Characterization of the 55-kb Mouse Histone Gene Cluster on Chromosome 3


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The histone gene cluster on mouse chromosome 3 has been isolated as a series of overlapping P1 clones, covering 110-120 kb, by probing with the histone H3-614 gene that had been mapped previously to mouse chromosome 3. There are genes for 10 core histone proteins present in a 55-kb cluster within this contig. There are three histone H3 genes, two of which are identical; four histone H2a genes, two of which are identical, one histone H4 gene; and two histone H2b genes. These histone H3 and H2a genes encode ~40% of the total H3 and H2a mRNA, whereas the histone H4 and histone H2b genes encode <10% of the total H4 and H2b mRNA. There are no histone H1 genes present in this cluster. All of the histone H2a genes encode histone H2a.2 proteins (or variants of H2a.2), and account for all the H2a.2 genes in the mouse genome. All three histone H3 genes encode the histone H3.2 protein. A 2.1-kb region containing the adjacent H3-614 and H2a-614 genes has been duplicated and is present in an inverted repeat separated by 4.5 kb. The other two H2a genes are adjacent, with the 3' ends of their mRNAs separated by only 49 nucleotides in the DNA and the U7 snRNP binding sites separated by only 20 nucleotides. One of the histone H2b genes has lost the stem-loop sequence characteristic of the replication-dependent histone mRNAs and encodes only polyadenylated mRNAs.

The histone proteins are encoded by a multigene family, consisting of 10-20 genes for each of the core histone proteins in mammals (Jacob 1976). Most of the histone genes are replication dependent, and their mRNAs are expressed coordinately with DNA replication. The replication-dependent histone mRNAs are the only metazoan mRNAs that lack a poly(A) tail, ending instead in a highly conserved 26 nucleotide sequence that can form a stem-loop (Birnstiel et al. 1985; Marzluff 1992). The histone genes in mammals are present in two clusters on separate chromosomes; chromosomes 1 and 6 in humans (Triputti et al. 1986) and chromosomes 3 and 13 in mouse (Graves et al. 1985b). The majority of the replication-dependent histone genes (Graves et al. 1985b), including all the histone H1 genes (Wang et al. 1996), are on mouse chromosome 13, and human chromosome 6 (Doenecke et al. 1994), whereas a smaller number of genes are present on mouse chromosome 3 and human chromosome 1. Two of the genes on mouse chromosome 3, H3-614 and H2a-614, encode a significant fraction of the histone H3 and H2a mRNA (Graves et al. 1985b), suggesting that the minor histone gene cluster makes an important contribution to the total histone mRNA population.

We have described a 2-Mb region from mouse chromosome 13 containing 45 histone genes (Wang et al. 1996). Here we describe a 120-kb region containing 10 histone genes clustered in 55 kb on mouse chromosome 3. There are genes for all four core histones present in this cluster. There are four H2a genes and three H3 genes, which produce a significant fraction of the total histone H3 and H2a RNA. The two H2b genes and single H4 gene, in contrast, are expressed at a low level.

RESULTS

We previously identified two histone genes, H2a-
614 and H3-614, which mapped to mouse chromosome 3 (Graves et al. 1985b). To see whether there were other histone genes linked to these genes, we screened a P1 library by PCR with specific primers flanking the histone H3-614 gene. Four overlapping P1 clones (P0, P1, P2, and P3), which formed a single contig, were obtained. Each of these P1 clones contained multiple histone genes. Because some of these genes were close to the end of this contig, the P1 library was rescreened with primers specific for the end of the P0 clone. Two additional P1 clones (P91 and P92) were obtained, but these extended the contig only an additional 9 kb. By use of a probe from the end of the P92 clone, we screened a genomic library and isolated a λ phage, λ111, which extended the contig an additional 12 kb. By use of gene specific probes we identified genes for all of the core histones in this contig by Southern blotting.

There were 10 histone genes identified in a 55-kb region, which was completely included in clone P0 (Fig. 1). Flanking these genes was at least 40 kb on one end and 22 kb on the other end with no additional histone genes. Southern blots identified three histone H3 genes (Fig. 2A), three fragments containing histone H2a genes (one of which contained two H2a genes; Fig. 2B), a single histone H4 gene, and two histone H2b genes (Fig. 2C). By use of specific probes for the H2a-614 and H2a-615 5' flanking regions and the H3-614 and H3-615 3' flanking regions, we determined that the 5' ends of the H2a-614 and H2a-615 genes were on the same SalI fragment, whereas the 3' ends of the H3-614 and H3-615 genes were on different SalI fragments (Fig. 2D). Thus, this cluster is oriented with the 5' ends of the H2a genes juxtaposed. Figure 1 shows a map of this region, with the location and orientation of the histone genes indicated.

