

LETTERS

Essential roles of PI(3)K-p110 β in cell growth, metabolism and tumorigenesis

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On activation by receptors, the ubiquitously expressed class IA isoforms (p110 α and p110 β) of phosphatidylinositol-3-OH kinase (PI(3)K) generate lipid second messengers, which initiate multiple signal transduction cascades^{1–5}. Recent studies have demonstrated specific functions for p110 α in growth factor and insulin signalling^{6–8}. To probe for distinct functions of p110 β , we constructed conditional knockout mice. Here we show that ablation of p110 β in the livers of the resulting mice leads to impaired insulin sensitivity and glucose homeostasis, while having little effect on phosphorylation of Akt, suggesting the involvement of a kinase-independent role of p110 β in insulin metabolic action. Using established mouse embryonic fibroblasts, we found that removal of p110 β also had little effect on Akt phosphorylation in response to stimulation by insulin and epidermal growth factor, but resulted in retarded cell proliferation. Reconstitution of p110 β -null cells with a wild-type or kinase-dead allele of p110 β demonstrated that p110 β possesses kinase-independent functions in regulating cell proliferation and trafficking. However, the kinase activity of p110 β was required for G-protein-coupled receptor signalling triggered by lysophosphatidic acid and had a function in oncogenic transformation. Most strikingly, in an animal model of prostate tumour formation induced by *Pten* loss, ablation of p110 β (also known as *Pik3cb*), but not that of p110 α (also known as *Pik3ca*), impeded tumorigenesis with a concomitant diminution of Akt phosphorylation. Taken together, our findings demonstrate both kinase-dependent and kinase-independent functions for p110 β , and strongly indicate the kinase-dependent functions of p110 β as a promising target in cancer therapy.

Class IA PI(3)Ks are heterodimeric lipid kinases consisting of a p110 catalytic subunit complexed to one of several regulatory subunits, collectively called p85 (refs 4, 5). In response to stimulation by growth factor, p110 subunits catalyse the production of the lipid second messenger phosphatidylinositol-3,4,5-trisphosphate (PtdIns(3,4,5)P₃) at the membrane^{1–4}. This second messenger in turn activates the serine/threonine kinase Akt and other downstream effectors^{9,10}. Knockout mice for either p110 α or p110 β die early in embryonic development^{11,12}. However, recent studies using conditional knockout strategies⁸ and isoform-specific small molecule inhibitors⁷ showed that p110 α is important in growth-factor signalling, whereas a kinase-inactive knock-in mouse model showed that insulin responses depended on the catalytic activity of p110 α (ref. 6).

To investigate the role(s) of p110 β in cell, tissue and organismal physiology and to examine it as a potential therapeutic target in cancer, we generated mice carrying a conditional *Pik3cb* allele (Supplementary Fig. 1). We first investigated the role of p110 β in insulin action. Because liver is the major insulin-responsive organ,

we examined the effects of p110 β loss on hepatic insulin function. To achieve liver-specific deletion of p110 β , we injected the tail veins of p110 β ^{flox/flox} mice with adenoviruses expressing β -galactosidase (Ade-LacZ) or Cre recombinase (Ade-Cre) to generate matched cohorts of control mice and mice with hepatocyte-specific deletion of p110 β . Additional cohorts of wild-type animals were subjected to Ade-Cre or Ade-LacZ, allowing us to rule out potential non-specific Cre effects (data not shown). A more than 90% decrease in p110 β protein was seen in the livers of Ade-Cre-injected mice, whereas p110 β expression remained unchanged in the livers of the control mice and muscle tissues from both groups as measured by western blotting (Supplementary Fig. 2a, b). Consistent with previous findings^{6,7} that the kinase activity of p110 β has only a minor function in insulin signalling, we saw no significant change in Akt phosphorylation in response to insulin challenge in livers lacking p110 β (Supplementary Fig. 2a). However, mice deficient in hepatic p110 β had higher levels of insulin in the blood than control animals when fasted (Fig. 1a). These animals also showed a lower tolerance of glucose and sensitivity to insulin on challenge by intraperitoneal injection of glucose or insulin (Fig. 1b, c). Mice deficient in hepatic p110 β produced more glucose than control animals did in a pyruvate challenge test (Fig. 1d). An analysis of lipogenesis showed no significant changes in serum triglycerides, fatty acids and cholesterol levels when p110 β was deleted from liver (Supplementary Fig. 3), but leptin levels were elevated compared with those in control animals, as was seen in p110 α kinase-dead knock-in animals⁶ (Supplementary Fig. 3). Of a panel of gluconeogenic genes, only that encoding phosphoenolpyruvate carboxykinase (PEPCK) was increased in p110 β -deficient livers (Supplementary Fig. 4). PEPCK promotes the production and synthesis of glucose in liver, resulting in a greater release of glucose into blood. This result therefore provides at least a partial explanation for the metabolic phenotypes observed. Although these findings indicate that p110 β might contribute to metabolic regulation through a kinase-independent mechanism, we cannot rule out the involvement of the catalytic role of p110 β in insulin responses. Our observations are in line with earlier work⁷ in which a p110 β -specific small-molecule inhibitor was used to demonstrate that acute blockage of the kinase activity of p110 β had little effect on insulin action. In addition, another study¹³ found that mice doubly heterozygous for knockout of p110 α and p110 β showed decreased sensitivity to insulin with no apparent changes in Akt phosphorylation.

