α-SMA Expression Increased Over Cell Passages and Decreased by Exogenous TGF-β1, In Vitro Studies on Myofibroblast Derived from Orbital Socket Contracture

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ABSTRACT

α-smooth muscle actin (α-SMA), a marker of myofibroblast, induces cytoskeleton reorganization, increases contractility and stimulates cell migration in TGF-β1 induced stress fibers. The aims of the present study were to determine the level of α-SMA expression and morphological cell changes in different passages of myofibroblasts with varied TGF-β1 concentrations. Myofibroblast cell cultures were derived from fibrotic tissues of fourth degree socket contracture. The α-SMA expression level was measured in myofibroblast cultures passage I, II, and III with and without 10 ng/mL TGF-β1, and in passage III with 2.5; 5; 10; and 20 ng/mL TGF-β1. Results: The levels of α-SMA expression level in passage I to III were I 31.42 ± 3.4; 40.34 ± 8.14 and 56.37 ± 7.57, respectively. Addition of 10 ng/mL TGF-β1 into passage I-III myofibroblast cultures resulted in α-SMA expression level of 31.24 ± 2.93; 36.81 ± 6.09; and 14.29 ± 2.72, respectively. Myofibroblasts passage III showed the lowest α-SMA expression level following exposure to TGF-β1 10 ng/mL (22.37 ± 12.86) and highest without TGF-β1 (48.34 ± 13.36), however no morphological changes detected. α-SMA expression level increased with cell passages, decreases with addition of TGF-β1 while not affecting morphology of myofibroblast derived from the orbital socket contracture.

Keywords: TGF-β1, α-SMA, fibrotic, eye, socket, contracture

INTRODUCTION

Orbital socket contracture is a wrinkled state of anophthalmic socket, thus, it cannot fit the prosthesis. The incidence of socket contracture was reported as 7.7% of the overall anophthalmic socket [1]. Our study, at dr. Saiful Anwar General Hospital Malang, Indonesia, in 2007, however, showed that the occurrence of socket contracture was 32.25% overall, mostly 5 years post-eye removal (60%) [2].

Our study showed that application of Mitomycin C during and after socket reconstruction surgery prevented eye socket contracture recurrences in 50% cases [3]. In one case report, the socket contracture did not recur on a patient undergoing 7 repair surgeries within one year with the use of 0.04% Mitomycin C intra- and post-operatively [4]. Prevention of recurrence of this case may be explained provided that the molecular mechanism of socket contracture has been known.

The socket contracture formed fibrotic tissue and granulation due to chronic irritation, one of which due to friction of the prosthesis with the socket, resulted in impairment of wound healing process [5, 6]. The formation of hypertrophic scars that dominates the socket contracture tissue is a deviation of wound healing in the form of hyperactivity of fibroblast, excessive formation of myofibroblasts and extracellular matrix (ECM) deposition [7, 8]. TGF-β plays roles in all phases of eye wound healing process by increasing growth factor secretion involved in cell migration, proliferation, ECM deposition and myofibroblast formation. Increased growth factor activity not only leads to an increase in wound healing rate, but also a higher risk for scar formation. Therefore, TGF-β is one of the most potential targets for scarring intervention [8, 9, 10, 11].

Currently, the definite pathogenesis of socket con-
tracture has not been clearly elucidated and may involve many factors. Studies on fibrosis were mostly performed using tissues other than socket contracture. Among few studies on socket contracture, one has demonstrated that myofibroblasts play an important role in modulating wound healing, tissue remodeling and organ deformation [12]. TGF-β1 is a major profibrotic cytokine and increased expression plays an important role in the deposition of collagen and extracellular matrix, wound healing and scar formation in the socket contracture tissue [6]. TGF-β1 is a key mediator that induces α-SMA [11, 12].

Alpha-smooth muscle actin (α-SMA) may serve as a reliable marker for myofibroblast cells [12], although others considered it as not a consistent marker for contractile fibroblasts and lots of collagen [13]. Alpha-SMA widely presents in stressed fibers induces cytoskeleton reorganization, increases contractility and stimulates cell migration. Stress fibers contracts myofibroblasts such as in muscle movements.

Myofibroblast cells morphologically resembling fibroblasts and smooth muscle cells, rich in cytoplasmic microfilament (actin-rich stress fibers) and α-SMA expression. Fibroblasts appear on day 2-3 following injury, while myofibroblasts appear on day 12 when wound contractions are nearly 80% complete [11, 12].

In the present study, we reported our in vitro study determining the level of α-SMA protein expression in myofibroblasts derived from orbital socket contracture and whether addition of TGF-β1 may affect α-SMA expression and morphology of such myofibroblasts. Since passage number of cells passed several times may affect protein expression, we also determined whether passage number alone, with or without addition of TGF-β1, affect the level of α-SMA expression in myofibroblast passages [14].

**Materials and Method**

**Sample of myofibroblast**

This study was conducted in November to December 2017 at Ophthalmology Department-dr. Saiful Anwar Hospital and Central Laboratory of Life Sciences – Brawijaya University, Malang. The study design was approved by the Ethics Committee of dr. Saiful Anwar General Hospital and followed the guidelines of the Declaration of Helsinki.

The inclusion criteria for the patient was orbital socket contracture (degree 1 – 5) with medical indication for surgical socket reconstruction. The selected patient was based on signed informed consent to participate in the study. The sample of this study was fibrotic tissue removed from a patient with fourth degree socket contracture.

