FGF signaling targets the pRb-related p107 and p130 proteins to induce chondrocyte growth arrest

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Introduction

Signaling through FGF receptors (FGFR)* plays a major role in development by eliciting a variety of responses in FGF-targeted cells, ranging from induction of proliferation and differentiation to growth inhibition. FGF signaling plays a role in bone development and affects the process of endochondral ossification, which accounts for the formation of most of the skeletal bones (Goldfarb, 1996; De Luca and Baron, 1999; Karsenty, 1999; Powers et al., 2000). In endochondral ossification, the initial formation of a cartilage anlage by the condensation of mesenchymal stem cells is followed by proliferation and differentiation of chondrocytes that provide the template for bone formation. Considerable evidence from human and mouse genetics indicates that longitudinal bone growth is regulated by FGF signaling. Gain of function mutations in FGFR3, which is expressed in growth plate chondrocytes, are responsible for several forms of human dwarfism including achondroplasia, whereas in mice, knockout of the FGFR3 gene causes overgrown and deformed long bones and unregulated FGF signaling produces a variety of chondrodysplasias (Deng et al., 1996; Naski et al., 1996; Webster and Donoghue, 1997; Karsenty, 1999).

In line with the genetic evidence that excessive FGF signaling impairs long bone growth (Naski and Ornitz, 1998), we previously showed that FGF signaling inhibited chondrocyte proliferation in vitro and in vivo (Sahni et al., 1999), suggesting that one major mechanism by which FGFs regulate bone development is through their inhibitory effect on chondrocyte proliferation. We also showed,
through the use of Stat1 knockout mice, that this phenomenon required the intervention of STAT1, a signal transducing molecule originally identified in the interferon pathway (Sahni et al., 1999). Furthermore, we showed that FGF signaling induced chondrocyte apoptosis both in vitro and in vivo, and this also required STAT1 function (Sahni et al., 2001). Although FGF induces a proliferative arrest through a STAT1-dependent process in chondrocytes, in most other cell types FGFs elicit increased proliferation (Basilico and Moscatelli, 1992). Presently, the basis for the FGF growth inhibition of chondrocytes, versus the FGF-proliferative effect in most other cell types, is not defined.

To gain information on the mechanisms through which FGF signaling induces growth arrest in chondrocytes, we sought to identify other downstream effectors of FGF-mediated growth inhibition. It had been previously shown that mice with targeted inactivation of two members of the retinoblastoma (Rb) gene family, p107 and p130, died at birth with developmental defects including defective endochondral bone development and abnormally high levels of chondrocyte proliferation (Cobrinik et al., 1996). The Rb family of pocket proteins includes three members (pRb, p107, and p130) that are known to be major regulators of cell cycle progression in essentially all cells and tissues (Ewen, 1998; Mulligan and Jacks, 1998). In the active underphosphorylated form, the Rb family proteins bind the E2F family of transcription factors, resulting in transcriptional inhibition of many cell cycle progression and DNA synthesis genes. Inactivation of Rb family proteins by phosphorylation allows freeing of E2Fs factors which then positively influence transcription of cell cycle progression genes (Dyson, 1998).

We have investigated the hypothesis that the high levels of chondrocyte proliferation observed in p107−/− p130−/− mouse embryos could have been due, at least in part, to an impaired response to the growth-inhibitory effects of FGF signaling. To this end, we have expressed in cultured chondrocytes the adenovirus E1A (E1A) and polyoma large T (PyLT)-Ag proteins, that are known to bind and inactivate Rb proteins (Dyson et al., 1990; Classon and Dyson, 2001), and studied the response to FGF treatment of chondrocyte micromass cultures and cartilaginous bone rudiments from pRb, p130, and p107 knockout mouse embryos. Here, we show that expression of E1A or PyLT in rat chondrosarcoma (RCS) chondrocytes produces cells that no longer respond to FGF with growth inhibition. Experiments with chondrocyte micromass cultures and bone rudiments from mutant embryos further revealed that p107 and p130, but not pRb, are required for the growth inhibitory effects of FGF. Our data demonstrate that p107 and p130 are essential downstream mediators of the FGF-induced growth arrest of chondrocytes.

