Exosome-educated macrophages and exosomes differentially improve ligament healing

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Abstract
Recently, our group used exosomes from mesenchymal stromal/stem cells (MSCs) to simulate an M2 macrophage phenotype, that is, exosome-educated macrophages (EEMs). These EEMs, when delivered in vivo, accelerated healing in a mouse Achilles tendon injury model. For the current study, we first tested the ability of EEMs to reproduce the beneficial healing effects in a different rodent model, that is, a rat medial collateral ligament (MCL) injury model. We hypothesized that treatment with EEMs would reduce inflammation and accelerate ligament healing, similar to our previous tendon results. Second, because of the translational advantages of a cell-free therapy, exosomes alone were also examined to promote MCL healing. We hypothesized that MSC-derived exosomes could also alter ligament healing to reduce scar formation. Similar to our previous Achilles tendon results, EEMs improved mechanical properties in the healing ligament and reduced inflammation, as indicated via a decreased endogenous M1/M2 macrophage ratio. We also showed that exosomes improved ligament remodeling as indicated by changes in collagen production and organization, and reduced scar formation but without improved mechanical behavior in healing tissue. Overall, our findings suggest EEMs and MSC-derived exosomes improve healing but via different mechanisms. EEMs and exosomes each have attractive characteristics as therapeutics. EEMs as a cell therapy are terminally differentiated and will not proliferate or differentiate. Alternatively, exosome therapy can be used as a cell free, shelf-stable therapeutic to deliver biologically active components. Results herein further support using EEMs and/or exosomes to improve ligament healing by modulating inflammation and promoting more advantageous tissue remodeling.

1 | INTRODUCTION

Despite surgical and physical therapy advancements designed to improve ligament healing, none eliminate scar formation, thereby making them prone to further injury. Recently, we delivered exosome-educated macrophages (EEMs) to simulate an M2-mediated anti-inflammatory response within a tendon wound.1,2 EEMs were generated by exposing CD14+ macrophages to mesenchymal stromal cell (MSC) derived exosomes.1,2 In
vitro tests showed EEMs exhibit M2-like behavior as indicated by an increase in M2 cell surface markers: CD206, PD-L1, and PD-L2. When tested in vivo, treatment of mouse Achilles tendon injuries with EEMs reduced inflammation and improved strength of the healing tendon, supporting the use of EEMs to promote healing. Indeed, results were superior when compared to treatment with MSCs. Further development of a more predictably beneficial therapy would be clinically significant.

Exosomes are small (40-200 nm) lipid membrane-bound vesicles that participate in cell-to-cell communication by transferring specific host cell-derived nucleic acid/protein to targeted recipient cells and reprogramming cell behavior, making the clinical potential of exosomes significant. Previous in vivo studies indicated that exposure of healing Achilles tendon to exosomes reduced the M1/M2 macrophage ratio and increased the number of endothelial cells 2 weeks after healing. However, the functional/mechanical benefit of exosomes was not as obvious as the EEMs in that mouse Achilles tendon model. The goals of the current study were two-fold. First, to reproduce the healing effects of EEMs in a different rodent model, that is, a rat medial collateral ligament (MCL). We hypothesized treatment with EEMs would reduce inflammation and accelerate ligament healing, similar to our previous tendon results. Second, to study the effects of exosome therapy in the rat MCL injury given the cell-free translational advantage of this therapy. We hypothesized that MSC-derived exosomes could similarly improve ligament healing and reduce scar formation.

2 | MATERIALS AND METHODS

2.1 | Cell culture

Human cell use was approved by the Health Sciences Institutional Review Board, University of Wisconsin (UW)-Madison, School of Medicine and Public Health (Protocol number: 2016-0298). MSCs were isolated from bone marrow of normal healthy donors as previously reported. Identity of MSCs was confirmed via cell adherence and flow cytometry. Passage 4-6 MSCs were used to isolate exosomes via differential ultracentrifugation. Exosome protein and RNA was characterized with a NanoDrop spectrophotometer. Particle concentration and diameter were measured using ISON qNano Nanoparticle (Zen-Bio, Inc, Research Triangle Park, North Carolina) analysis.

