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Long-Range Comparison of Human and Mouse *SCL* Loci: Localized Regions of Sensitivity to Restriction Endonucleases Correspond Precisely with Peaks of Conserved Noncoding Sequences

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Long-range comparative sequence analysis provides a powerful strategy for identifying conserved regulatory elements. The stem cell leukemia (*SCL*) gene encodes a bHLH transcription factor with a pivotal role in hemopoiesis and vasculogenesis, and it displays a highly conserved expression pattern. We present here a detailed sequence comparison of 193 kb of the human *SCL* locus to 234 kb of the mouse *SCL* locus. Four new genes have been identified together with an ancient mitochondrial insertion in the human locus. The *SCL* gene is flanked upstream by the *SIL* gene and downstream by the *MAP17* gene in both species, but the gene order is not collinear downstream from *MAP17*. To facilitate rapid identification of candidate regulatory elements, we have developed a new sequence analysis tool (SynPlot) that automates the graphical display of large-scale sequence alignments. Unlike existing programs, SynPlot can display the locus features of more than one sequence, thereby indicating the position of homology peaks relative to the structure of all sequences in the alignment. In addition, high-resolution analysis of the chromatin structure of the mouse *SCL* gene permitted the accurate positioning of localized zones accessible to restriction endonucleases. Zones known to be associated with functional regulatory regions were found to correspond precisely with peaks of human/mouse homology, thus demonstrating that long-range human/mouse sequence comparisons allow accurate prediction of the extent of accessible DNA associated with active regulatory regions.

One of the major challenges currently facing biological science concerns the characterization of transcriptional networks in higher organisms. The regulatory information must be present in the primary DNA sequence, but deciphering the code remains a formidable challenge. Long-range comparative sequence analysis provides an attractive strategy for the identification of functionally important gene regulatory regions, on the basis that their sequences are highly conserved during evolution (Hardison et al. 1997). Early studies used functional assays to identify homologous regulatory elements and subsequently used local sequence comparisons to identify conserved transcription factor binding sites (Aparicio et al. 1995; Popperl et al. 1995; Nonchev et al. 1996). More recently, the increasing availability of large tracts of genomic sequence allows a shift to long-range comparisons, and it has been suggested that the identification of regulatory elements through human/mouse sequence comparisons is suffi-

cient justification for sequencing the entire mouse genome (Hardison et al. 1997). However, only a limited number of long-range comparisons have been reported so far. Human/mouse comparisons were shown to be an efficient approach for the reliable prediction of coding exons (Ansari-Lari et al. 1998; Endrizzi et al. 1999; Jang et al. 1999). Besides, analysis of the human and murine *Bruton's tyrosine kinase* loci revealed a new enhancer with activity in transfection assays (Oeltjen et al. 1997). Comparison of human and murine β -globin loci showed that peaks of high homology corresponded with functional regulatory regions, including the well-characterized DNaseI hypersensitive sites within the locus control region (Jackson et al. 1996; Hardison et al. 1997). Comparative sequence analysis of the human and mouse *adenosine deaminase* genes showed that several known regulatory regions displayed higher sequence conservation than some of the coding exons (Brickner et al. 1999). Most recently, long-range sequence comparisons of the human and mouse *IL4/IL13/IL5* gene clusters led to the identification of a putative chromatin regulatory region, the activity of which was assayed in vivo using transgenic mice (Loots et al. 2000).

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The *SCL* gene encodes a bHLH transcription factor with a critical role in hemopoiesis and vasculogenesis. It was identified by virtue of its disruption in T-cell acute leukemia, and rearrangements of the *SCL* locus are perhaps the most frequent molecular pathology associated with this tumor (Barton et al. 1999; Begley and Green 1999). Targeted mutation of the *SCL* gene has shown that it is essential for the development of all hemopoietic lineages (Porcher et al. 1996; Robb et al. 1996) and also for normal yolk sac angiogenesis (Visvader et al. 1998). Ectopic *SCL* expression in zebrafish embryos specifies hemangioblast development from early mesoderm, results in disproportionate production of blood and endothelial progenitors, and can partially rescue endothelial and hemopoietic phenotypes of the *cloche* mutant (Gering et al. 1998; Liao et al. 1998).

SCL is normally expressed in hemopoietic cells, endothelium, and within specific regions of the CNS. This pattern of expression is highly conserved throughout vertebrate evolution from mammals to teleost fish (Green et al. 1992; Kallianpur et al. 1994; Gering et al. 1998; Mead et al. 1998; Sinclair et al. 1999; Drake and Fleming 2000). *SCL* expression is tightly regulated and involves two alternative promoters with lineage-specific activity in distinct hemopoietic cell types (Lecointe et al. 1994; Bockamp et al. 1995, 1997, 1998). In addition, a detailed analysis of the chromatin structure of the mouse *SCL* locus identified a number of DNaseI hypersensitive sites associated with enhancer or silencer activity (Göttgens et al. 1997). More recently, studies using transgenic mice have identified five separate enhancers, which direct reporter gene expression in vivo to endothelium, midbrain, hindbrain/spinal cord, or hemopoietic progenitor cells, all subdomains of the normal *SCL* expression pattern (Sanchez et al. 1999; Sinclair et al. 1999; Göttgens et al. 2000).

