

Rescue of osteoclast function by transgenic expression of kinase-deficient Src in *src*^{-/-} mutant mice

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The Src tyrosine kinase has been implicated in a wide variety of signal transduction pathways, yet despite the nearly ubiquitous expression of *c-src*, *src*^{-/-} mice show only one major phenotype—osteopetrosis caused by an intrinsic defect in osteoclasts, the cells responsible for resorbing bone. To explore further the role of Src both in osteoclasts and other cell types, we have generated transgenic mice that express the wild-type and mutated versions of the chicken *c-src* proto-oncogene from the promoter of *tartrate resistant acid phosphatase (TRAP)*, a gene that is expressed highly in osteoclasts. We demonstrate here that expression of a wild-type transgene in only a limited number of tissues can fully rescue the *src*^{-/-} phenotype. Surprisingly, expression of kinase-defective alleles of *c-src* also reduces osteopetrosis in *src*^{-/-} animals and partially rescues a defect in cytoskeletal organization observed in *src*^{-/-} osteoclasts. These results suggest that there are essential kinase-independent functions for Src *in vivo*. Biochemical examination of osteoclasts from these mice suggest that Src may function in part by recruiting or activating other tyrosine kinases.

[Key words: Src; tyrosine kinase; transgenic mice]

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The Src tyrosine kinase is a member of a family of nine closely related tyrosine kinases defined by a common structure, including SH2 and SH3 protein interaction domains, a catalytic region, and a negative-regulatory tyrosine located near the carboxyl terminus (Brown and Cooper 1996). Much of what is known about the function of these domains has come from studies of cellular transformation either by *v-src*, or by activated versions of the *c-src* proto-oncogene. However, the function of these domains in signal transduction pathways involving endogenous Src is not well defined. Src kinase is activated by a variety of signaling pathways including stimulation of growth factor receptors, G protein-coupled receptors, oxidative and UV stress, and integrin-mediated signal transduction (Brown and Cooper 1996). Furthermore, Src has been demonstrated to regulate the activity of both potassium and NMDA channels (Holmes et al. 1996; Yu et al. 1997). To date, however, critical downstream targets of the kinase have remained controversial in many of these pathways.

Targeted gene-disruption of *c-src* in mice leads to only one major phenotype—severe osteopetrosis—with re-

sulting thickened bone, decreased marrow space, and a failure of tooth eruption (Soriano et al. 1991). The severe morbidity and increased mortality resulting from osteopetrosis greatly complicate analysis of Src function in other tissues and organ systems in these mice. The *src*^{-/-} phenotype results from an intrinsic defect in osteoclasts, the cells that resorb bone (Lowe et al. 1993). Osteoclasts are highly polarized cells of the monocyte-macrophage lineage that break down bone through the actions of the ruffled border, a specialized region of the cell consisting of highly invaginated membranes surrounded by the sealing zone, where the osteoclast firmly attaches to the bone matrix (Baron et al. 1993). The ruffled border is the site of secretion of hydrogen ions and proteolytic enzymes that promote digestion of the bone matrix. Although osteoclasts are present in *src*^{-/-} mice in normal or increased numbers, they fail to form ruffled borders and consequently do not resorb bone (Boyce et al. 1992). The osteoclast therefore strictly requires Src activity for its proper morphology and function and provides a unique cell type in which to evaluate Src function in a physiological setting.

We have developed a genetic system for studying Src function in the osteoclasts of transgenic mice by driving expression of wild-type and mutant versions of Src with

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the promoter of *tartrate resistant acid phosphatase (TRAP)*, a gene expressed highly in osteoclasts. Using this system, we demonstrate that Src kinase activity is not essential for rescue of the *src*^{-/-} phenotype. These results suggest that there are essential kinase-independent functions for Src in vivo. Furthermore, by generating *src*^{-/-} mice that express *src* in limited cell lineages, this approach may allow the analysis of Src function in other tissues in mice free from the morbidity of osteopetrosis.

Results

Generation of transgenic mice expressing *c-src* in osteoclasts

To achieve expression of Src in osteoclasts, we used the promoter of the gene-encoding *TRAP*, an enzyme highly expressed in osteoclasts. The wild-type chicken *c-src* cDNA was fused to the ATG and ~1.8-kb of upstream sequences of the *TRAP* gene, including the first exon and intron (Reddy et al. 1995), and introduced into mice by pronuclear injection (Fig. 1A). We elected to use the chicken *c-src* cDNA in these experiments for several reasons. First, the availability of an avian-specific monoclonal antibody allows the specific identification of the protein encoded by the transgene in mouse tissues. Second, the chicken *c-src* gene has been studied widely and large numbers of mutants have been characterized in this background. Furthermore, our laboratory has demonstrated previously that the chicken *c-src* gene can rescue a cell-spreading phenotype observed in fibroblasts derived from *src*^{-/-} mice, suggesting that the chicken *c-src* gene is able to substitute for mouse *c-src* function, at least in this celltype (Kaplan et al. 1995).

Expression of the *TRAPsrc*^{WT} transgene was examined in bone marrow removed from transgenic mice and cultured in the presence of 1,25 dihydroxy vitamin D3 (1,25 D3) under conditions that induce the differentiation of

osteoclasts (Takahashi et al. 1988a). Four of six independent transgenic lines demonstrated 1,25 D3-dependent expression of the transgene as assayed by reverse transcriptase (RT)-PCR (Fig. 1B.) Immunohistochemistry of bone sections from *TRAPsrc* mice also demonstrates that expression of the transgene in bone is restricted to osteoclasts (L. Xing and B. Boyce, unpubl.). *TRAPsrc*^{WT} mRNA was also observed by Northern analysis in a limited number of other tissues, including small intestine (in all lines examined to date), liver and kidney (3/4 lines), and thymus (1/4 lines) (Table 2, below). This pattern of expression is consistent with that reported previously for the *TRAP* promoter in transgenic mice (Boyce 1995) and differs from the nearly ubiquitous expression of endogenous *c-src*.

