Human umbilical cord blood-mesenchymal stem cell-derived secretome in combination with atorvastatin enhances endothelial progenitor cells proliferation and migration
[version 1; peer review: 2 approved with reservations]

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Abstract

Background: Human umbilical cord blood-mesenchymal stem cell (hUCB-MSC)-derived secretome is known to be able to promote neovascularization and angiogenesis, so it is also thought to have a capability to modulate endothelial progenitor cell (EPC) functions. Atorvastatin is the cornerstone of coronary artery disease (CAD) treatment which can enhance EPCs proliferation and migration. This study aims to analyze the effect of the hUCB-MSC-derived secretome and its combination with atorvastatin toward EPCs proliferation and migration.

Methods: EPCs were isolated from a CAD patient’s peripheral blood. Cultured EPCs were divided into a control group and treatment group of 2.5 µM atorvastatin, hUCB-MSC-derived secretome (2%, 10%, and 20% concentration) and its combination. EPCs proliferation was evaluated using an MTT cell proliferation assay, and EPC migration was evaluated using a Transwell migration assay kit.

Results: This research showed that hUCB-MSC-derived secretomes significantly increase EPC proliferation and migration in a dose-dependent manner. The high concentration of hUCB-MSC-derived secretome were shown to be superior to atorvastatin in inducing EPC proliferation and migration (p<0.001). A combination of the hUCB-MSC-derived secretome and atorvastatin shown to improve EPCs proliferation and migration compared to hUCB-MSC-derived secretome treatment or atorvastatin alone (p<0.001).

Conclusions: This study concluded that the hUCB-MSC-derived secretome work synergistically with atorvastatin treatment in
improving EPCs proliferation and migration.

**Keywords**
coronary artery disease, endothelial progenitor cells, mesenchymal stem cells, secretome, statins
Introduction
Coronary artery disease (CAD) is the leading cause of mortality and morbidity worldwide. It is responsible for the deaths of 7.2 million people or 12.2% of total deaths per year worldwide. Despite advancement in CAD management (e.g. novel antiplatelet therapy, coronary stents, percutaneous coronary intervention techniques and devices, and coronary artery bypass surgery), there are some clinical subsets of CAD which remain untreatable such as ischemic cardiomyopathy, refractory angina, and patient which cannot undergo revascularization due to clinical and anatomical complexity.

It is already known that CAD is caused by atherosclerosis, which is followed by reduced levels of circulating endothelial progenitor cells (EPCs). EPCs can differentiate into mature endothelial cells and also promote endothelial repair. Hence, increasing circulating EPC levels is proven to improve endothelial function. EPCs also had a critical role in the stimulation of angiogenesis and vasculogenesis. Hence, increasing EPC proliferation and migration may reduce ischemia and improve myocardial performance.

Regenerative treatment for CAD using stem cells has been extensively studied in the last decade. However, these treatments faced challenges of low engraftment, poor survival, and low differentiation of the transplanted cells. Despite regenerative treatment shown to be promising in vitro, clinical studies showed unsatisfying results. Hence, the researcher started to shift regenerative treatment from cell-based treatment to cell-free treatment using paracrine stimulation. Nowadays, the usage of cell-free therapeutics as a regenerative therapy in cardiovascular diseases also started to be emerged.

The secretome is the wide array variety of paracrine factors produced by mesenchymal stem cells (MSCs). Human umbilical cord blood mesenchymal stem cells (hUCB-MSCs) derived secretome was proven could promote neovascularization, angiogenesis, and improved cardiac systolic function by protecting myocardial cells from apoptosis. However, using this approach to improve neovascularization is yet to be investigated. Hence, it is hypothesized that increasing EPCs proliferation and migration by the hUCB-MSC-derived secretome may be responsible for this effect. Statins, through their pleiotropic effect, are the cornerstone of CAD treatment. Atorvastatin is one of the most prescribed statins, whose ability to modulate EPCs proliferation and migration has already been well studied in both laboratory and clinical settings. Furthermore, this study aims to compare the effect of the hUCB-MSC-derived secretome, atorvastatin and the two in combination in modulating EPC proliferation and migration.