The Hist2 Gene Cluster Maps to Mouse Chromosome 3F1-2

By use of fluorescence in situ hybridization (FISH), we hybridized one λ clone and four phage P1 clones to mouse metaphase chromosomes. The signals from each probe showed exclusive hybridization to chromosome 3F1-2. No cross-hybridization to the Hist1 locus on chromosome 13 was detected. Examples of these images are shown in Figure 3, A and B. DAPI(G/Q) banding patterns place the Hist2 cluster within the proximal half of band 3F; fractional length measure-

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**Figure 1** Structure of the histone gene cluster on chromosome 3. A schematic of the histone gene cluster on chromosome 3, with the orientation of each gene, is shown. The structure of the P1 clones used to deduce the organization of the cluster is included and the position of the SalI sites (○) and SphI sites (■) on the clones are indicated. λ111 is the λ phage that overlaps the end of the P92 clone. (*) T7 promoter; (●) SP6 promoter. The position of the genes was determined by use of Southern blotting (Fig. 2). The orientation of the H2a-613 and H2b-613 genes was determined by cloning the end of the P0 clone and sequencing. The orientation of the H2b-616 and H3-616 genes was determined by analysis of SphI digests of the P0 and P3 clone DNA with probes specific to the 5'- and 3'-flanking regions of the H3-615 and H3-616 genes. The orientation of the H4 gene was determined by PCR amplification of a fragment from the 3' end of the H3-616 gene to the H4 gene and was confirmed by sequencing of this fragment.
ments were in good agreement with these results and allowed us to refine the locus to the F1-F2 subregion (see Fig. 3C; FLpter = 58–66%).

The Histone H2a-614 and H3-614 Genes Are Duplicated

The regions containing the histone genes were cloned and sequenced. Surprisingly, two of the linked histone H3 genes and histone H2a genes were nearly identical (24 changes out of 1720 nucleotides) in the coding and intragenic regions. The region of near identity extended for 2122 nucleotides, from 323 nucleotides 5' of the start of the H2a protein to just past the U7 snRNP-binding site in the H3 gene (Fig. 4B). One of these pairs was the previously cloned H2a-614 and H3-614 genes, and the other pair was named H2a-615 and H3-615. The position and orientation of these two sets of genes was unambiguously established by use of probes 3' of the H3 genes and 5' of the H2a genes in Southern blots (Fig. 2D). The H2a-614 and H2a-615 5' flanking probes hybridized to the same 8-kb SalI fragment, present on clones P0, P1, and P2 but not on P3, demonstrating that these genes were located next to each other (Fig. 2D, lanes 1–6). The H3-614 3' flanking probe hybridized to a 15-kb SalI fragment that was present only on clones P0, P1, and P2 but not on P3 (Fig. 2D, lanes 7–9). The H3-615 3' flanking region probe hybridized with a 17-kb SalI fragment present on all the P1 clones (Fig. 2D, lanes 10–12). The 3' end of the H3-615 gene was the probe that hybridized with clone P3, indicating that this clone ended between the H2a-615 and H3-615 genes.

This result unambiguously places the 3' end of the H3-615 gene near one end of the cluster defined by clone P3. These results also demonstrate

Figure 2 Localization of the genes by Southern analysis. (A) DNA from the P0–P3 clones were digested with EcoRI and hybridized with a probe containing the coding region and part of the 5'-flanking region of the H3-614 gene. The size of the hybridizing fragments is indicated. The 9-kb fragment is the H3-614 gene, the 4-kb fragment is the H3-615 gene, and the 1.4-kb fragment is the H3-616 gene. The 5-kb fragment on the P3 clone is from the 5' end of the H3-615 gene, a result of the P3 clone ending in the fragment containing the H2a-615 and H3-615 genes. (B) The P0–P3 clones were digested with EcoRI and hybridized with the H2a-614 gene (coding region plus 500 nucleotides of 5'- and 300 nucleotides of 3'-flanking region). The 9-kb fragment is the H2a-615 gene, the 4-kb fragment is the H2a-614 gene, and the 6-kb fragment is the H2a(A)-613 and H2a(B)-613 genes. The 5-kb fragment in the P3 clone is the H2a-615 gene, a result of the clone ending in the fragment containing the H2a and H3-615 genes. (C) The orientation of the 614 and 615 gene cluster. The P0, P2, and P3 clones were digested with EcoRI and hybridized with specific probes for the H2a-614 5'-flanking region (614-5', lanes 1–3), the H2a-615 5'-flanking region (615-5', lanes 4–6), the H3-614 3'-flanking region (614-3', lanes 7–9), and the H3-615 3'-flanking region (615-3', lanes 10–12). The size of the hybridizing fragments is indicated. The weak 17-kb band detected by the 614-3' probe in the P3 DNA is from contamination of the probe with a small amount of vector DNA, which hybridizes with the 17-kb P1 vector.
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Figure 3  Chromosomal location of the mouse Hist2 gene cluster. Biotin-labeled mouse histone clones P0 (A) and λ111 (B) map to chromosome 3 (green). DAPI staining generated the G/Q banding pattern in both panels (blue). (C) An ideogram of mouse chromosome 3. The bar (at right) indicates that the range of signals for these probes localizes to 3F1-2.

