ORIGINALE ARTICLE

Loss of the xeroderma pigmentosum group B protein binding site impairs p210 BCR/ABL1 leukemogenic activity

NL Pannucci1, D Li1, S Sahay1, EK Thomas2, R Chen1, I Tala1, T Hu1, BT Ciccarelli1, NJ Megjugorac1, HC Adams III1, PL Rodriguez1, ER Fitzpatrick1, D Lagunoff1, DA Williams3 and IP Whitehead1

Previous studies have demonstrated that p210 BCR/ABL interacts directly with the xeroderma pigmentosum group B (XPB) protein, and that XPB is phosphorylated on tyrosine in cells that express p210 BCR/ABL1. In the current study, we have constructed a p210 BCR/ABL1 mutant that can no longer bind to XPB. The mutant has normal kinase activity and interacts with GRB2, but can no longer phosphorylate XPB. Loss of XPB binding is associated with reduced expression of c-MYC and reduced transforming potential in ex-vivo clonogenicity assays, but does not affect nucleotide excision repair in lymphoid or myeloid cells. When examined in a bone marrow transplantation (BMT) model for chronic myelogenous leukemia, mice that express the mutant exhibit attenuated myeloproliferation and lymphoproliferation when compared with mice that express unmodified p210 BCR/ABL1. Thus, the mutant-transplanted mice show predominantly neutrophilic expansion and altered progenitor expansion, and have significantly extended lifespans. This was confirmed in a BMT model for B-cell acute lymphoblastic leukemia, wherein the majority of the mutant-transplanted mice remain disease free. These results suggest that the interaction between p210 BCR/ABL1 and XPB can contribute to disease progression by influencing the lineage commitment of lymphoid and myeloid progenitors.

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INTRODUCTION

BCR/ABL fusion proteins are the products of reciprocal translocations that are causally associated with Philadelphia chromosome-positive leukemias. Depending upon the position of the breakpoint within the BCR locus, different fusion proteins are generated, resulting in different clinical outcomes. Thus, p210 BCR/ABL1 is responsible for virtually all cases of chronic myelogenous leukemia (CML), whereas p190 BCR/ABL is associated with a subset of ALL.1 Although there is a general agreement that the tyrosine kinase activity residing within the ABL component of BCR/ABL is the principle driving force behind Philadelphia chromosome-positive leukemias, domains that are contained within the BCR sequences are also required for transformation.2–6 Efforts to understand the genetic instability, which usually accompanies progression of CML from the chronic phase to the blast phase, indicate that DNA repair pathways, such as homologous recombination repair, non-homologous end joining and mismatch repair, may be altered in hematopoietic cells expressing p210 BCR/ABL1.6–8 Such cells are resistant to apoptosis induced by chemotherapeutic agents and γ-irradiation,9–13 despite exhibiting a tendency to accumulate more DNA damage.14 Although the pathways downstream of p210 BCR/ABL1 responsible for this phenotype have not been clearly defined, there is evidence suggesting that p210 BCR/ABL1-positive cells may have enhanced DNA repair capability.15 Some of the DNA damage observed may therefore actually represent intermediates in an accelerated process of repair. Independent studies have also revealed a role for p210 BCR/ABL1 in the regulation of nucleotide excision repair (NER),15,16 a mammalian DNA repair system that removes a wide range of structurally unrelated lesions, including those induced by UV radiation and alkylating agents.17 In myeloid cells, the overexpression of p210 BCR/ABL1 increases NER activity and decreases UV electromagnetic radiation subtype C (UVC)-mediated cytotoxicity,15,16 whereas in lymphoid cells the overexpression results in decreased NER activity and increased UVC-mediated cytotoxicity.15 Although an exact mechanism for these changes in NER is not known, it has been shown that both BCR and p210 BCR/ABL1 bind to an essential protein in the repair process: xeroderma pigmentosum group B (XPB).18,19 XPB is a 3’–5’ DNA helicase with associated ATPase activity that forms part of the core subunit of the transcription factor TFIIH,7,20,21 and is required for NER and transcriptional initiation.22 Whether or not the interaction between XPB and p210 BCR/ABL1 is responsible for altered NER and whether it supports disease progression has not yet been determined. In this study we show that a p210 BCR/ABL1 mutant lacking the XPB-binding site is attenuated in its ability to drive myeloproliferation and lymphoproliferation in murine models for CML and B-cell acute lymphoblastic leukemia (B-ALL), but not in its ability to regulate NER.

