Overcharged lipid metabolism in mechanisms of antitumor by *Tremella fuciformis*-derived polysaccharide

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Abstract. *Tremella fuciformis*-derived polysaccharide (TFP) is a natural macromolecular compound that is well known for whitening skin, as well as for its ability to regulate lipids and immunity. However, its mechanism of action is not clear. In the present study, B16 cells treated with TFP were inoculated subcutaneously in the right flank of C57BL/6 mice to explore the effect of TFP on melanoma in vivo. Western blotting, immunohistochemistry, immunofluorescence staining and transcription analysis of tumors were utilized to assess the expression of key molecules in production of melanin, lipid metabolism and immunity. It was found that TFP promoted B16 cell apoptosis and induced G2/M cell cycle arrest, which was associated with activation of cell cycle-related pathways. TFP induced the expression of glucose transporter type 4 and CD36, thus resulting in an increase in the uptake of lipids, which markedly suppressed sterol regulatory element-binding transcription factor 1 and phosphorylated-AMP-activated protein kinase expression; increased the number of lipids in the cell membrane, endoplasmic reticulum and nucleus; and induced the RNA expression of molecules related to lipid metabolism, as revealed by RNA-sequencing in vivo. Increased lipid binding, upregulated lipid storage, and elevated triglyceride and lipid catabolism resulted in disruption of cell volume homeostasis and activated innate immune response, thus inhibiting melanoma development and progression. These data revealed a novel molecular mechanism involved in the antitumor effect of TFP via lipid metabolism.

Introduction

Malignant melanoma is one of the most aggressive tumor types, and is characterized by high proliferation rates, poor prognosis and low survival rate (1). The high rate of metastasis is the main cause of mortality in patients with melanoma (2). Previous evidence has shown that altered tumor metabolism suppresses the infiltration of immune cells and helps melanoma cells to escape the immunity of the host (3).

Metabolic reprogramming is a hallmark of cancer (4), and a lipogenic phenotype is a main characteristic of cancer. Sterol regulatory element binding protein (SREBP) has been reported to be highly activated in tumors, and their levels and activity are tightly controlled by endogenous sterol levels via a negative feedback loop. Sterols regulate SREBP by controlling its endoplasmic reticulum (ER)-to-Golgi apparatus transport (5). SREBPs control lipogenesis and lipid uptake, SREBP1 preferentially regulates the lipogenic process by activating genes involved in fatty acid and triglyceride synthesis (6), whereas SREBP2 primarily controls cholesterol homeostasis by activating genes required for cholesterol synthesis and uptake (7).

Cancer cells become independent of systemic regulation due to their autonomous generation of lipids. Cancer cells engage in extensive *de novo* lipogenesis to produce lipids (8). Three important enzymes in *de novo* fatty acid synthesis, namely ATP-citrate lyase (ACLY), acetyl coenzyme A (CoA) carboxylase (ACC) and fatty acid synthase (FAS), are upregulated in numerous types of cancer (9). ACLY produces acetyl CoA, in the cytoplasm and provides the precursor of lipid synthesis, while ACC controls the rate-limiting step of fatty acid synthesis. FAS has been shown to play a key role in
tumor malignant growth and at the terminal step in de novo synthesis of fatty acids (10). The FAS and ACC genes are directly regulated by SREBP1 in fatty acid synthesis. ACLY involved in cholesterol synthesis also participates in fatty acid biosynthesis (11), and is required for glucose transporter type 4 (GLUT4)-mediated glucose-dependent fatty acid synthesis (12).

Cancer cells have alternative ways to acquire nutrients, and certain tumors develop adaptations that allow them to intake fatty acids directly from plasma or surrounding adipose tissue (13). Increasing lipid biosynthesis occurs in cancer cells, which influences the tumor microenvironment, and SREBP1 transcription factors respond to diverse environmental changes, such as lipid supply and inflammatory signals (14). Cells deal with excess lipids such as fatty acids by esterifying them to form more inert neutral lipids, such as triacylglycerols or sterol esters. Peroxisome proliferator-activated receptor γ (PPARγ) plays a major role in adipogenesis and storage of fatty acids such as triglycerides (15). The formation of these neutral lipids occurs within membrane bilayers; however, since bilayers are not suitable for storing large quantities of neutral lipids, an emulsion of neutral-lipid droplets forms in the cytoplasm (16). Lipid trafficking is highly organized and regulated in space. The aforementioned neutral-lipid emulsion originates in the ER, which serves as a phospholipid reservoir in cells to generate stable emulsions (17). When ER emulsions are delayed, this leads to accumulation of neutral lipids in the ER and blebbing out into inappropriate sites such as the nucleus (18). Lipids in the nucleus can sequester transcription factors and chromatin components, as well as generate lipid ligands for certain nuclear receptors (17).

