Fibroblast growth factor-6 enhances \textit{CDK2} and \textit{MATK} expression in microvesicles derived from human stem cells extracted from exfoliated deciduous teeth [version 4; peer review: 1 approved, 1 approved with reservations, 1 not approved]

Previosuly titled: Fibroblast growth factor-6 enhances \textit{CDK2} and \textit{MATK} expression in microvesicles derived from human stem cells extracted from exfoliated deciduous teeth

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\textbf{Abstract}

\textbf{Background:} Stem cells from human exfoliated deciduous teeth (SHEDs) are considered one of the most convenient sources of adult stem cells. This study aimed to examine the effect of fibroblast growth factor 6 (FGF-6) on SHEDs and evaluate \textit{CDK2} and \textit{MATK} gene expression in SHED-derived microvesicles (MVs). SHEDs were cultured from deciduous teeth pulp.

\textbf{Methods:} SHEDs were divided into two groups: the control group and test groups, with and without FGF-6 supplementation, respectively. After the third passage, SHED proliferation was assessed by MTT assay. MVs were purified and \textit{CDK2} and \textit{MATK} gene expression was assessed by real-time polymerase chain reaction. SHEDs were identified by their positivity for CD90 and CD73, and negativity for CD45 and CD34.

\textbf{Results:} SHEDs proliferation in the test group was significantly higher than in the control group (P<0.001). mRNA from SHED-derived MVs from the test group exhibited a markedly elevated expression of \textit{CDK2} and \textit{MATK}, (P<0.002 and P<0.005, respectively) in comparison with those of the control group. FGF-6 enhanced the proliferation of SHEDs. Proliferation enhancement is favorable for the production of a large number of stem
cells, which will then be beneficial for cell-based therapies.

**Conclusions:** *CDK2* and *MATK* genes in SHED-derived MVs can be used as molecular biomarkers for SHED proliferation.

**Keywords**
Stem cells from human exfoliated deciduous teeth, fibroblast growth factor 6, microvesicles, CDK2, MAKT
Any reports and responses or comments on the article can be found at the end of the article.

Corresponding author: Ahmed Othman (ahmed.alsaid@dentistry.cu.edu.eg)

Author roles: Othman A: Writing – Original Draft Preparation, Writing – Review & Editing; Mubarak R: Supervision; Sabry D: Investigation, Methodology, Supervision, Validation

Competing interests: No competing interests were disclosed.

Grant information: The author(s) declared that no grants were involved in supporting this work.

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How to cite this article: Othman A, Mubarak R and Sabry D. Fibroblast growth factor-6 enhances CDK2 and MATK expression in microvesicles derived from human stem cells extracted from exfoliated deciduous teeth [version 4; peer review: 1 approved, 1 approved with reservations, 1 not approved] F1000Research 2020, 7:622 https://doi.org/10.12688/f1000research.14900.4

First published: 22 May 2018, 7:622 https://doi.org/10.12688/f1000research.14900.1
Amendments from Version 3

I have corresponded the aim of the study with the conclusion.
I have also amended the discussion part to be more focused on the main finding and mentioned previous work related to the study.
Unfortunately, there were no population gating images with our study. These are the only images of flow cytometry has given us.

Image 2 A contains 4 pictures: each one has two graphs the first one on the left is the isotype control of each CD.

Any further responses from the reviewers can be found at the end of the article.

Introduction

Stem cells from human exfoliated deciduous teeth (SHEDs) are a type of adult stem cell acquired from the dental pulp of human exfoliated deciduous teeth. SHEDs stand out from other types of adult stem cells since they possess a remarkable growth and proliferation rate, providing an adequate stem cell source for any prospective clinical or laboratory use. The natural exfoliation of deciduous teeth provides a good chance to procure and isolate SHEDs without effort or complications, and with little or no trauma1–3.