The four *TRAPsrc*^{WT} lines (A-D) were crossed to *src*^{+/-} mice, and progeny were interbred to generate *src*^{-/-}*TRAPsrc*^{WT} mice and evaluate the effects of the *TRAPsrc*^{WT} transgene on the *src*^{-/-} phenotype. Because functional osteoclasts are required to resorb a channel in jaw bones through which incisor teeth pass, one of the most obvious aspects of the *src*^{-/-} phenotype is a lack of tooth eruption (Soriano et al. 1991). Expression of the *TRAPsrc*^{WT} transgene in all four lines restored normal tooth eruption in *src*^{-/-} mice (Table 1), as illustrated radiographically in Figure 2a. Microscopic examination of long bones revealed that the transgene had restored normal bone histology (Fig. 2b) and volume (Fig. 3). Furthermore, expression of the transgene also rescued osteoclast morphology (Fig. 2c); osteoclasts in bone sections from rescued mice display normal ruffled borders.

The *TRAPsrc*^{WT} transgenic lines showed no major deleterious effects of expressing the chicken *c-src* cDNA. The rescued mice, including those from line C, in which transgene expression is limited to osteoclasts and intestine, were grossly indistinguishable from their wild-type and heterozygous littermates. The *src*^{-/-}*TRAPsrc*^{WT} mice appeared healthy, with no unexpected deaths up to 1.5 years of age, and both sexes were fully fertile. In

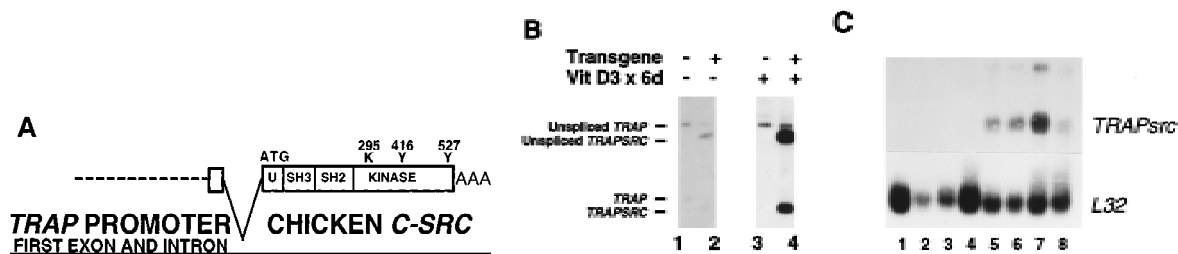


Figure 1. (A) Structure of the *TRAPsrc* transgene. The *TRAP* promoter, first exon, and intron were fused to the ATG of the chicken *c-src* cDNA. Encoded amino-acids important for activation of Src or mutated in transgenic constructs are indicated above the drawing. (U) Unique region; (SH2) Src homology 2 region; (SH3) Src homology 3. (B) Expression of the *TRAPsrc* transgene in cultured osteoclasts from transgenic mice. RNA samples isolated from fresh bone marrow and from bone marrow cultured in the presence of 10^{-9} M Vitamin D3 (Takahashi et al. 1988) were analyzed by RT-PCR, using primers that generate products extending from the *TRAP* first exon to both the chicken *c-src* and the endogenous *TRAP*-coding sequences. Products were hybridized with a P^{32} -labeled probe that recognizes the *TRAPsrc*-specific band as well as unspliced *TRAP* RNA. (Lanes 1,2) Fresh bone marrow; (lanes 3,4) bone marrow cultured in the presence of 10^{-9} M Vitamin D3. (Lanes 1,3) Non-transgenic mice. (C) Northern analysis of RNA derived from tissues of transgenic line A and C mice. (Lanes 1-7) Line A (Lane 1) Testes; (lane 2) brain; (lane 3) heart; (lane 4) spleen; (lane 5) kidney; (lane 6) liver; (lane 7) small intestine; (Lane 8) line C small intestine. (top) The filter was hybridized with a probe consisting of the first 200 bp of the chicken *c-src* gene; (bottom) the hybridized to a probe for RNA encoding the L32 ribosomal protein.

Table 1. Tooth eruption in *src* gene-targeted and transgenic mice

Mouse line	c- <i>src</i> genotype	Transgene	Percent tooth eruption
	+/,+/-	none	100
	-/-	none	<1 (n > 100)
A	-/-	TRAPsrc ^{wt}	100 (n > 100)
B-D	-/-	TRAPsrc ^{wt}	100 (n > 12/line)
G	-/-	TRAPsrc ^{Y416F}	100 (n = 23)
H	-/-	TRAPsrc ^{Y416F}	100 (n = 10)
J	-/-	TRAPsrc ^{K295M}	96 (n = 22)
N	-/-	TRAPsrc ^{K295M}	73 (n = 18)

contrast, *src*^{-/-} mice in this genetic background had greatly reduced fertility and high rates of morbidity and mortality, as described previously, with ~1/3 of animals dying between 3–6 weeks of age and continual loss thereafter (Soriano et al. 1991). Therefore, despite a relatively limited pattern of expression of this typically widely expressed gene, we have obtained full complementation of both the osteopetrosis and the increased morbidity of *src*^{-/-} mice. Abnormalities have been reported in several cell types, including neurons and fibroblasts, cultured from *src*^{-/-} mice (Ignelzi et al. 1994; Kaplan et al. 1995). However, our data suggest that the major cause of increased morbidity in *src*^{-/-} mice results directly from osteopetrosis and support earlier evidence that *src* deficiency in osteoclasts is responsible for the bone pathology.

Kinase-defective mutants of *Src* rescue the *src*^{-/-} phenotype

The ability of the wild-type transgene to rescue mice from osteopetrosis facilitates a genetic approach to the study of *Src* in osteoclasts in vivo, a physiologically relevant setting in which the lack of *Src* protein produces marked phenotypic effects. We have taken advantage of this system to investigate the requirement for the various structural regions of the *Src* protein in the osteoclast by generating *src*^{-/-} mice that express mutant versions of *Src* that affect kinase activity.