Mononuclear cells were obtained from granulocyte-colony stimulating factor-mobilized peripheral blood of healthy donors, using Ficoll Paque Plus density gradient separation (GE Healthcare Bio-Sciences, Piscataway, New Jersey). Red blood cells were lysed. Monocytes were isolated by via autoMACS Pro Separator (Miltenyi Biotech, Auburn, California) using anti-human CD14+ microbeads (Miltenyi Biotech). CD14+ monocytes were cultured 7 days for macrophage differentiation (Figure 1A). To generate EEMs, CD14+ macrophages were exposed to 3 × 10⁹ exosomes for 3 days.
HRP polymer or rabbit-on-rodent HRP polymer (Biocare Medical, Pacheco, California). Bound antibody complex was visualized using diaminobenzidine. H&E staining was performed and used for fractal analysis (to quantify collagen organization) and wound area measurements. Stained sections were imaged via camera-assisted microscope (Nikon Eclipse, model E6000 with Olympus camera, model DP79). Images (3-5 sections/animal) were captured within and outside of the granulation/scar tissue. The border of the wound was defined as the interface between the organized and disorganized collagen. Staining and wound size were quantified using ImageJ (NIH, Bethesda, Maryland). Scar formation was estimated at day 14 by quantifying type I and III collagen IHC staining, performing fractal analysis to measure collagen organization, and measuring wound size of H&E images.

2.4 Mechanical testing

Pull-to-failure testing was performed. Each MCL was removed with both femoral and tibial insertion sites intact. The surrounding tissue was excised. The femur-MCL-tibia complex was mounted in a custom testing bath and mechanical testing machine. A pre-load of 0.1 N was applied, cross-section was measured, and each MCL was preconditioned (cyclically loaded to approximately 1% strain for 10 cycles). The ligament was pulled to failure at a rate of 10% strain per second. Failure force, failure stress, ligament stiffness, and Young’s modulus were measured/computed to determine posttreatment MCL mechanical behavior.

2.5 Statistical analysis

A one-way analysis of variance (ANOVA) examined treatment differences of all data. If the overall P-value for the F-test in ANOVA was significant (P ≤ .05), Fisher’s LSD post hoc comparisons were performed. Data were presented as mean ± SD. All analyses were performed using KaleidaGraph, version 4.03 (Synergy Software, Inc, Reading, Pennsylvania).

3 RESULTS

3.1 EEM study

3.1.1 Mechanical testing

Compared to the control, EEM treatment significantly increased failure load (P = .03; Figure 2A) and max stress (P = .01; Figure 2B).
However, treatment differences in stiffness ($P = .64$; Figure 2C) and Young’s modulus ($P = .41$; Figure 2D) were not significant. Collectively, these results support the concept that EEM treatment improves the mechanical properties of the healing MCL.

### 3.1.2 | IHC of cellular factors

Macrophage immunophenotypes indicated that the number of day 7 endogenous M1 macrophages was significantly reduced after EEM treatment. Data are considered significantly different ($P \leq .05$) based on Fisher’s LSD post hoc pairwise analysis from four MCLs/treatment. Values are expressed as mean density/mm$^2 \pm$ SD. Scale bars = 50 $\mu$m.
FIGURE 4  Immunohistochemistry and histology results by the day 14 healing medial collateral ligament (MCL) after exosome treatment. Treatment of the injured MCL with exosomes (Exo) had no significant effect on the number of (A,B) M1 macrophages, or (C,D) M2 macrophages 14 days postinjury. E,F, In contrast, Exo treatment significantly (E) reduced size of scar tissue compared to the control. F, Representative H&E stained images indicate the reduction in scar size. Outlined region (ie, border between the organized and disorganized collagen fibers) indicates wound area. Exo treatment also significantly increased (G,H) type I and (I,J) type III collagen production within the granulation tissue. K, Exo treatment significantly improved collagen organization. L, Table showing results of tested factors that were not significantly different after Exo treatment. Data are considered significantly different (P < .05) based on Fisher’s LSD post hoc pairwise analysis from five MCLs/treatment. Values are expressed as mean area or density/mm² ± SD.
treatment \((P < .01)\) compared to the control (Figure 3A). In contrast, M2 macrophages were significantly increased by EEMs (Figure 3B). These changes in macrophage phenotypes by EEMs significantly reduced the M1/M2 macrophage ratio (Figure 3C,D; \(P = .01\)). No changes were noted in the number of endothelial cells or myofibroblasts \((P > .05);\) Figure 3E).

3.3.1 | Scar size, collagen production, and organization

No significant changes were noted in wound size, IHC quantified type I and type III collagen production, and fractal analysis of collagen organization \((P > .05);\) Figure 3E).

3.2 | Exosome study

3.2.1 | Mechanical testing and IHC of cellular factors

At the concentration tested, treatment with exosomes did not significantly improve mechanical function (Figure 1C), nor did they elicit changes in endogenous M1 and M2 macrophages (Figure 4A-D), endothelial cells, and myofibroblasts (Figure 4L).

