Atractyloside targets cancer-associated fibroblasts and inhibits the metastasis of colon cancer

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Background: Several evidences have proved that cancer-associated fibroblasts (CAFs) play a crucial role in tumor progression. In fact, CAFs form a major component of tumor microenvironment (TME). Therefore, the development and metastasis of tumors can be effectively inhibited by small molecular compounds that target CAFs.

Methods: In this study, we mainly analyzed the expression profile of colon cancer (CC). We determined the intensity of CAFs in CC tissues by using the immune cell infiltration score. Gene enrichment analysis and the screening of differentially expressed genes were performed on the basis of the intensity of CAFs in CC tissues. We screened the small molecular compounds that were converted from differentially expressed genes. The results indicated that atractyloside was a small molecular compound related to CAFs in CC tissues. We identified the relationship between atractylosides and CAFs through target protein analysis and network analysis, and verified the inhibition effect of atractylosides on CC cells (CCC) by migration assay and scratch wound-healing assays.

Results: We found that many target proteins of atractyloside, such as the matrix metalloproteinase family and integrin proteins, were related to the biological function of CAFs. By performing network analysis, we found that the target proteins FGF1, ITGB1, and EDNRA were closely related to tumor angiogenesis, while the target proteins MMP9 and ITGAV were correlated to an extracellular matrix (ECM) and cell motility. These findings which further confirmed the relationship between atractylosides and CAFs. In addition, transwell cell migration and scratch wound-healing assays proved that atractylosides could significantly inhibit the migration of CCCs.

Conclusions: The atractyloside might be a small molecular compound that potentially targets CAFs and inhibits the development as well as metastasis of CC by changing the TME.

Keywords: Cancer-associated fibroblasts (CAFs); colon cancer (CC); atractyloside; tumor microenvironment (TME)

doi: 10.21037/atm-20-1531
View this article at: http://dx.doi.org/10.21037/atm-20-1531
Introduction

The tumor microenvironment (TME) is a complex tissue environment, which comprises cancer cells, cancer-associated fibroblasts (CAFs), inflammatory cells, macrophages, myeloid-derived suppressor cells, mast cells, and extracellular matrix (ECM) (1). Since CAFs are major constituents of the TME, they play a crucial role in the occurrence, development, and metastasis of tumors. Furthermore, CAFs are activated fibroblasts in tumors and are mostly derived from fibroblasts, but they may be also derived from smooth muscle cells of vasculature, pericytes, and bone marrow-derived mesenchymal cells, or from epithelial or endothelial-mesenchymal transition. Fibroblast activation protein (FAP) and α-smooth muscle actin (α-SMA) are usually considered as the molecular markers of CAFs (2,3). However, the COX-2/PGE2 signal, TGF-β, and exosomes are the three main cross-talks, which facilitate extensive interactions between CAFs and cells of the TME. Moreover, CAFs have positive or negative interactions with all protumoral cells, including cancer cells, regulatory T cells, endothelial cells, M2 macrophages, cancer stem cells, myeloid-derived suppressor cells, tumor-associated dendritic cells, cytotoxic T lymphocyte, natural killer (NK) cells, etc. These cross-talks are usually responsible for the progression of tumor (4-6). Furthermore, CAFs modulate anti-tumor immunity responses at various levels, potentially affecting both innate and adaptive anti-tumor immunity responses. Thus, CAFs play a pivotal role in tumor progression (7).

The initial step of metastasis involves the migration and invasion of cancer cells derived from the primary site. Interestingly, CAFs promote the migration and invasion of cancer cells through paracrine signaling of cytokines. In this process, one of the major cytokines is TGF-β. Furthermore, CAFs alter the ECM environment to promote the migration and invasion of cancer cells (8). Because CAFs are frequently found in colon cancer (CC) tissues, they could be a potential marker in the prognosis of CC. In fact, several studies have reported about the relationship between CAFs and CC (9,10). In CC tissues, CAFs secrete cytokines and chemokines that target endothelial cells, such as VEGF, and induce angiogenesis (11). Most anti-cancer drugs target cancer cells, but the TME supports the growth and invasion of cancer cells. Therefore, there are still some targets that need to be investigated. Because CAFs are linked to tumor drug resistance, extensive research studies must be conducted to identify drugs targeting CAFs. This strategy would be useful in solving the issue of tumor drug resistance (12). However, CAFs create a physical barrier by synthesizing and secreting ECM components and by participating in the constitution of the TME. Therefore, drugs targeting CAFs should reorganize the ECM and enhance the permeability of anti-tumor drugs. By secreting proinflammatory cytokines and chemokines, CAFs induce immunosuppressive cells into the tumor stroma and reject effector T cells. Drugs targeting CAFs profoundly enhance the immune treatment of cancers (13).