We first generated mice expressing *src*^{Y416F}, an allele that reduces full activation of the kinase severalfold by changing the tyrosine of the major autophosphorylation site in pp60^{src} to phenylalanine (Kmiecik and Shalloway 1987; Piwnicka-Worms et al. 1987). Like wild-type *src*, this mutant also fully restored tooth eruption in two independent transgenic lines (lines G and H) when expressed as a transgene (TRAPsrc^{Y416F}). X-ray and histological analyses confirmed that the *src*^{Y416F} mutant transgene produced near normal bone histology and volumes in *src*^{-/-} mice (Figs. 2a,b and 3). Furthermore, like their wild-type-rescued counterparts, the *src*^{-/-}TRAPsrc^{Y416F} mice were fully fertile and have remained healthy up to 9 months of age.

As a more stringent test of the requirement for kinase activity, we examined the effects of expression of

src^{K295M}, an allele that encodes a protein deficient in kinase activity because the ATP-phosphotransferase site of pp60^{src} is disrupted (Kamps and Sefton 1986). Surprisingly, expression of this allele under the control of the TRAP promoter also restored tooth eruption in the *src*^{-/-} background. One transgenic line expressing the *src*^{K295M} mutant (line J) complemented the *src*^{-/-} phenotype almost completely. A second line (line N) restored tooth eruption in the majority (73%) of *src*^{-/-} mice (Table 1). Although the *src*^{-/-}TRAPsrc^{K295M} rescued mice were often smaller than their wild-type littermates, we have not observed increased mortality in the rescued animals up to 9 months of age. However, these mice are less fertile and exhibit more restricted patterns of expression of the transgene than the *src*^{-/-}TRAPsrc^{WT} transgenic mice (Table 2). As with the Y416F and wild-type transgenes, we observed no adverse effects of expression of the K295M mutant in *src*^{+/+} and heterozygote backgrounds.

Expression of the K295M mutant protein, with its severely impaired kinase activity, was associated with more variable effects on bone histology and density than was expression of the Y416F mutant (Fig. 2, bottom two panels). Nevertheless, bone volumes were improved significantly relative to *src*^{-/-} mice ($P < 0.05$; Fig. 3); in some of the mice (Fig. 2, bottom panel), the bones appeared entirely normal, indicating complete rescue of the bone phenotype. Osteoclasts from mice rescued with either the *src*^{K295M} or the *src*^{Y416F} transgenes formed ruffled borders, although these structures were present in fewer osteoclasts in animals rescued by the *Src*^{K295M} mutant compared with *Src*^{Y416F} (Fig. 2c). We also did not observe an increase in the number of osteoclasts in these mice, compared with wild-type animals, unlike the increase observed in *src*^{-/-} mice.

Functional capabilities of the kinase-inactive rescued osteoclasts

To test the functional capabilities of osteoclasts in *src*^{-/-} mice expressing the TRAPsrc^{K295M} transgene, we challenged these animals with interleukin-1 (IL-1), a potent stimulator of bone resorption that typically causes a transient increase in whole-blood-ionized calcium and an increase in osteoclast number in normal mice after several days treatment (Boyce et al. 1989). Osteoclasts from *src*^{-/-} mice treated with IL-1 increase in number, but do not resorb bone, and no change is seen in blood calcium levels (Boyce et al. 1992). In contrast, animals rescued by the TRAPsrc^{K295M} allele responded to a challenge of IL-1 with an increase in whole blood ionized calcium, indicative of appropriately activated osteoclasts (Table 3) and a generalized increase in bone resorption in vivo.

src^{-/-} osteoclasts have altered cell morphology

Despite complete or almost complete rescue of the osteopetrosis, osteoclasts in the *src*^{-/-}TRAPsrc^{K295M} mice had an elongated rather than a rounded morphology and

