Heterozygous splice mutation in PIK3R1 causes human immunodeficiency with lymphoproliferation due to dominant activation of PI3K

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Class IA phosphatidylinositol 3-kinases (PI3K), which generate PIP₃ as a signal for cell growth and proliferation, exist as an intracellular complex of a catalytic subunit bound to a regulatory subunit. We and others have previously reported that heterozygous mutations in PIK3CD encoding the p110α catalytic PI3K subunit cause a unique disorder termed p110α-activating mutations causing senescent T cells, lymphadenopathy, and immunodeficiency (PASLI) disease. We report four patients from three families with a similar disease who harbor a recently reported heterozygous splice site mutation in PIK3R1, which encodes the p85α, p55α, and p50α regulatory PI3K subunits. These patients suffer from recurrent sinopulmonary infections and lymphoproliferation, exhibit hyperactive PI3K signaling, and have prominent expansion and skewing of peripheral blood CD8⁺ T cells toward terminally differentiated senescent effector cells with short telomeres. The PIK3R1 splice site mutation causes skipping of an exon, corresponding to loss of amino acid residues 434–475 in the inter–SH2 domain. The mutant p85α protein is expressed at low levels in patient cells and activates PI3K signaling when overexpressed in T cells from healthy subjects due to qualitative and quantitative binding changes in the p85α–p110α complex and failure of the C-terminal region to properly inhibit p110α catalytic activity.

Primary human immunodeficiency diseases offer insights into genes and pathways critical for host defense and healthy immune homeostasis. We and others have recently described a unique immune disorder featuring recurrent sinopulmonary infections, predisposition to chronic EBV and CMV viremia, lymphoproliferation, and increased lymphoma susceptibility (Angulo et al., 2013; Crank et al., 2014; Kracker et al., 2014; Lucas et al., 2014). Heterozygous gain-of-function mutations in the PIK3CD gene encoding the leukocyte-restricted p110α catalytic subunit of phosphatidylinositol 3-kinase (PI3K) are responsible for this disorder, which we have termed p110α-activating mutations causing senescent T cells, lymphadenopathy, and immunodeficiency (PASLI) disease (Lucas et al., 2014). PASLI disease is caused by mutation of at least four different sites in PIK3CD that drive hyper-activation of PI3K signaling in immune cells (Crank et al., 2014; Lucas et al., 2014). Some of the disease-causing amino acid substitutions in
p110β are identical to those occurring in tumor cells at homologous sites in PIK3CA encoding p110α, suggesting a similar molecular mode of action. Indeed, PASLI patients exhibit increased lymphoma risk that is further compounded by immunodeficiency leading to poor control of EBV viral loads (Crank et al., 2014; Kracker et al., 2014). We are now aware of ~80 PASLI patients worldwide, and the number of patients diagnosed with this disorder is expected to increase. Our previous work clearly established that hyperactivation of the PI3K signaling pathway causes immune dysregulation and raised the question of whether or not mutations in other PI3K genes would cause similar clinical manifestations by augmenting this pathway.

The phosphoinositide 3-kinase (PI3K) pathway transduces cell growth and proliferation signals through generation of the PIP3 second messenger, which is important for recruitment and activation of pleckstrin homology (PH) domain–containing signaling proteins. The class IA PI3Ks include the catalytic p110α, p110β, and p110δ proteins and the regulatory p85α, p55α, p50α, p85β, and p55γ proteins. The complex becomes activated upon recruitment to tyrosine-phosphorylated YXXM motifs with major signaling roles downstream of the insulin receptor, insulin-like growth factor-1 receptor, cytokine receptors, T cell receptor, and others. The class IA PI3Ks exist as a dimer of a catalytic and a regulatory subunit. The major roles of the regulatory subunit are to bind and stabilize p110 (Conley et al., 2012), inhibit p110 kinase activity (Burke et al., 2011), and recruit the PI3K complex to phosphotyrosine where binding of the SH2 domains to phosphotyrosine relieves the inhibitory (but not dimerizing) contacts with the catalytic subunit (Yu et al., 1998). There is debate about the existence and potential roles for free monomeric p85α that is not bound to p110 and its possible function in regulating PI3K activity (Geering et al., 2007b). Evidence against roles for free p85α includes the observation that monomeric p85α is relatively unstable (Brachmann et al., 2005; Zhao et al., 2006) and that p85α and p110 are obligate heterodimers normally present in the cell at 1:1 ratio (Geering et al., 2007a). Whether or not p85α can exist unbound to p110 and whether or not free p85α exerts biological or pathological effects remain open questions.