In this study, our aim is to find small molecular compounds that target CAFs in CC tissues. Immune cell infiltration score was used to determine the intensity of CAFs in CC tissues. In the high-intensity CAFs group and the low-intensity CAFs group, the differentially expressed genes were screened and converted into small molecular compounds. To confirm the relationship between small molecular compounds and CAFs, we analyzed the potential mechanism of target proteins in small molecular compounds. In this study, we were able to find small molecular compounds that target CAFs in CC tissues. These molecules could be used to inhibit the development and metastasis of CC. We presented the following article in accordance with the MDAR checklist (available at http://dx.doi.org/10.21037/atm-20-1531).

Methods

Screening of small molecular compounds targeting CAFs

Estimate the Proportion of Immune and Cancer Cells

Screening of small molecular compounds targeting CAFs
groups (16,17). The analysis was based on the following parameters: biological processes, molecular functions, and cell components.

After comparing the biological processes of the high-intensity CAFs group and the low-intensity CAFs, 20 gene sets were selected with higher enrichment. From each gene set, genes that had enrichment function were collected. Thus, we obtained enrichment genes from the two groups.

After comparing the expression data of 239 CC cases in the high-intensity CAFs group and the low-intensity CAFs, we screened the differentially expressed genes. GeneSpring software was used for screening the differentially expressed genes. The parameters were compared with unpaired t-test, the P value was calculated, and multiple testing was corrected with Bonferroni FWER (18). We selected differentially expressed genes (P<0.05) that had a folder change of one time. In the two CAFs groups, intersections were analyzed between the screened differentially expressed genes and enriched genes. The genes obtained by intersection were imported into Connectivity Map. Then, small molecular compounds of higher relativity were screened by considering their enrichment score (Figure 1). Connectivity Map collected the expression profiles of tumor cells treated with 1,309 small molecular compounds (19,20). The data was used to determine the relationship between differentially expressed genes and small molecular compounds. Thus, we could screen small molecular compounds related to tumor cells by determining the changes in tumor tissue expression profiles, which were converted from the Connectivity Map.

**Prediction of target protein of atractyloside and the construction of signal-regulating network**

After screening all the small molecular compounds, we considered atractylosides as the target compounds. SwissTargetPrediction (21-23), a tool that can predict the potential target protein of small molecular compounds, was used to predict the binding target protein of atractyloside. It contains a library of 376,342 compounds known to be experimentally active on an extended set of 3,068 macromolecular targets. SwissTargetPrediction compares target compounds with 376,342 small molecules in the database. It uses a combination of 2D and 3D similarity measures for the comparison and predicts the binding target protein. ToppGene Suite database (24) was used to perform gene ontology enrichment analysis on the target protein of atractyloside. Thus, the biological function of the target protein was determined. First intersection was analyzed between the up-regulated genes in the high-intensity CAFs group and target proteins, while second intersection was analyzed with the enriched genes in the high-intensity CAFs group. Hence, the enrichment of up-regulated target proteins was achieved, and a signal regulating network was constructed with Cytoscape software. The network modules were calculated by using the plug-in MCODE in Cytoscope software (25-27). Thus, we determined the relationship between the target protein and the biological processes of the high-intensity CAFs group.

**CC cell (CCC) migration was inhibited by atractyloside**

To verify the effect of atractyloside on CCCs migration, two CCC lines (RKO and HCT116 purchased from the American Type Culture Collection) were selected to conduct the assay on cell scratch-wound healing. Cell culture medium and fetal bovine serum (FBS) were used for culturing the cells were purchased from Gibco Laboratories (Thermo Fisher Scientific, USA). The cell line was cultured in RPMI-1640 culture medium containing 10% FBS. The cell culture was maintained at 37 °C in a 5% CO₂ incubator.
Cells in good growing condition were removed and added into a six-well plate at 8×10^4 cells per well. After the growth of cells reached 90%, the tip of a 100 μL pipette was used to scratch the plate bottom, and the culture medium was replaced by atractyloside serum-free medium at a different concentration in each well to sequentially culture the cells. The concentrations of atractyloside were 0, 5, 25, 50, 75, and 100 μM. The six wells were photographed using an optical microscope at 0 and 24 h to record cell migration (at magnification of 10×).

