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Identification of the homologous beige and Chediak–Higashi syndrome genes

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Abstract

Vesicular transport to and from the lysosome and late endosome is defective in patients with Chediak–Higashi syndrome (CHS) and in mutant beige (*bg*) mice^{1–4}. CHS and *bg* cells have giant, perinuclear vesicles with characteristics of late endosomes and lysosomes that arise from dysregulated homotypic fusion^{3–5}. CHS and *bg* lysosomes also exhibit compartmental missorting of proteins, such as elastase, glucuronidase and cathepsin G^{2,3,6,7}. *Lyst*, a candidate gene for *bg*, was identified by direct complementary DNA selection from a yeast artificial chromosome (YAC) clone containing a 650-kilobase segment of the *bg*-critical region on mouse chromosome 13. *Lyst* is disrupted by a 5-kilobase deletion in *bg*^{11J} mice, and *Lyst* messenger RNA is markedly reduced in *bg*^{2J} homozygotes. The homologous human gene, *LYST*, is highly conserved with mouse *Lyst*, and contains a frame-shift mutation at nucleotides 117–118 of the coding domain in a CHS patient. Thus *bg* mice and human CHS patients have homologous disorders associated with *Lyst* mutations. *Lyst* encodes a protein with a carboxy-terminal prenylation motif and multiple potential phosphorylation sites. *Lyst* protein is predicted to form extended helical domains, and has a region of sequence similar to stathmin, a coiled-coil phosphoprotein thought to act as a relay integrating cellular signal response coupling^{8–10}.

We have previously localized the *bg* locus within a 0.24-centimorgan interval on mouse chromosome 13, and isolated contiguous arrays of YACs that cover 2,400 kb of this interval^{11–13}. Candidate cDNAs for *bg* were isolated from YAC 195A8, which contains 650 kb of the *bg* non-recombinant interval using direct cDNA selection with mouse spleen cDNA¹³ (Fig. 1). Of 56 candidate cDNA clones analysed from a direct-selection experiment, evidence for causality in *bg* was found in one (see below), and this gene was designated *Lyst* (lysosomal trafficking regulator). As this clone was 132 nucleotides long, additional *Lyst* sequences were sought by screening three mouse cDNA libraries and performing polymerase chain reaction (PCR) amplification of cDNA ends¹⁴. Ten overlapping *Lyst* clones were identified, representing ~7kb (Genbank accession number, L77889). These were physically assigned to mouse chromosome 13 with pulsed field gel electrophoresis (PFGE) Southern blots,

confirming that they were all derived from a single gene (mouse genome database accession number, MGD-PMEX-14). The *Lyst* probes identified the same polymorphic PFGE restriction fragments as *nidogen* (*Nid*)¹², indicating that *Lyst* and *Nid* are clustered within 650 kb. *Lyst* was also mapped genetically in 504 [C57BL/6J-*bg*^J × (C57BL/6J-*bg*^J × CAST/EiJ)F₁] backcross mice¹² by means of three *TaqI* restriction fragment length polymorphisms (RFLPs). The *Lyst* RFLPs cosegregated with *bg* (and *Nid*), confirming their colocalization on proximal mouse chromosome 13 (MGD accession number, MGD-CREX-615).

Evidence for *Lyst* mutations was found in two *bg* alleles. A 5-kb genomic deletion that contained the 3' end of *Lyst* exon β, and exons γ and δ, was identified in *bg*^{11J} DNA (Fig. 2). The *bg*^{11J} deletion corresponds to loss of ~400 internal amino acids of the predicted *Lyst* peptide. Furthermore, whereas the 5' end of the *bg*^{11J} deletion occurs within *Lyst* exon β, the 3' end is intronic. Therefore the truncated *Lyst* mRNA in *bg*^{11J} mice is also anticipated to splice incorrectly, terminate prematurely, and lack polyadenylation.

