A new human adipocyte model with PTEN haploinsufficiency

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ABSTRACT

Few human cell strains are suitable and readily available as in vitro adipocyte models. We used resected lipoma tissue from a patient with germline phosphatase and tensin homolog (PTEN) haploinsufficiency to establish a preadipocyte cell strain termed LipPD1 and aimed to characterize cellular functions and signalling pathway alterations in comparison to the established adipocyte model Simpson-Golabi-Behmel-Syndrome (SGBS) and to primary stromal-vascular fraction cells. We found that both cellular life span and the capacity for adipocyte differentiation as well as adipocyte-specific functions were preserved in LipPD1 and comparable to SGBS adipocytes. Basal and growth factor-stimulated activation of the PI3 K/AKT signalling pathway was increased in LipPD1 preadipocytes, corresponding to reduced PTEN levels in comparison to SGBS cells. Altogether, LipPD1 cells are a novel primary cell model with a defined genetic lesion suitable for the study of adipocyte biology.

Introduction

Obesity and concomitant diseases are a major public health issue [1]. In vitro models are needed to elucidate molecular mechanisms of adipose tissue dysfunction and identify novel drug targets for obesity treatment. Adipose precursor cells derived from the stromal-vascular fraction (SVF) of adipose tissues from various species including humans have been used for in vitro studies of adipocyte biology. These cell systems are more physiological than transformed murine preadipocyte models, such as 3T3-L1, but availability of tissue and supply of cells are limited. Moreover, there is high donor-to-donor variability of adipose tissue characteristics and function [2]. Only few human preadipocyte cell strains are available for the study of adipose tissue biology [3–5], among which the Simpson-Golabi-Behmel syndrome (SGBS) cell strain is the most widely used. SGBS cells are characterized by a retained capacity for adipogenic differentiation in long-term cell culture [5]. We used resected lipoma tissue from a patient with PTEN hamartoma tumour syndrome (PHTS) carrying a heterozygous germine microdeletion in the phosphatase and tensin homolog (PTEN) gene to establish a cell strain termed LipPD1 [6]. PTEN catalyses the conversion of the lipid messenger phosphatidylinositol-3,4,5-trisphosphate (PIP3) to phosphatidylinositol-4,5-bisphosphate (PIP2), counteracting the activity of the class I phosphatidylinositol 3-kinases (PI3 Ks) and is involved in the regulation of adipocyte function including insulin sensitivity [7]. PTEN inhibition was shown to increase glucose utilization by up-regulating intracellular PIP3 concentrations and activation of AKT in 3T3-L1 adipocytes [8] as well as to increase insulin sensitivity in 3T3-L1 preadipocytes [9]. Patients with PHTS were shown to be more insulin sensitive despite being more obese compared to matched controls [10]. We aimed to characterize adipocellular function and particularly responses to insulin of the newly established LipPD1 cell strain in comparison to PTEN wild type human primary SVF cells and the established adipocyte model SGBS.

Materials and methods

Ethics statement

This study was carried out in accordance with the recommendations of the ethics committee of the University of Leipzig. The protocol was approved by...
the Ethics Committee of the University of Leipzig (-425-12-1712201 and 2425-12-171220). All patients or patient’s parents gave written informed consent in accordance with the Declaration of Helsinki 1975.

**Histopathology and cellular composition of lipoma tissue**

Lipoma and adjacent adipose tissue (subcutaneous, peri- and prerenal adipose tissue) of an 11-month-old male PHTS patient were resected for diagnostic and therapeutic purposes. Parts of the resected lipoma were embedded in paraffin and stained with hematoxylin/eosin and a periodic acid-Schiff stain for tumour diagnosis. Furthermore, S-100-protein was detected immunohistochemically according to routine protocols to exclude malignancy [11].

