Allelic inactivation of rDNA loci

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Human cells contain several hundred ribosomal genes (rDNA) that are clustered into nucleolar organizer regions (NORs) on the short arms of five different acrocentric chromosomes. Only ~50% of the gene copies are actually expressed in somatic cells. Here, we used a new cytological technique to demonstrate that rDNA is regulated allelically in a regional manner, with one parental copy of each NOR being repressed in any individual cell. This process is similar to that of X-chromosome inactivation in females. Early in development, one copy of each NOR becomes late-replicating, thus probably marking it for inactivation and subsequent targeted de novo methylation at rDNA promoter regions. Once established, this multichromosomal allelic pattern is then maintained clonally in somatic cells. This pathway may serve as an epigenetic mechanism for controlling the number of available rDNA copies during development.

[Keywords: Ribosomal DNA; replication timing; monoallelic expression; early development; DNA methylation]

Supplemental material is available at http://www.genesdev.org.

Received June 18, 2009; revised version accepted August 27, 2009.

Despite the clear-cut organization of rDNA copies into specific loci on the human genome, little is known about the logic of ribosomal gene expression. All ribosomal RNA synthesis is carried out by RNA polymerase I (PolI) within a small number of distinct nucleoli that appear to be formed for this purpose [Boisvert et al. 2007; McStay and Grummt 2008]. Even though all of the rDNA copies are probably associated with these nucleoli, only ~50% are actually transcribed [Grummt 2007], with the remaining genes being inactivated through a combination of different epigenetic mechanisms that take advantage of specific repression factors [Grummt et al. 1986; Santoro et al. 2002], noncoding RNAi [Mayer et al. 2006, 2008], DNA methylation [Santoro and Grummt 2001; Ghoshal et al. 2004], and late replication timing [Li et al. 2005]. Despite this general understanding of silencing at the individual gene level, little is known about how genes are chosen for inactivation in each cell. In addition, it is not yet clear how the ribosomal genes are regulated during normal development.

DNA replication timing is a regional epigenetic marking mechanism that is correlated with gene expression. The entire genome is divided into replication time zones that are organized in a distinct banding pattern on each chromosome. Active gene regions usually replicate early during S phase, while inactive genes replicate late, with only a relatively small number of genes not adhering to this correlation [Hiratani and Gilbert 2009]. In keeping with this correlation, regions of monoallelic expression have been found to replicate asynchronously, with one allele copied earlier than the other [Goldmit and Bergman 2004]. In the case of imprinted gene regions, this differential replication pattern is actually established in the gametes with the paternal allele preferentially replicating early in S [Simon et al. 1999]. In contrast, for many other monoallelic regions, differential replication timing is set up randomly in the early embryo, with one allele, either the maternal or paternal, being early and the other allele being late. Once established, this binary pattern is then clonally maintained in somatic cells [Mostoslavsky et al. 2001]. This early marking system apparently plays a role in allele usage during development. The establishment of late replication timing, for example, represents one of the earliest events of X-chromosome inactivation in female embryos [Keohane et al. 1998], and for many autosomal regions, asynchronous replication is instrumental in determining the preferred expression of one allele in each cell [Gimelbrant et al. 2005]. This early differential mark also plays a role in the preselection of a single receptor allele for recombination in the immune system [Goldmit et al. 2005]. In light of the regional nature of replication timing, we reasoned that this mechanism may also be involved in the silencing of rDNA.

Results

Replication timing-specific hybridization (ReTiSH)

In order to determine how rDNA clusters replicate during S phase, we developed a new approach for visualizing
BrdU uptake specifically at these loci. In this method, cells are labeled with BrdU for various times and then blocked in metaphase. Incorporation into mitotic chromosomes is visualized by a modification of CO-FISH (Cornforth and Eberle 2001), a technique in which replicated regions (BrdU-labeled) are converted to ssDNA and then hybridized directly with specific probes (Fig. 1A). To test this new methodology, we analyzed the hybridization patterns of control genes that replicate at different times during S phase in mouse embryonic fibroblasts (MEFs). Hybridization using a probe for the Adyc8 gene region (Fig. 1B), for example, revealed a signal on both chromosome 11 homologs when BrdU was initiated in either early S (12 h) or late S (7 h), consistent with the fact that this region replicates in late S phase. In contrast, labeling of the early-replicating A2M gene is only observed when BrdU is present from the beginning of S phase, and no signal was obtained if BrdU was added for only 7 h (Fig. 1C). Analysis of an olfactory receptor locus in human lymphoblasts revealed that both alleles are labeled if BrdU is initiated during early S phase, but only one can be observed when BrdU is added during late S phase (Fig. 1D), consistent with the fact that this gene undergoes asynchronous replication (Chess et al. 1994). In all ReTiSH experiments, the labeling and detection procedures were calibrated to ensure that all signals are indeed BrdU-dependent. 

