Autism-associated PTEN missense mutation leads to enhanced nuclear localization and neurite outgrowth in an induced pluripotent stem cell line

Chi Wai Wong¹, Yubing Wang¹*, Tian Liu¹, Lisha Li¹, Stanley Kwok Kuen Cheung¹, Penelope Mei-Yu Or¹, Alfred Sze-Lok Cheng¹, Kwong Wai Choy², Johannes Peter Henri Burbach³, Bo Feng¹, Raymond Chuen Chung Chang⁴ and Andrew M. Chan¹,⁵

¹ School of Biomedical Sciences, The Chinese University of Hong Kong, Hong Kong SAR, China
² Department of Obstetrics and Gynaecology, Prince of Wales Hospital, The Chinese University of Hong Kong, Hong Kong SAR, China
³ Department of Translational Neuroscience, Brain Center Rudolf Magnus, University Medical Center Utrecht, The Netherlands
⁴ Laboratory of Neurodegenerative Diseases, School of Biomedical Sciences, Li Ka Shing Faculty of Medicine, The University of Hong Kong, Hong Kong SAR, China
⁵ Brain and Mind Institute, The Chinese University of Hong Kong, Hong Kong SAR, China

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Correspondence
A. M. Chan, School of Biomedical Sciences, Room 705, Lo Kwee-See Integrated Biomedical Sciences Building, The Chinese University of Hong Kong, Hong Kong SAR, China
Tel: +852 3943 3771
E-mail: andrewmchan@cuhk.edu.hk

*Present address
School of Bioscience and Technology, Weifang Medical University, China
Chi Wai Wong and Yubing Wang contributed equally to this work.

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Germline mutation in the PTEN gene is the genetic basis of PTEN hamartoma tumor syndrome with the affected individuals harboring features of autism spectrum disorders. Characterizing a panel of 14 autism-associated PTEN missense mutations revealed reduced protein stability, catalytic activity, and subcellular distribution. Nine out of 14 (64%) PTEN missense mutants had reduced protein expression with most mutations confined to the C2 domain. Selected mutants displayed enhanced polyubiquitination and shortened protein half-life, but that did not appear to involve the polyubiquitination sites at lysine residues at codon 13 or 289. Analyzing their intrinsic lipid phosphatase activities revealed that 78% (11 out of 14) of these mutants had twofold to 10-fold reduction in catalytic activity toward phosphatidylinositol phosphate substrates. Analyzing the subcellular localization of the PTEN missense mutants showed that 64% (nine out of 14) had altered nuclear-to-cytosol ratios with four mutants (G44D, H123Q, E157G, and D326N) showing greater nuclear localization. The E157G mutant was knocked-in to an induced pluripotent stem cell line and recapitulated a similar nuclear targeting preference. Furthermore, iPSCs expressing the E157G mutant were more proliferative at the neural progenitor cell stage but exhibited more extensive dendritic outgrowth. In summary, the combination of biological changes in PTEN is expected to contribute to the behavioral and cellular features of this neurodevelopmental disorder.

Introduction

Phosphatase and tensin homolog deleted on chromosome 10 hamartoma tumor syndrome (PHTS) represents a collection of four overgrowth syndromes including Cowden, Bannayan–Riley–Ruvalcaba, Proteus, and Proteus-like, with germline PTEN mutation detected at variable frequency from 7% to 85% [1].

Abbreviations
ASDs, autism spectrum disorders; PHTS, PTEN hamartoma tumor syndrome; PI3-K, phosphoinositide 3-kinase; PTEN, phosphatase and tensin homolog deleted on chromosome 10.
While having distinct diagnostic criteria, these overgrowth syndromes share common clinical features of multiple hamartomatous growth and increased risk for developing breast and thyroid cancers. In individuals with PTEN germline mutation, macrocephaly is diagnosed in 94% [2], and 22% of PHTS subjects have features of autism spectrum disorders (ASDs) [3]. ASD is a collection of neurodevelopmental disorders including autistic disorder, Asperger syndrome, pervasive developmental disorder not otherwise specified (PDD-NOS), and Rett syndrome [4]. The core symptoms are social and communication deficits, and sensory hypersensitivity. Affected individuals also display repetitive behavior, epileptic episodes, and mental retardation [5]. The early onset (< 3 years) reflects aberrant neurogenesis at a stage when sensory inputs are shaping excitatory and inhibitory synapses. Altered anatomy, function, and connectivity have been demonstrated in ASD brains. Anatomically, ASD is characterized by overgrowth early in life (2–4 years old) in the amygdala and frontal cortex (reviewed in Ref. [6]). These changes coincide with the developmental stage of intense acquisitions of language and social skills.