Methods
Sample collection
A 50–100 mL peripheral blood sample was obtained from a patient with CAD. The patient was recruited from the outpatient cardiovascular clinic at Pusat Pelayanan Jantung Terpadu, Dr. Soetomo General Hospital, Surabaya, in March 2020. The inclusion criteria were as follows: male, aged 40–59 years old, history of chronic ischemic heart disease as proven by coronary angiography results that showed >50% stenosis of left main coronary artery or >70% of other coronary arteries. The exclusion criteria were as follows: a history of percutaneous coronary intervention procedures or coronary artery bypass grafting surgery, acute coronary syndromes, and anemia.

This study protocol has an ethical clearance from the Health Research Ethics Committee of Dr. Soetomo General Hospital, Surabaya (No.1567/KEPK/X/2019, approved on 8 October 2019). The included subjects provided written informed consent before subject recruitment. All details which include personal information were omitted.

HUCB-MSCs-derived secretome preparation
The HUCB-MSCs-derived secretome was prepared according to the previous study. HUCB-MSCs (3H Biomedical AB, Uppsala, Sweden) was cultured in MesenCult™ MSC Basal medium, supplemented with MesenCult™ Stimulatory supplement (StemCell Technologies Inc., Vancouver, Canada), and also added with penicillin and streptomycin. Upon reaching 80% confluency, the media was replaced with MesenCult™ MSC Basal medium (supplement-free media) and incubated for 24 hours. The media was collected and centrifuged. The supernatant was used as a conditioned medium that contained hUCB-MSCs-derived secretome.

Isolation and culture of EPCs
Peripheral blood mononuclear cells (PBMCs) were isolated by density centrifugation of CAD patient’s peripheral blood using Histopaque-1077 (Sigma-Aldrich, USA). After centrifugation of peripheral blood, PBMCs then cultured with STEMLINE-II hematopoietic stem cell expansion medium (Sigma-Aldrich, USA) supplemented with stem cell factor, thrombopoietin, Flt-3 ligand, vascular endothelial growth factor, and interleukin-6. A total of 5×10^6 mononuclear cells/ml were seeded into fibronectin-coated 6-well plate dish and cultured at 37°C and 5% CO2, levels for 5 days. Non-adherent cells were then transferred for the proliferation and migration assay. After five days of culture, EPCs were confirmed using FITC-labeled anti-human CD34 antibody (animal source was mouse, 5 μL antibody was diluted at 500 μL per 1 × 10^6 cells; catalog number 60013FI, Gene ID: 947, by StemCell Technologies Inc., Vancouver, Canada) staining and examined with immunofluorescence microscopy.

Treatment of EPCs
Cultured EPCs were divided into eight treatment groups for each proliferation and migration assays. Those treatment include control group, 2.5 μM atorvastatin, low (2%), medium (10%) and high (20%) doses of hUCB-MSC-derived secretome, and combination of 2.5 μM atorvastatin with each dose of the hUCB-MSC-derived secretome. There were n=5 replications made from each treatment.

EPCs proliferation assay
The MTT cell proliferation assay kit (Sigma-Aldrich, St Louis, MO, USA) was used to measure EPCs proliferation as described previously. Treated EPCs were added with MTT reagent
and incubated in a 37°C incubator with 5% CO₂ for 4 hours. Proliferation was determined from the reduction of tetrazolium (MTT) into insoluble formazan product by viable EPCs mitochondria. Absorbance was measured with a microplate reader at 595 nm wavelength. EPCs proliferation was measured as optical density (OD). MTT assay was measured at day 3 after reagent addition.

**EPCs migration assay**

EPCs migration was evaluated using the 24-mm diameter insert, 3-μm pore size, 6-well Transwell migration assay kit (Corning, USA). A total of 5x10⁵ cultured EPCs were placed in the upper part of the Transwell migration assay kit. Next, 2 mL of EPC media and each treatment were added in the lower chamber compartment and then incubated for 24 hours at 37°C. Non-migratory cells were removed manually. On the receiver plate, the new basal medium was placed and added 500 μL of trypsin + EDTA solution 0.5%, followed by 10 minutes incubation. Then, cells on the bottom surface of the membrane were stained with Giemsa and cell images were obtained on a light microscope and counted manually in n=5 random fields/sample.

**Statistical analysis**

Statistical analyses were conducted using SPSS Statistics 23.0 to detect significance level at p<0.05. One-way ANOVA was used to compare groups, with Fisher’s least significant difference (LSD) post hoc test. Kruskal-Wallis test was used if there are violations to the assumption of normality and the assumption of homogeneity of variance. Correlation between variables was obtained using Spearman’s correlation followed by a linear regression test.

