Research Report

The hernia sac—A suitable source for obtaining mesenchymal stem cells☆☆☆

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A B S T R A C T

Background: Inguinal hernia sac, extended tissue from peritoneum, gradually enlarged in size with hernia disease time and prolapsed tissue volume. We hypothesize that mesenchymal stem cells are present in the development of hernia sac. The current study aimed to test the hypothesis that hernia sac, which is often resected and discarded as medical waste, contains mesenchymal stem cells and thus might be a suitable source to harvest mesenchymal stem cells.

Methods: Between July 2019 and June 2020, 4 hernia sacs were resected during hernia surgery and then obtained for mesenchymal extraction using the Miltenyi gentleMACS Dissociator. The presence of mesenchymal stem cells was determined by the markers CD105, CD73, and CD90, with assessment of the expressions ≥95%, whereas markers CD45, CD34, CD11b, CD19, and HLA-DR were used to assess lack expression (≤2%). Moreover, von Kossa staining, Alcian blue staining, and Oil Red O staining were used to verify the cells' ability for differentiation.

Results: Cells retrieved from the hernia sacs displayed a spindle-shaped morphology and exhibited adherence to plastics. The cell surface immunophenotypic profile was confirmed using surface markers APC-A (CD73), FITC-A (CD90), and PerCP-Cy5-5-A (CD105), with results showing 100%, 100%, and 99.2%, respectively, strongly indicating the presence of mesenchymal stem cells. Moreover, staining of in vitro cell cultures showed in vitro differentiation of precursor cells into osteoblasts, adipocytes, and chondroblasts, suggesting positive differentiation ability and identification of mesenchymal stem cells.

Conclusion: Inguinal hernia sac is a novel source of mesenchymal stem cells that can be easily obtained and stored for future usage.

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INTRODUCTION

Mesenchymal stem cells (MSCs), defined as multipotent stromal cells that can mature or differentiate into various cell types, have a myriad of clinical applications in tissue engineering, regenerative surgery, MSC-rich tissue scaffolding, etc. These cells can be found in a variety of places throughout the body, such as the bone marrow, umbilical cord tissue, adipose tissue, developing tooth bud of the mandibular third molar, and amniotic fluid [1–5]. The most common source for obtaining MSCs postnatally has been from the umbilical cord, including umbilical cord blood cells, amnion/placenta, umbilical cord vein, or umbilical cord matrix cells [6]. Among adults, however, MSCs are commonly derived from the bone marrow [3]. Other sources include adipose tissues, which could be a much easier and more inexpensive source compared to the bone marrow [7]. However, several other human body parts have been found to contain mature MSCs, with a recent study showing the peritoneum as one such source for MSCs. Peritoneal MSCs can be separated from peritoneal or peritoneal dialysis fluid [8,9]. Hernia sacs originate from peritoneum and gradually downward developed in size with hernia disease time and prolapsed tissue volume. The sac is often respected during hernia surgery procedures. It is our hypothesis that hernia sac contains MSCs as to develop the gradually larger pockets to contain and hang up the heavier and heavier weight of prolapsed herniated tissues in hernia sac.

If the hypothesis is proven, patients with inguinal hernia could have one more choice to store their MSCs. Instead of discarding the hernia sac as medical waste, surgeons could preserve the excised hernia sac without additional surgical procedures. MSCs from hernia sacs are easier and more practical to obtain compared to current common sources, such as umbilical cord tissue or bone marrow [10,11]. This study aimed to confirm whether MSCs can be harvested from hernia sacs discarded following hernia surgery.
MATERIALS AND METHODS

Cell Dissociation From Hernia Sacs. Surgery for adult inguinal hernia (Fig 1) involves performing the open method with inguinal incision. Fresh hernia sacs obtained from 4 randomly selected inguinal hernia surgeries were harvested and immediately sent to the laboratory. Details regarding these 4 patients are summarized in Table 1. MSCs were harvested within 48 h. Peritoneal tissue from hernia sac was first isolated and cut into small pieces, after which the mesenchymal cells within them were obtained using the Miltenyi gentleMACS Dissociator, which has been used for the semiautomated dissociation of tissues into single-cell suspensions or thorough homogenates (www.miltenyibiotec.com). The dissociated suspension was filtered through 40-μm strainer (BD Falcon) and cultured in alpha-Minimal Essential Medium (Thermo) plus 10% fetal bovine serum (Hyclone) at 37 °C, 5% CO2 and humidified atmosphere. This study had been reviewed and approved by institutional review board of Tung’s Taichung Metro Harbor Hospital. All patients/participants provided written informed consent prior to participation (IRB #109060).