MATERIALS AND METHODS

Molecular constructs and yeast two-hybrid analysis

The pAX142 mammalian expression vector and pAX142-bcr-abl have been previously described.23 pAX142-xpb contains a full-length cDNA for human XPB. pAX142-bcr-ablΔ674–695 encodes full-length,
hemagglutinin-epitope-tagged p210 BCR/ABL1, with an internal deletion of residues 674–695. The yeast two-hybrid constructs pGBT9-bcr(1–1271), pGBT9-bcr(1–413), pGBT9-bcr(491–880) and pGBT9-bcr(871–1271) have been previously described.24 pGBT9-bcr(491–668), pGBT9-bcr(491–681), pGBT9-bcr(491–691), pGBT9-bcr(491–700) and pGBT9-bcr(491–727) contain cDNAs that encode the indicated residues of BCR. pGBT9-bcr(674–695) encodes full-length BCR with an internal deletion of residues 674–695, pGBT9-dbs and pGBT9-ect2 contain full-length cDNAs for murine Dbs and Ect2, respectively. The yeast two-hybrid constructs for full-length MYC (PAG-medyc), XBP (pGAD-xbp) and ubiquitin (pGAD-ubq) have been previously described.23,24 The MSCV-IRES-gfp retroviral vector has been previously described.25 IRES-gfp, MSCV-bcr-abl/p210-ires-gfp and MSCV-bcr-ubq/p210(674–695)-ires-gfp contain full-length p210 BCR/ABL1 and the p210 BCR/ABL1 XBP-binding mutant, respectively. The pCLeco helper plasmid26 was kindly provided by Dr Sagi Ghaffari. All yeast-two-hybrid analysis was performed as previously described.24

RESULTS

Localization of the docking site for XBP within BCR and p210 BCR/ABL1

In a previous report, yeast two-hybrid analysis was used to demonstrate an interaction between full-length XBP and a fragment of BCR (residues 413–789).19 We used a similar approach to more precisely map the docking site for XBP to residues 681–691 of BCR (Figure 1a). A full-length BCR(674–695) mutant still interacts with two other BCR-binding partners, MYC24 and ubiquitin,23 thus confirming its structural integrity (Figure 1b). XBP does not interact with two other RhoGEF family members (Ect2 and Dbs, Figure 1c), confirming the specificity of the interaction. As the docking site for XBP is retained within p210 BCR/ABL1,19 a mutant lacking the putative XBP-binding site (p210 BCR/ABL1(674–695)) was constructed in a mammalian expression vector. Both mutant and wild-type p210 BCR/ABL1 were then co-expressed with XBP in 293T cells, and a coimmunoprecipitation assay was performed (Figure 1d). Whereas we are readily able to detect an interaction between XBP and p210 BCR/ABL1, only a weak interaction is observed between XBP and the mutant.

XBP binding is not required to support p210 BCR/ABL1 auto- and trans-kinase activity

To determine whether the tyrosine kinase activity of p210 BCR/ABL1 requires XBP binding, we expressed both wild-type and mutant proteins in 293T cells and performed western blot analyses to examine the phosphorylation levels of known substrates of p210 BCR/ABL1 tyrosine kinase activity.29 An equivalent level of phosphorylated CRKL was observed in cells that express p210 BCR/ABL1 and the mutant when compared with vector controls, suggesting that the trans-kinase activity is unaffected by loss of XBP binding (Figure 1e). Similarly, when we examined lysates with an antibody that recognizes the Tyr-245 autophosphorylated form of p210 BCR/ABL1, an equivalent level of auto-kinase activity was observed (Figure 1e). Next we determined whether p210 BCR/ABL1 and the mutant have an equivalent ability to interact with GRB2. It has been previously shown that p210 BCR/ABL1 autophosphorylates on Tyr-177, creating a docking site for GRB2.5 As shown by coimmunoprecipitation assays (Figure 1f), both p210 BCR/ABL1 and the mutant interact with GRB2 to an equivalent level. This indicates that the interaction with GRB2 does not require XBP binding and confirms that the auto-kinase activity of the mutant is unchanged.