Tremella fuciformis-derived polysaccharide (TFP) is extracted from Tremella fuciformis, which is a common nutritional food in China and also a traditional Chinese medicine due to its anti-inflammatory and anticancer properties, and its ability to strengthen immunity, and reduce hypertension and blood glucose (19). TFP has also been used clinically for whitening skin, cancer and anti-aging treatments in China. However, the protective mechanism of TFP is not clear yet. The present study explored the function of TFP on the lipid metabolism of B16 cells, and evaluated lipid metabolism changes influenced the tumor growth and microenvironment in vivo.

Materials and methods

Cell culture and treatment. The B16 mouse melanoma cells line was purchased from the cell bank of Chinese Academy of Sciences (Shanghai, China). Cells were cultured in DMEM medium with phenol red, supplemented with 10% FBS, 100 U/ml penicillin, and 100 μg/ml streptomycin in a dynamic incubation system at 37°C in a 5%–CO₂ humidified atmosphere. The cells were treated in the presence of series of dosages of TFP (0, 312.5, 625, 1,250, 2,500 and 5,000 μg/ml) for 24 h.

HPLC analysis of TFP. The content of TFP in B16 cells was detected by high-performance lipid chromatography (HPLC). Briefly, the B16 cells with or without TFP exposure were centrifuged at 1,000 x g for 3 min at room temperature and suspended with ultrapure water in ice bath for 10 min, then centrifuged for 10 min. The supernatant was collected for HPLC. HPLC separation was performed on sugar column (water ultra-hydrogel 500) and detected using differential detector 1200 (Agilent Technologies, Inc.). The mobile phase was ultrapure water. The column temperature was kept at 85°C and the flow rate was set at 0.6 ml/min.

Animal model and drug treatment. A total of 18 male C57BL/6 mice (6-8 weeks old; weight, 25 g) were purchased from Shanghai Experimental Animal Center, Chinese Academy of Sciences (Shanghai, China). The mice were housed in temperature (25°C) and humidity (30-40%)-controlled room, kept on a 12-h light/dark cycle and provided with unrestricted amount of rodent chow and water. The animal protocol was approved (approval no. 2020002) by the Animal Care and Use Committee of Minnan Normal University (Zhangzhou, China).

To study the direct effects of TFP in vivo tumor development, B16 cells were preincubated in vitro with two dosages of TFP (2,500 and 5,000 μg/ml). After 24 h, the cells were washed 3 times, and 1x10⁶ B16 cells/ml resuspended in 100 μl of PBS were inoculated subcutaneously in the right flank of the mice. The tumor volume 2,000 mm³ calculated according to length and width was a limit or the maximum permitted tumour burden in the present study. At the experiment terminal, the mice were sacrificed by cervical dislocation, sacrifice was confirmed when the mice had ceased breathing and did not respond to stimulation. The tumors were collected and weight was measured.

Cell proliferation assay. For determination of cell growth using Real Time Cellular Analysis (RTCA), plate 16 assemblies were seeded with B16 cells (1x10⁴ cells/well). The plate was then assembled on the RTCA DP analyzer, after 24 h cells were treated with five dosages of TFP (312.5, 625, 1,250, 2,500, 5,000 μg/ml), and data were gathered at 5-min intervals for 120 h at 37°C in 5% CO₂.

Flow cytometric analysis of cell cycle, apoptosis and co-localization. Cells were harvested by treatment with 0.25% trypsin, resuspended with PBS, then stained with propidium iodide and Annexin V. The cells were then assembled on the RTCA DP analyzer, after 24 h cells were treated with single cells (1x10⁴ cells/well). The plate was then assembled on the RTCA DP analyzer, after 24 h cells were treated with five dosages of TFP (312.5, 625, 1,250, 2,500, 5,000 μg/ml), and data were gathered at 5-min intervals for 120 h at 37°C in 5% CO₂.

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membrane, ER, Golgi, or nucleus with lipid was analyzed using colocalization analysis application Wizard in IDEAS software.

Nile Red staining. B16 cells cultured on confocal dish (Coverglass Bottom Dish) were treated with different concentrations of TFP for 24 h. Then the cells were stained by Nile red (1:500) and DAPI (1:1,000) for 30 min avoiding light, and then images were captured using confocal laser scanning microscope (Leica Microsystems GmbH).

