Living with Lethal PIP3 Levels: Viability of Flies Lacking PTEN Restored by a PH Domain Mutation in Akt/PKB

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The phosphoinositide phosphatase PTEN is mutated in many human cancers. Although the role of PTEN has been studied extensively, the relative contributions of its numerous potential downstream effectors to deregulated growth and tumorigenesis remain uncertain. We provide genetic evidence in Drosophila melanogaster for the paramount importance of the protein kinase Akt [also called protein kinase B (PKB)] in mediating the effects of increased phosphatidylinositol 3,4,5-trisphosphate (PIP3) concentrations that are caused by the loss of PTEN function. A mutation in the pleckstrin homology (PH) domain of Akt that reduces its affinity for PIP3 sufficed to rescue the lethality of flies devoid of PTEN activity. Thus, Akt appears to be the only critical target activated by increased PIP3 concentrations in Drosophila.

Mutations in the tumor suppressor gene PTEN (the phosphatase and tensin homolog on chromosome 10) are frequent in glioblastomas, endometrial carcinoma, melanomas, and prostate cancer (1). Furthermore, two dominant hamartoma syndromes, Cowden disease and Bannayan-Zonana syndrome, are linked to germ line mutations in PTEN (1). The PTEN protein carries a phosphatase domain resembling those of dual-specificity protein phosphatases (2–4). Although it can dephosphorylate protein substrates such as focal adhesion kinase (5) and the adapter protein Shc (6), PTEN’s predominant enzymatic activity appears to be the dephosphorylation of phosphoinositides at the D3 position. Because PTEN uses the second messenger PIP3 as a substrate, PTEN antagonizes the function of phosphatidylinositol-3 kinase (PI3K) (7, 8). Immortalized mouse embryonic fibroblasts or embryonic stem cells lacking PTEN function show an approximately twofold increase in PIP3 concentrations (9, 10). PIP3 interacts with a wide variety of PH domain-containing proteins, including the serine-threonine kinases Akt (also called PKB) and phosphoinositide-dependent kinase 1 (PDK1), Btk family tyrosine kinases, guanine nucleotide exchange factors for the Rho and Arf families of small guanosine triphosphatases, and phospholipase C

Fig. 1. Reduced kinase activity caused by an amino acid substitution in the PH domain of dAkt. (A) Effect of the G99S substitution in the PH domain on dAkt kinase activity from larval extracts (42). Activity from wild-type larvae was considered to be 100%. Inset, dAkt protein was detected in 40 μg of larval extracts using the same antiserum. (B) Reduced insulin-induced activation of the G99S mutant dAkt. The dAkt constructs were expressed in HEK 293 cells (43). Transfected cells were starved for 24 hours before stimulation with insulin for the indicated time periods, and dAkt kinase activity was determined (44). The activity of wild-type dAkt from unstimulated cells was considered to be relative activity = 1.

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additional phenotypes associated with mutations in PTEN that cannot easily be reconciled with an exclusive function of PTEN in insulin receptor signaling (for example, the burst vulva phenotype in C. elegans (13) and defects in the actin cytoskeleton in Drosophila (17)). To better understand the consequences of loss of PTEN function, it would be useful to know which important downstream effectors react to increased PIP3 concentrations and whether PTEN has other physiological substrates in addition to PIP3.

The protein kinase Akt is an important component of insulin receptor signaling (20). Akt is recruited to the plasma membrane by virtue of the interaction of its NH$_2$-terminally located PH domain with PIP3. At the membrane, subsequent phosphorylation events by PDK1 and an unidentified kinase lead to the full activation of Akt (21–23). In PTEN-deficient mouse embryonic fibroblasts and embryonic stem cells, Akt is phosphorylated and activated (9, 10). The phenotypes associated with Akt mutations in both C. elegans and Drosophila are consistent with its role in signal transduction downstream of the insulin receptor (24–27).