**Results**

Baseline characteristics and demography of CAD patient

Clinical examination, blood sampling, electrocardiography, echocardiography and coronary angiography was conducted and evaluated in order to examine the inclusion and exclusion criteria. Our sample had a 1-year history of coronary artery disease, he suffered from refractory chest pain despite the optimum medical therapy. The coronary angiography showed complex lesion (three-vessel disease with chronic total occlusion) which was not amenable to undergo revascularization. The baseline characteristics of the patient are presented in **Table 1**.

<table>
<thead>
<tr>
<th>Variables</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex</td>
<td>Male</td>
</tr>
<tr>
<td>Age</td>
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</tr>
<tr>
<td>Body Mass Index (BMI)</td>
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<td>Blood pressure</td>
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<td>Heart rate</td>
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<td>Sinus rhythm, pathological Q-waves at V-V6 leads.</td>
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<td>Laboratory</td>
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<tr>
<td>Triglyceride (mg/dL)</td>
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<td>LDL (mg/dL)</td>
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<td>HDL (mg/dL)</td>
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</tr>
<tr>
<td>Hemoglobin (mg/dL)</td>
<td>14.2</td>
</tr>
<tr>
<td>Blood glucose (mg/dL)</td>
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</tr>
<tr>
<td>Echocardiography</td>
<td></td>
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<td>Left ventricle ejection fraction</td>
<td>41% (teich); 36% (biplane)</td>
</tr>
<tr>
<td>Left ventricle internal diameter</td>
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</tr>
<tr>
<td>Wall motion</td>
<td>Hypokinesia at anterior, anteroseptal, inferoseptal, other segments kinetic was normal</td>
</tr>
<tr>
<td>Valves</td>
<td>mild mitral regurgitation</td>
</tr>
<tr>
<td>Coronary Angiography</td>
<td></td>
</tr>
<tr>
<td>Left main coronary artery (LMCA)</td>
<td>normal</td>
</tr>
<tr>
<td>Left anterior descending artery (LAD)</td>
<td>70% stenosis at ostegal, chronic total occlusion (CTO) at distal</td>
</tr>
<tr>
<td>Left circumflex coronary artery (LCX)</td>
<td>70% stenosis at proximal, CTO at distal, grade 2 collaterals from LCX to RCA</td>
</tr>
<tr>
<td>Right coronary artery (RCA)</td>
<td>CTO at proximal</td>
</tr>
</tbody>
</table>
**EPC characteristics**

EPCs were successfully isolated and cultured from the CAD patient’s peripheral blood. It was confirmed through light microscopy and an immunofluorescence assay (Figure 1). Raw images are available as Underlying data.

**HUCB-MSCs-derived secretome and atorvastatin increase EPCs proliferation**

EPCs were evaluated using the MTT proliferation assay. As shown in Figure 2, both atorvastatin and hUCB-MSCs-derived secretome treatment groups at all doses increase EPCs proliferation compared to the control (p<0.05, ANOVA). hUCB-MSC-derived secretome treatment showed a dose-dependent relationship with EPCs proliferation. At medium (10%) and high (20%) doses, hUCB-MSC-derived secretome was shown to elicit superior EPC proliferation than atorvastatin (OD 1.252±0.104 and 1.585±0.029, respectively, vs 0.738±0.025; p<0.01). Raw absorbance data for MTT assays are available as Underlying data.

Pearson’s correlation showed a significant and strong correlation between hUCB-MSCs-derived secretome treatment with EPC proliferation (r=0.954; p<0.001). The linear regression test showed an R² of 0.910.

**Combination of hUCB-MSCs-derived secretome and atorvastatin increase EPCs proliferation compared with single treatment**

Figure 2 showed the combination of atorvastatin and hUCB-MSC-derived secretome at the dose of 2%, 10% and 20% concentration have significantly higher EPCs proliferation compared to atorvastatin alone (OD 0.803±0.046, 1.298±0.075 and 1.761±0.419 vs 0.738±0.025, p<0.05). In addition, combination of hUCB-MSC-derived secretome at dose of 2%, 10% and 20% with atorvastatin showed higher EPCs proliferation compared to hUCB-MSC-derived secretome alone (OD 0.803±0.046 vs 0.713±0.049, 1.298±0.075 vs 1.252±0.104 and 1.761±0.419 vs 1.585±0.029, p<0.05). The combination group showed a significant and very strong correlation with EPC proliferation (r=0.973; p<0.001). Linear regression test showed R² of 0.947.

