 1:1 ratio of two-allele to oneallele nuclei. Genes with lower levels usually had only a single allele expressed (Fig. 3C). This showed that variable expression levels can be visualized in situ and scored for intensity, number of alleles, or dosage. Possible applications include monitoring allele silencing and determining the ploidy of heterogeneous samples.

The ability to collect a binary "snapshot" of genes that are "on" or "off" at a single moment in time for many nuclei offered new insights into transcript regulation. Of the 10 mRNAs analyzed, several pairs of genes were coexpressed with higher probability than would be predicted by random association (table S2). Relative to β -actin-negative cells, nuclei positive for β -actin expression were 5.2 times as likely to express γ -actin (99% CI, 1.6 to 16.7) and 3.2 times as likely to express EGR-1 (99% CI, 1.3 to 7.7). Remarkably, all three of these assayed genes contain one or more sequences to which serum response factor binds (17–20). This implies that genes with similar promoter elements exhibit correlated activation at the single-cell level. Such activity could result from nonhomogeneous distribution of a transcription factor in creased accessibility of similarly regulated



Fig. 1. Conceptual diagram of barcoding. (A) Schematic representation of the use of color groups to encode unique gene identities. Combinations shown are only two colors chosen from a total of four used throughout our experiments. A minimum of two colors is used to reduce the chance of single-color false positive signals. Each detected site is therefore representative of two or more independent hybridization events that are spatially colocalized, thus increasing fidelity. (B) Schema of the actual probe hybridizations at the transcription site leading to detection by barcode. The gene locus is represented (green) with several polymerases (blue) transcribing nascent messages (pink). A shotgun approach is used such that each specific probe may be of any of the colors in the barcode (red, yellow, magenta). The transcription site is shown below with hybridization data against the nuclear background (blue). Scale bar, 3 μ m. (C) An example of the signal readout interpreted by the transcription detection algorithm, showing the pixel intensities for the area of the transcription sites in all five color bandwidths. In this example, there is signal for each of the probe components used (red, yellow, and magenta) and only background levels for the color not in the barcode (purple).

genes to their activating factors. Except for the genes just described, cell expression profiles were variable, without a consistent pattern of inclusion and exclusion. If early serum-activated transcription resulted from a distinctly ordered pathway, subsets of cells would have been in different stages of the

Fig. 2. Simultaneous detection of many genes. (A) A single human colon adenocarcinoma cell (DLD-1) G2 nucleus (by DAPI signal) with a pseudo-colored representation of 17 transcription sites detected in situ. The image is "flattened" such that all 12 0.5- μm Z-sections are displayed on the background, which is the DAPI counterstain from the middle image of the stack. Gene identity is denoted by color and the Z-location is recorded by the adjoining number. Lower numbers represent closer proximity to the cover slip. Scale bar, 3 μ m. (B) Three G₁ DLD-1 nuclei from the same field, which together express all 10 genes assayed. Arrows indicate sites that are shown below magnified from the original data. From left to right, the 10 marked transcription sites are IL-8, MCL-1, DUSP-1, cyclin D1, γ-actin, EGR-1, TIEG-1, β -actin, c-myc, and c-jun. Scale bar, 3 μ m. (C) Chart of the 10 genes detected in (B). The "Pseudo" column shows the arbitrary pseudo-color used to denote the gene identity of each transcription site in the renderings above. Columns at right show the actual signal recorded at the appropriate

response. This was not the case. Pairwise analyses of transcription sites showed no distinct mutual exclusions or anticorrelations (the 99% CI for the odds ratio of each of the 45 measured gene pairs exceeded 1). Hierarchical log-linear models showed no higher order associations; cells were positive for different sets of transcripts. At the single-cell level, physiological and random variation, possibly based on the level or activity of a transcription factor, may cause the seruminduced genes to be expressed at different times.

To assess timed expression activation,



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Z-section for the transcription sites shown with an arrow in (B). Each band of data of fluorescein isothiocyanate (FITC), Cy3, Cy3.5, and Cy5 is shown, with the positively scored signals highlighted by surrounding boxes. Each area of the unprocessed image shown is $1 \,\mu m^2$. Observed misalignment is due to chromatic shift between filter sets.