**Results**

**FGF-induced growth inhibition of RCS cells is associated with dephosphorylation of Rb family proteins**

Treatment of RCS chondrocytes with FGF-1 induces a potent growth inhibitory response (Sahni et al., 1999; see Fig. 2 A). FACScan™ analysis revealed that FGF-1 treatment of RCS cells promotes an accumulation of cells in the G0/G1 phase of the cell cycle (Fig. 1 A). Because Rb proteins constitute major regulators of the G1 to S phase transition (Ewen, 1998), we examined the phosphorylation status of these proteins as a read-out of their activities. This was done by Western blotting and immunodetection of the different phosphorylated forms of the Rb proteins, based on their characteristic electrophoretic mobilities. Treatment of RCS cells with FGF-1 for different times over a 24-h period showed that all three Rb family members (pRb, p107, and p130) became dephosphorylated after FGF treatment (Fig. 1 B). The kinetics of dephosphorylation was rapid, particularly in the case of p107, which is totally dephosphorylated as early as 9 h after FGF treatment. After 18 h of FGF treatment, all three Rb proteins were mainly present in their underphosphorylated forms. The G1 arrest induced by FGF-1 in RCS chondrocytes is reversible. Resumption of cell proliferation is accompanied by an increase in Rb proteins phosphorylation (unpublished data).

Because Rb proteins phosphorylation is mediated by Cdk, whose activity in turn is regulated by several Cdk inhibitors, we measured the expression of several members of the Cip/Kip and Ink4 families of Cdk inhibitors in RCS cells after FGF treatment. As previously shown by us and others (Sahni et al., 1999; Aikawa et al., 2001), the expression of p21 was up-regulated rather rapidly after FGF addition. On the other hand, expression of p27 and p57 was increased slightly and at later times. No changes in the expression of p16 and p18 were detected (Fig. 1 C).

**Inactivation of Rb family proteins abolishes FGF-mediated growth inhibition of RCS cells**

The rapid dephosphorylation of Rb proteins after FGF treatment suggested that their dephosphorylation could be a cause and not an effect of the growth arrest. To investigate whether the function of Rb proteins was required for FGF-induced growth arrest of RCS cells, we expressed by retroviral-mediated gene transfer two viral oncoproteins, PyLT antigen and E1A, which are known to interact directly with Rb proteins, and as a consequence, inactivate their function (Classon and Dyson, 2001). We measured the frequency of DNA-synthesizing cells by counting cells that had incorporated BrdU from 18 to 24 h after FGF treatment, and compared parental RCS cells with RCS that stably expressed PyLT or E1A. As shown in Fig. 2 A, expression of PyLT or E1A into RCS cells led to a marked decrease in FGF-induced growth inhibition in comparison with wild-type or vector-transfected RCS cells. In the case of PyLT-expressing RCS cells, we further confirmed that the early effects of FGF signaling were unaffected by the introduction of PyLT protein as judged by FGF-induced tyrosine phosphorylation of FGFR3 (the major FGF receptor expressed in these cells) as well as tyrosine phosphorylation of several endogenous proteins (unpublished data).

To confirm that the Rb family proteins binding ability of PyLT or E1A was implicated in the loss of FGF response, we used well defined mutants of these proteins carrying either a specific deletion or a point mutation of the Rb binding site (PyLT-Δ141–158 and E1A-125,928, respectively; Wang et al., 1993; Pilon et al., 1996), and having impaired binding to each of the pRb family proteins. As shown in Fig. 2 A, expression of these mutants did not affect the growth inhibitory re-
response observed in FGF-treated RCS cells, indicating that inactivation of Rb proteins by PyLT or E1A was responsible for the lack of FGF response. To exclude the possibility that the effect produced by the expression of PyLT and E1A could be triggered by the introduction of any oncogene into RCS cells, we also expressed another unrelated oncogene, an activated form of H-ras (Dotto et al., 1985), in these cells. Proliferation of v-H-ras–expressing RCS cells was inhibited by FGF to the same extent as RCS-treated cells (unpublished data).