p210 BCR/ABL1 can phosphorylate XBP on tyrosine in vivo

It has been previously shown that XBP is tyrosine phosphoryl-ated in cells expressing p210 BCR/ABL1.19 To determine whether this requires a direct interaction, we expressed full-length XBP along with either p210 BCR/ABL1 or the XBP-binding mutant in 293T cells. We then immunoprecipitated XBP and performed a western blot with antibodies that recognize the total or phosphorylated form of the protein. As shown in Figures 1g, a high level of phosphorylated XBP is detected in lysates that contain p210 BCR/ABL1, but not in cells that express the mutant.

The interaction with XBP does not influence the effects of p210 BCR/ABL1 on NER

In myeloid cells, overexpression of p210 BCR/ABL1 has been shown to increase NER activity and decrease UVC-mediated cytotoxicity.15,16 Whereas in lymphoid cells the overexpression results in decreased NER activity and increased UVC-mediated cytotoxicity.15 We examined whether the XBP-binding mutant can similarly influence NER. For this analysis, we cloned both p210...
then irradiated with UVC (10 J/m²) and Comet assays were performed (Figure 2b). Cells were selected by culturing in the presence of granulocyte colony-stimulating factor, stem cell factor and thrombopoietin. Cells were selected by culturing in the presence of granulocyte colony-stimulating factor, stem cell factor and thrombopoietin. Cells were then irradiated with UVC (10 J/m²) and Comet assays were performed (Figure 2b). In control cells, a significant increase in NER is observed by 1 h post irradiation. Cells that express p210 BCR/ABL1 exhibit significantly reduced NER relative to control cells, which is consistent with previous observations.15 Cells that express the mutant have levels of repair equivalent to cells that express p210 BCR/ABL1 at both 1 h and 3 h post irradiation. To examine NER in myeloid cells, bone marrow cells were collected from BALB/C mice and infected with retrovirus that contain p210 BCR/ABL1, p210 BCR/ABL1(Δ674–695), or cognate vector. Myeloid cells were selected by culturing in the presence of granulocyte colony-stimulating factor, stem cell factor and thrombopoietin. Cells were then irradiated with UVC (10 J/m²) and Comet assays were performed (Figure 2c). In control cells, a significant increase in NER was observed at 1 h post irradiation and, consistent with previous results, repair was significantly enhanced in cells that express p210 BCR/ABL1.15 An equivalent and significant increase in repair activity was also observed in cells infected with the binding mutant.

**Figure 1.** Mapping the XPB-binding site in BCR and BCR/ABL1. (a) Yeast two-hybrid mapping shows the XPB-binding site to be within residues 681–691 of BCR (*). Upper schematic shows the domain structure of the full-length BCR protein, whereas the lines below indicate the regions of the protein included in the cDNA derivatives used for mapping. The breakpoints in BCR for p190 BCR/ABL and p210 BCR/ABL1 are indicated by arrows. (b) BCR(Δ674–695) binds to c-MYC and ubiquitin, but not to XPB. Interactions between proteins are demonstrated by the ability to grow on histidine-deficient plates (Leu/Trp/His). (c) BCR does not interact with full-length Dbs or Ect2. (d) p210 BCR/ABL1(Δ674–695) is impaired in XPB binding. Cells were transiently co-transfected with full-length XPB and the indicated hemagglutinin-tagged BCR/ABL1 constructs. Lysates collected at 48 h were immunoprecipitated (IP) and/or examined by western blot (WB) analysis with the indicated antibodies (LC, loading control). (e) p210 BCR/ABL1(Δ674–695) has normal auto- and trans-phosphorylation activity and (f) interacts with GRB2. Cell lysates were collected at 48 h and examined by WB analysis. (g) The interaction with p210 BCR/ABL1 supports the phosphorylation of XPB on tyrosine. Lysates collected at 48 h were IP and/or examined by WB analysis with the indicated antibodies.

