A dipose stromal vascular fraction (SVF) has moved further into the focus of stem cell research, regenerative medicine, and fat grafting with the development of new industries worldwide. Tissue engineering involving adipose stromal vascular cells (SVCs) represents an interesting research field for different diseases, including degenerative, congenital, or traumatic conditions, and bone, articular, and soft-tissue defects. In plastic surgery, these cells have been used mostly to supplement fat grafts, improving graft retention and long-term outcomes.1–5

Adipose SVF consists of a heterogeneous, mesenchymal population of cells that includes not only adipose stromal, hematopoietic stem, and progenitor cells but also endothelial cells, erythrocytes, fibroblasts, lymphocytes, monocyte/macrophages, and pericytes, among others.4 SVF can be isolated by enzymatic methods, digestion with collagenase, and centrifugation. When compared with enzymatic methods, mechanical isolation required less time but yielded fewer cells. Both case-control studies reported improved volume retention with cell-supplemented fat grafts for breast reconstruction.

Conclusions: Mechanical isolation methods are alternatives to circumvent safety issues posed by enzymatic protocols. However, randomized comparative studies with long-term clinical outcomes using mechanically isolated stromal vascular cells are needed to identify their ideal clinical applications. (Plast Reconstr Surg Glob Open 2016;4:e1017; doi: 10.1097/GOX.0000000000001017; Published online 7 September 2016.)

**Background:** Standard isolation of adipose stromal vascular fraction (SVF) requires the use of collagenase and is considered more than “minimally manipulated” by current good manufacturing practice requirements. Alternatively, nonenzymatic isolation methods have surfaced using physical forces to separate cells from the adipose matrix. The purpose of this study was to review the literature on the use of mechanical isolation protocols and compare the results. The implication for use as a standard procedure in practice is discussed.

**Methods:** A systematic review of the literature was performed on mechanical isolation of SVF with a search of six terms on PubMed and Medline databases. One thousand sixty-six articles were subject to evaluation by predetermined inclusion and exclusion criteria.

**Results:** Two level 2 evidence articles and 7 in vitro studies were selected. SVF was isolated using automated closed systems or by subjecting the lipoaspirate to centrifugation only or by shaking or vortexing followed by centrifugation. Six articles reported isolation in laboratory settings and three inside the operating room. Stromal vascular cells expressed CD34, and CD44, CD73, CD90, and CD105, and differentiated along adipogenic and osteogenic lineages. When compared with enzymatic methods, mechanical isolation required less time but yielded fewer cells. Both case-control studies reported improved volume retention with cell-supplemented fat grafts for breast reconstruction.

**Conclusions:** Mechanical isolation methods are alternatives to circumvent safety issues posed by enzymatic protocols. However, randomized comparative studies with long-term clinical outcomes using mechanically isolated stromal vascular cells are needed to identify their ideal clinical applications. (Plast Reconstr Surg Glob Open 2016;4:e1017; doi: 10.1097/GOX.0000000000001017; Published online 7 September 2016.)
cient, this enzymatic isolation protocol involves the use of xenogenic components that may pose certain risks and safety issues, such as exposure to infectious agents and immune reactions. Thus, xeno-free enzymatic products have been used and shown that they can replace the current research grade products effectively without any negative effect in the yield or function of human adipose stem cells (ASCs). To circumvent the need for manual and external manipulation, single devices have been used to separate and concentrate SVCs from the adipose matrix, which may be mixed with fat to improve results in fat-grafting procedures. Such systems may decrease the risk of infections and operator dependency. Still, the complexity of current good manufacturing practice requirements has created many obstacles to the translation of enzymatic SVF isolation protocols, whether manual or automated, to clinical scenarios.

Nonenzymatic protocols have been attempted consisting of mechanically dissociating SVF using different devices or an automated closed system, resulting in ready-to-use SVF or SVF-supplemented fat. The cellular composition of SVF can differ according to the isolation protocols used and may have an effect on its capabilities of differentiation, angiogenesis, and regeneration.