that the 5' ends of the H2a genes are juxtaposed and that these gene pairs are oriented as shown in Figure 1.

The Histone H3-614, H3-615, and H3-616 Genes
In addition to the H3-615 gene, there was an additional histone H3 gene, H3-616, located ~20 kb from the H3-615 gene. This gene also encodes a H3.2 protein. The coding region of the gene contains only 4-nucleotide changes from the H3-614 and H3-615 genes (Fig. 4A). The 5'-untranslated and flanking region is completely different from the H3-614 and H3-615 genes, but the 3'-untranslated region is nearly identical to the H3-614 and H3-615 sequence to the end of the mRNA (Fig. 4B). The U7 snRNP binding site and 3'-flanking region, however, are different from both the H3-614 and H3-615 genes (Fig. 4B).

The SalI site conserved in all other mouse histone H3 genes and used for S1 mapping of these genes (Wang et al. 1996; Graves et al. 1985) is missing from the H3-616 gene because of a single base change. Thus, to determine whether the H3-616 gene was expressed, we labeled the gene at the 5' end of the EagI site at codon 88. When the histone H3-616 probe was hybridized to total cell RNA, two fragments were protected. One (267 nucleotides) mapped to the ATG codon and results from protection by a number of different histone mRNAs. The minor protected fragment (280 nucleotides) mapped to the 5' end of the mRNA expressed from the H3-616 gene (Fig. 4C, lanes 1,4). The 5' end is located ~25 nucleotides from a TATAA sequence, consistent with this being the start site of transcription (Fig. 4A). The H3-616 gene encodes ~6% of the histone H3 mRNA, a lower amount than encoded by the H3-614 and H3-615 genes, which together encode ~30%–35% of the histone H3 mRNA (Graves et al. 1985b). To demonstrate the difference in expression levels, probes for the H3-614 gene, labeled at the SalI site, were used together with the H3-616 probe. The H3-614 probe protects fragments of 180 and 220 nucleotides. The two probes were mixed and hybridized to total RNA from mouse myeloma cells. The four expected protected fragments were seen (Fig. 4C, lanes 5,6). The fragments mapping to the initiation codon (H3_M) were of similar intensity with each probe, whereas the specific fragment derived from the H3-616 gene was much less intense than the fragment derived from the H3-614 gene, demonstrating that the H3-616 gene was expressed at a low level relative to the H3-614 and H3-615 genes. Because the H3-614, H3-615, and H3-616 genes all have nearly identical 3'-untranslated regions, it was not possible to unambiguously detect the 3' end of the H3-616 mRNA.

The Histone H2a Genes
There were four histone H2a genes present in this cluster. Two of these, H2a-614 and H2a-615, encode the H2a.2 protein (Hurt et al. 1989) and make up ~30%–35% of the histone H2a mRNA
Figure 4  Structure and expression of the histone H3 genes. (A) Sequences of the histone H3-614, H3-615, and H3-616 genes. The promoters of the H3-615 (614) and H3-616 genes are compared, and the start site of transcription, the TATAA box, and CCAAT boxes are underlined. Where the three genes are identical in the coding region, only the H3-614 sequence is shown. (B) The 5'-flanking region of the H2a-614 and H2a-615 genes are compared around the point of divergence that is 323 nucleotides from the transcription start site, demonstrating the abrupt end of the region of identity. The 3'-flanking regions of the H3-614, H3-615, and H3-616 genes are compared from the end of the coding region to the region of divergence past the U7 snRNP-binding site. The conserved stem-loop at the end of the mRNA and the U7 snRNP-binding site are underlined. (C) Expression of the H3-614 (615) and H3-616 genes. To map the 5' end of the H3-616 mRNA the H3-616 gene was labeled at the 5' end of the Eagl site (lanes 1, 4) or the 3' end of the Eagl site (lane 2) and the probe hybridized to 2.5 (lane 5) or 5 (lanes 1-4, 6) µg of total cell RNA from mouse myeloma cells. The hybrids were treated with S1 nuclease and the nuclease-resistant fragments were resolved by electrophoresis on 6% polyacrylamide/7 M urea gels and detected by autoradiography. The H3-614 gene was labeled at the 5' end of the SalI site at codon 58, and this probe was used to analyze the same RNA samples (lane 3). (Lanes 5, 6) Two 5' labeled probes were mixed prior to hybridization.

