

OSTEOCHONDRAL ALLOGRAFT TRANSPLANT SYSTEM

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Abstract

Osteochondral allograft transplantation is an increasingly popular procedure that repairs osteochondral defects by introducing mature cartilage and subchondral bone to facilitate defect healing. These defects can arise from trauma, osteonecrosis, osteoarthritis, and other degenerative cartilage disorders. Existing surgical systems are detrimental to chondrocyte viability and limit vertical graft adjustment, which are both crucial for successful surgical outcomes. To address both challenges, we developed a novel surgical system that creates threads on the graft and receiving site to produce a screw-in graft. Testing revealed a significant improvement in chondrocyte viability with the screw-in graft over the traditional impaction method. However, matching the surface of the graft with the surface of the receiving site was not fully addressed with our current device. Since our device relies on threading, the vertical and rotational alignment of the graft with the receiving site are coupled once the threads are defined. Aligning the graft correctly in the receiving site is important to avoid incongruencies in the receiving site surface. Therefore, further testing of the device is necessary to develop a threading procedure that ensures correct rotational and vertical alignment with each use of the device. To validate the device, we must measure the difference in height between the surface of the graft and surface of the native cartilage. We plan to use 3D laser scanners to obtain surface measurements for evaluating how well the graft surface matches the native surface.

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Introduction

Motivation

Osteochondral allograft (OCA) transplantation is a surgical procedure that fuses a healthy cartilage and subchondral bone implant from cadaver donor tissue into the patient's cartilage lesion site, particularly in young, active adults [1]. The rate of OCA transplantations performed is increasing by 5% annually, and is expected to reach 3500 procedures by the year 2020 [2]. Despite the prevalence of this procedure, the failure rate is as high as 18% due to unsuccessful integration of the donor and recipient tissues. Nevertheless, the benefit of this procedure over total knee arthroplasty is the promising possibility of restoring full-range of motion, and maintaining the patient's quality of life [3]. The motivation in this project, therefore, is to improve full-graft integration and long-term integrity by protecting chondrocyte viability—a significant factor in determining procedure success [4].

Existing Devices

Arthrex Osteochondral Allograft Transfer System (OATS)

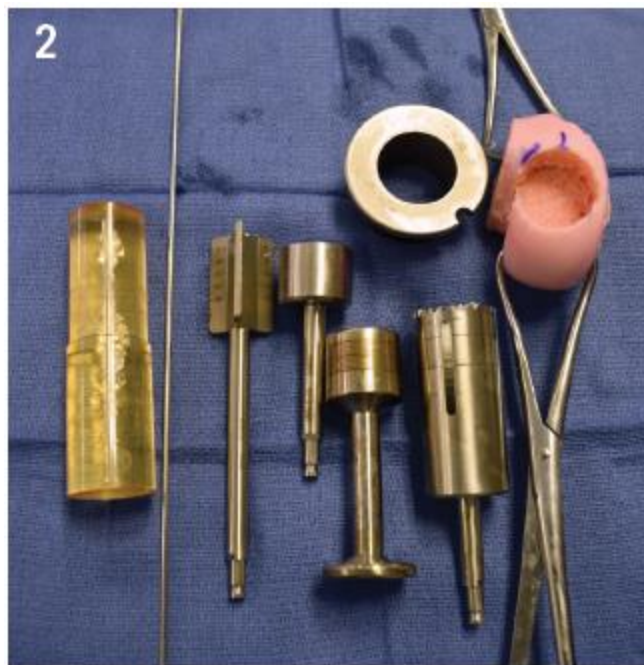


Figure 1: Arthrex Osteochondral Allograft System. (1A) Locating and sizing guide. (1B) Stainless steel guide wire. (1C) Cannulated reamer. (1D) Surgical hole saw guide ring. (1E) Surgical hole saw. (1F) Impacting rods.

The Arthrex Osteochondral Allograft Transfer System (OATS) uses several different tools to prepare the donor site, and harvest the graft before impacting it into the patient [5]. As shown in Figure 1, is a translucent plastic sizing guide that is used to determine how large of a graft must be placed to completely repair the defect. The surgeon places this guide over the defect to ensure that it is completely covered, selecting a larger or smaller size as needed. Once the proper size is determined, the sizing rod is held orthogonal to the surface of the defect and the guidewire (1B) is

inserted through the hole in the center of the sizing guide, and a drill screws the guidewire through the center of the defect and into the bone. After the guidewire is positioned, the cannulated reamer (1C) (with a diameter corresponding to the sizing guide) is inserted over the guidewire to drill a receiving hole to the proper depth (typically 7-14 mm). Miscellaneous tools (not pictured) are used to remove loose tissue from the bottom of the hole, as well as from the cartilage surrounding this hole.

To harvest the donor graft, the cadaver tissue is placed in a vice (not pictured) or another similar fixture to secure it for cutting. The shape of the condyle surrounding the prepared donor site is noted and the best geometric match on the donor tissue is selected. A surgical hole saw guide (1D) is held over the matched geometry of the cadaver graft and the hole saw (1E) is then used to cut the graft cylinder. The graft is inserted using the impaction rod (1F) and a surgical hammer until it sits flush with the surface.

Zimmer Chondrofix Osteochondral Allograft System

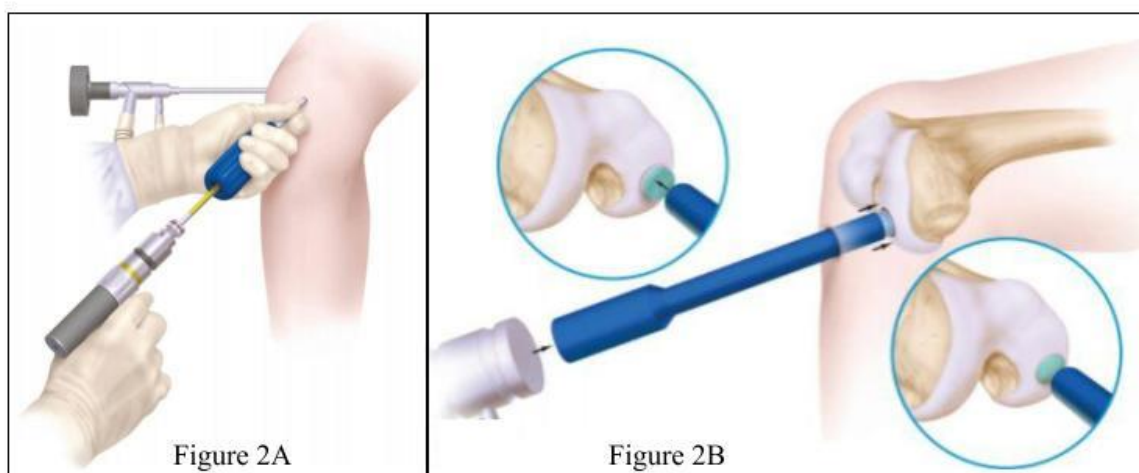


Figure 2: Zimmer Chondrofix Osteochondral Allograft System. (2A) Recipient site arthroscopic drill guide prepares the receiving site. (2B) Arthroscopic impactor secures the decellularized osteochondral allograft into the patient.