Next, we wanted to exclude the possibility that PyLT- or E1A-expressing RCS cells were resistant to all growth inhibitory signals. Dexamethasone, a glucocorticoid that has been shown to inhibit chondrocyte growth both in vitro and in vivo (Jux et al., 1998; Klaus et al., 2000), induced a dose-dependent inhibition of proliferation of RCS cells (Fig. 2 B). Importantly, PyLT-expressing RCS cells treated with dexamethasone were comparably growth inhibited. Thus, the expression of PyLT does not make RCS cells refractory to all growth inhibitory signals.

Inactivation of pRb is not sufficient to block FGF-mediated growth inhibition of RCS cells

As discussed in the Introduction, studies of knockout mice have indicated an important role of the pRb-related proteins (p107 and p130) during chondrogenesis and particularly in the growth control of chondrocytes. To determine whether pRb, p107, and p130 equally contributed to FGF-induced growth arrest, we took advantage of a PyLT deletion mutant (PyLT–H9004256–272) that has been described to possess a reduced ability to bind p107, while maintaining both the potential to interact with pRb and to immortalize fibroblasts in vitro (Pilon et al., 1996). Because the binding characteristics of this mutant were determined in vitro using recombinant proteins, we first asked whether this PyLT mutant behaved similarly when expressed in cells. We expressed PyLT–Δ256–272 in RCS cells and examined the interaction of this mutant with Rb proteins in coimmunoprecipitation experiments. For these experiments as well as for the proliferation experiments, we selected clones
expressing similar amounts of PyLT and PyLT-H9004 256–272 proteins, as judged by Western blot analysis (Fig. 3 B, inset). As shown in Fig. 3 A, immunoprecipitation of both p107 and p130 revealed a strong reduction in the amount of PyLT-H9004 256–272 in the immune complexes (Fig. 3 A, lanes 3), in comparison with wild-type PyLT (Fig. 3 A, lanes 2). On the other hand, immunoprecipitation of pRb produced bands of similar intensity for both wild-type and mutant PyLT. Thus, these results were in agreement with the published data regarding the pRb/p107 binding abilities of the mutant PyLT-H9004 256–272 protein (Pilon et al., 1996). They also revealed that the ability of PyLT-H9004 256–272 protein to interact with p130 is affected in parallel with its ability to bind p107, a finding perhaps predictable given the extensive protein sequence similarities between p130 and p107.

BrdU incorporation assays (Fig. 3 B) showed that PyLT-H9004 256–272–expressing RCS cells were growth-inhibited by FGF treatment in a dose-dependent manner, similarly to control RCS cells, whereas wild-type PyLT-expressing RCS cells were consistently resistant to FGF growth inhibition. The ability of a PyLT mutant that can bind pRb, but not p107 and p130, to block the FGF response indicates that inactivation of pRb alone is not sufficient to interfere with FGF-mediated growth inhibition of RCS cells, and that the FGF response is likely mediated through p107 and p130 function.

p107 and p130 mediate the FGF-induced growth arrest in micromass chondrocytes

Although the above analyses suggested the importance of p107 and p130 to FGF-induced arrest in RCS cells, they did not exclude the possibility that any one of the Rb family proteins was sufficient to mediate the FGF response. Moreover, because RCS are tumor cells, it was of interest to define the role of each of the Rb family proteins in the FGF response in nontransformed cells. Thus, to define the importance of p107, p130, or pRb to FGF-induced growth
To determine the effect of FGF on chondrocytes proliferation, day 3 cultures were treated with FGF for 24 h, and BrdU was added over the last 2 h. The cultures were stained simultaneously with an antibody against collagen II (to identify chondrocytes) and with an antibody against BrdU (to identify S phase cells), and the proportion of BrdU-positive chondrocytes was determined by confocal microscopy. This technique allows individual chondrocytes within the cartilage nodules to be examined by focusing on a single plane.