To obtain cells for detailed signalling studies, mouse embryonic fibroblasts (MEFs) were isolated from floxed embryos and their wild-type littermates, as described in Supplementary Information (Supplementary Fig. 5a–c). MEFs lacking p110 β proliferated significantly more slowly than parental (p110 β ^{flox/flox}) or wild-type

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($p110\beta^{+/+}$ after Cre) MEFs (Fig. 2a). To obtain a second, more easily renewed supply of knockout cells, we established immortalized $p110\beta^{\text{flox/flox}}$ and $p110\beta^{+/+}$ MEFs by means of infection with a retrovirus encoding a dominant-negative form of p53 (DNp53)¹⁴. We also generated an add-back line by introducing haemagglutinin (HA)-tagged human $p110\beta$ to DNp53-immortalized $p110\beta^{\text{flox/flox}}$ MEFs. These immortalized MEFs were then treated with Ade-Cre to yield the following MEF lines: β KO (from $p110\beta^{\text{flox/flox}}$) and β KO+ β (from the add-back). For wild-type control MEFs, designated WT, we used DNp53-immortalized $p110\beta^{\text{flox/flox}}$ cells without Cre treatment interchangeably with Cre-treated DNp53-immortalized $p110\beta^{+/+}$ MEFs, because no significant differences were ever seen between these two possible controls. Deletion of $p110\beta$ had no obvious negative effect on the phosphorylation of Akt in either primary MEFs or DNp53-immortalized MEFs in response to stimulation by insulin, epidermal growth factor (EGF) and platelet-derived growth factor (Fig. 2c and Supplementary Fig. 6a–c). However, a moderate diminution in the phosphorylation of the S6 ribosomal protein (S6RP) at Ser 235/Ser 236 was detected in these β KO cells in response to insulin or serum (Supplementary Fig. 7). Previous studies have implicated $p110\beta$ in signalling elicited by G-protein-coupled receptors (GPCRs)^{15,16}. We found consistently that both phospho-Akt and phospho-S6RP levels were decreased in response to lysophosphatidic acid (LPA) in cells lacking $p110\beta$ (Fig. 2d and Supplementary Fig. 8a).

To dissect the potential kinase-dependent and kinase-independent roles of $p110\beta$, we reconstituted β KO MEFs with a kinase-inactive allele of HA-tagged human $p110\beta$, using the previously reported K805R mutation (KR)¹⁷ to generate the β KO+KR MEF line. Though the expression of KR was lower than that of the WT add-back construct, it was expressed at a level slightly higher than endogenous $p110\beta$, and expression levels of $p110\alpha$ were unchanged (Supplementary Fig. 9a, and data not shown). Loss of lipid kinase activity in the KR cells was confirmed by lipid kinase assay^{8,18} after immunoprecipitation with anti- $p110\beta$ (Supplementary Fig. 9b). We