Fibroblast growth factors (FGFs) are a family of secreted cytokine proteins that have a role in the regulation and direction of numerous cellular processes, including proliferation, differentiation, migration or metabolism5. FGF-6 is part of the FGF-4 subfamily of canonical FGFs. Like the other members of FGF-4 subfamily, FGF-6 is a secreted protein with a cleavable N-terminal signal peptide that binds and activates FGF receptors as an extracellular mediator1. Despite the fact that FGF-6 expression is almost completely limited to myogenesis5, it was found to exert a proliferating effect on human osteoblasts under specific conditions6.

Recently, microvesicles (MVs) have been identified as an approach deployed by stem cells as a mean of mediating intercellular interactions7. These phospholipid membrane-bound MVs partake in intercellular interactions, such as proliferation, differentiation and gene expression alteration, via their content of mRNA, miRNA and transfer proteins8–9.

Human liver stem cell-derived MVs were found to have a role in hepatic regeneration, since they transfer proteins and mRNAs associated with the control of transcription, translation, proliferation, and apoptosis to hepatocytes8. Cyclin-dependent kinase 2 (CDK2) and megakaryocyte-associated tyrosine kinase (MATK) genes are associated with cellular proliferation as they were found in the mRNA of purified MVs at the time of microarray analysis and reverse transcription-quantitative PCR (RT-qPCR)9.

CDK2 is the catalytic subunit of the cyclin-dependent protein kinase complex, which controls advancement through the cell cycle via its involvement in the G1 to S phase transition10,11. MATK has been identified by Avraham et al. as an intracellular tyrosine kinase that participates in the proliferation and survival of megakaryocyte progenitors12. Furthermore, Findings by Herrera et al. demonstrated that MATK conveyed by MVs was one of the genes responsible of liver stem cell proliferation13.

The current study was performed to use SHEDs derived microvesicles as biomarker for cellular proliferation after FGF-6 supplementation by assessing the CDK2 and MATK gene expression in microvesicles’ mRNA.

Methods

Sample collection

A total of 28 deciduous teeth indicated for extraction were collected from 25 patients at the Pediatric Dentistry Department in Faculty of Dental Medicine, Cairo University. Patient age range from 7 to 12 years. Collection was done at the pediatric clinic over 3 days, we looked for deciduous teeth indicated for extraction due to their natural shedding time in order to make room for their permanent successors, so no ethical concerns would arise. Deciduous tooth collection was conducted after obtaining the guardians’ written informed consent at Pediatric Dentistry Department in the Faculty of Dental Medicine Cairo University, with the approval of the Ethics Committee of the Faculty of Oral and Dental Medicine, Cairo University. Subjects were identified by their treating physician, following which we contacted the guardians of the subjects for consent to use the extracted teeth. Stem cell propagation (at the Medical Biochemistry Department in the Faculty of Medicine Cairo University) was performed in accordance with recommendations and with the approval of the Ethics Committee of the Faculty of Oral and Dental Medicine, Cairo University.

Deciduous tooth surfaces were washed several times with Dulbecco’s PBS (Biowest, USA). Dental pulp was extracted delicately from teeth using a sterile endodontic barbed broach and placed in falcon tube containing PBS (Biowest, USA).

SHED culture and characterization

SHEDs culture and characterization were done after taking established procedures into account14. A total of 3 mg collagenase type II (Sigma Aldrich, USA) was dissolved in PBS to digest the extracted dental pulp tissues for 1 h at 37°C in a 5% CO2 incubator and shaken well at 10 min intervals until the tissues were fully digested. The samples were strained using a cell strainer (40 μm nylon PP) (Bio Basic, Inc., Canada) to remove tissue debris and then centrifuged for 10 min at 3000 rpm at 5°C to obtain pellets of isolated cells. The supernatant fluid was discarded and cell suspension was obtained by pipetting cells in RPMI 1640 culture medium (Biowest, USA). Next, the isolated cell pellets were seeded in 75 cm2 tissue culture flasks for cell culture propagation. Culture medium (RPMI 1640) (was supplemented with 1% Pen/Strep solution (Lonza, USA) and 10% fetal bovine serum (FBS) (Lonza, USA) were supplemented to the culture media to achieve cell propagation at 37°C in humidified CO2 incubator for 7–10 days, with medium changes every 3 days.