Western blotting (WB). The protein levels and phosphorylation status were examined by WB. The cells were washed twice with ice-cold PBS and lysed in RIPA protein lysis buffer containing 1 mM PMSF (cat. no. R0010, Beijing Solarbio Science & Technology Co., Ltd.). The samples were centrifuged at 13,000 x g for 10 min at 4˚C before the supernatant was collected and the protein concentration was measured using a BCA kit (cat. no. P0012, Beyotime Institute of Biotechnology). Samples were boiled for 5 min after being mixed with a 5X loading buffer solution. The protein lysates (30 µg) were resolved in 8-10% SDS-polyacrylamide gels and transferred to nitrocellulose membranes. The membranes were blocked using 5% skimmed milk for 1 h at room temperature, washed three times with TBST (0.1% Tween 20) for 10 min and then incubated overnight at 4˚C with various primary antibodies: anti-Adenosine 5'-monophosphate-activated protein kinase (AMPK; 1:2,000; cat. no. ab32047; Abcam), anti-phosphorylated-AMP-activated protein kinase (p-AMPK; 1:2,000; cat. no. EPR3051; Abcam), anti-SREBP-2 (1:500; cat. no. ab30682; Abcam), anti-PPARγ (1:1,000; cat. no. C26H12; Cell Signaling Technology, Inc.), anti-ACLY (1:1,000; cat. no. ab40793; Abcam), anti-Lysosomal-associated membrane protein1 (LAMP1; 1:100), and mouse anti-mouse antibody (PDL1, 1:100; cat. no. ab238697; CD68, 1:100, cat. no. NB600-985; CD86, 1:50, cat. no. ab220188; all from Abcam) at 4˚C overnight. The primary antibody was visualized by using anti-rabbit Alexa Fluor 647 (1:1,000; cat. no. P0180; Beyotime Institute of Biotechnology), and anti-mouse FITC conjugated antibody (1:1,000; cat. no. P0196; Beyotime Institute of Biotechnology). The images were captured using Leica TCS SP8 confocal laser microscope.

RNA-seq, data processing, and gene annotation. RNA of Model group and 5,000 µg/ml TFP group were extracted. Total RNA was extracted using TRIzol reagent kit (Invitrogen; Thermo Fisher scientific, Inc.). RNA quality was assessed on an Agilent 2100 Bioanalyzer (Agilent Technologies, Inc.). After total RNA was extracted, mRNA was enriched by Oligo (dT) beads. Then the enriched mRNA was fragmented into short fragments using fragmentation buffer and reversely transcribed into cDNA by using NEBNext Ultra RNA Library Prep Kit for Illumina (New England Biolabs). The purified double-stranded cDNA fragments were end repaired, a base added, and ligated to Illumina sequencing adapters. The ligation reaction was purified, ligated fragments were subjected to size selection (200 bp) by AMPure XP Beads (1.0X) and PCR amplification. The resulting cDNA library was sequenced using Illumina Novaseq6000 by Gene De novo Biotechnology Co. (Guangzhou, China).

High quality clean reads were further filtered by fastp (version 0.18.0). The gene expression level was normalized by using Fragments Per Kilobase of transcript per Million mapped reads. The genes with a fold change ≥2 and P<0.05 were considered as significant differentially expressed genes (DEGs). Three samples were prepared from each treatment.

Gene Ontology (GO) enrichment of DEGs was performed, the significantly enriched GO terms in DEGs was performed by comparing to the genome background using hypergeometric test. The biological pathways of the DEGs were enriched to the Kyoto Encyclopedia of Genes and Genomes (KEGG) using hypergeometric test as the above GO term enrich- ment (https://www.genome.jp/kegg/). Gene Set Enrichment Analysis (GSEA) (22) was used to identify whether a set of genes in specific GO terms/KEGG pathways/Reactome pathways shows significant differences in two groups. The sequences data reported in the present study was archived in the Sequence Read Archive (SRA) with the accession number BioProject ID PRJNA772896 (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA772896/).

Statistical analysis. The data was examined using the statistical analysis program SPSS 22.0 (IBM Corp.). The data are expressed as the mean ± SD. Tumors weight were analyzed by using one-way Analysis of Variance (ANOVA)

Hematoxylin and eosin (H&E) staining. Paraffin-embedded sections were cut at 3 µm thickness, mounted on microslides and processed for H&E staining, as previously described (21). The stained slides were dehydrated and mounted using Cytoseal. Images of the slides were captured (magnification, x200) with a light microscope (Olympus Corporation).