We have recently cloned and sequenced the *SCL*

locus from human, mouse, and chicken (Göttgens et al. 2000), but only limited sequence was available from the region downstream of the mouse *SCL* locus. To permit long-range human/mouse sequence comparisons, we have isolated and sequenced an additional 148 kb from the 3' region of the mouse *SCL* locus. The structures of the human and mouse *SCL* loci are very similar in the immediate vicinity of the *SCL* gene, but substantial differences are present downstream of the *SCL* flanking gene, *MAP17*. We have developed a new sequence analysis tool (SynPlot) to allow rapid identification of candidate regulatory regions. Finally, we show that for known regulatory regions, the extent of homology within an individual peak corresponds precisely with sites sensitive to restriction endonuclease digestion.

RESULTS

Structure of Human and Mouse *SCL* Loci

Our previous analysis of the human *SCL* locus showed that human *SCL* was flanked upstream by the *SIL* gene and downstream by the *MAP17* gene (Göttgens et al. 2000). However, the relative position of the original human and mouse genomic clones restricted the overlap between the human and mouse *SCL* loci to only 55.8 kb with only 11.0 kb of 3' flanking sequence, which did not extend to the 3' flanking gene. A new mouse *SCL* genomic clone was therefore isolated and completely sequenced (see Methods). This allowed, for the first time, a complete comparative analysis of 193 kb of the human and 234 kb of the mouse *SCL* loci, which extended to the 5' and 3' flanking genes in both species (Fig. 1).

Complete annotation of the two sequences led to the description of three new murine genes: mouse *MAP17* and two members of the cytochrome p450

Cyp4 family of genes. In addition, examination of the human *SCL* sequence downstream of *CYP4A11* revealed the 3' half of another member of the *CYP4* gene family, the as-yet-unpublished *CYP4Z1* gene (D. Bell, pers. comm.; Fig. 1). Comparison of the human and mouse loci showed that the order of the *CYP4* genes was not conserved, even though the *SCL* locus on human chromosome 1 forms part of a long region of synteny with mouse chromosome 4. The *CYP4* family of cytochrome p450 genes is divided into subfamilies based on sequence homology (i.e., *CYP4A*, *CYP4B*, etc.). The two new mouse genes have been allocated to subfamilies and named *Cyp4x1* and *Cyp4a21* (see Cytochrome p450 Nomenclature Committee at <http://drnelson.utmem.edu/CytochromeP450.html>). Surpris-

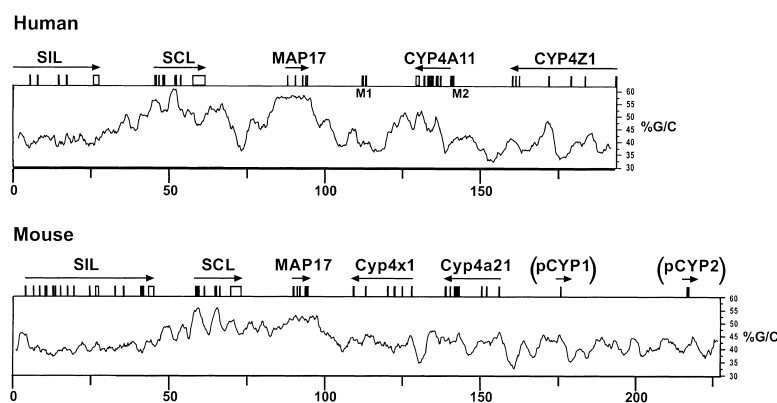


Figure 1 Structure of the human and murine *SCL* loci. The gene structure of the human and mouse *SCL* loci is shown above a profile displaying the respective G/C content. Arrows indicate the direction of genes. M1 and M2 refer to the sequences homologous to mitochondrial DNA, and (pCYP1) and (pCYP2) refer to partial segments of *CYP4* genes present in the mouse locus.

ingly, the 5' human gene, *CYP4A11*, is more similar to the 3' mouse gene *Cyp4a21* than to the 5' mouse gene *Cyp4x1*. Moreover, a human ortholog to mouse *Cyp4x1* has recently been mapped to the same region of human chromosome 1 and presumably lies further into the *CYP4* gene cluster (D. Bell, pers. comm.). These results show that the order of individual genes within the *CYP4* locus is not conserved in man and mouse. Consistent with this observation, the number of genes within each *CYP4* subfamily differs between mammals (see <http://drnelson.utmem.edu/CytochromeP450.html>), indicating that the *CYP4* locus is evolving by gene duplication. This view is further supported by the presence in 3' of mouse *Cyp4a21* of two regions homologous with the last exon and exons 4–5 of *CYP4* family members. These two regions (pCyp1 and pCyp2, respectively in Fig. 1) were both in the opposite transcriptional orientation to *Cyp4a21*, and are likely to represent remnants of *Cyp4* genes resulting from partial gene duplications/inversions. Therefore, our analysis showed that the gene structure and order of the *SIL*, *SCL*, and *MAP17* genes was highly conserved, whereas 3' of *MAP17* of the human and mouse loci were not colinear.

