

Research Article

Isolation of Skeletal Muscle-derived Cells Modeling Neural Crest-derived Stem Cells for Therapeutic use in Regenerative Periodontology

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Citation: Grimm WD, Benlidayi M, Didenko N, Dhingra K, Dolgalev A, et al. (2021) Isolation of Skeletal Muscle-derived Cells Modeling Neural Crest-derived Stem Cells for Therapeutic use in Regenerative Periodontology. Dent Adv Res 6: 178. DOI: 10.29011/2574-7347.100078

Received Date: April 01, 2021; **Accepted Date:** April 12, 2021; **Published Date:** April 16, 2021

Abstract

Periodontitis is microbial infection affecting periodontium, the tooth supporting structure and affects >743 million people worldwide. Neural crest-derived stem cells (NCSCs) hold the promise to regenerate the damaged periodontium. These cells have been identified within adult adipose tissue, periodontal ligament, and palatal tissue. Typical enzymatic isolation protocols are expensive, time consuming and often not clinically compliant. Enzyme-free, mechanical dissociation has been suggested as an alternative method of generating cell suspensions required for cell separation and subsequent expansion *ex vivo*. In our study, samples of rat skeletal muscle tissue were used to appraise the suitability of a novel mincing method of mechanical dissociation against enzymatic digestion with collagenase and dispase. Skeletal muscle is readily available and has been shown to contain NCSC populations. We used a Rigenera-Human Brain Wave[®] prototype mincer to produce a suspension of skeletal muscle-derived cells modeling NCSCs. We have compared the resulting cell cultures produced via mechanical dissociation and enzymatic dissociation, producing single cell suspensions suitable for Magnetic Cell Sorting (MACs) and Fluorescence-activated cell sorting (FACS). Despite the Countess Automated Cytometry data demonstrating that cell suspensions produced by mechanical dissociation (n=24) contain on average 26.8 times as many viable cells as enzymatic cell suspensions (n=18), enzymatic suspensions produced more successful cell cultures. Spheroids and subsequently adherent cells formed from 4 enzymatic cell suspensions (44.4%) vs. 1 mechanical cell suspension (8.3%). Enzymatic digestion protocols formed spheroids faster and more plentifully than mechanical cell suspensions. Adherent cells and spheroids isolated *via* both methods appear morphologically similarly to NCSCs from our previous studies.

Keywords: Isolation; skeletal muscle-derived cells; Neural crest-derived stem cells; Mechanical dissociation; Regenerative periodontology; Enzymatic digestion

Introduction

Periodontitis is a chronic multifactorial inflammatory disease associated with dysbiotic plaque biofilms and characterized by progressive destruction of the tooth-supporting apparatus [1]. In 2010, severe periodontitis was identified as the world's 6th most prevalent disease, affecting more than 743 million people [2]. The need for an effective treatment is further highlighted by the fact that periodontitis remains the leading cause of tooth loss worldwide [3]. Tooth loss *via* exfoliation or professional extraction is witnessed in advanced cases of periodontitis following extensive attachment loss and resorption of alveolar bone. Most current treatments for patients suffering from periodontitis focus on the conservation of remaining periodontal structures and reducing disease progression [4]. Despite continuing development and research into treatments for periodontitis, including tissue engineering and regenerative therapies, difficulty remains in restoring all the damaged structures of the periodontium [5]. Furthermore, even with current treatments, studies have shown worldwide prevalence of severe periodontitis stagnated between the years of 1990 and 2017 [2].

Periodontal ligament (PDL) cells are resident cells of the periodontium and have a critical role in tissue homeostasis, destruction, and regeneration by their ability to synthesize and degrade collagen and other matrix molecules [6]. The immune response is orchestrated by numerous cytokines, chemokines, and growth factors, which are produced by resident host cells. However, these cells can also participate in the immunoinflammatory processes of periodontitis [7,8]. Periodontal healing is determined by the type of cells that repopulate the root. By the application of regenerative treatment methods, which promote PDL cell proliferation, migration and attachment, the re-establishment of the initial periodontal tissue architecture is possible [9]. However, the outcomes of currently available regenerative treatment approaches are sometimes compromised by several factors and are not predictable [10,11]. Therefore, the search for new therapeutic methods with a regenerative potential are a major goal in periodontology [12].

Stem cells represent an exciting potential therapy for periodontitis thanks to their ability to differentiate into many different cell types, such as bone and connective tissues [13]. However, the appropriate type of stem cells must be selected to develop the most effective treatment for periodontitis. In 2004, the periodontal ligament was identified as a source of highly

proliferative, multipotent mesenchymal stem cells with potential for the treatment of periodontitis [14]. Conversely to mesenchymal PDL-SCs which differentiate into those cells and tissues related to the mesenchyme; neural crest cells (NCSCs) have displayed multipotent capabilities to differentiate into derivatives of both mesenchymal and neuronal lineage [15]. The most attractive sources for adult NCSC isolation are dental sources because of easy access and minimally invasive biopsy. NCSCs have been identified and isolated from several tissues and organs of mammals at later fetal and postnatal stages of development [16]. Adult NCSCs could undergo directed differentiation into adipocytes, osteoblasts, chondrocytes, melanocytes, neurons, and Schwann cells [15,17,18]. Moreover, adult cultured NCSC therapeutic application led to the restoration of bone [19], cartilage defects [20], damaged peripheral nerves [21], and spinal cord [22]. Therefore, selective isolation of adult oral NCSCs are important technical issues for the successful applications for regenerative medicine purposes. Various techniques were applied for the selective isolation of adult NCSCs:

- fluorescence-activated cell sorting [23,24]
- selective culturing conditions for growth as shown as neurosphere-like structures [25]
- explant technique [26]

Only pluripotent stem cells exceed the cellular plasticity displayed by isolated adult multipotent neural crest-derived stem cells (NC-SCs). The hard palate presents as an accessible source of cranio-facial tissue for the procurement of neural crest-related cells [25]. In 2009, populations of multipotent NC-SCs were identified within the ridges of the hard palate of adult Wistar rats [27]. Considering the highly regenerative nature of the hard palate and accessibility of palatal rugae, this procedure could act as a minimally invasive source of proliferative, therapeutically appropriate multipotent NC-SCs [25]. Following isolation and cultivation of nestin-positive and Sox2-positive NC-SCs, secondary neurosphere forming assays identified NC-SCs forming spheres at a frequency of 1.8%, compared to a frequency of 0.01% in PDL-SCs [27,28]. The NC-SCs are a heterogeneous population of stem/stromal cells isolated from subepithelial palatal tissue complex. Administration of NC-SCs holds a strong therapeutic potential for regenerative and wound healing medicine applications aimed at functional restoration of tissues damaged by chronic periodontal diseases [29]. NC-SCs are obtained from subepithelial palatal tissue complex by collagenase digestion, incubation/isolation, and pelletized by centrifugation. Enzymatic disaggregation may alter the relevant biological characteristics of subepithelial palatal tissue

complex, while providing release of complex, multi-attachment of cell-to-cell and cell-to-matrix, effectively eliminating the bioactive ECM and periadventitial attachments. The isolation of cellular elements is considered as a minimal-invasive access in periodontal surgery [17,30]. Several alternative, nonenzymatic methods of palatal tissue processing have been developed to obtain via minimal mechanical manipulation an autologous NC-SC product intended for delivery, reducing the procedure duration, decreasing regulatory burden, and shortening the translation into the clinical setting [26]. Ideally, these procedures might allow for the integration of harvesting and processing of palatal tissue for ease of transferring, in a single procedure utilizing a nonexpanded cellular product at the point of periodontal regeneration, while permitting intraoperative autologous NC cellular-based periodontal regenerative therapies. Therefore, palatal-derived NC-SCs present as more potent and more proliferative alternative to PDL-SCs. Here, we investigate the options, advantages, and limitations of the methods alternative to enzymatic processing currently developed for minimal manipulation of palatal tissue.