Cell Identification and Morphology. The presence of MSCs was determined by using the following minimum criteria for defining multipotent MSCs according to The International Society for Cellular Therapy [12]: (1) adherence to plastic, (2) specific surface antigen (Ag) expression, and (3) multipotent differentiation potential.

MSCs were placed in plastic culture plates to observe their morphology and ability for plastic adherence when maintained in standard culture conditions using tissue culture flasks. The hematopoietic cells, mature adipocytes, etc, would not attach to the plastic plate in this condition. Positive markers selected for our studies were based on existing literature and antibody availability for human MSCs [12]. The markers CD105, CD73, and CD90 were used to assess expressions ≥95%, whereas markers CD45, CD34, CD11b, CD19, and HLA-DR were used to assess lack of expression (≤2%). All positive CD markers used (CD105, CD73, and CD90) had an expression of ≥95% when tested.

Cell Morphology and Differentiation. When MSCs grew to 90% confluent, the medium was shifted to differentiation medium for the assay of trilineage capability. The differentiation media were StemMACS Osteodiff medium, StemMACS Chondrodiff medium, or StemMACS Adipodiff medium, respectively (Miltenyi Biotec). After 3 weeks, cells underwent von Kossa staining to verify osteogenesis, Alcian blue stain to verify ability to undergo chondroblasts, and Oil Red O staining to determine adipogenesis.

RESULTS

This study first assessed the morphology of the cells retrieved from hernia sacs obtained following inguinal hernia surgeries and their ability to adhere to plastics. All cells from the 4 patients displayed a spindle-shaped morphology and exhibited adherence to plastics (Fig 2), partly indicating the presence of MSCs [13].

Determining Reliable Positive Markers for Mesenchymal Stem Cell Identification. The cell surface immunophenotypic profile, along with the following positive CD markers, was determined: CD73 (100%), CD90 (100%), and CD105 (99.2%). The following negative CD markers were also assessed to further verify the presence of MSCs: CD11b, CD19, CD34, CD45, and HLA-DR (Fig 3). Accordingly, all negative CD markers tested (CD11b, CD19, CD34, CD45, and HLA-DR) had an expression of ≤2% (Fig 3), strongly suggesting the presence of MSCs.

Differentiation Ability of Cells. The in vitro differentiation of precursor cells into osteoblasts, adipocytes, and chondroblasts was determined through staining of in vitro cell cultures. Von Kossa staining confirmed

Table 1

Summary of patients’ characteristics.

| Case 1 | Case 2 | Case 3 | Case 4 |
|--------|--------|--------|--------|
| Age    | 54     | 36     | 71     | 42     |
| Sex    | Male   | Male   | Male   | Male   |
| Height | 165    | 166    | 156    | 174    |
| Weight | 70     | 77     | 56.2   | 87     |
| Smoking| No     | No     | No     | No     |
| Hernia size | 32 × 14.5 × 2.5 cm | 6 × 4 × 8 cm | 2 × 3 × 5 cm | 3 × 3 × 5 cm |
| Medical history | Hepatitis B, hypertension | No | Prostate cancer | Lung arteriovenous aneurysm, scrotal spermatic varicocele |
| Surgical history | No | No | Robot-assisted laparoscopic radical prostatectomy | Embolization |

Fig 1. Hernia sac and inguinal hernia.
differentiation into osteoblasts, Oil Red O staining demonstrated differentiation into adipocytes, and Alcian blue staining confirmed differentiation into chondroblasts [12]. Figure 4 revealed positive differentiation ability and identification of MSCs following von Kossa, Alcian blue, and Oil Red O staining (Fig 4). The uninduced cells showed no sign of differentiation.