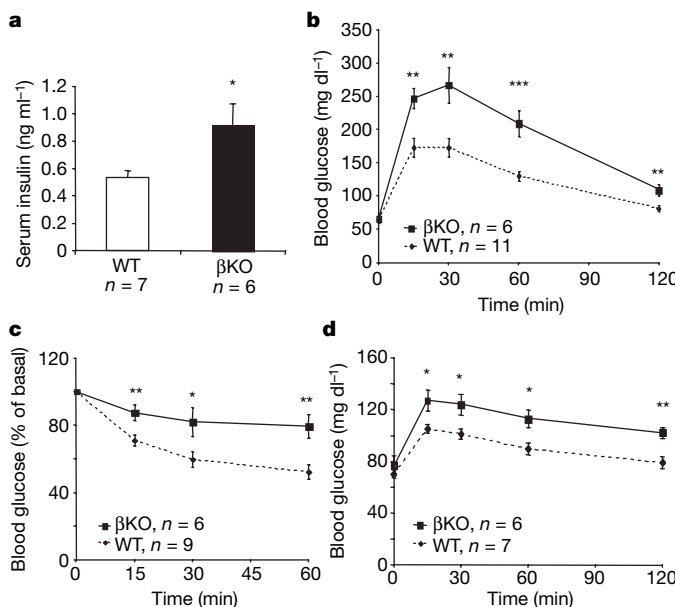


Figure 1 | Mice with liver-specific deletion of $p110\beta$ show resistance to insulin and intolerance of glucose. Mice 8–10 weeks old were injected with adenoviruses expressing LacZ or Cre recombinase. Two weeks after injection, metabolism was analysed as follows: fasted serum insulin levels (a); glucose tolerance test (b); insulin tolerance test (c) (results represent blood glucose concentrations as a percentage of starting value at time zero); pyruvate challenge (d). Data are shown as means and s.e.m. Asterisk, $P < 0.05$; two asterisks, $P < 0.01$; three asterisks, $P < 0.001$ (*t*-test).

then examined the effect of WT or KR add-back on the altered signalling seen after the loss of $p110\beta$. The decrease in both phospho-Akt and phospho-S6RP in response to stimulation with LPA observed in β KO cells was restored by adding back WT but not the KR allele of $p110\beta$ (Fig. 2d and Supplementary Fig. 8b), suggesting a catalytic function for $p110\beta$ in LPA signalling. This seems to be unique to $p110\beta$, because loss of $p110\alpha$ has no obvious effect on LPA signalling (Fig. 2e). The lower phospho-S6RP levels in β KO cells were restored by both WT and KR add-backs in response to insulin or fetal bovine serum (FBS) (Supplementary Fig. 7, and data not shown), suggesting a scaffolding role of $p110\beta$ in signalling by insulin and growth factors. However, our MEF data do not rule out a role for $p110\beta$ in classical signalling by PI(3)K in other circumstances. For instance, when we ablated $p110\alpha$ in our earlier work, residual phosphorylation by Akt was observed in response to growth factors⁸. Because MEFs express $p110\alpha$ and $p110\beta$ and not other class I PI(3)Ks, this residual signal was presumably transduced by $p110\beta$. We also note that $p110\beta$ ablation removes the protein as a competitor for $p110\alpha$ on receptors, which may allow any decrease in signalling caused by $p110\beta$ loss to be masked or compensated for by an increased signal flux through $p110\alpha$.

To test the kinase-dependent and/or kinase-independent effects of $p110\beta$ on cell proliferation, we studied cell cycle kinetics by first synchronizing cells by serum starvation and then measuring the proportion of cells in S phase with the incorporation of bromodeoxyuridine (BrdU) after re-feeding. Whereas β KO cells had a delayed peak of BrdU incorporation, KR reconstituted cells showed a similar incorporation of BrdU to that of WT and β KO+ β cells (Fig. 2f).