Studies in animal models have revealed a complex relationship between p110 and p85α (Vanhaesebroeck et al., 2005). The total PIK3R1 knockout mouse dies in the perinatal

Table 1. Clinical summary

<table>
<thead>
<tr>
<th>Features</th>
<th>A.1</th>
<th>B.1</th>
<th>B.II.1</th>
<th>C.1</th>
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<td>Recurrent lymphoma; HSCT</td>
<td>Cystic kidneys; poor growth</td>
<td>Hepatomegaly; sepsis; poor growth</td>
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LAD: lymphadenopathy; ct: count; arrows indicate high and low values, respectively.
period and shows secondary loss of p110 catalytic protein (Fruman et al., 2000). Mice heterozygous for p85α have normal levels of p110 and show greater insulin-stimulated PI3K activity than WT counterparts but display no overt immunological phenotypes (Ueki et al., 2002; 2003; Vanhaesebroeck et al., 2005). Two inherited human diseases have been associated with mutations in the PIK3R1 gene: (1) SHORT syndrome, a disease of short stature, hyperextensible joints, Rieger anomaly of the eye, teething delay, lipoatrophy, and often insulin resistance, caused by heterozygous PIK3R1 mutations (Chudasama et al., 2013; Dymant et al., 2013; Thauvin-Robinet et al., 2013; Bárcena et al., 2014); and (2) agammaglobulinemia due to absent B cells caused by a homozygous PIK3R1 mutation that leads to loss of p85α with secondary loss of p110 (Conley et al., 2012). Somatic, heterozygous mutations in PIK3R1 have also been found in human glioblastoma (Cancer Genome Atlas Research Network, 2008; Parsons et al., 2008) and colon cancer (Jaiswal et al., 2009). These mutations reduce inhibition of p110 by p85α, leading to hyperactive PI3K signaling and tumorigenesis (Jaiswal et al., 2009; Sun et al., 2010). More recently, somatic PIK3R1 mutations in endometrial carcinoma were discovered that cluster mostly within amino acid residues 434–475 in the inter-SH2 domain of p85α and augment PI3K signaling (Urick et al., 2011). These findings in mice and previously described human diseases shed light on the various physiological roles of PIK3R1 gene products and support the hypothesis that cancer-related PIK3R1 gene mutations could be a driver of PASLI-like disease.

We have now discovered heterozygous PIK3R1 splice site mutations in patients with PASLI-like disease characteristics and striking hyperactivation of PI3K signaling in immune cells. An independent report has also recently described similar patients with the same splice site mutation (Deau et al., 2014). Here, we not only describe the clinical findings and gene defect in these patients but also provide biochemical evidence that the mutant p85α protein is expressed in patient cells, associates normally with p110α, and dominantly drives constitutive PI3K signaling due to loss of inhibitory contacts, which results in cellular derangements that contribute to immunodeficiency in this patient population. These findings further provide a possible treatment option for these patients using approved or investigational drugs that target PI3K or its downstream effectors (i.e., mTOR inhibition with rapamycin).

RESULTS

Immunodeficiency with poor antibody production, lymphoproliferation, and inflammatory disease

Our evaluation of the index patient A.1, a 32-yr-old female, revealed recurrent sinopulmonary infections with associated bronchiectasis, diffuse lymphadenopathy with splenomegaly, refractory immune thrombocytopenia purpura treated with splenectomy, and development of arthritis and inflammatory bowel disease (IBD) in adulthood (Table 1). Analysis of her blood cells revealed T lymphocytosis as the primary abnormality, elevated erythrocyte sedimentation rate (ESR), and C-reactive protein (CRP). The patients in family B suffer from a similar disease with immunodeficiency, lymphoproliferation, and poor production of class-switched antibody. Patient B.1 developed Hodgkin lymphoma and has undergone hematopoietic stem cell transplantation (HSCT), and...
The three families in our cohort demonstrate a dominant or de novo disease inheritance pattern, and whole-exome sequencing (WES) uncovered a heterozygous point mutation in PIK3R1 at a splice donor site (G > C or G > A at chr5:67589663, which corresponds to c.1425+1 in NM_181523.2) in genomic DNA from PBMCs in patients A.1, B.II.1, and C.1 (Fig. 1 A). Due to her HSCT, genomic DNA was isolated from fingernail clippings from patient B.1 to confirm the presence of the mutation. The PIK3R1 mRNA variant encoding p85β contains 16 exons beginning with a noncoding exon 1 in the 5’ UTR, and the patient splice site mutation is in the first intronic nucleotide after exon 11. The mutation results in skipping of exon 11 (delE11), as revealed by sequencing of PIK3R1 cDNA (Fig. 1 B). As previously reported (Deau et al., 2014), loss of exon 11 led to an in-frame deletion of the nucleotides encoding amino acid residues 434–475 of p85β, which falls in the inter-SH2 domain shared by the p85β, p55α, and p50α PIK3R1 isoforms (Fig. 1 C). This deletion is immediately N-terminal to the mapped binding site (residues 478–513) of the p110 catalytic subunit (Dhand et al., 1994). Structural modeling of the truncated inter-SH2 domain showed loss of part of the coiled coil structure of the inter-SH2 domain and predicted a dramatic change in the position of the nSH2 (positioned in the lower left in WT and shifted to the top in delE11) and cSH2 domains that are required for inhibitory contacts with p110 and recruitment to phosphotyrosine motifs (Fig. 1 D). Such a marked alteration in the inter-SH2 domain and the positioning of the SH2 domains strongly suggests that there will be qualitative and/or quantitative alterations in the contacts made between delE11 p85β and the p110 catalytic subunit. Moreover, all three PIK3R1 gene products, the p85β, p55α, and p50α regulatory subunits, will be affected by this structural alteration that is predicted to impair p110 inhibition (Fig. 1 C).