The G/C content of the human and mouse sequences was 44.5% and 43%, respectively, which is typical of the relatively gene-poor isochore H1 (Bernardi 2000), and consistent with the location of the human *SCL* gene within a Giemsa light band. Neither the human nor the murine G/C profile followed the regular sinusoidal pattern described for the human α -globin locus (Flint et al. 1997), but the *SCL* and *MAP17* genes were in regions of high G/C content in both species. CpG islands were found to be associated with the promoters of *SIL* and *SCL*, but not the *MAP17* or *CYP4* genes (data not shown). Indeed, the CpG content of a 50-kb segment containing the *SCL* and *MAP17* genes is so high, that it would be part of the gene-rich isochore H2 (Bernardi 2000). This finding contrasts with the suggestion that isochores are units greater than 200 kb in size and characterized by a high compositional homogeneity (above a 3 kb size level) (Bernardi 2000).

A Mitochondrial Genome Insertion in the Human but Not the Mouse Locus

Sequence database searches with the human *SCL* locus sequence revealed the presence of two regions on either side of the *CYP4A11* gene that had high homology with segments of the mammalian mitochondrial genome (Fig. 2). The first region (112028–112388 and 113106–113425) matches the human mitochondrial genome from 2932 to 3651. The match is flanked by Alu and LTR repeats at the 5' and 3' ends, respectively, and the gap interrupting the match contains two partial Alu repeats. The second region (140989–141045

and 141372–141429) matches the human mitochondrial genome from 12213 to 12327, and was flanked by LINE repeats (L2 and L1PA16 repeats, respectively). No similarity to these or any other regions of vertebrate mitochondrial genomes were detected in the sequence of the mouse *SCL* locus.

The serial endosymbiosis theory postulates that symbiotic organelles such as mitochondria gradually transferred a large proportion of their genes to the eukaryotic/nuclear genome during early evolution. Remnants of more recent transposition events of mitochondrial DNA into the nuclear genome are also detectable by Southern hybridization (du Buy and Riley 1967; Tsuzuki et al. 1983). However, this is the first report describing the sequence of such an insertion. The two segments of nuclear-mitochondrial DNA described here were codirectional, consistent with the suggestion that they represent the remnants of a single integration event. Two observations indicate the relatively ancient nature of this mitochondrial-to-nuclear transposition. First, the transposed mitochondrial DNA has been disrupted through the integration of repeat elements and even an entire gene (*CYP4A11*). Second, the nuclear sequence is no more similar to human than to other mammalian mitochondrial sequences, and it is the most divergent sequence when aligned to mammalian mitochondrial DNA (Fig. 2B; data not shown). BLAST scores were higher for alignments with horse, cow, and sheep mitochondrial DNA (BLASTX scores ~180) than for alignments with human or mouse mitochondrial DNA (BLASTX scores ~160). Taken together, these data indicate that the transposition event occurred before the divergence of human and horse, cow or sheep.

Sequence Variation of Mouse Strains

The sequence of the murine *SCL* locus used in this study was assembled from P1 clone ICRFP703C1281Q and PAC clone PAC129.0.726 from the genetic backgrounds C57/Bl6 and 129SV/SvEvTac, respectively. We analyzed the overlap of 23.8 kb of the two insert sequences to assess the frequency of sequence length and single nucleotide polymorphisms (SNPs) between these two important laboratory strains. We found 18 sequence length polymorphisms, 10 of which were just one nucleotide, with the remaining eight covering 33 bp in total. The number of C \leftrightarrow T transitions (11) exceeded the number of transversions (6) consistent with findings from human SNP analysis. However, all six transversions were of the G \leftrightarrow T type, with no occurrence of the G \leftrightarrow C or A \leftrightarrow T alternatives, whereas the three possible transversions occur at similar frequencies in human (Brookes 1999). Therefore, it will be interesting to see if other regions of the mouse genome show a preference for the G \leftrightarrow T transversion. The total number of 17 SNPs in 23.8 kb is in good agreement

not possible to display the position of homology peaks within the test sequence relative to other sequence features (exons, repeats, other homology peaks, etc).