We monitored three properties of Drosophila Akt (dAkt) separately: kinase activity, abundance of the protein, and membrane localization. We relied entirely on mutations in the endogenous gene encoding dAkt to avoid potential side effects caused by overexpression of mutant proteins. dAkt encodes a catalytically inactive protein, dAktF327I (25). The viable dAkt allele contains a P-element insertion upstream of the dAkt gene and therefore results in the reduced expression of wild-type dAkt protein (19, 28). Finally, we characterized the viable hypomorphic mutation dAkt$^4$ (29) that selectively impairs the membrane recruitment of dAkt in response to increased concentrations of PIP3. Sequencing of genomic DNA extracted from dAkt$^4$ homozygous flies revealed a single nucleotide exchange resulting in the substitution of a serine residue for a nonconserved glycine at the end of the sixth $\beta$ sheet of the PH domain. To address the mechanism by which this Gly$^{99} \rightarrow$ Ser$^{99}$ (G99S) mutation in the PH domain affects dAkt, we compared the amount of dAkt protein and activity in wild-type and dAkt$^4$ mutant larvae. Whereas no apparent difference in expression of the protein was observed (Fig. 1A, inset), dAkt activity from the mutant larvae represented only 30% of that in wild-type larval extracts (Fig. 1A). We also expressed epitope-tagged forms of wild-type dAkt, catalytically inactive dAktF327I, and PH domain mutant dAktG99S in insulin-responsive human embryonic kidney (HEK) 293 cells. All three proteins were expressed in similar amounts (30). dAktG99S activity from insulin-stimulated cells was reduced by about 90% as compared to that of the wild-type kinase (Fig. 1B). All forms of dAkt proteins were detected in the cytosol of unstimulated cells (Fig. 2, A, D, and G). Stimulation of the cells with insulin for 5 min resulted in association of the wild-type and the catalytically inactive enzymes with the plasma membrane, but failed to recruit the dAktG99S mutant protein (Fig. 2, B, E, and H). In contrast, treatment of HEK 293 cells with the protein-tyrosine phosphatase inhibitor pervanadate, a potent activator of Akt (31), led to membrane recruitment of all dAkt proteins (Fig. 2, C, F, and J).

**Fig. 2.** Reduced membrane localization of the G99S mutation of dAkt. HEK 293 cells plated on coverslips were transfected with epitope-tagged wild-type (A to C), G99S (D to F), and F327I (G to J) dAkt and deprived of serum for 16 hours before stimulation with insulin (B, E, H) or pervanadate (C, F, J) for 5 min. Fixed and permeabilized cells were incubated first with the monoclonal antibody 12CA5 to the HA epitope and then with fluorescein isothiocyanate–conjugated secondary antibody. An analysis by confocal microscopy revealed the subcellular localization of the dAkt variants.

**Fig. 3.** Restored viability of flies lacking dPTEN function by the PH domain mutation in dAkt. (A) Morphology and weight of dPTEN mutant flies rescued by $dAkt^3/dAkt^1$. The left panel shows female flies, the middle panel shows male flies, and the right panel shows the weight of adult male flies. (B) PIP3 concentrations in flies devoid of dPTEN function rescued by the $dAkt^3$ mutation.
Consistently, pervanadate treatment stimulated dAktG99S activity to 80% of the wild-type level. However, pervanadate-induced activation of the mutant protein occurred more slowly than did that of the wild-type kinase (30). Taken together, these data indicate that the G99S substitution reduces the association of dAkt with the plasma membrane, probably by affecting the affinity of its PH domain for PIP3. Thus, dAkt<sup>G99S</sup> enabled us to study the consequences of impaired recruitment of dAkt to the plasma membrane.

We combined the dAkt alleles with null mutations in <i>dPTEN</i> (32). Animals lacking dPTEN function die during larval stages. A reduction in dAkt expression using the viable dAkt<sup>36</sup> allele did not rescue the lethality associated with <i>dPTEN</i>. Similarly, animals doubly mutant for <i>dPTEN</i> and dAkt<sup>36</sup> did not survive. Thus, either dAkt activation is not the sole reason for the lethality caused by loss of PTEN, or dAkt function is not dispensable in the absence of dPTEN. The latter hypothesis is strongly supported by results obtained with the dAkt allele that selectively impairs the membrane recruitment of dAkt. Flies devoid of functional dPTEN were rescued to viability by any dAkt allelic combination that included dAkt<sup>36</sup> (Fig. 3A) (33). The rescued flies did not display morphological defects that would be expected in light of the phenotypes ascribed to clones of <i>dPTEN</i> mutant cells (17). Tangential sections through compound eyes revealed essentially normal ommatidial and rhabdomeric structures, and the wings of the rescued flies showed no abnormalities in the venation, such as missing crossveins (34). We determined the PIP3/PiP2 ratio by metabolic labeling of phospholipids from larvae (35). PiP3 concentrations were increased in the <i>dPTEN</i> dAkt<sup>36</sup> doubly mutant larvae as compared to those of wild-type larvae (Fig. 3B), excluding the possibility that PiP3 concentrations remain within physiological limits by a feedback regulation mechanism involving dAkt. This suggests that the potential activation of a number of PH domain-containing proteins other than dAkt does not interfere with viability.

Our results indicate that the activation of dAkt is the only crucial outcome of the loss of dPTEN function. Activation of dAkt should therefore mimic the dPTEN loss-of-function phenotype. We expressed a constitutively activated membrane-anchored dAkt during eye development (36). The resulting eyes were increased in size due to enlarged ommatidia (Fig. 4B), a phenotype similar to that seen in clones of <i>dPTEN</i> mutant cells (17–19). This overgrowth phenotype is independent of upstream signals, because it was still evident in a chico or <i>Opal110/PiP3K</i> mutant background (Fig. 4D) (34).