### PI3K signaling is hyperactive and constitutive

We next sought to directly examine PI3K signaling in these patients. PI3K activity generates PIP3, which is recruited to the membrane at signaling sites by binding
The mutant p85α protein is detectable at low levels in patient T cell blasts and drives PI3K hyperactivation due to altered interaction with p110β

We next sought to examine expression of the mutant p85α protein to determine if the in-frame deletion produces a stable protein product. Detection of endogenous p85α by immunoblot in lysates from patient T cell blasts revealed a reduction in full-length p85α compared with controls (Fig. 3 A). Importantly, a band at the predicted size of 78 kD was detectable in patient lanes (Fig. 3 A, bottom arrow). However, the truncated delE11 mutant form of p50α was undetectable (Fig. 3 B).

The low-level mutant p50α protein expression despite the clear mRNA expression (unpublished data) suggests that the mutant protein is unstable. Examination of PI3K signaling after siRNA-mediated knockdown of PIK3R1 showed no evidence for haploinsufficiency of PIK3R1 contributing to hyperactivation of PI3K in primary human T cells (unpublished data). Moreover, we observed normal expression of the PTEN and SHIP1 phosphatases in patient T cell blasts, excluding the possibility that reduction in expression of the enzymes that counteract PI3K accounted for increased signaling (unpublished data). We next performed overexpression experiments in which WT versus delE11 5×-myc–tagged p85α or p50α were analyzed by immunoblot for myc tag, p110β, phospho-AKT, or total AKT, as indicated. Arrow marks phospho-AKT band, and asterisks mark residual signal from p50α–myc blot re-probed for phospho-AKT. Four (A and C) or three (B, D, and E) independent experiments were performed.

The delE11 PIK3R1 mutant p85α gene product is expressed at low levels in patient T cell blasts and drives PI3K hyperactivation via poor inhibition of p110β. (A) Western blot for p85α (Ab recognizing region just after SH3 domain), p110β, or β-tubulin, as indicated. Arrows point to full-length (top arrow) and truncated delE11 (bottom arrow) p85α. (B) Western blot analysis of p50α using an antibody recognizing the N-terminal SH2 (top). Levels of p110β protein (middle) and β-tubulin loading control (bottom) are also shown. (C) Primary T cells overexpressing EV, WT, or delE11 mutant 5x-myc–tagged p85α were analyzed by immunoblot for myc tag, p110β, phospho-AKT, or total AKT, as indicated. (D) p110β immunoprecipitates (IP, top) or input (In) and flow-through (FT, bottom) from T cells overexpressing EV, WT p85α–myc, or delE11 p85α–myc were probed with the indicated immunoblot (IB) antibodies. (E) Primary T cells overexpressing EV, WT, or delE11 mutant 5x-myc–tagged p85α or p50α were analyzed by immunoblot for myc tag, phospho-AKT, total AKT, or actin, as indicated. Arrow marks phospho-AKT band, and asterisks mark residual signal from p50α–myc blot re-probed for phospho-AKT. Four (A and C) or three (B, D, and E) independent experiments were performed.

Figure 3.
delE11 p85α in patient cells because the p85α monomer has been shown to be unstable (Brachmann et al., 2005; Zhao et al., 2006). These data, together with the observation that p110δ protein levels are relatively normal in these patients, indicate that the delE11 p85α protein expressed in patient cells can associate with p110δ and stabilize it, though the affinity of the interaction is reduced compared with WT because the complexes fail to remain intact in immunoprecipitation procedures.