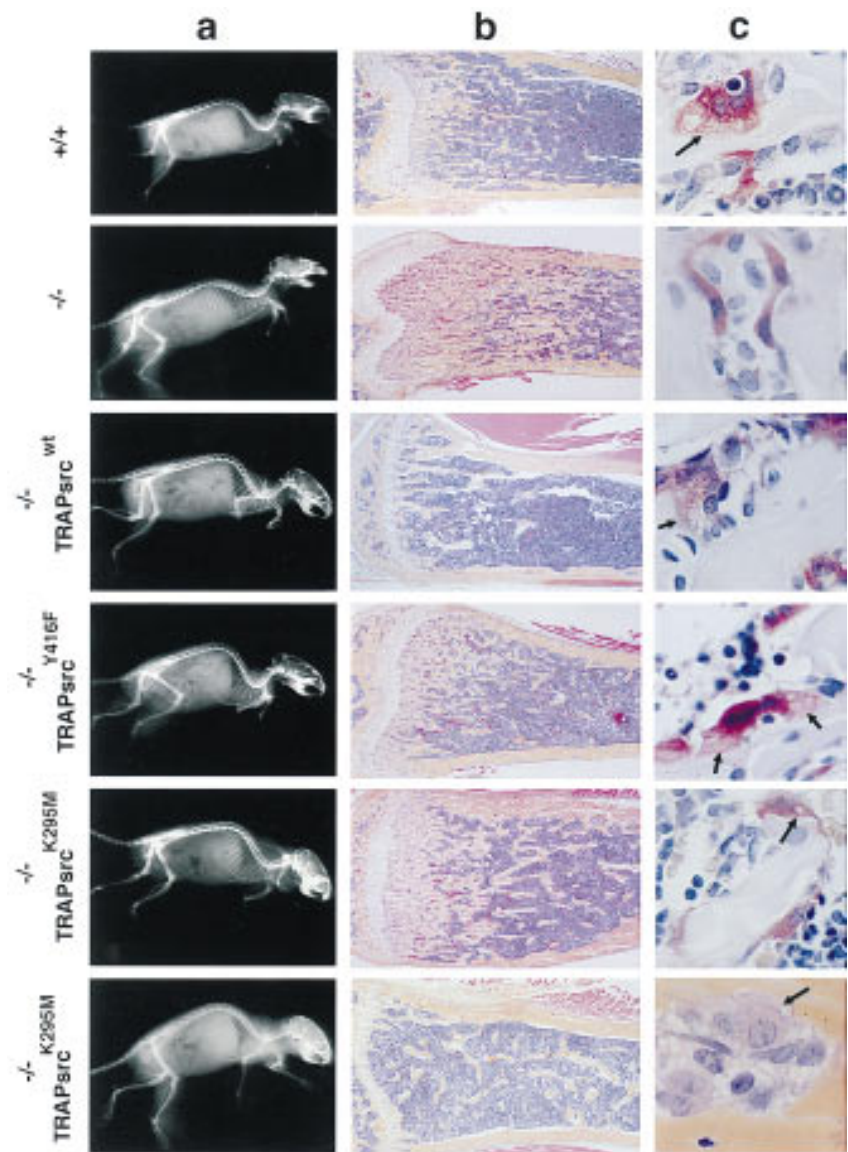


Figure 2. Radiographic and histological analysis of mice. (A) X-rays, lateral view. (B) Histological sections of tibia, low power (4 \times magnification). Sections were stained both with hematoxylin and eosin and for TRAP activity. (C) Histological sections, TRAP-stained osteoclasts in bone, high power (100 \times , note lacy ruffled border structures at the interface between osteoclasts and bone surface.) Genotypes of mice are indicated. Two examples of bones from *src*^{-/-} mice rescued by the K295M transgene (*TRAPsrc*^{K295M}) are shown to demonstrate the variability in phenotype. The *bottom* sample from a *src*^{-/-} *-TRAPsrc*^{K295M} mouse is derived from the calvarium after treatment with IL-1 and is stained with hematoxylin and eosin only.

as such appeared more similar to osteoclasts in *src*^{-/-} mice (Fig. 2). To explore further differences in the morphology of these cells, we examined osteoclasts differentiated from bone marrow cells in culture. Osteoclasts differentiated from bone marrow cells cultured from *src*^{-/-} mice in the presence of 1,25 D3 appear markedly different from osteoclasts derived from the bone marrow of *src*^{+/+} mice (Fig. 4A). In particular, after they form by fusion of TRAP-positive mononuclear precursors, these cells fail to assume the typical rounded appearance of wild-type osteoclasts and have an elongated appearance, somewhat resembling cells seen in bone sections from these mice.

To examine these differences further, we stained cells with rhodamine-phalloidin to highlight F-actin filaments and examined them by confocal microscopy. Normal osteoclasts adhere to plastic through a specialized structure called the podosome, found on motile cells like

macrophages as well as on fibroblasts transformed by *v-src*. (Marchisio et al. 1984). These adhesive structures are represented by dot-like aggregations of actin, which cluster in a ring around the periphery of wild-type osteoclasts plated on plastic (Lakkakorpi and Vaanenen 1991; Teti et al. 1991). *src*^{-/-} osteoclasts, in contrast, have a markedly different distribution of actin, with no peripheral podosome arrangement; the actin organization in these cells more closely resembles the stress fibers of fibroblasts (Neff et al. 1996). These observations suggest that Src has a fundamental role in organizing the actin cytoskeleton of osteoclasts.

Examination of osteoclasts from *src*^{-/-} mice rescued with a wild-type transgene demonstrated that actin is organized normally, providing further evidence of full rescue of the morphology of these cells (Fig. 4B). Osteoclasts from mice rescued with the Y416F mutant also exhibited a normal pattern of actin. However, part of

Table 2. Summary of expression patterns of transgenic mice

Tissue	Transgenic line					
	A	B	C	D	J	N
Cultured bone marrow	+	+	+	+	+	+
Small intestine	+	+	+	+	+	+
Liver	+	+	-	+	-	-
Kidney	+	+	-	+	-	-
Spleen	-	-	-	-	-	-
Heart	-	-	-	-	-	-
Thymus	-	+	-	-	-	-
Brain	-	-	-	-	-	+
Muscle	-	-	-	-	-	-
Testes	-	-	-	-	-	-
Ovary	-	-	-	-	-	-
Mammary	-	N.D.	N.D.	N.D.	-	-
Lung	-	-	-	-	-	-

Expression patterns were examined by Northern blotting, except for cultured osteoclasts, which were analyzed by RT-PCR as in Fig. 1B. Lines A-D express the *TRAPsrc*^{WT} transgene. Lines J and N express the *TRAPsrc*^{K295M} transgene. (N.D.) Not determined.

their actin ring was often diffuse, a feature that has been observed in osteoclasts moving across a surface (Baron et al. 1993). In contrast, many osteoclasts from *src*^{-/-} mice rescued with the *TRAPsrc*^{K295M} transgene appear very similar to *src*^{-/-} osteoclasts (Fig. 4B; bottom right). Some *src*^{-/-}*TRAPsrc*^{K295M} osteoclasts did have a more rounded appearance and a peripheral actin organization. However, the actin pattern in many of these cells was still not consistent with a normal podosome arrangement. Only relatively rare osteoclasts display the normal peripheral actin podosome pattern (Fig. 4B, bottom left), despite the ability of these cells to rescue bone resorption in mice. These differences may result from variation in the degree of rescue of the osteoclasts from these

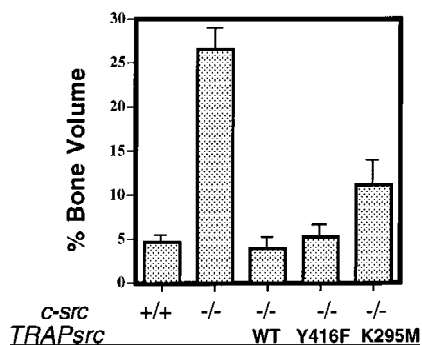


Figure 3. Histomorphometric analysis of rescued mice. Genotype of mice indicated on x axis. Percentage bone volume [amount of bone matrix as a percentage of the cancellous bone (Boyce et al. 1992)] is indicated on the y axis. A minimum of four samples were examined for each transgene. Histology was performed on samples from at least two transgenic lines for each construct, except for the *TRAPsrc*^{Y416F} rescued mice (histomorphometry was performed on line G only).