The Zimmer Chondrofix Osteochondral Allograft system (Figure 2) relies on a pre-made, decellularized osteochondral graft. This eliminates the need to prepare an allograft from cadaveric tissue during surgery. The steps leading up to graft insertion are similar to the Arthrex system. A plastic sizing rod determines the size of the graft that the surgeon will insert. A hollow punch of corresponding size is pounded into the bone over the defect while the surgeon keeps it perpendicular to the condyle surface. Depth markings on the side of the punch allow for greater control over the depth of the receiving hole. After punch insertion, the impacting handle is removed to expose a center hole that accepts a corresponding drill bit which removes the remaining bone inside the punch and leaves a perfectly sized graft receiving hole. Unlike the Arthrex system, this drilling system has a built-in depth stop allowing greater depth control, which can be challenging for surgeons. The drill bit and punch are removed, and the hole depth is verified before cutting the pre-made graft to length. The graft is inserted using the insertion tool, leaving it slightly proud of the surface, and the impaction tool pushes it flush with the surface. This system is designed for arthroscopic use, unlike with the Arthrex system [6]

Depuy Synthes COR® Precision Targeting System

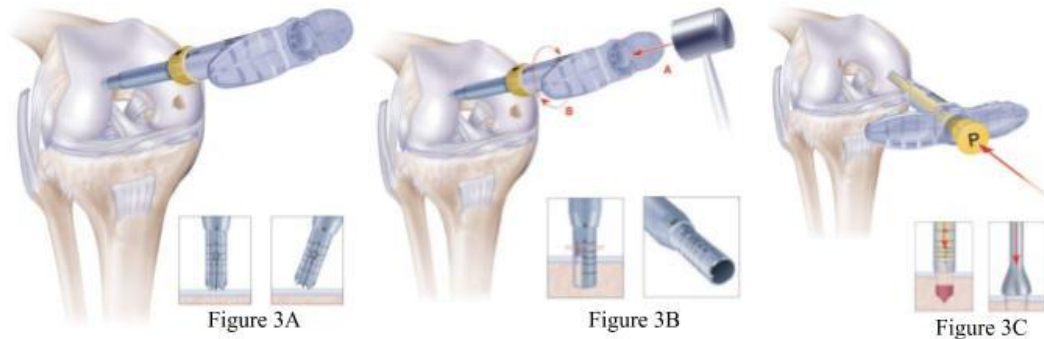


Figure 3: COR® Precision Targeting System. (3A) Graft harvesting tool placement. (3B) Graft harvesting tool impacted into bone and rotated to score the graft for removal from the patient. (3C) Graft transfer tube is placed over the receiving site, and a low impact insertion tool secures the graft into the patient.

The COR® Precision Targeting System boasts ease of use and improved accuracy, but its claim to protect chondrocyte viability defines it from other systems. Using “no-impact transfer” and “low-impact delivery”, it is designed to be used to surgically treat femoral articular cartilage lesions via autograft transplantation. However, the claims of improved chondrocyte viabilities is unsubstantiated by the provided literature. Use of an autograft is another concept unique to this system. To harvest the donor graft, the graft harvesting tool is placed on a non-weight-bearing articular surface (Figure 3A), and a mallet drives the cutter to the desired depth, indicated by measurements on the tip of the tool (Figure 3B). Rotating the tool scores the bottom of the graft to free it from the patient. The graft inside the graft transfer tube is then aligned with the recipient site and impacted until it is fully inserted (Figure 3C) [7].

These three systems indicate that there is little variation in methodology to OCA transplantation procedures. Every OCA system currently on the market relies on impaction to set the graft in place. This represents a significant gap in the market that an improved osteochondral grafting system can fill.

Problem Statement

Osteochondral transplantation procedures are becoming increasingly common but maintain a procedural failure rate of 18%. Current surgical methods involve impaction of an osteochondral allograft into the region of the defect. The goal of this treatment is to introduce mature hyaline cartilage and subchondral bone that will ultimately integrate with the native tissue and repair the defect. The main problem with current OCA surgical systems is that they all rely on impaction of the graft, which has been shown to be deleterious to chondrocyte viability, and this directly affects the success of the procedure. To address this concern, we developed a novel OCA surgical system that cuts matching threads on the graft and recipient site resulting in a screw-in graft.

Testing showed the chondrocyte viability was significantly improved using the screw in method compared to impaction. However, matching the surface of the graft with the surface of the receiving site was not fully addressed with our current device. Since our device relies on threading, the vertical and rotational alignment of the graft with the receiving site are coupled once the threads are defined. Aligning the graft correctly in the receiving site is important to avoid incongruencies in the receiving site surface, which can lead to overloaded joints and premature graft failure [8]–[10]. Therefore, further testing of the device is necessary to develop a threading procedure that ensures correct rotational and vertical alignment of the graft with each use of the device. To validate the device, we also must develop a measurement tool to assess how well the surfaces of the graft and receiving site match.

Background

Osteochondral Defect Etiology

Osteochondral defects arise from any type of pathology or injury that cause the bone and articular cartilage to separate; these include osteonecrosis, osteochondritis dissecans, and idiopathic developmental defects [1], [5], [11]. The leading concomitant knee pathology for this defect is a tear in the medial meniscus, which reduces support of the knee and results in greater joint contact forces [4]. Other pathologies leading to osteochondral defects include abnormal bone growth and excessive stress in the knee [12].

Osteochondral Allograft Transplant Procedure

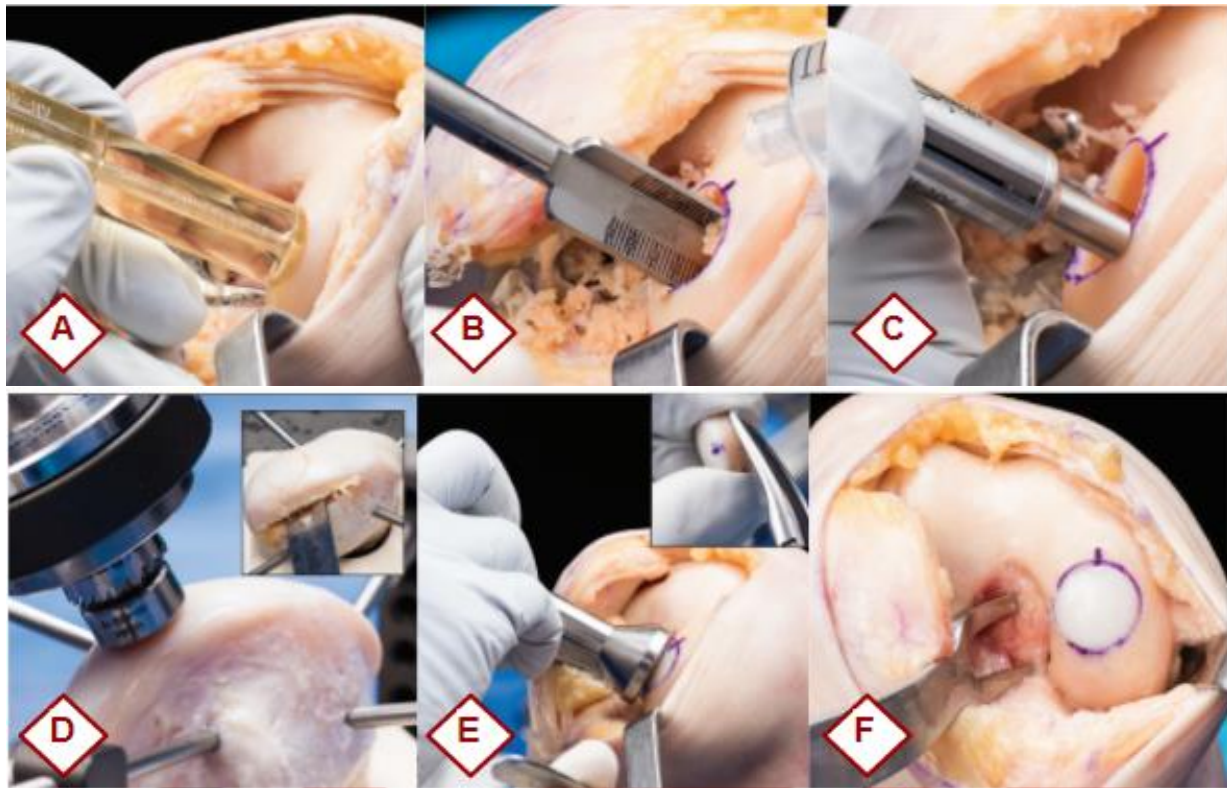


Figure 4: OCA transplant procedure as outlined by the current surgical guide. (4A) Sizing the defect with plastic sizing rod. (4B) Drilling the recipient site to desired depth with a cannulated reamer. (4C) Measuring depth of recipient site with plastic measuring rod. (4D) Cutting donor graft with surgical hole saw. (4E) Impacting donor graft into recipient site with impacting rod. (4F) A successfully implanted graft. [13]

The most common surgical approach to implanting an osteochondral allograft is the dowel technique. This procedure begins by preparing the recipient site for the allograft. The focus of this preparation is to create a cylindrical void that is perpendicular to the surrounding cartilage. To ensure perpendicularity, a guide wire is inserted orthogonal to the condyle at the defect site. A cannulated dowel reamer is passed down the guidewire and advanced to a depth of between 7 mm -14 mm, clearing a void 10 mm-25 mm in diameter.