(Graves et al. 1985b). The other two H2a genes, H2a(A)-613 and H2a(B)-613, are located immediately adjacent to each other with their 3' ends juxtaposed. We predict that the primary transcripts from these genes overlap, because the U7 snRNP-binding sites are only 20 nucleotides apart (Fig. 5A). Each of these genes encodes a protein similar to the H2a.2 protein, with the 4 amino acid differences characteristic of H2a.2 (shown in boldface type in Fig. 5A). However, their carboxyl termini differ from each other and from other H2a proteins. Thus, each of these genes encodes an H2a.2 protein variant that has not been described previously.

By use of S1 nuclease protection assays with the H2a(A)-613 gene labeled at the AccI site and the H2a(B)-613 gene labeled at the NarI site as probes, we identified mRNAs expressed from
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Figure 5 Structure and expression of the H2a genes. (A) The sequences of the H2a-614, H2a(A)-613, and H2a(B)-613 genes are compared, extending from the promoters of the gene to the region of overlap past the U7 snRNP-binding site. The overlapping sequences between these two genes are shown in bold italics. The transcription start site, the TATAA box, and CCAAT boxes are underlined as are the conserved stem–loop at the end of the mRNA and the U7 snRNP-binding site. (B) The expression of the H2a(A)-613 and H2a(B)-613 genes was determined by use of S1 nuclease mapping. The H2a-A gene was labeled at the 5' end (lane 1) or 3' end (lane 2) of the Accl site, and the H2a-B gene was labeled at the 5' end (lane 4) or the 3' end (lane 5) of the Ntll site. The probes were hybridized with 5 μg of total RNA from mouse myeloma cells and the protected fragments resolved by gel electrophoresis and detected by autoradiography.

Each of these genes (Fig. 5B). Specific fragments mapping to the 3' end and the 5' end of the two histone mRNAs were mapped. The level of expression of both of these genes was very low. The H2a(A)-613 gene protects two fragments; the major one (H2aM) derived from the H2a-614 and H2a-615 mRNA (151 nucleotides in the 5' map and 244 nucleotides in the 3' map), and the minor fragment (H2a, 200 nucleotides in the 5' map and 282 in the 3' map) protects the H2a(A)-613 mRNA. Note that because only the H2a-614 and H2a-615 genes contain an Accl site, which is absent in all of the histone H2a genes from chromosome 13, only these two mRNAs are protected to give the H2aM fragment. Thus, the expression of the H2a(A)-613 gene is much less than the closely linked H2a-614 and H2a-615 genes. There are two transcription start sites in the H2a(A)-613 gene, ∼5 nucleotides apart, corresponding to two A's located 22 and 27 nucleotides from the TATAA box (Fig. 5B, lane 1).

The H2a(B)-613 gene is also expressed at a...
low level. Use of the Ncol site, which is present in the H2a-614 and H2a-615 genes as well as the H2a genes on chromosome 13, generated two protected fragments with each probe (Fig. 5B, lanes 4,5). The major 5' fragment (H2aM, 135 nucleotides) maps to the AUG codon, whereas the minor fragment (H2a, 166 nucleotide) maps to the start site of transcription of the H2a(a)-613 mRNA (Fig. 5B, lane 4). The major fragment (H2aM) from the 3' labeled probe is only 120 nucleotides, and maps to the cluster of nucleotide changes unique to the H2a(B)-613 gene at amino acids 87 and 88 (Fig. 5B, lane 5). The minor 304-nucleotide fragment (H2a) maps to the end of the H2a(B)-613 mRNA, bearing in mind that the H2a-614 and H2a-615 genes account for ~30% of the H2a mRNA and the H2a(a)-613 and H2a(B)-613 genes each account for no more than 5% of the H2a mRNA. Given this low level of expression, it is possible that these two genes are not transcribed simultaneously, as there would then presumably be a collision between the two RNA polymerases. Alternatively, it is possible that the close proximity of the genes reduces the expression of the genes due to interference between two elongating RNA polymerase molecules.

