Characterizing the relationship between basic calcium phosphate particulates in the knee joint and the frictional coefficient of articular cartilage in the intact rat knee.

Senior Honors Thesis
Departments of Mechanical Engineering and Orthopaedics

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Abstract

In order to examine the role of calcium phosphate particulates in the development of joint degeneration, a pendulum apparatus for evaluating the whole-joint coefficient of friction of intact knees using the rat model was designed, constructed, and evaluated. The femur is mounted to the apparatus at a fixed angle, and a weighted pendulum is affixed to the tibia such that its weight places the joint in compression. The pendulum is displaced, and then allowed to oscillate freely until it returns to rest. The oscillation of the pendulum is recorded using an optical tracking system. A decay function is fitted to the oscillation in MATLAB, and the coefficient of friction is derived from this function. An initial sample population (n = 2 limbs) of pilot specimens was prepared, and the coefficient of friction of each specimen was evaluated in its naïve state, and following particulate injections at 0, 1, 10, and 100 mg/mL concentrations. The coefficient of friction of each specimen was evaluated after each injection. The 0 mg/mL treatment (phosphate buffered saline only) was found to have no statistically significant effect on coefficient of friction (p = 131); all other treatments investigated were found to cause statistically significant increases in coefficient of friction (p < 0.001). Based upon these data, a physiologically plausible treatment of 100 μL of 1 mg/mL suspension was selected for the main study. A sample population (n = 6 limbs) of specimens was prepared, and the coefficient of friction of each specimen was evaluated in its naïve state and after sham (saline) and loaded (1 mg/mL) treatments. The sham treatment was found to have no statistically significant effect (p = 0.367), while the loaded treatment was found to cause a 31% increase in coefficient of friction (p = 0.039).

Introduction

Osteoarthritis (OA) of the knee affects over 9 million adults over age 60 in the United States alone.21 OA is a painful condition characterized by wear, degradation, and loss of joint cartilage, and can seriously impact physical ability and quality of life. Treatment of OA is complicated by the fact that once damaged, cartilage heals poorly. Thus, prevention and mitigation of damage are critical. Improved understanding of the numerous risk factors associated with the development and progression of OA may eventually lead to better prediction, earlier intervention, novel therapies, and better overall care. Characterizing the effects of basic calcium phosphates on joint friction in the rat model may lead to a more complete understanding of their relationship to OA in humans.

The formation of mineralized cysts within the matrix of articular cartilage is a form of cartilage degradation associated with aging and OA.13 The deposition of basic calcium phosphates in articular cartilage has been associated with aging, decreased joint function, and severity of osteoarthritis in humans.8,9 Common mineral species found in cystic deposits include calcium pyrophosphate (associated with pseudogout) and a variety of other calcium phosphate species, including carbonate-substituted hydroxyapatite, tricalcium phosphates, octacalcium phosphate, and whitlockite; these species are collectively referred to as basic calcium phosphates (BCPs).18 As cartilage is worn away at the articulating surface of the joint or cysts increase in size, these cysts become exposed and ultimately rupture, releasing their contents into the synovial fluid. Cystic BCP deposits analogous to those found in human joints are also common in the joints of mature Sprague-Dawley rats.18
Little is currently known about the tribological effects of cartilage mineralization. The phenomenon has been characterized and mechanical analyses of affected tissue have been performed in the rat model, but the mechanisms of cyst formation and the tribological effects of cyst rupture remain unknown. In the human hip joint, the degree of BCP calcification has been found to correlate to clinical symptoms and histological markers of OA. Calcification has been linked to both aging and OA, and mineral particulates and wear debris have been found to contribute to inflammation and degeneration. It is possible that the presence of BCPs in the joint capsule contributes to OA through cartilage abrasion, as well as by inducing inflammation. From a mechanical perspective, the introduction of hard particulate matter into the lubricating fluid of any joint or bearing alters the interaction between contact surfaces, typically resulting in an increase in friction and ultimately in abrasive wear. However, the specific relationship between synovial BCPs and joint friction is currently unknown.

A variety of arthropathies are linked to the presence of mineral crystals in the joints. Pyrophosphate arthropathies such as calcium pyrophosphate dihydrate disease (CPPD) provide some insight into the possible effects of intracapsular mineral particulates. In CPPD, the presence of calcium pyrophosphate (CPP) in the synovial fluid leads to swelling and inflammation. While some crystalline arthropathies, such as gout, are caused by the formation of crystals in the joint fluid and surrounding soft tissues, CPPD is caused by the release of CPP into the joint capsule from cystic formations in the connective tissue, mechanically similar to the rupture of BCP cysts in OA. The in-vivo introduction of CPP into the synovial fluid of the knee joints of rabbits with preexisting induced OA has been found to increase the severity of the condition, suggesting that the presence of intracapsular particulates can exacerbate cartilage wear and inflammation.

Rats are widely used as model organisms in OA research, as their small size makes them easy and cost-effective to keep, and their rapid maturation allows efficient longitudinal study. However, their small size complicates surgical modeling procedures and specimen manipulation, and provides limited amounts of tissue for subsequent analysis. Due to these size constraints, pendulum testing is the preferred method for evaluating joint friction in small-animal models. Other commonly-used methods of evaluating friction, such as pin-on-disk, pin-on-plate, and cartilage plug analysis, examine the interaction between a metal or glass instrument and a cartilage sample. In rats, mice, and other small animals, however, these methods are impractical or impossible due to the difficulty of excising and manipulating cartilage samples on such a small scale. For example, the thickness of the articular cartilage layer in the rat knee is approximately 300 μm. By utilizing the whole, intact knee joint, the pendulum method eliminates the need to excise and manipulate such miniscule samples.

The measured coefficient of friction of a material also varies depending upon the lubricant used and the material with which it is interacting. Data obtained via this method reflect the frictional properties of synovially-lubricated cartilage-on-cartilage contact between geometrically compatible articular surfaces over an anatomically appropriate range of motion—that is to say, the data reflect the properties of the whole joint under conditions similar to those which exist in vivo. Furthermore, the pendulum method preserves the natural anatomical structures of the joint, ensuring that the articulating surfaces interact with one another much as they would in vivo. A further advantage of the pendulum method for this area of inquiry is that an intact joint capsule
can contain and retain fluids, allowing the introduction of a fluid particulate suspension into the joint space in order to simulate BCP release due to cyst rupture.

The small-animal pendulum testing methodology described by Drewniak et al. uses the whole joint and adjacent bones.\textsuperscript{2,3} The femur is mounted to the apparatus at a fixed angle; a weighted pendulum is affixed to the tibia such that its weight places the joint in compression. The flexion angle of the joint when the pendulum is at rest is chosen to approximate the mean flexion angle of the knee joint during the gait cycle in \textit{vivo}.\textsuperscript{2,3} In rats, this angle is \textasciitilde 67°.\textsuperscript{5,6} Pendulum weight is based upon percentage of mean specimen bodyweight, representing ordinary weight-bearing limb use. For skeletally mature Sprague-Dawley rat specimens with average body mass of approximately 600 g, the pendulum mass should be 300 g, or approximately 50% bodyweight, reflecting the mean joint loading during gait.\textsuperscript{22} Pendulum length is chosen to yield a pendulum frequency approximating the flexion frequency of a continuous walking gait; in rats, this frequency is \textasciitilde 1 Hz.\textsuperscript{1,22} The pendulum is displaced by an angle equal to half the mean range of motion during gait—12° in rats—then released and allowed to oscillate freely until it returns to rest.\textsuperscript{5,6,22} The oscillation of the pendulum is tracked using an optical tracking system, and the coefficient of friction is determined by analysis of the decay envelope of the oscillation.