In wild-type cultures, FGF induced a 50% decrease in chondrocyte BrdU incorporation, with chondrocytes in the middle of the foci responding more strongly to FGF than those at the edge (Fig. 5). Interestingly, Rb−/− chondrocytes responded to FGF to the same extent as wild-type chondrocytes (Fig. 5), indicating that pRb was not needed for FGF to inhibit proliferation. Similarly, p107−/−;p130−/− chondrocytes underwent an FGF-induced growth arrest (Fig. 5), implying that a single wild-type p107 and p130 allele is sufficient for the full FGF response. In contrast, p107−/−;p130−/− chondrocytes displayed a partial inhibition (Fig. 5), suggesting that p130 can mediate, at least in part, the FGF growth arrest response. Strikingly, p107/p130 double mutants chondrocytes did not arrest in response to FGF. Indeed, neither did p107−/−;p130−/− nor p107−/− chondrocytes (Fig. 5), indicating that p107 was crucial to the FGF-induced growth arrest in this system.

**p107 and p130 mediate FGF-induced growth inhibition in mouse bone rudiments**

Next, we sought to determine whether p107 and p130 were needed for the FGF response not only in cultured chondrocytes, but also in chondrocytes that are within developing bones that have a normal cartilage architecture. For this purpose, we used organ culture of metatarsal bone rudiments of mice with targeted p107 and/or p130 inactivation. In brief, bone rudiments from E15 mouse embryos were isolated and cultivated in vitro for 24 h and then treated with FGF-1 or left untreated for an additional 24 h. BrdU was added to the culture medium for the last 6 h. The rudiments were processed for immunohistochemistry, and BrdU incorporation was evaluated by immunostaining of the rudiment sections.

The frequency of chondrocytes incorporating BrdU in the proliferative zone of untreated rudiments was similar (Fig. 6) in all the different genotypes that we examined. Metatarsals from double heterozygous mice (p107+/−;p130+/−) showed strong inhibition of DNA synthesis upon FGF treatment (Fig. 6), similar to that observed in wild-type bone rudiments (not depicted). FGF treatment produced essentially no inhibition of DNA synthesis in p107+/−;p130+/− rudiments, and a slight but reproducible decrease in p107+/−;p130+/− rudiments, similar to that observed in chondrocyte micromass cultures. FGF treatment of bone rudiments from p107/p130-null rudiments resulted in no inhibition in DNA synthesis, and actually caused a small but reproducible increase in the rate of proliferation (Fig. 6 B), suggesting that blocking the growth inhibitory effects of FGF signaling in these cells could perhaps unmask its growth stimulatory potential. These results show that in developing bone rudiments, removal of p107 activity essentially abolishes chondrocyte growth inhibition by FGF, and that chondrocytes become completely resis-
tant when p130 is also absent. pRb, which is not mutated in these embryos, fails to compensate for the loss of p107 and p130. Thus, p107 and p130 are the major mediators of FGF growth inhibition in growth plate chondrocytes.

Discussion

The results described in this paper indicate that two members of the Rb family of pocket proteins (p107 and p130), but surprisingly not the prototype of the family (pRb), are essential effectors of the FGF signal cascade in chondrocytes that leads to inhibition of cell proliferation. Expression of viral proteins such as E1A and PyLT-Ag, which bind and inactivate Rb proteins, essentially suppress the growth inhibitory response of RCS chondrocytes to FGF. Not only was this effect not manifested in cells expressing mutant forms of PyLT and E1A incapable of binding to all Rb proteins, but more specifically a PyLT mutant that is selectively impaired in binding to p107 and p130, but not to pRb, was also incapable of blocking the FGF response. Notably, the loss of Rb family proteins’ function did not affect the response to other growth inhibitory agents.

The most conclusive demonstration of the p107 and p130 role in mediating the FGF effect in cell proliferation was provided by experiments with chondrocyte micromass cultures or cartilaginous metatarsal bone rudiments from murine embryos with targeted mutations of pRb, p107, and p130. Although the absence of pRb had no apparent effect on the FGF response, the loss of either p107 or p130 resulted in almost no inhibition of chondrocyte proliferation. Chondrocytes lacking both p107 and p130 showed no growth inhibition in response to FGF, and actually a small but reproducible increase in the rate of proliferation. This effect could be due to an “unmasking” of the growth stimulatory effects of FGF when growth inhibition is removed, and is reminiscent of what we previously observed in bone rudiments from STAT1 knockout mice (Sahni et al., 1999).