**HUCB-MSCs-derived secretome and atorvastatin increase EPCs migration**

EPCs migration from each treatment group was analyzed using the Transwell migration assay. As shown in Figure 3, EPC treatment with atorvastatin and all doses of hUCB-MSCs-derived secretome significantly increase EPC migration compared to the control group (p<0.05, ANOVA). Treatment with 2.5 μM atorvastatin has significantly higher EPCs migration than low (2%) and medium (10%) doses of hUCB-MSC-derived secretome (34.40±3.05 vs 17.20±1.92 and 27.00±4.00, p<0.05). However, high doses (20%) of hUCB-MSC-derived secretome showed significantly higher migrated EPCs than atorvastatin (51.00±5.15 vs 34.40±3.05, p<0.001). Raw cell counts used to assess migration are available as Underlying data.

Pearson’s correlation showed a significant and very strong correlation between hUCB-MSCs-derived secretome treatment with EPC migration (r=0.968; p<0.001). The linear regression test showed an R² of 0.937.

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**Figure 1. Immunofluorescence characterization of cultured EPCs.** (A) DAPI staining of cultured EPCs showed viable cells through blue fluorescent of cells nuclei. (B) EPCs were confirmed, using FITC-labeled anti-human CD34 expression on immunofluorescence microscope. (C) Merged view of DAPI and FITC stained cells. (D) The light microscope view showed the spindle shape of EPCs. The white bar represents 50 μm.
Figure 2. Comparison of EPC proliferation effects among all treatment groups (see text).

- Significant difference compared to the control group (p < 0.001).
- Significant difference compared to the 2.5 µM atorvastatin group (p < 0.001).
- Significant difference compared to the 2% hUCB-MSC-derived secretome group (p < 0.001).
- Significant difference compared to the 10% hUCB-MSC-derived secretome group (p < 0.001).
- Significant difference compared to the 20% hUCB-MSC-derived secretome group (p < 0.001).
- Significant difference compared to the combination of 2% hUCB-MSC-derived secretome and atorvastatin group (p < 0.001).
- Significant difference compared to the combination of 10% hUCB-MSC-derived secretome and atorvastatin group (p < 0.001).
- Significant difference compared to the combination of 20% hUCB-MSC-derived secretome and atorvastatin group (p < 0.001).

Figure 3. Comparison of EPCs migration effects among all treatment groups (see text).

- Significant difference compared to the control group (p < 0.001).
- Significant difference compared to the 2.5 µM atorvastatin group (p < 0.001).
- Significant difference compared to the 2% hUCB-MSC-derived secretome group (p < 0.001).
- Significant difference compared to the 10% hUCB-MSC-derived secretome group (p < 0.001).
- Significant difference compared to the 20% hUCB-MSC-derived secretome group (p < 0.001).
- Significant difference compared to the combination of 2% hUCB-MSC-derived secretome and atorvastatin group (p < 0.001).
- Significant difference compared to the combination of 10% hUCB-MSC-derived secretome and atorvastatin group (p < 0.001).
- Significant difference compared to the combination of 20% hUCB-MSC-derived secretome and atorvastatin group (p < 0.001).
Combination of hUCB-MSCs-derived secretome and atorvastatin increase EPCs migration compared with single treatment

In Figure 3, EPCs migration was significantly higher in combination treatment groups (atorvastatin and hUCB-MSC-derived secretome) at 2%, 10%, and 20% doses compared to the atorvastatin alone (38.20±3.49, 50.20±5.31 and 76.40±7.50 vs 34.40±3.05, p<0.001).

Combination of hUCB-MSC-derived secretome with atorvastatin also showed higher EPCs migration than the hUCB-MSC-derived secretome-only group at 2%, 10% and 20% concentrations (38.3±3.49 vs 17.2±1.92, 50.2±5.31 vs 27.0±4.00 and 76.4±7.50 vs 51.0±5.15, respectively; all p<0.001). The combination of high-dose (20%) hUCB-MSC-derived secretome and atorvastatin had the highest number of migrated EPC (76.4±7.50 × 10^3) cells. The combination group had a significant and very strong correlation with EPC migration (r=0.970; p<0.001). The linear regression test showed an R^2 of 0.942.

Discussion

This research showed that treatment with hUCB-MSC-derived secretome, atorvastatin and a combination of the two increased the proliferation and migration of EPCs (isolated from CAD patient’s peripheral blood). HUCB-MSC-derived secretome enhances EPCs proliferation and migration in a dose-dependent manner. The combination of hUCB-MSC-derived secretome and atorvastatin was shown to be superior to atorvastatin or hUCB-MSC-derived secretome alone.