Quantitative reverse transcription (RT)–PCR demonstrated a moderate decrease in *Lyst* mRNA in *bg* and *bg*^J liver, and a gross reduction in *bg*^{2J} (*Lyst* ΔOD after normalization for β-actin mRNA: +/+, 1.00; *bg*^{2J}/*bg*^{2J} 0.19; *bg*/*bg*, 0.28; *bg*^J/*bg*^J, 0.40). A commensurate reduction in *bg*^{2J} transcript abundance was noted by using several primer pairs derived from different regions of the *Lyst* cDNA. Aberrant *Lyst* RT–PCR products were not observed. The particularly striking (more than fivefold) reduction in *Lyst* expression evident in *bg*^{2J} homozygotes suggested the existence of a mutation in *bg*^{2J} *Lyst* that results in decreased transcription or mRNA instability. The molecular basis of the decrease in *Lyst* mRNA in *bg*^{2J} is not yet known, but it is reminiscent of the leaky ablation of mature message associated with an intronic retrotransposition event¹⁴.

The predicted open reading frame (ORF) of *Lyst* was 4,635 nucleotides, encoding a protein of 1,545 amino acids and relative molecular mass 172,500 (*M*_r 172.5K) (Fig. 3a). Nucleotides 51–74 are rich in CG dinucleotides, a common feature of the 5' region of housekeeping genes. Comparison with DNA databases indicated that *Lyst* is novel, and resembles only uncharacterized human expressed sequence tags (ESTs) (L. Hillier *et al.*, unpublished data; J. Macke *et al.*, unpublished data). The sequence of a cDNA clone corresponding to one such human EST (Genbank accession number L77889) matched the 5' region of mouse *Lyst* (nucleotide identity was 76% in the 5' untranslated region (UTR), 91% in the ORF, and amino-acid identity was 97%; Fig. 3c); another human EST matched the 3' region of the mouse *Lyst* coding domain (Genbank accession number W26957). On hybridization to PFGE Southern blots of mouse DNA¹², the human clones identified restriction fragments that were indistinguishable from mouse *Lyst*; physical mapping of the human clones to the same region of the mouse genome as *Lyst* indicates that they are indeed homologous to *Lyst*.

It has been suggested that CHS and *bg* represent homologous disorders, as their clinical features¹ and defects in lysosomal transport⁴ are identical. Homology of *bg* and CHS is supported by genetic complementation studies: fusion of fibroblasts from *bg* mice and CHS patients fails to reverse lysosomal abnormalities, in contrast to fusions with normal cells¹⁵. Furthermore, recent genetic linkage studies have shown that CHS maps within a linkage group conserved between human Chromosome 1q43 and the *bg* region on mouse Chromosome 13 (F. J. Barrat *et al.*, manuscript submitted). Therefore LYST mutations in CHS patients were sought by sequencing LYST lymphoblast and fibroblast cDNAs corresponding to these ESTs from 10 CHS patients. In one patient, a single-base insertional mutation was found at nucleotides 117–118 of the LYST coding domain, resulting in a frame shift and termination after amino acid 62 (Fig. 3c).

Previous studies showing spontaneous aggregation of membrane-bound concanavalin A (capping) suggest that there is a defect in microtubule dynamics in *bg* cells^{16,17}. In a search of the SWISSPROT database, using BLITZ and BLASTP, a similarity was found between a domain in *Lyst* and stathmin (oncoprotein 18), a phosphoprotein that may regulate polymerization of micro-tubules¹⁰ (27% identity from residues 463 to 536; best expected occurrence by chance, 4.36×10^{-6}). The domain in stathmin that matches *Lyst* is helical and has heptad repeats that participate in coiled-coil interactions with other proteins⁸⁻⁹. The stathmin-like region of *Lyst* is also predicted to be helical and form coiled coils. However, it is the charged residues, rather than the hydrophobic ones, that are conserved between *Lyst* and stathmin, suggesting that the sequence similarity is not primarily due to conserved secondary structure. Thus this region of *Lyst* potentially encodes a coiled-coil protein-interaction domain that may regulate microtubule-mediated lysosome transport. Although *Lyst* is not predicted to have transmembrane helices, the C-terminal tetrapeptide (CYSP; amino acids 1,542–1,545) is strikingly similar to known prenylation sites¹⁸, which could provide attachment to lysosomal/late endosomal membranes through thioester linkage with the cysteine.