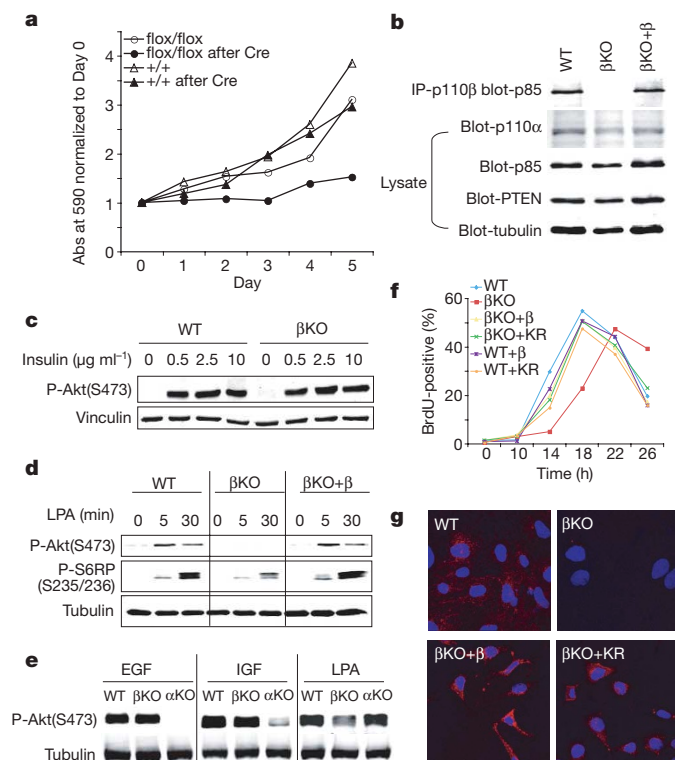


Figure 2 | Analyses of the effects of $p110\beta$ deletion on cell growth and signalling. a, Loss of $p110\beta$ retards cell growth of primary MEFs. Representative data are shown from triplicate experiments. b, Loss of endogenous $p110\beta$ protein in immortalized MEFs. IP, immunoprecipitation. c, Loss of $p110\beta$ has no negative effect on insulin signalling. d, Loss of $p110\beta$ impairs LPA-induced signalling. e, Comparison of the responses of β KO, β KO and WT MEFs to stimulation with EGF, insulin-like growth factor (IGF) or LPA. f, BrdU incorporation assay. g, Transferrin uptake in various MEF lines is shown as indicated. Transferrin is labelled red and 4,6-diamidino-2-phenylindole blue.

Consistently, β KO+KR MEFs showed proliferation rates similar to those of WT and β KO+ β cells (Supplementary Fig. 9c), suggesting a kinase-independent role of p110 β in cell proliferation.

Because previous studies have found p110 β associated with members of the Rab family of small G proteins and clathrin-coated vesicles¹⁹, we measured transferrin uptake in β KO MEFs and found it to be defective compared with that in WT and β KO+ β MEFs (Fig. 2g). Interestingly, normal uptake of transferrin was restored by the KR construct (Fig. 2g). Although there is ample published evidence indicating the importance of transferrin uptake for the growth of a variety of cell types²⁰, it is not clear whether the defect in transferrin uptake is a primary cause of the growth defect observed here.

Class IA PI(3)Ks have been clearly implicated in cancer^{21–24}, with much recent work delineating the role of p110 α in cancer^{25–27}. To study a potential role of p110 β in oncogenic transformation, we performed focus formation assays by infecting monolayers of DNp53-immortalized MEFs with retroviruses expressing various oncogenes. Oncogenic HRas-G12V and EGFR-Del (ΔL747–E749, A750P) retroviruses efficiently raised foci in WT cells but failed to transform β KO MEFs (Fig. 3a). The decreases in foci seen in β KO MEFs were actually more pronounced than those seen in p110 α KO MEFs (Supplementary Fig. 10). Transformation was fully restored in

β KO+ β cells but partly restored in β KO+KR cells (Fig. 3a), suggesting that both the kinase activity and kinase-independent functions of p110 β may have a function in oncogene-induced transformation.

PTEN, a lipid phosphatase, functions to oppose class IA PI(3)K kinase activity. Loss of *Pten* expression is a common event in many solid tumours²⁸. The key challenge is to identify which p110 isoform's catalytic activity is unshackled by *Pten* loss in any given tumour. To test for a role of p110 β in tumorigenesis driven by *Pten* loss, we generated mice that carried the *Pten*^{fl/fl} (ref. 29) and p110 β ^{fl/fl} alleles, as well as a probasin-driven Cre transgene³⁰, to specifically delete *Pten* and p110 β in prostatic epithelium. Prostates had a normal appearance in the absence of p110 β (Fig. 3b, c). Prostate tissue lacking *Pten* expression showed universal high-grade prostatic intraepithelial neoplasia in the anterior lobe by 12 weeks. Ablation of p110 β blocked the tumorigenesis caused by *Pten* loss in the anterior prostate (Table 1 and Fig. 3b, c). The loss of *Pten* was confirmed by genomic DNA analysis after laser-capture-assisted microdissection of single epithelial layers and by western blotting (Supplementary Fig. 11, and data not shown). Whereas Cre-mediated loss of *Pten* efficiently activated Akt in the prostate as judged by its phosphorylation on Ser 473, additional ablation of p110 β diminished the phospho-Akt levels (Fig. 3b), suggesting that p110 β catalytic activity contributes to tumorigenesis. More surprisingly, when we performed the same set of experiments using p110 α ablation, we saw no changes either in tumour formation or in Akt phosphorylation (Table 1 and Fig. 3b, c). Again, the complete excision of p110 α in tumour tissues was confirmed by multiple measures (Supplementary Fig. 12, and data not shown). It has been suggested that p110 α and p110 β generate distinct pools of PtdIns(3,4,5)P₃ (ref. 7). In response to insulin or other stimuli, an acute flux of PtdIns(3,4,5)P₃ is produced largely by p110 α and is efficiently coupled to Akt phosphorylation. In contrast, p110 β has been proposed to generate a basal level of PtdIns(3,4,5)P₃ with little effect on Akt phosphorylation⁷. It was shown that Akt phosphorylation induced by *Pten* loss *in vitro* was sensitive to p110 β -specific inhibitors⁷. We propose that it is this basal PtdIns(3,4,5)P₃ signal that has been enhanced to drive transformation and Akt activation by *Pten* loss in the murine prostate (Fig. 3d). Alternatively, the differential effects of p110 α and p110 β ablation may arise because the signal activating PI(3)K is generated by an as yet unidentified GPCR or a receptor tyrosine kinase that functions through p110 β .