Ribosomal genes replicate bimodally

The ReTiSH technique was then applied for analyzing replication timing at the five pairs of rDNA loci in the nucleus by labeling human lymphoblast cells with BrdU for various times. Strikingly, replication at these sites is seen to occur in a bimodal pattern, with approximately five loci replicating early in S phase and the remaining five replicating late (Fig. 2). Similar results were observed in primary human foreskin fibroblasts, as well. While this experiment confirms that ~50% of the rDNA copies replicate late in S phase (Li et al. 2005), it also indicates for the first time that the control of rDNA replication timing is carried out in a regional manner with each individual locus undergoing uniform replication either early or late in the cell cycle. These results are consistent with previous FISH experiments showing the presence of both replicated and unreplicated rDNA loci in the same interphase nucleus (Haaf 1997).

Figure 1. ReTiSH. (A) Scheme of ReTiSH method showing how BrdU-labeled regions are converted to ssDNA so that they can be identified by hybridization with specific probes. MEFs were labeled with BrdU for 7 or 12 h, arrested in metaphase, and subjected to ReTiSH using BAC probes for the late-replicating gene Adyc8 (RP23-352F10) (B) or the early-replicating gene A2M (RP23-191C6) (C). (D) Human lymphoblasts were labeled with BrdU for 7 or 12 h, arrested in metaphase, and subjected to ReTiSH using a BAC probe (RP11-55G7) for the asynchronously replicating olfactory receptor (family 4) gene cluster on chromosome 14 (OR4). Arrows indicate relevant gene signals.
rDNA replication is allelic

In order to test whether replication timing is regulated in an allelic manner, we labeled cells for 7 h with BrdU and carried out ReTiSH using one probe for detecting rDNA and the second specific for the subcentromeric region on chromosome 14. Under these conditions, we observed five labeled rDNA loci in each cell. We then counted the percentage of cells that have only a single copy of rDNA replicated on chromosome 14 [Fig. 3A]. If replication timing occurs in a completely random manner, one would expect to find a single copy of rDNA from chromosome 14 among the five late-replicating loci in only 55% of the metaphase spreads. In contrast, our analysis showed that this occurred in ≥80% of the cases. A similar pattern was observed for rDNA replication on other chromosomes (15, 21, and 22), as well. Altogether, we analyzed 430 metaphase spreads, with 86% of them showing a single replicated rDNA locus for each chromosome. This indicates \( P < 10^{-41} \) that replication of rDNA is controlled allelically, with one allele replicating early and one replicating late in almost every cell. This pattern clearly accounts for the observation that ≥50% of rDNA copies are late-replicating and inactive in any given cell [Li et al. 2005].

rDNA asynchrony is clonally maintained

In all of the other examples of asynchronous replication in the genome, once set up in the early embryo, the allelic pattern is clonally inherited, with the maternal allele replicating early in some clones and the paternal allele replicating early in others [Mostoslavsky et al. 2001; Singh et al. 2003; Ensminger and Chess 2004]. In order to determine whether this is also true for the rDNA loci, we carried out ReTiSH on single-cell-derived clones from human lymphoblasts containing a polymorphically enlarged centromeric region [Kitsberg et al. 1993b] on one homolog of chromosomes 14, 15, and 22. This enabled us to distinguish between the two parental rDNA alleles on these chromosomes [Fig. 3B]. When this technique was applied to a pool of human lymphoblasts using a 7-h BrdU incubation, we found that rDNA on chromosome 15 or 22 replicates late on the large centromeric allele in some cells while the other copy replicates late in others [Fig. 3C], and this is consistent with observations on other asynchronous replicating regions in the genome. For some unknown reason, the rDNA locus on chromosome 14 appears to have a fixed allelic pattern in this particular individual.