Autism spectrum disorders has a strong genetic component. Almost 80% of ASD cases are sporadic with the remaining 20% being syndromic [7]. Syndromic ASD patients have autistic phenotypes characterized by the pathologies of some well-established neurodevelopmental syndromes such as fragile X (FMR1), Rett syndrome (MECP2), Angelman syndrome (UBE3A), Phelan–McDermid syndrome (SHANK3), and PHTS (PTEN).

PTEN was originally isolated as a tumor suppressor gene inactivated in 30% of human cancers (reviewed in Ref. [8]). It encodes a lipid phosphatase and dephosphorylates phosphatidylinositol 3,4,5-triphosphate (PIP3), a second messenger that activates the phosphoinositide 3-kinase (PI3-K). Thus, PTEN loss hyperactivates PI3-K and a host of downstream effectors. Structurally, PTEN is composed of a catalytic domain in the amino (N) terminus and a lipid-binding C2 domain in the carboxyl (C) terminus (Fig. 1). This is followed by a flexible ‘tail’ region composed of multiple phosphorylation sites, which ends in a postsynaptic density/Digl/ZO-1-binding motif (PDZ-BM). PTEN is mainly localized to microtubule-associated vesicles in the cytoplasm [9]. It is also localized in the nucleus where it plays a role in genome stability [10,11]. Post-translational modifications involving ubiquitination at Lys13 and Lys289 or sumoylation at Lys254 and Lys266 have been shown to regulate PTEN proteolysis and nuclear translocation, respectively (reviewed in Ref. [12]).

Independently, knockout of Pten gene in differentiated neurons results in viable mice with macrocephaly and abnormal social interaction and learning [13]. Mice that are germline heterozygous for PTEN resemble those with PHTS, harboring macrocephaly, impaired social interactions, and repetitive behaviors [14,15]. In this report, we comprehensively characterized a list of PTEN missense mutants associated with ASD and identified diverse changes in biochemical properties that may perturb the physiological functions of PTEN.

**Results**

**ASD-associated PTEN missense mutants possess reduced protein expression**

To investigate the ASD-associated functional defects in PTEN mutants, a total of 14 missense mutations were identified based on a literature search for PHTS patients with autistic features (Fig. 1). Eight mutants were located in the N-terminal half of PTEN where the catalytic domain resides, while six were in the C2 domain, with none reported in the PEST domain and PDZ-BM. As a control, the PTEN R130Q mutant frequently detected in human endometrial carcinoma was also included [16]. To examine the properties of these PTEN missense mutants, similar amounts of expression plasmids were transfected into three different mammalian cell lines. Specifically, 293T cells expressed wild-type (WT) PTEN, while both U87MG and PC3 were PTEN-null. Western blotting analysis clearly revealed two distinct groups of mutants. The first group was composed of five mutants (H93R, H123Q, I135R, E157G, and Y176C) that had protein expression plasmids were transfected into three different mammalian cell lines. Specifically, 293T cells expressed wild-type (WT) PTEN, while both U87MG and PC3 were PTEN-null. Western blotting analysis clearly revealed two distinct groups of mutants. The first group was composed of five mutants (H93R, H123Q, I135R, E157G, and Y176C) that had protein expression levels very similar to that of WT PTEN (Fig. 2). The second group was composed of nine mutants that had a greater than threefold decrease in the 57-kDa protein species of PTEN (Fig. 2). This drastic reduction in PTEN protein expression was observed in all of the C2 domain mutants, while only three out of eight N-terminal domain mutants (H118P, M134T, and I135R) were affected (Fig. 2A). Interestingly, the catalytic domain mutant, H118P, had the lowest expression among all of the missense mutants. Very similar observations were also detected in 293T and U87MG cells (Fig. 2B,C). In addition, there were subtle differences in the molecular mass with the H93R mutant showing a band of faster mobility, while the I135R mutant had reduced mobility compared with the WT (Fig. 2A–C). These differences are likely to be mutation-specific rather than cell type-specific as similar changes were observed in all three cell lines tested.
Reduced protein half-life and enhanced polyubiquitination of ASD-associated PTEN mutants

Phosphatase and tensin homolog deleted on chromosome 10 is a relatively stable protein with an average half-life of ~9 h [17]. To test whether the reduced protein levels in the PTEN missense mutants were due to reduced protein half-life, protein expression was evaluated following cycloheximide treatment. We selected two low-PTEN expression mutants each from the catalytic domain (H118P, I135R) and C2 domain (D252G, N276S). As a control, we selected the E157G mutant, which has comparable protein expression to the WT. The WT PTEN expressed in 293T cells had an average half-life of ~6 h (Fig. 3). All four mutants, H118P, I135R, D252G, and N276S, with reduced PTEN protein levels had shorter half-lives of 3.0–4.0 h (Fig. 3). As expected, the E157G mutant, which retained 80% PTEN protein expression, had a half-life resembling that of the WT.