Thus, our data suggest distinct functions for p110 β and p110 α in cell signalling and transformation. We have shown that p110 β has an important physiological function in metabolic regulation and glucose homeostasis, perhaps involving a kinase-independent mechanism. A kinase-independent function of p110 β was further suggested in controlling cell proliferation and trafficking in p110 β KO MEFs and MEFs reconstituted with a WT or kinase-dead allele of p110 β . It would clearly be a mistake to overlook the contributions of p110 β as a kinase. The basal PtdIns(3,4,5)P₃ pool catalysed by p110 β seems to be 'silent' in response to stimulation by insulin and other growth factors, but becomes a 'powerhouse' to drive oncogenic

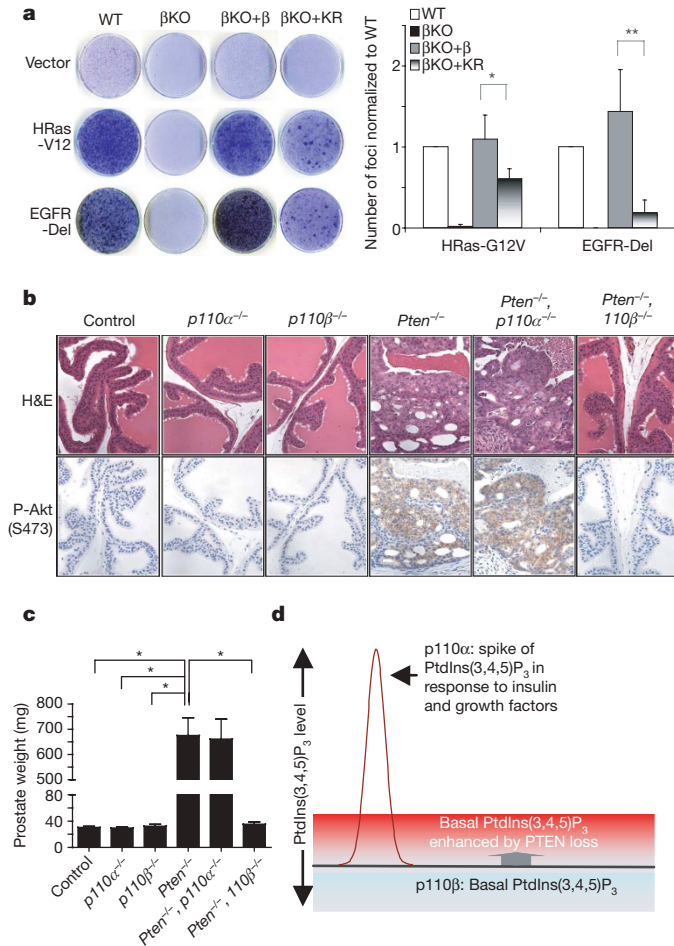


Figure 3 | Kinase activity of p110 β contributes to transformation both *in vitro* and *in vivo*. **a**, Focus formation assay in KO and reconstituted MEFs. Results are shown as means and s.e.m. for four independent experiments (asterisk, $P < 0.05$; two asterisks, $P < 0.01$, *t*-test). **b**, Effects of p110 β or p110 α ablation on tumorigenesis caused by PTEN loss in the anterior prostate. **c**, Quantification of the weight (means and s.e.m.) of anterior prostate tissues of the indicated strain ($n = 10$ per group; asterisk, $P < 0.001$, *t*-test). **d**, A model for the elevation of basal PtdIns(3,4,5)P₃ signals derived from p110 β catalytic activity induced by PTEN loss.

Table 1 | Effects of ablation of p110 α or p110 β on prostate tumorigenesis induced by PTEN loss

Abbreviation	Full name	Animals positive for PIN/number of animals used
Control	Floxed littermates	0/20
p110 α ^{-/-}	p110 α ^{fl/fl} , PbCre4	0/20
p110 β ^{-/-}	p110 β ^{fl/fl} , PbCre4	0/14
Pten ^{-/-}	Pten ^{fl/fl} , PbCre4	20/20
Pten ^{-/-} ; p110 α ^{-/-}	Pten ^{fl/fl} , p110 α ^{fl/fl} , PbCre4	15/15
Pten ^{-/-} ; p110 β ^{-/-}	Pten ^{fl/fl} , p110 β ^{fl/fl} , PbCre4	0/16

Paraffin sections of anterior prostates from the indicated strains at 12 weeks were stained with haematoxylin/eosin. The pathological phenotype was uniformly observed within each genotype and is summarized in the table. PIN, prostatic intraepithelial neoplasia.

transformation in the absence of PTEN, as evident in our mouse prostate tumour model. Taken together, our findings indicate that p110 β may be an attractive target for kinase inhibitors in cancer treatment with minor metabolic disturbances.