The model most commonly used to derive coefficient of friction from pendulum data is the Stanton, or linear, fit. This model approximates the decay of the oscillation as a linear function, and derives a linear coefficient of friction \( \mu \) from the slope of the line as

\[ \mu = l m / 4 r \]

where \( l \) is the distance the fulcrum to the center of mass of the pendulum, \( m \) is the slope of the linear-fit decay function, and \( r \) is the epicondylar radius.\textsuperscript{1} This linear approach does not accurately reflect the curved profile of the decay envelope, however, and does not account for viscous damping effects due to soft tissues or air resistance. A second approach is to fit a function with both exponential and linear terms to the data to obtain a coefficient of viscous damping \( c \) and a coefficient of friction \( u \) as

\[ u = 4 \delta \pi^2 l_0 / m g r T^2 \text{ and } c = 4 j \pi l / T \]

where \( \delta \) is derived from a coefficient of the model, \( j = c T / 4 \pi l \), \( \theta_0 \) is the angular displacement at cycle 0, \( T \) is the time between cycles, \( I \) is the moment of inertia of the pendulum, \( m \) is the mass of the pendulum, \( g \) is the gravitational constant, and \( r \) is the epicondylar radius.\textsuperscript{25}

A comparative study applied both Stanton (“linear”) and combined linear-exponential (“combination”) decay models to the same experimental data, and concluded that although the combination model fits the experimental data more closely and accounts for viscous damping, and thus more accurately reflects the behavior of the specimen, the linear model has greater measurement precision.\textsuperscript{5} Furthermore, the linear model has been widely used, and thus linear fit analyses are more directly comparable to existing research.\textsuperscript{2} Due to the differing advantages of both methods, however, it is prudent to analyze experimental data using both methods.
The aims of this research are a) to develop an experimental apparatus for the purpose of determining the whole-joint coefficient of friction of a specimen in vitro, and b) to determine whether the presence of basic calcium phosphate particulate in the joint capsule has a significant effect upon whole-joint coefficient of friction. The primary hypothesis is that the whole-joint coefficient of friction will increase with the introduction of basic calcium phosphate particulate into the intracapsular space.

Apparatus

Pendulum Device

![Figure 1: Full Pendulum Device with Specimen and Trigger Bar](image)

The pendulum device consists of three main elements: the frame, the mounting block, and the pendulum. The frame is constructed of 25 mm (1”) square T-slotted extruded aluminum tubing. T-slotted tubing was selected for its strength, light weight, and ease of assembly. Because of the modular nature of the tubing, the frame can be fully disassembled for storage, and is easily reconfigurable to accommodate additions and modifications. A wide rectangular footprint provides stability. The height of the frame can accommodate pendulum of up to 45 cm in length, making it possible to test larger specimens or at lower frequencies if desired. The two front upright members, located at opposite sides of the base, create an uninterrupted viewing window within which the pendulum can be optically tracked without marker occlusion. The single rear upright member and associated crosspieces support the mounting block. The entire frame is painted matte black to minimize reflective interference with the optical tracking system.
The mounting block is fabricated of high-density polyethylene. It is removable, and can be replaced with another block or other mounting platform, allowing the frame to be used with larger or smaller model organisms. A removable angle block sets the specimen mounting angle; this block can be replaced to allow reuse of the mounting block with other similarly-sized model organisms and different ranges of motion. Two set screws fix the position of the specimen when mounted in the block. An acrylic viewing window allows visual observation of the specimen when mounted and aids in orientation during mounting. A reservoir allows the specimen to be immersed in saline solution or cell culture medium to maintain hydration during extended testing. The exterior of the mounting block is painted matte black to minimize reflective interference with the optical tracking system; the viewing window and interior are unpainted for ease of observation.

The pendulum is fabricated of aluminum stock, and has an overall length of 28 cm below the upper crosspiece. A milled socket in the upper crosspiece fits the 11 mm (7/16”) potting tube used for the tibial segments. Pairs of holes in the side pieces allow adjustment of the lower crosspiece position (and hence the center of mass) by increments of 10 mm. Holes in the lower crosspiece allow the addition of plates to adjust the mass, and a bearing on the lower crosspiece interfaces with a linear actuation system to allow cyclic loading of the specimen. The pendulum is painted matte black to minimize reflective interference with the optical tracking system. Four 4 mm self-adhesive retroreflective markers (NaturalPoint, Corvallis OR) are affixed noncollinearly to one side of the pendulum, allowing it to be defined as a rigid body by the optical tracking software.

The trigger bar is an auxiliary tool used to uniformly and reproducibly deflect the mounted specimen for motion trials. It consists of two telescoping lengths of square aluminum tubing with a slot and thumbscrew for fixing the length. Crosspieces at either end allow one end to be placed flush with an element of the frame base, while the other end catches the bottom of the pendulum. The deflection angle is set by adjusting the length of the trigger bar such that it deflects the pendulum to the desired degree. Pendulum deflection is confirmed visually using a goniometer, and the length of the trigger bar is fixed.
Potting Jig

A jig is used to control the orientation of specimens during the potting process. It consists of a grooved polyethylene block and two threaded spring plungers. The block is cut at a 67° angle, reflecting the mean flexion angle of the rat hind limb during gait.\textsuperscript{5,6,22} The vertex of the block is removed to create a plateau which accommodates the joint capsule. Two walls, on either side of this plateau, support the spring plungers. The spring plungers pass through threaded holes in these walls, located where the vertex of the block would be. The specimen is placed over the block, and the spring plungers are “dialed in” on anatomical landmarks on the specimen, fixing it in place and aligning the transepicondylar axis perpendicular to the block. Grooves cut into the narrow sides of the block accommodate potting tubes, which are fitted into place below the specimen, then slid up to cover the free ends of the long bones. Slots perpendicular to these grooves allow the placement of stops, preventing the potting tubes from sliding back down and holding them in place while the potting medium cures.

Optical Tracking System

Optical tracking technology is widely used in laboratory and clinical settings to capture and analyze motion. A theoretical minimum of two cameras are required, but the addition of a third camera provides redundancy and improves accuracy. Typically, a minimum of three infrared cameras are arranged in noncoplanar fixed positions, such that their fields of view overlap, forming a “capture volume.” Objects within the capture volume are thus viewed from three separate perspectives. A wand or other calibration device of known geometry, with retroreflective markers placed at known positions, is used as a basis for calibration; based upon
the observed relative position of the markers and the geometry of the calibration tool, a spatial coordinate system is derived. Retroreflective markers are applied to the objects to be tracked, and their motion within the capture volume is recorded. A number of tracking systems were evaluated, and OptiTrack Flex 13 cameras with compatible Tracking Tools software package (NaturalPoint, Corvallis OR) were selected based on the high rated translational accuracy (0.1 mm) and the relatively low cost of the system. A calibration tool, ground plane tool (used to define axes within capture volume), and self-adhesive retroreflective markers were also obtained from NaturalPoint.

Methods

Optical Tracking System

Cameras were mounted on small tripods for stability and arrayed noncoplanarly on a bench ~5 feet from experimental apparatus, and all associated cables were plugged in and arranged to prevent future disturbance of camera position. Individual camera orientation was adjusted such that the capture volume encompassed a space sufficient to accommodate the experimental apparatus. Using the motion tracking software package (Tracking Tools, NaturalPoint, Corvallis OR), a full camera calibration was run. Wanding was performed uniformly and methodically throughout the capture volume. The frame of the pendulum apparatus was then placed on the bench, and more intensive wanding was performed within the volume defined by the frame, where accuracy is most critical. After calibration, the critical region of the capture volume was evaluated using the calibration wand. Overall wand error within the frame was found to be less than 0.1% (less than 0.25 mm). The ground plane tool (NaturalPoint) was placed on the bench and aligned with the frame, and the ground plane was set in Tracking Tools and the calibration was saved for later use. The ground plane tool was then removed to minimize unnecessary marker capture during trials.