Primary culture of myofibroblast was prepared from patient with socket contracture surgery [15]. Each tissue was cut into pieces approximately 1A – 1 mm in size, placed in 12 well plastic culture plates in DMEM with 50 U/mL of penicillin, 50 g/mL streptomycin and 10% FCS, and incubated at 37°C in a humidified (95%) incubator under 5% CO2. Cell culture media were changed thrice weekly. Upon reaching confluence, cells were trypsinized with 0.25% trypsin/EDTA solution in phosphate buffered saline (PBS) and passed in 25 cm2 plastic cell culture flasks. Cells were cultured further for this study, or passaged 2-3 times and used directly for experiments, or harvested, resuspended in freezing media and stored frozen in -80°C until used. For experiments, a series of cells passaged I-III were cultured for 3 weeks and incubated 24 hours with or without the addition of 10 ng/mL TGF-β1. Another series of experiments were performed using cells passage III, cultured for 3 weeks and incubated 24 hours with the addition of 2.5; 5; 10; or 20 ng/mL TGF-β1.

**Measurement of α-SMA level and myofibroblast morphology**

The expression levels of α-SMA were measured in passage I, II, III without and with the administration of TGF-β1 10 ng/mL as well as in passage III with TGF-β1 2.5, 5 ng/mL, 10 and 20 ng/mL.

The α-SMA expression level was measured with anti α-SMA antibody [1A4, Cat. number ab7817, Abcam]. Cells were fixed with paraformaldehyde, permeabilized with 0.1% Triton X-100 and blocked with 3% BSA for 30 minutes at 25°C. Samples were incubated with primary antibody (1/400 in 1x TBS) for 3 hours at 25°C. An Alexa Fluor®633-conjugated Goat anti-mouse IgG polyclonal (1/500) was used as the secondary antibody. Fluorescent images of α-SMA myofibroblasts were acquired in red channel using 400x magnification, 0.5% power of excitation lines, and gain set at 850 of the laser scanning confocal microscope (CLSM FV1000, Olympus Corp., Waltham, MA). Confocal images of the fluorophores were analyzed using Olympus Fluoview ver.4.2a software. α-SMA expression was measured in arbitrary unit (AU). Cells were observed also morphologically using 100x magnification with inverted microscope Olympus IX 71 and 400x magnification with CLSM FV1000.
**Data analysis**

The results were analysed using IBM SPSS version 2.0 statistical analysis application. The data analysis in the passage group with TGF-β1 exposure using Shapiro-Wilk normality test, followed by ANOVA or Kruskal-Wallis test, and post-hoc Tukey or Mann-Whitney test to know differences between each group. Results were considered statistically significant when p < 0.05.

**RESULTS AND DISCUSSION**

**Effect of the number of passage on α–SMA level**

The level of α–SMA expression increased with increasing passage number, with values of 31.42 ± 3.4, 40.34 ± 8.14 and 56.37±7.57 arbitrary unit (Figure 1).

Addition of 10 ng/mL TGF-β1 in cell culture decreased α–SMA level only in passage III (p = 0.000). Values of α–SMA level in each group was 31.24 ± 2.93; 36.81 ± 6.09; and 14.29 ± 2.72 arbitrary unit for passage I–III, respectively (Figure 2).

Microscopically, different passage with or without addition of TGF-β1 did not change cell morphology (Figure 3).

When different passages were compared between with and without addition of TGF-β1, our results showed that the most pronounced effects were seen on passage III and with addition of TGF-β1 (Figure 4). On passage III, addition of TGF-β1 decreased a-SMA expression up to three folds.

Effect of TGF-β1 concentration added on a-SMA expression was studied using passage 3 of myofibroblast primary culture from socket contracture. The results showed that compared to the myofibroblast without TGF-β1, only 10 ng/mL TGF-β1 decreased significantly α–SMA expression level (p=0.000). The α-SMA expression level was 48.34 ± 13.36; 46.45 ± 47.04; 38.07 ± 9.54; 22.37 ± 12.86; and 29.80 ± 13.33 in cell culture with TGF-β1 0; 2.5; 5; 10; and 20 ng/mL, respectively (Figure 5).

Our study demonstrated that passaging influence α–SMA level on myofibroblast primary culture derived from orbital socket contracture tissue. Similar results were reported by Kinner et.al. (2001) that levels of α–SMA are increasing following the number of passages or passages performed. One of the main factors of physiological change from fibroblasts to myofibroblast is a mechanical tension. [16] According to Wakatsuki et.al. (2000) there is progressive increase in matrix rigidity overtime if the tissue experienced a tension, including in our present study during passaging process. [17] Our study demonstrated that following passage and cultured

![Figure 1. Level of α-SMA in cell culture with different passage (P1, P2, P3) without TGF-β1](image1)

![Figure 2. Level of α-SMA III in cell culture with different passage (P1, P2, P3) with addition of 10 ng/mL TGF-β1. * denotes significantly different compared to P1 (p=0.000)](image2)
Figure 3. Confocal microscopic observations. Bright field view (left), red filter immunofluorescent view for α-SMA (middle), and the merged view of both (right). Passage I (A), passage I with TGF-β1 10ng/ml (B), passage II (C), passage II with TGF-β1 10 ng/mL (D), passage III (E), passage III with TGF-β1 10 ng/mL (F). Different passages and addition of TGF-β1 did not resulted in observable changes on cells morphology.
myofibroblast differentiation was inhibited without decreasing TGF-β1 receptor expression. [25]. Further studies are required to clarify whether FAK activation causes α-SMA degradation and apoptosis of myofibroblasts, or other pathways causing TGF-β1 decreased α-SMA.

CONCLUSION

Passage number increased whereas TGF-β1 could decrease α-SMA expression levels in dose-dependent fashion, however, both did not affect morphology of myofibroblast culture derived from orbital socket contracture. Therefore, in vitro experiments to study myofibroblast from orbital socket contracture should consider passage number and level of TGF-β1.

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REFERENCES