In this research, hUCB-MSC-derived secretome treatment increased EPC proliferation in a dose-dependent manner, with the concentrations of 10% and 20% shown to be superior to atorvastatin. Previous studies showed that atorvastatin treatment is superior to other statins at improving EPC proliferation according to recent studies. The combination of hUCB-MSC-derived secretome and atorvastatin was shown to be superior to atorvastatin or hUCB-MSC-derived secretome alone.

A mitogen-activated protein kinase (MAPK) pathway has been known to play a role in increasing EPCs proliferation through increased Cyclin D1 expression mediated by PI3K/Akt and MAPK pathway also involved in EPCs proliferation. While increasing microRNA 221/222 expression shown to reduce EPCs proliferation capabilities. HUCB-MSC-derived secretome treatment was speculated to increase EPCs proliferation through MAPK/ERK and PI3K/protein kinase B pathway. While atorvastatin improves EPCs proliferation through downregulation of microRNA 221/222 expression. The involvement of these multiple pathways may result in higher EPCs proliferation in the combination group.

The synergistic effect of the HUCB-MSC-derived secretome with atorvastatin in enhancing EPCs proliferation and migration was demonstrated in this study. These combinations significantly increase EPCs proliferation and migratory activity by up to two-fold. Previously, The combination of MSCs with another compound, including statins, was shown to have beneficial effects in angiogenesis and neovascularization. Co-culture of MSCs and EPCs have been shown to have improved EPC proliferation and migration, and enhance their angiogenic capacity. However, the exact mechanism of these combinations to improve EPCs proliferation and migration is yet to be investigated. It is speculated that the involvement of multiple pathways may be responsible for its superiority against HUCB-MSCs-derived secretome or atorvastatin alone.
oxidative stress that impairs EPCs migration. Statin also could prevent EPCs senescence by upregulating TRF2 of EPCs, hence enhance migratory capacity. Those facts suggested that the combination of hUCB-MSC-derived secretome and atorvastatin will have superior EPCs migration through the involvement of multiple pathways.

In summary, hUCB-MSC-derived secretome may be developed and combined with atorvastatin treatment in CAD patients to improve EPCs proliferation and migration. However, this research did not measure the exact composition of the hUCB-MSC-derived secretome. The previous study showed that the secretome from another type of MSC can increase EPCs migration but not EPCs proliferation. Hence, further research should be directed to identify the substance within the hUCB-MSC-derived secretome which is responsible for increasing EPC proliferation and migration, and compare it with other MSC secretomes. Further research should also verify the multiple pathways which may be responsible for the improvement of EPCs proliferation and migration in the combination group.

Conclusions
High dose hUCB-MSC-derived secretome outperforms atorvastatin to improve EPC proliferation and migration. A combination of hUCB-MSC-derived secretome with atorvastatin seems to be beneficial in promoting neovascularization through improved EPCs proliferation and migration effect compared to hUCB-MSC-derived secretome or atorvastatin alone.

Data availability
Underlying data
Figshare: Human umbilical cord blood-mesenchymal stem cell-derived secretome in combination with atorvastatin enhances endothelial progenitor cells proliferation and migration. https://doi.org/10.6084/m9.figshare.12186507.v2

This project contains the following underlying data:
- RAW Data - F1000 revis2 by SAH (XLSX). (Raw absorbance data from MTT proliferation assay and cell counts from the Transwell migration assay.)
- RAW DATA f1000 (ZIP). (Raw images generated in this study, including images used to generate cell counts and raw immunofluorescence images.)

Data are available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY 4.0).

Acknowledgements
The authors are thankful to Christian Pramudita, MD., Ilma Alfia Isaridha, MD., Melly Susanti MD., and Dwi Fachrul MD., for invaluable support in this project.

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John David Symmons
University of Utah School of Medicine, Salt Lake City, UT, USA

Anwar Tandar
Division of Cardiovascular Medicine, University of Utah School of Medicine, Salt Lake City, UT, USA

This is an interesting research involving the evaluation of EPCs in CAD and association with statin.