The experiments above shed light on the stability of the association between p85α and p110δ, which dimerize via stable contacts mediated by the inter-SH2 domain of p85α and the adaptor-binding domain of p110δ. However, the data above do not address the additional inhibitory contacts normally made between the SH2/inter-SH2 domains of p85α and various regions of p110δ. We assessed the contribution of the SH2/inter-SH2 domains of the delE11 p85α mutant in driving PI3K signaling by overexpressing myc-tagged p50α protein containing only these domains. When WT and delE11 forms of the C-terminal p50α fragment were overexpressed, we observed that delE11 p50α was sufficient to increase AKT phosphorylation on serine 473 (Fig. 3 E). These data rule out a role for the N-terminal domains (i.e., the SH3, PRR, and BH domains) of p85α in mediating the dominant gain-of-function effect on PI3K activity and instead point to a failure of delE11 p85α to inhibit p110δ activity due to alterations in the structure of the SH2 and inter-SH2 domains that normally form inhibitory contacts with the p110 subunit. Thus, the heterozygous PIK3R1 splice mutation results in expression of mutant p85α that is sufficient to dominantly hyperactivate the PI3K pathway due to qualitatively and quantitatively different binding and impaired inhibition of p110δ.

Patient PBMCs are enriched with CD8 T cells that are terminally differentiated and have shortened telomeres

Because the PIK3R1 mutation in these patients drives PI3K signaling, we expected to see similar cellular derangements as previously observed in PASLI patients with gain-of-function mutations in PIK3CD encoding p110δ. We therefore evaluated patient PBMCs and found several of the key cellular features of PASLI disease. CD8 T cells were increased in frequency and deficient in naive (CD45RA+CCR7+) cells with features of PASLI disease. CD8 T cells were increased in frequency in patient PBMCs and found several of the key cellular mutations in previously observed in PASLI patients with gain-of-function signaling, we expected to see similar cellular derangements as based fluorescence in situ hybridization (flow-FISH) and mere length in the lymphocyte population by flow cytometry– (Fig. 4 C). To extend these cellular findings, we assessed telomere differentiation and senescence of the CD8 T cell population (Fig. 4 C). To extend these cellular findings, we assessed telomere length in the lymphocyte population by flow cytometry– based fluorescence in situ hybridization (flow–FISH) and observed severely shortened telomere length in PIK3R1 patients (Fig. 4 D). Thus, in PIK3R1–driven disease as in p110δ PASLI disease, hyperactive PI3K signaling in lymphocytes causes expansion, terminal differentiation, and senescence of CD8 T cells.

The p110δ catalytic subunit accounts for hyperactivation of PI3K signaling and its inhibition offers potential for therapeutic benefit

To further validate our hypothesis that the PIK3R1 mutation underlies the disease pathogenesis in these patients and causes hyperactivation of PI3K and mTOR in lymphocytes, we next
DISCUSSION

The PI3K signaling pathway plays critical roles in virtually all cell types and has been intensively studied because of its prominent role in promoting tumorigenesis. It has become clear that either too little or too much PI3K signaling can be pathological. Mice completely deficient in the PIK3R1 gene products die in the perinatal period and exhibit liver necrosis and hypoglycemia, whereas mice lacking just the p85α isoform are viable but suffer from hypoglycemia and increased insulin sensitivity linked to up-regulation of the p55α and p50α isoforms (Fruman et al., 2000). Hyperactivation of PI3K signaling has been identified as a cause of human diseases including megalencephaly syndromes (Lee et al., 2012; Rivière et al., 2012) and overgrowth syndromes (Lindhurst et al., 2012; Keppeler-Noreuil et al., 2014). A vast literature also places PI3K among the top pathways driving cancer cell growth and survival through gain-of-function mutation of PI3K genes, most commonly PIK3CA encoding the p110α catalytic subunit, or mutations activating upstream initiators of PI3K signaling (Wong et al., 2010).

Although most studies of PI3K signaling in disease states focus on the catalytic subunit, recent reports have clearly highlighted the important role that the regulatory subunit can play in modulating PI3K activity. Recently, several independent groups have identified heterozygous mutations in PIK3R1 in patients with SHORT syndrome, a rare disorder characterized by short stature, hyperextensible joints, ocular depression, Rieger anomaly of the eye, teething delays, and characteristic facial gestalt (Chudasama et al., 2013; Dyment et al., 2013; Thauvin-Robinet et al., 2013; Bárcena et al., 2014). The PIK3R1 mutations in these patients cluster predominantly in the cSH2 domain and the biochemical effect of these heterozygous mutations appears to be a reduction in insulin-stimulated PI3K activity (Chudasama et al., 2013; Dyment et al., 2013; Thauvin-Robinet et al., 2013; Bárcena et al., 2014). No severe immune abnormalities have been reported in patients with SHORT syndrome, and the clinical features of SHORT syndrome are not found in our PIK3R1 patients with PASLI-like immune disease. These observations raise the possibility that the C-terminal SH2 domain of p85α plays a unique role in binding tyrosine-phosphorylated insulin receptor substrate-1 (IRS-1) during insulin signaling, leading to SHORT syndrome when mutated. In contrast, the N-terminal SH2 and inter-SH2 domains of p85α appear to be particularly important in inhibiting catalytic activity of p110β because their alteration in our heterozygous PIK3R1 patients causes PASLI-like immune disease via hyperactivation of p110β.