The transwell cell-migration assay was conducted to verify the effect of atractylosides on CCC migration. Twenty percent of the culture medium was added to a 24-well plate at 650 μL per well. RKO/HCT116 cells cultured in serum-free medium for 24 h were extracted for 500 μL (with a density of 2×10^5 cells/mL) and centrifuged to collect sediment. To each of the tubes was added 200 μL atractyloside solution with a final concentration set to 0, 5, 25, 50, 75, and 100 μM, respectively. After mixing evenly, each of these solutions was added into the upper chamber of a transwell membrane. The control group contained an equal volume of PBS and three duplicate wells were set for each concentration. The membrane was immersed in a culture medium in a 24-well plate and incubated in a 5% CO_2 incubator for 24 h, after which it was fixed in 4% paraformaldehyde for 30 min and received Giemsa staining for 30 min. Finally, the membrane was rinsed three times with PBS. A cotton swab was used to remove the cells that had not passed through the upper layer of the membrane. The cells were placed on glass slides and three fields of view were randomly selected for counting the cells under an optical microscope and photographed at 20× magnification.

Confirming the effect of atractylosides on the microtubule cytoskeleton of CCCs

Changes in the cell microtubule cytoskeleton could affect the movement and morphology of cancer cells, which is closely related to tumor metastasis. Therefore, we observed the effect of atractylosides on the microtubule cytoskeletons of CCCs. Atractylosides were used to treat CCCs followed by immunofluorescence staining, and the distribution of α-tubulin in the cells was observed under a laser confocal microscope. The details were as follows: RKO/HCT116 cells were placed in a confocal dish (Wuxi NEST Biotechnology Co., Ltd., Jiangsu, China), treated with atractylosides at different concentrations (0.1% dimethyl sulfoxide and 0, 25, and 50 μM atractyloside) for 24 h, fixed with 4% paraformaldehyde, treated with 0.25% Triton-X, blocked with 5% bovine serum albumin, incubated with the first antibody (α-tubulin, dilution 1:1,000; ProteinTech America) in 4 ℃ overnight, washed three times with PBS for 5 min, incubated with the second antibody (Alexa Fluor 594-labeled goat anti-mouse immunoglobulin, dilution 1:100; Beijing Zhongshan Golden Bridge Biotechnology Co., Ltd., Beijing, China) for 30 min in the dark, washed with PBS and stained with 4′,6-diamidino-2-phenylindole (DAPI) for 5 min, washed again with PBS, and observed under a laser confocal microscope (at magnification of 120×).

The study was conducted in accordance with the Declaration of Helsinki (as is revised in 2013).

Statistical analyses

SPSS 19.0 (IBM Corp., Armonk, NY, USA) was used to analyze the data. Data between the two groups were analyzed using the Student’s t-test for independent samples. ANOVA was used to analyze the data for multiple comparisons.

Results

GSEA enrichment analysis of CAFs in CC

For the importance of CAFs in the TME and tumor immunity, we used the EPIC method to perform immune infiltration analysis on the CC expression profiles of 239 cases that were included in the dataset GSE2109. After obtaining the infiltration proportion of CAFs in each tumor sample, the data of 239 cases was classified into the high-intensity CAFs group and the low-intensity CAFs group. By performing GSEA enrichment analysis, we found that CC tissues of the high-intensity CAFs group were involved in various biological processes, such as extracellular structure organization, angiogenesis, ossification, and endothelial cell migration (Figure 2). The CC tissues of the high-intensity CAFs group were also included as cellular components in the composition of ECM, collagen, vesicle, endoplasmic reticulum, lysosome, the granule of platelets, and the lumen of secretory granules. In the high-intensity CAFs group, the CC tissues performed the molecular function of ECM structural constituent, binding with glycosaminoglycan, heparin, integrin, glycoprotein, cell adhesion molecule, cytokine, collagen, fibronectin, laminin, and proteoglycan. The above results indicate that the high-intensity CAFs group was related to the biological function of the TME, thereby confirming the role of CAFs in the development of...
Figure 2 GSEA enrichment results of the high-intensity CAFs group and the low-intensity CAFs group. CAFs, cancer-associated fibroblasts; GSEA, Gene Set Enrichment Analysis.
tumors. Thus, we also verified the accuracy of the results of immune infiltration analysis.