Specimen Preparation

Hind limbs were disarticulated at the hip and ankle joints, and the soft tissues were dissected away to expose the joint capsule and the medial and lateral collateral ligaments. A small portion of the quadriceps femoris was preserved in order to anchor the patellar tendon and maintain the integrity of the joint capsule. On each specimen, a hole was drilled axially through the distal end of the tibia, and a fine wire approximately 40 mm in length (“tibial pin”) was inserted. Tibial pins were fixed in place with a fast-curing cyanoacrylate adhesive, and then visually aligned with the central axis of the tibiofibular segment. Prepared specimens were wrapped in saline-soaked gauze to prevent dehydration during storage.

Prepared and thawed specimens were patted dry, and the femoral insertion points of the medial and lateral collateral ligaments were visually identified and marked with a dot of ink. Location of insertion points was then confirmed by probing for non-visible bony landmarks (“divots”) using fine-tipped dissecting forceps. Specimens were placed over potting jig in correct orientation relative to grooves, and spring plungers were “dialed in” on landmark points, fixing the orientation of the specimen relative to the jig, such that the transepicondylar axis of the
specimen was oriented perpendicular to the grooves, and the long axis of the tibiofibular segment
was parallel to the grooves.

Potting tubes (11 mm (7/16”) square for tibia, 19 mm (3/4”) square for femur, 35 mm length)
were sealed at one end with masking tape. Tape was folded down onto the sides of the tube for
adhesion, and any excess tape where the tube interfaces with the potting jig was trimmed away
with a razor blade. The center of the sealed end of the tibial tube was geometrically located, and
a small puncture was made. The tibial tube was then fitted into the tibial groove and the free end
of the tibial pin was aligned such that it passes through the center of the sealed end. The tibial
tube was then slid up to cover the free end of the tibiofibular segment, and fixed in place with a
stop. The femoral tube was applied to the free end of the femoral segment and fixed in place in
the same manner, but without controlling axial alignment.

The potting jig was placed under a fume hood (or other adequate ventilation), and the base was
propped up at an angle of ~35 degrees, such that the tibial tube was approximately vertical. A
small batch (~2.5 g solid) of polymethylmethacrylate (PMMA) cement (Acraweld Repair Resin
and Self Cure Acrylic Liquid, Henry Schein #101-1105 and #101-7550) was prepared at a ratio
of 1 g solid to 1 mL liquid, and injected into the tibial tube while still fluid. When the surface of
the tibial potting had cured sufficiently to allow repositioning of the jig, the position of the base
was reversed such that the femoral tube was vertical. A larger batch of PMMA (~6 g solid) was
prepared at the same solid-to-liquid ratio and injected into the femoral tube. The joint capsule
was shrouded in saline-soaked gauze to prevent dehydration, and the potting was allowed to cure
under the fume hood until hard (~50 minutes) before specimen was removed from jig.

BCP Formulation and Injection

Calcium phosphate powder (Ca₃(PO₄)₂, particle size < 45 μm, Sigma Aldrich #21218) was
obtained, and suspensions were prepared in phosphate buffered saline (PBS) at concentrations of
0, 1, 10, and 100 mg/mL. Suspensions were vortex mixed immediately prior to loading of
syringe, and were not permitted to settle prior to injection. Suspensions were delivered into joint
via a 26 gauge needle, using a central anterior approach through the patellar tendon. Delivery
was confirmed by visual observation of joint capsule expansion upon injection.

General Testing Procedure

Masking tape was removed from potting tubes, and tubes were swabbed with acetone or other
appropriate solvent if necessary to remove any residual adhesive and/or PMMA. Some potting
tubing also required light filing to remove rough edges. The protruding portion of the tibial pin
was trimmed ~5 mm from end of tube, then bent such that it lay flush with the PMMA. The
tibial tube was then fitted into the socket in the upper crosspiece of the pendulum. The femoral
tube was fitted into the mounting block and positioned such that it was flush with both the rear
wall and the angle block. The tube was then slid down the angle block until its front edge was
flush with the lip of the reservoir, and the set screws were tightened to fix the specimen in place.
After mounting, the specimen was allowed to hang until stationary. The tracking system was
then initialized and the saved calibration state was loaded. The pendulum markers were selected
and used to define a rigid body in Tracking Tools. In order to define a zero position for use in
data processing, a recording was made of the stationary pendulum in its resting position each
time a specimen was mounted.

One crosspiece of the trigger bar was placed flush with inside of the lower left base member, and
the pendulum was manually deflected. The trigger bar is positioned, and the lower extremities of
the pendulum were allowed to rest against the second crosspiece of the trigger bar. The trigger
bar was then adjusted, maintaining alignment with the base, until the pendulum was uniformly
supported but only barely in contact with the crosspiece. The recording was started, and the
trigger bar was gently lowered to the bench, allowing the pendulum to swing freely.

Data Analysis

Position data for each trial were exported from Tracking Tools and read into MATLAB.
Position data were initialized by subtracting the mean position of the resting pendulum, and
pendulum displacement over time in both the x direction (anterior-posterior motion only) and in
the x-z plane (both anterior-posterior and medial-lateral motion) were calculated and plotted.
Points of maximum deflection in the positive direction ("peaks") were identified and validated\(^\dagger\),
and these spatial deflections were converted to angular deflections based upon the geometry of
the pendulum. Due to substantial medial-lateral motion in many specimens, x-z displacements
were used in the analysis of study data. Linear and combination fits were applied to the set of all
valid angular peaks, and values for linear-model COF, combination-model viscous damping,
combination-model COF, and root-mean-squared error values and other goodness-of-fit data for
each model were exported to a spreadsheet. Plots showing each stage of the data analysis
process were exported to a PostScript file. All MATLAB code is provided in Appendix II. Plots
illustrating each phase of the data analysis process are provided in Appendix III.

Means and standard deviations for the preliminary data (Specimens P3 and P4, and total) were
calculated for all treatment levels. Total means and standard deviations were used to calculate
the Cohen’s \(d\) effect size for each treatment level. Univariate analysis of variance was performed
to examine differences among treatments for the main study data (Specimens E1 through E6, and
total). Post-hoc LSD was used to perform pairwise comparisons between treatment levels.

Preliminary Study: Effect Size

In order to obtain an estimate of treatment effect size, a preliminary study was performed.
Specimens were treated with a saline sham injection and a series of BCP treatments, spanning a
range of concentrations based on observations of BCP concentration in human synovial fluid.\(^{23,24}\)

A skeletally mature male Wistar or Sprague-Dawley rat cadaver was sourced from ongoing
IACUC-approved research within the institution. A sample population (\(n = 2\)) of pilot specimens
was prepared, consisting of the left and right limbs of a single animal. The COF of each
specimen was evaluated in its naïve state. An injection of 100 \(\mu\)L of phosphate buffered saline
(PBS) was delivered into the joint space, and the specimen was shrouded in PBS-soaked gauze

\(^\dagger\) Validation algorithm identifies changes of slope and evaluates the time intervals between them. Peaks occurring in
rapid succession are discarded as invalid. This eliminates peaks attributable to system noise when pendulum is
stationary, as well as occasional “hiccups” in tracking data.
and allowed to equilibrate for at least 10 minutes prior to further testing. A series of further treatments, at 1, 10, and 100 mg/mL concentrations, were delivered following the same procedure, and COF was evaluated after each treatment. For all trials, pendulum was deflected 12°. Six motion trials were recorded at each treatment level for each specimen.

Main Study: Impact of BCPs on Joint Friction

Three skeletally mature male Wistar and Sprague-Dawley rat cadavers were sourced from ongoing IACUC-approved research within the institution. A sample population (n = 6) of pilot specimens was prepared, consisting of the left and right limbs of three animals. The COF of each specimen was evaluated in its naïve state. An injection of 100 μL of phosphate buffered saline (PBS) was delivered into the joint space, and the specimen was shrouded in PBS-soaked gauze and allowed to equilibrate for at least 10 minutes before reevaluation of COF. A second treatment consisting of 100 μL of 1 mg/mL BCP suspension was delivered following the same procedure, and COF was reevaluated. For all trials, pendulum was deflected 12°. Six motion trials were recorded at each treatment level for each specimen.