Cells were identified as being mesenchymal stem cells (MSCs) by their morphology and adherence to the plastic flask. In
addition, quantification of several expressed MSCs markers was conducted using flow cytometry analysis. Adherent cells were trypsinized and subjected to centrifugation to form cell pellet. Next, 1x10^6 cells were incubated with 10 µl monoclonal CD90 PE (catalog number FAB2067A; R&D Systems), CD73PE (catologue number FAB5795P; R&D systems) CD34 PE (catologue number FAB72271P; R&D Systems) and CD45 PE (catalog number DAB1430P; R&D Systems) antibodies, at 4°C in the dark. Same plate isotypes served as a negative control. Mouse IgG1 PE conjugated antibody (catalog number IC002P; R&D Systems). After a 20 min incubation, 2 ml PBS containing 2% FBS was added to a tube of monoclonal treated cells. The mixtures were then centrifuged for 5 min at 2500 rpm, followed by discarding the supernantant and re-suspending cells in 500 µl PBS containing 2% FBS. Cell analysis was performed using a CYTOMICS FC 500 Flow Cytometer and analyzed using CXP Software version 2.2.

SHEDs proliferation process and passaging
Passaging of SHEDs was done according to established protocols, with modifications for this experiment. Sub-culturing and passaging was done when adherent cells primary culture (passage zero) have reached 80% confluence. 10^3~10^4 cells were seeded into 24-well plates prior to grouping and subsequent passaging till reached third passages. Seeded cells were divided into two groups: control group (SHEDs untreated with FGF-6) and test group (SHEDs treated with FGF6). FGF-6 was added at concentration 20 ng/ml for test group.

Cell viability
MTT reagent, supplied ready for use after the third passage of the SHEDs, was obtained from Tacs Trevigen (Gaithersburg, USA). For the cell viability assay, the two cell groups were seeded in three 96-well tissue culture plates each, at 10^4 cells/ml per well. The MTT reagent was added and the plate was incubated in the dark for 2–4 h. Detergent reagent (catalog number # 4890-25-02, TACS) was added to each well to solubilize formazan dye prior to absorbance measurement. The absorbance in each well was measured at a range from 490 to 630 nm using an enzyme-linked immunosorbent assay plate reader (Stat Fax 2200, Awareness Technologies, Florida, USA).

MV isolation
MVs were obtained from supernatants of third-passage MSCs (5x10^6 cells/ml) cultured in RPMI-1640 deprived of FBS and supplemented with 0.5% of bovine serum albumin (BSA) (Sigma Aldrich, USA). After centrifugation at 2000g for 20 min to remove debris, cell-free supernatants were centrifuged at 100,000g for 1 h at 4°C, washed in serum-free medium 199 containing 25 mM HEPES (Sigma) and submitted to a second ultracentrifugation under the same conditions. MVs were then prepared for electron microscopy characterization. Briefly, MVs were diluted in 145 µL PBS containing 0.2% paraformaldehyde (w/v). 10 µl was administered to a formvar-carbon-coated 300 mesh grid (Electron Microscopy Sciences, Hatfield, USA) for 7 min, followed by staining with 1.75% uranyl acetate (w/v). Samples were left to dry at room temperature for 2 h and imaged by transmission electron microscopy (TEM) (CM-10, Philips, Eindhoven, The Netherlands) at 100 kV afterwards.