This review article summarizes the published literature on nonenzymatic isolation of adipose SVF and compares both the techniques and the results. The purpose of this systematic review of the literature is to improve our understanding of the current, available mechanical protocols and to potentially provide guidance for improvements of the methods going forward.

**METHODS**

A comprehensive search of the Pubmed and MEDLINE databases was conducted in January 2016 using the following search terms: “isolation,” “dissociation,” “adipose,” “fat,” “stromal vascular fraction,” and “stem cells.” The inclusion criteria were studies in the English literature, documenting the use of mechanical methods for isolating SVF of human adipose tissue. Articles that described enzymatic methods or mechanical dissociation combined with enzymatic digestion to obtain SVF were excluded. Not only articles that described fat-processing methods lacking steps to specifically separate SVF but also those that used explant culture to extract only mesenchymal stem cells (MSCs) or isolated cells from the lipoaspirate fluid (infranatant or bottom layer) were excluded.

Data collected included the following: donor information (age, sex, and body mass index), fat-harvesting technique, processing techniques, characterization studies, such as multilineage properties of the isolated cells, phenotyping of markers associated with SVF, specific gene expression, and in vivo outcomes.

**RESULTS**

The primary search yielded 1,066 articles; of which, 754 titles passed initial screening. After duplicates were removed, 450 articles remained, and their abstracts were reviewed. The method sections of 278 articles were read in their entirety. Nine articles met our predetermined inclusion and exclusion criteria and were selected (Fig. 1). The journal types in which these articles were published were diverse. Four articles were published in plastic surgery journals, three articles in cellular therapy journals, and two articles in basic biology journals. Countries that contributed articles were Brazil (3), Italy (3), United States (2), and Russia (1). There were 2 prospective comparative studies of level 2 evidence and 7 basic science studies (Table 1). Summaries of the findings in the discussed articles are presented in Tables 2 and 3.

**Prospective Comparative Studies**

The effects of commercially available SVF-isolation methods on in vitro activity and in vivo outcomes were studied in 2 prospective comparative studies. Eighty-six patients underwent fat grafting for breast reconstruction with or without SVF enrichment, 26 of whom received injection of fat enriched with mechanically isolated SVF.

**Statistical Analysis**

A formal statistical analysis of the eligible studies was not performed because of the methodological heterogeneity and novel nature of these methods. A detailed systematic review and comparison of the diverse findings was undertaken instead.
Domenis et al.\textsuperscript{10} used an automatic system using filtration and centrifugation, the Fastem (CORIOS Soc. Coop, San Giuliano Milanese MI, Italy), to isolate SVF. Characterization of the mechanically isolated cells determined the presence of MSCs. Immunophenotyping studies indicated positivity for CD44, CD73, CD90, and CD105 and negativity for CD45. The cells expressed the pluripotent-linked genes, NANOG, Oct-4, Sox-2, and c-kit. These cells were used for fat-graft enrichment in patients undergoing breast reconstruction. The in vitro and in vivo results were compared with those of fat grafting enriched with enzymatically isolated SVF (Cytori Celution System; Cytori Ltd, Deeside, United Kingdom, and Lipokit Medikhan System; Medikan International Inc, Pusan, Korea) and nonenriched fat. No significant difference in the frequency of CD31\textsuperscript{+}, CD73\textsuperscript{+}, and CD90\textsuperscript{+} was observed among the 3 isolation methods. The proportion of CD45\textsuperscript{−}/CD31\textsuperscript{−}/CD34\textsuperscript{+} cells was lower in Fastem-enriched samples than in enzymatically enriched samples. In Fastem-isolated ASCs, the CD45\textsuperscript{−}/CD31\textsuperscript{−}/CD34\textsuperscript{+} fractions demonstrated reduced ability to differentiate along adipogenic, myogenic, and vasculogenic lineages compared with enzymatically isolated ASCs. The gain in breast subcutaneous thickness observed with Fastem cell–enriched fat was not significantly different from fat enriched with enzymatically isolated cells. After 12 months, significant improvement in volume maintenance was observed with enriched fat from both enzymatic and mechanical methods. This study was limited by the small sample sizes of each group (Fastem, n = 6; Cytori, n = 9; Lipokit, n = 5; nonenriched, n = 16).