In order to liberate stem cells from tissue samples, the tissue must be dissociated using mechanical or enzymatic methods. In the reports mentioned previously, samples of periodontal ligament and palatal rugae were dissociated using enzymatic methods, involving treatments with enzymes such as trypsin, dispase, and collagenase I and II [14,25,27,31]. The limitations of enzymatic digestion surround the facts that these enzymes are both expensive, and often not “xeno-free” [32]. Enzymes sourced from different species of animals, or bacteria could result in serious adverse reactions for the patient, including infection or anaphylaxis [32]. Furthermore, inconsistencies with enzyme concentration, purity and activity have been noted between different batches and manufacturers. These issues from the common production method of enzymes *via* recombinant bacteria [33] required adaptation of digestion protocols depending on varying w/v % concentrations of available enzymes. Mechanical dissociation aims to provide an alternative, enzyme- and xeno-free process for the liberation of stem cells from tissue samples. Methods utilizing centrifugal force, vibration, pressure, shearing force, and radiation force have been developed to replace the requirement of an enzymatic digestion step. However, these methods have displayed varying successes [33,34].

This study aimed to compare dissociation of mammalian tissue achieved via novel mechanical mincing techniques against conventional enzymatic dissociation methods utilizing dispase I and collagenase I. This is a *proof of principle* study, which as a model for palatal-derived NC-SCs compares the mechanical

isolation of skeletal muscle-derived cells with enzymatic methods. Muscle tissue samples taken from rats were selected to assess the differences in cell suspensions produced by these mechanical and enzymatic dissociation methods. Recently, Mellows et al. [35] and Mitchell et al. [28] reported that the secretome from human amniotic fluid stem cells (AFSCs) facilitates tissue regeneration in a cardiotoxin (CTX)-induced model of muscle degeneration [28,35]. In accordance with these publications rat skeletal muscle contain a variety of different cells displaying the characteristics of mesenchymal stem cells. Hence, rat skeletal muscle presents as a suitable, readily available sample tissue for the initial assessment of a novel mechanical dissociation protocol against proven enzymatic digestion methods. The research hypothesis for this study was that clinically viable skeletal muscle-derived cells modeling NC-derived stem cells can be isolated from samples of mammalian tissue using mechanical mincing techniques, instead of enzymatic digestion for the treatment of periodontitis.

Materials and Methods

Rat muscle tissue sample

Fresh samples of rat muscle tissue specifically, samples of the fibularis, tibialis and extensors were isolated the day before scheduled experiments. A beaker of Male Wistar Kyoto (WKY, 8–12 weeks, 200–300 g) rats were killed according to schedule one of the Animals (Scientific Procedures) Act 1986 and thus was given an ethical approval waiver by the University of Reading Animal Welfare and Ethical Review Board (AWERB). Tissue samples in Krebs solution was provided, stored at a temperature of approximately 4°C.

Cell medium

Rat smooth muscle cells were cultured in high glucose DMEM supplemented with 20% FBS, 1% L-glutamine and additionally with 1% Penicillin-Streptomycin (Sigma-Aldrich, Germany), 77% Dulbecco’s Modified Eagle’s Medium – High Glucose (Sigma-Aldrich, Germany), 0.00005% FGF and 1% Amphotericin B. Unless otherwise stated, cell cultures were fed every 3 days. Successful cultures in T25 flasks would have two additions of 3ml of media for the first two 3-day cycles, before supernatant was removed and adherent cells were split.

Weighing tissue samples

Tissue sample mass was measured using a digital precision balance (Sartorius, R200D Digital Analytical Lab Balance) in grams, providing accuracy to five decimal places, facilitating recording in milligrams. The mass of each sample was maintained within a 10% range between lightest to heaviest, for each set of samples for each experiment. Before treatment with either

mechanical or enzymatic dissociation methods, tissue samples were manually minced into divisions of 6, 12, or 18 (Figure 1). Once manually minced, tissue samples were gently rinsed in 1% Penicillin-Streptomycin in PBS solution, comprising of Dulbecco's PBS (Modified, without calcium chloride and magnesium chloride) (Sigma-Aldrich, Germany) and Penicillin- Streptomycin (Sigma-Aldrich, Germany) in order to minimize microbial contamination during dissociation procedures.

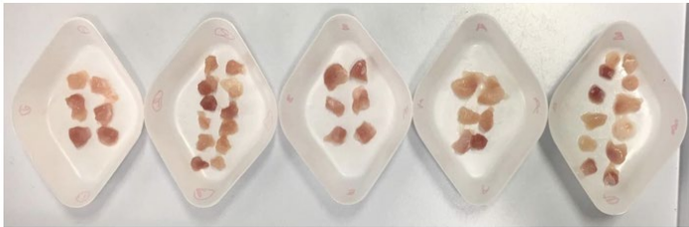


Figure 1: Manually minced rat muscle tissue prior to enzymatic and mechanical dissociation.

Enzymatic digestion methods

Following manual mincing and washing, a select cohort of rat tissue samples were subject to enzymatic dissociation. Firstly, tissue samples were treated with 2ml of 1U/2ml Dispase I in PBS solution, comprising of 10 μ l of 200U/ml Dispase I protease (Sigma-Aldrich, Germany) with 1990 μ l sterile Dulbecco's PBS (Sigma-Aldrich, lot no. Germany). The tissue sample in Dispase-PBS solution were vortexed together in a 10ml falcon tube for 30 seconds and refrigerated for 24-hours at a temperature of 4°C; completing the first stage of the dissociation procedure. Following 24-hours of refrigeration, samples were centrifuged at 300G for 10 minutes (Eppendorf, Centrifuge 5810R) to create a pellet, so that the Dispase-PBS supernatant could be removed. The resultant pellet was then re-suspended in 2ml of a 1U/2ml Collagenase I in PBS solution, comprising of 200 μ l 5U/ml Collagenase I (Collagenase from *Clostridium histolyticum*) (Sigma-Aldrich, Germany) with 1800 μ l sterile Dulbecco's PBS (Modified, without calcium chloride and magnesium chloride) (Sigma-Aldrich, Germany). The resultant tissue in Collagenase-PBS suspension was incubated in a water bath at 37.5°C for 90 minutes. Following incubation, the suspension was made up to 12ml with sterile Dulbecco's PBS and transferred to a 50ml falcon tube. Once transferred, the suspension was filtered using a 70 μ m cell strainer into a 10ml falcon tube to remove any debris. The resultant suspension was centrifuged at 300G for 10 minutes (Eppendorf, Centrifuge 5810R) to create a pellet so the collagenase-PBS supernatant can be removed. The resultant pellet was re-suspended in 1ml of media, containing 20% Foetal Bovine Serum (Sigma-Aldrich, Germany) 1% L-Glutamine (Sigma-Aldrich, Germany), 1% Penicillin-Streptomycin (Sigma-

Aldrich, Germany) 78% Dulbecco's Modified Eagle's Medium – High Glucose (Sigma-Aldrich, Germany) and 0.00005% FGF before counting and seeding.