BCR/ABL1 and the mutant into MSCV-IRES-gfp. Retrovirus was then produced and was used to infect Ba/F3 cells. Consistent with the 293T cells, an equal and elevated level of phosphorylated CrkL is detected in lysates that contain p210 BCR/ABL1 or the mutant (Figure 2a), suggesting that the tyrosine kinase activity is unaltered. Cells that express p210 BCR/ABL1 also show elevated levels of endogenous, phosphorylated XPB and this is not seen in cells expressing the binding mutant. Cells were then irradiated with UVC (10 J/m²) and Comet assays were performed (Figure 2b). In control cells, a significant increase in NER is observed by 1 h post irradiation. Cells that express p210 BCR/ABL1 exhibit significantly reduced NER relative to control cells, which is consistent with previous observations.15 Cells that express the mutant have levels of repair equivalent to cells that express p210 BCR/ABL1 at both 1 h and 3 h post irradiation. To examine NER in myeloid cells, bone marrow cells were collected from BALB/C mice and infected with retrovirus that contain p210 BCR/ABL1, p210 BCR/ABL1(Δ674–695), or cognate vector. Myeloid cells were selected by culturing in the presence of granulocyte colony-stimulating factor, stem cell factor and thrombopoietin. Cells were then irradiated with UVC (10 J/m²) and Comet assays were performed (Figure 2c). In control cells, a significant increase in NER was observed at 1 h post irradiation and, consistent with previous results, repair was significantly enhanced in cells that express p210 BCR/ABL1.15 An equivalent and significant increase in repair activity was also observed in cells infected with the binding mutant.

Loss of XPB binding is associated with reduced expression of c-MYC. It has been shown previously that c-MYC is stabilized by p210 BCR/ABL1(ref 30) and is required for p210 BCR/ABL1 transformation.31 As c-MYC expression is known to be directly regulated by XPB,32,33 we also examined c-MYC expression in the Ba/F3 cells that stably express p210 BCR/ABL1, or the mutant (Figure 2d). As expected, the c-Myc levels were elevated in cells expressing p210 BCR/ABL1 relative to vector controls. In contrast, c-MYC levels were significantly diminished relative to vector controls in cells that express the mutant.

The interaction with XPB influences transformation in murine bone marrow ex-vivo assays. To explore the role of the XPB interaction in the transformation of murine hematopoietic progenitor cells, bone marrow was collected from BALB/C mice and infected with retrovirus that contain p210 BCR/ABL1, p210 BCR/ABL1(Δ674–695), or cognate vector. Bone marrow colony formation was assessed on media that supports growth of granulocyte–macrophage progenitors (GMP) (M3534), erythroid progenitors (BFU-E) (M3434) and B-cell progenitor cells (CFU-preB) (M3630). When we compare p210 BCR/ABL1, and the mutant, with vector on the M3434 media, we observe an equivalent number of BFU-E colonies (Figure 3a). In contrast, whereas p210 BCR/ABL1 shows enhanced growth of GMPs on M3534 media, the mutant is impaired in its ability to
Lung capillaries in the mutant mice contained numerous granulocytes, but there was considerably less hemorrhage than in the p210 BCR/ABL1 mice. Vector-transplanted mice (n = 20 total) had normal WBC counts (≥13,000/μl) and remained disease free through 6 months post BMT (not shown).

Immunophenotyping at different points after transplantation reveal differences in disease progression

To directly compare disease progression, three mice from each group (including vector) were killed at days 16 and 30 post BMT and immunophenotyping was performed (Figure 4). At day 16 post BMT, over 50% of WBCs in the p210 BCR/ABL1 mice were GFP-positive compared with ~25% for vector and mutant expressing mice (Figure 4a). Both the p210 BCR/ABL1 and mutant mice exhibited an approximately twofold increase in WBCs expressing the myeloid-specific marker, CD11b, relative to vector mice.