METHODS SUMMARY

Mice carrying floxed *p110 β* (generated in this work), floxed *p110 α* (ref. 8), floxed *Pten*²⁹ and probasin-driven Cre transgene³⁰ (Mouse Models of Human Cancers Consortium (MMHCC), National Cancer Institute) were used in this study. All animals were housed and treated in accordance with protocols approved by the Institutional Animal Care and Use Committees of Dana-Farber Cancer Institute and Harvard Medical School. The generation, culture and immortalization of MEFs, growth factor signalling study, retroviral infection, cell growth, cell cycle, lipid kinase assay, transferrin internalization, focus formation, glucose tolerance testing, insulin tolerance tests, pyruvate challenge, immunoprecipitation, immunoblotting, immunohistochemical and histological analyses were performed in accordance with standard or published protocols. Statistical analyses were performed with Student's *t*-test unless otherwise indicated.

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

Received 27 February; accepted 15 May 2008.

Published online 25 June 2008.

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Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

Acknowledgements We thank C. D. Stiles and J. D. Igglehart for advice, and H. Wu for providing floxed PTEN mice. This work was supported by grants from the National Institutes of Health (M.L., T.M.R. and J.J.Z.), the Department of Defense for Cancer Research (J.J.Z.), the V Foundation (J.J.Z.) and the Claudia Barr Program (J.J.Z.). In compliance with Harvard Medical School guidelines, we disclose the consulting relationships: Novartis Pharmaceuticals, Inc. (M.L., T.M.R. and J.J.Z.).

Author Contributions Z.L., S.Z. and S.L. generated the floxed *p110 β* mouse. S.J. carried out mouse tumorigenesis studies. Z.L. and S.Z. performed MEF studies. P.L. performed *in vivo* metabolic studies. L.Z. performed transferrin uptake assays. S.S. and M.L. assisted in focus formation and BrdU incorporation experiments. S.S. and M.L. performed and interpreted pathological analyses of mouse prostate tumors. T.M.R. and J.J.Z. supervised the research, interpreted the data and wrote the paper. S.J., Z.L., S.Z., P.L., L.Z., S.L. and M.L. participated in the writing of the paper.

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METHODS

Mice for metabolic and tumour studies. Conditional knockout mice of *p110 β* were generated using the Cre-LoxP system. In brief, the targeting construct was assembled by isolating a 7.5-kilobase genomic fragment of *Pik3cb* from 129SvEv mouse strain and inserting two LoxP sites to flank exon 2 of *Pik3cb*. The targeting construct was electroporated into embryonic stem (ES) cells of 129SvEv mouse. ES clones carrying floxed *Pik3cb* were injected into the blastocysts of C57BL/6 mice. Male chimaeras were bred to C57BL/6 females to establish germline transmission of the conditional allele. The resulting heterozygous line (*p110 β* ^{flx/+}) was intercrossed to yield a homozygous line (*p110 β* ^{flx/flx}).

For metabolic studies, 8–10-week-old male *p110 β* ^{flx/flx} littermates were tail-vein injected with 75 μ l of adenovirus CMV-lacZ and CMV-cre (titre between 10^{10} and 4×10^{10} plaque-forming units ml $^{-1}$; University of Iowa Gene Transfer Vector Core). Two weeks after injection of adenoviruses, glucose tolerance test, insulin tolerance test, pyruvate challenge and *in vivo* insulin signalling were performed as described previously³¹. Blood glucose values were determined with an Accu-Chek AVIVA glucose monitor (Roche). Serum insulin and leptin (Crystal Chem Inc.), serum-free fatty acids and triacylglycerols (Wako) and serum cholesterol (Thermo) were measured by ELISA in accordance with the manufacturer's instructions.