Mechanical dissociation

Following manual mincing and washing, samples selected for mechanical dissociation were stored in 2ml PBS (Sigma-Aldrich, Germany) with 1% Penicillin-Streptomycin (Sigma-Aldrich, Germany) whilst waiting for mechanical mincing. The prototype mincer provided by was utilized for the mincing of rat muscle samples (Figure 2). Tissue samples were placed into the Rigenera Cons (Rigenera-HBW, 2018) mincing pot with approximately 5ml PBS (Sigma-Aldrich, Germany) and secured with laboratory grade Parafilm M. Mincing was carried out in sessions of 5-10 minutes followed by removal of supernatant via a sterile syringe with the supplied filter. Filtrate was dispensed into a 10ml falcon tube, and replacement PBS was instilled into the Rigenera Cons device, and re-secured with Parafilm M. An additional 5-10 minutes of mincing was carried out, and the resultant supernatant removed and instilled into the filtrate 10ml falcon tube. The filtrate solution was made up to 12ml with sterile Dulbecco's PBS and transferred to a 50ml falcon tube. Once transferred, the solution was filtered using a 70 μ m cell strainer into a 15ml falcon tube to remove any debris. The resultant solution was centrifuged at 300G for 10 minutes to create a pellet so the PBS supernatant can be removed. The resultant pellet was resuspended in 1 ml of media, containing 20% Foetal Bovine Serum (Sigma-Aldrich, Germany) 1% L-Glutamine (Sigma-Aldrich, Germany), 1% Penicillin-Streptomycin (Sigma-Aldrich, Germany) 78% Dulbecco's Modified Eagle's Medium – High Glucose (Sigma-Aldrich, Germany) and 0.00005% FGF before counting and seeding.



Figure 2: Prototype Mincing Apparatus for Mechanical Dissociation Experiments. (L and M) Rigenera Cons mincing apparatus. (M) Rigenera Cons together with supplied filter. (R) Handheld motor component of the prototype mincing apparatus, including foot pedal control.

Cell population counts

Cell counts for enzymatic and mechanical dissociation were carried out before seeding, using the Countess II FL Automated

Cell Counter. 10µl of the 1ml cell suspension in culture medium was sampled and placed into a 50µl Eppendorf microcentrifuge tube. 10µl of Trypan Blue cell stain was then added to the 10µl cell suspension sample, creating a total volume of 20µl. 10µl of this 20µl Trypan Blue-cell suspension was removed from the 50µl Eppendorf microcentrifuge tube and instilled into each chamber of a sterile countess cell counting chamber slide. Each end of the slide can then be inserted into the automated hemocytometer, providing multiple readings for each dissociation experiment.

Passaging cultures

Following assessment of confluence using transmitted light microscopy, cell cultures at approximately 80% confluency were “passaged”, or split, in the sterile environment of the laminar flow cabinet. For cell cultures with adherent cells: old cell culture media was removed from the flask using a sterile stripette and disposed of using Biocleanse solution. For cell cultures with suspended cells: old cell culture media was removed from the flask using a sterile stripette and transferred into a 10ml falcon tube. This cell suspension was centrifuged at 300G for 10 minutes to create a pellet suitable for resuspension. After removing old cell culture media, the pellet was resuspended in 1ml fresh media, ready for a cell count or seeding in a new cell culture flask.

The remaining adherent cells and flask were washed with Dulbecco’s PBS (Modified, without calcium chloride and magnesium chloride) (Sigma-Aldrich, Germany) in sterile conditions 2-3 times, before the addition of Trypsin-EDTA Solution (Sigma-Aldrich, Germany). 4 ml trypsin was used for a T75 flask, whereas 1.3 ml was used for a T25 flask. Trypsin was distributed over the base of the flask, ensuring all adherent cells are covered. Cell cultures were then incubated for 5 minutes in a 10% CO₂ humidified incubator, at 37°C. Following incubation, cells were viewed using transmitted light microscopy to assess whether all adherent cells were now detached from the base of the culture flask; additional incubation was applied if gently tapping the side container did not dislodge the remaining adherent cells. A volume of standard cell media, equal to that of the added trypsin solution, was added to the flask using a stripette in the sterile environment of the laminar flow cabinet. The suspended cells present in the trypsin-media solution was centrifuged at 300G for 10 minutes to create a pellet, so that the trypsin-media supernatant could be removed. Following removal of supernatant, the pellet of cells was resuspended in 1ml of media, ready for a cell count or for reseeding in a T25 or T75 cell culture flask, with additional fresh media.

Transmitted light microscopy

Invitrogen™ EVOS™ XL microscopes were used to assess cell cultures under transmitted light microscopy. Together with an optical magnification of 10-times, imaging of cell cultures was carried out at 4-, 10-, 20- and 40-times objectives.

DAPI/Phalloidin staining and microscopy

Cells were seeded into an adherent 24-well plate prior to the assay with 1ml standard cell media. Once cells had adhered, media was removed and replaced with 1ml of 4 % paraformaldehyde solution for 20 minutes at room temperature. After 20 minutes, paraformaldehyde solution was removed, and the cells washed with PBS three times, at 5-minute intervals. Finally, 150µl of 0.1% Triton™ X-100 (Sigma-Aldrich, Germany) in PBS was added to permeabilize adherent cells and left to set at room temperature for half- an-hour and then removed using a pipette. Residual Triton X-100 solution was washed away with three PBS washes, at 5-minute intervals. Each well of cells selected for DAPI/Phalloidin staining were then treated with 250µl of 1% phalloidin, 99% DAPI treatment stain solution. The 24-well plate was subsequently protected from light using foil and refrigerated at 4°C overnight. Invitrogen™ EVOS™ FL microscopes were used to assess cell cultures following DAPI/Phalloidin staining protocols using fluorescent light microscopy.

Live/Dead staining and microscopy

Cells were seeded into an adherent 24-well plate prior to the assay with 1ml standard cell media. Once cells had adhered, media was removed and replaced with 1ml of Calcein AM (CalAM) and ethidium homodimer-1 (EthD-1) staining solution. This solution was made by combining 10ml of standard growth media with 20µl of EthD-1 and 5µl of CalAM. Once desired wells have been treated with the EthD-1 and CalAM stain, the plate was incubated for 40 minutes before imaging. Invitrogen™ EVOS™ FL microscopes were used to assess cell cultures following live/dead staining protocols using fluorescent light microscopy.

Cleaning protocol

The “Rigenera Cons” mincing apparatus was cleaned using a novel cleaning method. Firstly, the mincer was stored in ethanol 70% v/v overnight. Following this, the ethanol was drained off and replaced with a 10% w/v detergent solution of ‘Tergazyme’ enzyme activated detergent with distilled water in a 150ml beaker into which the apparatus that needed to be cleaned was placed. Tergazyme 10% solution was added until all contents of the

beaker were completely submerged. The beaker was then secured on a see-saw rocker (Stuart, SSL4) at a rate of 40 oscillations-per-minute for 2 hours. Finally, Tergazyme 10% w/v solution was drained off and a series of washes were carried out using water, double distilled water and finally 70% v/v ethanol.