Surprisingly, however, over 30% of the WBCs in the p210 BCR/ABL1-transplanted mice expressed a B-cell marker (B220), whereas the B-cell counts in the mutant-transplanted mice did not exceed 3%. It is likely that this expansion of B cells in the p210 BCR/ABL1-transplanted mice accounts for the difference in total number of GFP+ cells. Histologic examination performed at day 16 post BMT revealed the beginning of disease progression with little significant difference between the mice (Supplementary Figure 1).

Comparison of mice on day 30 post BMT revealed more dramatic differences in disease progression. Greater than 95% of the WBCs from the p210 BCR/ABL1 mice were GFP-positive and 70–80% of the cells stained positive for myeloid markers (Figure 4b). In comparison, only 50% of the WBCs from mutant mice were GFP-positive and only 35% stained positive for myeloid markers.
markers. Expansion of the B-cell population was less apparent in the p210 BCR/ABL1-transplanted mice at day 30. Organ histology at day 30 also revealed differences between the groups, with mutant-transplanted mice having better preservation of liver and lung architecture than p210 BCR/ABL1-transplanted mice (Supplementary Figure 1).

Mice were also subject to immunophenotyping when they became moribund (Table 1). In order to determine whether there is a qualitative difference in myeloid expansion, cells were also examined for the expression of a granulocyte-specific marker (Gr1^+). At death, all of the mice have predominantly myeloproliferative disease, although most still have slightly elevated B-cell counts. Although the percentage of GFP^+ Gr1^+ cells is equivalent in all tissues examined, the percentage of cells that are GFP^+ Gr1^+ is significantly higher in mice transplanted with the mutant than in mice transplanted with p210 BCR/ABL1. This suggests that myeloid expansion in the mutant-transplanted mice is primarily restricted to neutrophils.

Loss of XPB binding alters progenitor expansion

In order to determine whether the impairment in disease progression and myeloid expansion could be attributed to differences in progenitor expansion, GFP-positive cells were examined from the bone marrow of p210 BCR/ABL1 and p210 BCR/ABL1(Δ674–695)-transplanted mice at death (Figures 5a and b). Surprisingly, the total number of progenitors is significantly increased in the mutant-transplanted mice, which is attributable to a large increase in GMP. Cell cycle analysis performed on the GMs, CMPs and MEPs revealed no significant difference in either proliferative potential or sensitivity to apoptosis (Figure 5c).

XPB-binding supports B-cell proliferation in a BMT model for B-ALL

As mice transplanted with the XPB-binding mutant did not show the early B-cell proliferation seen in p210 BCR/ABL1-transplanted mice, we determined whether the mutant could drive lymphoproliferation in a murine model for B-ALL. Consistent with previous reports, mice transplanted with p210 BCR/ABL1 exhibited B-cell lymphocytosis when killed at day 20 and 38 post BMT (Table 2, p210 BCR/ABL1 mice 1–6). Between days 37 and 75, 10 of 13 mice succumbed to disease (Figure 6a), and necropsies performed at death revealed characteristic signs of lymphadenopathy and moderate splenomegaly (spleen weight = 0.2–0.5 g).

Flow cytometry performed on tissues from randomly selected mice showed a predominance of GFP^+/B220^+ cells (Table 2, p210 BCR/ABL1 mice 7–10), a large proportion of which were IgM^+ /BP-1^−, suggesting immaturity (data not shown). On day 76, the survival study was terminated and the three remaining p210 BCR/ABL1 mice were killed and analyzed. One mouse showed clear signs of B-ALL (Figure 6b, Table 2, p210 BCR/ABL1 mice 11), one showed no evidence of disease (not shown) and one showed expansion of CD11b^−, B220^+ and CD3^− cells (not shown). In summary, out of 13 mice examined, 11 had immunophenotypes consistent with B-ALL.