Table 3. *TRAPsrc*^{K295M} restores osteoclast bone resorption in response to IL-1

Genotype	Blood Ca ²⁺ (mmole/liter)		
	pre-IL-1	post IL-1	difference
Wild type	1.25 ± 0.03	1.36 ± 0.05	0.11
<i>src</i> ^{-/-} <i>TRAPsrc</i> ^{K295M}	1.25 ± 0.02	1.33 ± 0.03	0.08
Wild type	1.28 ± 0.01	1.44 ± 0.03	0.16
<i>src</i> ^{-/-}	1.29 ± 0.02	1.28 ± 0.01	-0.01

Mice (three to five per group) were treated with IL-1 four times daily for 3 days. Whole Blood Ca²⁺ was measured 2 hr after the last injection. Ca²⁺ levels in *src*^{-/-} animals post-IL-1 treatment were statistically lower than those in the wild-type mice ($P < 0.01$), whereas Ca²⁺ levels in the *src*^{-/-}*TRAPsrc*^{K295M} rescued mice were not statistically different than those of wild-type mice ($P > 0.25$). Results from two different experiments are shown (Boyce et al. 1992).

mice. But, overall our results suggest that, whereas expression of the K295M mutant permits bone resorption in vivo, it does not fully restore osteoclasts morphology, and only restores partial cytoskeleton organization in osteoclasts plated on glass.

Biochemical analysis of osteoclasts derived from kinase-defective *Src* mice

Given the surprising finding that a kinase-defective allele rescues *Src*-deficient animals from osteopetrosis, we confirmed that the point mutations in the kinase-defective *src* allele had not reverted to normal sequence by examining both DNA and protein from the transgenic mice. DNA from *TRAP-src* transgenic mice was amplified by PCR and the expected products were detected by allele-specific hybridization with oligonucleotides demonstrating retention of the mutations (data not shown). To examine the enzymatic activity of the *Src* proteins, in vitro kinase assays were performed on immunoprecipitated pp60^{Src} from osteoclasts cultures derived from the bone marrow of *src*^{+/+}, *src*^{-/-}, *src*^{-/-}*TRAP-src*^{K295M} and *src*^{-/-}*TRAPsrc*^{Y416F} mice. Despite similar levels of protein relative to endogenous *Src* (within two-to four-fold), no kinase activity above background was detected with the *Src*^{K295M} mutant on the exogenous substrate, enolase (Fig. 5A), whereas the *Src*^{Y416F} mutant displayed reduced kinase activity relative to that of wild-type *Src*.

We then examined patterns of total cellular tyrosine phosphorylation in osteoclasts cultured from mice rescued by the kinase-inactive *src*^{K295M} allele. Basal levels of phosphotyrosine were consistently lower in osteoclasts derived from *src*^{-/-} bone marrow than in cells derived from wild-type mice (Fig. 5B). However, the overall extent of tyrosine phosphorylation was restored in osteoclasts derived from mice rescued by the *TRAPsrc*^{K295M} transgene. Hunter and colleagues have detected an extremely low level of residual kinase activity of *Src*^{K295M} when overexpressed by baculovirus (<0.003% of wild-type levels) (Broome 1996). We think it

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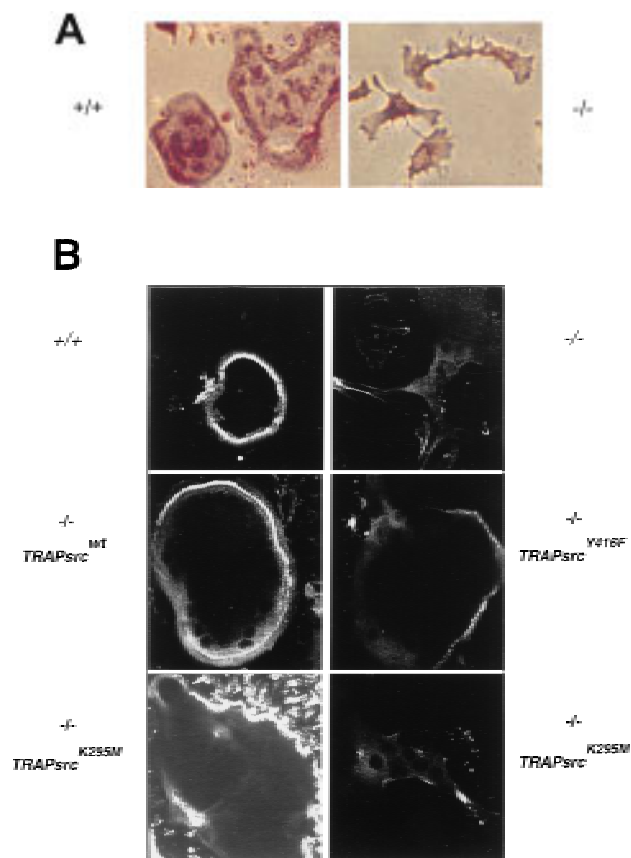


Figure 4. Rescue of abnormal morphology of *src*^{-/-} osteoclasts by expression of *TRAPsrc* transgenes. (A) Morphology of *src*^{-/-} osteoclasts. Osteoclasts were differentiated on plastic, partially purified, and stained for TRAP activity. (B) Actin organization of osteoclasts expressing *TRAPsrc* transgenes. Osteoclasts were stained with Rhodamine-phalloidin and examined by confocal microscopy. Genotypes of Osteoclasts are indicated. Note lack of peripheral actin ring in *src*^{-/-} osteoclasts.

unlikely that this small amount of kinase activity could account for the increase in cellular tyrosine phosphorylation we observe, particularly because the amount of Src^{K295M} protein present in cultured osteoclasts is lower (two to fivefold) than that of endogenous Src in cells derived from normal mice. Because the intrinsic kinase activity of Src^{K295M} is defective and kinase activity is undetectable in osteoclasts from *src*^{-/-}*TRAPsrc*^{K295M} mice (Fig. 5A), the increase in tyrosine phosphorylation we observe suggests that Src^{K295M} may function in part by activating or recruiting other tyrosine kinases.