Fibroblasts and assessment of cleaning protocol

Adult Human Dermal Fibroblasts (NHDF-Ad) passages 8, 9 and 10 were used to assess the Rigenera Cons cleaning protocol. Control fibroblasts were seeded into a T25 adherent cell culture flask at a cell density of 8.75×10^4 cells in 3ml of standard cell culture medium. Test fibroblasts were passed through the Rigenera Cons mincing device multiple times using a sterile micropipette within a sterile laminar flow cabinet. Following five minutes of exposure, cells were removed and seeded at a density of 8.75×10^4 cells in 3ml. This process was repeated three times.

Statistical analysis

GraphPad Prism 8 software was utilized for statistical analysis, including Student’s t-test, two-tailed, with confidence level 95%. $P < 0.05$ was used as the definition of statistical significance.

Results

More cells are liberated from tissue samples using mechanical dissociation compared to enzymatic digestion

Analysis of the cell suspensions produced by mechanical and enzymatic protocols indicates mechanical dissociation as producing significantly larger populations of living cells in comparison to enzymatic digestion (Figure 3).

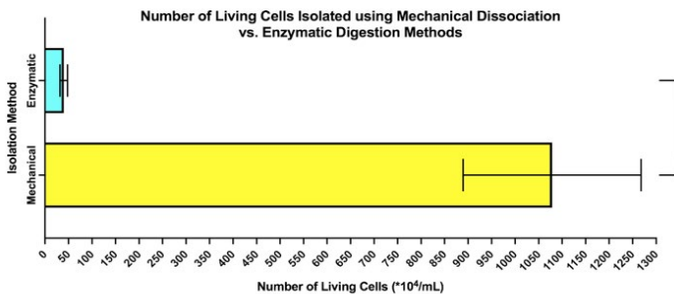


Figure 3: Quantification of the Living Cell Numbers Reveals Higher Yield of Mechanical Dissociation. These cell numbers were quantified using the Countess Automated Cytometer. Data represented as mean and SEM. Student’s t-test analysis set at ($P < 0.05$) indicates extreme statistical significance ($P < 0.0001$). Enzymatic n=18, Mechanical n=24.

This difference in populations of liberated live cells is despite equal masses of sample tissue used with each technique (Figure 4).

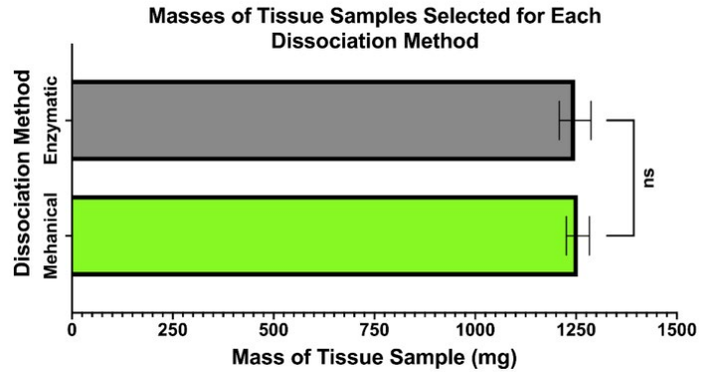


Figure 4: Tissue Samples Selected for Mechanical and Enzymatic Dissociation were of Equal Mass. Data represented as mean and standard error of the mean (SEM). Analysis of tissue sample masses using Student’s t-test analysis set at ($P < 0.05$) indicates no statistical significance between masses of tissue samples selected for each dissociation method. Enzymatic n=9, mechanical n=12.

Figures 5 and 6 further illustrate the differences between populations of living and dead cells present in cell suspensions produced by enzymatic and mechanical dissociative techniques.

Using data from the Countess Automated Cytometer, on an average, enzymatic digestion produced cell suspensions with 89.07% of cells identified as “alive”, compared to 99.64% of liberated cells identified as “alive” from mechanical dissociation experiments. Considering mean data for sample masses and cell suspension populations, further calculations can identify the average numbers of living cells liberated from 1000mg of tissue sample, using each dissociative protocol.

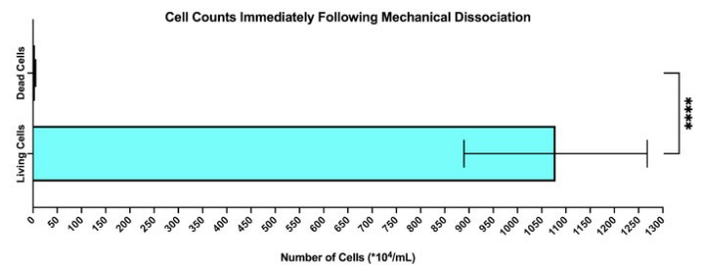


Figure 5: Comparison of living cell and dead cell populations immediately following mechanical dissociation shows on average 99.64% of cells are alive. These cell numbers were quantified using the Countess Automated Cytometer. Data represented as mean and SEM. Student’s t-test analysis set at ($P < 0.05$) indicates extreme statistical significance ($P < 0.0001$). Mean values for dead cells = 3.934×10^4 /ml and mean values for living cells = 1079×10^4 /ml. Living Cells n=24, Dead Cells n=24.

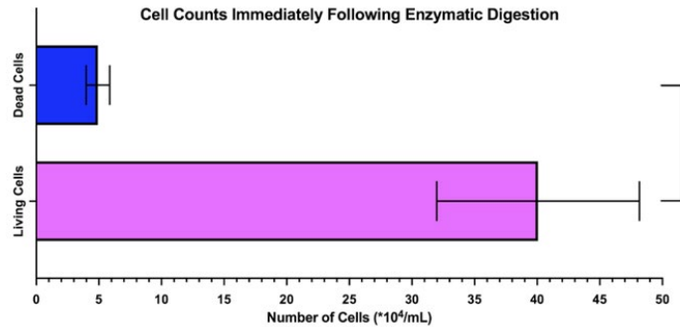


Figure 6: Comparison of living cell and dead cell populations immediately following enzymatic digestion shows on average 89.07% of cells are alive. These cell numbers were quantified using the Countess Automated Cytometer. Data represented as mean and SEM. Student’s t-test analysis set at ($P < 0.05$) indicates extreme statistical significance ($0.0001 < P < 0.001$). Mean values for dead cells = $4.919 \times 10^4/\text{ml}$ and mean values for living cells = $40.07 \times 10^4/\text{ml}$. Living Cells $n= 18$, Dead Cells $n=18$.

Data in Table 1 suggests that mechanical dissociation should produce 26.8 times the number of living cells compared to enzymatically produced cell suspensions, using the same mass of tissue sample.

Dissociation Method	Average Sample Size	Average living cell population	Average number of cells per 1000mg tissue sample
Mechanical Dissociation	1254 mg	1079×10^4	860.45×10^4
Enzymatic Digestion	1248 mg	40.07×10^4	32.11×10^4

Table 1: Average number of living cells liberated using mechanical or enzymatic protocols per 1000 mg of tissue sample.

Figure 7 shows the visual differences between cell suspensions produced by mechanical and enzymatic protocols. Images 1 and 2 clearly show a typical fine layer of occlusive debris produced by mechanical mincing, compared to the clearer cell suspensions produced by enzymatic digestion (images A and B). Furthermore, suspensions produced from enzymatic digestion (images A and B) included presence of larger fragments of muscle tissue, absent from suspensions produced by mechanical dissociation (images 1 and 2).