In comparison with the p210 BCR/ABL1-transplanted mice, those transplanted with the XPB-binding mutant showed a significant increase in lifespan, with only one mouse dying within...
the time period covered by the survival study (Figure 6a, Table 2, p210 BCR/ABL1(Δ674–695) mouse 7). This mouse developed a large cervical tumor, which stained negative for CD11b, B220 and CD3 (not shown). Mice that were killed and examined at day 20 and 38 exhibited no signs of lymphocytosis (Table 2, p210 BCR/ABL1(Δ674–695) mice 1–6). Between days 75 and 89 post BMT, seven of the remaining mutant-transplanted mice were killed and necropsies were performed. Five of these mice showed no obvious signs of disease, having normal WBC counts and spleen weights, and no observable lymphadenopathy (Figure 6b, Table 2, p210 BCR/ABL1(Δ674–695) mice 8–10, 12 and 13). GFP⁺ cells comprised less than 2% of the bone marrow, spleen and peripheral blood, which is equivalent to what we observed for vector-transplanted mice (Table 2). Although two of the killed mice showed evidence of disease progression (mice 11 and 14), their phenotype was consistent with T-cell leukemia. These mice presented with significant ascites and one had a large abdominal tumor (mouse 14). Flow cytometry performed on the peripheral blood, spleen, ascitic fluid and tumor of this mouse demonstrated an expansion of GFP⁺ cells (Figure 6c). Although all of these cells were negative for CD11b and B220 expression, a proportion did express the T-cell marker CD3. Two additional

**Figure 4.** Comparison of immunophenotypes during early-stage disease progression in mice transplanted with p210 BCR/ABL1 or p210 BCR/ABL1(Δ674–695). WBCs were collected from p210 BCR/ABL1, p210 BCR/ABL1(Δ674–695) and vector-transplanted mice following elective killing on day 16 (a) and day 30 (b) post BMT, and were examined by flow cytometry for GFP expression. Cells were stained for CD11b, B220 and CD3 as indicated.
Table 1. Immunophenotyping of disease progression in a BMT assay for CML

|                      | Peripheral blood | Bone marrow | Spleen |
|----------------------|------------------|-------------|--------|
|                      | GFP⁺             | GFP⁺/CD11b⁺ | GFP⁺/Gr1⁺ | GFP⁺/CD3⁺ | GFP⁺/B220⁺ |
| Vector               | 8.69 ± 6.51      | 4.78 ± 1.44 | 1.81 ± 0.07 | 0.046 ± 0.04 | 0.002 ± 0.005 |
| BCR/ABL              | 82.12 ± 8.53     | 69.20 ± 16.69 | 16.78 ± 2.63 | 0.51 ± 0.22 | 6.72 ± 6.35 |
| Δ674–695             | 67.00 ± 4.69     | 57.50 ± 5.97 | 37.70 ± 2.08**| 0.75 ± 0.14 | 4.38 ± 1.01 |
|                      | 9.80 ± 9.09      | 7.41 ± 3.97 | 4.11 ± 1.96 | 0.64 ± 0.30 | 0.003 ± 0.002 |
|                      | 69.19 ± 17.23    | 55.38 ± 16.17 | 16.3 ± 3.58 | 0.82 ± 0.60 | 3.61 ± 3.58 |
|                      | 72.56 ± 21.26    | 62.86 ± 24.86 | 40.34 ± 15.68*| 0.70 ± 0.29 | 3.58 ± 0.85 |
|                      | 5.67 ± 3.74      | 4.74 ± 1.73 | 2.37 ± 0.88 | 1.85 ± 0.59 | 0.005 ± 0.003 |
|                      | 65.12 ± 9.07     | 39.97 ± 17.11 | 15.95 ± 2.36 | 2.94 ± 1.44 | 4.61 ± 2.29 |
|                      | 50.00 ± 15.36    | 47.05 ± 14.56 | 33.55 ± 6.89*| 1.08 ± 0.52 | 3.97 ± 0.82 |

Abbreviations: BMT, bone marrow transplantation; CML, chronic myelogenous leukemia. Immunophenotyping was performed at death as described in Materials and Methods. Vector mice were electively killed on day 30 post BMT. Data shown are an average of a minimum of five mice with s.d. (*P < 0.05, **P < 0.01, relative to BCR/ABL).