Gene expression profile
Total RNA was isolated from MVs using an RNA purification kit (Gene JET, Kit, #K0731, Thermo Fisher Scientific, Inc.). RNA quantification using RT-qPCR was achieved using a one-step reaction (SensiFAST™ SYBR® Hi-ROX One-Step Kit, catalog no. PI-50217 V; Bioline, UK). Sequence-specific primers (Bio Basic, USA) for the studied target genes (CDK2 and MATK) and reference housekeeping gene (β-actin) were used. The prepared reaction mix samples were applied in real time PCR (StepOne Applied Biosystem, Foster city, USA). The cDNA was subsequently amplified using a SYBRGreen I PCR Master kit (Fermentas) in a 48-well plate as follows: 10 min at 95°C for enzyme activation, followed by 40 cycles of 15 s at 95°C, 20 s at 60°C and 30 s at 72°C for the amplification step. Changes in the expression of each target were normalized relative to the mean Cq values of β-actin as housekeeping gene by the 2^{-\Delta\Delta Cq} method. We used 1 µM of both primers specific for each target gene. Primers sequences were as follows: CDK2 sense, 5'-AATCCGCCCTGGACACTGAGA-3' and antisense, 5'-CCAGCAGCTTGACAATATTAGGA-3' (Genbank accession number XM011537321); MATK sense, 5'-CCCGGACGCTCATCCACTAC-3' and antisense, 5'-TTGTAATGCTCCACCATGTCCAT-3' (Genbank accession number AH006874); and GAPDH sense, 5'-GGGCGGACCACCATGTACCT-3' and antisense, 5'-AGGGGCCGACTCGTACA-3' (Genbank accession number AAG01996).

Statistical analysis
Data were coded and entered using SPSS version 23. Data are presented as the median and interquartile range for quantitative data. Comparisons between quantitative variables were done using the non-parametric Mann-Whitney test. Correlations between quantitative variables were done using Spearman’s correlation coefficient. P-values less than 0.05 were considered as statistically significant.

Results
SHED characterization
Cultured SHEDs exhibited fusiform fibroblast like appearance for both groups. During culture and passaging, SHEDs in the test group proliferated more than SHEDs in the control group (Figure 1). Flow cytometric analysis for SHEDs was negative for CD34 and CD45 and positive for CD90 and CD73 (Figure 2A).

Cell viability
The viability of the cells in the test group was significantly higher (P<0.001) in comparison with that of the control group (Table 1).

TEM
TEM detected MVs purified from SHED after ultracentrifugation (Figure 2B). MVs were characterized by their size (500 nm), as detected by TEM.

RT-qPCR
Purified MVs demonstrated a significant positive expression intensity of CDK2 (P=0.002) and MATK (P=0.005) in the test
Figure 1. Isolation, morphological observation of stem cells from human exfoliated deciduous teeth through phase contrast microscopy. (A) Passage one shows stem cells with spindle-like morphology as grow from human exfoliated deciduous teeth pulp in few number. (B and C) Passages two (B) and three (C) show an increase in number of stem cells with spindle-like morphology. Isolation, morphological observation of stem cells from human exfoliated deciduous teeth in the test group through phase contrast microscopy. (D) Passage one shows a marked increase in number and confluency of stem cells with spindle-like morphology in comparison with control group in passage one. (E and F) Passages two (E) and three (F) show a pronounced, confluent and expanded SHED with fibroblast like morphology in relation to control groups of second and third passages.

Figure 2. Flow cytometry and transmission electron microscopy. (A) Flow cytometry analysis of CD90, CD73, CD34 and CD45 for stem cell characterization. (B) Electron microscopy ultrastructure of released microvesicles (black arrow) from the mesenchymal stem cells of dental pulp.
group in relation with the control group. A box plot (Figure 3) shows that expression of CDK2 and MATK is higher in the test group than the control group, as they display a higher interquartile range (IQR) and median. The correlation coefficients of CDK2 and MATK expression to cell viability were positive for both genes in the test group (Figure 4).

**Table 1. Cell proliferation assay for both studied groups.** Data presented as median (IQR).

<table>
<thead>
<tr>
<th>Variable</th>
<th>SHEDs</th>
<th>SHEDs supplemented with FGF-6</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Absorbance (450 nm)</td>
<td>0.90 (0.77-1.36)</td>
<td>2.55 (1.63-2.98)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

SHEDs, stem cells from human exfoliated deciduous teeth; FGF-6, fibroblast growth factor-6.

![Figure 3](image1.png)  
**Figure 3.** Box-and-whiskers plot showing number distribution for CDK2 and MATK expression in both test and control groups. (A and B) Expression of each gene is higher in the FGF-6-supplemented group than in the control group, since higher interquartile range (IQR) and median values are observed.