Gentile et al.\textsuperscript{11} also used an automatic system with washing and filtration cycles, the Mystem (Mystem LLC, Wilmington, Del.) and Fastem, to isolate SVF. Cell yields from these 2 mechanical methods and 3 enzymatic methods (Cytori, Medikhan, and manual collagenase digestion) were compared. There was no further analysis of the isolated cell population throughout the methods. The focus of this study was on clinical outcomes in breast reconstruction with fat enriched by each method (Mystem, n = 10; Fastem, n = 10; Cytori, n = 10; Medikhan, n = 10; and nonenriched, n = 10). Enrichment with Fastem and Cytori provided significantly greater contour and volume maintenance compared with nonenriched controls. Comparison of outcomes between the different isolation methods was not reported. The authors observed the presence of oil cysts and cytosteatonecrotic areas at 12 months without specifying which method contributed the most to these complications.

### In Vitro Studies
Seven articles reported the in vitro properties of mechanically isolated SVF. The protocols involved manual separation of SVC population as opposed to separation in automated closed systems. Three articles described sub-

| References | No. of Donors for SVF | Level of Evidence | Journals |
|-------------|-----------------------|------------------|----------|
| Domenis et al\textsuperscript{10} | 6 | II | Stem Cell Research and Therapy |
| Gentile et al\textsuperscript{11} | 20 | II | Plastic and Reconstructive Surgery Global Open Biotechnology Letters |
| Markarian et al\textsuperscript{14} | 10 | — | Plastic and Reconstructive Surgery |
| Raposio et al\textsuperscript{19} | — | — | Cytotherapy |
| Condé-Green et al\textsuperscript{17} | 9 | — | Aesthetic Surgery Journal |
| Shah et al\textsuperscript{12} | 13 | — | Cytotherapy |
| Condé-Green et al\textsuperscript{13} | 10 | — | Bulletin of Experimental Biology and Medicine |
| Baptista et al\textsuperscript{12} | 13 | — | |
| Romanov et al\textsuperscript{16} | — | — | |

| Method | References | Stromal Vascular Cell Yield | Adipose-derived Stromal Cell Yield |
|--------|------------|-----------------------------|----------------------------------|
| Fastem | Domenis et al\textsuperscript{10} | n/a | n/a |
| Mystem | Gentile et al\textsuperscript{11} | \(=3\times10^4\) per mL | n/a |
| RBC lysis and centrifugation | Gentile et al\textsuperscript{11} | \(=8\times10^4\) per mL | n/a |
| | Baptista et al\textsuperscript{12} | \(240\pm7.4\times10^3\) per mL | 1.2 \(\pm\) 3.7 \(\times\) 10\(^4\) per mL |
| | Condé-Green et al\textsuperscript{17} | \(2.3\times10^5\) per mL | n/a |
| | Markarian et al\textsuperscript{14} | \(2.5\times10^5\) per 10 mL | n/a |
| | Markarian et al\textsuperscript{14} | \(7\times10^4\) per 10 mL | n/a |
| Centrifugation (800g \(\times\) 15 min) | Markarian et al\textsuperscript{14} | \(1.5\times10^4\) per 10 mL | n/a |
| RBC lysis, and centrifugation | Shah et al\textsuperscript{15} | \(2.5\times10^4\) per mL (after a mean of 13 d of culture) | n/a |
| Vigorous washing, hand-shaking, and centrifugation | Romanov et al\textsuperscript{16} | n/a | n/a |
| Dilution, vortexing \(\times\) 2 to 3 min, centrifugation | Romanov et al\textsuperscript{16} | n/a | n/a |
| Vortexing \(\times\) 3 min, RBC lysis, and centrifugation | Raposio et al\textsuperscript{18} | \(1\times10^7\) per 80 mL | \(5\pm1\times10^7\) per 80 mL |