Figure 5. XPB binding limits the self-renewal of myeloid progenitors in p210 BCR/ABL1-transplanted mice. BMCs isolated from diseased mice at death were used for immunophenotypic and cell cycle analysis of progenitor populations. (a) Representative fluorescence-activated cell sorting staining profiles of progenitor populations. (b) Percentages of each progenitor populations (GMP, CMP and MEP) relative to total GFP⁺ cells. Values were derived from at least five mice per group and are represented as averages. Data show significant increase of GMP in BCR/ABL1(Δ674–695) mice relative to BCR/ABL1 mice (P < 0.05)). (c) Cell cycle analysis of CMPs, GMPs and MEPs in p210 BCR/ABL1 mouse and p210 BCR/ABL1(Δ674–695) mice. Values were derived from at least five mice per group and are represented as averages.

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In the current study, we have demonstrated that disruption of the XPB interaction results in disease attenuation in BMT models of CML and B-ALL. In the model for CML, mice transplanted with the XBP-binding mutant exhibit increased survival. Myeloid expansion is primarily restricted to Gr1⁺ cells, which, in turn, is driving the disease phenotype. Although the number of GMPs at death in the mutant-transplanted mice is elevated, they show no increase in their proliferative potential. This is consistent with the ex-vivo clonogenic assays wherein the mutant is actually more limited in its ability to support the growth of GMP. As the disease phenotype appears to be restricted to a sub-lineage of GMPs, a larger number of GMPs may need to accumulate in order to achieve a tumor burden, resulting in morbidity. This would account for the substantially increased lifespan.

At day 16 post BMT, we observed an expansion of the B220⁺ cells in the p210 BCR/ABL1-transplanted mice but not in the mutant. This expansion suggests that transformed lymphoid progenitors are engrafting earlier than myeloid progenitors and may be enjoying an early proliferative advantage. However, as the myeloid lineages expand, they may limit further expansion of the lymphoid progenitors. The failure of the mutant to drive lymphoproliferation was confirmed in a BMT model for B-ALL. Whereas the p210 BCR/ABL1-transplanted mice develop and succumb to a disease resembling human B-ALL, the mutant-transplanted mice either remained disease free or developed T-cell leukemias with long latencies. The ability of p210 BCR/ABL1 to drive T-cell leukemias has not been previously observed in the BMT model and suggests that it may be transforming a common lymphoid progenitor. Whereas the interaction with XBP in these cells supports lymphoid expansion, the loss of the interaction may favor T-cell expansion.

As cells that express the XPB-binding mutant show lower levels of XBP phosphorylation on tyrosine, it is possible that p210 BCR/ABL1 transformation is influenced directly by XBP-associated activities, which may be altered by phosphorylation. It has been shown that tyrosine phosphorylation of XBP by p210 BCR/ABL1 reduces its ATPase and helicase activities in vitro,¹⁹ which is likely to result in both transcriptional and repair defects. Several studies indicate that the rate of NER is influenced by p210 BCR/ABL1, although opposing effects have been observed in lymphoid and myeloid cells.¹⁵,¹⁶ Although we observe similar effects of p210 BCR/ABL1 on NER in Ba/F3 cells and primary murine myeloid cells, these effects do not appear to be dependent upon the interaction with XBP.

p210 BCR/ABL1 expression may also interfere with the transcriptional functions of TFIIH. A number of studies have documented altered transcription of specific target genes in response to p210 BCR/ABL1 expression, including c-MYC,³⁷ Bcl-Xl,³⁸ PKC,³⁹ and TRAIL.⁴⁰ In addition, global changes in gene expression have been observed in 32DC13 myeloid cells that stably express p210 BCR/ABL1.⁴¹ Although some of these transcriptional changes can be attributed to alterations in STAT-regulated pathways,³⁸ the interaction with XBP may represent a separate mechanism through which p210 BCR/ABL1 can regulate transcriptional events. For example, recent studies suggest that expression of the c-myc gene, which is frequently upregulated in CML, is controlled by a transcriptional complex that contains components of TFIIH, including XBP.³²,³³,³⁴ We have previously shown that BCR is a nuclear protein that binds directly to c-MYC and inhibits its expression, thus supporting leukemic expansion. This in turn may provide unique opportunities for therapeutic intervention.