We then analyzed single-cell subclones by ReTiSH using a 7-h incubation with BrdU. In contrast to the pattern seen in the overall cell population, metaphase spreads showed that one particular allele (the maternal or paternal) in each clone is preferentially replicated in late S phase. In order to confirm this result, we carried out a BrdU pulse-chase experiment that specifically labels DNA undergoing replication during early S phase. In this way, we were able to demonstrate that it is indeed the opposite centromeric allele that is early-replicating in these same cells [Fig. 3B]. As expected, for the nucleolar organizer regions [NORs] located on chromosomes 15 and 22, we identified clones having either the paternal or maternal allele late-replicating, and these loci appear to segregate independently. Overall, this epigenetic
clonality is consistent with previous experiments showing that the same rDNA molecules are late-replicating and methylated in succeeding cell generations (Li et al. 2005).

Histone modification is monoallelic at each rDNA locus

In light of the general correlation between replication timing and gene expression profiles, our results showing rDNA allelic replication strongly suggest that these genes may be selectively transcribed in a regional manner from a single allele at each locus. Preliminary experiments indicated that both primary RNA transcripts as well as the RNA polIII machinery are retained at rDNA loci during metaphase, making it difficult to assay expression in a direct manner. Thus, in order to identify active and inactive rDNA regions, we took advantage of the fact that actively transcribed gene regions are packaged with acetylated histones, while repressed genes are not. This turns out to be an excellent indicator of gene expression, since these structural features are known to be maintained on metaphase chromosomes (Jeppesen and Turner 1993). As a first step, we prepared metaphase nuclei from human lymphoblast cell lines, using FISH to label all of the rDNA loci and immunohistochemistry with anti-H4Ac or anti-H3Ac to detect regions of histone acetylation (Fig. 4A). In these initial assays, we were able to demonstrate that approximately half of the rDNA loci are packaged with acetylated histones and methylated H3K4 (H3K4me), another structural marker of active gene expression (Fig. 4A). This regional pattern is consistent with previous experiments showing that ~50% of individual rDNA copies are packaged in an open chromatin conformation (Grummt 2007).

In order to confirm the relationship between early replication and histone acetylation at rDNA loci, we then analyzed individual lymphoblast clones carrying a centromeric polymorphism on both chromosomes 14 and 15, thereby allowing us to distinguish between the two parental alleles. Our results already showed that in clone 2 the maternal (small centromere) rDNA allele on chromosome 15 replicates early (Fig. 3). Using anti-H4Ac, we were able to demonstrate that this same allele is
preferentially packaged with acetylated histones, suggesting that this rDNA gene cluster is in an active chromatin conformation, as opposed to the unlabeled paternal allele (Fig. 4B), which replicates late. In contrast, in clone 127, it is the paternal allele (large centromere) that is marked by H4Ac, and this is in keeping with it being early-replicating in these cells (Fig. 3). Similar results were obtained using an antibody to H3K4me. These markers are also correlated with early replication at the rDNA locus on chromosome 14 [legend for Fig. 4]. Taken together, these observations show that early replication at rDNA loci is associated with an open chromatin structure, strongly suggesting that these regions are expressed monoallelically.

rDNA replication is coordinated with other monoallelic markers

Previous studies have shown that asynchronous replication of many monoallelic gene regions is coordinated at the chromosomal level (Singh et al. 2003; Ensminger and Chess 2004). Thus, for example, both the Igκ and Tcrβ loci on chromosome 6 replicate asynchronously, but the early allele of one is always on the same parental chromosome that has the early allele of the other. This coordination has also been seen for many olfactory receptor loci as well as other monoallelic markers. These results suggest that random asynchronous replication may be controlled by independent master regulators associated with each individual chromosome. In order to test whether rDNA loci also replicate in coordination with other markers on the same chromosome, we performed several double-label experiments (Fig. 5A). As a first step, we carried out standard FISH analysis on interphase nuclei from a single clone containing a pericentromeric polymorphism on chromosome 14, using a probe for an olfactory receptor gene located nearby on the same chromosome. Analysis of nuclei with a single/double hybridization pattern for the olfactory receptor locus revealed that the early allele (double dot) is almost always associated with the enlarged pericentric heterochromatin region (paternal allele), and this is the same chromosome that was found to have the early-replicating rDNA locus in these same cells (Fig. 3C). Similar results were also obtained using the ReTISH technique. In this case, a single clone was labeled with BrdU exclusively in late S (7 h), and resulting metaphase chromosomes were analyzed by hybridization with probes for the centromeric region of chromosome 15, rDNA, and an olfactory receptor region located on this same chromosome. These studies showed that whenever the rDNA is labeled, the olfactory receptor locus is also labeled on the same chromosome (Fig. 5B,C), indicating that both regions are late-replicating in a coordinate manner. A similar pattern was observed for an olfactory receptor gene region on chromosome 21 (Fig. 5D).