Staining for PTEN was performed after demasking the tissues in Target Retrieval solution pH 9.0 (Agilent, #S2367) using a rabbit anti-PTEN antibody (Abcam, #ab32199) and the LSAB+ Dako Real Detection Systems, AP/RED, Rabbit/Mouse detection kit (Agilent, #K5005) according to manufacturer’s instructions. The anti-PTEN antibody was used at a dilution of 1:50 and incubated for 1 h at room temperature.

**Cell culture**

SGBS cells were established and cultured as described previously [5]. In our culture studies, we used SGBS cells starting from 35 population doublings which corresponds to approximately 70 d in culture. LipPD1 cells were obtained from lipomatous tissue of an 11 month-old male PHTS patient. SVF#6 (male donor, 27 y) and SVF#10 (female donor, 29 y) cells were established from visceral adipose tissue resected during bariatric surgery of obese, non-diabetic adult patients. SVF cells were isolated from the patient’s lipoma (LipPD1) or from visceral adipose tissue of healthy donors (SVF#6 and SVF#10) and transferred into culture as described previously [6]. For the isolation of preadipocytes, the tissue (~5–20 g) was cut into pieces of about 0.5 cm³ and washed twice with sterile PBS (Sigma, #D8537). After digestion in 200 U/ml Collagenase-HBSS-solution (Sigma, #C5138) for 90 minutes in a 37°C shaking water bath, the material was filtrated through a nylon mesh (pore size 400 µm) and centrifuged at 1,100 rpm at room temperature for 5 min. The pellet was washed and centrifuged again at 1,500 rpm. Erythrocytes were eliminated by lysis (150 mM ammonium chloride (Sigma, #A9434), 10 mM potassium hydrogen carbonate (Roth, #3904.1), 0.1 mM EDTA (Roth, #1410.2), pH 7.29, room temperature, 10 min) and subsequent centrifugation at 1,500 rpm. The pellet was resuspended in culture medium (DMEM/F12 (Thermo Fisher Scientific, #11330032), supplemented with 10% FCS (Merck, #S0615), 33 mM biotin (Sigma, #B4639) and 17 µM pantothenic acid (Sigma, #P5155) and filtered once more (pore size 30 µm). Cells were seeded as adherent cultures and subcultured continuously at a ratio of 1:4 before reaching confluence, usually twice a week. For estimating the capacity for adipocyte differentiation during long-term culture, adipocyte differentiation was induced every three passages.

For 3D culture, 10,000 cells per well were seeded in 96-well ultra-low attachment plates (Greiner Bio-One, #650970) according to a modified protocol from Klingelhutz et al. [12] as previously described [13]. Microscope images were taken daily using an EVOS FL Auto 2 Cell Imaging System (Thermo Fisher Scientific, #AMAFD2000). Medium change was performed by aspirating and replacing half of the medium every other day.

**Adipocyte differentiation**

Lipoma cells and controls were differentiated into mature adipocytes following standard protocols as described previously [5]. Before inducing adipocyte differentiation, LipPD1 and SVF were passaged for approximately 40–60 d, while SGBS were passaged for approximately 70–90 d. Cells were seeded at high density (~10,000 cells/cm²) on 6-well plates and reached full confluence after one or 2 d. Differentiation was then induced by incubation with differentiation medium (DMEM/F12, Thermo Fisher Scientific, #11330032), 33 mM biotin (Sigma, #B4639) 17 µM pantothenic acid (Sigma, #P5155), 0.13 mM apo-transferrin (Sigma, #T1147), 20 nM insulin (Sigma, #I2643), 100 nM hydrocortisone (Sigma, #H0888) and 0.2 mM triiodothyronine (Sigma, #T6397). For the first 4 d, cells were incubated in QD medium differentiation medium supplemented with 25 nM dexamethasone (Sigma, #D1756), 500 µM 1-methyl-3-isobutyl-1-methylxanthine (Sigma, #I5879) and 2 µM rosiglitazone (Sigma, #R2408). Medium was changed every other day until day 12. To confirm and quantify differentiation efficiency, cells were fixed with Roti-HistoFix 4% PFA (Roth, #P087.4), double-stained with Nile Red (Sigma, #N3013) and Hoechst 33342 (Sigma, #14533) and analysed by fluorescence microscopy. The percentage of lipid accumulating cells in proportion to total cell number was determined by enumerating five defined, randomly chosen areas per well using fluorescence microscopy. Experiments were done three times in quadruplicates. For comparison of
LipPD1, SVF#6, SVF#10 and SGBS cells were plated at ~13,000 cells/cm² on 6-well plates or ~45,000 cells/cm² on 96-well plates and differentiation was induced the next day with QD medium. Cells were kept in QD medium for 8 d and were either frozen for RNA extraction or fixed for lipid staining. Undifferentiated controls were kept in serum-free basal medium for 24 h prior to freezing for RNA extraction. Fixed cells were first double-stained with Nile Red and Hoechst 33342 followed by a staining with Oil Red O solution (0.03% in 60% isopropanol, Sigma, #O0625) for 15 min at 37°C.