### DISCUSSION

In the current study, we have demonstrated that disruption of the XBP interaction results in disease attenuation in BMT models of CML and B-ALL. In the model for CML, mice transplanted with the XBP-binding mutant exhibit increased survival. Myeloid expansion is primarily restricted to Gr1⁺ cells, which, in turn, is driving the disease phenotype. Although the number of GMPs at death in the mutant-transplanted mice is elevated, they show no increase in their proliferative potential.

*Table 2. Immunophenotyping of disease progression in a BMT model for ALL*  

| Mouse (day at death or killing) | Peripheral blood (% of total cells) |
|-------------------------------|------------------------------------|
|                               | GFP⁺ / CD11b⁺ | GFP⁺ / B220⁺ | GFP⁺ / CD3⁺ |
| **Vector**                    |                      |                |
| No. 1 (day 20)               | 13                  | 12             | 1           |
| No. 2 (day 38)               | 8                   | 6              | 1           |
| No. 3 (day 93)               | 8                   | 1              | 6           |
| p210 BCR/ABL1                |                      |                |
| No. 1 (day 20)               | 34                  | 5              | 31          |
| No. 2 (day 20)               | 21                  | 15             | 15          |
| No. 3 (day 20)               | 76                  | 5              | 72          |
| No. 4 (day 38)               | 51                  | 15             | 35          |
| No. 5 (day 38)               | 6                   | 3              | 4           |
| No. 6 (day 38)               | 69                  | 2              | 69          |
| No. 7 (day 45)               | 15                  | 4              | 13          |
| No. 8 (day 47)               | 42                  | 5              | 40          |
| No. 9 (day 54)               | 71                  | 2              | 65          |
| No. 10 (day 64)              | 96                  | 5              | 89          |
| No. 11 (day 76)              | 85                  | <1             | 85          |

**Table 2.** Immunophenotyping of disease progression in a BMT model for ALL

Abbreviations: ALL, acute lymphoblastic leukemia; BMT, bone marrow transplantation. Immunophenotyping was performed as described in Materials and Methods. *Mice that succumbed due to disease. All other mice were electively killed.

mutant-transplanted mice were killed at day 106, and neither exhibited any signs of disease progression (Table 2, p210 BCR/ABL1(Δ674–695) mice 15 and 16). To summarize, in all of the 16 mutant-transplanted mice that were killed and immunophenotyped, there was no evidence of B-cells proliferation in any organ that was examined.

Recipients receiving marrow infected with vector (n = 20 total) exhibited no sign of disease throughout the survival study (Table 2).
Figure 6. XPB binding supports lymphoproliferation in a murine model for B-ALL. (a) Survival of mice transplanted with p210 BCR/ABL1 or p210 BCR/ABL1(Δ674–695). A Kaplan–Meier curve was generated for the first 75 days following BMT. A Mantel–Cox test yielded values of P = 0.0002 and χ² test = 13.91. (b and c) Hematopoietic tissues were collected at the time of death and were examined by flow cytometry. Cells were stained for CD11b, B220 and CD3 as indicated. (b) Comparison of disease progression. On day 76, a p210 BCR/ABL1 (Table 2, p210 BCR/ABL1(Δ674–695)-transplanted mouse was killed for analysis. Results shown are from the bone marrow and are representative of both the blood and spleen. (c) A subset of the mutant-transplanted mice develop T-cell leukemias with long latency. On day 89, a p210 BCR/ABL1(Δ674–695)-transplanted mouse was killed for analysis (Table 2, p210 BCR/ABL1(Δ674–695) mouse 14). Tumor and ascitic fluid samples were also analyzed.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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