Previous studies of *bg* leukocytes have shown correction of microtubule function (as assessed by Concanavalin A capping) and natural killer activity when treated with inhibitors of protein kinase C (PKC) breakdown^{19,20}, suggesting that *bg* might be regulated by phosphorylation. *Lyst* contains 25 sites of potential phosphorylation by PKC, 36 by casein kinase II (CKII) (many of which overlap those of PKC), two by cAMP-dependent protein kinase, and one by tyrosine kinase (Fig. 3b). Almost half of the predicted helices outside the stathmin-like region (14 of 30) have a PKC- or CKII-phosphorylation signal at their amino terminus, and eight of them form consecutive helical pairs. Thus *Lyst* seems to contain helical bundles with clusters of phosphorylation sites at either end. Stathmin also has an N-terminal phosphorylation site and helix motif, and these *Lyst* domains may have a similar ‘signal-relaying’ function to stathmin^{8,9}. Furthermore, phosphorylation of these positions could provide a control mechanism by causing a conformational shift in the bundles, thereby affecting interactions with other molecules.

Northern analysis and RT-PCR indicated that *Lyst* is ubiquitously transcribed, both temporally and spatially, in mouse and human tissues (Fig. 4). Northern blot analysis also revealed complex alternative splicing of *Lyst* mRNA, with both constitutive and anatomically restricted *Lyst* mRNA isoforms. The largest *Lyst* transcript in human and mouse was 12–14 kb, but this transcript was not constitutively expressed. In mRNA from mouse spleen, human peripheral blood leukocytes, promyelocytic leukaemia HL-60, and several leukaemia lines, the 12–14-kb isoform was either undetectable or barely detectable, but smaller *Lyst* transcripts were abundant (Fig. 4). Given the significance for *bg* mice and CHS patients of defects in the lysosomal and late-endosomal compartments of granulocytes, NK cells and cytolytic T lymphocytes^{21–24}, it is likely that these *Lyst* mRNAs of ~3 kb and 4 kb represent the transcripts of primary functional significance. Probes derived from the 5' or 3' ends of the *Lyst* ORF detected multiple bands on northern blots (Fig. 4a, c), whereas probes that corresponded to more central exons (β and γ) identified only the 12–14-kb transcript (Fig. 4b). Assuming that mature, cytoplasmic *Lyst* transcripts were detected, these results suggest modular assembly of *Lyst* mRNAs, with obligatory usage of the exons that contain start and termination codons, and adaptability in retention of central exons. The possible coexistence of several *Lyst* protein isoforms within cells would allow precise control of functions.

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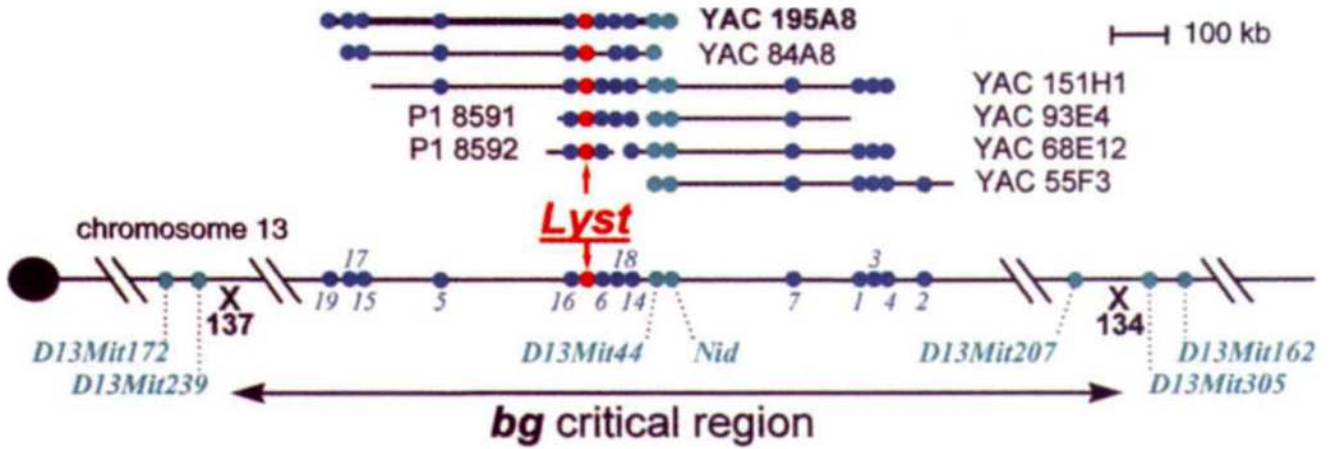


FIG. 1.