In the low-intensity CAFs group, the CC tissues were involved in the biological processes of nucleoside metabolism, cellular respiration, oxidative phosphorylation, and mitochondrial respiratory process. In the low-intensity CAFs group, the CC tissues were also included as cellular components in the composition of mitochondrial matrix, microbody, cytochrome complex, respiratory chain, and oxidoreductase complex. In the low-intensity CAFs group, the CC tissues performed the molecular function of nucleotide kinase activity, oxidoreductase activity, transferase activity, and coenzyme binding. The above results indicate that the low-intensity CAFs group was involved in cellular respiration and mitochondrial ATP synthesis. The early study (28) reported that in the metastatic stage of CC, up-regulated genes were related to ECM and inflammatory cell infiltration, and down-regulated genes were related to mitochondrial respiratory chain function. This indicated that the intensity of CAFs was closely related to the metastasis of CC. Hence, small molecular compounds that target CAFs in CC tissues can also effectively inhibit the metastasis of CC.

**Screening of small molecular compounds targeting CAFs**

By performing GSEA enrichment analysis, we obtained enriched genes that were related to various biological processes in the high-intensity CAFs group and the low-intensity CAFs group. After deleting duplicate genes, we collected 545 enriched genes from the high-intensity CAFs group and 325 enriched genes from the low-intensity CAFs group. To further verify the reliability of differentially expressed of enriched genes, we compared the expression data of the high-intensity CAFs group and the low-intensity CAFs group. Then, we screened the differentially expressed genes and obtained 1,572 genes up-regulated expression in the high-intensity CAFs group and 668 genes down-regulated expression in the low-intensity CAFs group. An intersection was analyzed between these differentially expressed genes and enriched genes, and we obtained 350 intersected genes in the high-intensity CAFs group and 78 intersected genes in the low-intensity CAFs group. Then, these genes were imported into the Connectivity Map. According to the hint of Connectivity Map, compounds with negative enrichment score could reverse the expression of differentially expressed genes, which correlated with anti-tumor CAFs in CC tissues. Therefore, compounds with negative enrichment score were selected with a P value of less than 0.01 and experiment was repeated more than 5 times. Finally, 25 small molecular compounds were obtained. After considering the enrichment score, P value, and the results of reported studies, we decided that atractylsides would be the target compounds in this study.

**Target protein prediction of atractylsides and network analysis**

Based on the intensity of CAFs in CC tissues, we obtained the small molecular compounds from the Connectivity Map. Hence, the biological function of the ideal small molecular compounds should be closely related to CAFs. The SwissTargetPrediction tool was used to predict the target protein that would bind with atractylsides. In this process, we screened 97 target proteins, including matrix metalloproteinase family proteins (MMP1, MMP2, MMP3, MMP8, MMP9, MMP10, MMP12, and MMP13), integrin family proteins (ITGB1, ITGA2B, ITGA4, ITGB5, ITGB7, ITGA5, and ITGAL) and caspase family proteins (CASP2, CASP1, CASP3, CASP7 and CASP8).

Gene ontology enrichment analysis was performed with ToppGene Suite; the enrichment analysis results indicated that target proteins were mainly involved in cell migration and cell apoptosis. Moreover, included as cellular components in the ECM, cell membrane, and some target proteins showed matrix metalloproteinase activity and fibronectin activity. An intersection was analyzed between 97 target proteins and 350 intersected genes in the high-intensity CAFs group. Thus, we identified 13 target proteins that had an up-regulated expression in the high-intensity CAFs group and were believed to be involved in the biological processes of CAFs. The 13 target proteins were as follows: FGF1, MMP13, MMP2, EDNRA, MMP9, PTPRC, ITGB1, ITGA1, ITGAV, ITGB5, CTSB, BMP1, and ITGA4. Obviously, these 13 target proteins belong to the following families: matrix metalloproteinase, integrin, endothelin receptor, fibroblast growth factor, bone morphogenetic protein, cathepsin, and protein tyrosine phosphatase receptor. Thus, they were closely related to the biological function of CAFs, which confirms the results of GSEA enrichment analysis.