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\textbf{Table 1. List of Retained Publications on Mechanical Isolation of Adipose-derived Stromal Vascular Fraction with Their Study Population and Journal Distribution}
Table 3. Summary of Articles on Mechanical Isolation of Adipose-derived Stromal Vascular Fraction Including the Mechanical Methods Used and the Relevant Findings

| References          | Mechanical Isolation Method                                                                 | Method Duration | Yield                                                                 | Differentiation | Gene Expression | Immunofluorescence | Impact of SVF on Fat Grafting |
|---------------------|-----------------------------------------------------------------------------------------------|----------------|------------------------------------------------------------------------|-----------------|-----------------|---------------------|-------------------------------|
| Domenis et al10     | Automatic filtration, centrifugation (1,700 rpm × 10 min) (Fastem)                           | n/a            | Cell yield with Fastem < Lipokit < Cytori                             | n/a             | Adipogenic      | Myogenic Vascuogenic      | Increased subcutaneous thickness in breast in all enriched grafts Contour/volume maintenance at 12 mo: Fastem (52±4.6%) Mystem (43±3.8%) Fastem and Cytori greater volume maintenance |
| Gentile et al11     | Automatic washing, filtration cycles (Mystem) and Automatic filtration, centrifugation (1,700 rpm × 10 min) (Fastem) | n/a            | Cell yield with Mystem< Fastem (p<0.01)                               | n/a             | n/a             | n/a                 | n/a                          |
| Raposio et al18     | Vibration (6,000/min × 6 min), centrifugation (1,600 rpm × 6 min), and collection of pellet | ≈15 min        | 1 × 10^7 SVC per 80 mL 5 × 1 × 10^6 ASCs per 80mL CD45−/CD31−/CD34− and CD90+ cells present | n/a             | n/a             | n/a                 | n/a                          |
| Condé-Green et al13 | Centrifugation at high speed, vortexing × 5 min, or centrifugation and RBC lysis            | n/a            | CD45/CD34: 6–13% CD14: collagenase < mechanical CD31: collagenase > mechanical Collagenase > mechanical Collagenase > trypsin | n/a             | n/a             | n/a                 | n/a                          |
| Markarian et al14   | RBC lysis of lipoaspirate, centrifugation (600g × 10 min), resuspension in DMEM + FBS      | n/a            | Viable cells isolated mechanically after additional centrifugation step did not proliferate after 14 d | n/a             | n/a             | n/a                 | n/a                          |
| Shah et al15        | Vigorous washing and hand-shaking, centrifugation (1,200 rpm × 5 min), and resuspension    | ≈1 h via mechanical ≥3h for collagenase | 19-fold fewer cells via mechanical method CD29 (48.5 ± 32.0) CD44 (4.8 ± 2.9) CD73 (8.8 ± 6.4) CD90 (23.2 ± 24.5) CD105 (3.9 ± 3.5) CD34 (25.7 ± 21.2) Percentage of SVC Decantation: CD45/CD31 (3.8 ± 3.1) CD45/CD34 (2.8 ± 2.7) CD45/CD105 (2.9 ± 2.4) Centrifugation: CD45/CD31 (2.9 ± 0.7) CD45/CD34 (2.3 ± 1.7) CD45/CD105 (2.1 ± 1.5) Pellet: CD45/CD31 (7.1 ± 3.6) CD45/CD34 (4.3 ± 2.3) CD45/CD105 (4.7 ± 1.6) | Adipogenic: 13.6% Osteogenic: 65.6% | n/a             | n/a                 | n/a                          |

(Continued)
jecting lipoaspirate to centrifugation, and four articles described primarily using shaking by hand or electronically or vortexing plus centrifugation to obtain SVCs.