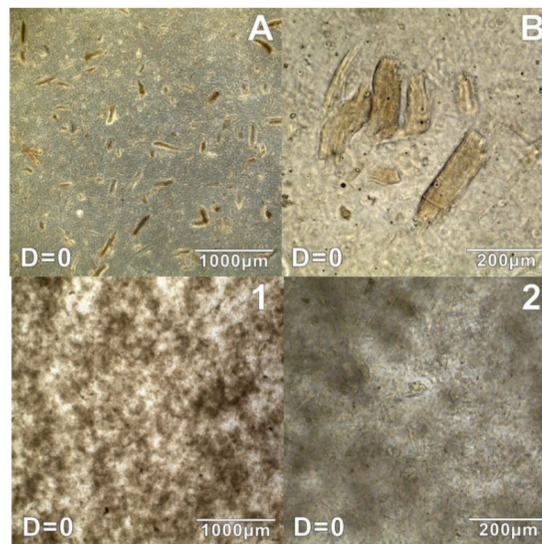


Figure 7: Comparison of cell suspensions immediately after seeding cell suspensions produced by mechanical and enzymatic dissociation techniques shows debris from mincing process. (A and B) Successful culture ‘Enzymatic Dissociation B’. (1 and 2) Successful culture ‘Mechanical Dissociation 2’.

Adhesion of cells following mechanical dissociation and enzymatic digestion

Mechanical dissociation and enzymatic digestion successfully produced cultures of adherent cells. Formation of large cellular spheroids, similar in morphology to neurospheres produced by neural crest-derived stem cells, occurred quickly in enzymatic cultures (Figure 8). Successful mechanical cultures produced far fewer numbers of spheroids than seen in enzymatic cultures. Those spheroids that did occur were much smaller and sparsely distributed throughout the flask (images A, D and E, Figure 9). Successful mechanical cultures did not reach comparable densities of adhesive cells seen in enzymatic cultures (D=13) until day 19.

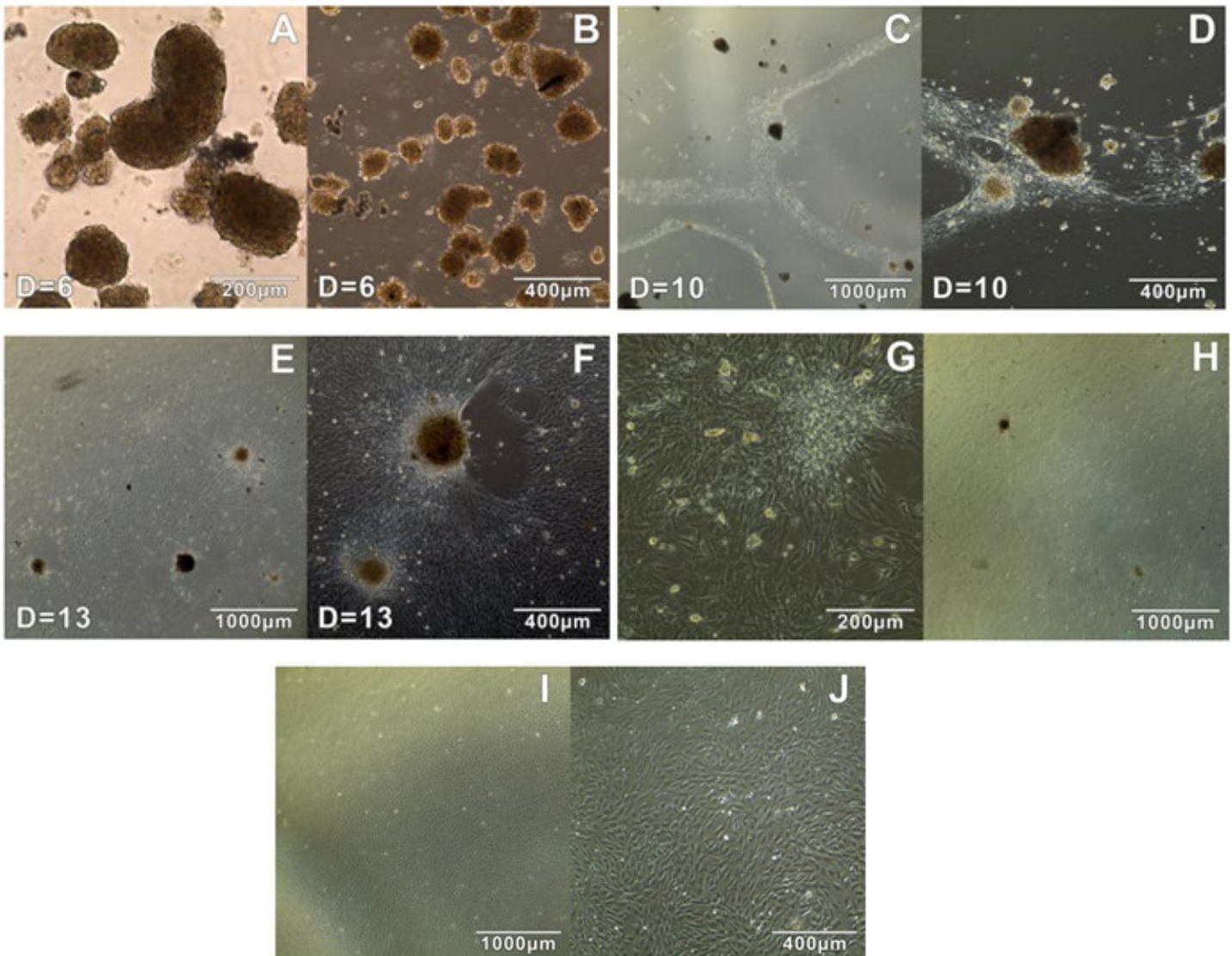


Figure 8: Enzymatic dissociation methods successfully produce spheres, from which adherent cells migrate outwards. (A and B) Multiple cell spheroids, similar in morphology to NCC neurospheres, identified in cell culture 6 days (D=6) after initial cell seed on 13/11/18. (C and D) Migration of adherent cells out of spheres. (E and F) Proliferation of adherent cells, following migration out of cell spheres. Images taken prior to first split, following treatment with Amphotericin B on 23/11/18. (G and H) Adherent cells continue to proliferate following first split and Amphotericin B intervention (I and J). Adherent cells continue to proliferate in the absence of spheres, following second split of cell culture.