Several concerns should be addressed:
1. It is derived from one patient. The conclusion and validity need to be addressed seriously when there is sample from one patient. In addition, since there is only one patient, why is there even a need for inclusion and exclusion criteria

2. Typographical error in the Introduction... refractory angina, and patients

3. Reference needed for the statement: HUCB-MSCs-derived secretome preparation. The media was collected......(reference 3).

4. Treatment of EPCs...Please describe the rationale for 5 replications. Please describe the rationale behind the selected number 5.4.

5. All abbreviations need to be spelled out in the beginning or first time used

6. Similar to the rational for EPCs treatment, in EPC migrations assay section: Please describe the rationale for 5 fields and describe the sizes of the field.

7. This interesting research should investigate more patients to allow a stronger interpretation and to allow more sound conclusion

8. Other statins should also be investigated as they are being used widely in real world.

9. The exact composition of the hUCB-MSc derived secretome should be described.
In summary, this research needs more diligent work especially with the number of samples from more robust patients population.

**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?**
Partly

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

**If applicable, is the statistical analysis and its interpretation appropriate?**
I cannot comment. A qualified statistician is required.

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

**Are the conclusions drawn adequately supported by the results?**
Partly

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

**Reviewer Expertise:** Anwar Tandar, MD: Clinician, Interventional Cardiologist. John D. Symmons, PhD: Basic Scientist

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

Author Response 30 Nov 2020

**Yudi Her Oktaviono**, Universitas Airlangga, Soetomo General Academic Hospital, Surabaya, Indonesia

Dear Dr Anwar Tandar

Thank you very much for your valuable feedback
I am truly honoured with your feedback

In this opportunity, please let me answer the question

**Q:** It is derived from one patient. The conclusion and validity need to be addressed seriously when there is sample from one patient. In addition, since there is only one patient, why is there even a need for inclusion and exclusion criteria.

**A:** In this experimental laboratory research, the sample used was EPCs derived from single patient with coronary artery disease. We have ensured the diagnosis of coronary artery disease from the ECG, echocardiography and coronary angiography. The purpose of the
single sample being used in this research is to ensure the homogeneity of the sample while inclusion and exclusion criteria was used to identify the best EPCs quality that represent patient with CAD. Mixing EPCs from the various patient may cause EPCs failed to grow efficiently due to incompatibility with other source of EPCs. Inclusion and exclusion criteria is still relevant because we want to ensure that the patient was really diagnosed with coronary artery disease with specific criteria without any other significant disease. Without clear inclusion and exclusion criteria, we may select patient with multiple disease which EPCs function may differ from patient with coronary artery disease only.

Q: Typographical error in the Introduction... refractory angina, and patients
A: Thank you for the feedback, we will revise accordingly

Q: Reference needed for the statement: HUCB-MSCs-derived secretome preparation. The media was collected......(reference 3).
A: We have added the reference, thank you

Q: Treatment of EPCs...Please describe the rationale for 5 replications. Please describe the rationale behind the selected number 5
A: The 5 replication was based on Federer's formula: \((t-1)(n-1) \geq 15\), where \(t\) is the number of treatments and \(n\) is the number of replication.

Q: All abbreviations need to be spelt out in the beginning or first time used
A: Thank you for the feedback, we will revise accordingly

Q: Similar to the rationale for EPCs treatment, in EPC migrations essay section: Please describe the rationale for 5 fields and describe the sizes of the field.
A: The usage of the 5 fields was the standard protocol of cell migration calculation using transwell migration assay based on the Transwell protocol. In this research, we used 0.04mm in diameter as the sizes of the field.

Q: This interesting research should investigate more patients to allow a stronger interpretation and to allow more sound conclusion
A: Thank you very much for your genuine feedback. This research was part of our preliminary research which explore the possibilities of secretome as an alternative regenerative treatment other than cell-based regeneration treatment for the patient with coronary artery disease. In this early research we used limited patient but we will expand further the research with involving more samples and aim to have clinical trials should we achieve satisfying results. We also consider to combine the secretome treatment with existing coronary artery disease treatment, to ensure the usage of secretome combined with current medical treatment will benefit synergistically.

Q: Other statins should also be investigated as they are being used widely in real world.
A: Thank you for the feedback, we will consider that as our future research suggestion. However, we first prefer atorvastatin since it is readily available in our country, Indonesia. Previously, we have compared the effect of atorvastatin, rosuvastatin and simvastatin effect on the EPCs (without secretome) and conclude that atorvastatin was the most superior in
inducing EPCs migration. Thus, this research prefers to use atorvastatin and combine it with secretome.