Results

Preliminary Study

Delivery of 100 μL of PBS into the joint space did not result in a significant increase in combination fit COF $u$ ($p = 0.131$). Subsequent treatments with 100 μL of 1, 10, and 100 mg/mL BCP suspension all resulted in significant increases in COF ($p < 0.001$ for all BCP treatments). Treatment at 1 mg/mL level resulted in a 69.8% increase in COF over the preceding treatment (PBS), from 0.023±0.002 (mean±SD) to 0.039±0.012 (+97.6% over naïve condition). Treatment at 10 mg/mL level resulted in a 70.2% increase over the preceding treatment (1 mg/mL), from 0.039±0.012 to 0.054±0.020 (+169.9% over naïve condition). Treatment at 100 mg/mL level resulted in a 38.4% increase over the preceding treatment (10 mg/mL), from 0.054±0.020 to 0.075±0.022 (+273.6% over naïve condition). Values of $u$ and $\mu$ for all specimens and treatment levels are presented in Tables I and II, respectively.

Table I : Combination Fit Coefficient of Friction $u$

Mean values and standard deviations of $u$ for individual specimens and total at all treatment levels. * denotes values significantly higher than preceding treatment level. For each specimen, n = 6 trials per treatment level. Totals combine all trials across all specimens at each treatment level (n = 12).

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Naïve</th>
<th>St.Dev.</th>
<th>PBS</th>
<th>St.Dev.</th>
<th>1</th>
<th>St.Dev.</th>
<th>10</th>
<th>St.Dev.</th>
<th>100</th>
<th>St.Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>P3</td>
<td>0.018</td>
<td>0.001</td>
<td>0.022</td>
<td>0.002</td>
<td>0.049*</td>
<td>0.008</td>
<td>0.072*</td>
<td>0.006</td>
<td>0.094*</td>
<td>0.010</td>
</tr>
<tr>
<td>P4</td>
<td>0.022</td>
<td>0.001</td>
<td>0.024</td>
<td>0.002</td>
<td>0.030*</td>
<td>0.001</td>
<td>0.035*</td>
<td>0.001</td>
<td>0.055*</td>
<td>0.008</td>
</tr>
<tr>
<td>Total</td>
<td>0.020</td>
<td>0.002</td>
<td>0.023</td>
<td>0.002</td>
<td>0.039*</td>
<td>0.012</td>
<td>0.054*</td>
<td>0.020</td>
<td>0.075*</td>
<td>0.022</td>
</tr>
</tbody>
</table>
Figure IV: Combination Fit COF $u$ vs. Concentration, Individual Specimens

Figure V: Combination Fit COF $u$ vs. Concentration, Total * denotes statistically significant interval.
Effect size and observed power calculations were performed in order to obtain an estimate of the experimental group size needed for the main study. These calculations used the linear fit COF $\mu$, as the code for the combination fit was not working properly at the time. Based upon these calculations, an experimental group size of $n = 6$ limbs was selected. For $\mu$, the Cohen’s $d$ effect size at the 1 mg/mL treatment level (vs. PBS) was found to be 2.239, and the observed experimental power at 95% confidence and $n = 6$ was found to be 0.753. When recalculated later, using the combination fit COF $u$, the effect size at the 1 mg/mL treatment level (vs. PBS) was found to be 1.949, and the observed experimental power at 95% confidence and $n = 6$ was found to be 0.649. However, the estimated number of specimens required to achieve an experimental power greater than 0.80 ($n = 10$) exceeded the number of specimens available within the project timeline, and the main study was conducted with $n = 6$. Effect sizes for $\mu$ and $u$ at all treatment levels are presented in Table III.

Table II: Linear Fit Coefficient of Friction $\mu$
Mean values and standard deviations of $\mu$ for individual specimens and total at all treatment levels. * denotes values significantly higher than preceding treatment level. For each specimen, $n = 6$ trials per treatment level. Totals combine all trials across all specimens at each treatment level ($n = 12$).

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Naïve</th>
<th>St.Dev.</th>
<th>PBS</th>
<th>St.Dev.</th>
<th>1</th>
<th>St.Dev.</th>
<th>10</th>
<th>St.Dev.</th>
<th>100</th>
<th>St.Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>P3</td>
<td>0.028</td>
<td>0.001</td>
<td>0.029</td>
<td>0.001</td>
<td>0.055*</td>
<td>0.005</td>
<td>0.076*</td>
<td>0.007</td>
<td>0.107*</td>
<td>0.005</td>
</tr>
<tr>
<td>P4</td>
<td>0.033</td>
<td>0.001</td>
<td>0.032</td>
<td>0.000</td>
<td>0.038*</td>
<td>0.002</td>
<td>0.041*</td>
<td>0.001</td>
<td>0.064*</td>
<td>0.006</td>
</tr>
<tr>
<td>Total</td>
<td>0.031</td>
<td>0.003</td>
<td>0.031</td>
<td>0.002</td>
<td>0.046*</td>
<td>0.010</td>
<td>0.058*</td>
<td>0.019</td>
<td>0.085*</td>
<td>0.023</td>
</tr>
</tbody>
</table>

Table III: Cohen’s $d$ Effect Sizes vs. PBS
Cohen’s $d$ effect size relative to PBS for all other treatment levels.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>$\mu$</th>
<th>$u$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Naïve</td>
<td>0.021</td>
<td>1.418</td>
</tr>
<tr>
<td>1</td>
<td>2.239</td>
<td>1.949</td>
</tr>
<tr>
<td>10</td>
<td>2.078</td>
<td>2.165</td>
</tr>
<tr>
<td>100</td>
<td>3.834</td>
<td>3.271</td>
</tr>
</tbody>
</table>

Main Study

Delivery of 100 $\mu$L of PBS into the joint space did not result in a significant increase in combination fit COF $u$ ($p = 0.905$). Subsequent treatment with 100 $\mu$L of 1 mg/mL BCP suspension resulted in a significant increase in $u$ ($p < 0.001$). Treatment at 1 mg/mL level resulted in a 30.3% increase in $u$ over the preceding treatment (PBS), from 0.031±0.009 to 0.040±0.011 (+31.0% over naïve condition). Values of $u$ and $\mu$ for all specimens and treatment levels are presented in Tables IV and V, respectively.
Table IV: Combination Fit Coefficient of Friction $u$
Mean values and standard deviations of $u$ for individual specimens and total at all treatment levels. * denotes values significantly higher than preceding treatment level. † 3 trials removed from E5 at PBS level due to processing error. For all other specimens, n = 6 trials per treatment level. Totals combine all trials across all specimens at each treatment level (n = 33)

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Naïve</th>
<th>St.Dev.</th>
<th>PBS</th>
<th>St.Dev.</th>
<th>1</th>
<th>St.Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>E1</td>
<td>0.030</td>
<td>0.001</td>
<td>0.038</td>
<td>0.002</td>
<td>0.053*</td>
<td>0.005</td>
</tr>
<tr>
<td>E2</td>
<td>0.012</td>
<td>0.002</td>
<td>0.018</td>
<td>0.006</td>
<td>0.026*</td>
<td>0.001</td>
</tr>
<tr>
<td>E3</td>
<td>0.045</td>
<td>0.004</td>
<td>0.040</td>
<td>0.004</td>
<td>0.053*</td>
<td>0.004</td>
</tr>
<tr>
<td>E4</td>
<td>0.036</td>
<td>0.021</td>
<td>0.027</td>
<td>0.001</td>
<td>0.034*</td>
<td>0.001</td>
</tr>
<tr>
<td>E5</td>
<td>0.032</td>
<td>0.005</td>
<td>0.038†</td>
<td>0.009†</td>
<td>0.043*</td>
<td>0.007</td>
</tr>
<tr>
<td>E6</td>
<td>0.029</td>
<td>0.002</td>
<td>0.028</td>
<td>0.002</td>
<td>0.032*</td>
<td>0.002</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>0.031</td>
<td>0.013</td>
<td>0.031</td>
<td>0.009</td>
<td>0.040*</td>
<td>0.011</td>
</tr>
</tbody>
</table>

Figure VI: Combination Fit COF $u$ vs. Treatment, Individual Specimens
Figure VII: Combination Fit COF $\mu$ vs. Treatment, Total  * denotes statistically significant interval.