Other PIK3R1 findings particularly relevant for the studies described here are reports describing patients with homozygous loss of p85α (Conley et al., 2012) and tumors with heterozygous gain-of-function PIK3R1 variants that augment PI3K signaling (Jaiswal et al., 2009; Sun et al., 2010; Urick et al., 2011). Conley et al. (2012) described a patient with colitis and absent B cells with normal T cells who harbored a homozygous loss-of-function mutation in PIK3R1 resulting in absence of p85α protein with normal expression of the p50α and p55α splice variants. The mutation results in a premature stop codon in the BH domain that is present in the p85α but not the p55α or p50α isoforms of PIK3R1. Due to the lack of the regulatory p85α protein, a loss in expression of the catalytic p110β subunit was observed, emphasizing the requirement for p85α to stabilize p110β even when p55α and p50α are present. Interestingly, loss of p85α led to a dramatic decrease in p110β with a more subtle decrease in expression of p110α and p110β (Conley et al., 2012). Thus, p85α/p110β are required for B cell development, and the
N-terminal region of p85α plays a particularly important role in stability of p110δ (more so than the other class IA catalytic subunits). The heterozygous parents of this patient were not reported to be clinically affected, providing further support that haploinsufficiency of p85α is not a driver of PI3K hyperactivation in the immune system. In contrast to p85α loss of function, heterozygous mutations in PIK3R1 that drive hyperactivation of PI3K have been identified in human cancers and found to have no effect on p110 protein expression levels (Jaiswal et al., 2009; Sun et al., 2010; Urick et al., 2011). In endometrial cancer, these mutations cluster in the inter-SH2 domain of p85α around amino acid residues 434–475, which are precisely the residues deleted by skipping of exon 11 in our patients, and the PIK3R1 mutants drive hyperactivation of PI3K signaling (Urick et al., 2011).

Here, we describe patients who suffer from immunodeficiency with lymphoproliferation, antibody defects, and terminal differentiation and senescence of CD8 T cells. We identified a heterozygous splice site mutation that leads to in-frame deletion of exon 11 of PIK3R1 and hyperactivation of PI3K and AKT. Similar patients with the same splice site mutation have now been independently reported (Deau et al., 2014). As in Deau et al. (2014), we describe a similar patient phenotype, mutation site, exon skipping, hyperphosphorylation of AKT and S6, and ability of overexpressed delE11 p85α to induce hyperphosphorylation of AKT. We further show that other AKT substrates including FOXO and GSK3 proteins are hyperphosphorylated in patient T cells. Most of the clinical features of the disease observed in our patients and those reported by Deau et al. (2014) are consistent, with lymphoproliferation and autoimmune/inflammatory disease being the two exceptions. We observed marked lymphoproliferation and reduced CD4/CD8 ratios in our patients, whereas only one of four patients reported by Deau et al. (2014) is reported to have evidence of lymphoproliferation (enlarged tonsils) and two of four patients have reduced CD4/CD8 ratios. Additionally, two of our four patients had arthritis, whereas none was reported by Deau et al. (2014). In both patient cohorts, recurrent sinopulmonary infections, immunoglobulin defects, and reduction in naive T cells are key characteristics of the disease. The molecular basis of PI3K hyperactivation and metabolic effects of cellular phenotype were not elaborated in Deau et al. (2014). We now report the mechanism by which deletion of amino acid residues 434–475 of p85α augments PI3K signaling and alters T cell phenotype and function, and our major conclusions are discussed below.

Normal T cells robustly express the p85α and p50α isoforms but not the p55α isoform of PIK3R1. Analysis of patient T cell lysates by immunoblot for PIK3R1 protein products using an antibody against the nSH2 domain revealed, as expected, that full-length WT p85α and p50α were reduced in protein abundance. A truncated delE11 p85α band was observed at the expected size of 78 kD; however, the delE11 p50α mutant protein could not be detected despite clear mRNA expression (unpublished mRNA data). These observations left open the possibilities of gain-of-function effects on PI3K signaling mediated by low-level expression of delE11 p85α or a haploinsufficiency effect due to the reduction in expression of full-length WT p85α and p50α. Although haploinsufficiency of PIK3R1 has been shown previously to increase PI3K signaling in response to insulin (Mauvais-Jarvis et al., 2002), we ruled out a major contribution of PIK3R1 haploinsufficiency in this immune cell phenotype using siRNA-mediated knockdown of PIK3R1 in T cells from healthy control subjects. These experiments showed no effect on levels of basal AKT phosphorylation, which is consistent with previous reports that insulin signaling is unique in its hyperresponsiveness caused by PIK3R1 haploinsufficiency (Vanhaesebroeck et al., 2005). In contrast, when delE11 p85α was overexpressed in control T cells, we observed a clear increase in basal AKT phosphorylation relative to the WT p85α transfectants. This finding supports the conclusion that expression of the delE11 variant of p85α in patient immune cells is a dominant driver of PI3K hyperactivation.