It is well established that PTEN protein turnover is regulated by ubiquitin-mediated proteolysis events. To test whether missense mutants with shortened protein half-life had greater ubiquitination, cells expressing these mutants were first treated with a proteasome inhibitor, MG132, for 6 h. The WT PTEN showed almost undetectable polyubiquitination, and ubiquitination only became observable following MG132 treatment (Fig. 4A). All of the missense mutants except E157G had readily detectable basal polyubiquitinated protein species, and polyubiquitination was greatly potentiated by MG132 (Fig. 4A,B). Among the missense mutants tested, I135R had the greatest polyubiquitination, followed by D252G, H1118P, and N276S (Fig. 4B). To elucidate the polyubiquitination sites responsible, two known PTEN ubiquitination sites at K13 and K289 were mutated in the two unstable mutants, I135R and N276S, and in the relatively stable E157G mutant. In addition, because PTEN sumoylation and ubiquitination events have previously been shown to display crosstalk [18], we also eliminated the two known sumoylation sites at K254 and K266 through substitutions. Unexpectedly, none of the ubiquitination- and sumoylation-site mutants could rescue the protein expression levels of either the I135R or N276S mutants (Fig. 5A). In contrast, the expression levels of the more stable E157G mutant could be elevated by mutating the K13 but not the K289 ubiquitination site (Fig. 5B). Furthermore, mutations in the K254 but not the K266 sumoylation site significantly reduced the expression levels of the E157G mutant.
mutant (Fig. 5B). Therefore, individual ASD-associated PTEN missense mutants are differentially affected by specific ubiquitination and sumoylation events, which affect their steady-state protein expression.

Previous studies also demonstrated that phosphorylation at the C-terminal PEST domain could confer protein stability. In particular, the cluster of serine and threonine residues at S362, T366, S379, S380, T382, T383, and S385 has been shown to be phosphorylated by glycogen synthase kinase 3-β, polo-like kinase 3, and casein kinase II (CKII) [19]. We examined the phosphorylation states of the CKII phosphorylation cluster of S380/T382/T383 and the PTEN auto-phosphorylation site at T366 with phospho-specific antibodies. However, no significant differences were observed among all of the PTEN missense mutants (Fig. 6).

### Partial inactivation of lipid phosphatase activity of PTEN missense mutants

For mutants with comparable protein half-lives as WT PTEN, attempts were made to unravel the biological properties that were disrupted. PTEN is a lipid phosphatase with specificity toward PI(3,4,5)P3, and the catalytic pocket of PTEN is located in the N-terminal portion of the protein [20]. To test the relative ability of the PTEN missense mutants in suppressing the PI3-K signaling pathway, similar amounts of expression plasmids were transfected into the PTEN-null cell line U87MG, which harbors appreciable levels of p-Akt. The WT PTEN strongly suppressed p-Akt by 80% (Fig. 7A). While almost all of the N-terminal PTEN missense mutants, except I135R, retained > 80% of the WT’s activity, only half of the C-terminal mutants retained some capacity to suppress Akt phosphorylation. As expected, the catalytic-dead mutant R130Q was completely devoid of suppressive activity (Fig. 7A). These data were, however, confounded by the variable expression levels of individual PTEN mutants expressed in U87MG cells.

To clearly define the intrinsic phosphatase activity of these missense mutants, AU5 epitope-tagged mutants were transfected into 293T cells and similar amounts of PTEN proteins were immunoprecipitated with an anti-AU5 antibody. Lipid phosphatase reaction was initiated by the addition of PI(3,4,5)P3 to PTEN proteins immobilized on sepharose beads followed by incubation for 10 min. The levels of phosphate released were measured by the standard malachite green assay. As shown in Fig. 7B, all of the PTEN missense mutants with the exception of H118P, E157G, Y176C, and T202I had a 60–80% decrease in phosphatase activity compared with the WT. In contrast, the cancer-associated catalytic-dead mutant R130Q was almost completely inactivated. Thus, among the five stable missense mutants, only E157G, and Y176C, retains significant lipid phosphatase activity.