Table V: Linear Fit Coefficient of friction $\mu$
Mean values and standard deviations of $\mu$ for individual specimens and total at all treatment levels. * denotes values significantly higher than preceding treatment level. † 3 trials removed from E5 at PBS level due to processing error. For all other specimens, $n = 6$ trials per treatment level. Totals combine all trials across all specimens at each treatment level ($n = 33$)

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Naïve</th>
<th>St.Dev.</th>
<th>PBS</th>
<th>St.Dev.</th>
<th>Total</th>
<th>St.Dev.</th>
</tr>
</thead>
<tbody>
<tr>
<td>E1</td>
<td>0.041</td>
<td>0.001</td>
<td>0.049</td>
<td>0.001</td>
<td>0.062*</td>
<td>0.002</td>
</tr>
<tr>
<td>E2</td>
<td>0.042</td>
<td>0.001</td>
<td>0.039</td>
<td>0.001</td>
<td>0.045*</td>
<td>0.002</td>
</tr>
<tr>
<td>E3</td>
<td>0.052</td>
<td>0.002</td>
<td>0.048</td>
<td>0.002</td>
<td>0.065*</td>
<td>0.002</td>
</tr>
<tr>
<td>E4</td>
<td>0.050</td>
<td>0.011</td>
<td>0.041</td>
<td>0.001</td>
<td>0.050*</td>
<td>0.000</td>
</tr>
<tr>
<td>E5</td>
<td>0.051</td>
<td>0.004</td>
<td>0.055†</td>
<td>0.005†</td>
<td>0.063*</td>
<td>0.005</td>
</tr>
<tr>
<td>E6</td>
<td>0.036</td>
<td>0.001</td>
<td>0.033</td>
<td>0.001</td>
<td>0.037*</td>
<td>0.001</td>
</tr>
<tr>
<td>Total</td>
<td>0.045</td>
<td>0.008</td>
<td>0.043</td>
<td>0.007</td>
<td>0.053*</td>
<td>0.011</td>
</tr>
</tbody>
</table>
Discussion and Conclusions

In both the preliminary and the main studies, the whole-joint coefficient of friction was found to increase with the introduction of basic calcium phosphate particulates into the joint space, and COF was found to increase with increasing BCP concentration. The range of naïve combination fit COF values (\(u\)) observed in these studies (preliminary: \(u = 0.020\pm0.002\), main: \(u = 0.031\pm0.013\)) are similar to those reported by Drewniak in BL6 mice (\(u = 0.022\pm0.007\), range 0.016-0.037), and within an order of magnitude of those reported by Akelman in Hartley guinea pigs with a similar pendulum mass (\(u = 0.08\) at 300 g, range 0.042-0.117).²,²⁵

In the main study group, linear fit COF \(\mu\) was 0.045 0.008 in naïve specimens, increasing to 0.053±0.011 at the 1 mg/mL treatment level. In the preliminary study, higher concentrations resulted in \(\mu\)-values as high as 0.085±0.023 (100 mg/mL). In mice, Drewniak reports a baseline linear fit COF (\(\mu\)) of 0.036±0.010 in lubricin-producing (Prg4+/+) animals and a baseline \(\mu\) of 0.077±0.001 in totally lubricin-deficient animals (Prg4-/−).³ Prg4-/− mice display precocious joint disease, as do congenitally lubricin-deficient humans (persons with camptodactylarthropathy-coxa vara-pericarditis syndrome), suggesting a link between high joint friction and arthropathy.³

Due to the large size of the introduced BCP particles relative to the thickness of the articular cartilage layer—more than 10% of the overall cartilage thickness—and the immediacy of the effect, it seems plausible that the observed increase in friction in these studies is attributable to third-body wear between the hard particulate and the softer articulating surfaces of the joint. If this is in fact the case, then when subjected to sustained loading and motion, either in vitro or in vivo, the presence of particulate would likely contribute substantially to mechanical wear of cartilage and the production of wear debris. As is seen in mechanical wear in joint replacements, wear debris can cause inflammation, and also contributes to additional wear. Some calcium phosphate species, such as octacalcium phosphate, can also cause inflammatory responses in vivo.²⁷,²⁸,²⁹ Inflammation, in turn, also contributes to cartilage degradation. Thus, there are multiple mechanisms by which the presence of BCP particulate may contribute to the progression of OA.

The specimens used in these studies comprise a sample of convenience, as they were sourced from other research ongoing within the institution. Thus, there were substantial differences in bodyweight, age at death, age postmortem, and time and temperature of storage between animals. Variations in specimen storage time and temperature after preparation contribute further to this variability. Drewniak reports changes in frictional properties after a single freeze-thaw cycle; an unknown and uncontrolled number of freeze-thaw cycles is likely a substantial confounding factor.²⁶ Variability in specimen size may contribute to variability in potting alignment, and the potting process itself is not yet fully reproducible even when fixing the same specimen multiple times. As constructed, the potting jig obscures the positions of the bones from many angles during radiography and fluoroscopy, making it difficult to confirm specimen position prior to injection of potting medium. A revision of the jig design allowing for greater ease of fluoroscopy, employing more radiolucent materials, would contribute substantially to potting quality and reproducibility.
Another consideration is the use of a single type and size of chemically-produced BCP particulate. If the frictional effects observed are indeed due to third body wear interactions, then the size and shape of the particles will directly impact their frictional effects. Furthermore, the BCP species which occur in vivo are not necessarily identical in composition or form to those available from laboratory supply companies. The extent to which BCP size, shape, and provenance influence their effects on joint friction are unknown as of yet, and are beyond the scope of this study. In spite of these limitations, however, this work comprises the first known investigation of the effects of BCP particulates on joint tribology in any organism. In light of the dramatic result obtained, this area of research certainly merits further investigation. Of especial interest are investigation of wear as a result of cyclic loading in the presence of introduced particles, and perhaps ultimately measurement of COF and cartilage wear and degradation after in-vivo introduction of particles (as studied in rabbits by Fam et al.) or loading-induced OA.4

Acknowledgements

My sincerest thanks to:

Professor Bruce Beynon, for taking me in and finding me something to do when I showed up out of the blue a year ago, bewildered and nervous.

Professor Maria Roehmoldt and Mack Gardner-Morse, for working with me on this project and for teaching me so many things. This has been as much your project as mine.

Professor Rachael Oldinski, for mentoring, advice, and moral support over the past two years.

My parents, for all their love and support.