The allograft is created from fresh cadaver tissue, and its geometry is matched to the recipient site on the patient. To harvest the graft, a surgical hole-saw is passed through a guide ring on the articular cartilage creating a cylindrical dowel. Then, the measurements of the recipient site depths are used to guide the surgeon as they cut the graft to a complementary length with an oscillating saw. The allograft is then positioned directly above the recipient site, and impacted until the graft lies flush with the surrounding cartilage [13].

Physiology

Impaction force used to press fit osteochondral allografts into place during a transplant procedure induces cell death in the superficial portion of the articular cartilage. The impaction impulse deforms mechanoreceptors in the cell. This initiates an intracellular signaling cascade ultimately activating executioner caspases, triggering cell apoptosis (Figure 5).

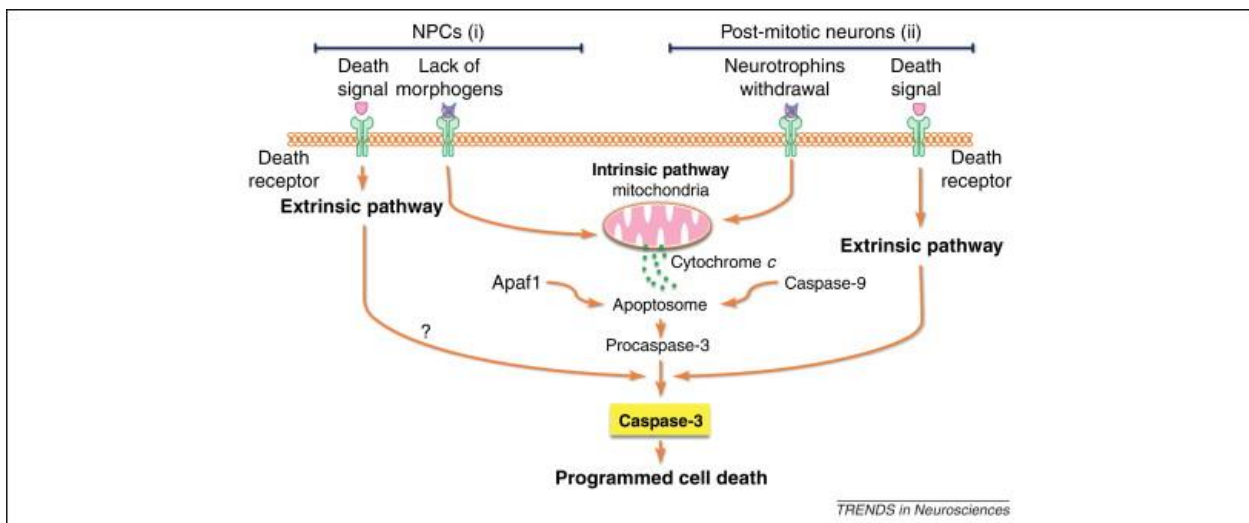


Figure 5: Bio-signaling pathway leading to chondrocyte death following impaction. Mechanoreceptors initiate a signal cascade ultimately activating executioner caspases and leading to apoptosis [14].

This mechanism was discovered in a study to assess the effects of impaction on chondrocyte viability during OCA transplantation. In this study, grafts were taken from the distal aspect of the femoral head and inserted into their recipient sites. Additional grafts were taken from each donor knee and used as controls. The grafts were assessed after forty-eight hours, and the impacted grafts had an average of 47% greater cell death, particularly on the superficial layer of the cartilage (Figure 6). The impacted grafts showed increased levels of caspase 3 activity which is a known enzyme involved in programmed cell death [14].

A separate study was conducted to assess the optimal ratio between the number of impacts, and the total force required for graft implantation. Allografts were impacted with 37.5, 75, 150, and 300 N loads 74, 37, 21, and 11 times respectively. One unimpacted allograft was kept as a control. The researchers found a direct relationship between cell viability and the force to strike ratio: lower impulses with more strikes yielded higher cell viability. The unimpacted control allograft had little to no cellular death [15]. This study demonstrated that graft impaction forces during OCA are deleterious to chondrocyte viability.

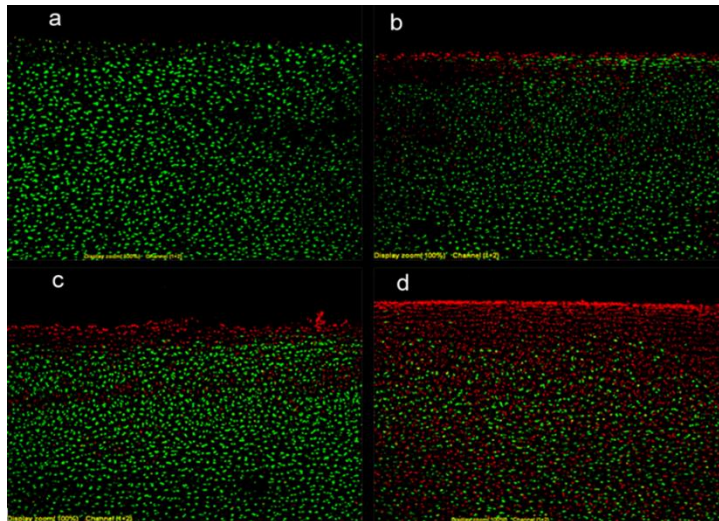


Figure 6: Live/dead chondrocyte cell staining following impactation at varying loads. Red indicates cell death; green indicates viable cells. (a) control (b) 75 N (c) 150 N (d) 300 N [15].

The effects of impactation on chondrocyte viability is an important medical concern for this procedure as chondrocyte viability at the time of impactation is the primary determinant of allograft success. A study was performed in canine models to assess the effects of chondrocyte viability at the time of impactation on allograft success. Subjects received an osteochondral allograft and graft cell viability was assessed at the time of impactation where viability ranged from 23-99%. Six months post-surgery, procedural success was compared to initial chondrocyte viability. The researchers found that no graft with an initial chondrocyte viability below 70% was successful [2]. While other factors contributed to procedural success, none were as significant as initial chondrocyte viability.

Previous Design Work

Overview of Prototype

The current prototype consists of three components: a tap, a die and die base, and graft screwdriver. The die base, shown in Figure 7, is made of aluminum but could easily be transitioned to stainless steel for application in a surgical setting. It consists of two parallel plates separated by vertical stainless-steel pins. In the bottom plate, a removable supporting cup holds the graft. Two thumb screws tighten down the graft and prevent it from rotating when the die is threading it. In the top guiding platform, there is a hole cut through it that matches the size of the die. This hole lies directly over the supporting cup, which ensures axial alignment between the threads and the graft.