![Figure 4](image2.png)  
**Figure 4.** Assessment of cell proliferation. (A) Expression of CDK2 is positively correlated with cell proliferation in the test group. (B) Expression of MATK is positively correlated with cell proliferation in the test group.
Discussion
We performed this experiment to deal with difficulties sourcing stem cells and a lack of sufficient stem cells for reliable tissue formation. The study focused on stem cells isolated from human deciduous teeth (SHEDs) and tissue-inducing substances, which in this case is FGF-6. The reason we chose SHEDs for the isolated stem cells is that they present an opportune source of adult stem cells; the deciduous teeth are naturally exfoliating so there are no ethical problems surrounding their use, and the isolation of cells is simple, painless, convenient and time-efficient.\(^1\)

There are several criteria for SHED identification; we identified stem cells by their morphology under the inverted microscope, as they appeared as fibroblast-like cells. Another feature exhibited was that they have a plastic adherence feature under our normal culture conditions.\(^2\) SHEDs were also identified as ectomesenchymal stem cells through the quantification of several expressed mesenchymal stem cell markers using flow cytometry; they were shown to be positive for CD90 and CD73, and negative for CD45 and CD34.\(^3\)

In this case, the stem cells are quiescent, unlike progenitor cells, meaning growth factor treatment is required to produce a large amount of cells. In our experiment, FGF-6 was chosen as it has, to our knowledge, never used on SHEDs before, and would avoid the conflicting reports of the effects of bFGF on SHEDs.\(^4,5\)

SHEDs group with added FGF-6 demonstrated increased cells vitality and number in comparison with the control group. It was evidenced by the MTT assay results and increased vitality and number in comparison with the control group. SHEDs with added FGF-6 demonstrated increased cells vitality and number in comparison with the control group. It was evidenced by the MTT assay results and increased vitality and number in comparison with the control group.

Megakaryocyte-associated tyrosine kinase is the enzyme which is encoded by $MATK$ in humans. This enzyme possesses a similar amino acid sequence to tyrosine-protein kinase CSK. It was chosen for our experiment as it is not frequently used for the assessment of SHED proliferation, to evaluate whether this ambiguous gene can be studied in further research to assess the proliferation rate of SHEDs and other types of stem cells it is known to be capable of phosphorylating and inactivating Src family kinases, and may inhibit T-cell proliferation.\(^6\)

Conclusion
The present study showed an increased expression of $CDK2$ and $MATK$ genes present in RNA of microvesicles we purified from SHEDs after FGF-6 supplementation. Thus, MVs derived from SHEDs can be used as a biomarker for cellular proliferation. We recommend further experimentation of differentiation potential of these microvesicles and compared them to the SHEDs themselves before and after growth factor application, utilize and compare different isolation protocols for microvesicles, and test more cargo genes and CDs.

Data availability

**Dataset 1.** Raw data for the MTT cell viability assay and for reverse transcription-quantitative PCR. DOI: https://doi.org/10.6084/m9.figshare.11666460.v1\(^7\).

**Underlying data**

**References**


Abbas Ali Khademi  
Department of Endodontics, School of Dentistry, Isfahan University of Medical Sciences, Isfahan, Iran

This study is about evaluation of the effect of FGF-6 on SHEDs and assessment of CDK2 and MATK gene expression, in which the authors reported that CDK2 and MATK genes in SHED-derived can be used as molecular biomarkers for proliferation.

1. Add mean and SD of patients and add female to male ratio or male to female ratio.

2. Please provide approval code or public link.

3. For characterization only flow cytometry is not enough, RT-PCR on genes that show mesenchymal stem cells have been isolated would be interested, e.g. STRO1, NANOG, SOX family, OCT4. Or differentiation to osteoblast, chondroblast and adipocytes.