To understand how delE11 p85α drives exuberant PI3K signaling, we examined the ability of this mutant protein to associate with the p110δ catalytic subunit because the deleted exon encodes residues adjacent to the p110 binding site. These experiments demonstrated a detectable but reduced association between endogenous p110δ and overexpressed delE11 p85α, whereas the association with overexpressed WT p85α was robust. This reduced association offers an explanation for the low-level expression of delE11 p85α in patient lysates because p85α is unstable if not bound to p110 (Brachmann et al., 2005; Zhao et al., 2006). How can delE11 p85α promote p110δ hyperactivation despite less stable interaction with p110δ? We evaluated three major possibilities: (1) structural effects of the deletion result in increased recruitment of the PI3K complex to phosphotyrosine; (2) delE11 p85α that is free from p110 interacts with other signaling molecules via its N-terminal SH3 domain, proline-rich regions, and/or BH domain to promote PI3K activity; and/or (3) the ability of delE11 p85α to form inhibitory contacts with p110δ is impaired. To address (1), we evaluated association of myc-tagged WT or delE11 p85α with LAT, which recruits p85α once it is tyrosine phosphorylated upon TCR engagement, and found that WT and delE11 p85α were similarly associated with LAT (unpublished data). We next addressed possibilities (2) and (3) by cloning and overexpressing the C-terminal fragment of WT and delE11 p85α containing only the nSH2, inter-SH2, and cSH2 domains (i.e., p50α) that normally form inhibitory contacts with p110. In T cells from healthy subjects, we found that overexpression of delE11 p50α was sufficient to cause hyperphosphorylation of AKT at serine 473. This result rules out a requirement for the N-terminal SH3, PRR, and/or BH domains of “free” delE11 p85α in promoting PI3K activity. Instead, our findings are consistent with a failure of the SH2 and inter-SH2 domains of delE11 p85α to properly inhibit p110δ due to structural derangements that interfere with formation of inhibitory contacts.

We now recognize two classes of PASLI disease distinguished by the gene underlying hyperactive PI3K signaling. We term these classes PASLI-CD and PASLI-R1 for the forms...
caused by mutations in PIK3CD and PIK3R1, respectively. The clinical and cellular consequences of hyperactive PI3K signaling in immune cells were revealed by studies of PASLI-CD patients harboring heterozygous, gain-of-function mutations in PIK3CD encoding p110δ (Angulo et al., 2013; Lucas et al., 2014). The clinical presentations of the PASLI-R1 patients described here and in a recently published report (Deau et al., 2014) are remarkably similar to PASLI-CD patients, with recurrent sinopulmonary infections, poor antibody responses requiring IVIG, susceptibility to EBV and CMV, lymphoproliferation, and increased incidence of lymphoma. At the cellular level, we now show that, as in PASLI-CD patients, this new cohort of PASLI-R1 patients have increased glucose uptake, expanded CD8 T cells with significant reduction in the frequency of naive cells, and concomitant increase in the frequency of terminally differentiated effector-type T cells (Lucas et al., 2014). The CD57 senescence marker is also highly expressed on CD8 T cells in PIK3R1 patients, and we extended this finding by determining that telomeres in lymphocytes are markedly shorter than expected for the age of the patient. Although presently these diseases appear to be very similar, following the natural history of these patients may reveal subtle clinical and/or molecular differences.

Given the molecular etiology of this disease, treatment with drugs inhibiting the PI3K signaling pathway would offer the most ideal therapeutic option for these patients. Indeed, we found that inhibition of mTOR using rapamycin in vitro robustly inhibited the hyperphosphorylation of S6 but had no effect on AKT hyperphosphorylation in PASLI-R1 T cell blasts. Because other targets of AKT besides mTOR (e.g., FOXO and GSK3α/β) are also hyperphosphorylated, a more ideal approach would be to specifically target select PI3K isoforms while avoiding negative consequences of global PI3K inhibition. Although T cells express all three class IA catalytic subunits, we were intrigued to find that inhibition of the leukocyte-restricted p110δ catalytic subunit alone was sufficient to reduce the hyperphosphorylation of AKT and S6 in PIK3R1 patient T cell blasts. This has the interesting biological implication of p85α-mediated regulation of p110δ being unique and selectively affected by loss of PIK3R1 exon 11. Given these findings and the predominantly immunological clinical disease in these patients, specific targeting of p110δ using inhibitors such as those currently in clinical trials for hematological malignancy (Lannutti et al., 2011; Brown et al., 2014; Flinn et al., 2014; Gilbert, 2014; Kahl et al., 2014) offers a potentially safe and effective treatment option to restore immune homeostasis while limiting unwanted effects in nonimmune cells that do not express p110δ.