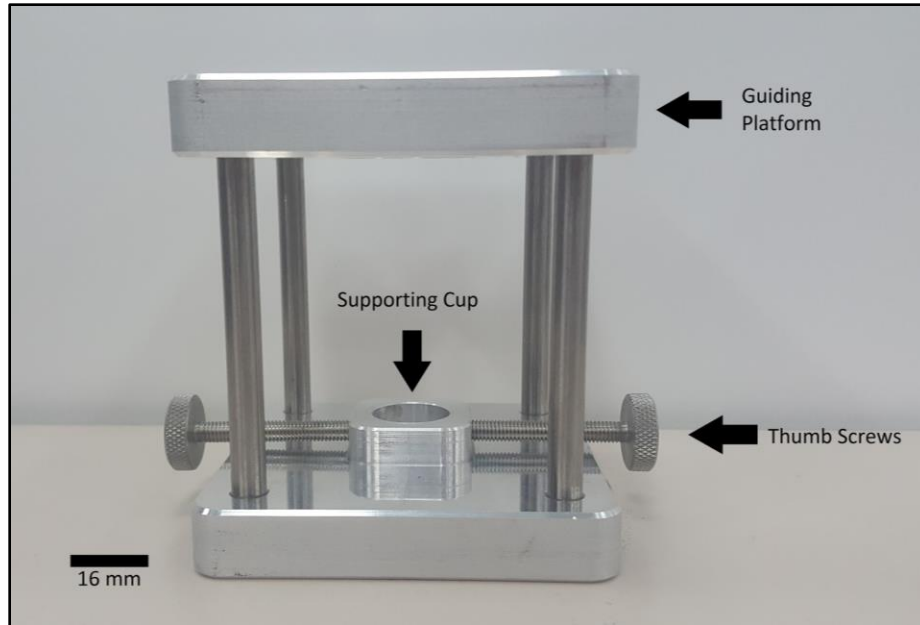


Figure 7: The above image is the final prototype of the stand used to hold the allograft in place while external threads are created. The guiding platform ensures axial alignment. The allograft would be inserted cartilage side up into the supporting cup, and the thumb screws would tighten around the allograft.

The die, as depicted in Figure 8 consists of a stainless steel body and handle. The handle is removable and offers the surgeon a comfortable grip when using the tool. The die body consists of an open-ended cylinder. The open end has 4 flutes built in to allow the bone shavings created during the threading process to escape. The threads have a 1.5 mm pitch, allowing the surface of the graft to always remain 0.75 mm of the native surface. A previous iteration of this prototype used a 2 mm thread pitch. Finally, the die threads begin as a taper and lead in to allow more consistency during the threading process while requiring less pressure from the surgeon.

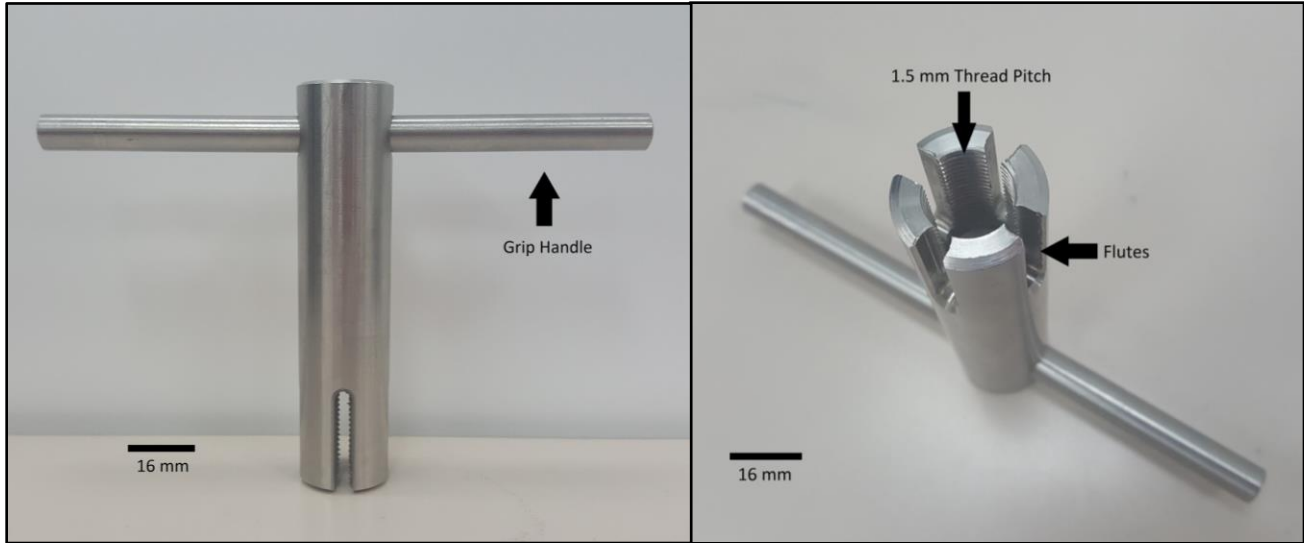


Figure 8: Depicted is the die system used to create threads on the external profile of the graft before insertion into the recipient site. The die would be inserted through the guiding platform to maintain alignment as it creates the external threads in the cartilage and subchondral bone.

The tap, as depicted in Figure 9 consists of a stainless steel body and handle. The die body consists of a cylinder with a hole along the central axis, and threads protruding from working end. The central hole matches the guidewire currently used in surgical systems and is used to slide the tap along said guidewire. This ensures the threading axis is perpendicular to articular surface. The tap has 4 flutes built in to the threads that allow the bone shavings created during the threading process to escape. The threads have a 1.5 mm pitch, matching that of the die above. Finally, the tap threads begin as a taper and lead in to allow more consistency during the threading process while requiring less pressure from the surgeon. The handle is removable and also has a guide hole to slide over the guide wire.

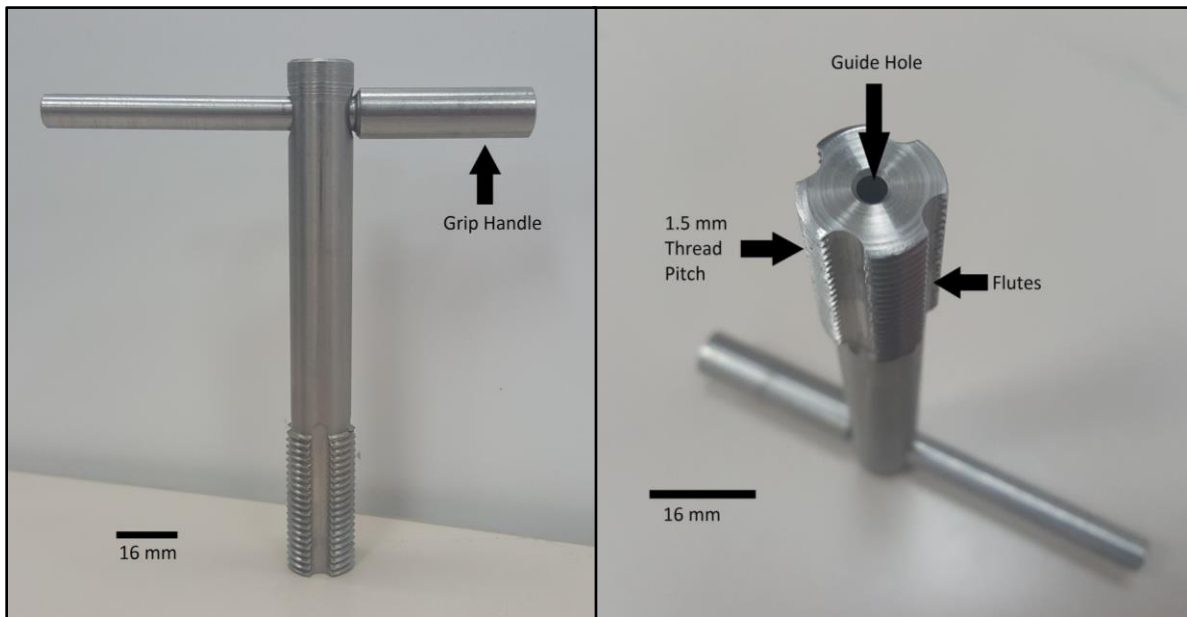


Figure 9: Seen above is the tap system used to create sister-threads within the recipient site when preparing it for graft receipt. A guide wire is to be slid through the guide hole and inserted into the center of the recipient site to ensure proper alignment.