3.2.2 | Scar size, collagen production, and organization

Unlike EEMs, treatment with exosomes significantly reduced scar formation 14 days postinjury compared to the control (Figure 4E,F). Exosome treatment also increased type I and type III collagen production within the granulation tissue (Figure 4G-J). Collagen organization, determined via fractal analysis, was also significantly improved (Figure 4K).

4 | DISCUSSION

To our knowledge, this is the first study to test the effects of EEMs on ligament healing. Similar to our previous Achilles tendon results, EEMs improved the rat MCL’s mechanical properties and reduced the endogenous M1/M2 macrophage ratio indicating less inflammation,\(^1\) whereas exosomes improved ligament remodeling through changes in collagen production, improved organization, and reduced scar without improvements in mechanical function.

Few studies have reported the effects of direct macrophage application on the healing process.\(^{17-21}\) Activated macrophages delivered to a cutaneous guinea pig wound, reduced wound size.\(^{17}\) Likewise, treatment of human ulcers with macrophages were reported to stimulate wound repair.\(^{19,20}\) More recently, a phase I in-human trial was conducted to evaluate the safety and feasibility of autologous macrophage delivery in patients with liver fibrosis.\(^{22}\) One year post-treatment, patients were alive and remained transplant free. In our study, direct delivery of EEMs to the transected MCL was able to suppress/prevent inflammation normally present after injury, by increasing the presence of endogenous M2 macrophages and reducing the M1 macrophages. Our earlier experiments tracking EEMs in vivo, indicated that the majority of exogenously administered EEMs are gone from the wound by day 7, thereby suggesting the effects are likely via paracrine or chemotactic action.\(^1\) Altogether, these studies support the concept that exogenous administration of M2-like macrophages are able to accelerate healing in various healing models and thus hold a much broader orthopedic promise.

Similar to our Achilles tendon results, EEMs improved strength of the MCL without increased production of type I collagen or improved collagen organization.\(^1\) The known mechanism for improved strength after EEM treatment extends beyond the scope of this manuscript but may involve changes in collagen fibril size, molecular cross-linking, and/or MMP/TIMP activity.\(^{22,23}\) For instance, one study demonstrated that collagen fibril formation was markedly reduced and cardiac rupture increased following M2 macrophages depletion after injury.\(^{22}\) Additional research is underway to elucidate the molecular mechanisms of EEM therapy on ligament healing.

In our study, exosomes elicited a significant biological response, indicated by an increase in type I and III collagen, improved collagen organization, and a reduction in scar size 14 days after injury. The ability for exosomes to reduce scar formation and upregulate collagen type I and III expression has been documented in tendon healing models, but this is the first study to our knowledge to report this in ligaments.\(^1,24\) We hypothesize that the postinjury timing of exosome delivery is important. At the time of injury and exosome administration, few macrophages are present, thereby reducing the likelihood for exosomes to target macrophages and control inflammation. Instead, exosomes would target fibroblasts and/or tissue stem cells to affect ECM production and remodeling.\(^{25}\) Future studies will include treatment of injuries with both EEMs and exosomes in a temporal manner where EEMs would reduce inflammation and exosomes would stimulate remodeling.

5 | CONCLUSION

We demonstrate that EEMs as well as MSC-derived exosomes have the potential to improve ligament healing via different mechanisms. EEMs and exosomes each have attractive characteristics as therapeutics. As a cell therapy EEMs are terminally differentiated and will not proliferate or differentiate to undesirable cell types, which remain a concern for many stem cell therapies. Moreover, EEMs could be generated from a patient’s monocytes using off-the-shelf exosomes, resulting in a faster and more facile process compared to autologous MCS. Alternatively, exosome therapy could be a cell free, shelf-stable therapeutic to deliver biologically active components. Altogether, results herein support the use of EEMs and/or exosomes to improve ligament healing by modulating inflammation and tissue remodeling.
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C.S.C.: conception and design, collection and/or assembly of data, data analysis and interpretation, manuscript writing; J.A.K.: conception and design, collection and/or assembly of data, provision of study material or patients; L.A.W.: collection and/or assembly of data, manuscript writing; M.M., K.H.: collection and/or assembly of data; A.M.S.: revision and final approval of manuscript; M.A.H.: financial support, revision and final approval of manuscript; P.H.: conception and design, provision of study material or patients, final approval of manuscript; R.V.: conception and design, provision of study material or patients, final approval of manuscript.

DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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