4. Please add a paragraph that show how your results would be helpful to clinical research.

5. Increase the quality of figures.

6. In Figure 3 for CDK2 there are two outliers data.

7. Figure 4 can be removed. Authors can report the correlation as follows in the results section: Expression of CDK2 is positively correlated with cell proliferation in the test group (r=?, P_value:?)

Is the work clearly and accurately presented and does it cite the current literature?  
Yes

Is the study design appropriate and is the work technically sound?  
Yes

Are sufficient details of methods and analysis provided to allow replication by others?  
Yes
If applicable, is the statistical analysis and its interpretation appropriate? 
Yes

Are all the source data underlying the results available to ensure full reproducibility? 
Yes

Are the conclusions drawn adequately supported by the results? 
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Endodontics.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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Author Response 21 May 2020

Ahmed Othman, Cairo University, Cairo, Egypt

Dear Dr. Abbas Ali Khademi,

I would like to thank you very much for your review. I truly appreciate your time and effort. We did the following modifications to the article as requested,

The Recommendation section was created to state any future experiment carried out should include a multipotential differentiation test and more genes testing to correctly identify adult stem cells to avoid confusion with dental pulp fibroblasts.
Additionally, a paragraph about potential use in clinical research has been added.
In the correlation section, we added (r) and (P) values.
We added sample size number (n) in the cells viability section.
We rectified the mistake of adding the GAPDH sequence instead of a beta-actin sequence, by adding beta-actin sequence and accession number.
We also recommend in the future to take into consideration male to female ratio and standard deviation during sample collection.
We tried to increase the quality of the figures but unfortunately, we could not, because of the status of equipment during the experiment. In future experiments, we will certainly take that into consideration.
The ethics approval has been provided to F1000Research editors before publishing.

Unfortunately, the funding for the experiment was very limited, so I could not do the multipotential differentiation test or test the presence of STRO1, NANOG, SOX family, OCT4 genes. In the future, we will apply for a much larger fund to conduct the experiments more properly.

My best regards

**Competing Interests:** No competing interest
The present article described the effect of FGF6 on RNA contents of microvesicles. In addition, the correlation of detected RNA on cell proliferation was described. There are many issues required to address in order to improve scientific content of this article.

1. According to Dominici et al. (2006)\(^1\), mesenchymal stem cells can be characterized by 1) plastic adherence properties, 2) surface marker expression, and 3) multipotential differentiation ability. This article lacks multipotential differentiation test. Hence, the claim of these cells as stem cells isolated from human exfoliated deciduous teeth was not accurate. Indeed, these cells may be dental pulp fibroblast cells.

2. Authors did not mention which time points that they evaluated for cell viability and proliferation. To determine cell proliferation, at least 2 time points are required and the second time points should be longer than the normal doubling time of these cells.

3. Please provide more detail on those reagent used in the study. The catalog number as well as company is crucial information for others to replicate these finding. The main reagents (for example FGF6) were lacking this information.

4. How many cells were seeded in the tissue culture plates and treated with FGF6 for further cell viability test and MV isolation? Authors mentioned \(10^3\)-\(10^5\) cells and this is a wide range. Different cell density could reflect the different results obtained.

5. In PCR, authors indicated the use of beta-actin as reference gene but provided GAPDH sequence.

6. In TEM results, it seems that there were a lot of contaminated substance in MV collection. Would this effect the evaluation of gene expression since the RNA from the cells might also contaminated with MVs RNA?

7. In PCR results, it seems that there was no statistically significant difference. This might be due to the different baseline expression among donors. Plots demonstrated the individual data could help to visualize the trend between control and treatment in each donors.

8. What is the \(r\) for correlation analysis? Please indicate the statistical number.

9. Please indicate \(n\) in each experiment.
10. Overall, the results seems preliminary and required further improvement to strongly support the conclusion claimed by authors.

References

Is the work clearly and accurately presented and does it cite the current literature?
No

Is the study design appropriate and is the work technically sound?
No

Are sufficient details of methods and analysis provided to allow replication by others?
No

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
No

Are the conclusions drawn adequately supported by the results?
No

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Dental stem cells.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to state that I do not consider it to be of an acceptable scientific standard, for reasons outlined above.