MATERIALS AND METHODS

Human subjects. All human subjects (or their legal guardians) in this study provided written informed consent in accordance with Helsinki principles for enrollment in research protocols that were approved by the Institutional Review Board of the National Institute of Allergy and Infectious Diseases, National Institutes of Health (NIH). Blood from healthy donors was obtained at the NIH Clinical Center under approved protocols. Mutations will be automatically archived by Online Mendelian inheritance in Man (OMIM) once the paper is published, and whole-exome data will be submitted in dbGaP. We will also create a webpage through National Center for Biotechnology Information (NCBI) to accumulate patient mutation data in the format of the Leiden Online Variant Database (LOVD) as patients are identified.

DNA sequencing. Genomic DNA was isolated from PBMC for proband A.1 and family members. SureSelect Human All Exon 50 Mb kit (Agilent Technologies) coupled with massively parallel sequencing by Illumina HiSeq Sequencing System was performed using the collected DNA. For individual samples, WES produced ~50–100× sequence coverage for targeted regions. Similar WES was also performed for patient B.II.1 and C.1. For confirmation of mutations in the patients, genomic DNA was PCR amplified and Sanger sequencing of purified PCR amplified products was performed. For cDNA preparation, total RNA was extracted from T cell blasts and cDNA was produced using an oligo-dT primer. The PIK3R1 gene was amplified from the cDNA and the full-length and truncated bands were excised from an agarose gel for Sanger sequencing to detect joining of exon 10 with exon 12.

Whole-exome sequence analyses. All sequenced DNA reads were mapped to the hg19 human genome reference by Burrows-Wheeler Aligner with default parameters. Single nucleotide variant and indel calling were performed using the Genome Analysis Toolkit (the Broad Institute). All SNVs/indels were annotated by SeattleSeq Annotation and an in-house custom analysis pipeline was used to filter and prioritize for autosomal recessive or de novo disease-causal variants based on the clinical pedigree for patient A.1. For patient B.II.1 and C.1, the mutations were identified by targeted gene screening of the WES database on the similarity of clinical phenotype in the cohort.

Protein structure modeling. The structure of the delE11 p85α N-SH2 and inter-SH2 domains was generated by manual editing of p85α coordinates obtained from the complex of p110α and p85α (PDB ID: 3HHM), which were regularized (i.e., disordered amino acids were interpolated) using SWISS-MODEL. The p85α N-SH2 domain and extant region of the coiled-coil domain were treated as rigid bodies, and were fused together by manual transposition of the SH2 domain to the new terminal residue of the coiled-coil. The relative orientations of the domains were then modified to restore steric clashes with the cognate p110δ subunit of PI3K, modeled using I-TASSER and ModRefiner. All manual modeling and subsequent rendering was performed using PyMOL.

Cell culture and transfection. Human PBMCs were isolated by Ficoll-Paque PLUS (GE Healthcare) density gradient centrifugation, washed twice in PBS, and resuspended at 106 cells/ml complete RPMI 1640 (rRPMI) medium (Lonza) containing 10% FBS, 2 mM glutamine, and 100 U/ml each of penicillin and streptomycin (Invitrogen). Cells were activated with 1 µg/ml anti-CD3 (clone HIT3a; BD) and 1 µg/ml anti-CD28 (clone CD28.2; BD). After 3 d, activated T cells were washed and then cultured in complete rRPMI-1640 medium supplemented with 100 U/ml recombinant human IL-2 (shl-2; R&D Systems). Transfection was performed with Amaxa Nucleofection kits (Lonza) for primary cells, using C-terminally 5×-myc-tagged constructs in pcDNA3.1.