Proliferation of lipoma cells

Cells were seeded at low density (~1,000 cells/cm²) in multiwell-plates with serum containing medium. Cells were fixed, nuclei were stained with Hoechst-33342 at several time points (24–96 h) and counted using fluorescence microscopy in defined areas in random order. In three independent experiments population doubling (PD) time was estimated in the exponential growth phase using the formula PD = 3.32 (log Xe – log Xb) + S (Xb: cell number at the beginning of the incubation time, Xe: cell number at the end of the incubation time, S: starting PD level, animal cell culture guide; www.atcc.org)

Quantification of mRNA expression

Adipocyte and other cell-type markers from different adipose tissue depots, lipoma tissue, 3D- and 2D cell cultures were quantified via quantitative PCR (qPCR). For RNA extraction from tissue or 3D cultures we used TRIzol Reagent (Thermo Fisher Scientific, #15596026) and homogenized the material using a TissueLyser II (QIAGEN). RNA from 2D cell cultures was extracted using the RNeasy Mini Kit (QIAGEN, #74106). Reverse transcription was done according to standard protocols using the M-MLV Reverse Transcriptase (Thermo Fisher Scientific, #28025013). Following the producer’s instruction and protocols, gene expression was quantified using TaqMan-Technology (Thermo Fisher Scientific). Primers and probes were designed with the software PrimerExpress (Version 3.0, Thermo Fisher Scientific). All measurements were normalized on housekeeping genes hypoxanthine phosphoribosyltransferase (HPRT) and Tata-box binding protein (TBP). Analysed genes and primer sequences are listed in Supplementary Table 1.

Insulin/IGF-I signalling pathway

The amount of PTEN protein and phosphorylated AKT was detected as described previously [6]. For the detection of AKT phosphorylation after overnight serum starvation, cells were stimulated with increasing doses of IGF-I (GroPep, #IU100) or insulin (Sigma, #I2643). After lysis, electrophoretic separation of proteins, and transfer onto nitrocellulose membranes, immunoblotting was performed to detect the amount of PTEN (PTEN (138G6) Rabbit mAb, #9559), phosphorylated (Phospho-Akt (Thr308) (224F9) Rabbit mAb, #4056; Phospho-Akt (Ser473) (193H12) Rabbit mAb, #4058) and total AKT (Akt Antibody #9272). All antibodies were purchased from Cell Signalling Technologies and used according to standard protocols. GAPDH (Merck, #MAB374) or β-Actin (Sigma, A5316) was used as loading control. For semi-quantification, densitometric analysis was carried out using the open-source software ImageJ (Version 1.41 [14]),

Lipolysis

Adipocytes (LipPD1 and SGBS, differentiated until day 13) were incubated in basal medium (DMEM/F12 (Thermo Fisher Scientific, #11330032), 33 μM biotin (Sigma, #B4639) 17 μM pantothenic acid) for 5 h and subsequently stimulated with different concentrations of isoproterenol (Sigma, #I6504, 10 pmol/l – 100 pmol/l) for 22 h. The release of glycerol as measure for an induced lipolysis was quantified using a glycerol assay kit (Sigma, #MAK117-1KT) and normalized to the amount of protein.