Therefore, we developed a new software tool (SynPlot), which uses the Dialign algorithm (Morgenstern et al. 1998) for computation of a long-range global alignment. Dialign scans sequences of unlimited length for areas of high local similarity and uses these as anchor points for a global alignment. The SynPlot graphical output includes a similarity profile of the long-range alignment together with a diagrammatic representation of both loci, including the position and size of all gaps inserted to permit optimum global alignment. Consequently, the positions of all features in the individual sequences have to be transformed into their new positions within the aligned file. This transformation is performed automatically by SynPlot, which calculates the new positions using the locations and lengths of all gaps. Moreover, feature files generated during annotation in ACeDB (Durbin and Mieg 1991), and that contain the positions of exons and repeat elements, can be directly imported into the graphical output. Therefore, the SynPlot output conveys comparative gene structure, repeat patterns (plus any other user-defined patterns), and relative sequence homology in a single linear plot with the upper limit being the length of the region of synteny.

The SynPlot output file (Fig. 3) displays a comparative analysis of the human and mouse *SCL* loci from within the *SIL* gene to beyond the *CYP4A11* and *Cyp4a21* genes, respectively. Although the latter two genes may not be orthologs (because there are three more murine members of the *Cyp4a* subfamily, *Cyp4a10/12/14*), their subfamily relationship is recognized by Dialign. As a result, the two genes are aligned and long gaps introduced in the human sequence to accommodate the region occupied by mouse *Cyp4x1*. Display of repeat elements in the profile shows that most other gaps inserted for optimum sequence alignment coincide with species-specific repeat elements in either locus.

All coding exons clearly stand out as homology peaks. Noncoding homology peaks are seen only in the region of the *SIL*, *SCL*, and *MAP17* genes. These peaks represent candidate regulatory regions, or may serve other conserved aspects of genome function such as replication or domain structure. The distinct absence of noncoding homology peaks in the *CYP4* gene region is consistent with our suggestion that the genes present in human and mouse are not orthologous.

Local Regions of Sensitivity to Restriction Endonucleases Correspond to Peaks of Human/Mouse Homology

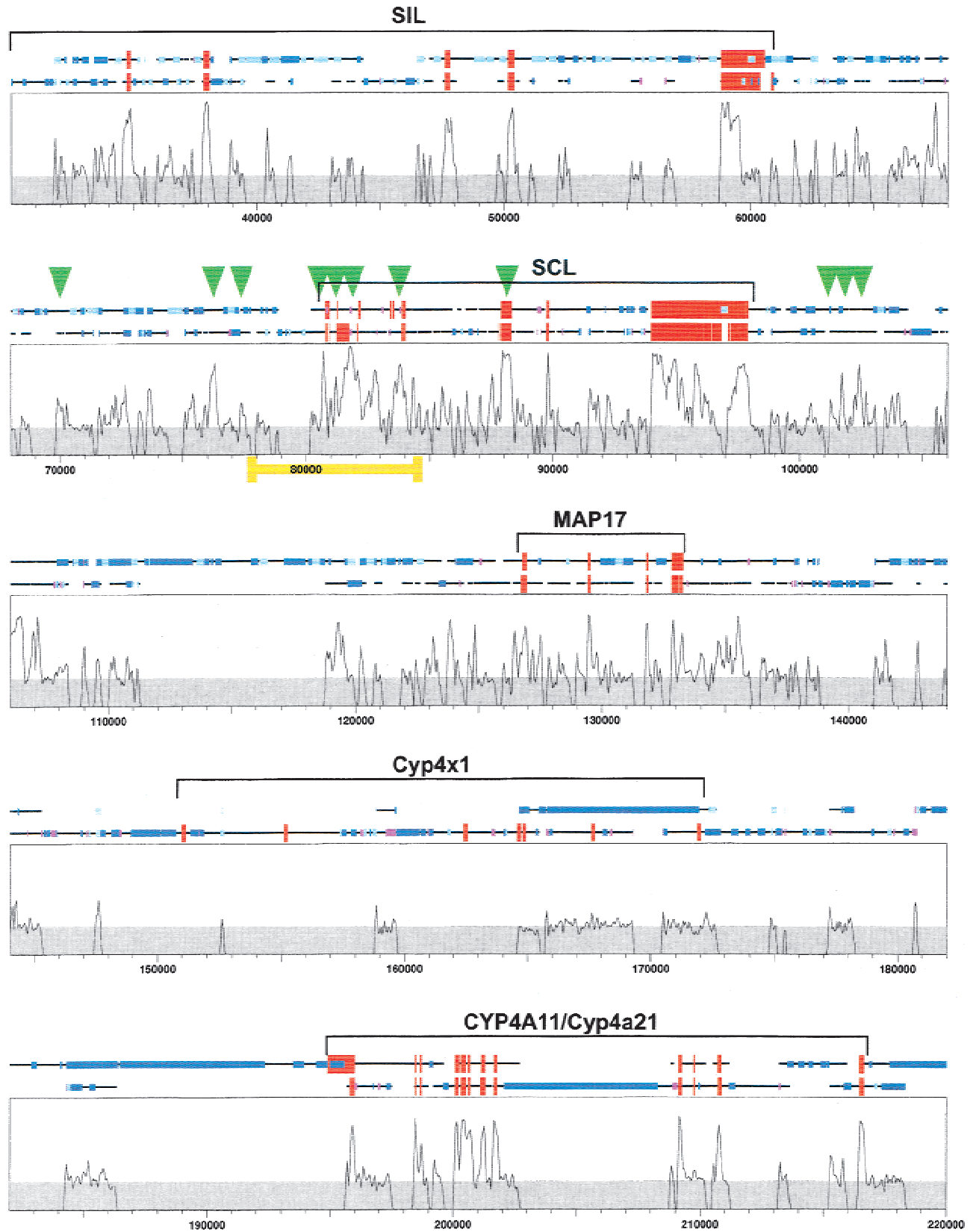
Many active enhancers are associated with regions of hypersensitivity to DNaseI digestion. Therefore, analy-