Number of alleles / genes

REPORTS

we analyzed the transcription of 11 serumresponsive genes over 14 time points in normal human fibroblasts (21). We profiled 2199 nuclei that contained 10,134 transcription sites. For each time point, 55 gene-gene comparisons were made to explore the possibility of ordering of transcription timing (a total of 770 pairwise comparisons). As in the DLD-1 experiments, there were no anticorrelated or mutually excluding gene pairs, indicating that a linear pathway of gene induction is highly unlikely among the early response genes.

All the induced genes showed similar transcription site activation and deactivation kinetics over a 90-min period (Fig. 4). Several transcripts we visualized easily showed no changes by microarray [less than a factor of 2.2, the detection limit of the technology (16)]. These include β -actin (Fig. 4A), γ -actin, and c-junall shown to be serum-responsive genes (22). Incremental changes may not be detectable using total mRNA. However, with transcription site analysis, small changes in expression could be observed, irrespective of the total abundance of the RNA. Therefore, microarray and in situ assays yield complementary information. A metaphor for the difference is that transcription site profiling measures the transcriptional "thermostat" and microarrays view the "ambient temperature" of whole-cell RNA levels.

All genes showed measurable activation in the population of cells at 5 or 10 min after induction, relative to starved, unstimulated cells (Fig. 4B). Over the first 10 min of induction, one allele was activated. The average cell assayed had 3.2 different genes activated with a total of only 4.0 transcription sites; single allele expression outnumbered the expression of both alleles by a ratio of 3 to 1. By 40 min of induction, the distribution was even: Cells averaged 3.4 genes with a single allele and 2.8 with both alleles active, a total of 9.0 sites per nucleus. From 60 min to 90 min, the number of sites dropped to 2.9. By 120 min, nearly all sites had returned to baseline prevalence (Fig. 4C). This time course fit well with the previous results from a single time point in DLD-1 cells. At 30 min of stimulation, DLD-1 cells showed an average of 6.5 genes and fibroblasts showed 6.1 genes. This implies that serum-responsive transcription follows similar kinetics in different cell types. Differences were also noted, as DLD-1 cells expressed considerably more β -actin (1.2 sites per cell versus 0.33) and somewhat more γ -actin (0.69 sites per cell versus 0.45) than fibroblasts.

Lags were apparent between the kinetics of transcription site activations and their effect on cellular levels of transcripts as assessed by microarray (14). This is a function of transcription rate, message stability, and abundance level at the start of induction. Previous analyses measured a combination of these factors, whereas FISH detects only new transcription. Levels of cyclin D1 transcripts reportedly increase later than the early immediate response genes, and it was thus classified as a delayed immediate response gene (23). In our studies, cyclin D1 was just as early as other genes with maximal activation at 50 min after stimulation, as compared to the 11gene average of 46 min. C-myc showed fairly low levels of induction on microarray (16), although we have observed frequent c-myc sites (14). Levels of cmyc are therefore likely to be controlled posttranscriptionally.

These studies demonstrate the power of cellular transcriptional profiling. The instantaneous transcriptional activity of genes in single cells allows observation of causes and effects of expression. Eventually, the physiological state of cells within tissues will become synonymous with a pattern of gene expression. This will provide a quantitative approach to factors influencing gene expression patterns, such as occur in cytopathology, development and cell differentiation, infectious disease, or response to drug treatment. Investigation of functional genomics may now be approached at the cellular level. We expect that the enormous information inherent to the expression of many genes in large cell populations will aid the understanding of relationships among genes in single nuclei and their cooperative and cumulative roles in physiology and disease.



Fig. 4. Transcriptional response of fibroblasts to serum. (A) The kinetics of activation of transcription for the 11 genes assayed was relatively similar, in contrast with results obtained through microarray analysis. Here, two serum-responsive genes, β -actin and c-fos, are shown to be similar, yet only c-fos activation was detected on microarray (16). (B) Increases in visualized transcription sites 10 min after serum induction, relative to starved controls. (C) Activation curve for all assayed genes over a time course of serum induction, showing how allele and gene prevalence change as the cascade progresses. In (A) and (C), extrapolated time points are shown in brackets.



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NM_021960), transforming growth factor β immediate early gene (TIEC, NM_005655), and dual specificity kinase 1/MAP kinase phosphatase (DUSP, NM_004417). These genes were first detected singly, using Cy3 and Cy5 probes. Then they were barcoded to distinguishable combinations of fluorophores for multiplexed detection (coding scheme in Fig. 2C). This allows comparison of sensitivity of detection with different color codes to ensure reliability independent of color code. Full sequences of probes are provided in supporting online material.