The Histone H2b and H4 Genes

Two H2b genes, H2b-616 and H2b-613, are present, one at each end of the cluster. Each of these genes encodes an H2b protein, similar in sequence to the reported mammalian H2b protein sequence. Note that the predicted amino acid sequence of each of the seven mouse H2b genes that we have isolated (Liu et al. 1987; Brown et al. 1996) is different. The consensus sequence from this sequence agrees with the reported mammalian H2b protein sequence, which has heterogeneity in at least two positions (Ohe et al. 1979). On the basis of the results from cloned H2b genes, the mouse H2b proteins are a mixture of at least seven closely related polypeptides.

The promoters of the H2b genes have a structure similar to the other mouse (and metazoan) H2b genes (Harvey et al. 1982; Liu et al. 1987; Ito et al. 1989; Bell et al. 1992). There are three elements in the proximal promoter: a TATAA box, a CCAAT box, and an octamer sequence, which binds Oct-1 (Fletcher et al. 1987; Das and Herr 1993) located between the CCAAT and TATAA boxes. The octamer sequence is present in all known metazoan H2b genes and is necessary for expression of the genes (LaBella et al. 1988; Sturm et al. 1988; Bell et al. 1992). Although the CCAAT, octamer, and TATAA boxes are identically spaced from each other in the H2b-613 and H2b-616 genes, there are no other similarities between these promoters.

The mRNA expressed from the H2b-616 gene was detected by S1 nuclease mapping with the gene labeled at the EcoRI site at codon 62 as a probe. Because the EcoRI site is not present in any of the other H2b genes we have isolated, there was a single protected fragment with each probe, which extended to the end of the mRNA (Fig. 6B, lanes 3,4). To estimate the level of expression of the H2b-616 gene, we analyzed the same RNA samples with the H3-616 gene labeled at the 5' end of the Eagl site to the same specific activity (Fig. 6B, lane 2). The H2b-616 and the H3-616 genes were expressed at a similar level (Fig. 6B, lanes 1,3).

The H2b-613 gene was <250 nucleotides from the H2a(a)-613 gene, and their 5' ends are juxtaposed. This is an arrangement that is very common in the mouse H2a and H2b genes on chromosome 13 (Liu et al. 1987; Brown et al. 1996) and in the H2a–H2b genes in other species (Grandy and Dodgson 1987). The distance between these genes is essentially identical with the five H2a–H2b pairs from mouse chromosome 13 (Brown et al. 1996; Liu et al. 1987; Brown et al. 1996). The H2b-613 gene encodes a complete H2b protein, although it lacks the stem–loop normally present at the 3' end of all replication dependent histone mRNAs. There is a sequence with similarity to the stem–loop present at the normal location, ~50 nucleotides from the stop codon (underlined in Fig. 6A), suggesting that this gene previously contained a stem–loop sequence.

Despite the fact that the H2b-613 gene lacks a stem–loop, this gene was clearly expressed. Total RNA from mouse myeloma cells or from newborn mice was analyzed by S1 nuclease mapping with the H2b-613 gene labeled at the 3' end of the Ncol site as a probe (Fig. 6B, lanes 9,10). The probe used extended to the SalI site 313 nucleotides past the stop codon. The Ncol–SalI fragment was cloned into pGEM5zf, and the resulting clone digested with Ncol and 3' end-labeled at the Ncol site. Any RNAs extending past the SalI site would result in a 528 nucleotide protected fragment. There clearly is a protected fragment of the appropriate size indicating that the H2b-613 gene is expressed (Fig. 6B, lanes 9,10). There was
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Figure 6 Structure and expression of the H2b genes. (A) Sequences of the H2b-613 and H2b-616 gene. Where the genes differ in amino acid sequence, the amino acid in bold is the one that is not found in H2b consensus sequence. The promoters are compared with the Oct-1, CCAAT, and TATAA sequences aligned. The 3'-flanking regions are shown with the conserved stem-loop at the end of the mRNA and the U7 snRNP-binding site underlined in the H2b-616 gene. The H2b-613 gene lacks these sequences, but there are possible remnants of each of these sequences that are underlined. (B) The expression of the H2b genes was determined by 5' nuclease mapping. The H2b-616 gene was labeled at the 5' end (lane 3) or the 3' end (lane 4) of the EcoRI site at codon 62. The H3-616 gene labeled at the 5' end (lane 1) or 3' end (lane 2) of the Eagl site at the same specific activity was used as a control. These probes were hybridized with 5 μg of total RNA from mouse myeloma cells (lanes 1-4). The H2b-613 gene was labeled at the 3' of the Ncol site (lanes 9, 10) and the probe extended to the Sal site in the 3'-untranslated region. This probe was hybridized with 30 μg of total RNA from adult testis (lane 9) or neonatal mice (lane 10). As a control for the level of expression, the H2a(A)-613 gene was labeled to the same specific activity at the 3' end of the Eagl site (lanes 7, 8). This probe was hybridized with 5 μg of total RNA from testis (lane 7) or 5 μg of total RNA from newborn mice (lane 8). (Lanes 5, 6) Marker pUC18 digested with HpaII.