Discussion

Transgenic rescue of *src*^{-/-} mice

Despite the ubiquitous expression of *c-src*, mice deficient in *src* demonstrate only one major phenotype—moderately severe osteopetrosis, with resulting high rates of morbidity and mortality attributable to an in-

trinsic defect in osteoclasts. We have now generated transgenic mice that express *c-src* in limited cell lineages. These mice support earlier conclusions that the defect in osteoclasts alone accounts for the *src*^{-/-} phenotype and that osteopetrosis is specifically due to the lack of Src expression in these cells. The availability of healthy, fertile *src*^{-/-} *TRAPsrc* mice with limited patterns of *src* expression should now facilitate the study of Src deficiency in other tissues, as well as the generation of *src*^{-/-} cell lines, and the breeding of Src deficiency into a number of genetic backgrounds. In particular, these mice may be useful for the generation of mice and cell lines containing loss-of-function mutations in multiple *src* family members and for the study of the actions of mutant forms of Src family proteins in signaling pathways.

Kinase-independent functions of Src

In this paper, we have used the *TRAPsrc* transgenic system to examine the functional capabilities of mutant versions of Src. Our observation that kinase-defective versions of Src protect *src*^{-/-} mice against osteopetrosis indicates that the intrinsic kinase activity of Src itself is not required for certain critical functions of Src in osteoclasts. These results suggest that there are essential Src

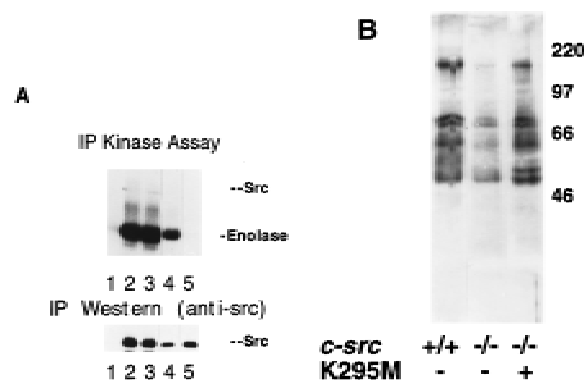


Figure 5. Osteoclasts from mice rescued with the *TRAPsrc*^{K295M} transgene lack detectable Src kinase activity but have normal levels of tyrosine phosphorylation. (A) In vitro kinase assays. Osteoclasts derived from bone marrow were lysed in RIPA buffer, and extracts were normalized for both protein concentration (Biorad) and TRAP enzymatic activity (Sigma); after immunoprecipitation with anti-Src monoclonal 327 (Calbiochem), in vitro kinase assays with the exogenous substrate enolase were performed as described previously (Kaplan et al. 1995). One-half of each immunoprecipitate was analyzed by Western blotting probing with the Src2 antisera (Santa Cruz). (Lane 1) *src*^{-/-}; (lane 2) *src*^{+/+}; (lane 3) *src*^{-/-}*TRAPsrc*^{WT} line A; (lane 4) *src*^{-/-}*TRAPsrc*^{WT} line C; (lane 5) *src*^{-/-}*TRAPsrc*^{K295M}. (B) Tyrosine-phosphorylated proteins from cultured osteoclasts from wild-type *src*^{-/-} and kinase-defective rescued mice. Protein extracts from cultured osteoclasts were normalized for both protein concentration (Bio-Rad) and TRAP enzymatic activity (Sigma), and analyzed with the 4G10, anti-phosphotyrosine antibody (UBI). Lanes are derived from the same exposure of a single gel.

signaling pathways in osteoclasts that are independent of Src kinase activity, a finding that contrasts with previous observations that certain signaling pathways in cultured cells appear to require Src kinase activity. For example, overexpression of the same kinase-defective mutant of Src used in our studies (K295M) interferes with growth factor signaling in isolated fibroblasts (Twamley-Stein et al. 1993; Broome and Hunter 1996). Kinase-negative alleles also adversely affect stress-activation of Jun kinase and hypoxia-induced expression of vascular endothelial growth factor (VEGF) (Devary et al. 1992; Mukhopadhyay et al. 1995), whereas other kinase-defective Src family members expressed previously in transgenic animals have acted in a dominant-negative manner (Cooke et al. 1991; Levin et al. 1993). Given the dominant-negative properties of overexpression of the K295M mutant in multiple pathways, it is remarkable that not only do we not see adverse effects of expression of this mutant in wild-type and heterozygote animals, but the same mutant rescues the *src*^{-/-} phenotype. It is yet another indication that the osteoclast may have distinctive signaling pathways that require the Src protein.

The global decrease in tyrosine phosphorylation we observe in the osteoclasts from *src*^{-/-} mice suggests that Src may be required for proper phosphorylation of a large number of proteins involved in many cellular functions. This observation is similar to the decrease in tyrosine phosphorylation found in the brains of *fyn*^{-/-} mice (Grant et al. 1995) and supports roles for Src family kinases in multiple signaling pathways within a given cell. Our present results suggest further that at least some of these pathways may be independent of Src's own intrinsic kinase activity.