Immunohistochemistry. For immunohistochemistry, tissue sections were deparaffinized with xylene and stepwise rehydrated with serial dilutions of ethanol. After epitope retrieval with citric acid buffer under high temperature and pressure for 1-2 min, and endogenous horseradish peroxidase activity quenching with 3% H2O2 for 10 min, the tissue sections were then blocked in 5% goat serum (cat. no. C0265; Beyotime Institute of Biotechnology) at room temperature for 1 h and incubated with rabbit anti-mouse (ACLY, 1:100; PPARγ, 1:100; LAMP1, 1:100), and mouse anti-mouse antibody (PDLI, 1:100, cat. no. ab238697; CD68, 1:100, cat. no. NB600-985; CD86, 1:50, cat. no. ab220188; all from Abcam) at 4˚C overnight. The primary antibody was visualized by using anti-rabbit Alexa Fluor 647 (1:1,000; cat. no. P0180; Beyotime Institute of Biotechnology), and anti-mouse FITC conjugated antibody (1:1,000; cat. no. P0196; Beyotime Institute of Biotechnology). The images were captured using Leica TCS SP8 confocal laser microscope.
followed by Fisher's least significant difference post hoc test for comparison between two groups. P<0.05 was considered to indicate a statistically significant difference.

**Results**

**TFP structure and HPLC analysis.** TFP is a polysaccharide whose main chain is mannan, which is a linear (1→3)-linked backbone, and its side chain is composed of mannose, glucuronic acid and xylose in uncertain order, as shown in Fig. 1A. The results of HPLC analysis results showed that there was no obvious TFP peak, since the TFP standard occurred at 8.183 min in B16 cells exposed to TFP, as shown in Fig. 1B. These data suggested that there was no TFP in B16 cells within the detection capability of HPLC.

**Effects of TFP on anti-proliferation, pro-apoptosis and cell cycle arrest in vitro.** The viability of B16 cells treated with TFP decreased in a dose-dependent manner, particularly at 1,250, 2,500 and 5,000 µg/ml TFP, which significantly inhibited the rate of B16 cell proliferation compared with the effects of other dosages of TFP (Fig. 2A). The effect of TFP on the relative proliferation rate of B16 cells was measured by its IC$_{50}$, which was calculated to be 1.836x10$^7$ µg/ml.
To determine whether TFP inhibits B16 cell proliferation via cell cycle arrest, the DNA contents of the cells were measured by flow cytometry upon staining the cell nuclei with DAPI (Figs. 2B and C, and S1). The data showed that high dosages of TFP increased the number of cells in the G2/M phase from 9.5% (0 µg/ml TFP) to 10.6% (1,250 µg/ml TFP), 19.1% (2,500 µg/ml TFP) and 24.6% (5,000 µg/ml TFP), while reducing the number of cells in the G1 phase from 88.7% (0 µg/ml TFP) to 85.8% (1,250 µg/ml TFP), 77.1% (2,500 µg/ml TFP) and 70.9% (5,000 µg/ml TFP). In addition, TFP slightly increased the number of cells in S phase from 1.27% to 1.44, 1.58, 2.32, 2.17 or 1.73% depending on the concentration employed. These results suggested that TFP inhibited cell proliferation mainly by inducing G2/M cell cycle arrest without cycle progression to G1 phase through mitosis in B16 cells.

Flow cytometry was used to evaluate the state of cell apoptosis (Figs. 2D and S1). The results showed that the proportion of early apoptotic cells in untreated B16 cells was 1.13%, which slightly decreased with increasing concentration of TFP (312.5, 625, 1,250 and 2,500 µg/ml). When the concentration of TFP reached 5,000 µg/ml, 2.16% of cells were markedly early apoptotic cells. B16 cells exposed to 0, 312.5, 625, 1,250, 2,500 and 5,000 µg/ml of TFP for 24 h experienced an induction of 4.05, 7.02, 5.32, 7.19, 13.15, 12.40% in late apoptosis, respectively. In addition, the proportion of necrotic cells increased gradually with increasing concentrations of TFP. These data revealed that TFP at suitable concentrations could effectively induce B16 cell death following late apoptosis, rather than early apoptosis, and necrosis.