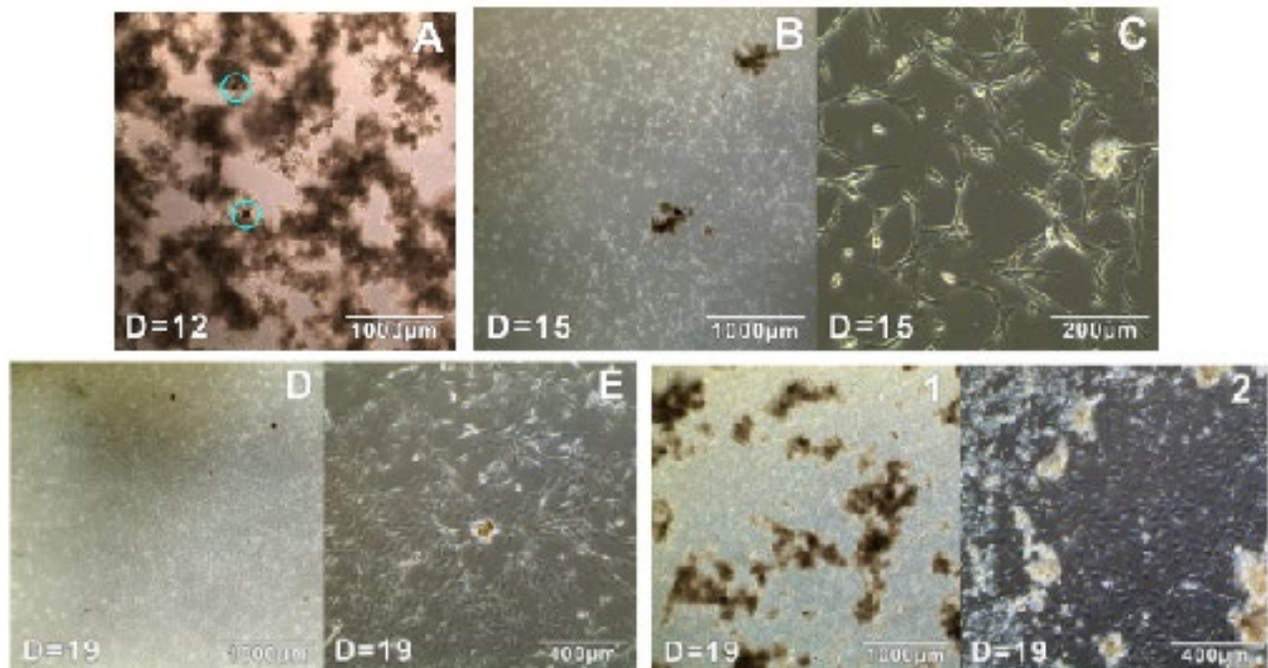


Figure 9: Mechanical Dissociation Methods Displays Similar Capability to Produce Spheres, from which Adherent Cells Migrate Outwards. Possible spheroids identified within tissue sample debris following mechanical dissociation 12 days before (D=12), (B and C) Adherent cells identified 15 days after initial seed; following removal of supernatant (1 and 2), (D and E) Adherent cells continue to proliferate following three cycles of media changes/addition. (1 and 2) Supernatant and non-adherent tissue debris produced proliferative cultures of adherent cells when seeded; 19 days after initial isolation.

Both successful mechanical and enzymatic cultures produced adherent cells that responded well to splitting *via* typical trypsinization methods. Two cell proliferation assays were carried out over a 7-day course to assess the cell proliferation of adherent cells isolated via enzymatic dissociation methods. Over 7 days with no addition of fresh media, cell populations grew from populations of approximately 10,000 cells per well to an average of 71,300 cells per well (Figure 10).

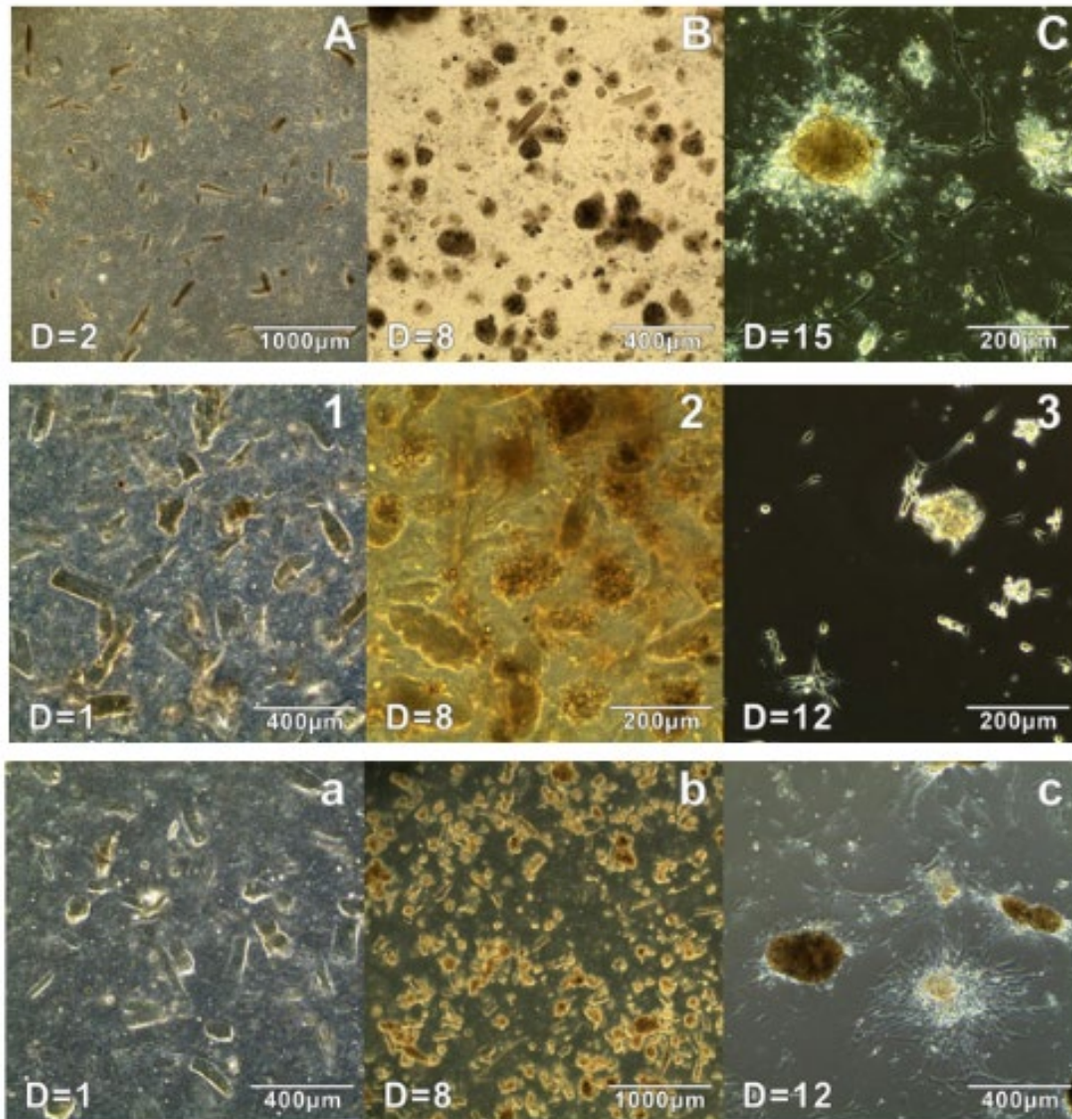


Figure 10: Trend in the Development of Adherent Cells from Cell Suspensions Produced by Enzymatic Digestion. Culture ‘Enzymatic B’: (A) Cell suspension 2 days post-seeding (D=2), Spheroids begin to form on day 8 (C) Non-adherent cells in supernatant produce spheroids and subsequent adherent cells following cultivation. Culture ‘Enzymatic 3’: (a) Cell suspension 1 day post-seeding (b) Spheroids begin to form within the cell suspension on day 8 (c) Non-adherent cells in supernatant produce spheroids and subsequent adherent cells following cultivation, Culture ‘Enzymatic 4’: (1) Cell suspension 1 day post-seeding (2) Spheroids begin to form within the cell suspension on day 8 (3) Adherent cells present in original flask, following removal of non-adherent supernatant and cultivation with cell media including 1% amphotericin B.

LIVE/DEAD staining assay

Results from live/dead staining assays using calcein-AM and ethidium homodimer-1 stains indicate that enzymatic and mechanical dissociation methods both display capacity to produce healthy cultures of adherent cells from samples of mammalian tissue (Figure 11). Furthermore, those cultures of adherent cells remain healthy, following repeated splitting and re-seeding.