¹⁴C-2-deoxy-glucose uptake

For glucose uptake assays, fully differentiated LipPD1 and SGBS adipocytes (differentiated until day 12) were incubated as described before [15]. Cells were incubated in culture medium overnight, washed and incubated in 20 mM Hepes Krebs Ringer/0.1% BSA buffer, pH 7.4 (136 mmol/l NaCl, 4.7 mmol/l KCl, 1.25 mmol/l MgSO₄, 1.25 mmol/l CaCl₂) for 2 h at 37°C. Afterwards, cells were stimulated with 1–10⁶ pM insulin for 2 h at 37°C. Glucose uptake was determined by adding [¹⁴ C]-2-deoxyglucose (#NEC720A050UC, Perkin Elmer) to a final concentration of 0.5 μCi/ml for 5 min. The reaction was stopped by adding 10 mmol/l 2-deoxyglucose and subsequent washing in ice-cold PBS, and cell lysates were prepared in 0.1% SDS (ROTH, #14102.2). Radioactivity was determined after adding scintillation fluid (OptiPhase-SuperMix, #1200-439, Perkin Elmer) using a WALLACE 1409 DSA Liquid Scintillation counter. Analyses were performed in quadruplicates.
**PI3 K activity measurement**

To quantify the amount of PIP₃ at the plasma membrane as a measure for PI3 K activity, LipPD1 or SGBS cells were transfected with a plasmid encoding the marker protein for PIP₃, YFP-GRP1(PH) [16] using the Amaxa nucleofection technique (Cell Line Nucleofector Kit V, Lonza, #VVCA-1003, program T-30). Nucleofected cells were seeded onto glass cover slips in normal culture medium and medium was replaced after 24 h with serum-free medium to reduce basal PI3-kinase activity. Total internal reflection/fluorescence redistribution after photobleaching (TIR/FRAP) measurements were performed after 3–6 h incubation in serum-free medium. Wortmannin (400 nmol/l, LC Laboratories, #W-2990) in serum-free medium for 3–6 h was used as negative control. Repetitive TIR/FRAP measurements at one spot of the cell membrane were performed as described previously using a setup that enabled image acquisition under TIR illumination in combination with spot photobleaching [16].

**Statistics**

Unless otherwise stated, experiments were performed ≥3 times. GraphPad®Prism 6 was used for data visualization and statistical calculations. Data were presented as mean±SEM. Statistical significance was tested by one-way or two-way analysis of variance (ANOVA) followed by Sidak’s multiple comparisons test for comparing three or more groups or Student’s *t*-test for comparing two groups.

**Results**

**Lipoma tissue is not different from white adipose tissue**

Lipoma and adjacent adipose tissues from a patient with PTEN hamartoma tumour syndrome (PHTS) and a heterozygous germline microdeletion of exons 2–9 of the PTEN gene were resected for diagnostic purposes and to reduce tumour burden [6]. Histological analysis was performed to exclude malignancy and showed about 80% normal white adipocytes with insular spots of multilocular fat containing cells. PTEN protein was similarly detected in lipoma tissue (Figure 1(b)) compared to subcutaneous (Figure 1(a)) and perirenal tissue (Figure 1(c)). No mitotic activities, nuclear polymorphisms or myxoid changes were observed. The investigation was repeated twice to confirm the somewhat unexpected result showing normal adipose tissue histology in the presence of severe lipomatosis. A comparative genomic hybridization (CGH) array was performed to detect the boundaries of the heterozygous PTEN deletion. Exons 2–9 of PTEN and additional 53,000 bp downstream were found to be deleted (schematic depiction in Supplementary Figure 1E). No other genes than PTEN were affected by the deletion [6].