Lipids uptake reprograms lipogenesis profile in vitro. TFP markedly suppressed SREBP1 nuclear translocation, and suppressed SREBP1 target proteins, such as ACC and FAS, leading to reduced de novo lipogenesis (Fig. 3A). Unexpectedly, the ACLY and plasma lipids levels increased...
Figure 3. Uptake of exogenous lipids and reprogrammed lipogenesis profile in vitro. (A) The levels of lipogenesis-related protein were detected by western blotting. (B) Fluorescence analysis of lipids. SREBP, sterol regulatory element binding protein; GLUT4, glucose transport 4; ACC, Aacetyl CoA carboxylase; FAS, fatty acid synthase; ACLY, ATP-citrate lyase.
according to increasing TFP dosages (Fig. 3B). To explain why these apparent contradictions coexisted in B16 treated by TFP, the levels of GLUT4 and CD36 (lipid transport receptor) were analyzed, since cancer cells use glucose as well as fatty acids for ATP production, and GLUT4 and CD36 migrate together in response to either stimuli, i.e. glucose or fatty acids (23). The results confirmed that TFP induced high expression of GLUT4 and CD36 (Fig. 3A), which resulted in an increased uptake of exogenous lipids, which were present at higher lipid fluorescence intensity in the cell membrane (Fig. 4A; panels a and b) and plasma, including ER (Fig. 4B; panels a and b), nucleus (Fig. 4D; panels a and b) as compared with that of control group. However, the uptake of lipids present in the Golgi apparatus was decreased at dosage of TFP5000 (Fig. 4C; panels a and b). The aforementioned data was in accordance with higher co-localization ratios of lipids with the cellular membrane and nucleus (Fig. 4E), and higher lipid intensity of the whole cell compared with that of the vehicle group (Fig. 4F).

The intracellular accumulation of lipids, particularly in the ER, inhibited the maturation of SREBP1 and the expression of full-length SREBP1 (Fig. 3A). Lipids are important molecules in the nucleus, there is emerging evidence (24) for the existence of a nuclear population of lipids, which has been proposed to directly modulate lipid metabolism in the nucleus. In the present study, the lipids in the nucleus may have the functions of generating lipid ligands for PPARγ, or cause chromosome segregation abnormalities, or modulate exogeneous lipid metabolism and storage.

**TFP delays B16 allograft tumor growth and reduces tumor weight.** Above results have shown that TFP induces the regulation of lipogenesis in B16 cells. To explore whether this response is capable of preventing or slowing tumor growth, 1x10⁶ B16 cells treated with or without TFP were subcutaneously injected in the right flank of C57 BL/6 mice (Fig. 5A). There were no significant differences in body weight of TFP treated groups referring to model group (Fig. 5B), while tumors grew more slowly and the tumor weights of the TFP-treated groups were less than those of B16 tumor-bearing mice (Fig. 5C). The tumor weights in the model and TFP 2,500 and 5,000 µg/ml groups were analyzed by one-way ANOVA followed by Fisher's least significant difference post hoc test (Fig. 5D), which revealed a significant difference among the three groups [F(2,14)=13.595, P=0.001]. The tumor weight of the TFP 2,500 µg/ml group exhibited a significant decrease compared with that of the model group (P=0.007), as well as the TFP 5,000 µg/ml group (P=0.001).

The histological characteristics were evaluated by (H&E) staining (Fig. 5E), which revealed increased pigmentation located in the model tumor section (as indicated by more white arrows), but less pigmentation (less white arrows) and scattered vacuoles in the TFP-treated tumor section (red arrows), which may be due to lipid accumulation, and was dissolved by xylene during H&E staining. These data suggested that TFP-treated B16 cells reprogrammed their lipid metabolism profile, and TFP promoted lipid accumulation in B16 cells.

**WB analysis showed upregulation of ACLY and PPARγ** (Fig. 5F and G). Upregulated ACLY and PPARγ activity increased lipid formation. As revealed by immunohistochemistry (Figs. 6A and S2), TFP upregulated the expression of ACLY and PPARγ, and promoted macrophage infiltration (as indicated by the increase in CD86 expression). Notably, LAMP1 expression increased according to the increase in TFP dosage. In certain areas of the TFP-treated groups, the formation of certain strip (long and linear) and round vacuoles was observed (the round vacuoles gathered at one end, while the strip vacuoles aligned orderly, similar to a ‘balsam pear’ surface), which was accompanied by the presence of numerous macrophages (Fig. 6B). Increased macrophage infiltration occurred when there was no formation of these structures. Not far from those structures, a number of bigger vacuoles could be observed, such as lipid accumulation dissolved by xylene in the process of staining, similar to the vacuoles noted during H&E staining (Fig. 5E). These data indicated that TFP controlled B16 allograft tumor growth through lipid metabolism regulation, and enhanced the antitumor immune response through macrophage infiltration and upregulation of LAMP1.