### Altered subcellular distribution of PTEN missense mutants

To explore what further defects could be associated with the E157G and Y176C mutants, we examined their subcellular localization. PTEN has been shown to be localized to various subcellular compartments including plasma membrane, cell–cell junctions, microvesicles in the cytosol, and mitotic spindles of the nucleus [21]. To

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**Fig. 3.** Reduced protein stability of ASD-associated PTEN mutants. 293T cells were transfected with the indicated expression plasmids. The cells were treated with cycloheximide (80 μg·mL⁻¹) for the indicated durations. The cells were solubilized in Laemmli buffer, and the levels of PTEN were determined by western blotting analysis and normalized to the actin levels. The results from three independent experiments were quantified. Bars, SD.
investigate whether the PTEN missense mutants had altered subcellular localization, expression plasmids were transfected into 293T cells and immunofluorescence was analyzed using an anti-AU5 antibody. To minimize the confounding effects of cells in different phases of the cell cycle, the cultures were synchronized in the G1/S phase by double thymidine blocking. The majority of cells in the G1/S phase had flat morphology with the cytosolic and nuclear compartments clearly demarcated. As previously reported, WT PTEN had a more cytosolic rather than nuclear localization (Fig. 8A). Six missense mutants, H118P, M134T, I135R, F241S, D252G, and N276S, had significant reduction in nuclear localization compared with the WT (Fig. 8A,B). Four mutants, G44D, H123Q, E157G, and D326N, displayed greater nuclear localization compared with the WT. They were mostly confined to the nucleoplasm and absent in nucleoli with the exception of the D326N mutant, which also populated the nucleoli (Fig. 8A). Furthermore, in four mutants, H93R, E157G, Y176C, and R130Q, readily detectable membrane extensions resembling pseudopodia or membrane ruffles were observed (Fig. 8A).

To delineate the underlying mechanism responsible for the preferential nuclear distribution of the E157G mutant, we generated additional mutations in this missense mutant. Of note, an E157K mutant that carried an opposing polar group also displayed greater nuclear
Localization, suggesting the importance of this position, rather than the charge, being a critical determinant (unpublished data). Because both ubiquitination and sumoylation have been implicated in PTEN nuclear localization, we tested a panel of E157G mutants carrying mutations in lysine residues that mediated these two post-translational events. As shown in Fig. 8C, D, mutating either of the two sumoylation sites at K254 and K266 alone failed to disrupt the nuclear targeting capacity of the E157G mutant. Unexpectedly, however, mutating both sumoylation sites increased the E157 G nuclear targeting. More importantly, mutating the K13 and K289 ubiquitination sites greatly diminished the nuclear-to-cytosol ratio of the E157G mutants (Fig. 8C, D). We further examined the overall sumoylation state of PTEN missense mutants by a pull-down assay. E157G mutant had a similar sumoylation level compared to WT PTEN protein (Fig. 9). Another mutant, H123Q, which had greater nuclear localization, displayed a slight reduction in sumoylation level. However, out of the 6 mutants with reduced nuclear localization, H118P, M134T, I135R, F241S, D252G, and N276S, all have elevated sumoylation levels (Fig. 9). Thus, there appears to be an inverse relationship between nuclear localization and sumoylation level in certain PTEN mutants. From these analyses, we assigned a tentative defect for the E157G mutant in displaying a preferential localization in the cell nucleus, which may involve ubiquitination events at K13 and K289.