It took two steps to generate compound mice for tumour studies. All floxed mice used were originally in the 129-C57BL/6 background and had been back-crossed once to C57BL/6. The probasin Cre mice were in the C57BL/6 background. In step 1, male PbCre4 mice were first crossed to the female *Pten*^{flx/flx}, *p110 α* ^{flx/flx} or *p110 β* ^{flx/flx} mice, and the male offspring carrying PbCre4 were then crossed to female *Pten*^{flx/flx}, *p110 α* ^{flx/flx} or *p110 β* ^{flx/flx} mice to obtain *Pten*^{flx/flx}; PbCre4 (hereafter termed '*Pten*^{-/-}'), *p110 α* ^{flx/flx}; PbCre4 (hereafter termed '*p110 α* ^{-/-}') or *p110 β* ^{flx/flx}; PbCre4 (hereafter termed '*p110 β* ^{-/-}') mice, respectively. We used cohorts of littermates lacking the probasin Cre transgene as wild-type controls (*Pten*^{flx/flx}, *p110 α* ^{flx/flx} or *p110 β* ^{flx/flx}), and no significant differences were observed between these groups.

In step 2, *Pten*^{-/-} male mice were crossed with female *p110 α* ^{flx/flx} or *p110 β* ^{flx/flx} mice, and their male offspring carrying PbCre were crossed further with *Pten*^{flx/wt}; *p110 α* ^{flx/wt} or *Pten*^{flx/wt}; *p110 β* ^{flx/wt} littermates to obtain the desired *Pten*^{flx/flx}; *p110 α* ^{flx/flx}; PbCre4 (hereafter termed '*Pten*^{-/-}; *p110 α* ^{-/-}') or *Pten*^{flx/flx}; *p110 β* ^{flx/flx}; PbCre4 (hereafter termed '*Pten*^{-/-}; *p110 β* ^{-/-}') compound strains. The resultant *Pten*^{flx/flx}; *p110 α* ^{flx/flx}; PbCre4 or *Pten*^{flx/flx}; *p110 β* ^{flx/wt}; PbCre4 mice from the same litters displayed high-grade prostatic intraepithelial neoplasia comparable to that of *Pten*^{flx/flx}; PbCre4 mice described in step 1, and were used interchangeably as tumour controls. Cohorts of littermates lacking the probasin Cre transgene were used as wild-type controls showing no significant differences from wild-type controls in step 1.

Alternatively, in step 2 *Pten*^{-/-}; *p110 α* ^{-/-} or *Pten*^{-/-}; *p110 β* ^{-/-} males were crossed with female littermates of *Pten*^{flx/flx}; *p110 α* ^{flx/flx} or *Pten*^{flx/flx}; *p110 β* ^{flx/flx} to generate larger numbers of the desired compound strains. No differences were seen in phenotypes in compound animals generated by either version of step 2.

Primary and immortalized MEFs. MEFs were prepared from embryos derived from intercrossing *p110 β* ^{flx/+} heterozygotes at embryonic day 13.5 after fertilization. Primary WT or floxed MEFs, and DNp53 immortalized WT or floxed MEFs were treated with Ade-Cre to generate WT control and *p110 β* -null (β KO) cells. Additional control cells used in our study were floxed MEFs without Ade-Cre treatment. β KO+ β and β KO+KR lines were generated by introducing either HA-tagged WT human *p110 β* or a kinase-inactive mutant K805R into β KO MEFs and then treated with Ade-Cre. Genotyping of MEFs was done by PCR using primer sets: LLF with LLR, and SLF with LLR (Supplementary Fig. 1).

Growth factors and western blotting. Cells were starved either for 2 h or overnight followed by stimulation with insulin (2.5 μ g ml $^{-1}$; Sigma I2767), EGF (10 ng ml $^{-1}$; Sigma E9644), IGF1 (5 ng ml $^{-1}$, Upstate 01-208), LPA (10 μ M; Sigma L7260) or 10% FBS for various periods as indicated in the corresponding figures. Western blot assays were performed as described previously⁸ with antibodies against PTEN (no. 9552), *p110 α* (no. 4255), phospho-Akt (Ser 473 (no. 9271) or Thr 308, no. 9275), Akt (no. 9272), phospho-p70 S6 kinase (Thr 389,

no. 9205), phospho-p44/42 MAP kinase (no. 9101), phospho-S6 ribosomal protein (Ser 235/Ser 236, no. 2211) and S6 ribosomal protein (no. 2217) (Cell Signalling), p85 (Upstate 06-195), *p110 β* (Santa Cruz, sc-602), Tubulin and Vinculin (Sigma T6199 and V9131). Immunofluorescently labelled anti-mouse IgG (Rockland Immunochemicals 610-132-003) and anti-rabbit IgG (Molecular Probes) were used to visualize western blots on an Odyssey scanner. The quantification of western blots was performed with Odyssey software version 2.0.