**RNA-sequencing (RNA-seq) analysis of differentiated gene expression and enriched pathways.** To further uncover the molecular mechanism of TFP, tumor gene expression profile analyses were conducted with the aim of uncovering valuable information for pharmacological research on therapeutic targets. Gene Ontology pathway enrichment analysis showed that negative regulation of mitotic cell cycle, regulation of cell proliferation and cell proliferation were significantly altered by TFP (Fig. 7A). The enrichment in pathways of cell cycle/proliferation confirmed the results of cell experiments.

Gene set enrichment analysis (GSEA) of the cell cycle-related RNA-seq results (Fig. 7B) showed that TFP treatment was strongly associated with the gene signature of cell cycle arrest, cell cycle and cycle phase transition. Furthermore, negative regulation of cell process, cell cycle and cell cycle phase transition were significantly positively associated with TFP. These data indicated that cell cycle regulation played an important role in the antitumor activity of TFP.

To determine whether the antitumor effect of TFP occurred through activation of lipid metabolism, gene expression was examined. Gene microarray analysis demonstrated the upregulation of mRNAs encoding genes involved in lipid metabolism (Fig. 8A), including glucolipid receptors (GLUT4), lipogenesis-related genes (ACLY), a set of mRNAs involved in the tricarboxylic acid cycle (Fig. 8B), and a number of lipolysis-related genes (such as LPL: lipoprotein lipase). GSEA of regulation of lipolysis in adipocytes further provided clues on the sources of lipids: upregulation of lipolysis of adipocyte in tumor indicated the possible sources of more lipids deposit in tumor (Fig. 8C). The omics effect of TFP on lipid metabolism is consistent with the aforementioned phenotypic results associated with increased lipid deposits in mitochondria, nucleus, ER and membrane, as well as the WB results related to increased glycolipid transport and increased lipid accumulation in allografts.

Subsequently, to identify the potential reason responsible for the activated lipid metabolism and to uncover the underlying molecular mechanisms of TFP involved in lipid metabolic processes, GSEA was used to mine the RNA-seq data. The results revealed that TFP treatment was associated...
Figure 4. TFP regulates lipid distribution in B16 cells. Flow cytometric analysis of co-localization of lipids and (A) membrane, (B) ER, (C) Golgi apparatus, and (D) nucleus. Lipid intensity and representative images of (A panels a and b) membrane, (B panels a and b) ER, (C panels a and b) Golgi apparatus, and (D panels a and b) nucleus. (E) Statistical analysis of the results of panels A-D. (F) Lipid intensity of the whole cell. ER, endoplasmic reticulum; TFP, *Tremella fuciformis*-derived polysaccharide.
with the gene signatures of ‘positive regulation of lipid binding’ (Fig. 9A), ‘positive regulation of lipid storage’ (Fig. 9B), ‘triglyceride catabolism’ (Fig. 9C) and ‘lipid catabolism’ (Fig. 9D). These results indicated that TFP not only promoted lipid uptake in cells, but also provoked lipid catabolism.

Next, it was hypothesized that metabolic changes could induce the immune system. It was found that TFP induced the enrichment
greatly increases the viscosity of liquids, and the glucuronic acids in TFP have the ability to entrap cholesterol (25). These combined TFP with cell membrane or intercellular cholesterol were easily removed by centrifugation in the process of intracellular fluid extraction. This may explain why no TFP was detected in B16 cells by HPLC analysis. The receptors of TFP have not been identified to date. The antitumor mechanism of TFP may involve the upregulation of glucolipid receptors, such as CD36 and GLUT4, which were markedly elevated in vitro and in vivo in the present study.

The current study demonstrated the influence of TFP on the proliferation, cell cycle and apoptosis of B16 cells. TFP inhibited cell proliferation, induced cell apoptosis and altered cell cycle distribution. In addition, TFP downregulated SREBP, which regulates cellular lipogenesis and lipid homeostasis (26), and has been shown to be involved in other cellular functions, such as regulation of cell cycle and cell proliferation (27). TFP inhibited de novo lipogenesis through SREBP1, thereby inhibiting cell proliferation. TFP could prevent cells from completing mitosis, arresting them at a ‘lipogenic’ G2/M phase. AMPK has been reported to exert anti-proliferative action (28). AMPK is a cell-intrinsic regulator of the cell cycle that coordinates cellular proliferation with energy source availability, and regulates a number of key cellular metabolic enzymes (29). The inhibitor effect of TFP on phosphorylated (p)-AMPK suggested that the anti-proliferative effect of TFP was AMPK independent. The nutrient sensor AMPK negatively regulates SREBP to limit lipogenesis under nutrient-limiting conditions (30). Cancer cell metabolism could be changed if sufficient ATP and the building blocks needed to sustain molecular synthesis were available. In the present study, the expression levels of AMPK and particularly those of p-AMPK were decreased by TFP, and AMPK positively regulated SREBP when CD36 and GLUT4 were notably upregulated by TFP. Loss of AMPK or downregulation of AMPK results in energy stress-driven apoptosis, and slows down tumor progression (31); AMPK inhibitors rather than activators could preferentially trigger cancer cell death in the context of metabolic stress (32). PPARγ is a transcription factor capable of promoting pleiotropic effects, including the control of cell proliferation. Pharmacological agonists of PPARγ were used in cancer therapy with the aim of promoting cell cycle deceleration (33). TFP as AMPK inhibitor and PPARγ agonist facilitated to control tumor growth, which was confirmed by our present studies.