Genetic and physical map of the *bg* non-recombinant interval on mouse chromosome 13 showing the location of *Lyst*. Mouse chromosome 13 is shown by a horizontal line with the centromere on the left. The *bg* critical region is delineated by chromosome crossovers (denoted with an X) in animals 134 and 137 of an intersub-specific mouse backcross¹² [C57BL/6J – *bg*^J × (C57BL/6J – *bg*^J × CAST/EiJ)F₁]. Microsatellite markers *D13MM172* and *D13Mit239* flank *bg* proximally; *D13Mit162* and *D13Mit305* lie distal to *bg* (indicated by turquoise circles). YAC and P1 clones identified by PCR screening^{25,26} with oligo-nucleotides corresponding to *Nid* or *D13Sfkl3* are shown above the chromosome. Novel sequence-tagged sites (STS, indicated by dark blue circles), generated by inverse repetitive element PCR or direct cDNA selection¹³, were used to order clones within the contiguous array. Novel mouse chromosome 13 STSs are numbered 1–18, corresponding to *D13Sfkl1* to *D13Sfkl18*, respectively¹³. *Lyst* was isolated from YAC 195A8, a 650-kb clone, by using direct cDNA selection¹³. The physical location of *Lyst*-associated STSs on YAC and P1 clones are shown in red (MGD accession number MGD-PMEX-13).