Endogenous E157G mutation displayed greater nuclear localization

To examine how the E157G mutation could impact on neurogenesis, this mutation was knocked-in to a
human iPSC cell line, ASE-9203, using CRISPR/Cas9. Successful homozygous knock-in in both alleles was confirmed by sequencing analysis (data not shown), and the identity was authenticated by short tandem repeat analysis (data not shown). Morphologically, the WT and E157G iPSCs were very similar (Fig. 10A). Next, neural induction was initiated in the iPSCs using neural induction medium and the neural progenitor cells (NPCs) were expanded in neural progenitor medium. Neuronal precursors were generated from the NPCs and matured for 1–2 weeks. As expected, the cells acquired neurite-like extensions following differentiation and maturation processes (Fig. 10A). However, more extensive and robust neurite extensions were observed in E157G cells compared with WT cells. Also, E157G at the NPC stage proliferated at a much greater rate than the WT. Western blotting analysis revealed upregulation of PTEN in both the WT and the E157G, both in NPC and during neuronal differentiation (Fig. 10B). This was correlated with a
corresponding downregulation of p-Akt. However, the levels of WT and E157G PTEN were very similar as we expected based on our results in PC3 cells (Fig. 1B). As expected, high levels of the reprogramming factors NANOGEN, SOX2, and OCT4 were observed in the iPSCs and their expression was downregulated upon neural induction and differentiation (Fig. 10B). However, the levels of SOX2 remained detectable in the E157G cells. More strikingly, the levels of PAX6 were upregulated following neural induction and were silenced following the induction of neuronal differentiation. Interestingly, PAX6 was more downregulated in E157G than in the WT (Fig. 10B,C). For the neuronal-specific marker, TuJ, its expression was higher in E157G than in WT cells during the entire differentiation process. Immunofluorescence analysis also revealed more MAP2-positive staining of microtubule structures in E157G when compared to WT in matured neurons (Fig. 10D). Finally, immunofluorescence analysis of NPC using an anti-PTEN antibody revealed a fourfold increase in nuclear-to-cytosol ratio for the E157G mutant compared with the WT (Fig. 10E,F). Thus, the E157G protein expressed in the iPSC line has properties very similar to those expressed in PC3 cells.

**Discussion**

Spinelli et al. [22] had previously characterized seven PTEN mutants associated with ASD. The present study expanded on this panel by adding an additional seven mutants. Overall, the present study also observed high fraction (> 60%) of PTEN missense mutation would lead to protein instability. Our study also uncovered aberrant subcellular distribution of PTEN missense mutants that was not investigated in previous studies [22,23]. Furthermore, the present study expanded on the previous studies by offering mechanistic insights into specific post-translational modifications to explain the defects associated with individual missense mutations. The overall findings of the present study are summarized in Fig. 11A. Two recent studies had investigated genotype–phenotype relationship of PTEN missense mutants using saturated mutagenesis in yeast cells followed by massively parallel sequencing [24,25]. In terms of catalytic activity, the data by Mighell et al. [24] showed striking consistency with our data with the exception of the H93R and Y176C (Fig. 11B). In terms of protein level, the report by Matreyek et al. [25] also showed high consistency with our data (Fig. 11C). Indeed, these two studies revealed that the E157G mutant has WT-like catalytic activity and protein abundance.

Expression analysis of PTEN mutants in three mammalian cell lines revealed that reduced level of this protein was the most prominent defect of these ASD mutants. This was particularly striking for C2 mutants. Because the same amount of DNA was transfected in each case, it is unlikely that the observed result was due to reduced transcriptional or translational events. Both MG132 treatment and cycloheximide chase experiments indicated that reduced protein stability was the major culprit for the lowered expression levels of ASD-associated PTEN mutants. One possibility is that some of the mutations cause conformational changes to PTEN and hence lead to structural instability. Another observation is that only three out of eight phosphatase domain mutants were unstable, while all six of the C2 domain mutants had drastic reductions in protein expression levels. This suggests that PTEN stability may be more sensitive to C2 domain mutations. Indeed, four out of six C2 domain mutants, D252, W274, N276, and D326, are located along the interphase between the C2 and phosphatase domain, with some partaking in hydrogen bond formation [26]. Mutations in these residues may disrupt these intramolecular interactions and confer structural instability. Our results resemble those reported by Spinelli et al. [22], who also characterized unstable ASD-associated PTEN mutants. Our present study identified two mutants, H118P, and N276S, with the most reduced protein expression. The heightened polyubiquitination in the unstable mutants may explain their shortened half-lives. Unexpectedly, however, the two
major ubiquitination sites at K13 and K289 were not responsible for the half-life shortening, at least for the unstable I135R and N276S mutants. It is possible that cryptic polyubiquitination sites are present. In contrast, the E157G mutant protein is relatively stable and retains ~77% of WT protein expression. This decrease in protein level was rescued by mutating the K13 ubiquitination site suggesting protein degradation mediated by K13 polyubiquitination may be responsible. The reduced expression of E157G in the K254R and K254R/K266R mutants may be related to an increase in polyubiquitination following the elimination of the K254 site. Similar crosstalk has been reported previously [18].