The graft screwdriver, as shown in Figure 10 is designed to aid in screwing the graft into the receiving site because hand screwing was found to be difficult. The device is made from two easily sterilizable materials: stainless steel and silicone. It utilizes a hex-bit to attach to a standard screwdriver handle, which is a familiar tool for most people. The working end utilizes two 1 mm diameter tines and a disposable silicone cap to protect the chondrocytes from overhead force when the device is in use. The tines are tapped through the cartilage into the subchondral bone, securing the graft for the surgeon to screw into the receiving site. Additional damage to the chondrocytes due to the tines was found to be minimal. There was localized death, but the viability returned to above the 70% threshold within 400 microns and the overall viability was not significantly altered from the control samples. Additionally, in the current system when particularly large defects requiring multiple grafts, similar pins are used to secure the first graft while the second is being inserted. This appears to have minimal effect on the outcome of the procedure, further justifying their use in this device.

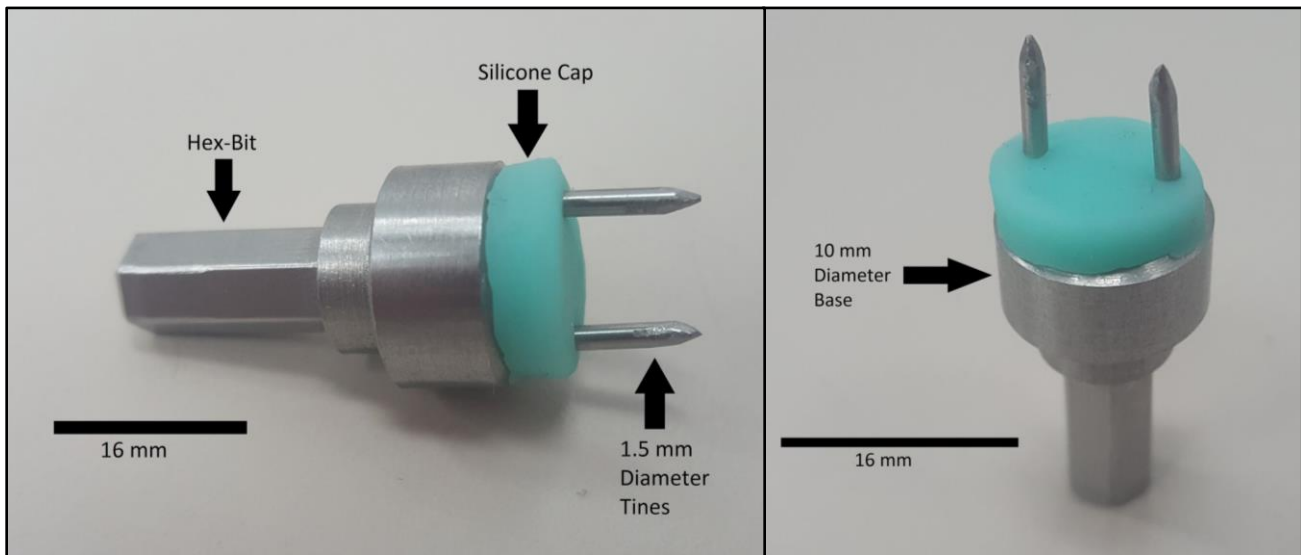


Figure 10: The novel bident tool design depicted above attaches to a standard screwdriver via the hex-bit extrusion. The tines are small enough to cause minimal damage to the cartilage and large enough to maintain the mechanical strength necessary to effectively insert the graft. The silicon cap is a failsafe intended to protect the cartilage from unwarranted impact in the case of accidental over-insertion of the bident into the cartilage.

Prototype Shortcomings

The primary challenge with the current prototype is that it lacks a robust protocol for ensuring that the graft is properly aligned with the native tissue upon insertion into the recipient site. The surgeon aims to insert graft such that it sits flush with the articular cartilage when fully inserted, however the surface geometry of the native cartilage surrounding the graft insertion site is non-planar. Thus, the distance from the base of the recipient site to the top of the articular cartilage varies throughout the circumference of this area. Therefore, it is imperative that the graft

is inserted in a specific orientation such that local graft height is complementary to recipient site depth.

In the traditional osteochondral allograft transplant procedure, a surgeon has two degrees of freedom, when inserting the graft: rotation and vertical translation. This allows the surgeon to first place the graft in the in the proper rotational alignment, so that throughout the circumference of the graft the local height of the graft is the same as the local depth of the recipient site. Then, the surgeon uses an impaction rod to drive the graft to a depth equal to the recipient site.

Due to the threaded nature of our system, rotation and vertical adjustment of the graft are coupled. Thus, we are limited to one degree of freedom when inserting the graft. As a result, threading of the graft must be both precise and accurate, to ensure that the graft sits flush and properly aligned with the articular cartilage.

Required Project Research

3D Laser Scanners

Measuring the geometry of the threaded graft and receiving site presents a unique engineering challenge. As it is difficult to accurately measure the point where threading starts on both components using conventional methods (i.e. calipers, ruler, protractor), and even more difficult to full characterize the size of the components using these techniques, we were forced to investigate more robust measurement techniques. 3D laser scanning provides a convenient method for obtaining a complete and accurate characterization of the surface geometry of the threaded receiving site and threaded graft. 3D laser scanners can be used to compile a highly accurate digital recreation of our threaded graft and receiving site, which will allow us to quantitatively determine how the two components will align, and ultimately allow our team to determine a method for properly aligning the surfaces of both components.

The University of Wisconsin-Madison Makerspace has two 3D laser scanners available to students. The first of these scanners is the Creaform Handyscan 700. This laser scanner is handheld and collects measurements of a component as it is passed over the object by the user. This scanner has a theoretical maximum resolution of 0.05 mm. However, the practical resolution of the scanner is limited by the stability and speed of the user's arm as the collect measurements, and rarely achieves the theoretical resolution.

The second laser scanner that the Makerspace offers is the Einscan SP. The Einscan SP reports a resolution of <0.05 mm, which is similar to the Handyscan. However, the Einscan SP connects the scanner to a measurement stage with a support arm. This feature of the Einscan SP fixes the relative point of reference of the system and makes the system independent of user technique. Thus, it is possible for the Einscan SP to consistently achieve the maximum limit of resolution.

3D Point Cloud Analysis

For analysis of different laser scans, even when collected with the same scanner, it is necessary to register the coordinate systems of the scans to ensure that any measurements are not affected by global rotations or translations during scan measurements. There are two algorithms

that have been implemented in MATLAB that should allow for easy registration between the different scans.

The normal distribution transform (NDT) algorithm was developed to reconstruct 3D renderings of rooms given 2D scans from images, or more importantly from LIDAR range finders. Individual points are grouped into 2D objects called cells. Once the point cloud data are split into these cells, mean position values are calculated for each cell and this mean is termed q . Once the mean is found, the convergence matrix is found given equation 1.

$$\Sigma = \frac{1}{n} \sum_i (x_i - q)(x_i - q)^t \quad (1)$$

The convergence matrix is used in an optimization to find convergence of the system through varying rotation angles and translations within this 2D plane. These standard rigid transformations yield potential solutions to the registration x' and y' as in equation 2.

$$\begin{pmatrix} x' \\ y' \end{pmatrix} = \begin{pmatrix} \cos \emptyset & -\sin \emptyset \\ \sin \emptyset & \cos \emptyset \end{pmatrix} \begin{pmatrix} x \\ y \end{pmatrix} + \{t\} \quad (2)$$

A score of p is used as the optimizing parameter given the transformed coordinates x' and y' as in equation 3 [16].

$$score(p) = \sum_i \exp\left(\frac{-(x'_i - q_i)^t \Sigma_i^{-1} (x'_i - q_i)}{2}\right) \quad (3)$$

Once convergence is found, the MATLAB function returns the point cloud data that have been transformed to the reference coordinate system for analysis.

A different registration algorithm used in MATLAB is the iterative closest point (ICP) algorithm. This algorithm works to find the closest corresponding point between the reference data X and un-registered data P where the difference between individual points x and p is calculated by equation (4).

$$d(p, x) = \min \|x - p\| \quad x \in X \quad p \in P \quad (4)$$

The points p having minimum distance to X are stored as the closest points in Y and represents the registration of P with respect to X using a least squares registration method until convergence of the mean-square error [17].