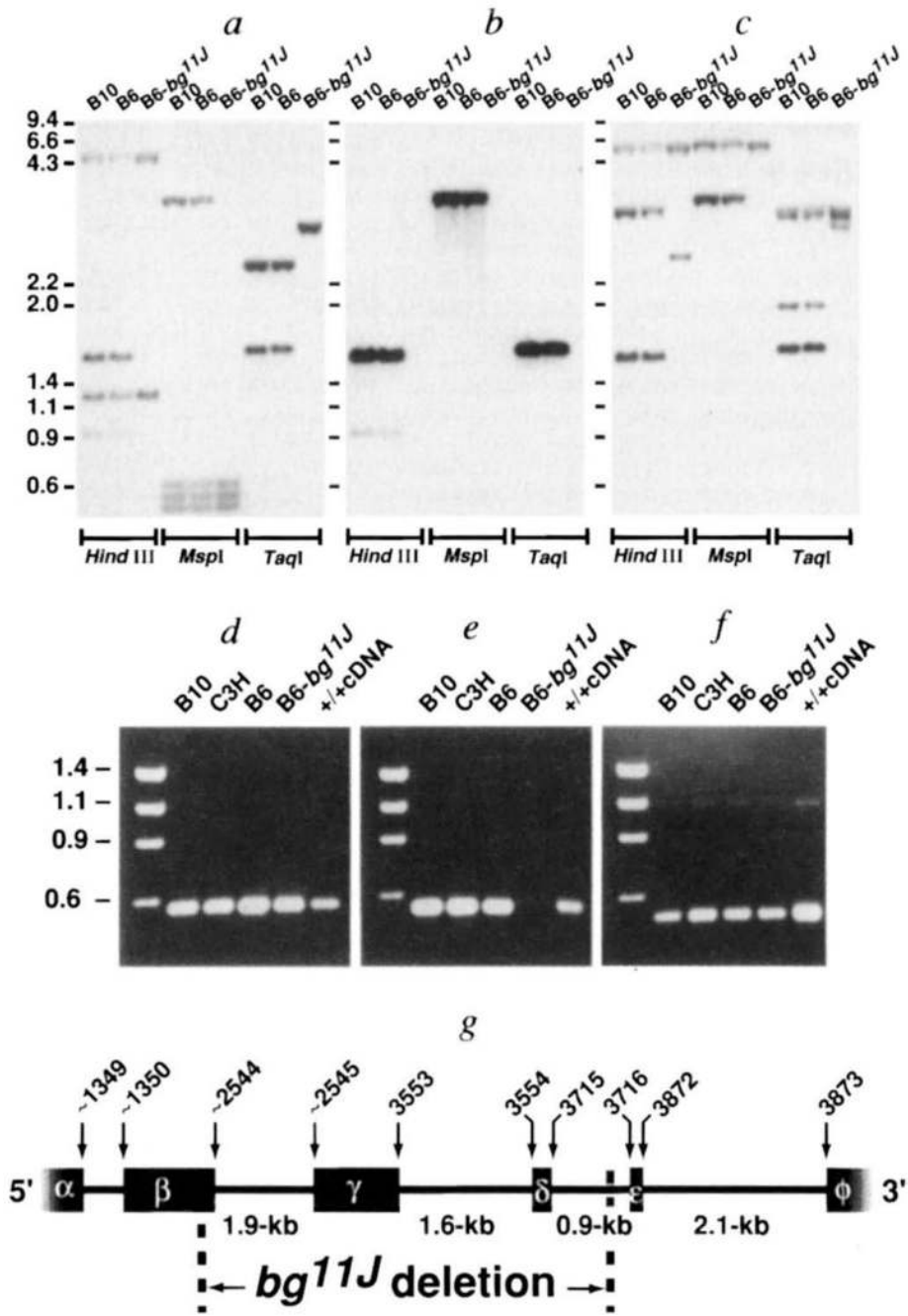
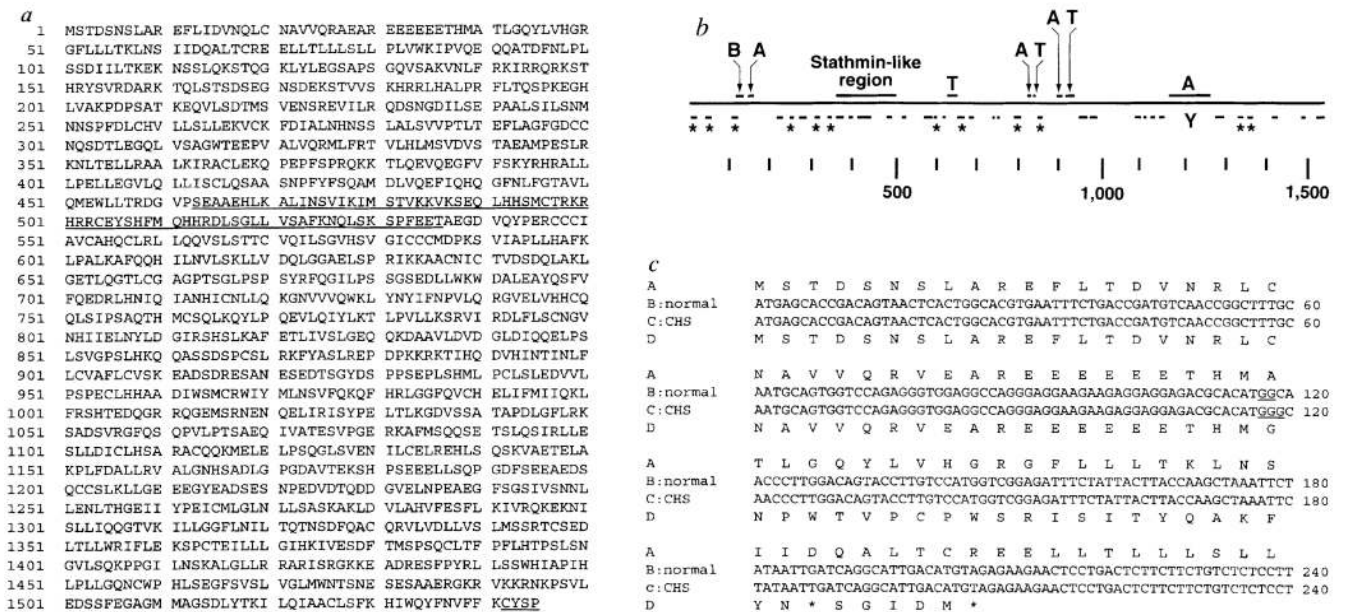


FIG. 2.

Intragenic deletion of *Lyst* in *bg^{11J}* DNA. *a-c*, Southern blot identification of an intragenic *Lyst* deletion in *bg^{11J}* DNA. A Southern blot was sequentially hybridized¹¹ with 3 *Lyst* probes: *a*, the probe (nucleotides 1,262–3,433 of *Lyst* cDNA) extends upstream from the *bg^{11J}* deletion; *b*, the probe (nucleotides 2,835–3,433 of *Lyst* cDNA) is completely deleted; *c*, the probe (nucleotides 3,594–4,237 of *Lyst* cDNA) extends downstream from the *bg^{11J}* deletion. Restriction endonucleases are indicated at the bottom of each panel, and molecular size standards (in kb) are shown to the left. Similar results were obtained with 3 additional restriction endonucleases (data not shown). The *bg^{11J}* mutation was discovered in 1992 at The Jackson Laboratory in a C57BL/6J-*jb* mouse at generation N4 after transfer from B6C3Fe-*a/a*. The