In line with the histological results, PTEN mRNA expression was similar in all examined adipose tissue depots of the patient (Figure 1(d)). To further characterize the lipoma tissue, we compared the expression of white adipocyte marker genes peroxisome proliferator-activated receptor γ (PPARγ, Figure 1(e)), adipocyte binding protein 2 (aP2, Figure 1(f)), fatty acid synthase (FASN, Figure 1(g)), adiponectin (Figure 1(h)) and brown adipocyte marker uncoupling protein 1 (UCP-1, Figure 1(i)). There was a trend towards higher expression of PPARγ, aP2 and higher expression of UCP 1 in the peri- and prerenal adipose depots compared to lipoma or subcutaneous adipose tissue of the patient. No significant differences in gene expression were detected between lipoma tissue and subcutaneous adipose tissue of the patient. This was also seen for the brown adipocyte marker PR Domain 16 (PRDM16), the endothelial cell marker cadherin 5 (CDH5), the macrophage marker cluster of differentiation 163 (CD163) and the fibroblast marker cartilage oligomeric matrix protein (COMP) (Supplementary Figure 1A-D).

**LipPD1 show enhanced capacity for adipose differentiation during long-term culture**

Lipoma tissue was used to establish a cell strain termed LipPD1 [6]. Doubling time was determined and was found to be approximately 25 h (Supplementary Figure 2A) until cells were approximately 70 d in culture. From this time point onwards, cells grew only very slowly and stopped dividing at approximately 90 d in culture (Supplementary Figure 2B).

Lipoma cells phenotypically resembled preadipocytes. Interestingly, they could be frozen and re-thawed and showed a high potential for adipose differentiation *in vitro* similar to the SGBS adipocyte model. This capacity was preserved under long-term culture conditions (Figure 2(a)) [5]. In line with this, LipPD1 cells passed for 40–60 d showed an amount of lipid accumulation in response to *in vitro* adipocyte differentiation comparable to SGBS cells. In contrast, control primary stromal-vascular fraction (SVF) cells incubated for the same time period completely lost their capacity to accumulate lipids (Figure 2(b)).

To determine whether the characteristic changes in gene expression patterns during adipocyte differentiation were retained during long-term culture both in SGBS and
LipPD1 cells, mRNA expression of PPARγ, aP2, FASN, leptin and adiponectin (Figure 2(c–g)) was quantified over 8 d of differentiation and compared with SVF cells. Expression of white adipocyte differentiation markers was significantly higher in SGBS after 8 d of differentiation compared to undifferentiated SGBS (p < 0.05). Except for the expression of PPARγ and leptin, this was the case also in LipPD1 (p < 0.05). SGBS also showed a higher expression of PPARγ and leptin than LipPD1 (p < 0.05). These results were supported when comparing gene expression time courses until day 12 of adipocyte differentiation (Supplementary Figure 2B-E). In accordance with the inability to accumulate lipids, long-term cultured SVFs did not show differences of adipocyte marker expression before and after differentiation (Figure 2(c–f)).

Figure 1. Characterization of lipoma tissue from a patient with phosphatase and tensin homolog (PTEN) hamartoma tumour syndrome.

Representative images of PTEN detection by immune histochemistry of (a) subcutaneous adipose tissue, (b) lipoma tissue and (c) perirenal adipose tissue of the patient with PHTS; bar = 500 µm; gene expression of (d) PTEN, and white adipose tissue markers (e) peroxisome proliferator-activated receptor γ (PPARγ), (f) adipocyte binding protein 2 (aP2), (g) fatty acid synthase (FASN), (h) adiponectin and (i) the brown adipocyte marker uncoupling protein 1 (UCP 1) was quantified in different adipose tissue depots. Data were normalized to TATA-box binding protein (TBP) and hypoxanthine phosphoribosyltransferase (HPRT) expression and presented as mean±SEM. n = 4 for PTEN, n = 3 for other genes. subc: subcutaneous, peri: perirenal, pre: prerenal
Figure 2. Growth and adipocyte differentiation of LipPD1 cells during long-term culture.