Centrifugation

Baptista et al\(^{12}\) isolated SVF by performing red blood cell (RBC) lysis of the lipoaspirate followed by centrifugation at 900g for 15 minutes and resuspension of the SVF-containing pellet. A mean of 24.0 ± 7.4 × 10^4 mechanically processed lipoaspirate cells per millimeter were obtained, 1.2 ± 0.37 × 10^4 cells per millimeter of which showed plastic adherence. The latter cells were CD45⁻, CD73⁻, CD31⁺, CD44⁺, CD90⁺, CD105⁺, and CD34⁺. This method was compared with manual enzymatic isolation, which showed a greater yield of total and plastic-adherent cells (58.4 ± 17.8 × 10^4 cells per millimeter and 8.5 ± 6.7 × 10^4 cells per millimeter, respectively) but required a greater amount of time. They also performed cryopreservation at −196°C, which was associated with a decreased cell yield.

Condé-Green et al\(^{13}\) isolated SVF after the previously described method by Baptista et al\(^{12}\) and compared the population of mechanically processed lipoaspirate cells in centrifuged and decanted lipoaspirates. They looked at the population of cells isolated mechanically from common fat-processing methods to be used in fat grafting. Samples of each processed lipoaspirate and the pellet issued from centrifugation were analyzed. The pellet showed a significantly higher quantity of MSCs and endothelial cells than the other processed samples. However, there was no comparison with enzymatic isolation and limited characterization and differentiation of cells.

Markarian et al\(^{14}\) reported 3 modified versions of the mechanical isolation method described by Baptista et al\(^{12}\). The first one subjected lipoaspirates to RBC lysis, centrifugation at 600g for 10 minutes. The second and third methods added a centrifugation step at 800g and 1,280g, respectively, for 15 minutes followed by RBC lysis, centrifugation at 600g for 10 minutes. Viable cells were isolated from all 3 methods; however, those from the second and third protocols did not proliferate after 14 days. These 3 mechanical methods were compared with collagenase and trypsin isolation methods. Collagenase isolation provided a significantly greater cell yield than trypsin digestion or the 3 mechanical isolation methods. However, no significant difference in cell yield was observed between the first mechanical isolation and trypsin digestion. No further analysis was performed on mechanically isolated SVCs because of limited growth in cell culture.

Shaking or Vortexing and Centrifugation

Shah et al\(^{15}\) performed repeated cycles of washing with PBS, vigorous hand shaking, and centrifugation at 1,200rpm for 5 minutes to isolate SVF. This method was compared with collagenase-based isolation. In non-enzymatically isolated samples, there was an increase in CD45⁻ cells with a decrease in expression of markers for both MSCs and ASCs. However, the changes in the phenotype were not clear since no significant difference was noted for CD29 between enzymatically and nonenzymatically isolated SVF. A significant increase in CD44⁺ was
observed in nonenzymatically isolated SVF. At passage 0, there were significantly greater numbers of MSC markers, based on phenotype, for nonenzymatically isolated cells. The increase in MSC markers correlated with a decrease in contaminating hematopoietic cells, as evidenced by decreases in cells positive for CD34 and CD45. These cells demonstrated comparable adipogenic and osteogenic differentiation to enzymatically isolated cells. Mechanical isolation required, at most, a third of the time required for collagenase isolation yielding 19-fold fewer cells. This study only reported flow cytometry but no additional studies on freshly isolated SVF.