mutation *jb* had, in turn, been discovered 14 generations earlier in B6C3Fe-*a/a-hyh* mice at generation N3 after transfer from C57BL/10J. The *hyh* mutation arose in C57BL/10J mice, and was maintained in that strain until transfer at F15. Thus the possible contributors of genetic information to *bg^{11J}* include C57BL/6J, C3HeB/FeJ and C57BL/10J. Southern blots were prepared from genomic DNA of all potential progenitor mouse strains, but only C57BL/10J, C57BL/6J and C57BL/6J-*bg^{11J}* are shown, *d-f*, PCR analysis of the *bg^{11J}* deletion. C57BL/10J, C3HeB/FeJ, C57BL/6J and C57BL/6J-*bg^{11J}* genomic DNA and *Lyst* cDNA were used as templates in the PCR reactions. Amplicons illustrated correspond to: *d*, *Lyst* cDNA nucleotides 1,337–1,837, which represent exon β and are upstream from the deletion; *e*, nucleotides 2,670–3,210, which represent exon γ , which is deleted in *bg^{11J}* DNA; and *f*, nucleotides 4,913–5,433, which represents an exon downstream from the deletion. No amplicon was observed in control PCR reactions performed without template. More than 30 other STSs that had been localized within the *bg* non-recombinant interval PCR amplified normally from *bg^{11J}* DNA. *g*, Genomic structure of *Lyst* in the vicinity of the *bg^{11J}* deletion. *Lyst* exons (α , β , γ , δ , ϵ and ϕ) are depicted by black boxes, and intervening introns by a solid line. Nucleotides of the mouse *Lyst* cDNA that correspond to exonic boundaries are indicated above the boxes. The 3' end of exon β , and all of exons γ and δ , are deleted in *bg^{11J}* DNA. The region of *Lyst* protein that is deleted in *bg^{11J}* contains a pair of helices with N-terminal phosphorylation sites. Genomic structure and intronic sequences were ascertained by sequence analysis of nested PCR products, performed with exonic primers and P1 clone DNA as template¹⁴. Boundaries of the *bg^{11J}* deletion were determined by PCR of genomic DNA.

**FIG. 3.**

Amino-acid sequence and predicted peptide structure of mouse and human *Lyst* cDNAs. *a*, Predicted amino-acid sequence of the mouse *Lyst* cDNA. The region (residues 463–536) with sequence similarity to stathmin is underlined, as is a C-terminal prenylation motif, *b*, Predicted secondary structure^{27–30} and motifs of mouse *Lyst* protein. The solid line denotes the protein sequence. Above the line are the stathmin-like region and areas rich in acid (A; Asp, Glu), basic (B; His, Arg, Lys), and turn (T; Pro, Gly, Ser, Thr, Asp) residues, the latter, especially at regions 600–625 and 875–895, are too long to be a simple turn between secondary structural elements, and are either very flexible or very rigid (in a manner reminiscent of collagen, although they do not fit the collagen consensus motifs). Helices predicted with >80% confidence are shown as solid lines below the protein sequence, and asterisks identify helices with N-terminal phosphorylation motifs. A tyrosine phosphorylation motif (residue 1,214) is designated Y. *c*, Nucleotide and predicted amino-acid sequence of the 5' end of the human *LYST* coding domain showing a frame-shift mutation in a patient with CHS. Line A illustrates the predicted normal human amino-acid sequence; line B shows the normal human nucleotide sequence; line C shows the nucleotide sequence from a patient with CHS; line D shows the predicted amino-acid sequence from the CHS patient. The CHS nucleotide sequence contains an insertional mutation at nucleotide 117–118 that results in a frame shift and premature termination (asterisk).