A network was constructed from the 13 target proteins and their corresponding biological processes related with CAFs. Out of the 20 biological processes related to CAFs, we found that 16 were closely linked with the 13 target proteins (Figure 3). This indicated that the 13 target proteins...
proteins were mainly involved in the biological processes related to CAFs. Gene expression analysis showed that the expression of 13 genes in the high-intensity CAFs group was higher than that in the low-intensity CAFs group (Figure 4). Using the plug-in MCODE in Cytoscape software, we analyzed the network modules and obtained two subnetworks. In the subnetwork [1], we found that the target proteins FGF1, ITGB1, and EDNRA were closely related to angiogenesis. This indicated that atractylosides could inhibit the angiogenesis of CC tissues by binding with the three target proteins. The other subnetwork [2] revealed that the target proteins MMP9 and ITGAV were associated with the ECM and cell motility, which indicated that atractylosides could inhibit cell motility by binding with these two target proteins.

**The atractyloside inhibits CCC migration**

Because there is a close relationship between CAFs and cancer metastasis, our analyses suggested a relationship between the target protein of atractylosides and CCC migration. To examine the effect of atractylosides on CCC migration, transwell cell migration and scratch-wound healing assays were conducted on RKO and HCT116 cells using different atractyloside concentrations (0, 5, 25, 50, 75, and 100 μM). After the cells were cultured in different concentrations of atractyloside for 24 h, the number of CCCs that passed through the transwell membrane significantly decreased with increasing atractyloside concentration (Figure 5). We observed that inhibition of CCCs migration first appeared at an atractyloside
Figure 4 Gene expressions of 13 target proteins in the high-intensity CAFs group and the low-intensity CAFs group. CAFs, cancer-associated fibroblasts.

Figure 5 Effect of different concentrations of atractyloside on CCC migration ability. Magnification, ×20. CCC, colon cancer cell.
concentration of 5 μM, while significant inhibition was observed at a concentration of 50 μM. One-way ANOVA tests were conducted to compare the differences among the concentrations of RKO/HCT116 cells, and the results were statistically significant (P<0.001). Similarly, we observed using the scratch wound-healing assay that the healing speed distinctly decreased with an increase in concentration, which was significant at an atractyloside concentration of 50 μM too (Figure 6). By calculating the healing rate after 24 h [healing rate = (1 − area after healing)/area before healing], the healing rate obviously decreased with an increase in concentration. All of these results suggested that the atractyloside could effectively inhibit the migration of CCCs.

**Effects of atractylosides on the CC microtubule cytoskeleton**

Immunofluorescence staining and observation under the laser confocal microscope exhibited the effects of atractylosides at concentrations of 0, 25, and 50 μM on the microtubule cytoskeletons of RKO and HCT116 cells. We...
found that with an increasing concentration of atractyloside, the microtubule cytoskeletons located in the cytoplasm of the CCCs significantly changed. Compared with that in the cells of the control group, the microtubule cytoskeletons in the CCCs became sparse and disordered. In particular, at a concentration of 50 μM, obvious cell damage was observed and many nuclei were exposed (Figure 7). The above results suggested that atractylosides affected the microtubule cytoskeleton of CCCs and was closely related to the CCC migration.

**Discussion**

An atractyloside is a diterpenoid glycoside that occurs naturally in some plants, which are found in Europe, Africa, South America, Asia, and the Far East. The atractyloside was first isolated from the rhizomes of *Atractylis gummifera*. The atractyloside significantly inhibits mitochondrial oxidative phosphorylation and affects the metabolism of sugars and lipids (29). Massive necrosis can occur with high doses of atractylosides, while apoptosis can occur with low doses of atractylosides (30). Therefore, the atractyloside is used as an apoptosis inducer in animal model of tumor (31). The acquired drug resistance hampers the efficacy of chemotherapy and immunotherapy, which are appeared in the treatment of osteosarcoma (32,33) and malignant melanoma (34). The atractylosides can open mitochondrial permeability transition pore (MPTP), and it can enhance the cytotoxicity of TNF-related apoptosis-inducing ligand (TRAIL) in more than two apoptosis-resistant cells. Thus, it is beneficial to the treatment of cancer (35). In liver cancer patients, multidrug resistance (36) develops due to defects in apoptosis and it hampers the efficacy of chemotherapy. According to previous studies (37), the atractyloside acts as an MPTP opener and incubates with drug-resistance cells. Moreover, it reduces the IC50 in cis-diaminedichloroplatinum (CDDP) and enhances the anti-tumor effect. For the treatment of CC tissues, a combination of 5-fluorouracil and atractylosides may be used to promote the effect of chemotherapeutic agent and to inhibit the proliferation of tumor cells (38).