Romanov et al\textsuperscript{16} isolated SVF by diluting lipoaspirate, vortexing for 2 to 3 minutes, centrifuging at 600g for 10 minutes. Cultured cells homogenously expressed anti-\textalpha smooth actin and produced type 1 collagen and fibro-nectin. Endothelial markers, high positivity for human leukocyte antigen-1, and vimentin were noted. Cells differentiated toward adipogenic, osteogenic, and neurogenic lineages. Analysis of freshly isolated SVF and comparison with enzymatic isolation were not performed.

Condé-Green et al\textsuperscript{17} isolated SVF using 2 different mechanical methods, subjecting lipoaspirate to centrifugation or vortexing for 3 minutes followed by centrifugation. Overall cell yield from centrifugation was double that of vortexing, but high cell viability was observed with both methods. Comparison was performed with collagenase digestion. SVC yield from collagenase isolation was 10-fold greater than that from centrifugation. In addition, mechanically isolated SVF contained a greater proportion of hematopoietic cells and monocytes/macrophages and fewer ASCs and endothelial cells. There was a lack of quantitative reporting of flow cytometry findings and further in vitro studies as the authors published this study as an abstract with few details on the entire protocol. The findings are nonetheless still relevant and were referenced in related studies.

Raposio et al\textsuperscript{18} reported isolation of SVF in the operating room, submitting lipoaspirate to vibration in a shaker for 30 minutes. Lipoaspirate was then centrifuged at 1,600 rpm for 6 minutes. A mean of $1 \times 10^7$ SVCs for 80 mL lipoaspirate was obtained, 5% of which were ASCs. There was no comparison with other isolation methods, only limited in vitro characterization and no in vivo outcome measurements, thus limiting conclusions of this method’s clinical benefits.

**DISCUSSION**

Enzymatic SVF isolation is standard but limits the volume of lipoaspirate to be processed as it is a lengthy process. In vitro manipulation with costly enzymes. Adipose tissue exposed to collagenase has also been considered more than “minimally manipulated,” which has been defined by U.S. Food and Drug Administration guidance documents as “processing that does not alter the original relevant biological characteristics of cells.”\textsuperscript{19} Furthermore, when considering the clinical use of SVCs, one must be cognizant of the fact that the use of tissue processing and cell-based product use pose risks for contamination and damage to cells.\textsuperscript{20} Numerous mechanical cell isolation systems have been developed and commercialized\textsuperscript{21,22} (Fig. 2). However, the small number of published studies limits their use and credibility. Our analysis of the 9 articles describing mechanical SVF isolation methods has shown promising results. These methods require substantially less time to perform than enzymatic methods.\textsuperscript{12,15} Although enzymatic digestion yields significantly more SVCs, mechanical protocols isolate a similar population of cells with great cell viability and pluripotency.\textsuperscript{15,16} Centrifugation achieved the highest cell yield\textsuperscript{12} followed by vortexing and centrifugation,\textsuperscript{17} followed by manual shaking.\textsuperscript{13} Some protocols were performed inside the operating room,\textsuperscript{10,11,18} suggesting potential easy application to clinical practice. Automated mechanical devices, such as Fastem and Mystem, as opposed to manual manipulation, isolate cells entirely within a single device and were used to enrich fat in patients undergoing breast reconstruction, resulting in significant volume maintenance improvement.\textsuperscript{10,11}

Despite these promising results, determining the most efficient mechanical isolation method among these studies is challenging, as a diverse range of mechanical forces was featured throughout the methods and only three articles directly compared different mechanical isolation methods. Repeated mechanical manipulation of lipoaspirate may negatively impact cell yield\textsuperscript{14} and growth.\textsuperscript{14} Mechanically isolated SVF seems to contain a larger proportion of CD45$^+$ cells, representing an increase in potentially contaminating cells, and a lower number of CD34$^+$ cells representing ASCs, which could negatively impact fat-grafting outcomes.\textsuperscript{10,15,17} Some studies have suggested that CD34$^+$ cells play an important role in promoting fat-graft retention, highlighted by their high degree of proliferation in the first 2 to 4 weeks after fat grafting.\textsuperscript{23,24} In addition, only 3 studies reported the ASC yield as opposed to the total SVC yield.\textsuperscript{12,15,18} ASCs have been proposed to play important roles in fat-graft retention because of their immunomodulatory, angiogenic, and multipotent characteristics.\textsuperscript{25,26} However, it is important to note that other cell types in the heterogeneous SVC population contribute important interactions, which contribute to favorable outcomes.