(a) Quantification of lipid accumulation in LipPD1 cells cultured for 80 population doublings and in vitro differentiated into adipocytes every three passages. Nile Red stained cells were counted and normalized to cell number as determined by DNA staining. Data are presented as mean±SEM, n = 3 independent experiments in quadruplicates. *Data for SGBS and controls 1–3 were taken from [5].

(b) Quantification of lipid accumulation in SGBS (generation/population doublings 42.5 and 43.5, cultured for approximately 40 d) and stromal-vascular fraction cells from two different healthy donors (SVF#6 generation/population doublings 30 and 31, cultured for approximately 60 d) and SVF#10, generation/population doublings 27.5 and 28.5, cultured for approximately 55 d) in vitro differentiated into adipocytes until day 8. Nile Red was normalized to Hoechst-33324 fluorescence intensity. Representative images of differentiated cells stained with Oil Red O. bar = 500 µm; Data are presented as mean±SEM, n = 2 independent experiments in 9-tuplicates. *p < 0.05 determined by one-way ANOVA, 0 F: undifferentiated LipPD1 cells expression of adipocyte marker genes (c) peroxisome proliferator-activated receptor γ (PPARγ), (d) fatty acid synthase (FASN), (e) adipocyte binding protein 2 (aP2), (f) leptin and (g) adiponectin, in LipPD1 (cultured 40–60 d) compared to SGBS (cultured 80–90 d) and stromal-vascular fraction cells from two healthy donors (SVF#6, cultured 40–50 d and #10, cultured 30–50 d), on day 0 and 8 of adipocyte differentiation. Data were normalized to TATA-box binding protein (TBP) and presented as mean±SEM, n = 3 independent experiments in triplicates.
**LipPD1 and SGBS adipocytes are functionally comparable**

We next tested whether typical adipocyte functional properties were similar in LipPD1 and SGBS adipocytes (Figure 3(a–c)). Lipolytic capacity was comparable between LipPD1 and SGBS adipocytes with a dose-dependent increase in lipolysis after isoproterenol stimulation both for LipPD1 and SGBS adipocytes with a maximum at 10 μmol/l isoproterenol (Figure 3(a)). Insulin stimulation led to a significant dose-dependent increase in 14C-2-deoxy-glucose uptake in both LipPD1 and SGBS adipocytes (p < 0.05) with no significant differences between LipPD1 and SGBS (Figure 3(b)). Both cell

![Figure 3](image-url)

**Figure 3.** Functional characteristics of LipPD1 compared to SGBS adipocytes. (a) Isoproterenol-induced lipolysis and (b) insulin-stimulated 14C-deoxyglucose uptake is comparable between LipPD1 and SGBS adipocytes. (c) LipPD1 and SGBS cells were cultured as spheroids in 3D differentiation culture for 9 d, photos were taken every 3 d. Representative images (100x) are shown of LipPD1 and SGBS spheroids differentiated to day 9, fixed in 4% paraformaldehyde and stained with Nile Red (lipids) and Hoechst-33324 (DNA). Expression of adipose differentiation marker genes in 3D in vitro differentiation: (d) peroxisome proliferator-activated receptor γ (PPARγ) and (E) adiponectin. TATA-box binding protein (Tbp) was used for normalization. Data were normalized to the mean of SGBS d4 differentiated cells. Data are presented as mean±SEM, n = 3 independent experiments  *p < 0.05 determined by two-way ANOVA.
strains grew well in 3D culture; spheroids increased in diameter during in vitro differentiation indicating lipid accumulation [12]. On day 6 and 9, SGBS spheroids were larger than LipPD1, reflecting enhanced adipocyte differentiation and therefore adipocyte size (Figure 3(c)). Expression of differentiation markers PPARγ (Figure 3(d)) and adiponectin (Figure 3(e)) was significantly higher in SGBS than in LipPD1 spheroids (p < 0.05).