In this study, we found that most target proteins of
Atractyloside were related to CAFs, especially in the two subnetworks. These target proteins included FGF1, ITGB1, EDNRA, MMP9, and ITGAV. The migration and invasion of CC was closely linked to CAFs. In a previous study (39), it was reported that fibroblast growth factor 1/fibroblast growth factor receptor 3 (FGF1/FGFR-3) signaling were mediators that caused an increased migration and invasion of CC. In an in vitro study, it was reported that FGF1 was released by CAFs, and it acted on tumor epithelial cells to promote the migration of tumor cells. Hence, the inhibitor of FGF1 or a neutralizing antibody may inhibit the metastasis of CC. The target protein ITGB1 is a member of the integrin family. This protein is used for cell adhesion to ECM proteins. In large samples, the immunohistochemistry and the clinical pathology of CC showed that a high expression of ITGB1 was associated with lymph node metastasis and liver metastasis. Moreover, an increased expression of ITGB1 was related to poor prognosis of CC (40). The target protein EDNRA is an endothelin receptor. A previous study (41) analyzed the data of lymph node metastasis in bladder cancer patients. It also included immunohistochemistry results of large samples. In addition, it was found that EDNRA was highly expressed in bladder cancer patients with lymph node metastasis and distant metastasis. Meanwhile, a high expression of EDNRA was associated with decreased cancer-specific survival. MMPs have been extensively investigated to establish its relationship with cancer. Type IV and V collagen can be degraded by enzymes, which are encoded with MMP-9 gene. In CC, the target proteins MMP-2, MMP-7, and MMP-9 were up-regulated and co-expressed together with trypsin. Thus, they promoted the progression and invasion of CC (42). The target protein ITGAV is a member of the integrin family, and its expression is associated with tumor angiogenesis and metastasis. In CC, a high expression of ITGAV protein is correlated with neural invasion (43), indicating that target proteins of the atractyloside are closely related to the development and metastasis of tumor. This finding further confirms that the screening of small molecular compounds can be based on the intensity of CAFs.

In this study, by performing immune cell infiltration analysis, we determined the intensity of CAFs in the expression profile of CC. After converting the differentially expressed genes, we obtained the small molecular compounds of anti-CAFs, atractylosides. By analyzing the target proteins of the atractyloside, we further verified its association with CAFs. In addition, the transwell cell migration and scratch wound-healing assays proved that atractylosides could significantly inhibit CCC migration. Immunofluorescence staining and observation under a laser confocal microscope confirmed that atractylosides can affect the distribution of the microtubule cytoskeleton in CCCs. Thus, the atractyloside might be a small molecular compound that can potentially target CAFs and inhibit the development and metastasis of CC by changing the TME.

**Acknowledgments**

**Funding:** This work was supported by the National Key Basic Research Program of China (973 program, 2015CB554002), the National Natural Science Foundation of China (81972754,81773101), the Scientific Research Starting Foundation of Southern Medical University (PY2018N003).

**Footnote**

**Reporting Checklist:** The authors have completed the MDAR checklist (available at http://dx.doi.org/10.21037/atm-20-1531).

**Data Sharing Statement:** Available at http://dx.doi.org/10.21037/atm-20-1531

**Conflicts of Interest:** All authors have completed the ICMJE uniform disclosure form (available at http://dx.doi.org/10.21037/atm-20-1531). The authors have no conflicts of interest to declare.

**Ethical Statement:** The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. This study was conducted in accordance with the Declaration of Helsinki (as is revised in 2013).

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Cite this article as: Qi L, Song F, Han Y, Zhang Y, Ding Y. Atractyloside targets cancer-associated fibroblasts and inhibits the metastasis of colon cancer. Ann Transl Med 2020;8(21):1443. doi: 10.21037/atm-20-1531