Q: The exact composition of the hUCB-MSc derived secretome should be described.
A: Thanks, we would like to admit that the limitation of this research was the inability to exactly describe the molecule in the hUCB-MSc derived secretome. Further research is required through proteomic analysis to determine the molecule in the hUCB-MSc derived secretome. This will also help to standardize the composition of hUCB-MSc derived secretome.

Again, many thanks for the review,

Best Regards
Yudi Her

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

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Author Response 22 Apr 2021

**Yudi Her Oktaviono,** Universitas Airlangga, Soetomo General Academic Hospital, Surabaya, Indonesia

Dear Dr Anwar Tandar

Thank you very much for your valuable feedback
I am truly honoured with your feedback

In this opportunity, please let me answer the question
Q: It is derived from one patient. The conclusion and validity need to be addressed seriously when there is sample from one patient. In addition, since there is only one patient, why is there even a need for inclusion and exclusion criteria.
A: In this experimental laboratory research, the sample used was EPCs derived from single patient with coronary artery disease. We have ensured the diagnosis of coronary artery disease from the ECG, echocardiography and coronary angiography. The purpose of the single sample being used in this research is to ensure the homogeneity of the sample while inclusion and exclusion criteria was used to identify the best EPCs quality that represent patient with CAD. Mixing EPCs from the various patient may cause EPCs failed to grow efficiently due to incompatibility with other source of EPCs. Inclusion and exclusion criteria is still relevant because we want to ensure that the patient was really diagnosed with coronary artery disease with specific criteria without any other significant disease. Without clear inclusion and exclusion criteria, we may select patient with multiple disease which EPCs function may differ from patient with coronary artery disease only.

Q: Typographical error in the Introduction... refractory angina, and patients
A: Thank you for the feedback, we will revise accordingly
Q: Reference needed for the statement: HUCB-MSCs-derived secretome preparation. The media was collected......(reference 3).
A: We have added the reference, thank you

Q: Treatment of EPCs...Please describe the rationale for 5 replications. Please describe the rationale behind the selected number 5
A: The 5 replication was based on Federer's formula: \((t-1)(n-1) \geq 15\), where \(t\) is the number of treatments and \(n\) is the number of replication.

Q: All abbreviations need to be spelt out in the beginning or first time used
A: Thank you for the feedback, we will revise accordingly

Q: Similar to the rationale for EPCs treatment, in EPC migrations essay section: Please describe the rationale for 5 fields and describe the sizes of the field.
A: The usage of the 5 fields was the standard protocol of cell migration calculation using transwell migration assay based on the Transwell protocol. In this research, we used 0.04mm in diameter as the sizes of the field.

Q: This interesting research should investigate more patients to allow a stronger interpretation and to allow more sound conclusion
A: Thank you very much for your genuine feedback. This research was part of our preliminary research which explore the possibilities of secretome as an alternative regenerative treatment other than cell-based regeneration treatment for the patient with coronary artery disease. In this early research we used limited patient but we will expand further the research with involving more samples and aim to have clinical trials should we achieve satisfying results. We also consider to combine the secretome treatment with existing coronary artery disease treatment, to ensure the usage of secretome combined with current medical treatment will benefit synergistically.

Q: Other statins should also be investigated as they are being used widely in real world.
A: Thank you for the feedback, we will consider that as our future research suggestion. However, we first prefer atorvastatin since it is readily available in our country, Indonesia. Previously, we have compared the effect of atorvastatin, rosuvastatin and simvastatin effect on the EPCs (without secretome) and conclude that atorvastatin was the most superior in inducing EPCs migration. Thus, this research prefers to use atorvastatin and combine it with secretome.

Q: The exact composition of the hUCB-MSc derived secretome should be described.
A: Thanks, we would like to admit that the limitation of this research was the inability to exactly describe the molecule in the hUCB-MSc derived secretome. Further research is required through proteomic analysis to determine the molecule in the hUCB-MSc derived secretome. This will also help to standardize the composition of hUCB-MSc derived secretome.

Again, many thanks for the review,
The study aims to compare the effect of hUCB-MSC-derived secretome, atorvastatin, and the two combinations in modulating EPC proliferation and migration. The study addresses the novel issues in refractory angina, whether the atorvastatin and secretome derived mesenchymal stem cells improve EPC expression. There are similar studies available addressing these issues. Zhang X et al. demonstrated that intravenous transplantation of huC-MSCs at an early stage could improve hypoxic-ischemic rats' behavior and decreased gliosis, this study was measured in other target disorders 1.