also a fragment of 201 nucleotides mapping to the stop codon. This is the result of protection of a minority of the other H2b mRNAs up to the stop codon. The 528-nucleotide fragment is from a polyadenylated mRNA (not shown), suggesting that this gene is expressed as a polyadenylated mRNA ending at a polyadenylation signal located at an unknown distance 3' of the coding region. The same pattern of expression was observed in both RNA from neonatal mice and mouse testis, a tissue that often expresses polyadenylated histone mRNAs (Challoner et al. 1989; Moss and Orth 1993; Moss et al. 1994). The level of expression of the H2b-613 gene was comparable to the expression of the H2a(A)-613 gene analyzed with the same RNA samples (Fig. 6B, GENOME RESEARCH @ 709 Cold Spring Harbor Laboratory Press on October 19, 2017 - Published by genome.cshlp.org Downloaded from
The H2b-613 gene was also expressed in a number of adult mouse tissues (data not shown) at a similarly low level. Thus, both of these H2b genes are expressed at low levels, contributing only a minority of the histone H2b mRNA.

There is a single H4 gene present in this cluster. This gene was first identified by Birnstiel and co-workers (Seiler-Tuyns and Birnstiel 1981). We identified the orientation and precise location of this gene by PCR with a primer from the 5′-flanking region of the adjacent H3-616 gene and with sense and antisense primers from the coding region of the histone H4 gene. Only the antisense fragment from the H4 gene gave a product that was sequenced and contained the H4 gene, indicating that the H3-616 gene and H4 gene are oriented in opposite directions. In agreement with the results of Seiler-Tuyns and Birnstiel (1981) in mouse L cells, this H4 gene is expressed at a very low level, contributing only a tiny fraction of the total mouse histone H4 mRNA (not shown). Thus, the great majority of the histone H4 mRNA is derived from the nine histone H4 genes on mouse chromosome 13.

The 10 histone genes on chromosome 3 encode a significant fraction of the histone H3 and H2a mRNAs but only a small fraction of the H2b and H4 mRNAs. There were no histone H1 genes present in this cluster.

**DISCUSSION**

The core histone proteins must be produced in equimolar amounts during S phase as the newly replicated DNA is packaged into nucleosomes. The high demand for histone mRNA to synthesize the 10^6 histone proteins required in each S phase presumably requires multiple copies of the histone genes. In addition, there are a number of nonallelic variants of the core histone proteins, which are present in all vertebrates (Zweidler 1984). Whether these variants are functionally significant is not known.

There are changes in the relative proportions of the different histone variants, both in different tissues and at different developmental stages (Zweidler 1984). The replacement histone H3.3 is synthesized constitutively and accumulates to high levels in the chromatin of terminally differentiated cells as they age (Zweidler 1984). The H3.3 protein is encoded by genes that contain introns and produce polyadenylated mRNAs (Brush et al. 1985; Wells and Kedes 1985), and these mRNAs are expressed constitutively. The histone H3.2 and H2a.2 proteins have been classified as partially replication dependent (Zweidler 1984), as they accumulate at higher levels in some terminally differentiated cells, although they are synthesized solely in S phase. Sittman and co-workers showed that the accumulation of the H3.2 and H2a.2 proteins in terminally differentiating erythroleukemic cells is the result of preferential expression of the H2a-614 and H3-614 (and presumably also the H3-615 and H2a-615) genes during the last S phase (Brown et al. 1985; Brown et al. 1988). They postulated that this accumulation might be a result of shutting off expression of histone genes on chromosome 13 (Brown et al. 1988) as cells terminally differentiate. Thus, it is somewhat surprising that the histone gene cluster on chromosome 3 does not contain any highly expressed histone genes for the other core histones. The histone H4 and H2b mRNAs are almost entirely derived (>90%) from the genes on mouse chromosome 13. The preferential expression of the H3-614 and H2a-614 (and H3-615 and H2a-615) genes must account for the accumulation of the H3.2 and H2a.2 protein variants as cells terminally differentiate. The H2b and H4 proteins, however, as well as the histone H1 proteins, continue to be produced primarily from mRNAs encoded on chromosome 13.