sis of chromatin structure is frequently used to identify the location of candidate regulatory regions. However, this strategy is constrained by the fact that the activity of individual enhancers may be restricted to rare cell types, thus precluding biochemical analysis. Furthermore the resolution of Southern blotting is limited. Therefore, we investigated whether the peaks of human/mouse homology can be used to predict the position and extent of localized zones of altered chromatin structure associated with regulatory regions of the murine *SCL* gene.

A systematic analysis of the chromatin structure of the mouse *SCL* locus previously identified a number of DNaseI hypersensitive sites associated with enhancer or silencer activity (Göttgens et al. 1997). However, DNaseI hypersensitive site mapping has a limited resolution and the precise location or extent of each DNaseI hypersensitive site was not known. To compare the extent of accessible DNA in each case with the human/mouse alignment, a restriction endonuclease accessibility assay (Boyes and Felsenfeld 1996) was modified to pinpoint the location of sites sensitive to restriction endonuclease digestion within the DNA sequence.

Restriction endonuclease accessibility studies were performed in two primitive myeloid cell lines (M1 and 416B) known to express the *SCL* gene (Fig. 4). Our results show that seven previously described DNaseI hypersensitive sites (promoter 1a, promoter 1b, +1, +3, Fig. 4; +7, -3, -4, data not shown; numbering represents the distance [kb] from the start of murine exon 1a) were associated with localized zones of increased endonuclease accessibility. All of these DNaseI hypersensitive sites have been shown to be associated with enhancer or silencer activity using transfection or transgenic reporter assays (Göttgens et al. 1997; Fordham et al. 1999; Sanchez et al. 1999; Sinclair et al. 1999). The use of multiple restriction endonucleases allowed the extent of the regions of endonuclease accessibility to be mapped in detail. In each case, these regions correlated precisely with a peak of homology in the human/mouse alignment (Fig. 4B), thus identifying these peaks as candidate regulatory regions.

We have previously identified 11 DNaseI hypersensitive sites in a ~40 kb portion of the mouse *SCL* locus (see arrowheads in Fig. 3) (Göttgens et al. 1997). All of these corresponded to regions of endonuclease accessibility and peaks in the homology profile. These 11 sites represent approximately one-third of all geographically distinct regions of high sequence similarity. A further four peak regions correspond to coding exons and the sequence surrounding the polyadenylation site. The significance of the remaining peaks of homology remains unclear. However, it is likely that at least some of these peaks will prove to mark the sites of enhancers associated with DNaseI hypersensitive sites in tissues in which the chromatin structure of the *SCL*



(See following page for legend.)