**PI3 K and AKT are constitutively activated in LipPD1 compared to SGBS preadipocytes**

To check whether the detected mutation in PTEN influenced cellular signalling, we analysed the PI3 K/ AKT signalling pathway. LipPD1 had lower PTEN protein compared to SGBS (Figure 4(a), 0.50 ± 0.11Iold in LipPD1, p < 0.05 in preadipocytes). Next, we tested whether PI3 K/AKT signalling was different in LipPD1 compared to SGBS preadipocytes. A higher basal and IGF-1-stimulated AKT phosphorylation both at Thr308 (p < 0.05; Figure 4(b), Supplementary Figure 3A) and at Ser473 (Supplementary Figure 3B [17]) was detected in LipPD1 compared to SGBS preadipocytes. The amount of a fluorescent PIP3 analogue at the cell membrane was quantified under serum free conditions or stimulated with IGF-1 (10 nmol/l). LipPD1 had a higher basal amount of membrane-bound PIP3 analogue than SGBS preadipocytes (p < 0.05, Figure 4(c)). PIP3 analogue measurements after stimulation with IGF-1 were not different in LipPD1 and SGBS cells (Figure 4(c)). In line with the similar functional responses to insulin stimulation, no differences in basal and insulin-stimulated AKT phosphorylation were found when comparing LipPD1 and SGBS adipocytes (Figure 4(d), Supplementary figure 4A, B).

Other phosphatases acting upon PIP3 or AKT could potentially be upregulated and substitute for PTEN haploinsufficiency. We checked mRNA expression of candidate phosphatases PH domain leucine-rich repeat protein phosphatases (PHLPP) 1 and 2, protein-tyrosine phosphatase (PTP)-1B or PTP receptor type F (PTP-RF). While expression of PHLPP 1 and 2 was not detectable in our cell models (data not shown), we found higher expression of PTP-RF in LipPD1 cells during adipocyte differentiation compared to SGBS cells (Figure 4(e)). PTP-1B expression was not different (Figure 4(f)).

**Discussion**

Human adipocyte models, which are genetically and functionally well characterized, are urgently needed for in vitro obesity research. We evaluated a human adipocyte model which was established from lipoma tissue of a patient with heterozygous germline PTEN deletion and was termed LipPD1 [6]. Germline PTEN mutations are frequently found in patients with PTEN hamartoma tumour syndrome (PHTS). Besides having an increased risk for malignancies [18], children with PHTS often develop multiple lipomas or non-malignant adipose tissue overgrowth, which can cause organ dysfunction due to rapid and invasive expansion [6,19]. This phenotype is specific for paediatric patients. Adults with PHTS rarely develop lipomas but were shown to have increased insulin sensitivity despite being more obese than age- and sex-matched controls [10]. PTEN is a lipid and protein phosphatase and a negative regulator of the PI3 K/AKT pathway, one of the major signalling pathways mediating cellular growth in response to insulin and other growth factors [20]. PTEN is also a tumour suppressor and frequently mutated in human malignancies.

Although being tumoural tissues, spontaneous non-syndromal lipomas have been shown to resemble dysfunctional adipose tissue, similar to what is seen in obesity [21], but lacking infiltration of inflammatory cells. This was also the case in our study, in which lipoma tissue morphology and gene expression were similar to subcutaneous adipose tissue of the same patient. Mutations or deletions in other tumour suppressor genes were excluded by Sanger sequencing, multiplex ligation-dependent probe amplification and comparative genomic hybridization both in the lipoma tissue and in lipoma-derived LipPD1 cells [6]. We can, however, not exclude the possibility that LipPD1 cells acquired somatic mutations during cell culture. A potential reason could be the accumulation of replication stress, as is described for the culture of human mesenchymal stromal cells [22].