I could not have done this without all of you. Thank you.
References


Appendix I: SolidWorks Drawings

Figure Ia: Base
Figure Ib: Mounting Block
Figure Ic: Pendulum, Original Design Used in Preliminary Study

- Upper crosspiece, with socket for potted specimen
- Side member, with paired holes for adjustment of lower crosspiece
- Lower crosspiece, with through holes for weight plates
Figure Id: Pendulum, Revised Design Used in Main Study
Appendix II: MATLAB Code

% RATS knee Coefficient of Friction analysis *
% M-File which calculates the rat knee coefficient of friction
% from pendulum data given file and directory names.
% NOTES: 1. Both pendulum resting and pendulum swinging trial
% data files must be available.
% 2. Data for different rat knees must be in different
% subdirectories of the current directory.
% 3. The M-files cof_pend_lin_and_exp.m, cof_pend_lin_and_exp2d.m, cycle.m, exp_fit.m, np_read.m,
% regres1.m and slope.m must be in the current path or
% directory.
% 28-Mar-2013 * Mack Gardner-Morse and Charlotte Morgan
%
% Directory (Knee) Names
% dnams = ['Pilot Specimen 1'
%          'Pilot Specimen 2'];
% ndir = size(dnams,1);
% Resting Data File Names
% fnamrs = ['P1_T2_rest.csv'
%          'P2_T1_rest.csv'];
% Pendulum Data File Names
% fnams = {'P1_T2_1.csv'
%          'P1_T2_2.csv'
%          'P1_T2_3.csv'
%          'P1_T2_4.csv'
%          'P1_T2_5.csv'
%          'P1_T2_6.csv';
%          'P2_T1_1.csv'
%          'P2_T1_2.csv'
%          'P2_T1_3.csv'
%          'P2_T1_4.csv'
%          'P2_T1_5.csv'
%          'P2_T1_6.csv'};
% Loop through Directories
% for k = 1:ndir
%    ddir = dnams(k,:);
%    fnamr = fullfile(ddir,fnamrs(k,:));
%    ntrials = size(fnams{k},1);
% % Loop through Trials
% 
for l = 1:ntrials
    fnam = fullfile(ddir,fnams{k}(1,:));
    cof_pend_lin_and_exp(fnamr,fnam);
    cof_pend_lin_and_exp2d(fnamr,fnam);
pause;
close all;
end
end

************************************************************************
%######################################################################
%             * COF pendulum linear and exponential *
%     M-file which performs linear and exponential fits and calculates
%       the rat knee coefficient of friction for AP pendulum data.
%     NOTES: 1. The M-files cycle.m, exp_fit.m, np_read.m,
%             regres1.m and slope.m must be in the current path or
%             directory.
%######################################################################

function [cofp,c,u,r2,rmse,exp_rmse] =
cof_pend_lin_and_exp2d(fnamr,fnam,istart);
% % Clear Workspace
% %
% clear all;
% close all;
% %
% Check Inputs
% if nargin<3||isempty(istart)||isnan(istart)||istart<1
    istart = 1;
end
% % Global Variables
% %
global g l m Ip k r theta_0
% %
g = 9806.65/1000;        % Acceleration of gravity in m/s^2
l = 115.37/1000;        % Distance between the pendulum center of rotation
and center of gravity in m
m = 291.14/1000;        % Pendulum mass in kg
Ip = 7524736.22/1.0e+9;  % Pendulum inertia about center of rotation in kgm^2
k = 0.0;                % Assume joint stiffness in physiological range is
minimal
r = 2.79/1000;          % Radius of femoral condyles in m
% % Distance between Pendulum Center of Rotation and Center of Markers
%
lm = 155.30; \hspace{1cm} \% Distance between the pendulum center of rotation and center of markers in mm

\% Read NP Data
\% t - time
\% pos - XYZ position in m
\% Determine zero position from resting data
\% fnam = 'P1_T2_rest.csv';
[t_rest,pos_rest] = np_read(fnamr);
\% Read in tracking data
\% fnam = 'P1_T2_6.csv';
[t,pos]=np_read(fnam);
idot = strfind(fnam,'.');
tnam = fnam(1:idot(end)-1);
% Subtract Initial Position (Zero)
% offset = mean(pos_rest); \hspace{1cm} \% Initial position - averaged to remove noise
pos0 = pos-repmat(offset,size(pos,1),1);
pos0 = pos0*1000; \hspace{1cm} \% Convert to millimeters (mm)
% Plot Raw Data
% hfl1 = figure;
orient landscape;
plot(t,pos0(:,1)); \hspace{1cm} \% Plot AP position (X)
% Find and Plot Changes in Slope
% idx = slope(pos0,1,1); \hspace{1cm} \% vector containing indices to all slope changes
figure(hfl1);
hold on;
plot(t(idx),pos0(idx,1),'go');
% Find Where Changes in Slope are Greater Than Zero
% ip = find(pos0(idx,1)>0); \hspace{1cm} \% vector containing indices to maxima in idx
idxp = idx(ip); \hspace{1cm} \% vector containing indices to maxima in pos0
plot(t(idxp),pos0(idxp,1),'rs');
% Find Valid Maxima - At Least 0.20 s Apart
% dt = diff(t(idxp)); \hspace{1cm} \% time differences between all maxima
idt0 = find(dt>0.6&dt<1.4); \hspace{1cm} \% Find peaks ~1 s apart
idt0 = idt0(1); \hspace{1cm} \% Start of valid peaks
idt = find(dt<0.2); \hspace{1cm} \% Not valid peaks
il = find(idt>idt0); \hspace{1cm} \% Find not valid peaks past start of valid peaks
if ~isempty(il)
    idt1 = idt(il(1))-1; \hspace{1cm} \% Stop of valid peaks - one before first
    % not valid peak
else
    idt1 = length(idxp);
end
id = (id0:idt1)'; % Index to slope changes greater than zero
idp = idxp(id); % Index to all valid peaks in pos0
idp = idp(istart:end); % Start at istart peak
plot(t(idp),pos0(idp,1), 'kx');

xlabel('Time (s)', 'FontSize', 12, 'FontWeight', 'bold');
ylabel('AP Displacement (mm)', 'FontSize', 12, 'FontWeight', 'bold');
title(tnam, 'FontSize', 16, 'FontWeight', 'bold', 'interpreter', 'none');
legend('AP Data', 'Negative Peaks', 'Positive Peaks', 'Valid Peaks');

% Convert Lateral Displacements to Angles in Degrees
% angp = atan(pos0(idp,1)/l); % Computes displacement angle from pendulum length l, displacement
angp = asin(pos0(idp,1)/lm); % Computes displacement angle from pendulum length l, displacement
theta_0 = angp(1); % Initial angular peak (theta_0) in radians

% Fit Linear Line to Peaks
np = (1:length(angp))'; % Cycle number for positive slope changes
[cp,r2,df,msse] = regres1(np,angp);
rmse = sqrt(msse); % Root mean square error of the residuals in radians
rmse = rmse*180/pi; % Root mean square error of the residuals in degrees
angp = angp*180/pi; % Angular peaks in degrees for exp_fit

% Fit Exponential Line to Peaks
options = optimset('MaxFunEvals',10000,'MaxIter',10000,'TolFun', 1e-20, 'TolX',1e-8); %,'TolX',1e-8
[u_c,resnorm] = lsqcurvefit(@exp_fit,[0.0001 0.0001],np,angp,[0 0],[1 1],options);

u = u_c(1);
c = u_c(2);
w = sqrt((k + m*g*l)/Ip);
period = 2*pi/w;
z = c./(2*w*Ip);
gam = z.*pi./((1-z.^2).^0.5);
delta = u.*m*g*r/(Ip*w^2*theta_0);
beta = (1+exp(-gam))./(1-exp(-gam));
exp_fit_peaks = theta_0*(exp(-2.*gam.*np) - delta.*beta.*(1 - exp(-2.*gam.*np)))*180/pi; %function of cycle number
exp_rmse = (sum((exp_fit_peaks - angp).^2)/(length(angp)-2))^0.5;

hf2 = figure;
orient landscape;
plot(np,angp,'bo','MarkerFaceColor','b');
hold on;
plot(np,polyval(cp,np)*180/pi,'g-','Color',[0 0.5 0]);
plot(np,exp_fit_peaks,'r-');

xlabel('Cycle Number', 'FontSize',12,'FontWeight','bold');
ylabel('AP Angular Peaks (degrees)'); title(tnam,'FontSize',16,'FontWeight','bold','interpreter','none'); legend('AP Angular Peaks','Linear Fit','Combination Fit');

% Use Angle versus Cycle Number Slope to Calculate Friction Coefficient

% Required Inputs:
% 1. Distance from Center of Rotation to Center of Gravity (l)
% 2. Epicondylar Radius (r)

% COF = l*slope/4*r

d = 4*r; % Denominator for Stanton equation
cofp = l*abs(cp(1))/d;