Flow cytometry. For standard surface staining, PBMCs (106 cells/sample), sorted cells, expanded T cells, or cell lines were washed with PBS and incubated for 30 min at 4°C (dark) in 100 µl 5% FBS in PBS with indicated fluorochrome-labeled monoclonal antibodies or their isotype controls. After washing with PBS two times, 10,000–50,000 live cells were analyzed by flow cytometry. For phospho-flow staining, unless otherwise indicated, cells were kept in rRPMI while alive, fixed directly in rRPMI using Lyse/Fix (BD), and then permeabilized with perm buffer III (BD) according to manufacturer’s instructions. The indicated antibodies were purchased from BD, eBioscience, BioLegend, or Cell Signaling Technology. For phospho-flow analyses, anti-pAKT S473 Alexa Fluor 647 (J9E; Cell Signaling Technology), anti-pAKT T308 Alexa Fluor 488 (C31E5E; Cell Signaling Technology), anti-pS6 S235/236 PE (N7-548; BD), and/or anti-pS6 S240/244 Alexa Fluor 647 (D68F8; Cell Signaling Technology) were used. Rabbit isotype control staining was performed for each experiment (Fig. 2 B, left, with Fig. 2 C, top; Fig. 2 B, right, with Fig. 2 C, bottom; and Fig. 5).
Immunoblotting and immunoprecipitation. Cells were washed in PBS or RIPA buffer with no FCS and immediately lysed on ice in 1% Triton X-100, 50 mM Tris-Cl, pH 8, 150 mM NaCl, 2 mM EDTA, 10% glycerol, complete protease inhibitor cocktail (Roche), and phosphatase inhibitor cocktails (Sigma-Aldrich). Protein was quantitated by BCA assay (Thermo Fisher Scientific). The lysates were then clarified by centrifugation at 15,000 g at 4°C for 10 min. Supernatants were transferred to a separate tube and used for subsequent experimentation. Approximately 20 μg of total protein was separated by SDS-PAGE and transferred to a nitrocellulose membrane (Bio-Rad Laboratories). Membranes were blocked with 5% nonfat dry milk in Tris-buffered saline (TBS) with 0.01% Tween-20 (TBST) for 1 h at room temperature before incubating with primary antibody overnight at 4°C. After washing with TBST for 1 h at room temperature, HRP-conjugated secondary antibody was added for an additional hour at room temperature. After a final 1-h wash step, HRP substrate (Luminata Forte; Millipore) was added to the membranes, which were then subjected to chemiluminescent imaging. Validated primary antibodies were purchased from Cell Signaling Technology, Santa Cruz Biotechnology, Inc., or Millipore, and secondary antibodies were from SouthernBiotech. For immunoprecipitation (IP) of p110δ, cells overexpressing empty vector (EV), WT p85α, or delE11 p85α were lysed as described above, precleared of proteins that nonspecifically bind anti-rabbit IgG IP beads (Rockland Immunochemicals), and incubated overnight with rabbit anti-p110δ IgG (Santa Cruz Biotechnology, Inc.). Anti-rabbit IgG IP beads were then added to capture the immune complexes, and samples were rotated for 1 h at 4°C. Captured immune complexes were pelleted by centrifugation, washed three times with lys buffer, and eluted by boiling after addition of SDS-containing sample buffer with reducing agent.

Glucose uptake. Cells were starved of glucose by incubation in PBS for 1 h before incubation with 100 μg/ml 2-deoxy-2-[(7-nitro-2,1,3-benzoxadiazol-4-yl)amino]-d-glucose (2-NBDG) for 20 min. Uptake was measured by flow cytometric evaluation of signal in the fluorescein channel.

Flow-FISH for telomere length assessment. Fresh blood from patient A.1 was shipped directly to Repeat Diagnostics for measurement of telomere length by flow cytometry, gating on either lymphocytes or granulocytes. A fluorescently labeled nucleic acid probe was used to hybridize with the TTAGGG repeats in telomeres, and fluorescence intensity provided a measure of telomere length. Telomere length measurements for patient lymphocytes and granulocytes were plotted relative to telomere lengths established for healthy control subjects with percentiles indicated.

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Contributions: C.L. Lucas performed experiments, analyzed data, and developed and wrote the manuscript. Y. Zhang analyzed WES data and discovered PIK3R1 mutations. A. Venida assisted with experiments and analyzed data. Y. Wang provided clinical care for and analysis of patient C.1. J. Hughes performed/organized WES studies and analyzed data. J. McElwee performed/organized WES studies and analyzed data. M. Buttrick assisted with clinical management of patient A.1. H. Matthews assisted with clinical management of patients A.1, B.1, and B.II.1. S. Price assisted with clinical care of patient A.1. M. Biancalana performed structural modeling analyses. X. Wang provided clinical care for and supervised analysis of patient C.1. M. Richards provided clinical care for and evaluation of patient B.II.1. T. Pozos provided clinical care for and evaluation of patient B.II.1. I. Barlan provided clinical care for and evaluation of patient A.1. V.K. Rao provided clinical care for and evaluation of patient A.1. H.C. Su provided clinical and research advice and supervised research and data analysis. M.J. Lenardo supervised research and data analysis, provided advice, and edited the manuscript. All authors discussed and reviewed the manuscript.

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