Developmental control of rDNA replication timing

In other examples of monoallelic expression, asynchronous replication is first established in the early embryo at about the time of implantation (Mostoslavsky et al. 2001). In order to test if this is also true for rDNA, we carried out ReTISH kinetics on early blastocysts grown from fertilized mouse oocytes in vitro. This analysis
revealed that, unlike the asynchronous pattern seen in somatic cells, in blastocysts, replication of all 10 rDNA loci is highly synchronized and takes place early in the cell cycle (Fig. 6A). In contrast to these very early embryos, rDNA replication was already found to be asynchronous in mouse embryonic stem (ES) cells. Even though these experiments were carried out in culture, the results strongly suggest that, in vivo, the switch to differential replication actually occurs at about the time of implantation [Tesar et al. 2007].

Methylation of rDNA

Although it is well known that ~50% of the rDNA copies are methylated at selected sites within the promoter region [Santoro and Grummt 2001; Ghoshal et al. 2004], the exact developmental pattern of methylation at these loci has never been determined. To this end, we carried out bisulfite analysis on rDNA from early embryos, ES cells, and somatic tissues [Fig. 6B; Supplemental Fig. 1]. The results show that while some regions of the rDNA repeat domain are completely methylated in spleen cells, a small window covering the gene promoter and extending ∼700 base pairs (bp) downstream appears to have a differential pattern, with ~50% of the molecules being completely unmethylated. In contrast, this same region is unmethylated on all alleles both in the early embryo [blastocysts] and in ES cells, and may still remain unmodified in MEFs [Schmittz et al. 2009]. This strongly suggests that the differential pattern seen in somatic tissues is established by de novo methylation of one allele in each cell, in a process that apparently takes place post-implantation. Previous studies have already demonstrated that this DNA modification is always associated with the late-replicating copies of rDNA [Li et al. 2005].

**rRNA expression during early development**

The fact that rDNA is early-replicating and completely unmethylated in the blastocyst as opposed to somatic tissues raises the possibility that the expression pattern of these genes may be developmentally regulated. To determine if this might be the case, we measured rRNA levels in blastocysts as compared with somatic tissues, such as brain or spleen. This was carried out by quantitative RT–PCR analysis using a set of three housekeeping genes to normalize the data. These studies show that...
there is approximately twofold more rRNA expression in the early embryo (Fig. 6C). This is consistent with studies showing that rRNA levels are elevated in undifferentiated ES cells, and are then down-regulated approximately twofold following differentiation to neural precursors (Efroni et al. 2008). In confirmation of this idea, we also found that, as opposed to somatic cells, >90% of the rDNA loci in ES cells are marked with both H3Ac and H3K4me. These experiments suggest that, during very early development, rRNA may be initially transcribed from all 10 loci and then undergo allelic repression at some later stage of differentiation.

Discussion

Although it has been known for a long time that a large percentage of the rDNA genes are inactivated in every somatic cell, the fundamental mechanism and molecular logic for this developmental process have remained elusive, mainly because it is difficult to study the epigenetics of repeated sequences. In order to overcome this problem, we developed a new cytological technique [ReTiSH] for measuring the time of replication at any sequence in the genome. This method is highly accurate, can be carried out on a very small number of cells in vivo, and, unlike the FISH method, is not affected by chromatid segregation (Azuara et al. 2003), so it yields easily interpretable results. In addition, this approach can be used to relate gene-specific replication timing to other chromosome markers.

Our studies indicate that rDNA repression takes place in an ordered, preprogrammed manner as part of a developmentally regulated pathway for distinguishing between chromosomal homologs in each cell. Thus, rDNA loci appear to be controlled in a manner similar to the X chromosome in female embryos. In the blastocyst, individual clusters [NORs] on both parental alleles replicate synchronously in early S phase and are all probably in an open conformation that allows transcription. At about the time of implantation, as seen in ES cells, one allele of each locus is stochastically chosen to undergo late replication. In some cells it is the paternal allele that is...