While both the NDT and ICP registrations appear to be applicable to our laser scanning application, the ICD algorithm was developed for registering distinct 3D objects, whereas the NDT algorithm was developed to create a 3D shape out of a series of 2D images. Given the parallel

between the ICP algorithm and our laser scanning application, we intend to pursue this algorithm for registering the different laser scans.

Threaded Graft Mechanical Integrity

Given the novel method of using a threading system to secure the graft into the patient, it is critical to characterize its mechanical strengths and ensure that the graft will not fail unexpectedly. In this case, the graft is usually unsupported at the bottom of the hole—this space is left to afford the surgeon a degree of adjustment to the vertical graft placement. Consequently, the only portion of the graft supporting tibiofemoral contact forces is the thread. Given contact forces applied to the axis of the graft, the threads are most likely to experience shear-stress failure.

Shear stress at the threads can be modeled based on the applied axial compressive load, and the geometry of the thread [18]. In this case, the thread shear area (AS_s in mm^2) is related the length of engagement (LE); thread pitch (p); the maximum minor diameter of the internal thread (D_{1max}); and the minimum pitch diameter of the external thread (d_{2min}) (equation 5). The diameter and pitch specifications are easily gathered from a table of thread dimension standards for each given thread size [19].

$$AS_s = \frac{\pi * LE * D_{1max}}{p} \left[\frac{p}{2} + 0.57735 (d_{2min} - D_{1max}) \right] \quad (5)$$

Shear stress V can be calculated by dividing the thread shear area by the applied force F (equation 6). The applied force F was estimated based on numerous assumptions of extreme loading circumstances. The graft was assumed to have been placed on the femoral condyle and sitting proud of the surface so that it bears the entirety of any tibiofemoral contact force. Such forces have been found to exceed 6.2 body-weights during large loading activities such as stair climbing [20]. Assuming the individual weighs 150-pounds (667 N), this corresponds to a simulated tibiofemoral contact force of over 4100 N.

$$V = \frac{F}{AS_s} \quad (6)$$

Given that $F = 4100$ N, the shear stress V was calculated for numerous graft sizes from 10-25 mm encompassing the most common sizes of osteochondral allografts across typical graft insertion depths (represented by the length of engagement LE in the equation). The results were plotted in Figure 11.

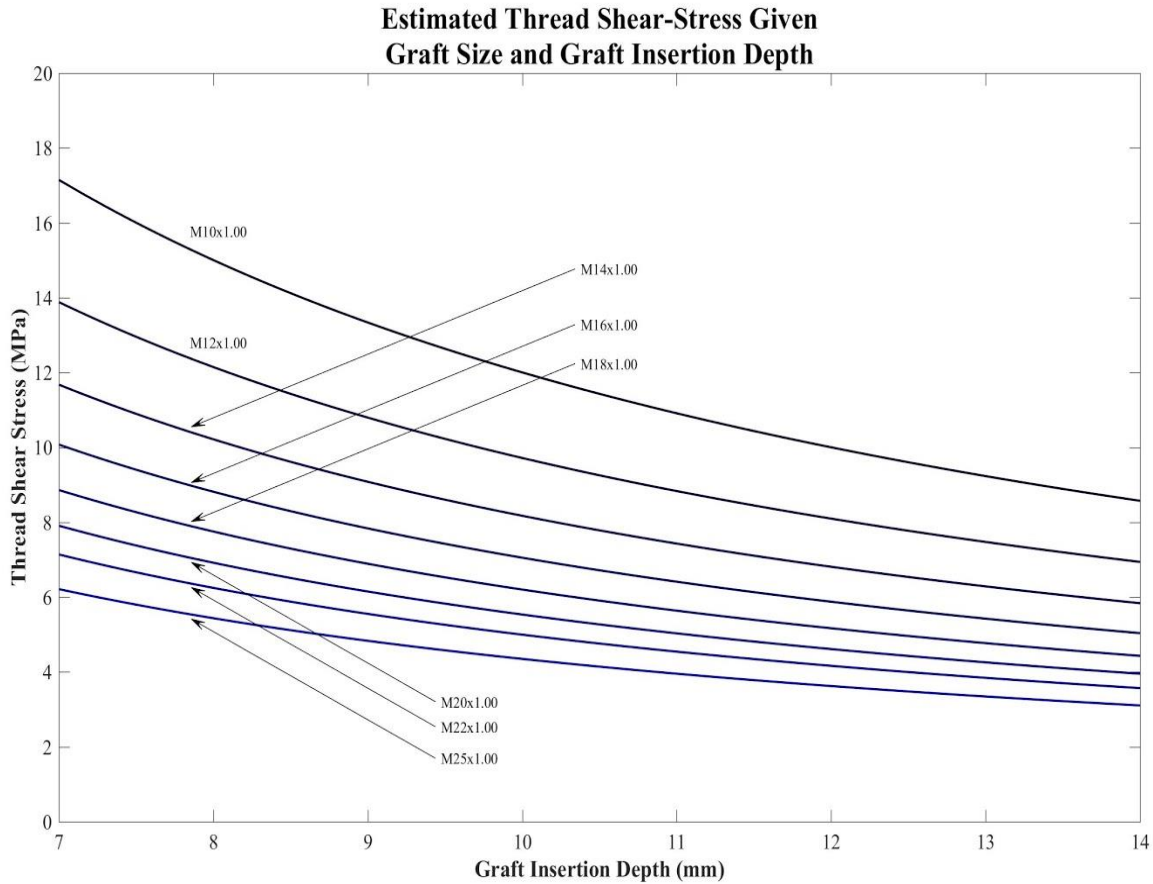


Figure 11: Plot of thread shear stress with various thread geometries varying with graft insertion depth. The simulated load comes from a 150-pound individual climbing stars generating a tibiofemoral contact force of 4100 N.

Cortical bone, such as that present surrounding the receiving hole for an osteochondral allograft, can support a shear stress of approximately 50 MPa [21]. Given the results of the simulation in Figure 11, shear stress in the smallest graft (a 10-mm graft with an M10x1.00 thread) at the minimum insertion of 7-mm only experiences a shear stress of 17 MPa—this is well below the prescribed failure criterion of 50 MPa. Given the extreme (and very unlikely) loading parameters described in this simulation, the contact forces acting directly on the graft will result in shear stress far below the failure stress. Ultimately, these data indicate that the graft can readily support moderate loads until the donor bone can integrate with native bone and reform a solid foundation.

Additionally, threads with a finer pitch exhibit a decreased shear stress and thus are less likely to fail under extreme loading. (The thread pitch p decreases in equation 5, which results in an increases shear area and consequently decreases the shear stress on the graft demonstrated with equation 6). Considering the application of the grafts, the finer threads also allow for finer adjustment by the surgeon to match the surface geometries. Previous testing with different thread types showed that the finer threaded tap and dies initiate the threading process with less force required by the operator. However, the finer pitches were also found to be less consistent than the

coarser threads and tend to experience more friction between the graft and the donor site. The testing results may be attributable to the differences in bone we used for each thread pitch. The coarser thread was tested on hard, mature bovine tissue while the finer thread was tested on softer, adolescent porcine tissue. One goal of this semester is to resolve these inconsistencies and determine a balance between the thread pitch, graft adjustability, thread quality, and initial threading location and difficulty.

FDA Manual Orthopedic Device Standards

The U.S. Food and Drug Administration outlines medical device regulations in CFR Title 21- Subchapter H [22]. There are particular exemptions to the requirement of sending premarket notifications to the FDA, provided that the device has existing characteristics of commercially distributed devices of that generic type [23]. In the case of intention to use a device for a different purpose than that of pre-existing devices of the same type, notification is still required. In addition, a modified device operating on a different fundamental technology requires notification of the FDA. For the purposes of manual orthopedic surgical instruments, exemptions apply in the same manner, so long as they are classified within a particular group, as well as adhere to specific limitations [24]. A generic device, such as a bone tap with minor modifications, would likely necessitate little regulation, and perhaps qualify for exemption, in contrast to a novel instrument for threading donor tissue.