Although our analysis indicates that pRb is dispensable for FGF-induced growth arrest, it is difficult at this point to determine whether p130 or p107 is the major mediator of the FGF response. Although the experiments performed with micromass chondrocytes seem to suggest that p107 plays the major role, it is clear from the deregulated proliferation of p107+/-;p130+/- cultures, the majority of the BrdU(+)/ nuclei were at the periphery of the foci, and that growth inhibition was more substantial for cells within the cores of the foci.
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sufficient to almost abolish the effect of FGF treatment, whereas loss of both is needed to substantially deregulate proliferation in vivo. Furthermore, other factors may counteract the action of FGF in vivo, which would have to be completely blocked to produce a distinct phenotype.

Although further work will be needed to resolve this issue, it is perhaps more interesting to ask why pRb does not seem to participate in the FGF response of chondrocytes. pRb is equally dephosphorylated in RCS cells upon FGF treatment, but alone it does not seem to be capable of affecting FGF-mediated growth inhibition. pRb may not be an important regulator of cell cycle progression in chondrocytes. It has been long suspected that the function of the Rb family proteins is overlapping, but also shows tissue specificity (Lipinski and Jacks, 1999). The loss of all three Rb proteins is necessary to fully immortalize fibroblasts (Dannenberg et al., 2000; Sage et al., 2000), but loss of pRb alone is sufficient to deregulate proliferation of liver, lens, neural, muscle, and pituitary cells (Clarke et al., 1992; Jacks et al., 1992; Lee et al., 1992). Similarly, the loss of pRb, but not of p107 or p130, inhibits late osteoblast differentiation in vitro (Thomas et al., 2001). On the other hand, in the Ba/F3 hematopoietic cell line, cell cycle progression appears to be mainly controlled by p130 (Hoshikawa et al., 1998). An alternative hypothesis is that our results could reflect a more specific role of p107 and p130 in the FGF response, such as if FGF signaling targeted functions of p107 and/or p130 that are not possessed by pRb. p107 and p130 are known to form complexes with E2F4 and E2F5, whereas pRb preferentially regulates E2F1, E2F2, and E2F3 (Sardet et al., 1995; Dyson, 1998; Classon and Dyson, 2001). E2F/p107 and p130 complexes bind specific DNA sequences and inhibit the transcription of targeted genes (Hurford et al., 1997). It is possible that FGF-mediated growth inhibition depends strictly on the repression of specific genes, which would be regulated by the E2F/p107' or p130 complexes. As another possibility, the p107- and p130-dependent inhibition of chondrocyte proliferation may depend upon the ability of these proteins to directly bind and inhibit Cdk2 kinase, a property that is not shared with pRb (Hannon et al., 1993; Classon and Dyson, 2001).

A further issue that is raised by the current work is whether the role of p107 and p130 in the FGF response is linked to their roles in chondrocyte differentiation. Analysis of p107−/−;p130−/− embryos clearly showed a block to chondrocyte hypertrophic differentiation (Rossi et al., 2002), and this defect was also observed in organ cultures of the metatarsal rudiments, regardless of FGF treatment. Sim-