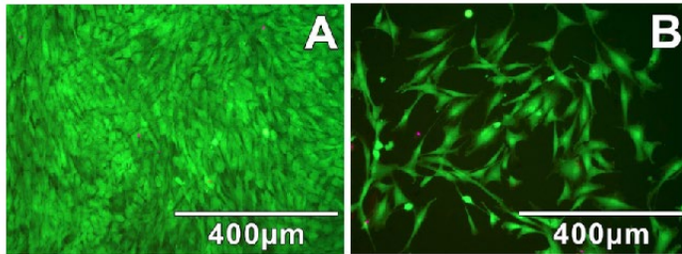


Figure 11: Results from LIVE/DEAD Staining Assays Indicate Both Mechanical and Enzymatic Dissociation Methods Produce Cultures of Healthy Adherent Cells. (A) Densely populated cultures of adherent cells isolated *via* enzymatic digestion; (B) More sparsely populated cultures of adherent cells isolated via mechanical digestion.

DAPI/Phalloidin staining assay

Images of cells stained with DAPI and phalloidin (Figure 12) were taken to gauge the morphology of adherent cells produced from both mechanical and enzymatic dissociation experiments. Staining with phalloidin and counter staining with DAPI allowed for clear identification of filamentous actin structures and nuclei of the cultivated adherent cells. Originally seeded at density of approximately 6.05×10^4 cells per well, enzymatically isolated cells proliferated rapidly over the three days they were in culture to form the dense sheet of cells seen in image A, Figure 12. Due to the slow proliferation rate of adherent cells gained from mechanical dissociation, the entire population of cells isolated from the successful “Mechanical 2” culture and supernatant culture were pooled and seeded into 20 wells of a 24-well plate (Figure 12, B). Those cells isolated using mechanical dissociation have not proliferated to the density as seen in image A. Instead, they present as more stellate structures, each cell covering a far increased area compared to those in image A.

Results from live/dead staining assays using calcein-AM and ethidium homodimer-1 stains indicate that enzymatic and mechanical dissociation methods both display capacity to produce healthy cultures of adherent cells from samples of mammalian tissue (Figure 11). Furthermore, those cultures of adherent cells

remain healthy, following repeated splitting and re-seeding.

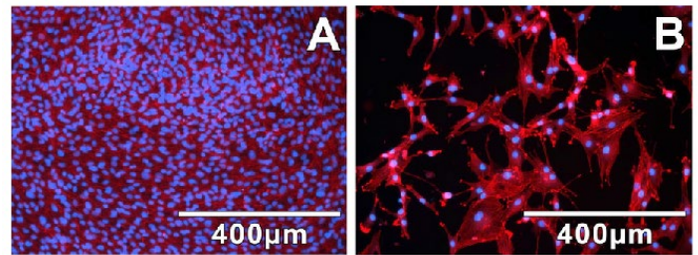


Figure 12: DAPI/Phalloidin staining clearly shows the morphology and density of adherent cells isolated by enzymatic. (A) and mechanical (B) dissociation methods

Discussion

The limitations of currently available treatments for periodontitis lie with the difficulty surrounding regeneration of the complex array of tissues that constitute the periodontium [36]. Hence, attention has turned to stem cell therapies, thanks to their abilities to differentiate into a wide variety of cell types, including those bone, ligament and cementum cells which form the supportive elements of the healthy periodontium [13].

Varying mechanical methods have been proposed over recent years, each aiming to produce comparable cell suspensions to those gained from tried and tested enzymatic protocols [33,37]. Development of these mechanical methods has been driven by the desire to eliminate enzymatic requirement for both health and economic reasons [32.] The possibility for eliminating the necessity of overnight incubation with expensive enzymes acts as a considerable financial and time saving incentive. For example, the enzymatic digestion protocol used in this study required tissue samples to be exposed to enzymes for a total of 25.5 hours, whereas mechanical dissociation of a single tissue sample takes less than half-an-hour.

In this study, skeletal muscle-derived cell suspensions modeling NCSCs gained from novel mechanical mincing and enzymatic digestion protocols were compared for the first time. The results show how spheroids and adherent cells similar in morphology to neurospheres and resultant NC-SCs isolated from heterogeneous suspensions of NCCs [17,25], can be derived from samples of mammalian skeletal muscle treated with both mechanical and enzymatic techniques.

Within this *proof-of-concept* study, we used multiple state-of-the-art methods to test the structural and visual properties of skeletal muscle-derived cell suspensions modeling NCSCs.

The handling and preparation of skeletal muscle-derived

cells was performed analogous to the preparation of oral-derived NCSCs as detailed in publications of our research group [25-27,29,30].

Before addressing the hypothesis and research question of this project; data from this report and similar research must be considered to assess whether mechanical dissociation is a viable alternative to enzymatic digestion for the isolation of therapeutic stem cells. Firstly, analysis of cell suspensions immediately following mechanical or enzymatic dissociation was carried out using the Countess Automated Cytometer. This assessment highlighted samples treated with mechanical dissociation as producing an extremely significant increase in liberated living cell populations, compared to those samples treated with enzymatic digestion. Further interpretation of these results revealed, on average, mechanical dissociation produced living cell populations 26.8 times those produced by enzymatic protocols (Table 1).

However, despite indications from similar research suggesting non-enzymatic dissociation yields 2.7-times the number of mesenchymal stem cells compared to collagenase-based dissociation [38], the living cell populations recorded using automated cytometry from this project proved inconsistent following microscopy and cell culture results.

Increased populations of living cells did not correlate with an increase in successful cell cultures. Only 8.3% of mechanical cultures produced spheroids and subsequent adherent cells, whereas 44.4% of tissue samples treated with enzyme digestion produced cultures of adherent cells, despite equal masses of starting material (see Figure 3) and culture conditions. Whilst tissue samples treated with enzymatic digestion produced cell suspensions containing larger fragments of muscle tissue, those cell suspensions treated with mechanical dissociation consisted of significantly finer tissue fragments.

Following dissociation, successful cell cultures seeded with standard cell medium (20% Foetal Bovine Serum, 1% L-Glutamine, 1% Penicillin-Streptomycin, 78% Dulbecco's Modified Eagle's Medium – High Glucose, 0.00005% FGF) began to form spheroids and subsequent adherent cells. Large numbers of spheroids appeared in enzymatic dissociation cultures 6-8 days after seeding cell suspension (see A and B, Figure 8; and 2, Figure 9). In comparison, the only successful mechanical dissociation culture produced fewer numbers of spheroid structures 12 days following seeding; identified within the tissue debris caused by mechanical dissociation (A, Figure 8).

The formation of spheroids and subsequent migration of adherent cells out of these spheroids is mirrored in the formation

of neurospheres as seen in studies focusing on NC-SCs isolated from the palate following culture with FGF enriched medium [25,27]. Additionally, research focusing on the enzymatic isolation of multipotent stem cells from PDL documented spheroid production over a similar 7-day course from cell suspensions [31]. However, unlike the studies mentioned above, spheroids cultured in this project were not cultivated in serum-free conditions.