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White Collar–1, a DNA Binding Transcription Factor and a Light Sensor

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Blue light regulates many physiological processes in fungi, but their photoreceptors are not known. In *Neurospora crassa*, all light responses depend on the Per-Arnt-Sim (PAS) domain–containing transcription factor *white collar–1* (*wc-1*). By removing the WC-1 light, oxygen, or voltage domain, a specialized PAS domain that binds flavin mononucleotide in plant phototropins, we show that light responses are abolished, including light entrainment of the circadian clock. However, the WC-1–mediated dark activation of *frq* remains normal in this mutant, and the circadian clock can be entrained by temperature. Furthermore, we demonstrate that the purified *Neurospora* WC-1–WC-2 protein complex is associated with stoichiometric amounts of the chromophore flavin-adenine dinucleotide. Together, these observations suggest that WC-1 is the blue-light photoreceptor for the circadian clock and other light responses in *Neurospora*.

Blue and near-ultraviolet light are known to regulate many physiological processes in a large number of organisms, including circadian clock functions from fungi to mammals. Currently, phototropins, cryptochromes, and a flavin-adenine dinucleotide (FAD)-containing adenylyl cyclase in *Euglena* are the three known types of eukaryotic blue-light photoreceptors, all of which are flavin-containing proteins (1-3). Most fungal photoresponses, ranging from growth responses to phototropism to carotenoid induction to circadian clock entrainment, are mediated by blue light (4). For *Neurospora crassa*, previous evidence has suggested that a flavincontaining blue-light photoreceptor is responsible for mediating all known light responses [reviewed in (4)].

The white collar-1 (wc-1) and wc-2 genes are required for all aspects of the known Neurospora blue-light responses, including induction of the circadian clock gene frequency (frq), which has been shown to mediate the light entrainment of the circadian clock (4-10). WC-1 and WC-2 are transcriptional activators that can bind to promoter elements upstream of light-inducible genes via their GATA-type zinc-finger DNA binding domains (5, 6, 11). Both proteins also contain Per-Arnt-Sim (PAS) domains, which are important for the in vivo formation of a nuclear complex between WC-1 and WC-2 1 (Fra-1, NM_005438) in the place of IL-8 and TIEG. Full sequences of probes are provided in supporting online material. Experiments were performed for logarithmically growing cells, starved (0 min), and 5, 10, 15, 20, 25, 35, 40, 45, 50, 60, 90, and 120 min after induction. Data were extrapolated for the 30-min time point and points between 60, 90, and 120 min.

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Supporting Online Material

www.sciencemag.org/cgi/content/full/297/5582/836/ DC1

Materials and Methods SOM Text References and Notes

Tables S1 and S2

26 March 2002; accepted 14 June 2002

(WCC) (9, 12). In addition to their critical roles in light responses, WC-1 and WC-2 are also essential components of the *Neurospora* frq-wc-based circadian feedback loops because they activate the transcription of frq in the dark (7, 13-16).

Sequence analysis of the WC-1 protein revealed that one of its three PAS domains (Fig. 1A) belongs to a specialized class of these domains known as a light, oxygen, or voltage (LOV) domain. These LOV domains have been best characterized in plant phototropins, in which each LOV domain binds a flavin mononucleotide (FMN) molecule and is able to undergo fully reversible photocycles (1, 17–19). The crystal structure of a LOV-FMN complex revealed 11 residues in the vicinity of FMN that are highly conserved in LOV domains (20), including that of WC-1, suggesting that WC-1 may bind FMN and function as a blue-light photoreceptor.

We made a construct in which the LOV domain was deleted from the wild-type wc-1 gene (Fig. 1A) and introduced it into a wc-1 null strain (wc-1.lov) (21). The apparent molecular weight of mutant WC-1 was smaller than that of wild-type WC-1, and mutant WC-1 was present in a lower amount than wild-type WC-1 (Fig. 1B). The amount of FRQ in the wc-1.lov mutant was near normal in 22 hours of constant darkness (DD22) but did not increase in cultures grown in constant light (LL), indicating that the light induction of frq is lost in the mutant. This result was confirmed with Northern blot analyses of frq and al-3 (a gene required for carotenoid biosynthesis) RNA amounts (Fig. 1C). As for FRQ in the dark, the amounts of frq mRNA were comparable in both strains (Fig. 1C, middle panel). These data indicate that the WC-1 LOV domain is essential for light-activated transcription of frq but not frq transcription activation in the dark.

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