The rationale and scientific background of this manuscript were justified, to disclose the role of secretomes and paracrine stimulation on EPC expression.

In the #Method section, the authors mentioned the inclusion criteria: male, aged 40-59 years old, history of chronic ischemic heart disease as proven by CAG (coronary angiography). The authors should quote the diagnostic criterion. Additionally, the authors should clearly state this is an experimental study of atorvastatin and hUCB-MSC-derived secretome and their combinations on EPC proliferation and migration.

Umbilical cord blood-derived EPC established in a previous study displayed cobblestone-like morphology; this is a typical feature of the EPC. The authors did not state this in the manuscript, except they confirmed using FITC-labelled anti-human CD 34+ expression. The authors should clearly explain it [ref-2].

There was another immunophenotype of EPC as determined by flow cytometry, VEGFR2-PE, vWF-FITC, and CD31-PE 2. Is there any reason why the authors only demonstrated with CD34+ expression. The authors should provide their ideas on it.

In the #EPCs proliferation assay, the authors explain EPCs proliferation measured using OD; there was no explanation of how OD transferred in a measurement scale in Figure 2 (Y-
How did the authors determine the percentage of hUCB-MSC-derived secretome? Is there any control over the measurement?

In the #Table 1 – Characteristics of the patient, the authors should explain "Left ventricle internal diameter" = 5.8 cm; I wonder whether that is either "end-systolic dimension or end-diastolic dimension"?

In #summary, the authors should open the opportunity on the horizon, whether the cell-based therapy or cell-free measures that win the future game?

Again, I would express my appreciation to all authors to address these evolving issues in regenerative medicine.

References

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?
Partly

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

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

Are the conclusions drawn adequately supported by the results?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: 1. Cardiovascular disease (particularly ischemic heart disease) 2. Lipidology and diabetes mellitus 3. Hypertension 4. Stem cell and regenerative medicine
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.

Yudi Her Oktaviono, Universitas Airlangga, Soetomo General Academic Hospital, Surabaya, Indonesia

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A. Thank you for the references and comparison, we will note this feedback and put it on the paper as other research which did similar intervention

2. The rationale and scientific background of this manuscript were justified, to disclose the role of secretomes and paracrine stimulation on EPC expression. Indeed, that was our research aim

3. In the Method section, the authors mentioned the inclusion criteria: male, aged 40-59 years old, history of chronic ischemic heart disease as proven by CAG (coronary angiography). The authors should quote the diagnostic criterion. Additionally, the authors should clearly state this is an experimental study of atorvastatin and hUCB-MSC-derived secretome and their combinations on EPC proliferation and migration.

A. We have added the diagnostic criteria of chronic ischemic heart disease as proven by coronary angiography results that showed >50% stenosis of left main coronary artery or >70% of other coronary arteries. We also explain that this is an experimental study of atorvastatin and hUCB-MSC-derived secretome and their combinations on EPC proliferation and migration.

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morphology before staining the cells with CD34 antibody. We already put these information on the method section.

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A. In this study, the use of FITC CD34+ only to confirmed the EPCs are sufficient, as the same EPCs culture method was used in authors previous research (which mentioned on references number 19, 21-22,45). There was also uncertainty about the use of another immunophenotype of EPC as determined by flow cytometry (VEGFR2-PE, vWF-FITC, and CD31-PE), caused by heterogenous types of EPCs (mentioned on the references of 51,52).

6. Q. In the EPCs proliferation assay, the authors explain EPCs proliferation measured using OD; there was no explanation of how OD transferred in a measurement scale in Figure 2 (Y-ordinate)?

A. EPCs proliferation can be determined through various method, In the previous study, proliferation can be determined relatively through OD (references no 21-22,45). Thus, we use OD as proliferation measurement scale in Figure 2 (Y-ordinate).

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A. The percentage of hUCB-MSC-derived secretome was determined from the dilution level of hUCB-MSC-derived secretome. For example, 1 mL of hUCB-MSC-derived secretome with 49 mL of the phosphate buffer saline (PBS) plus 2% fetal bovine serum (FBS). However, no control over measurement.

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A. It should be Left ventricle end-diastolic dimension, thanks for correcting this phrase.

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

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Author Response 22 Apr 2021

**Yudi Her Oktaviono**, Universitas Airlangga, Soetomo General Academic Hospital, Surabaya, Indonesia

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