Not all pathways involving Src may require kinase activity. Activation and cross-linking of integrin receptors is associated with a transient increase in Src kinase activity and translocation of Src to the focal adhesion complex where Src interacts with and phosphorylates the focal adhesion kinase (FAK) (Schaller et al. 1994; Schaller and Parsons 1994; Schlaepfer et al. 1994; Clark and Brugge 1995). Kaplan et al. (1995) have demonstrated previously that cultured fibroblasts from *src*^{-/-} mice have a defect in cell spreading when plated on fibronectin—a defect presumably related to the activation of Src by integrin-mediated signaling. Complementation of this defect may not require the kinase activity of Src, as cell spreading is restored by re-addition of alleles encoding either wild-type or kinase-defective Src. Similarly, we and others have shown that the Src-dependent tyrosine phosphorylation of pp130^{cas} in fibroblasts in response to fibronectin binding (Vuori et al. 1996) does not require intrinsic Src kinase activity (Schlaepfer et al. 1997; P. Schwartzberg, unpubl.). Our results in transgenic mice now show that such kinase-independent signaling pathways for Src are physiologically important for the function of osteoclasts in intact animals and, furthermore, suggest that some of the defects in these cells may be related to such kinase-independent pathways as integrin signaling.

Therefore, Src^{K295M} may provide signaling function in

some pathways and act in other pathways as a dominant-negative block to signal transduction. These differing actions of Src may in part account for the variability we observe in the phenotype of the rescued mice. In addition to beneficial effects of the transgene on osteoclast function, there may be detrimental effects; for example, high levels of expression of the *src*^{K295M} allele might block kinase-dependent signaling pathways as observed in other cell types and therefore prevent full rescue of the osteopetrosis. In cultured osteoclasts from these transgenic mice, our average levels of expression are similar to that of endogenous Src, potentially limiting adverse effects from overexpression of the kinase-dead allele. Alternatively, the mixed genetic background of the rescued mice (129/Sv/Ev × C57BL6/J × FVB/N) may partially account for the variation observed. Additional backcrossing to pure genetic strains should help resolve these issues.

Parallels with other tyrosine kinases

The related Src family member, Lck, has also been thought to have both kinase-dependent and kinase-independent signaling pathways in an isolated T lymphocyte cell line (Xu and Littman 1993). In the studies of Lck, however, kinase-independent functions were demonstrated with molecules fused to the upstream effector CD4 in cells that contained endogenous Lck. Recent evidence suggests that endogenous Lck may be required for the signaling by kinase-inactive Lck fusion proteins and that a multimeric complex may be involved in this pathway (Lee-Fruman et al. 1996). In *src*^{-/-} mice rescued with the *TRAPsrc*^{K295M} transgene, there is no endogenous Src. Furthermore, the kinase-independent rescue that we observe in osteoclasts was achieved without additional alteration of the Src molecule, suggesting that the kinase-inactive Src molecule itself can provide function. Our results more closely resemble the partial rescue observed with kinase-inactive D-abl of the *Dab*^{-/-} phenotype in *Drosophila melanogaster* (Henkemeyer et al. 1990). However, unlike our observations with *c-src*, a transgene-encoding kinase-inactive *c-abl* does not complement the *abl*^{-/-} phenotype in mice (Hardin et al. 1996).

Src functions in the osteoclast

Osteoclasts from *src*^{-/-} mice lack ruffled borders and normal organization of the actin cytoskeleton, and they do not resorb bone, but the biochemical and physiological defects that account for this complex phenotype remain unclear. Our findings with the *TRAP-src*^{K295M} mice suggest that Src may have multiple roles within the osteoclast. Proper intracellular transport, exocytosis, cell polarization, adhesion, migration, and cytoskeletal organization are all candidates for functions that might be impaired by defective signaling in *src*^{-/-} cells. It is clear that at least one pathway that requires Src function is that involved in cytoskeletal organization (Neff et al. 1996).

Restoration of osteoclast function by kinase-defective versions of Src suggests that at least some critical functions of Src in these cells may be related to other properties, such as the protein interaction and localization functions of the SH2 and SH3 domains. Complementation of the adhesion defect in *src*^{-/-} fibroblasts by a truncated version of Src requires intact SH2 and SH3 domains (Kaplan et al. 1995). These domains have been demonstrated to be important both for interactions with other molecules and for proper redistribution of Src protein mutants to focal adhesions on activation in fibroblasts (Pawson and Gish 1992; Kaplan et al. 1994). Furthermore, the increase in total tyrosine phosphorylation in osteoclasts rescued by kinase-inactive Src suggests that a critical function of Src may be in part to act as an adaptor molecule that recruits or activates other tyrosine kinases and their substrates. Particular targets for activation by Src include FAK and the related kinase Pyk2 in the osteoclast (Lev et al. 1995). These kinases may, in turn, phosphorylate downstream targets crucial for osteoclast function, such as the recently described Cbl molecule (Tanaka et al. 1996) or integrin-stimulated substrates, such as Cas (Vuori et al. 1996). It is interesting to note that integrin engagement is required for proper bone-resorption by osteoclasts (Davies et al. 1989); therefore, integrin-mediated signaling may be one pathway that is abnormal in *src*^{-/-} osteoclasts. Elucidation of such pathways may help delineate the functional abnormalities in osteoclasts and in other cells that lack the Src protein.

Materials and methods

Plasmid construction

The plasmid pKB5 containing the TRAP promoter was a kind gift from Drs. S. Reddy and G.D. Roodman (Reddy et al. 1995). The *TRAPsrc* plasmid was constructed by fusing the *src* ATG to the ATG of *TRAP* by PCR using primers TRAP1 (CCACCTC-CATGGTAGTGCC), Src1 (GTCTTGTTGGGGTCTGCG), TRAPSRC1 (CCCTCTCTTCCACAGATGGGGAGCAG CAAGAGCA and SRCTRAP (TGCTCTTGCTGCTCCCCATC TGTGAGGAAGAGAGGG). A 148-bp fragment was cloned to the Sph1 site of the *TRAP* promoter and the *NaeI* site of the chicken *c-src* gene using a three-fragment ligation. A *Bam*HI-*NotI* fragment containing the polyA site from plasmid pGKTK-polA was introduced into the *Bgl*II to *NotI* site of the *TRAPsrc* construct. The K295M mutant and Y416F mutant were rederived by PCR and subcloned into the *TRAPsrc* construct.