Surgical Instrument Material Standards

Various grades of stainless steel are used in biomedical applications. Corrosion resistance is an essential aspect of any surgical instrument. The International Organization for Standardization (ISO) specifies metals commonly used to manufacture standard surgical instruments [25]. There are many alloys of stainless steel available, however martensitic alloys are generally chosen for surgical instruments, due to its substantial hardness [26]. This grade of surgical steel meets the requirements of ISO product standards, passing corrosion tests based on the methods of sterilization normally encountered by these products (i.e. autoclaving) [27].

Client Information

Dr. Brian Walczak is a faculty member at the University of Wisconsin School of Medicine and Public Health. Dr. Walczak is an orthopedic surgeon specializing in sports medicine, pediatric sports medicine, and joint preservation. He is experienced with the OCA procedure and proposed the mechanism of a screw-in graft to address numerous shortcomings.

Design Specifications

We have developed a device for orthopedic surgeons performing osteochondral allograft transplantation which allows them to thread the donor graft and corresponding recipient site. The chief aim of the system is to improve chondrocyte viability, which has a positive relationship with procedure success. The system must therefore maintain chondrocyte viability above 70%, which has been shown to be a threshold for procedure success. Any damage to the graft beyond current surgical techniques should be avoided. Additionally, the surface of the graft should match the surface of the receiving site, and the total height difference between the two surfaces must not exceed 1mm. Rotational and vertical alignment between the graft and receiving site should be optimized to minimize irregularities in the implant surface. Furthermore, the procedure for

threading the graft into the donor site should be easy for the surgeon and should integrate with the current surgical technique. Ideally, the system will require minimum skilled input from the surgeon to prevent avoidable errors and to promote widespread adoption of the device. The entire system must be easily sterilizable, and operable in a surgical environment. For more detailed product specifications, refer to Appendix A.

Prototype Evaluation

Threading Protocol

Threading Consistency

The first thing we need to evaluate with our device is how consistently we can thread the graft and receiving site. Consistent threading is important because if we are unable to consistently define threads where we want them, then it will be impossible to develop a reliable procedure that ensures rotational alignment of the graft. Essentially, we need to evaluate if we can start threading where we want. We plan to evaluate this by first marking a spot on the top surface of the graft where we intend to start threading, then we will thread the graft and mark the spot where the threads actually start. Using ImageJ we can measure the angle between the two markings and then evaluate how large and how consistent this angle difference is.

Threading Procedure

We need to develop a protocol for using the device that ensures both rotational and vertical alignment of the graft in the receiving site. For a given thread size, diameter, and depth we need to determine where to start threading the graft so that the rotation of the graft is correct when fully screwed into the receiving site. We will start to develop this procedure using plastic pieces, first with a flat receiving site and then slanted surface receiving sites. We will be able to evaluate rotational alignment by marking where the graft should align and measuring the angle difference between where the graft ends up and where it should be. In this simple setup, we will likely be able to measure height difference of the surfaces with a caliper, and rotational alignment angle with ImageJ. Next, we will use a bone model to determine if our procedure developed in plastic works with the geometry and mechanical properties of bone. We plan to use either non-viable porcine tissue or SawBone as bone models. One issue we may face is that smaller threads don't work well with soft bone samples. While smaller threads allow greater vertical adjustment of the graft, smaller threads are harder to define and may break down in a softer material. Since the surface geometry of bone is more complex, we will likely need to utilize 3D laser scanning to validate our procedure in model bone.

3D Laser Scanning

To characterize the height differences in the implanted grafts from the native tissue, 3D laser scans and resulting point cloud analysis will be used. To start, a laser scan will be taken of the exposed joint without any modification. This scan will serve as a reference coordinate system for registration, and as a ground-truth for graft-height comparisons (i.e. how far from this native surface does the graft lie after implantation?). The grafting procedure will be performed with our threading method, as well as with the traditional impaction method. After the grafting is complete, the articular surfaces will be scanned again to measure any geometry changes. These scan data will be imported to MATLAB and registered to the unaltered joint scan using the ICP algorithm. Using plane fitting and interpolation features also built into MATLAB will define the articular surface and allow for a comparison between the native and grafted joints.

Viable Tissue Testing

Finally, we will perform the full OCA transplant procedure with viable tissue to simultaneously evaluate geometrical alignment and chondrocyte viability in our device. We will conduct a series of comparative surgeries in porcine models obtained from the Clinical Sciences

Center at the University of Wisconsin-Madison. Surgeries will be performed using both the standard impaction protocol and our new threading protocol. If possible, we will have an experienced surgeon, such as our client Dr. Walczak, to perform the procedures as they would be performed in a clinical setting.

A single biopsy of cartilage will be taken from the center of each allograft. These biopsies are intended to be a relative sample of the gross tissue viability of impacted grafts. An additional biopsy of cartilage that has not been implanted will be taken from each of the knees. This biopsy will be used to normalize the initial tissue viability of each sample.

All biopsies will be stained with Calcein AM and Ethidium Homodimer-1. This stain is a form of a live/dead assay which is intended to characterize tissue viability. Calcein AM is a green fluorochrome that binds to the membrane of living cells and will fluoresce green when excited using confocal microscopy. Ethidium Homodimer-1 is a red fluorochrome that integrates into dead cells and will fluoresce red when excited using confocal microscopy. All samples will then be imaged using an AIRS confocal microscope at the Wisconsin Institute for Medical Research Imaging Core. Analysis of cell viability from these images will then be performed using ImageJ.

Power stats

To calculate the significance of the threading angle difference testing results, a one-sided, one sample t-test with a significance level of $\alpha = 0.05$ will be used. The testing results will be compared to the null hypothesis of a 0-degree difference between intended and actual thread starting locations. The alternate hypothesis will be that our threads have a greater start angle than zero. The t-test can be calculated using equation 7.

$$t_{n-1} = \frac{X\sqrt{n}}{s} \quad (7)$$

In this equation, n is the number of samples, X is the mean sample angle difference, s is the sample standard deviation, and t is a test statistic which can be compared to a standard T-table to obtain a p-value. With our data, we can also create a $1-\alpha = 95\%$ confidence interval of where the observed threads begin relative to the intended beginning using equation 8.

$$1 - \alpha = \left[X - \frac{t_{n-1, \frac{\alpha}{2}}(s)}{\sqrt{n}}, X + \frac{t_{n-1, \frac{\alpha}{2}}(s)}{\sqrt{n}} \right] \quad (8)$$

For the surface matching tests, the mean difference between the observed and initial surfaces can be calculated. Equation (7) can be used again to compare the mean difference in surface height. However, for this test, a two-sided t-test will be used because the graft can be sit both proud and below the initial surface. Equation (8) can then be used to create a 95% confidence interval of the mean surface height difference. If the mean difference is greater in magnitude than +/-1 mm, our design must be reconsidered and modified. In addition, if the confidence interval or individual points in the data exhibit the same criteria, the threading and insertion process may require modification such that there is an overall surface height difference of less than 1 mm.

Conclusion

OCA transplantation corrects osteochondral defects through the implantation of a donor graft. This procedure is becoming increasingly common but maintains a relatively high failure rate. Current surgical methods impart high forces on the graft through impaction, which is deleterious to chondrocyte viability and negatively affects procedural outcome. We previously designed a device that utilizes a screw system, which aims to eliminate the force applied to the graft by the current impaction method. Testing showed that our device significantly improves chondrocyte viability compared to the standard impaction method. However, our current design does not address how well the surface of the graft matches the recipient site surface. Since we are using a screw system, the rotational and vertical alignment are coupled in the graft. Therefore, we must develop a threading procedure to ensure rotational alignment of the graft in the recipient site with a given thread size, diameter, and depth. We plan to test our device by first evaluating how consistently we can thread the graft. Next, we will develop a threading procedure using plastic pieces to test how consistently we can match rotational alignment of the graft. Next, we will evaluate how well this procedure translates to the geometry and mechanical properties of bone by using either non-viable porcine tissue or SawBone. Additionally, we plan to develop a 3D laser scanning method of measuring the difference in height between the surface of the graft and recipient site. Lastly, we will perform viable tissue testing to evaluate geometry and chondrocyte viability considerations simultaneously to determine if our procedure is a viable model to improve OCA transplantation procedure outcomes.