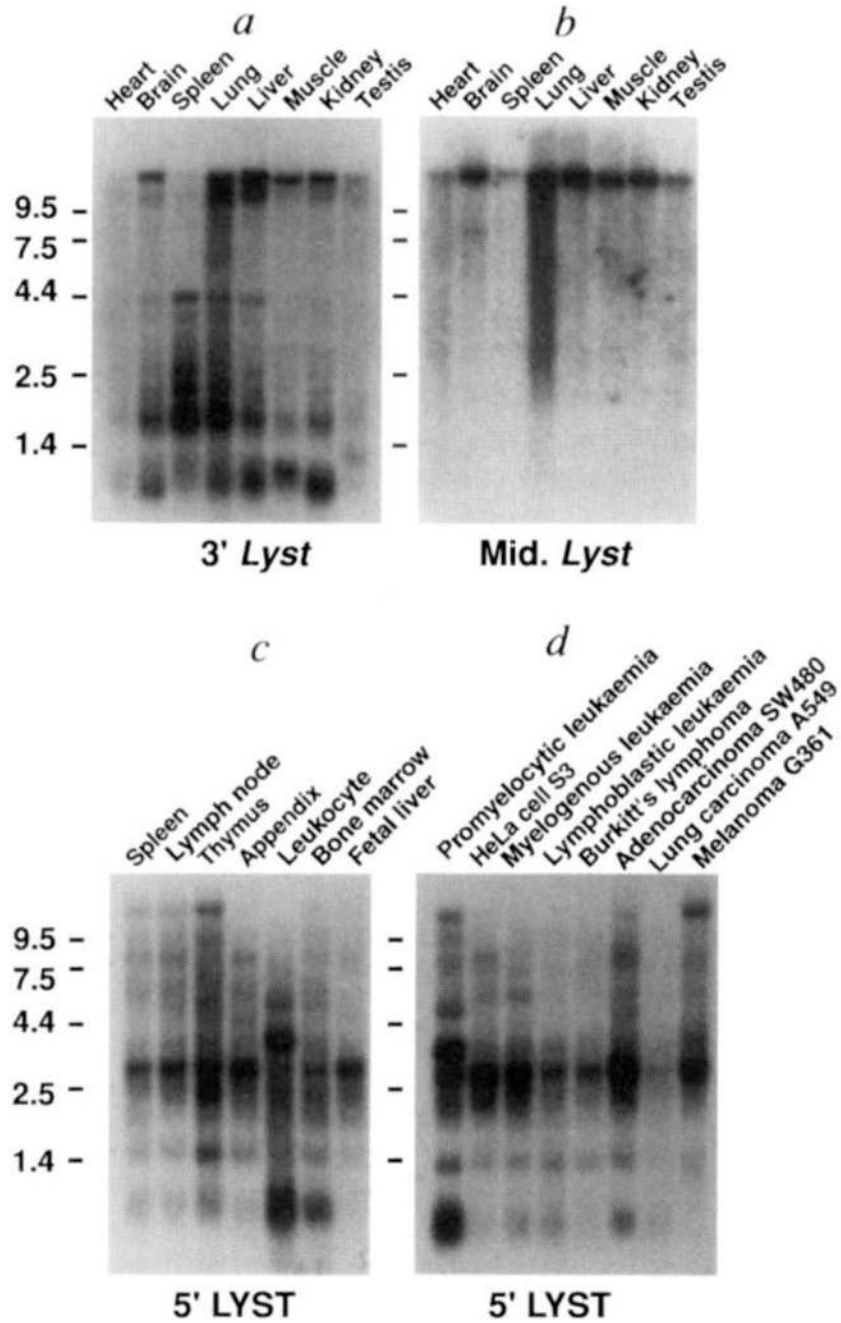


FIG. 4. Northern blot analysis of mouse and human *Lyst*. *a, b*, Northern blots of 2 μ g poly(A)⁺ RNA from various mouse tissues (Clontech) hybridized with probes that correspond to *a*, nucleotides 4,423–4,631, and *b*, nucleotides 1,430–2,457 (exon β) of mouse *Lyst* cDNA. *c, d*, Northern blot of 2 μ g poly(A)⁺ RNA from *c*, various human lymphoid tissues, or *d*, human cancer cell lines, hybridized with a probe that corresponds to nucleotides 357–800 of human *LYST* cDNA. Molecular size standards (in kb) are shown to the left. Hybridization of mouse mRNA with probes from mouse *Lyst* exons α and γ gave identical results to those shown with exon β (*b*), whereas probes from exons δ , ϵ , and ϕ gave results identical to those shown in *a*.