Figure 6. Regulation of rDNA loci during embryogenesis. [A] MEFs, mouse ES cells, and mouse blastocyst cells were labeled with BrdU for various times; arrested in metaphase; and subjected to ReTiSH using a probe for rDNA. The number of rDNA signals in each metaphase spread (n = 193) was counted, and the standard deviation was calculated for each data point (error bars). The data were adjusted to account for the shorter G2 (2 h) in blastocysts, as determined by the time at which 50% of metaphases are labeled with BrdU. [B] Diagram showing the stereotypic rDNA repeat indicating the transcription start site and the internal [ITS] and external [ETS] spacer regions. Methylation of the 5′ rDNA region was determined by bisulfite analysis on individual molecules (Supplemental Fig. 1), and the results are recorded as a function of position. The numbers of unmethylated (tan) and fully methylated [brown] molecules are shown. [C] Levels (±SD) of rRNA from mouse blastocysts and adult brain or spleen were measured by quantitative RT–PCR and normalized against a set of three housekeeping genes. [D] Metaphase spreads from mouse cells were used for immunohistochemistry with anti-H3Ac or anti-H3K4me and then hybridized. Results show the percent of rDNA loci in each nucleus (n = 127) that are labeled with H3Ac or H3K4me.
chosen, while in others it is the maternal copy, but each of the rDNA-containing chromosomes makes this decision independently. As demonstrated previously, asynchronous replication of multiple markers is coordinated on each chromosome (Singh et al. 2003; Ensminger and Chess 2004). Once these multichromosomal asynchronous replication patterns are established, they are then maintained in a clonal manner in differentiated cells. DNA methylation of rDNA appears to represent a secondary event that occurs post-implantation in a manner similar to what happens to the inactive X chromosome (Lock et al. 1987; Keohane et al. 1998) as well as genes responsible for pluripotency (Feldman et al. 2006; Epsztejn-Litman et al. 2008).

It should be noted that although the rDNA loci replicate synchronously in the blastula embryo, replication is seen to be asynchronous in ES cells that have been shown to have a developmental epigenetic pattern representative of the implantation stage. Despite this major change in the replication timing profile, almost all of these loci are packaged with H3Ac and H3K4me, and are probably fully expressed in these cells [Efroni et al. 2008]. It thus appears that the switch to asynchronous replication occurs prior to the silencing of rDNA. Since there is a clear-cut allelic association between late replication timing and repressed chromatin structure in somatic cells (Fig. 4), these results suggest, but do not prove, that replication timing patterns at an early stage may serve as an epigenetic template for choosing which allele is marked for inactivation.

This concept may also be true for other examples of monoallelic choice. Several studies, for example, have demonstrated that the X chromosomes in female ES cells replicate asynchronously prior to inactivation (Gribnau et al. 2005; Dutta et al. 2009), and the same is true for Igκ in pre-B cells, where this locus replicates asynchronously in a manner that predicts which allele is ultimately selected for the first rearrangement reaction (Goldmit et al. 2005). Differential replication timing of the parental alleles at imprinted gene loci is set up in the gametes (Simon et al. 1999), and this appears to occur prior to allele-specific methylation, which many times is actually established following fertilization (Birger et al. 1999; El-Maarri et al. 2001). All of these examples clearly indicate that asynchronous replication may represent an early epigenetic mark that can be recognized and used for allelic choice.

While this regional mechanism may be involved in establishing the initial epigenetic state of rDNA loci during development, it is clear that ribosomal RNA synthesis may also be controlled in a dynamic manner (Hammond and Bowman 1988). It is known that NORs may be subject to physiological influences mediated by both positive and negative transcription factors that operate within the nucleolus (Grummt 2007). Interestingly, overexpression of TIP5, a key component of the nucleolar remodeling complex [NoRC], is actually capable of altering the epigenetic structure of individual rDNA copies by its ability to reorganize nucleosome positioning and to recruit Dnmts that methylate key sites in the promoter region, and through the action of histone deacetylases that may be responsible for shifting the DNA to a late replication profile, perhaps by modifying nucleosomes at nearby replication origins (Santoro et al. 2002; Santoro and Grummt 2005). Recent studies further suggest that inactive copies of rDNA can even be induced to undergo demethylation (Schmmitz et al. 2009). Other factors may be involved in maintenance of rDNA epigenetic structure (McStay and Grummt 2008).