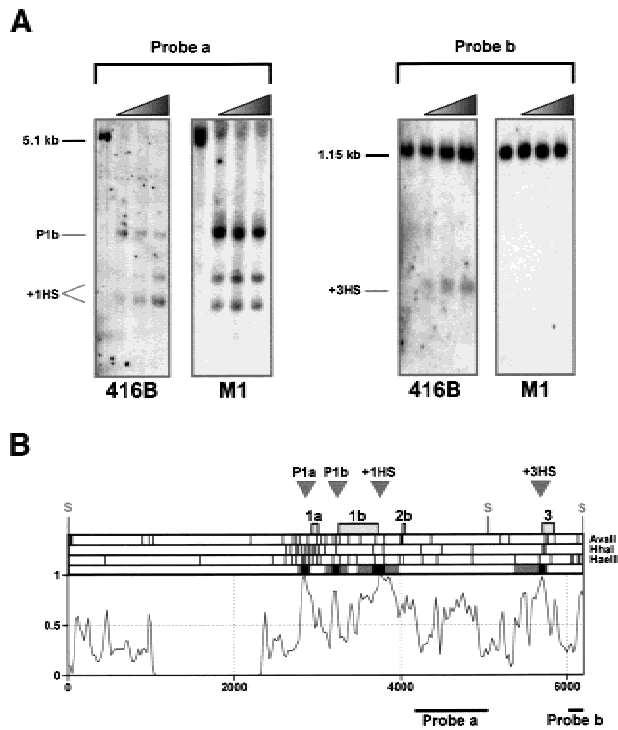


Figure 4 Localized regions of sensitivity to restriction endonucleases correspond to peaks of human/mouse homology. (A) Restriction endonuclease accessibility assay showing the mapping of the hypersensitive sites at promoter 1b, +1 (+1HS), and +3 (+3HS) (numbering corresponds to distances in kb from the start of exon 1a) in the 416B and M1 primitive myeloid cell lines. Nuclei were incubated with *HaeIII*. DNA was subsequently extracted, digested with *SacI*, and hybridized with probes indicated in B. The absence of the +3 hypersensitive site in M1 is consistent with previous DNaseI hypersensitive site analysis (Göttgens et al. 1997). (B) Summary of restriction endonuclease data for the 5' region of the mouse *SCL* gene. The top part of the diagram shows the approximate locations of previously mapped DNaseI hypersensitive sites (gray arrowheads labeled P1a, P1b, +1HS, and +3HS) followed by the positions of mouse *SCL* exons 1a to 3 and the *SacI* sites used for the Southern blots shown in part A. This is followed by restriction maps for the three enzymes (*Avall*, *Hhal*, and *HaeIII*) used to determine endonuclease sensitivity, and a summary of the endonuclease sensitivity experiments in which black and gray boxes represent the minimum and maximum regions of endonuclease accessibility in 416B and/or M1 cells. The profile of the mouse/human alignment underneath is a 6250 nucleotide section of the alignment from Fig. 3 (see yellow bar in Fig. 3) and shows the concordance of endonuclease accessibility and sequence conservation.

locus has not yet been characterized. Alternatively, they may mark the sites of regulatory elements not

linked with DNaseI hypersensitive sites, but which control other conserved processes such as domain structure, chromatin architecture, or DNA replication.

DISCUSSION

In this paper, we present a detailed comparison of 193 kb of the human to 234 kb of the mouse *SCL* locus. Four new genes and an ancient mitochondrial transposition have been identified, and a new program to support long-range comparative analysis has been developed. In addition, our results show for the first time that not only do long-range human/mouse sequence comparisons help locate chromatin sites associated with regulatory regions, but also that peaks of human/mouse homology accurately predict the extent of such regions of accessible DNA.

Correlation of Sequence Homology and Sites of Endonuclease Accessibility

A major objective of genome sequence analysis is to identify transcriptional regulatory elements in DNA sequences, and from this information to understand transcriptional networks in higher organisms. The *SCL* gene encodes a critical transcriptional regulator of hemopoiesis and vasculogenesis, and the molecular basis for the tissue-specific regulation of the murine *SCL* gene has been characterized in considerable detail (Göttgens et al. 1997; Sanchez et al. 1999; Sinclair et al. 1999; Göttgens et al. 2000). Here we describe detailed sequence comparisons of the human and mouse *SCL* loci, which together with a comprehensive chromatin structure analysis, have allowed us to draw several important conclusions.

First, all DNaseI hypersensitive sites linked to known *SCL* regulatory regions were associated with localized zones of restriction endonuclease accessibility that coincided with regions of high human/mouse sequence homology. It is not clear whether both DNaseI and restriction endonucleases reveal the same structural features, and the question remains regarding the pattern of endonuclease sensitivity that might be observed in a cell that does not express *SCL*. However, our data are consistent with a model in which both assays detect aspects of a localized region of an altered chromatin structure, which is associated with active *SCL* regulatory regions. There are several possible explanations for the peaks of homology that do not corre-

Figure 3 SynPlot analysis of the human and mouse *SCL* loci. Human and mouse clones starting with the last five exons of *SIL*, and ranging to beyond the *CYP4A11/Cyp4a21* genes, were aligned using Dialign. The alignment, together with locus features, was displayed using SynPlot. Numbers on the horizontal axis represent distance (nucleotides) from the beginning of the aligned file. Numbers on the vertical axis represent the proportion of identical nucleotides within a 49 nt window, moved by 25 nt increments across the entire alignment. Hence, regions with gaps of >50 bp show 0% identity. The horizontal lines above the profile represent the human and mouse sequences and illustrate the position of gaps introduced to permit optimum alignment. Red boxes show exon positions, and the smaller boxes represent repeats as follows: (dark blue) LINES, (light blue) SINEs, (magenta) tandem repeats. Green arrowheads indicate the positions of previously mapped DNaseI hypersensitive sites, and the yellow bar delimits the portion of the profile shown in Figure 4. Gray shading indicates background similarity of $\leq 25\%$.

spond to zones of DNaseI hypersensitivity or endonuclease accessibility. The chromatin studies have only been performed in a limited number of hemopoietic lineages and additional localized regions of altered chromatin structure may exist in other *SCL*-expressing cell types. Alternatively, some categories of enhancers may not be associated with such regions, as may other classes of regulatory regions that may have important roles in conserved processes such as gene silencing, domain structure, and chromosome architecture.