Finally, the cell cycle arrest of B16 cells treated with TFP could be due to both the inhibition of de novo fatty acid synthesis (causing inhibition of SREBP1) as well as the accumulation of exogeneous lipids in the nucleus (causing induction of PPARγ and impairment of cell proliferation through chromosome segregation abnormalities).

The present findings indicated that TFP activated lipid metabolism and the immune system. A balance between lipogenesis, lipid uptake and intracellular lipolysis is required to maintain lipid homeostasis (34). Cancer-associated adipocytes have been found to promote lipid accumulation (35). In the case of malignant melanoma, the tumor secreted an inhibitor of adipose LPL termed melanoma cachexia factor, which caused the inhibition of extracellular lipolysis (36). Activation of the transmembrane channel for exogenous free fatty acid uptake...
(CD36) and LPL (a key enzyme for extracellular lipolysis) confirmed the disposition of extracellular lipid stores and lipolysis in cells. The levels of free fatty acids, monoacylglycerides and diacylglycerides were not significantly elevated (data not shown), which suggested that lipid metabolism existed in cancer cells and cancer-associated adipocytes.

Figure 7. RNA-sequencing analysis confirms the effect of TFP on cell cycle. (A) Top 20 results of the GO pathway enrichment based on the DEGs identified in TFP-treated B16 allograft. In the scatter plot, the RichFactor corresponds to the ratio of the number of DEGs noted in a pathway term to all the genes noted in such pathway term, and a greater RichFactor means greater intensity. The Q-value is the corrected P-value ranging from 0 to 1, and a lower Q-value represents greater intensity. (B) GSEA revealed that cell cycle-related genes were significantly regulated. TFP, Tremella fuciformis-derived polysaccharide; GO, Gene Ontology; DEGs, differentially expressed genes; GSEA, Gene set enrichment analysis.
Elevated SREBP1 in cancer cells increased de novo lipogenic gene expression, enhanced fatty acid synthesis, and accelerated triglyceride accumulation; however, these metabolic abnormalities were reversed by the treatment of B16 cells with TFP. TFP inhibited SREBP1 and AMPK, an enzyme that inhibits lipid synthesis through the phosphorylation and inactivation of key lipogenic enzymes. The net effect on lipogenic metabolism depends on a balance between opposing effects. In the present study, the net result was a decrease in SREBP1 and AMPK, while certain lipid enzymes were reduced (FAS and ACC) and certain others activated, particularly those participating in the tricarboxylic acid (TCA) cycle. Importantly, several signaling molecules involved in the processes of immune cell activation and transformation are derived from metabolites of the TCA cycle (37). These results indicated the crucial role of lipid metabolism in the regulation of the tumor immune microenvironment and the feasibility of targeting lipid metabolism to regulate tumor immunity.

Increasing evidence indicated that aberrant de novo lipogenesis is being increasingly recognized as an important feature of malignant transformation (38), by contrast, the majority of normal human tissues use exogenous lipids, while de novo fatty acid synthesis is generally suppressed (39). De novo lipogenesis also determines the sensitivity of cancer cells to oxidative stress-induced cell death, and de novo synthesis was the preferred mechanism for fatty acid acquisition over lipolysis or receptor-mediated endocytosis (40). Tumor cells do not solely rely on de novo lipogenesis, but also utilize exogenous fatty acids for generating lipids for proliferation. Thus, despite the dominant role of de novo lipogenesis, exogenous fatty acids may also play an important role in cancer pathogenesis (41).