Previous studies conducted in *Saccharomyces cerevisiae* have shown that ASD-associated PTEN mutants suffered partial loss of PTEN activity compared with tumor-associated mutants [23]. In the present study, we performed both *in vivo* and *in vitro* assays to take into account the protein instability of PTEN mutants. Our *in vivo* assay also showed that most ASD-associated PTEN mutants had partial loss of lipid phosphatase activity in U87MG. This is expected as mutations within the N-terminal phosphatase domain are predicted to disrupt catalysis, while mutations in the C2 domain may decrease membrane binding or perturb structural integrity. Also, in most of the mutants with partial loss of *in vivo* phosphatase activity, the loss was not caused by their low protein expression because their *in vitro* lipid phosphatase activities using purified proteins were clearly compromised. Thus, the extent of loss of PTEN activities in PHTS patients reflects a

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**Fig. 10.** Aberrant neuronal differentiation in PTEN E157G iPSc. (A) Morphology of iPSC and different stages of neuronal development. Note the higher neurite outgrowth in E157G. Bars, 60 μm. A similar experiment was repeated once. (B) Western blotting analysis of lineage-specific markers at different stages of neurogenesis. Immunofluorescence (IF) analysis of lineage-specific markers in (C) NPC and (D) maturation stages. Bars, 30 μm. (E) IF of PTEN in NPC. Bars, 15 μm. Similar results were obtained in two additional experiments using iPSC cells. (F) The extent of nuclear localization of PTEN was quantified from a single experiment. Bars, SD. n = 13. ***P < 0.001. Student’s t-test, unpaired. DIFF, differentiation; MAT, maturation; NSC, neural stem cells.
combination of defects in catalytic activity, membrane-binding ability, and protein stability.

The only two mutants with significant lipid phosphatase activity in vivo are E157G and Y176C. One possibility is that these two residues are not located in the core of the phosphatase and C2 domains. In addition, both have comparable expression levels to that of the WT. Our findings are consistent with a previous study reporting the thermostability of PTEN variants in ASD [27]. These results imply that autistic features found in patients harboring E157G and Y176C mutation may be due to defects in PTEN biochemical functions independent of lipid phosphatase activity and expression level. It is uncertain whether ‘less-defective’ PTEN mutants are associated with milder symptoms. A patient with the Y176C mutation had difficulties in social interaction and delayed speech and was diagnosed by the Childhood Autism Rating Scale as severe [28]. The E157G mutation was discovered in a male subject diagnosed with PDD-NOS, and his identical twin was also diagnosed with ASD [29]. These diagnostic features are not remarkably different from those PTEN mutants that have relatively significant decreases in lipid phosphatase activity and protein expression level. Attempts to link specific PTEN defects to disease diagnosis are currently hampered by the rarity of PHTS cases.

The role of nuclear PTEN in ASD is not clear. Disruption of PTEN nuclear translocation has been shown to cause autism in a transgenic mouse model [30]. Mingo et al. [31] had previously reported a Q17E germline PTEN mutant with preferential nuclear localization derived from a PHTS patient with ASD. The present study represents a more comprehensive analysis of the subcellular distribution of ASD-associated PTEN mutants. We showed that the ASD-associated PTEN mutants varied in their abilities in nuclear targeting, but this variation was not obviously associated with their enzymatic activities or protein expression levels. Aberrantly increased or reduced nuclear localization was observed in the E157G and I135R mutants, respectively, which highlights the importance of balanced PTEN levels in cytosolic and nuclear compartments. Mechanistically, ubiquitination events at K13 and K289 are involved in enhancing E157G nuclear localization. Indeed, K13 [32] and K289 [11] were previously suggested to be involved in nuclear translocation of PTEN. Furthermore, we observed an increase in ubiquitination of the E157G mutant (Fig. 4). We are currently testing whether the E157G mutant can interact with PTEN-specific ubiquitin ligase.

The I135R mutant, in contrast, was excluded from the cell nucleus. Thus, it is not surprising that disrupting all of the known ubiquitination and sumoylation sites failed to restore nuclear targeting capacity. As discussed above, the I135R mutant is heavily polyubiquitinated, possibly on cryptic sites. This may disrupt ubiquitination and sumoylation events that are normally required for nuclear targeting.

Our iPSC data imply that the E157G mutation promotes proliferation of NPC but confers greater neuronal differentiation. This is consistent with the long-held notion that loss of PTEN function increases

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**Fig. 11.** Biological properties of PTEN missense mutants. (A) Results obtained in this study are summarized by using indicated color codes. The relative levels of biological readout for individual mutants were calculated based on the mean values from the respective assays and compared with WT protein. (B) Fitness scores extracted from the study by Mighell et al. [24] for the indicated mutants are shown. (C) Protein-level classification based on data from the study by Matreyek et al. [25] for the indicated mutant is shown.
dendritic arborization and hypertrophic growth. Indeed, hyperactivation of the mTOR pathway is directly linked to increased dendritic spine density in layer V pyramidal neurons in postmortem ASD temporal lobes [33]. The higher SOX2 expression in E157G NPC may explain the higher rate of proliferation. The persistence of SOX2 expression in E157G may also suggest that a subpopulation of neural stem cells persists following differentiation. It is currently unclear whether the enhanced neurite outgrowth seen in E157G is due to reduced cytosolic or enhanced nuclear PTEN functions. Whether the E157G mutant can directly regulate gene expression in the nucleus is not clear. Future studies to resolve these issues may yield new insights into the molecular pathogenesis of ASD.