Growth curves. MEFs were seeded in 12-well tissue culture plates and stained with crystal violet at each indicated time point. The dye was extracted with 10% acetic acid followed by plate reading at 590 nm. The values were normalized to the absorbance at day 0. Data shown are the average of at least two independent experiments.

Cell cycle analysis. Cells were synchronized by starvation in DMEM supplemented with 0.1% FBS for 48 h before being released into the cell cycle by stimulation with 10% FBS. Cells were pulse-labelled with BrdU at each indicated time point and analysed in accordance with the manufacturer's protocol (BD Biosciences).

Focus formation assays. MEFs at 40–50% confluence were infected with various retroviruses: pBabe–Vector, pBabe–HRas-G12V or pBabe–EGFR-Del (Δ L747–E749, A750P) and then cultured for 14–21 days for WT, β KO+ β and β KO+KR cells, but for 30–40 days for β KO cells. Confluent monolayers with foci were fixed in ethanol and stained with crystal violet.

Histology and immunohistochemistry. Prostate tissues were processed and stained as described previously³². Primary antibody used in immunohistochemistry was directed against phospho-Akt (Ser 473) (no. 3787, Cell Signalling).

Lipid kinase assays. *In vitro* lipid kinase assays were performed as described previously^{8,18}. In brief, anti-*p110 β* (Santa Cruz) immunoprecipitates from freshly prepared cell lysates were subjected to an *in vitro* lipid kinase assay with phosphatidylinositol (Avanti Polar Lipids) as the substrate. The phosphorylated lipids were resolved by thin-layer chromatography, detected by autoradiography and quantified with Adobe Photoshop.

Transferrin internalization assays. Cells were seeded on 10% poly-(L-lysine)-coated coverslips and grown overnight in DMEM medium supplemented with 10% FBS. The assay was performed as described previously³³ with Alexa Fluor555-conjugated human transferrin (Invitrogen), counterstained with 4,6-diamidino-2-phenylindole (1 μ g ml $^{-1}$; Sigma) and mounted with mounting medium (Fisher Scientific). Cells were observed with a Zeiss confocal microscope LSM510META/NLO at $\times 63$ magnification, and images were captured with Zeiss confocal microscope software 3.2.

Laser capture microdissection and DNA extraction. Laser capture microdissection was performed as described previously³⁴. Genomic DNA of microdissected prostate epithelium was extracted with phenol/chloroform before PCR analysis.

Quantitative reverse transcription PCR analysis. Liver total RNAs were extracted with the use of the RNeasy kit (Qiagen). The following gene expression assay probes (Applied Biosystems) were used for real-time RT-PCR quantification: *phosphoenolpyruvate carboxykinase 1* (*PEPCK/Pck1*, ID no. Mm0440636_m1), *glucose-6-phosphatase* (*G6Pase*, ID no. Mm00839363_m1), *fructose bisphosphatase 1* (*Fbp1*, ID no. Mm00490181_m1), *hepatic nuclear factor 4* (*Hnf4a*, ID no. Mm0433964_m1) and *glyceraldehyde-3-phosphate dehydrogenase* (*GAPDH*, catalogue no. 4352339E). Expression was normalized to mRNA for *GAPDH* and results were expressed as fold change in mRNA compared with the indicated control mice.

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CORRECTIONS & AMENDMENTS

CORRIGENDUM

doi:10.1038/nature16543

Corrigendum: Essential roles of PI(3)K-p110 β in cell growth, metabolism and tumorigenesis

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Nature **454**, 776–779 (2008); doi:10.1038/nature07091

In Fig. 3b of this Letter we inadvertently used the wrong images (partial duplicates of images representing *p110 $\alpha^{-/-}$* mice) in the panels representing *p110 $\beta^{-/-}$* mice. The corrected Fig. 3b is shown in Fig. 1 of this Corrigendum. The conclusions of the paper are not affected; knocking out either *p110 α* or *p110 β* does not affect the growth of the normal prostate.

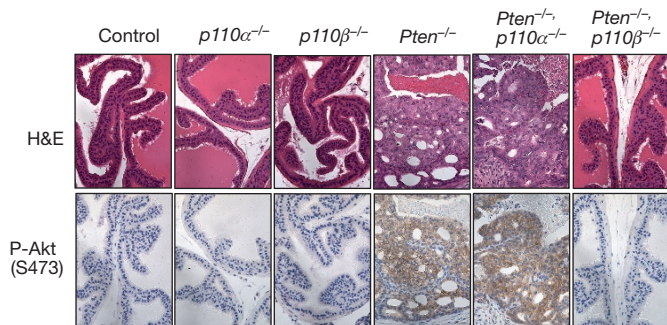


Figure 1 | This figure shows the corrected Fig. 3b of the original Letter.