DISCUSSION

The current study demonstrated, for the first time, that hernia sacs removed during inguinal hernia surgeries can be a good source for MSCs in terms of both quality and quantity. The morphology of the cultured cells (Fig 2), as well as the different types of staining used to identify the cells’ ability to differentiate, was indicative of MSCs. As observed in the images following von Kossa, Oil Red O, and Alcian blue staining, all 3 were indicative of adipogenesis, chondrogenesis, and osteogenesis, respectively (Fig 4). This further confirmed that MSCs are abundantly present in adipose tissue derived from hernia sacs obtained following inguinal hernia repair surgeries [14]. The most common source for MSCs postnatally has been from the umbilical cord, which includes the umbilical cord vein, blood cells, amnion/placenta, or matrix cells [6]. In the absence of umbilical cord stem

| Surface Marker | %  |
|---------------|----|
| CD 73         | 100|
| CD 90         | 100|
| CD 105        | 99.2|
| Negative (CD11b, 19, 34, 45, HLA-DR) | 1.7 |

Fig 3. Positive and negative markers used to identify the presence of mesenchymal stem cells in the hernia sac.
cells after birth, the bone marrow can be another common source of MSCs. However, bone marrow MSCs are particularly difficult to harvest in most patients and require invasive procedures. The present study confirmed that hernia sacs could provide enough volume of mesenchymal tissues with sufficient quality. In the current era, short- or long-term storage facilities for hernia sacs removed following surgery are widely available worldwide. As such, these MSCs could be harvested and stored for further or additional practical use (e.g., tissue engineering; tissue expansion; MSC-rich tissue scaffolding; and MSC cultures for functional tissues, such as insulin-secreting tissues, cartilage, and bone). Therefore, rather than discarding the entire hernia sac as a medical waste, it could be harvested and stored for future isolation of MSCs to be used for replacing and restoring damaged or dysfunctional cells. MSCs could also be engineered to deliver therapeutic proteins.

The current study suggested the presence of MSCs in the hernia sac excised following surgery, which can be used for a variety of important tasks, such as regenerative medicine and disease therapeutics [15–17]. Some aged patients who develop a hernia present with degenerative diseases, such as osteoarthritis of knee or coronary heart diseases, which may benefit from MSC treatment [18]. As such, future studies aimed at treating these degenerative diseases using MSCs harvested from the autologous hernia sac are warranted. MSCs can also be easily isolated during a variety of surgical procedures. These approaches are safer and can obtain larger amounts of stem cells compared to that used for extracting bone marrow MSCs. One such source is adipose tissue removed during liposuction and resected fat, which are feasible sources of MSCs for regenerative medicine [19,20]. However, given that adipose tissue can also be found in the inguinal region of the body, new developments and observations indicate that hernia sacs removed during inguinal hernia repair surgeries can be a good source of MSCs.

Although other stem cells and bone marrow MSCs have many similar biological characteristics, some differences in their immunophenotype, differentiation potential, transcriptome, proteome, and immunomodulatory activity have been identified [10]. Some of these differences may represent specific features of bone marrow MSCs and other stem cells, whereas others are suggestive of the inherent heterogeneity [21]. Different isolation and culture protocols may induce different characteristics. MSCs displayed certain enhanced functionalities when derived from specific places [13]. Most importantly, however, despite the minor differences between these MSC populations, stem cells from some other sources seem to be as effective as those derived from the bone marrow for clinical application and may even promote better outcomes in certain applications [22–24]. Further studies are needed to explore the clinical application of MSC obtained from hernia sacs and compare their efficacy with MSCs from other sources. There are different methods of inguinal hernia surgery; basically, there are 2 routes to approach: open method and laparoscopic method. The current study applied open hernia surgery to harvest the hernia sac without additional procedures. There is no increase in surgical risk, operation time, or anesthesia loading. Different from throwing the waste, aseptic preservation should be alerted in the scrubbing table. For more extensive applications in the future, it is necessary to investigate further the harvesting of hernia sac in laparoscopic hernia surgery.

In conclusion, our study confirmed that hernia sacs could provide sufficient quality and quantity of MSCs, indicating that the hernia sac could be a promising source of MSCs for further tissue engineering, regenerative surgeries, and further possible applications in patients with hernia.

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AD: collection and assembly of data, data interpretation, manuscript writing, and final approval of the manuscript. MC: data analysis and interpretation, final approval of the manuscript. CH: data analysis and interpretation, manuscript editing, manuscript revision, and final approval of manuscript. All authors contributed to the article and approved the submitted version.

Disclosure

The authors have no disclosures and competing interests.
Conflict of Interest

The authors declare no commercial or financial relationships that could be construed as a potential conflict of interest.

Credit Author Statement

Alpha Dian-Yu Lin: collection and assembly of data, data interpretation, manuscript writing, and final approval of the manuscript. Min-Che Tung: data analysis and interpretation and final approval of the manuscript. Chin-Heng Lu: data analysis and interpretation, manuscript editing, manuscript revision, and final approval of manuscript. All authors contributed to the article and approved the submitted version.

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