Second, high-resolution analysis of the chromatin structure of *SCL* regulatory regions showed that the extent of each individual homology peak corresponded precisely with the region accessible to endonucleases. This observation indicates that human/mouse sequence comparisons will greatly reduce the experimental studies required to define minimal functional units associated with enhancers. Moreover, incorporation of more distant species into the comparisons permits the generation of phylogenetic footprints (Aparicio et al. 1995; Popperl et al. 1995; Nonchev et al. 1996), which can identify conserved transcription factor binding sites, and thus provide rapid insight into the detailed architecture of an enhancer.

Third, our results confirm the utility of comparative sequence analysis as a way of identifying regulatory regions within the reams of sequence information generated by genome sequencing projects. The combination of comparative sequence analysis with a rapid and high throughput transgenic *Xenopus* assay (Kroll and Amaya 1996) provides a powerful strategy, particularly if focused on critical developmental genes with functions and expression patterns conserved throughout vertebrate evolution (Göttgens et al. 2000).

Interpretation of Comparative Sequence Analysis

Widespread application of long-range comparative sequence analysis will require the use of intuitive computational tools. At present, the most widely used approach is that of percentage identity plots (PIPs) generated by the PIPMaker program (Schwartz et al. 2000). These do not require a global alignment, but instead are based on local alignments and display areas of high local similarity between a reference and the test sequence, with the advantage that PIPs can display matches if the two sequences are not colinear. However, PIPs can display only the features (exons, repeats, etc.) of one of the two loci (i.e., the reference sequence), and the relative position of high-scoring local alignments within the test sequence cannot be displayed. A comparison of human, sheep, and mouse PrP loci (Lee et al. 1998) has provided an example of how locus features can be displayed in combination with a sequence similarity profile. However, this approach involves multiple steps, including the removal and reinsertion of species-specific repeats. Moreover, neither

the alignment program nor the display tools have been published or been made available on the Internet.

Therefore, we have developed a new software tool (SynPlot), which is based on a global alignment and has several novel features. First, the nature and positions of features, such as exons, repeat elements, or CpG islands, can be shown for both sequences, thereby indicating the position of homology peaks relative to the gene structure of both loci. Second, SynPlot allows large-scale sequence alignments of multiple (i.e., >2) sequences to be presented in a single graphical display. Third, the strategy used by SynPlot to transform the coordinates of sequence features to their new positions in the global alignment can be incorporated into future sequence analysis applications. These coordinates are read into SynPlot from a file in General Feature Format or GFF (see <http://www.sanger.ac.uk/Software/formats/GFF/>), which will facilitate the display of sequence features from other analysis programs and databases (e.g., the ENSEMBL project; see <http://www.ensembl.org/>).

The Dialign algorithm (Morgenstern et al. 1998) used in this manuscript does not employ gap penalties, but instead collects small gap-free alignments, which are weighted depending on their percent identity and length. The global alignment is constructed by assembling these local alignments in an optimized colinear configuration. Whereas this approach overcomes many problems of generating global alignments, its utility is restricted to colinear loci. Importantly, however, SynPlot and PIPMaker produce similar results on such loci when used for the identification of conserved noncoding sequences (unpubl.) even though they are based on very different principles.

When performing comparative analysis, it is our practice to start by assessing collinearity, e.g., by dotter analysis (Sonnhammer and Durbin 1995). "Working draft" sequence is noncontiguous, and therefore not suitable for long-range global alignments. Consequently, in the absence of colinearity and for working draft sequence, PIPs remain the preferred approach. However, if colinearity is shown, SynPlot analysis is performed to take advantage of its ability to present the position of homology peaks relative to other features in all of the loci being compared, together with its ability to display comparisons between multiple sequences.

Comparative Analysis of Evolutionarily Active Loci

To date, long-range human/mouse comparative sequence analyses have focused on loci conserved during mammalian evolution. The *SCL* locus on human chromosome 1 is part of a long region of synteny with mouse chromosome 4. However, our results show that, within this long region, the *CYP4* loci have evolved substantially because the divergence of man and

mouse and are not colinear. The cytochrome P450 monooxygenases (*CYPs*) are a large and ancient superfamily of proteins, often involved in the metabolism of hormones or foreign compounds such as toxins or drugs. Therefore, the acquisition of different sets of *CYP* genes may partially explain the different sensitivity of humans and mice to drugs and carcinogens.