The H3.2 protein is the major histone H3 variant in rodents (Zweidler 1984), although it is a minor variant in cows (Marzluff et al. 1972; Patthy and Smith 1975) and humans (Wu and Bonner 1981). Of the nine histone H3 genes on mouse chromosome 13, five encode histone H3.2 and four encode histone H3.1 (Wang et al. 1996). The predominance of the H3.2 protein variant caused by the expression of ~35% of the histone H3 mRNA from chromosome 3, all of which encodes the H3.2 variant. In other mammals there is a much higher proportion of the H3.1 than H3.2 protein. It is likely either that the genes on the minor cluster in other mammals encode H3.1 proteins, or that all the genes on the major cluster encode histone H3.1 proteins, accounting for the change in relative proportions of the protein variants in different mammals.

All of the histone H2a genes that have been cloned from mouse chromosome 13 encode the H2a.1 protein (Liu et al. 1987; Kosciessa and Doecke 1989; Gruber et al. 1990; Brown et al. 1996). Thus, it is very likely that the H2a.2 protein is exclusively encoded by the genes on chromosome 3. The H2a.2 protein makes up >30% of
the H2a protein in most mouse tissues, consistent with the high level of expression of the genes on chromosome 3. A number of human H2a mRNAs have been characterized. The four human H2a.1 mRNAs each are derived from human chromosome 6 (Mannironi et al. 1994). The human H2a.2 mRNA, is derived from human chromosome 1 (Mannironi et al. 1994). The single human H2a.2 mRNA, like the mouse H2a.2 mRNA encoded by the H2a-614 and H2a-615 genes, was expressed at a higher level than the other human H2a mRNAs, encoding 35%-45% of the H2a mRNA (Mannironi et al. 1994). Thus, the overall organization and expression level of the individual histone genes has probably been conserved between mice and men.

The H2a-614 and H3-614 genes have been duplicated in this cluster. The region of near identity extends for >2.1 kb, ending precisely after the U7 snRNP binding site 3' of the H3-614 gene and starting 323 nucleotides 5' of the H2a genes. The H2a-614 gene has recently been found expressed in testis as a polyadenylated mRNA utilizing a cryptic polyadenylation site that overlaps with the U7 snRNP-binding site (Moss et al. 1994). This sequence is identical in the H2a-615 gene, which is presumably also expressed as a polyadenylated mRNA in testis. Thus, in addition to providing a large fraction of the H2a mRNA in somatic cells, this gene pair expresses a unique transcript in the testis. These genes also encode most of the H2a and H3 mRNA present in mouse oocytes, and early cleavage embryos (Graves et al. 1985a).

The two H2a.2-like genes, H2a(A)-613 and H2a(B)-613, and the H2b-613 gene each encode a very small amount of mRNA. Although these three genes are clearly functional, it is not at all clear whether these H2a protein variants are important. They may represent genes that have accumulated mutations in the protein but have still remained functional. The H2b-613 gene also encodes an H2b-like protein that differs significantly from other mouse histone H2b proteins. Moreover, all the mRNAs from this gene are polyadenylated, as the stem–loop sequence is missing. We did not see high-level expression of this gene in a number of different mouse tissues (Z.F. Wang and W.F. Marzluff, unpubl.), including testis (Fig. 6B), reducing the possibility that this is a tissue-specific gene.

There are closely linked histone H2a and H2b genes encoded on human chromosome 1. It is possible that these genes are orthologous to the H2b-613 and H2a-613 genes described here, particularly because the human H2b gene also expresses a polyadenylated mRNA as well as an RNA ending in a stem–loop (Collart et al. 1991, 1992). The function of the polyadenylated RNA is not clear. Normally, it is produced in small amounts and increases when the normal histone 3'-end formation is inactive, in terminally differentiated cells and in cells in which DNA replication has been inhibited (Collart et al. 1991). The predicted amino acid sequence of the human H2b protein differs from the H2b-616 protein, and only one of the amino acids (Val-39) unique to the H2b-613 gene is conserved in the human gene on chromosome 1. The linked human H2a gene has not been completely sequenced so it is not known whether it encodes the same protein as the mouse H2a(A)-613 gene. Also, it is not clear whether there is an additional H2a gene present in humans.