**Materials and methods**

**Chemicals**

All chemicals used in this study were purchased from Sigma-Aldrich (St. Louis, MO, USA) unless otherwise indicated.

**Antibody reagents**

Antibodies against PTEN (#9552, #9559), p-Akt-S473 (#3271), c-Akt (#9272), PTEN-pS380/pT382/pT383 (#9554), Nanog (#4903), Pax6 (#60433), Oct4 (#2750), Sox2 (#3579), GFAP (#3670), GAPDH (#5174), and horseradish peroxidase (HRP)-conjugated anti-rabbit or anti-mouse IgG (#7074, #7076) were from Cell Signaling (Danvers, MA, USA). The anti-AU5 antibody (#ab130113) and anti-Vimentin (#ab8978) were from Abcam (Cambridge, UK), and anti-actin antibody (#I-19) was from Santa Cruz Inc. (Dallas, TX, USA). An antibody directed against PTEN-pT366 was a generous gift provided by Nick Leslie (Heriot-Watt University, UK) [34].

**Compilation of PTEN missense mutants**

A PubMed search was conducted using the search terms—PTEN, autism, and mutation. A list of germ-line PTEN missense mutations found in PHTS patients with ASD was compiled. The source references used were as follows: G44D [29], H93R [3], H118P [28], H123Q [35], M134T [35], I135R [36], E157G [29], Y176C [28], T202I [29], F241S [3], D252G [3], W274L [35], N276S [28], and D326N [37].

**Expression plasmids**

All PTEN missense mutants were generated by PCR-based site-directed mutagenesis techniques. All cDNAs generated were cloned into pEF-AU5 mammalian expression vector and their authenticity confirmed by sequencing analysis as described previously [38].

**Cell cultures**

The HEK293T, PC3, and U87MG cell lines were from the cell bank at the National Cancer Institute, USA. Cultures were maintained in DMEM supplemented with 10% FBS and penicillin–streptomycin (pen–strept; Thermo Fisher Scientific, Waltham, MA, USA). For lipofection, ~1 × 10^7 cells were plated in a well of a six-well culture dish. Samples of 0.1–3.0 µg of DNA constructs were mixed with 2 µL of Lipofectamine 2000 in 0.2 mL of Opti-MEM (Thermo Fisher Scientific). This mixture was added to 2 mL of DMEM + 5% FBS medium without pen–strept and incubated for 16 h. The cells were analyzed 1–2 days later.

**Lipid phosphatase assay**

An immunoprecipitation-based lipid phosphatase assay was conducted as previously described [39]. Around 3 × 10^5 293T cells in 100-mm plates were transfected by lipofection using 10 µg of various PTEN expression constructs. After 36 h, the cells were washed with cold TBS (25 mM Tris/HCl pH 8.0, 150 mM NaCl) and resuspended in 550 µL of lysis buffer (25 mM Tris/HCl pH 8.0, 150 mM NaCl, 1% Triton X-100, 2 mM DTT, and a protease inhibitor cocktail). PTEN was immunoprecipitated with 1 µg of an anti-AU5 epitope antibody for 3 h at 4 °C followed by the addition of 35 µL of GammaBind G sepharose beads (Millipore, Billerica, MA, USA) for 1 h. Phosphatase reactions were initiated with the addition of 30 µL of 25 mM Tris/HCl pH 8.0, 10 mM DTT, 50 µM PIP3 (Echelon, Salt Lake City, UT, USA). After incubating at 37 °C for 10 min, 100 µL of malachite green (Echelon) was added to each reaction. The absorbance at OD 620 nm was measured in a SpectraMax i3X microplate reader (Molecular Devices, San Jose, CA, USA). The enzymatic activity of PTEN was presented as pmol pyrophosphate (PPi) released ng⁻¹.min⁻¹.