Figure 6. Absence of FGF-induced growth inhibition in bone rudiment chondrocytes lacking p107 and/or p130. Metatarsal bone rudiments from E15 embryos were isolated and cultivated in vitro for 48 h with or without FGF treatment (100 ng/ml for 24 h). BrdU was added during the last 6 h of the culture. After fixation, rudiments were processed for immunodetection of BrdU. Counterstaining with Alcian blue was done in order to stain the cartilage-specific sulfate proteoglycans present in the extracellular matrix. (A) Representative pictures of BrdU incorporation in the proliferative zone of the growth plate for the different genotypes analyzed. R, reserve zone; P, proliferating zone; H, hypertrophic zone. (B) Quantification. Note that error bars in Fig. 6 B represent the SD calculated from at least six counts (percentage of cells positive for BrdU) of 80–160 cells each. Organ culture experiments were performed twice and gave similar results.
ilar to the in vivo situation, the appearance of hypertrophic chondrocytes could be easily detected in p107
\textsuperscript{-/-}/p130
\textsuperscript{-/-}
rudiments as well as in p130
\textsuperscript{-/-}/p107
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rudiments, but was rarely detected in p107
\textsuperscript{-/-}/p130
\textsuperscript{-/-}
rudiments and not detected at all in rudiments from p107
\textsuperscript{-/-}/p130
\textsuperscript{-/-}
embryos (data not shown). As shown recently (Rossi et al., 2002), the inhibition of chondrocyte differentiation observed in p130/p107 knockout embryos could be due to the impaired expression of Cbfα 1, a transcription factor that is important for chondrocyte hypertrophic differentiation (Inada et al., 1999; Kim et al., 1999; Takeda et al., 2001). Whether this is in turn related to the resistance of these cells to FGF treatment is currently unclear. The question of whether FGF treatment accelerates or inhibits chondrocyte terminal differentiation and whether this effect may require cell cycle arrest and the intervention of the Rb proteins is presently undergoing investigation.

Finally, our results are consistent with a relatively simple mechanism by which FGF would mediate growth inhibition of chondrocytes. FGF treatment could, through the activation of STAT1, induce the expression of Cdk inhibitors, leading to inhibition of Rb family proteins phosphorylation. Indeed, it was previously shown that p21 is induced by FGF treatment in chondrocytes (Sahni et al., 1999; Aikawa et al., 2001). As shown here, the expression of p27 and p57 is also induced by FGF in chondrocytes, but weakly and with delayed kinetics. On the other hand, we found that p21 is strongly induced by FGF in many cell types where its major biological effect is growth stimulation, and not growth inhibition. Furthermore, p21
\textsuperscript{-/-} mice have not been reported as having a clear bone phenotype (Deng et al., 1995), and p21 deletion only partially reduced the ability of FGF to inhibit chondrocyte proliferation in organ culture (Aikawa et al., 2001). In addition to inducing expression of Cdk inhibitors, FGF signaling could also induce the expression or the activity of the phosphatases specifically targeting the Rb proteins. This activity might be particularly relevant to p107, because p107 appeared to undergo dephosphorylation with kinetics that were more rapid than was the accumulation of G1 cells (our unpublished results). In this regard, p107 dephosphorylation has been reported to be rapidly induced by DNA damaging agents (Voorheoeve et al., 1999; Kondo et al., 2001), suggesting that rapid p107 dephosphorylation might similarly be induced in FGF-treated chondrocytes. At this time, we believe that induction of phosphatases and Cdk inhibitors could participate, probably together with other mechanisms, in producing the profound growth-inhibiting effect of FGF in chondrocytes.

### Materials and methods

**Cell culture and proliferation assay**

RCS cells were cultured on 100-mm cell culture dishes (Corning Incorporated) in DME containing 10% FCS. For proliferation assays, cells either cultivated on glass coverslips or Lab-Tek® chamber slides (Nunc) were incubated for 3 h with 10
\textsuperscript{-4} M BrdU and were fixed with 3.7% PFA. In addition to inducing expression of Cdk inhibitors, FGF signaling could also induce the expression or the activity of the phosphatases specifically targeting the Rb proteins. This activity might be particularly relevant to p107, because p107 appeared to undergo dephosphorylation with kinetics that were more rapid than was the accumulation of G1 cells (our unpublished results). In this regard, p107 dephosphorylation has been reported to be rapidly induced by DNA damaging agents (Voorheoeve et al., 1999; Kondo et al., 2001), suggesting that rapid p107 dephosphorylation might similarly be induced in FGF-treated chondrocytes. At this time, we believe that induction of phosphatases and Cdk inhibitors could participate, probably together with other mechanisms, in producing the profound growth-inhibiting effect of FGF in chondrocytes.