% Print Out COF

fid = fopen('cof.csv','at');
fprintf(fid,'%s,%s,%i,%12.8f,%12.8f,%12.8f,%12.4f,%12.8f,%12.8f\n', ... 
  fnam,'AP positive slope changes',istart,cofp,c,u,r2,rmse, ... 
  exp_rmse);
fclose(fid);

% Print Graphs

pnam = [tnam '.ps'];
figure(hf1);
if exist(pnam,'file')
  print('-dpsc2','-append',pnam);
else
  print('-dpsc2',pnam);
end

figure(hf2);
print('-dpsc2','-append',pnam);

return

**************************************************************************
% * COF pendulum linear and exponential 2D*
% M-File which performs linear and exponential fits and calculates
% the rat knee coefficient of friction for 2D pendulum data.
% NOTES: 1. The M-files cycle.m, exp_fit.m, np_read.m,
% regres1.m and slope.m must be in the current path or
directory.

function [cofp,c,u,r2,rmse,exp_rmse] =
cof_pend_lin_and_exp2d(fnamr,fnam,istart);
% Clear Workspace
% clear all;
% close all;
% Check Inputs
% if nargin<3||isempty(istart)||isnan(istart)||istart<1
  istart = 1;
end
% Global Variables
% global g l m Ip k r theta_0
% g = 9806.65/1000; % Acceleration of gravity in m/s^2
l = 115.37/1000; % Distance between the pendulum center of
% rotation and center of gravity in m
m = 291.14/1000; % Pendulum mass in kg
Ip = 7524736.22/1.0e+9; % Inertia about center of rotation in kgm^2
k = 0.0; % Assume joint stiffness in physiological range is minimal
r = 2.79/1000; % Radius of femoral condyles in m
% Distance between Pendulum Center of Rotation and Center of Markers
% lm = 155.30; % Distance between the pendulum center of
% rotation amd center of markers in mm
% Read NP Data
% t - time
% pos - XYZ position in m
% Determine zero position from resting data
% fnam = 'P1_T2_rest.csv';
% [t_rest,pos_rest] = np_read(fnamr);
% Read in tracking data
% fnam = 'P1_T2_6.csv';
% [t,pos]=np_read(fnam);
idot = strfind(fnam,'.');
tnam = fnam(1:idot(end)-1);

% Subtract Initial Position (Zero)
offset = mean(pos_rest); % Initial position - averaged to remove noise
pos0 = pos-repmat(offset,size(pos,1),1);
pos0 = pos0*1000; % Convert to millimeters (mm)

% Plot Raw Data
hf1 = figure;
orient landscape;
plot(t,pos0(:,1),'b.-'); % Plot AP position (X)
hold on;
plot(t,pos0(:,3),'g.-','Color',[0 0.5 0]); % Plot lateral position (Z)

hf2 = figure;
orient landscape;
plot(pos0(:,1),pos0(:,3),'g.-','Color',[0 0.5 0]);
hold on;
axis equal; % Plot Lateral by AP positions

% Get Vector Length
v = sign(pos0(:,1)).*sqrt(sum(pos0(:,[1 3]).*pos0(:,[1 3]),2));

% Plot Vector Data
hf3 = figure;
orient landscape;
plot(t,v(:,1)); % Plot 2D position (X and Z)

% Find and Plot Changes in Slope
idx = slope(v,1,1); % vector containing indices to all slope changes
figure(hf3);
hold on;
plot(t(idx),v(idx,1),'go');

% Find Where Changes in Slope are Greater Than Zero
ip = find(v(idx,1)>0); % vector containing indices to maxima in idx
idxp = idx(ip); % vector containing indices to maxima in v
plot(t(idxp),v(idxp,1),'rs');

% Find Valid Maxima - At Least 0.20 s Apart
dt = diff(t(idxp)); % time differences between all maxima
idt0 = find(dt>0.6 & dt<1.4); % Find peaks ~1 s apart
idt0 = idt0(1); % Start of valid peaks
idt = find(dt<0.2); % Not valid peaks
il = find(idt=idt0); % Find not valid peaks past start of valid peaks
if ~isempty(il)
    idt1 = idt(il(1))-1; % Stop of valid peaks - one before first
else
    idt1 = idt0(1)-1; % not valid peak

idt1 = length(idxp);
end
id = (idt0:idt1)';
% Index to slope changes greater than zero
idp = idxp(id);
% Index to all valid peaks in v
idp = idp(istart:end);
% Start at istart peak
plot(t(idp),v(idp,1),'kx');
%
xlabel('Time (s)','FontSize',12,'FontWeight','bold');
ylabel('2D Displacement (mm)','FontSize',12,'FontWeight','bold');
title(tnam,'FontSize',16,'FontWeight','bold','interpreter','none');
legend('2D Data','Negative Peaks','Positive Peaks','Valid Peaks');
%
figure(hf1);
plot(t(idp),pos0(idp,[1 3]),'rs');
xlabel('Time (s)','FontSize',12,'FontWeight','bold');
ylabel('Displacements (mm)','FontSize',12,'FontWeight','bold');
title(tnam,'FontSize',16,'FontWeight','bold','interpreter','none');
legend('AP Data','Lateral Data','Valid Peaks');
%
figure(hf2);
hl = plot(pos0(idp,1),pos0(idp,3),'rs');
text(pos0(idp,1)+0.2,pos0(idp,3),int2str((1:length(idp))'),...
     'FontSize',12,'FontWeight','bold');
xlabel('AP Displacement (mm)','FontSize',12,'FontWeight','bold');
ylabel('Lateral Displacement (mm)','FontSize',12,'FontWeight','bold');
title(tnam,'FontSize',16,'FontWeight','bold','interpreter','none');
legend(hl,'Valid Peaks');
%
% Convert Lateral Displacements to Angles in Degrees
% angp = atan(v(idp,1)/l); % Computes displacement angle from pendulum length l, displacement
angp = asin(v(idp,1)/lm); % Computes displacement angle from pendulum length l, displacement
theta_0 = angp(1); % Initial angular peak (theta_0) in radians
%
% Fit Linear Line to Peaks
% np = (1:length(angp))'; % Cycle number for positive slope changes
[cp,r2,df,msse] = regres1(np,angp);
rmse = sqrt(msse); % Root mean square error of the residuals in radians
rmse = rmse*180/pi; % Root mean square error of the residuals in degrees
angp = angp*180/pi; % Angular peaks in degrees for exp_fit
%
% Fit Exponential Line to Peaks
% options = optimset('MaxFunEvals',10000,'MaxIter',10000,'TolFun',... %
     1e-20,'TolX',1e-8); %,'TolX',1e-8
[u_c,resnorm] = lsqcurvefit(@exp_fit,[0.0001 0.0001],np,angp,[0 0],[1
1],options); %
u = u_c(1);
c = u_c(2);
w = sqrt((k + m*g*l)/Ip); %
period = 2*pi/w;
\[
z = c. / (2 \cdot w \cdot Ip);
\]
\[
gam = z. * \pi / ((1 - z.^2) ^ .5);
\]
\[
delta = u. * m * g * r / (Ip \cdot w^2 \cdot \theta_0);
\]
\[
beta = (1 + \exp(-\gamma)) / (1 - \exp(-\gamma));
\]
\[
\exp \_fit \_peaks = \theta_0 * (\exp(-2. \cdot \gamma \cdot np) - \delta * \beta \cdot (1 - \exp(-2. \cdot \gamma \cdot np))) \cdot 180 / \pi; \quad \text{%function of cycle number}
\]
\[
\exp \_rmse = (\text{sum((exp_fit_peaks - angp).^2)} / (\text{length(angp)} - 2)) ^ .5;
\]
\[
\% hf4 = figure;
\]
\[
\text{orient landscape};
\]
\[
\text{plot(np,angp,'bo','MarkerFaceColor','b')};
\]
\[
\text{hold on};
\]
\[
\text{plot(np,polyval(cp,np)*180/pi,'g-','Color',[0 0.5 0]);}
\]
\[
\text{plot(np,exp_fit_peaks,'r-')};
\]
\[
\% xlabel('Cycle Number','FontSize',12,'FontWeight','bold');
\]
\[
\text{ylabel('2D Angular Peaks (degrees)','FontSize',12,'FontWeight','bold');}
\]
\[
\text{title(tnam,'FontSize',16,'FontWeight','bold','interpreter','none');}
\]
\[
\text{legend('2D Angular Peaks','Linear Fit','Combination Fit');}
\]
\[
\% \text{Use Angle versus Cycle Number Slope to Calculate Friction Coefficient}
\]
\[
\% \text{Required Inputs:}
\]
\[
\% 1. Distance from Center of Rotation to Center of Gravity (l)
\]
\[
\% 2. Epicondylar Radius (r)
\]
\[
\% \text{COF} = l * \text{slope} / 4 * r
\]
\[
\text{d} = 4 * r; \quad \% \text{Denominator for Stanton equation}
\]
\[
\text{cofp} = l * \text{abs(cp(l))} / d;
\]
\[
\% \text{Print Out COF}
\]
\[
\text{fid = fopen('cof.csv','at');}
\]
\[
\text{fprintf(fid,'s,s,1,%12.8f,%12.8f,%12.8f,%12.8f,%12.8f\n',...}
\]
\[
\text{fnam,'2D positive slope changes',istart,cofp,c,u,r2,rmse, ...}
\]
\[
\text{exp_rmse)};
\]
\[
\text{fclose(fid);}
\]
\[
\% \text{Print Graphs}
\]
\[
\text{pnam = [tnam '.ps'];}
\]
\[
\text{figure(hf1);}
\]
\[
\text{if exist(pnam,'file')}
\]
\[
\text{print('-dpsc2','-append',pnam);}
\]
\[
\text{else}
\]
\[
\text{print('-dpsc2',pnam);}
\]
\[
\text{end}
\]
\[
\text{figure(hf2);}
\]
\[
\text{print('-dpsc2','-append',pnam);}
\]
\[
\text{figure(hf3);}
\]
\[
\text{print('-dpsc2','-append',pnam);}
\]
\[
\text{figure(hf4);}
\]
function [istart,iend,ncyc] = cycle(x,icol,iplot)
%CYLE  Returns the indices to the start of each cycle in a
%       a column of the input matrix.
%       ISTART = CYCLE(X,ICOL,IPlot) returns the index ISTART to the
%       start of each cycle in a column of the input matrix.  ICOL is
%       the column to be used (defaults is the first column [=1]).
%       IPlot plots the column data and the cycle start points
%       (default is no plotting [=0]).  The first and last points are
%       assumed to be the start of the first cycle and the end of the
%       last cycle.
%       [ISTART,IEND] = CYCLE(X,ICOL,IPlot) returns the index IEND to
%       end points of each cycle in a column of the input matrix.
%       [ISTART,IEND,NCYC] = CYCLE(X,ICOL,IPlot) returns the number of
%       cycles found in a column of the input matrix.
%       30-Apr-01