There is a lack of comparison between the automated, mechanical isolation methods, Fastem and Mystem, and with enzymatic isolation in terms of clinical outcomes and complications. These comparisons are needed to demonstrate if the improved cell yield and cell population composition observed with enzymatic methods translate to improved outcomes and justify the increased time and cost of these equipment (Table 4).

Adipose SVF offers tremendous potential for aesthetic and reconstructive applications. In specific circumstances, patients having autologous fat transfer may benefit greatly from enrichment of fat with SVCs to accelerate the regeneration process, further improving outcomes of the procedure.\textsuperscript{27,28} Mechanical processing of lipoaspirate with steps other than standard electronic centrifugation\textsuperscript{29} has been described to further fragment adipose tissue particles, aiming to improve fat-graft take. Tonnard et al\textsuperscript{30} used repeated shear forces, passing lipoaspirate between 2 syringes 30
times followed by filtration, to obtain a product with altered adipocytes and intact viable MSCs. Bianchi et al\textsuperscript{31} used a closed, handheld system using filtration with simultaneous emulsification and washing of the lipoaspirate. Although these methods have been demonstrated to maintain or increase the proportion of ASCs in the resulting adipose tissue product to be grafted, there are no reports of isolating SVF as a final product through these methods but rather a ready-to-use fat-grafting material instead.

Concerns regarding the safety of collagenase, centered around potential residual enzyme activity after injection, have not been supported by the published literature. Therefore, mechanical isolation methods most likely do not provide additional safety benefits through the avoidance of xenogenic enzyme. To ensure the highest possible safety for patients, a precisely defined procedure with high-quality control is required instead. Automation of the process within a closed system separates the cells from the external environment and reduces opportunity for error and contamination, further improving the safety of the isolation procedure. Moving forward, we must incorporate principles of evidence-based medicine into all aspects of our studies, with adequate control groups, and insist on reporting our fat grafting or cell grafting procedures on the GRAFT registry\textsuperscript{32} a U.S.-based nation-wide registry of fat grafting, so that these procedures can become more versatile and reliable in the hands of all plastic surgeons.

Fig. 2. The equipment used for mechanical dissociation of adipose-derived stromal vascular fraction in the discussed articles.
CONCLUSIONS

The standard protocol for SVF isolation, although effective, suffers from high costs, long protocol durations, and regulatory scrutiny. Nonenzymatic alternatives address these concerns and isolate populations containing adipose-derived regenerative cells. Mechanically isolated SVCs have demonstrated clinical benefit through contribution to improved volume retention of fat grafts. However, there is a lack of literature comparing different mechanical isolation methods, and published methods yield fewer SVCs than enzymatic isolation. Larger proportions of hematopoietic cells and fewer regenerative cells are isolated when using mechanical protocols. Future studies should compare and refine these protocols to improve cell yield and quality and reduce the proportion of contaminating cell populations. In addition, development of closed, automated systems may improve standardization and reproducibility of results, reducing operator-dependent variations, and simplify the techniques, making them more approachable in a clinical practice. Randomized control studies are needed to analyze long-term outcomes in volume retention and tissue quality after fat grafting supplemented with mechanically isolated cells. Further analysis of the duration and cost of these methods would be of great benefit. These techniques for nonenzymatic SVF isolation are still in infancy with much to be learned about their potential efficacy for research and clinical applications. They may carry significant implications for advancing and making SVC-based therapies more accessible.

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