**Preparation of chondrocyte micromass cultures**

To generate embryos of the desired genotypes, matings were conducted between p107
\textsuperscript{-/-}/p130
\textsuperscript{-/-} and p107
\textsuperscript{-/-}/p130
\textsuperscript{-/-} mice or by intercrossing p107
\textsuperscript{-/-} or Rb
\textsuperscript{-/-} mice, and were assumed for timing purposes to occur at midnight. Micromass cultures were performed as in Edwall-Arvindsson and Wrede-Bergsten (1996) with the following modifications: forelimb and hindlimb buds were removed from E11.5 embryos and pooled in an Eppendorf tube. After two washes in PBS, the limbs were centrifuged for 1 min at 3,000 rpm in a microcentrifuge, resuspended in 1 mg/ml collagenase A (Roche) in Ham’s F12 medium and incubated at 37°C for 1 h. The tissues were gently triturated to obtain a single cell suspension, washed two times in Ham’s F12 medium, and resuspended at 1.0 × 10^5 cells/ml in Ham’s F12 medium supplemented with 1% FCS (HyClone). High density (supraconfluent) micromass cultures were initiated by spotting 20-μl drops onto the surface of tissue culture dishes (6-well dish; Nunc), or onto a coverslip for experiments requiring confocal microscopy, and incubated at 37°C for 2 h to allow the cells to attach. Ham’s F12 medium containing 1% FCS and antibiotics was then added to the culture. The medium was changed after 2 days of culture or just before FGF treatment. At various times after plating, micromass cultures were washed in PBS and fixed for 30 min in 4% PFA in PBS. To determine the accumulation of extracellular cartilage matrix, fixed cultures were stained overnight with incubation with 0.5 μg/ml Hoechst (Sigma-Aldrich) before mounting on slides. Pictures were taken with a microscope (Axioplan 2; Carl Zeiss Microimaging, Inc.) equipped with a digital camera.

For FACScan™ analysis, trypan blue cells were fixed in 70% ice-cold ethanol. After centrifugation, the cell pellets were washed with PBS and treated overnight with 50 μg/ml propidium iodide in the presence of 100 μg/ml RNase in order to stain the nuclei. Flow cytometry was performed using a FACScan™ (Becton Dickinson) equipped with a doublet discrimination module. Doublets were gated out on a plot of signal area versus signal width and analyzed using ModFit LT™ (Verity Software House).

For proliferation assays, FACScan™ analysis, and biochemical assays, RCS cells cultivated in DME containing 10% FCS were treated with recombinant human FGF-1 at the indicated concentrations together with 10 μg/ml heparin. Dexamethasone was purchased from Sigma-Aldrich.

**Cloning and retroviral infections**

PyLT antigen cDNA cloned in the pBabe-Puro retroviral expression vector was described previously (Mansukhani et al., 2000). This vector contains a resistance gene for puromycin. Mutant PyLT cDNAs (PyLT-D141–158 and PyLT-D256–272, a gift from Dr. A. Mes-Masson, McGill University, Montreal, Canada), as well as wild-type E1A and mutant E1A (E1A-∆S2-928) cDNAs were subcloned in pBabe-Puro. For retroviral infection, 293T cells were first transfected by calcium phosphate precipitation method by mixing equal amounts of retroviral construct together with helper virus vector pCL-eco (Naviaux et al., 1996). 48–72 h after transfection, supernatant was collected and used to infect RCS cells for 16 h in the presence of 10 μg/ml polybrene. Cells were cultivated for two additional days before selection with 2 μg/ml puromycin. PyLT- and PyLT-D256–272-expressing RCS cells were further subcloned and at least two clones of each line were analyzed.