Author Response 21 May 2020

Ahmed Othman, Cairo University, Cairo, Egypt

Dear Dr. Thanaphum Osathanon,

I would like to thank you very much for your review. I truly appreciate your time and effort. We did the following modifications to the article as requested,

The Recommendation section was created to state any future experiment carried out should include a multipotential differentiation test and more genes testing to correctly identify adult stem cells to avoid confusion with dental pulp fibroblasts.
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Unfortunately, the funding for the experiment was very limited, so I could not do the multipotential differentiation test or test the presence of STRO1, NANOG, SOX family, OCT4 genes. In the future, we will apply for a much larger fund to conduct the experiments more properly. We contacted the lab to query about FGF-6, but unfortunately, they are unreachable because of lockdown in the university. As soon as I know the manufacturer and catalog number, I will update the article.

My best regards

**Competing Interests:** no competing interest

Reviewer Report 04 February 2020

https://doi.org/10.5256/f1000research.24301.r59229

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Asma Abdullah Nurul 🌐
School of Health Sciences, University of Science Malaysia, Kubang Kerian, Malaysia

The authors have responded to the comments given in the previous review.

Is the work clearly and accurately presented and does it cite the current literature?
No

Is the study design appropriate and is the work technically sound?
No

Are sufficient details of methods and analysis provided to allow replication by others?
No

If applicable, is the statistical analysis and its interpretation appropriate?
No

Are all the source data underlying the results available to ensure full reproducibility?
No
Are the conclusions drawn adequately supported by the results?
No

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Stem cell, regenerative medicine, molecular immunology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 3

Reviewer Report 18 September 2019

https://doi.org/10.5256/f1000research.21423.r52495

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Asma Abdullah Nurul
School of Health Sciences, University of Science Malaysia, Kubang Kerian, Malaysia

- In the revision 3, the authors have amended some part of the manuscript, but ignored some of the reviewer's comments.
- The conclusion in the abstract is not corresponded to the main aim of the study.
- Why the authors did not include mRNA expression of CDK2 and MATK from SHED or treated SHED, instead of SHED-derived MVs alone.
- Discussion sounds more like rationale of study, not discussing about the main findings and relate with previous works.
- The current write up is not so impressive, so I recommend re-writing of the manuscript.

Is the work clearly and accurately presented and does it cite the current literature?
No

Is the study design appropriate and is the work technically sound?
No

Are sufficient details of methods and analysis provided to allow replication by others?
No

If applicable, is the statistical analysis and its interpretation appropriate?
Are all the source data underlying the results available to ensure full reproducibility?
No

Are the conclusions drawn adequately supported by the results?
No

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Stem cell, regenerative medicine, molecular immunology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to state that I do not consider it to be of an acceptable scientific standard, for reasons outlined above.

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**Reviewer Report 16 July 2018**

https://doi.org/10.5256/f1000research.16430.r35756

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Asma Abdullah Nurul
School of Health Sciences, University of Science Malaysia, Kubang Kerian, Malaysia

The authors have presented a work on the effects of fibroblast growth factor-6 on the expression of CDK2 and MATK in microvesicles derived from stem cells from exfoliated deciduous teeth (SHED). In the study, the authors have shown that treatment with FBF-6 enhances the proliferative activity as well as the expressions of CDK2 and MATK in MV derived from SHED. This study is somewhat preliminary and the manuscript requires modifications to further improve it.

1. There is inconsistency in the usage of terminology for example MATK was misspelled with MAKT many times (also in the title).

2. Figure 2 should include population gating and isotype control.

3. The authors should provide r value of the correlation coefficients of the CDK2 and MATK expressions to cell proliferative activity as shown in Figure 4.

4. Conclusion need to be rephrased.

5. The language use in the manuscript is unsuitable for publication. I would like to suggest submitting the manuscript for professional English editing.
Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Partly

Are all the source data underlying the results available to ensure full reproducibility?
Partly

Are the conclusions drawn adequately supported by the results?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: stem cell, regenerative medicine, molecular immunology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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