It should be noted that this allelic mode of rDNA inactivation clearly provides a molecular mechanism for ensuring that the same number of rDNA loci are repressed in each cell. Furthermore, since each individual locus carries variable numbers of rDNA gene copies, this system also guarantees that ~50% of the genes are repressed—the same number in each cell. Although rRNA levels can certainly be regulated by a number of different nucleolar transcription factors, maximal production of nontranslated RNA products of this nature is often controlled by gene copy number [Brown and Dawid 1968; Gall 1968]. Our studies now demonstrate, for the first time, that mammalian ribosomal genes may indeed be subject to an epigenetic mechanism that allows for an increase in effective copy number during early embryogenesis. Considering the strict coordination between rRNA and ribosomal protein levels (LaMarca and Wassarman 1979; Hammond and Bowman 1988), this could play a role in providing additional ribosomes to keep up with the increase in protein synthesis that occurs during blastulation [LaMarca and Wassarman 1979].

Materials and methods

Cell culture

Primary lymphocytes were prepared from peripheral blood of YC [Kitsberg et al. 1993a] and transformed by EBV. Single-cell clones were then isolated by serial limiting dilutions. Mouse ES cells [ESD3] were grown in DMEM supplemented with 1000 U/mL LIF on 0.1% gelatin-coated plates. MEFs were cultured from 13.5-d embryos. All cells were grown in DMEM or RPMI medium supplemented with 20% inactivated fetal calf serum [HyClone], 2 mM glutamine, 1000 U/mL penicillin, 100 μg/mL streptomycin, 0.1 μg/mL β-mercaptoethanol, 5 μg/mL Na pyruvate, and 5 μg/mL nonessential amino acids.

ReTiSH

Unsynchronized, exponentially growing cells were labeled for various times with 3 × 10⁻⁶ M BrdU [Sigma], and Colcemid [Bet Haemek] was then added to a final concentration of 0.1 μg/mL for 1 h at 37°C. Cells were harvested and treated with prewarmed hypotonic KCl solution [0.075 M] for 30 min at 37°C, fixed, washed with methanol-glacial acetic acid [3:1], dropped gently on cold slides, and air-dried. Slides were then washed with PBS, fixed again with 2% formaldehyde for 2 min, washed in PBS [3×], and treated with pepsin [1 mg/mL in 2N HCl] for 10 min at 37°C. After additional washes with PBS and 2× SSC, the slides were then treated with Hoechst 33258 [0.5 μg/mL] for 15 min, exposed to 365-nm UV light for 20 min using a UV Stratalinker 2400 transilluminator [Stratagene], washed again in 2× SSC, and dried. Exoll [0.1 U per slide] [Promega] digestion was carried out in buffer for 10 min at 37°C and the slides were then washed
in 2× SSC, dehydrated through a series of cold ethanol washes, and air-dried. In BrdU pulse-chase experiments, cells were labeled with 1×10^{-6} M BrdU for 1 h, washed three times in PBS, and then grown for 10–12 h before treatment with Colcemid and fixation.

BAC probes were labeled directly with DIG or Biotin-conjugated nucleotides using a nick translation mix (Roche). The rDNA probe was labeled using a standard nick translation protocol substituting dTTP with dioxigenin-11-dUTP in a ratio of 2:1 (Selig et al. 1992). Labeled DNA probes (20–50 ng) were mixed with human placental DNA or mouse genomic sonicated DNA [10 μg] and human or mouse Cot-1 DNA [10 μg, Gibco], ethanol-precipitated, and resuspended in Hybridization Buffer (50% deionized formamide, 10% dextran sulfate, 2× SSC). After denaturation (5 min at 80°C), the probe mixture was preannealed for 10–30 min at 37°C in order to eliminate repeated sequences and then applied to undenatured specimens. Following hybridization overnight and subsequent washes, the slides were treated as described (Selig et al. 1992) with blocking solution (5% BSA, 4× SSC) for 30–10 min at 37°C, incubated with detection reagents for 30 min at 37°C in 1% BSA, 4× SSC, and 0.1% Tween 20, and then washed three times for 3 min each in 4× SSC and 0.1% Tween 20. Biotin-labeled probes were detected with rhodamine-conjugated avidin DCS [1:500 dilution, Vector Laboratories], and digoxigenin-labeled probes were detected with an anti-digoxigenin antibody conjugated to FITC (1:100 dilution; Roche). Counterstaining was done using diamidinophenylidole (DAPI; 200 ng/mL) in VectaShield. Visualization was accomplished by conventional fluorescence microscopy, and images were captured using a Nikon 90i C1 fluorescence microscope at 1000× magnification.