A potential of obtaining cell cultures with neural crest resemblance (neural crest-derived stem cells [NCSCs]) from dental-related tissues, has been discussed in the literature. However, most reports include the use of serum-rich conditions and do not describe the potential for periodontal regeneration, slowing translation to the clinic. Therefore, we aimed to culture and characterize skeletal muscle-derived cell suspensions modeling NCSCs and evaluate their ability to differentiate into periodontal-like cells. Cultures were established from a varied cohort of rat tissue samples and grown, as monolayers, serum-free, and under sphere-aggregation conditions to induce and identify a NCSC-like phenotype. We have characterized oral-derived NCSC cultures from different tissue sources by immunocytochemistry and reverse transcription quantitative polymerase chain reaction in our earlier publications [25-27,29,30]. Here, we tried to handle and prepare skeletal muscle-derived cells performed analogous to the preparation of oral-derived NCSCs. Indeed, skeletal muscle-derived monolayer cultures expressed stem-related markers as shown by Mellow et al. 2019 [35] and by Michell et al. [28]. Our results describe the establishment of skeletal muscle-derived monolayer culture in serum-free medium modeling NCSC that differentiate into periodontal-like cells, as well as an important effect of serum free media in enhancing the cell phenotype and differentiation comparable to oral-derived NCSCs.

Additionally, research papers investigating multipotent cells present within the adipose tissue of mice isolated and cultured mesenchymal surface antigen-positive/CD271-positive adherent cells in similar culture media as used in this study [39,40]. Those isolated adherent mesenchymal cells exhibit similar morphology and proliferation rates to adherent cells isolated via both enzymatic and mechanical process in this project. Despite this indicative evidence, this is not sufficient to classify these adherent cells as stem cells. There are three sets of criteria that cells must fulfil to classify as mesenchymal stem cells. Designed by the Mesenchymal and Tissue Stem Cell Committee, these criteria have been implemented in order to standardize classification of human mesenchymal stem cells between studies [41,42]. Cell cultures isolated from this project using mechanical and enzymatic protocols already fulfil the first standard for classification, as cells

have shown a preference for adhering to the surface of plastic culture flasks under standard culture conditions. However, further assessment is required into specific expression of surface antigens and the potential for cells to display multipotent differentiation [41]. 95% of the suspected mesenchymal stem cells must show expression of surface antigens including CD105, CD90 and CD73 [41]. Furthermore, $\leq 2\%$ of cells should positively express antigens CD45, CD34, CD14/CD11b, CD79-alpha/CD19, or HLA-DR [41]. Finally, multipotent differentiation into adipocytes, chondroblasts and osteoblasts must be displayed in-vitro, under the standard conditions for differentiation [41].

Alternatively, identification of mesenchymal stem cells based on positive expression of cell surface antigen CD271 has been shown as an effective method of classification, both immediately after isolation and following cell culture [42]. Previous research has proven positive screening for antigen CD271 using CD271 microbeads together with MACS is an effective method for the isolation of mesenchymal stem cells from samples of human adipose tissue [43].

Development of a novel cleaning procedure was required to facilitate the continued use of the single-use sterile Rigenera Cons mincing apparatus. The enzyme activated detergent Tergazyme was utilized as the main component of the washing protocol. Given the enzymatic activity of this detergent, a cytotoxicity assessment was required to assess suitability with this project. Fibroblasts compared against a control group of an equal concentration were passed through the washed Rigenera Cons mincer and seeded at the same concentration. The resultant cell culture showed no difference in proliferation or cell density in comparison to the control group. Therefore, this novel cleaning procedure proved efficacious at facilitating the cleaning and reuse of the required mincing apparatus, without impacting cell proliferation or survival.

Fluorescent microscopy represented the final assays conducted during this project. The results from live/dead staining using fluorescent markers calcein-AM and ethidium homodimer-1 further display how the adherent cells cultivated from cell suspensions produced by both mechanical and enzymatic cultures were healthy and proliferative.

DAPI/Phalloidin staining revealed additional morphological information about adherent cells isolated from mechanically and enzymatically produced cell suspensions. Despite differences in density of the imaged cells, morphology of both enzymatic and mechanical isolated adherent cells showed similarities to actin-stained examples of undifferentiated human MSCs from previous studies [44]. Furthermore, these stained cells showed similar

characteristics to those adherent NC-SCs isolated from ovine palate, imaged with nuclear stain DAPI and the neural crest marker Nestin [25].

Future directions

Stem cells continue to hold an exciting potential for the most complete treatment of periodontitis, with MSCs and NC-SCs remaining a focus in this field of research. Advancements are being made towards the culture of stem cells in xeno-free conditions, with use of reagents including Gentle Cell Dissociation Reagent proving useful in studies surrounding human pluripotent stem cells [45,46]. The ease of harvest with minimal donor morbidity, and plentiful access, makes palatal tissue a convenient source for autologous NC cell-based therapies for regenerative periodontal purposes. Optimization of these methods has the potential to offer unprecedented opportunities to further bring effective regenerative therapies at the periodontal surgery in a widely variable applications in oral wound healing, and periodontal plastic-reconstructive fields. At this point in time, the limited characterization of the processed material and the inconsistent methods to determine the effect of NC cell transplant represent a limitation for the enormous therapeutic potential of oral-derived neural crest-derived stem cells in a plethora of regenerative applications. Therefore, increased efforts to achieve optimized NCSCs isolation yield and more standardized methods for palatal tissue manipulation for clinical purposes and analysis of grafting are needed.

Conclusion

Cells with characteristics of putative mesenchymal stem cells were found in regenerating periodontal tissues, implying their involvement in periodontal regeneration. Although periodontal-derived neural crest-derived stem cells have been shown to possess a wide therapeutic range *in vivo*, there is still no standard protocol with uniform parameters for the isolation and also unclear aspects about the identity of cells isolated from periodontal and/or from palatal tissues. The formation of spheroids and subsequent migration of adherent cells out of these spheroids is mirrored in the formation of neurospheres as seen in studies focusing on NC-SCs isolated from the palate following culture with FGF enriched medium [25,30]. Publication of study outcomes, comparative studies as well as standardization of cell products will allow the field to bring further effective therapy to the clinics. Mechanical dissociation has proven a viable non-enzymatic method of producing cellular suspensions from skeletal muscle, capable of forming adherent cells; the first classification required for identification as stem cells. Furthermore, mechanical dissociation also produced cell suspensions containing considerably higher

populations of living cells compared to enzymatic digestion. Rat muscle tissue has proven capable of producing large number of spheroids following enzymatic digestion protocols, with similar proliferation and migration as displayed in PDL-SCs [31] and NC-SCs [25,27]. This data further reinforce reports of skeletal muscle containing a variety of cells displaying mesenchymal stem cell characteristics [47]. In order to classify isolated adherent cells from this study, colonies should be tested for positive presentation of stem cell surface antigens CD105, CD90 and CD73. Furthermore, assessment of multipotent differentiation into adipocytes, chondroblasts and osteoblasts under standard conditions would confirm stemness of isolated adherent cells [41]. Alternatively, availability of availability of CD271 microbeads used in conjunction with MACS would facilitate identification of cells within the cell suspension. Considering the morphological similarity of spheroids and adherent cells to neurospheres and NC-SCs, testing adherent cells for NC-SC markers p75 (CD271), Nestin and Sox2 would also be appropriate [27]. Finally, the therapeutic suitability of isolated adherent cells could be assessed using a suitable animal model. Moreover, a necessary condition for further clinical translation is represented by standardization of the procedures as well as of the clinical results of the transplantation studies. Publication of study outcomes, comparative studies as well as standardization of cell products will allow the field to bring further effective therapy to the clinics.

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