Rapidly evolving loci, such as the human and mouse *CYP4* clusters, highlight two challenges for comparative analysis. First, unless the individual sequences are annotated very carefully, incorrect homology relationships between human and mouse genes may be inferred. Second, the usefulness of comparative analysis for the prediction of regulatory regions may be severely impaired. Such predictions are based on common ancestry, and true orthologs may be difficult to establish. Moreover, rapid evolution may be accomplished by the acquisition of distinct expression patterns by individual members of a gene cluster.

Interpreting the Genome Code

Genome projects are producing a vast data resource which will have to be made accessible to the scientific community. It is clear that, in addition to the primary DNA sequence, genome-wide data on gene expression patterns, protein structures, and protein-protein interactions will also be available in the near future. Our own interest lies in how gene regulatory networks are encoded in the primary DNA sequence, and we use comparative sequence analysis to address this interest. The data presented here indicate that a very careful analysis of individual loci is necessary to establish true homology relationships before proceeding with the comparative analysis. We also show that human/mouse comparisons can give insights into recent genome evolution. Most importantly, we show that comparative sequence analysis, combined with an intuitive graphical display, will be very useful for the prediction of the location and size of gene regulatory regions.

METHODS

Isolation and Sequencing of 129/SvEvTac Murine *SCL* Locus

The RPCI-21 (129S6/SvEvTac) mouse PAC library (segment 2) (Osoegawa et al. 2000) was screened with a probe from exon 6 of the mouse *SCL* gene. Eight positive clones were identified and further analyzed by PFGE and Southern analysis. One clone (PAC129.0.726) did not hybridize to a probe from the 5' end of the mouse *SCL* gene and contained a large insert (>150 kb), thereby maximizing the extension of the known mouse *SCL* sequence. The insert of clone PAC129.0.726 was fully sequenced as described (Göttgens et al. 1998). The insert was found to be 171,863 bp, and it extended the previously known mouse *SCL* sequence by 148,000 bp. The human and mouse sequences used in this study, together with their respective feature files, are available at <http://www.sanger.ac.uk/~jgrg/SynPlot>.

Sequence Analysis

Interactive annotation of genomic sequences was performed within ACeDB (Eeckman and Durbin 1995) as described (Göttgens et al. 1998). Global alignments were calculated using Dialign version 1 (Morgenstern et al. 1998) with the threshold for diagonals (T) set to 20. We wrote an application, SynPlot (see <http://www.sanger.ac.uk/~jgrg/SynPlot>), to visualize global alignments of syntenic genomic sequence. SynPlot is a set of Perl modules and a driver script and is freely available from the SynPlot Web site. It takes as its input aligned sequences in fasta format, with gaps in the sequences introduced by the alignment represented by '-' characters. We have also written the necessary modules to feed the output of the recently published GLASS global sequence alignment algorithm (Batzoglu et al. 2000) into SynPlot (see <http://www.sanger.ac.uk/~jgrg/SynPlot>). The percentage identity is calculated from the alignment within a sliding window, the width of which can be specified by the user. This information is used to draw a picture of the alignment in postscript format. The sequences are rendered as lines interrupted by spaces corresponding to the gaps introduced by the alignment, with a plot of the percentage identity underneath. Features can also be drawn on the sequence lines. This uses a GFF format file (see <http://www.sanger.ac.uk/Software/formats/GFF>) output by ACeDB from the annotated genomic sequence, and a configuration file, which specifies the color, height, and order in which the rectangles representing the features are drawn. Small-scale DNA alignments were performed and displayed as described (Göttgens et al. 1998). G/C profiles were calculated and displayed using the GCG sequence analysis package (Wisconsin package version 8.0, Genetics Computer Group) using a window size of 2000 bp with an incremental move of 320 bp. BLASTX scores for mitochondrial sequences were calculated using the vertebrate mitochondrial genetic code.

Restriction Endonuclease Accessibility Assay

M1 and 416B cells were maintained as described previously (Bockamp et al. 1997). For the restriction endonuclease assay, 2×10^8 cells were washed twice in PBS before resuspending in 8 mL of lysis buffer (50 mM KCl, 10 mM MgSO₄, 3 mM DTT, 5 mM HEPES at pH 7.2, 0.05% NP-40, and 1 mM PMSF), followed by a 60 min incubation at room temperature. Nuclei were collected by 5 min centrifugation at 5000 g, washed in RSB (10 mM NaCl, 10 mM Tris at pH 7.4, 3 mM MgCl₂), and resuspended in 10 mL restriction enzyme buffer 2 (New England Biolabs, Hitchin, UK). Four hundred μ L aliquots were digested with 0, 80, 160, or 320 units of *Pall* (*Hae*III), *Ava*II, or *Hha*I for 60 min at 37°C, followed by conventional DNA isolation and Southern blot analysis using *Sac*I digestion and the following probes: a = 900 bp *Sma*I/*Sac*I fragment; b = 180 bp *Sma*I/*Sac*I fragment (Fig. 4B).

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