Transgenic mice

Transgenic mice were generated by standard pronuclear injection methods into FVB/N strain. Tail DNA was isolated and mice genotyped by PCR using primers TRAP1, SRC 1, and TRAP2 (GTGGAATGGGGCATTGGGG) for the *TRAPsrc* alleles. F₁ mice were bred to *src*^{+/-} mice (129/Sv/Ev × C57Bl6/J) generously provided by Dr. P. Soriano. Primers used for genotyping the endogenous *c-src* allele were SrcF (CAGCAA-CAAGAGCAAGCCCAAGGACG), NeoS (CAGTCATAGCC-GAATAGCCTCTCCACC), NSrc3 (GGGAGGTGACGGTGT CCGAGGA). These primers differ from primers published previously to eliminate spurious amplification from the *TRAPsrc*

allele. Products were visualized on a 10% polyacrylamide TBE gel.

Histology

Bones (calvaria, fore- and hindlimbs, and lumbar vertebrae) were fixed in 10% buffered formalin, decalcified in 14% EDTA, and embedded in paraffin. The sections were stained both with hematoxylin and eosin and for TRAP activity and the bone volume (amount of bone matrix as a percentage of the cancellous space), and osteoclast numbers (expressed per millimeter of the cancellous bone surface) in the cancellous space of the long bones were measured, as described previously (Boyce et al. 1992). For histology, a minimum of four samples were examined for each transgene. Histology was performed on samples from at least two transgenic lines for each construct except for the *TRAPsrc*^{Y416F} rescued mice where histomorphometry was performed on line G only.

IL-1 treatment

Human recombinant IL-1a (a gift from Dr. R. Chizzonite, Hoffman LaRoche, Nutley, NJ) (0.2 μg in 10 μl) was injected four times daily for 3 days into the loose subcutaneous tissue overlying the calvaria of the mice. Whole-blood-ionized calcium was measured in retro-orbital venous samples using a Ciba Corning 634 ISE Ca⁺⁺/pH analyzer (Medfield, MA) before and 2 hr after the last IL-1a injection. A minimum of three mice per group were examined. Results from the different sets of animals were compared using unpaired (two-tailed) student *t*-tests.

Cell culture

Osteoclasts were generated either by plating bone marrow directly in the presence of vitamin D3 (10⁻⁹ M) for 7 days (Takahashi et al. 1988a) or by plating with primary calvarial osteoblasts derived from *src*^{-/-} mice for 5–7 days (Takahashi et al. 1988b). Newborn litters of *src*^{+/-} intercrosses were genotyped by PCR at birth and calvaria cells from the *src*^{-/-} progeny were isolated as described (Takahashi et al. 1988b).

RNA analysis

Total cellular RNA was isolated using RNazol (Tel-test) from osteoclasts derived from bone marrow cultured in the presence of vitamin D3. RNA was reverse-transcribed (cDNA cycle Kit, Invitrogen) and transgene-specific primers, TRAP4 (TGCAGCC TCTCTGACCACCTGTGC) and SRC1, were used to amplify a fragment extending from the TRAP first exon to the chicken *c-src*-coding region. The TRAP1 primer was included in the same reaction to amplify a control TRAP-specific band. Products from amplification were visualized on a 10% polyacrylamide TBE gel and stained with ethidium bromide. The gel was transferred to Zetaprobe membrane (Bio-Rad) at 0.4 amps in 0.5× TBE, fixed to the membrane with UV and hybridized with the P³²-labeled kinased oligonucleotide probe TRAPSRC1. This probe recognizes the *TRAPsrc*-specific band as well as unspliced TRAP RNA. For Northern analysis, 30 μg of total RNA was analyzed on 1.2% agarose formaldehyde gels, transferred to Hybond-N and hybridized to a 200-bp probe corresponding to the first 200 bp of coding sequence of chicken *c-src*. Filters were stripped and reprobated according to the manufacturer's recommendations.

Protein

Osteoclasts from bone marrow cultures were partially purified by washing off the osteoblast layer with PBS. Cells were lysed in RIPA buffer (50 mM HEPES, 1% dexoycholic acid, 1% Triton, 0.1% SDS, 150 mM NaCl, 1 mM EDTA plus AEBBSF, leupeptin, and pepstatin). Extracts were normalized for both protein concentration (Biorad) and by TRAP enzymatic activity (Sigma) assayed at 405 nm. Equivalent amounts of protein were immunoprecipitated with 0.5 µg monoclonal 327 (Lipsich et al. 1983) (Oncogene Science) and anti-mouse immunoglobulin-agarose beads (Sigma), washed extensively with RIPA and then washed three times with kinase buffer (20 mM Tris at pH 7.4, 5 mM MnCl₂). Reactions were incubated in kinase buffer containing ~2 µg acid-denatured enolase for 5 min at room temperature, as described previously (Kaplan et al. 1995). Half of the immunoprecipitation was analyzed by 10% SDS-PAGE, transferring to nitrocellulose and blotted using either monoclonal 327 or Src2 sera (Santa Cruz). Phosphotyrosine-containing proteins were examined in normalized whole-cell extracts from osteoclasts purified from co-cultures in which the osteoblast feeder layer was removed completely by careful washing with PBS. Under these conditions, the majority of remaining cells were multinucleated osteoclasts, as determined by light microscopy. Only highly purified cultures of osteoclasts were used in these experiments, and under these conditions, there was a good correlation between protein levels and TRAP enzymatic activity. After electrophoresis and blotting of cell extracts, phosphotyrosine-containing proteins were detected with the 4G10 anti-phosphotyrosine-monoclonal antibody (UBI).

Immunofluorescence

Osteoclasts derived from the coculture of bone marrow cells and osteoblasts were grown on serum-coated coverslips, partially purified and fixed with 10% formalin for 10 min at room temperature. Cells were either stained for TRAP enzymatic activity or dehydrated with acetone for 2 min, then stained with Rhodamine-phalloidin (Molecular Probes) according to the manufacturer's recommendation. Cells were visualized by confocal laser scanning microscopy (emission wavelength 568 nm) using a Biorad MRC 1024.

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