We compared LipPD1 cells to primary SVF cells from healthy adult donors that had been cultured for the same time period and to the functionally well-characterized adipocyte model Simpson-Golabi-Behmel Syndrome (SGBS) cells [5,23] with amounts of PTEN comparable to primary SVF cells from healthy donors [17]. SGBS cells have been utilized to study white and brown adipocyte development, beiging of white adipocytes as well as effects of co-culture in a multitude of studies [23–34]. While the syndrome is associated with mutations in glypican 3 and 4 as well as in oral-facial-digital syndrome (OFD1) [35], the specific mutation which is carried by the SGBS cells and is causative for the SGBS phenotype of increased cellular lifespan and preserved capacity for adipocyte differentiation during long-term culture is not yet defined [23].

LipPD1 cells are haploinsufficient for PTEN function, which presumably is the cause for their increased cellular life span and enhanced in vitro adipocyte differentiation.
capacity. This is supported by earlier results using alpelisib, a specific PIK3CA inhibitor on LipPD1 cells. Alpelisib-treated LipPD1 cells differentiated into adipocytes less well and showed a decrease in proliferation. We did not detect induction of apoptosis or cell death, but found an upregulation of senescence marker
p16 and increased β-galactosidase staining indicating induction of senescence [13]. PTEN’s role in senescence seems to be cell-type specific and can be enhancing or suppressing. While an upregulation of PTEN in alveolar epithelial cells caused senescence-associated fibrosis [36], PTEN overexpression in glioblastoma neurospheres led to increased senescence [37]. A recent study in a mouse model with conditional PTEN knockout in adipose tissue depots showed a phenotype of adipose tissue redistribution with massive expansion of the PTEN knockout depot and reduced fat mass in other adipose tissue depots. We saw a similar phenotype in the PHTS patient from whose lipomas the LipPD1 cells were derived [6], pointing to a direct role for PTEN in regulating whole-body adipose tissue distribution.

Because of the reduced PTEN function, we expected LipPD1 cells to be more sensitive both in cellular responses towards insulin and in insulin signalling pathway activation compared to SGBS which express PTEN at a similar level as primary human SVF cells [38]. LipPD1 responses to insulin stimulation, however, were comparable to SGBS, with similar glucose uptake and capacity for adipocyte differentiation. These findings were supported by the fact that insulin-stimulated AKT activation was similar in both cell cultures after in vitro adipocyte differentiation. One reason for this could be the upregulation of other phosphatases during adipocyte differentiation that substitute for the reduced PTEN function. Possible candidates for this role could be PH domain leucine-rich repeat protein phosphatases (PHLPP) 1 and 2 [39], protein-tyrosine phosphatase 1B [40] or protein-tyrosine phosphatase receptor type F (PTP-RF) [41] as they target the insulin signalling pathway and were shown to be implicated in the development of insulin resistance. We found a higher expression of PTP-RF in LipPD1 compared to SGBS cells during adipocyte differentiation.

In contrast to the situation in adipocytes, we detected a significantly higher basal and IGF-I stimulated AKT activation in LipPD1 compared to SGBS preadipocytes, which corresponded to a higher amount of membrane-bound PIP3 analogue under basal conditions. This result recapitulates the findings of increased AKT phosphorylation in adipose tissue and mucosa-associated lymphoid tissue of patients with PHTS [10,39,42].

Conclusion

LipPD1 cells were comparable, but not equal to, SGBS cells in their capacity for adipocyte differentiation after long-term culture, expression of white adipocyte marker genes and adipocyte function. SGBS cells with slightly longer duration in continuous culture showed higher expression of some white adipocyte markers as well as a bigger size of 3D spheroids during in vitro adipocyte differentiation. In contrast to SGBS cells with unknown genetic alterations [23], the genetic lesion in LipPD1 cells has been characterized [6]. The usage of LipPD1 cells as adipocellular model is limited by the fact that they are no cell line with infinite capacity for passaging as they are not immortalized and no unlimited supply is possible. However, they can be easily cultured, differentiated into adipocytes by established protocols and function as mature adipocytes in key functional assays. LipPD1 cells can therefore be considered a new primary cell model with a prolonged life span, which is suitable to study adipocyte biology.

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Disclosure statement

All authors declare no conflict of interest.

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