**Immunoprecipitation and Western blotting**

Cell lysis was done either in RIPA buffer or immunoprecipitation lysis buffer (50 mM Tris, pH 7.5, 250 mM NaCl, 0.1% Triton X-100, 1 mM EDTA) in the presence of a cocktail of protease and phosphatase inhibitors. For immunoprecipitation assays, 1 mg of total protein extracts was incubated overnight at 4°C with 10 μg antibodies. Protein G Plus–Agarose (Oncogene Research Products) was added the next day for 2 h at 4°C, and the immune complexes were washed three to four times in immunoprecipitation lysis buffer. Immunoprecipitates and total protein extracts (20 μg) were separated by SDS-PAGE, transferred onto nitrocellulose membranes (Protran; Schleicher & Schuell) and analyzed by ECL detection system (Amer sham Biosciences). The following antibodies were used: anti-pRb mAb (G3–245; BD Biosciences), anti-p107 pAb (C-18; Santa Cruz Biotechnology, Inc.), p130 mAb (R27020; Transduction Laboratories), anti-p16, anti-p15, anti-p18, and anti-p57 pAbs (M-156, N-20, C-19, and H-91, respectively; Santa Cruz Biotechnology, Inc.), anti-p27 mAb (K25020; Transduction Laboratories), anti-Grb2 mAb (16G270; Transduction Laboratories), anti-PyLT antigen mAb (PN116; a gift from Dr. B. Schaffhausen, Tufts University Medical School, Boston, MA), anti-FGF3 pAb (C-15; Santa Cruz Biotechnology, Inc.), and anti-phosphotyrosine mAb (4G10; Upstate Biotechnology). HRP-conjugated anti-mouse and anti-rabbit secondary antibodies were purchased from Promega.
1% Alcian blue (Sigma-Aldrich) in 3% acetic acid (Leonard et al., 1991), and were washed for 6 h in 3% acetic acid.

Confocal microscopy analysis
Micromass cultures were washed in PBS and fixed in 100% methanol. After three washes in PBS, the cultures were blocked in 4% horse serum in PBS for 1 h. Antibodies anti-BrdU (Beckton Dickinson) at 1/100 dilution, anti–collagen II (Santa-Cruz Biotechnology, Inc.) at 1/100 dilution, or anti–FGFR3 (sc-123; Santa Cruz Biotechnology, Inc.) at 1/100 dilution were added in 4% horse serum in PBS, at 4°C overnight. After three washes with PBS, the secondary antibodies (anti–mouse Cy3, anti–goat FITC, and anti–rabbit Cy3; Jackson ImmunoResearch) were added at 1/2000 dilution, for 45 min at RT. After three washes with PBS, the nuclei were stained with DAPI, and the coverslip was mounted on a glass slide with Vectashield (Vector Laboratories). Foci of ~20 μm diameter were selected and analyzed over a series of 1 μm confocal sections using a confocal laser scanning microscope (model LSM410; Carl Zeiss Micro imaging, Inc.). The percentage of BrdU-positive cells was determined by analysis of each confocal section. The results are a mean of at least five foci for each genotype and each condition.

Organ cultures
Metatarsal long bone rudiments from E15 mouse embryo (generated as described in the chondrocyte micromass section) heterozygotes or nulls for the p107 and/or p130 genes were dissected under sterile conditions. The cartilaginous long bones were left intact. Organ culture medium consisted of MEM without nucleosides (GIBCO BRL) supplemented with 50 μg/ml ascorbic acid, 300 μg/ml l-glutamine, 50 μg/ml gentamicine, 250 μg/ml Fungizone, 1 mM β-glycerophosphate, and 0.2% BSA Cohn fraction V (complete medium; Sigma-Aldrich). Each long bone was cultured individually in Transwells® (Costar) containing 400 μm diameter were selected and analyzed for endogenous TGF-beta-like activity.

We wish to thank Lisa Dailey, Alka Mansukhani, and H. Skopicki for advice and helpful discussion, Milton Gomez for help with the histology preparations, and Anne Marie Mes-Masson and Paolo Dorato for the gift of plasmids. This investigation was supported by National Institutes of Health grants DE13745 (to C. Basilico) and HD35156 (D. Cobrinik), The Columbia University Optical Microscopy Facility is supported by grant P30CA13696.

Submitted: 7 May 2002
Revised: 14 June 2002
Accepted: 18 June 2002

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