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Appendix

Appendix A: Product Design Specifications

Osteochondral Allograft Tapping System Product Design Specifications

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Function: Osteochondral allografts (OCAs) are used to repair chondral defects in young, active patients. The current procedure involves cutting the graft from cadaveric tissue, then using impaction to drive the graft into a low-clearance receiving hole drilled over the defect. The large impulse associated with graft impaction often leads to decreases in grafted chondrocyte viability, and negatively affects procedure outcomes [1]. To avoid deleterious impaction, we created a screw-in system which taps the patient receiving site and threads the donor graft allowing the graft to be screwed into the patient. Testing revealed that this new system has significantly higher implanted chondrocyte viability when compared to the impaction protocol. A challenge unique to our system, however, is that the one degree-of-freedom (DOF) nature of a screw mechanism limits graft adjustment relative to the traditional two DOF impacted graft. Therefore, the aim of this project is to develop a protocol for threading the graft and receiving site such that desired graft rotation and height can be achieved simultaneously when the graft is fully inserted into the patient.

Client Requirements

1. The protocol must permit a graft height offset from native tissue of no more than $\pm 1.0\text{mm}$.
2. After graft preparation and insertion, chondrocyte viability must be consistently greater than 70%, which has been shown to be a threshold to successful graft integration [1].
3. The entire system must be sterilized before use in surgery.
4. The threading protocol must be quick and easy to learn so as not to drastically alter the current surgical practice.
5. Damage to the chondral surface must be no greater than what presently occurs during OCA transplantation.

Design Requirements

- 1) Physical and Operational Characteristics
 - a) Performance Requirements

- i) Threading the graft and receiving site should not damage the articular cartilage
 - (1) It should limit gouging, scratching, and other mechanical alterations to the native, or graft cartilage.
 - (2) It should not result in significant chondrocyte death after use
 - ii) Insertion of the graft must be easily executed so as to minimize the risk of tissue damage.
 - iii) During the procedure, the graft should be easy to insert and remove allowing the surgeon to adjust the graft depth.
 - iv) The threading protocol must cut threads in the graft and receiving site that result in predictable graft placement.
- b) Safety
- i) The threading system should not increase the chances of postoperative complications, including (but not limited to) infection, tissue death, or graft dislocation.
 - ii) Long term, the threaded graft must not lead to an associated cartilage disorder, significant fissuring or fibrous tissue infiltration, or improper tissue integration.
- c) Accuracy and Reliability
- i) The threading protocol should allow for successful graft integration into the recipient site. This means that the procedure should maintain at least 70% chondrocyte viability after implantation.
 - ii) The measurement protocol should ensure that, after graft insertion, the donor curvature closely matches that of the recipient site within ± 1.0 mm of height difference.
- d) Life in Service
- i) Non-disposable components must be serializable to allow for repeated use
 - ii) Life of device materials will vary depending on chosen stainless steel alloy.
 - iii) Disposable components should be minimized in the design to prevent excessive recurring costs.
- e) Shelf Life
- i) Capable of storage at room temperature.
 - ii) Must be compliant with hospital regulations of storage.
 - iii) Shelf life is not likely to present as a significant design consideration.
- f) Operating Environment
- i) Protocol must not compromise sterility of the device or surgical field.

- ii) Must function within range of operating room temperatures, in addition to *in vivo* conditions.
 - iii) Must be usable in concurrence with all other orthopedic tools and materials.
- g) Ergonomics
- i) The devices must be designed for comfortable handheld use by the orthopedic surgeon during the procedure.
 - ii) To promote easy rotation, the tool must be easy to locate over the central-axis of the graft.
- h) Size
- i) Tools will be appropriately sized for handheld usage by orthopedic surgeon.
 - ii) The device should accommodate bone graft sizes 10 mm - 25 mm in diameter and 7 mm - 14 mm deep.
- i) Weight
- i) Since the device will be hand-held, its total weight should not be so heavy that it is cumbersome or fatigues the surgeon during use.
- j) Materials
- i) All materials must pass ISO regulations to corrosion resistance and excessive wear from use [2].
 - ii) Tools involved in the procedure must be sterilizable or disposable.
- k) Aesthetics
- i) Aesthetics will serve as a secondary initiative to the function of the final product.
- 2) Production Characteristics
- a) Quantity
- i) One prototype capable of inserting the graft into the patient.
 - (1) The prototype may have more than one component.
- b) Components
- i) The final product must consist of a mechanism for inserting the graft into the recipient hole.
 - (1) A component must hold the graft in place and align a threading mechanism.
 - (2) An external threading component must create threads on a harvested graft.
 - (3) An internal threading component must create threads in the patient receiving site.

- (4) A component will function as a screwdriver to screw the graft into the recipient site.
- (5) A final component must define the starting threading position on the graft threading component to ultimately allow for predictable graft placement.

3) Miscellaneous

a) Standards and Specifications

- i) The final product must comply with the FDA standard for manual surgical instruments as stated by CFR 21 - Subchapter H - Medical Devices [2]

b) Customer

- i) Orthopedic surgeons implanting an osteochondral allograft.

c) Patient Related Concerns

- i) Decreasing chondrocytes cell viability leads to diminished graft integrity.
- ii) Unwanted debris and fragments of the graft may be released into the synovial fluid environment and cause other complications.
- iii) A graft with an articular surface homologous to the native tissue is necessary for long term grafting success and patient health.

4) Current Systems

- a) Arthrex Osteochondral Allograft Transfer System (OATS). This system is the prototypical system used in osteochondral transplant procedures (and is most similar to the system Dr. Walczak uses). It uses a sizing guide, guide wire, and cannulating reamer to size, locate, and ream the chondral defect. The allograft is prepared using the hole saw which is guided by a manually held ring. The impaction rods forces the graft into the receiving hole [3].
- b) Zimmer Chondrofix Osteochondral Allograft. This system uses a hollow punch hammered into the bone to guide the drill bit during receiving site preparation. There is no need to prepare an allograft since it comes with a pre-made, decellularized allograft that fits precisely in the hole created by the punch and drill bit. The graft is inserted the majority of the way using the insertion tool, and is pounded in the remainder of the way using an impaction rod [4].
- c) COR Precision Targeting System. This is the only surgical system that claims to address chondrocyte viability concerns associated with OCA transplantation. The tool encloses the graft during harvesting and insertion to protect it from mishandling. The surgical guide also claims to use “low impaction insertion” but does not describe how impaction forces are minimized relative to traditional tools. Despite the promise with the system, it is not currently in use in human OCA transplantation. [5]
- d) There are no direct competitors, and of the ones currently in use, all rely on graft impaction.

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Appendix B: Fabrication and Testing Material Expenses

Use	Product	Part Number	Supplier	Link	Quantity	Unit Price	Total Price
Mock graft for geometric fitting in plastic	Rod Stock, HDPE, $\frac{5}{8}$ in., 48 in.	22JL48	Grainger	https://www.grainger.com/product/POLYMERSHAPES-Rod-Stock-22JL48	1	\$9.40	\$9.40
Mock receiving site for geometric fitting in plastic	Sheet Stock, 12" LX 12" W X 1.000" Thick, 176 Max. Temp. (F), Off-White	1ZAH3	Grainger	https://www.grainger.com/product/POLYMERSHAPES-Sheet-Stock-1ZAH3	1	\$22.15	\$22.15
						Material Total:	\$31.55
						Tax:	\$2.48
						Shipping:	\$13.59
						Total:	\$47.62

Table 1: Complete list of all materials used to make the prototype. Total project expenses are \$47.62.