Evolution of the Histone Genes on Chromosome 3

The almost complete identity between the H3-614 and H3-615 genes and the H2a-614 and H2a-615 genes might indicate that there has been a very recent duplication event involving this region of chromosome 3. At least one line of evidence argues against that interpretation, however. Duplication of one of the most highly expressed H3 and H2a genes but not of any H2b or H4 genes would likely upset the stoichiometric relationship among the mRNA and protein levels of the four core histone (although this argument does presuppose that equal production of the four core histones is important, and that there is no feedback mechanism for autoregulation of histone expression levels).

An alternative mechanism to explain the high level of sequence identity among these genes, as well as between the 614/615 pairs and the H3-616 gene and between the two H2a-613 genes, is gene conversion. Clearly, some histone genes on chromosome 13 have been involved in gene conversion (Liu et al. 1987; Brown et al. 1996; Wang et al. 1996), and much of the difference between the chromosome 3 H3 genes compared to the chromosome 13 H3 genes appears to be caused by gene conversion among the chromosome 13 genes (Debry and Marzluff 1994). Differences in the promoter region between the H3-616 gene compared to the H3-614/H3-615 pair strongly suggest that gene conversion of the coding region, and not a recent duplication, is
responsible for the extreme similarity among these three genes. These data agree with the conclusion that gene conversion is very rare between the histone genes on chromosomes 3 and 13 (Debry and Marzluff 1994).

The two H2a-613 genes also show an interesting pattern that is likely attributable to gene conversion. Within the first five codons, the H2a(A)-613 and H2a(B)-613 genes share three silent differences compared to H2a-614. For most of the remaining sequence, however, the H2a(A)-613 gene is virtually identical to H2a-614, with only three silent differences over 113 contiguous codons. Over that same stretch, the H2a(B)-613 gene has a total of 34 differences, including 2 replacement differences. The implication is that the H2a-614 gene may have recently converted the H2a(A)-613 gene over most of the coding sequence.

In contrast, the two H2b genes on chromosome 3 differ greatly at the nucleotide sequence level, although they encode quite similar proteins, suggesting that these two genes have not been involved in a recent gene conversion event. It is certainly not clear whether all of the histone genes are essential, and in particular, whether the different minor protein variants encoded by the H2a and H2b genes are physiologically important.

Although we cannot unequivocally rule out the presence of additional histone genes on mouse chromosome 3 located farther than 25 kb from any of these genes, in situ hybridization experiments with coding region probes clearly show that most of the core histone genes are present on chromosome 13 (W.F. Marzluff and J. Hozier, unpubl.). There are about nine copies of each of the core histone genes on chromosome 13 (Wang et al. 1996), and the additional two to four copies present on chromosome 3 probably account for the complete complement of the replication-dependent mouse histone genes.

METHODS

Isolation of P1 and Phage λ Clones

A series of overlapping P1 clones (P0, P1, P2, and P3) was isolated by Genome Systems (St. Louis, MO) by PCR with a set of primers flanking the H3-614 gene. After ordering these P1 clones, the end of clone P0 was sequenced and a set of primers were designed to isolate a second set of P1 clones (P91 and P92). Because both clones ended in the same region, a probe was synthesized by use of T7 RNA polymerase from the end of clone P92, which extended the furthest from one end of the histone cluster, and this probe was used to screen a mouse genomic library in λFII. A single phage that extended this cluster an additional 12 kb was isolated in this screen.

Mapping of the P1 Clones

Probes to the ends of each of the P1 clones were synthesized by use of SP6 or T7 RNA polymerase and these probes were used to orient and order the clones by Southern blotting. The size of the inserts was estimated after digestion with NotI, which cuts at the junction of the vector and the insert. There were no NotI sites in any of the inserts. The position of each gene was determined by Southern blotting with probes from each of the core histone genes. Specific probes from the 3'-flanking region of the H3-614 and H3-615 genes, and the 5'-flanking region of the H2a-614 and H2a-615 genes were used in Southern blots to determine the orientation of these two duplicated genes.

In Situ Hybridization

Mouse metaphase spreads were prepared from cultured spleenocytes by use of a standard methanol/acetic acid fixation protocol. The following clones were used for FISH: P0, P3, P91, P92, and λ111. Mapping was done as described in Wang et al. (this issue).

DNA Sequencing

The genes were subcloned in plasmid vectors and the sequences determined in the UNC DNA Sequencing Facility.

SI Nuclease Mapping

Total RNA was prepared from cultured mouse myeloma cells or from neonatal mice (1 day old) and the transcripts detected by S1 nuclease mapping essentially as described (Graves et al. 1985; Wang et al., this issue).

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