We conclude that flies devoid of the tumor suppressor dPTEN can live with abnormally high concentrations of PiP3 if only the affinity of dAkt for PiP3 is decreased. Thus, the PH domain-mediated translocation of dAkt to the membrane and its subsequent activation is the only lethal event triggered by increased PiP3 concentrations. Because the PH domain of Akt interacts with the substrate of PTEN’s lipid phosphatase activity, we also conclude that PTEN does not exert any essential function other than the dephosphorylation of PiP3.

References and Notes

9. V. Stambolic et al., <i>Cell</i> 95, 29 (1998).

Fig. 4. Growth in the developing <i>Drosophila</i> eye promoted by activated dAkt. (A to D) Scanning electron micrographs of compound eyes of female flies; the anterior is to the left. (A) Wild type. (B) Overexpressing a membrane-tethered version of dAkt (GMR-Gal4 UAS-myr-dAkt). (C) chico mutant. (D) GMR-Gal4 UAS-myr-dAkt in a chico mutant background (45).

18. H. Huang et al., <i>Development</i> 126, 5365 (1999).
27. S. E. Scanga et al., <i>Onconeogene</i> 19, 3971 (2000).
29. The dAkt<sup>36</sup> allele was found in a collection of homozygous viable mutations from Ch. Zuber and K. Koundakjian (University of California, San Diego) because it yields flies of severely reduced body size.
32. The dPTEN alleles used are dPTEN<sup>17/17</sup> (19) and dPTEN<sup>17/17</sup> (37). Both are presumably null alleles. We also tested them over a deficiency lacking the dPTEN locus, Df(2L)7170B. Because this deficiency also lacks chico, an upstream component of insulin receptor signaling, we reintroduced chico by means of a genomic rescue construct. In all cases, we obtained similar results.
33. Whereas the strongest allelic combination dAkt<sup>7</sup>/dAkt<sup>2</sup> completely rescued the dPTEN mutant flies to wild-type size, the combinations dAkt<sup>7</sup>/dAkt<sup>21</sup> could rescue the lethality associated with loss of PTEN function, but the resulting flies were slightly enlarged. Furthermore, we observed a variability of the phenotypes in all combinations. Consistently, flies that emerged earlier showed a tendency to be of increased size, whereas some retarded flies were of smaller size.
34. H. Stocker, E. Hafen, unpublished results.
35. Nonwandering third instar larvae were phosphate-starved in phosphate-free Schneider S2 medium and then labeled with 2 mCi per sample of inorganic <sup>32</sup>P (50 mCi/ml). PiP3 and PiP2 levels were determined according to (38).
36. To anchor dAkt to the plasma membrane, the myristoylation-palmitoylation motif from the Lck tyrosine kinase was fused to the NH<sub>2</sub>-terminus of hemagglutinin (HA) epitope–tagged dAkt, as previously described for mammalian Akt (39). Analysis of the subcellular localization by immunofluorescence using the epitope antibody to HA confirmed the constitutive membrane localization of myr-dAkt.
41. Single-letter abbreviations for the amino acid residues are as follows: A, Ala; E, Glu; F, Phe; G, Gly; I, Ile; P, Pro; R, Arg; S, Ser; and T, Thr.
42. dAkt was immunoprecipitated from wild-type and dAkt3\(^3\) larvae with a rabbit polyclonal antibody raised against recombinant dAkt-66 (40). In vitro kinase assays were performed, as described, using the peptide CRPRTSSAEG (41) as a substrate (39).
43. HEK 293 cells were transfected by a modified calcium phosphate method, as described (39). The transfection mixture was removed after a 16-hour incubation, and cells were serum-starved for 24 hours before stimulation with 0.5 to 1 \(\mu\)M insulin (Boehringer Mannheim).
44. The HA epitope–tagged dAkt proteins were immunoprecipitated from 100 \(\mu\)g of cell-free extracts using the monoclonal antibody 12CA5 coupled to protein A–Sepharose. In vitro kinase assays were performed, as described (39).
45. The size difference of the eyes in Fig. 4, B and D, is entirely due to varying numbers of ommatidia. The size of the ommatidia, however, is comparable (insets) and massively larger than in chico mutants (inset in 4C). The failure of myr-dAkt to compensate for the reduced number of ommatidia in chico mutants is consistent with the late onset of expression driven by GMR-Gal4.
46. We thank Ch. Zuker and E. Koundakjian for the dAkt\(^3\) allele; T. Radimerski for the 5Z labeling medium; T. Gutjahr, Ch. Hugentobler, R. Bopp, P. Zipperlen, P. Cron, P. Müller, and H. Angliker for technical support; K. Basler and P. Gallant for critical reading of the manuscript and valuable suggestions; and D. Pan and S. Leevers for providing fly stocks. Supported by grants from the Schweizerische Krebsliga (B.A.H. and M.P.W.) and the Swiss National Science Foundation (E.H.).