After B16 cells were exposed to TFP, their lipid metabolism was altered. TFP treatment resulted in the downregulation of de novo fatty acid synthesis due to feedback suppression with concomitant increase in exogenous fatty acid uptake. This metabolic change was facilitated by the upregulation of several key transporters, enzymes and signaling pathways. Specifically, CD36 and GLUT4 were remarkably upregulated; de novo lipogenesis was completely abolished; the lipid distribution in the cytoplasm was rearranged; and the lipid content in the cell membrane, ER and nucleus was elevated, which was accompanied by an increase in ACLY and PPARγ expression. The RNA-seq data precisely depicted the effect of TFP on lipid metabolism and processing, showing its antitumor effects. TFP caused excess lipid metabolism and dysregulated lipid storage in B16 cells, which resulted in the irreversible disruption of cell volume homeostasis and activation of the immune system.

There were a number of limitations to the present study. Firstly, functional experiments to demonstrate the effect of TFP in differentiated adipocytes should be performed to

Figure 8. RNA-sequencing analysis confirms the effect of TFP on lipid metabolism. (A) Heatmap representation of differentially expressed genes associated with lipid metabolism in tumors treated with TFP. (B) Gene set enrichment analysis of the RNA-seq data was used to identify enriched biological pathway ‘TCA cycle’. (C) Gene set enrichment analysis of the RNA-seq data was used to identify enriched biological pathway ‘regulation of lipolysis in adipocyte’. TFP, Tremella fuciformis-derived polysaccharide.
Figure 9. RNA-sequencing analysis confirms the mechanism of TFP on lipid metabolism and the immune system. Gene set enrichment analysis identified (A) positively enriched lipid binding, (B) lipid storage, (C) triglyceride, (D) lipid catabolic process, (E) cell volume homeostasis, (F) response to reactive oxygen species and (G) innate immune system. Heatmap representation of differentially expressed genes associated with immune response in tumors treated with TFP. (H) The vertical black lines indicated the position of each gene in the studied gene set. The normalized enrichment score and false discovery rate values are shown for each pathway. Heatmap representation of differentially expressed genes associated with the immune system in tumors. TFP, *Tremella fuciformis*-derived polysaccharide.
demonstrate a specific role in TFP augmenting lipolysis of adipocytes. Secondly, mechanism of interactions between adipocytes and B16 cells should be explored. Thirdly, tumor cells manipulated ex vivo may serve as a uniquely effective carriers for the therapeutic delivery of bioactive cytokines to parental tumor by their homing properties especially highly metastatic cells (42). It is rarely reported that tumor cells modified with polysaccharides are used to treat tumors. The following antitumor effect and mechanism of B16 cells reprogrammed by TFP in vein in lower dosage was urgently needed in B16 subcutaneously implanted mice.

In conclusion, the present study identified the function of TFP on B16 cells, along with its unexpected molecular mechanism of action responsible for its antitumor effects, as summarized in the schematic diagram of Fig. 10. TFP inhibited de novo lipogenesis through downregulation of sterol regulatory element-binding transcription factor 1 while inducing the presence of lipids in the cytoplasm of B16 cells. This apparent contradiction was resolved by transcriptomic analysis in vivo, which revealed that the lipolysis of adipocytes supplied the lipids for B16 cells, which enhanced lipid binding, storage and catabolism. Overcharged lipid mechanisms depleted B16 cells and induced cell volume homeostasis disruption, thus activating the cellular response to reactive oxygen species and the innate immune response. TFP, Tremella fuciformis -derived polysaccharide.

Figure 10. Scheme showing the potential mechanism of action of TFP, including lipid changes in vitro and transcriptional profile in vivo. TFP inhibited de novo lipogenesis through downregulation of sterol regulatory element-binding transcription factor 1 while inducing the presence of lipids in the cytoplasm of B16 cells. This apparent contradiction was resolved by transcriptomic analysis in vivo, which revealed that the lipolysis of adipocytes supplied the lipids for B16 cells, which enhanced lipid binding, storage and catabolism. Overcharged lipid mechanisms depleted B16 cells and induced cell volume homeostasis disruption, thus activating the cellular response to reactive oxygen species and the innate immune response. TFP, Tremella fuciformis -derived polysaccharide.

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Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions
XL and YP conceived and designed the study, QS and XL performed the experiments. XL analyzed the data and wrote the manuscript. XL and YP confirm the authenticity of all the raw data. All authors read and approved the final version of the manuscript.

Ethics approval and consent to participate
The present study was approved (approval no. 2020002) and supervised by Experimental research Ethics Committee of Minna Normal University (Zhangzhou, China). All animal experiments were performed according to the relevant regulatory standards and were performed in accordance with the Experimental animal research Ethics Committee of Minna Normal University.

Patient consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

References


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