Standard FISH was performed as described previously [Lichter et al. 1990; Selig et al. 1992] by culturing cells for 1 h with BrdU (3×10^{-3} M), isolating nuclei using hypotonic KC1 (0.5%), treatment, and fixing them with methanol:acetic acid (3:1). Briefly, denaturation was carried out by incubation in 70% formamide and 2× SSC for 2 min at 72°C, and then slides were dehydrated by a series of ice-cold ethanol washes (70%, 90%, and 100% for 5 min each). Probes were labeled as described above. BrdU was detected by an anti-BrdU antibody (1:100; NeoMarkers) followed by Amca or rhodamine-conjugated antimouse antibody (1:100; Jackson Immunoresearch Laboratories).

**Preparation of blastocysts for ReTiSH**

Superovulated C57Black BALB/C F1 female mice were mated with males, and zygotes were removed from the oviducts 14–20 h after hormone injection. Zygotes were incubated at 37°C for 24 h and then applied to undenatured specimens. Following hybridization, overnight and subsequent washes, the slides were treated as described (Selig et al. 1992) with blocking solution (5% BSA, 4× SSC) for 30–10 min at 37°C, incubated with detection reagents for 30 min at 37°C in 1% BSA, 4× SSC, and 0.1% Tween 20, and then washed three times for 3 min each in 4× SSC and 0.1% Tween 20. Biotin-labeled probes were detected with rhodamine-conjugated avidin DCS [1:500 dilution, Vector Laboratories], and digoxigenin-labeled probes were detected with an anti-digoxigenin antibody conjugated to FITC (1:100 dilution; Roche). Counterstaining was done using diamidinophenylidole (DAPI; 200 ng/mL) in VectaShield. Visualization was accomplished by conventional fluorescence microscopy, and images were captured using a Nikon 90i C1 fluorescence microscope at 1000× magnification.

Bisulfite analysis

Bisulfite conversion of genomic DNA was carried out using the Qiagen EpiTect Bisulfite Kit according to the manufacturer’s instructions. PCR primers were designed using Methyl Primer Express software version 1.0 (https://www2.appliedbiosystems.com). Primer sequences are available on request. After PCR amplification, products were extracted from gels using the Qiagen MinElute Gel Extraction Kit, cloned into the Promega pGEM-T vector system, transfected into Promega JM109 competent bacteria, and plated on LB-Agar + Amp + X-gal and IPTG. The DNA from 10–12 white clones of each ligation was amplified using the Illustra Templiphi 100 amplification kit and subjected to sequencing with T7 or Sp6 primers.

Expression

Total RNA (250 ng) was prepared with the TriPure isolation reagent (Roche) and converted to cDNA using the M-MLV reverse transcriptase (Promega) and random hexanucleotide pd(N)6 primers (GE Healthcare) in a reaction volume of 20 μL under conditions recommended by the manufacturer. In some cases, RNA was first treated with DNase I (Promega). Control reactions lacking reverse transcriptase were systematically verified for the absence of products. Quantitative real-time PCR (Q-PCR) was performed (Finnzymes) in a 20-μL reaction mix, and primer–dimer products were distinguished from specific products by dissociation analysis. For each primer pair designed for Q-PCR, a no-template control was also included. Measurements were performed with SYBR Green Universal Mix in triplicate, and three control genes were used to normalize between samples. For each cDNA sample, cycle thresholds for the target gene were calculated relative to that of the three control genes (HP1-α, Cyclophilin-A, and Cyclophilin-B) (see Supplemental Table 1).

Acknowledgments

We are grateful to B. McStay for the gift of the human rDNA intergenic spacer probe, to A. Rahat for the chromosome 15 α satellite probe, and to Y. Kaufman for his assistance in analyzing blastocyst DNA methylation. This work was supported by grants from the Israel Academy of Sciences (to Y.B. and H.C.), the NIH, European Community 5th Framework Quality of Life Program (to Y.B.), the Israel Cancer Research Fund (to Y.B. and H.C.), and a research award from Lew Sanders (to H.C.).

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*Genes Dev.* 2009, 23:
Access the most recent version at doi:10.1101/gad.544509

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