**Western blotting analysis**

Cells were solubilized in 100 µL of either 2 × Laemmli buffer or RIPA buffer composed of 50 mM Tris/HCl pH 7.5, 1% Triton X-100, 0.5% sodium deoxycholate, 0.1% SDS, 150 mM NaCl, 10 mM MgCl₂, and protease and phosphatase inhibitors. Around 5–20 µg of total cell lysates were resolved on 10–15% SDS/PAGE gels,
transferred onto nitrocellulose membranes, and probed with selected primary antibodies. Bound antibodies were detected with either HRP-conjugated anti-mouse or anti-rabbit secondary antibodies. The expression levels of protein were visualized by the standard ECL method (Life Technologies, Carlsbad, CA, USA) followed by exposure to X-ray films. Band intensity was quantified by the GS-900 Calibrated Densitometer System (Bio-Rad, Hercules, CA, USA).

**Immunofluorescence analysis**

Around $1 \times 10^4$ cells were seeded on a glass coverslip coated with 0.5% Matrigel®Matrix (CORNING #356231, Corning, NY, USA) in a 12-well plate. The cells were fixed with 4% paraformaldehyde for 15 min and permeabilized with 0.2% of Triton X-100 in PBS for 12 min at room temperature. The cells were then incubated with antibodies (1 : 100) at 4°C overnight. They were washed with PBS and incubated with Alexa Fluor® 488-conjugated donkey anti-mouse IgG (H + L) antibody (1 : 200; Thermo Fisher Scientific) for 1 h. They were mounted with 300 nM DAPI on glass slides. They were then imaged with a confocal microscope (Olympus FV1200, Tokyo, Japan) using the METAMORPH imaging software (San Jose, CA, USA), and the fluorescence intensity was quantified by the IMAGEJ software (NIH, MD, USA). The nuclear-to-cytosol ratios were calculated from 15 to 20 cells using the mid-sections of the z-stacks of each cell.

**SUMO pull-down assay**

HEK293T cells were plated at a density of $1 \times 10^6$ on a 100-mm dish and transfected with 7 μg of PTEN expressing plasmids, 5 μg of His$_6$-SUMO2 plasmid, 3 μg of UBC9 plasmid, and 12 μL of Lipofectamine 2000 (Invitrogen, Carlsbad, CA, USA) for 16 h. After 48 h, cells were lysed by denaturing cell lysis buffer (6 M guanidinium-HCl, 10 mM Tris, 100 mM sodium phosphate buffer pH 8.0, and 5 mM β-mercaptoethanol) and imidazole to 5 mM and then sonicated briefly. After centrifugation at 3000 g for 15 min, the supernatant was incubated with 50 μL prewashed Ni$^{2+}$ NTA sepharose for 4 h at 4°C. Beads were washed once with 5 mL cell lysis buffer supplemented with 0.1% Triton X-100, once with 4 mL of pH 8.0 wash buffer (8 M Urea, 10 mM Tris, 100 mM sodium phosphate buffer pH 8.0, 5 mM β-mercaptoethanol, 0.1% Triton X-100), and three times with 4 mL of pH 6.3 wash buffer (8 M Urea, 10 mM Tris, 100 mM sodium phosphate buffer pH 6.3, 5 mM β-mercaptoethanol, 0.1% Triton X-100). SUMO-conjugated proteins were eluted with an elution buffer (200 mM imidazole, 5% SDS, 150 mM Tris/HCl pH 6.7, 30% glycerol, 720 mM β-mercaptoethanol, 0.0025% bromophenol blue).

**Knock-in K157G mutation in iPSC**

A knocked-in PTEN K157G iPSC line was generated using an Applied StemCell kit (Milpitas, CA, USA). Briefly, ASE-9203 human iPSC cells were electroporated with the following gRNA: 5’-TAGATTCTATGGG-GAACTAAGG-3’ in the cloning vector, pBT-U6-Cas9-2A-GFP. The CRISPR-Cas9 positive clone was confirmed by Sanger sequencing analysis. The iPSC cells were cultured and differentiated according to the manufacturer’s instructions (STEMCELL Technologies, Vancouver, BC, Canada).

**Statistical analysis**

All values are expressed as means ± SD. Statistical significance was analyzed using a two-tailed Student’s t-test for two-group comparisons. For more than two groups, analysis of variance (ANOVA) with Dunnett’s post-tests was used. Statistical significance was considered to be $P < 0.05$.

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**Conflict of interest**

The authors declare no conflict of interest.

**Author contributions**

CWW, YW, TL, LL, KKC, and PMO conducted the experiments; CWW, YW, and AMC planned, conducted the experiment, and wrote the manuscript; ASC, KWC, JPHB, BF, RCCC, and AMC provided expertise in planning key experiments reported in this study.
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