%************************************************************************
% Check for Inputs
% if (nargin<3)
%    iplot = 0;
% end
% if (nargin<2)
%    icol = 1;
% end
% if isempty(icol)
%    icol = 1;
% end
% if (nargin<1)
%    error(' *** ERROR in CYCLE:  An input argument is required.');
% end
% Check that Last Two Inputs are Scalar
% l = length(icol);
% l2 = length(iplot);
% if ((l~=1)&&(l2~=1))
%    error(' *** ERROR in CYCLE: Column and plotting arguments must be scalar.');
% end
% Find Points about Zero
y = x(:,icol);
n = length(y);
% del = 1.02*max(abs(diff(y)));
% idc = find(abs(y)<del);
% Find Half and Full Cycles
% ide = find(diff(idc)>1);
% idb = ide+1;
% ide = ide(3:2:end); % Skip first cycle
% idb = idb(2:2:end); % Skip first cycle (Index is 1)
% ncycm1 = min([length(idb); length(ide)]);
ncyc = ncycm1+1;
% idb = idb(1:ncycm1);
ide = ide(1:ncycm1);
% id1 = idc(idb);
id2 = idc(ide);
% Find Start of each Cycle
% istart = zeros(ncyc,1);
% Start of first cycle
% for k = 1:ncycm1
% id = id1(k):id2(k);
% yc = y(id);
% idp = find(yc>=0);
% if isempty(idp)
% [val,i] = max(yc);
% istart(k+1) = id(i);
% else
% [val,i] = min(yc(idp));
% istart(k+1) = id(idp(i));
% end
% end
% End Points of Cycles
% iend = [istart(2:end)-1; n];
% Plot Cycles
% if (iplot)
% figure
plot(y,'b.-');
hold on;
plot(istart,y(istart),'go');
if nargout>1
    plot(iend,y(iend),'ro');
end
axlim = axis;
plot(axlim(1:2),[0 0],'k:');
title ([int2str(ncyc) ' Cycles']);

pause
close
end

************************************************************************

function t = exp_fit(x,xdata);
% xdata is cycle number
global g l m Ip k r theta_0
    u = x(1);  c = x(2);
    w = sqrt((k + m*g*l)/Ip);
    period = (2*pi)/w;
    z = c/(2*w*Ip);
    gam = z*(pi/((1-z^2)^0.5));
    delta = (u*m*g*r)/(Ip*w^2*theta_0);
    beta = (1+exp(-gam))/(1-exp(-gam));
    t = theta_0*(exp((-2*gam)*xdata) - delta*beta*(1 - exp((-2*gam)*xdata)))*180/pi;
return
************************************************************************

function [time,xyz] = np_read(fnam)
% NP_READ  Opens a Natural Point CSV tracking file and reads the time
% and centroid coordinates of the first trackable rigid body.

37
[TIME,XYZ] = NP_READ(FNAM) reads the CSV file, FNAM, and returns the time in a vector TIME and the X, Y and Z coordinates of the centroid of the first rigid body into a three column matrix, XYZ.

NOTES: 1. Currently only handles first rigid body.
2. For use with Natural Point CSV tracking output files.

26-Sep-2012 * Mack Gardner-Morse

%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
% Check for Input
% if (nargin<1)
%   error(' *** ERROR in np_read:  No CSV file name!');
% end
% Open Input File
% fid = fopen(fnam,'rt');
% % Get Frame Count
% lin = 1;
% while lin~=1
%    lin = fgetl(fid);
%    if length(lin>5)
%       if strmatch(lin(1:4),'info')
%          if strmatch(lin(6:15),'framecount')
%             nframe = eval(lin(17:end));
%             lin = -1;
%          end
%       end
%    end
% end
% % Read Lines from File
% time = zeros(nframe,1);
% xyz = zeros(nframe,3);
% k = 0;
% lin = fgetl(fid);
% while lin~=1
%    if length(lin>5)
%       if strmatch(lin(1:5),'frame')
%          k = k+1;
%          ic = findstr(',' ,lin);
%          time(k) = eval(lin(ic(2)+1:ic(3)-1));
%          xyz(k,:) = eval(['[' lin(ic(5)+1:ic(8)-1) ' ]']);
%       end
%    end
% end
lin = fgetl(fid);
end
%
% Close Input File
% fclose(fid);
%
return

************************************************************************

function [b,r2,df,sse,f,b1se,t] = regres1(x,y);
%REGRES1 Computes least squares linear regression for the model Y =
%        B1*X+B0.
%        B = REGRES1(X,Y) returns the slope B(1,1) and intercept B(1,2)
in vector B (1x2) given the independent X vector (nx1) and
dependent Y vector (nx1).
% [B,R2,DF,SSE,F] = REGRES1(X,Y) returns the coefficient of
determination (R2), the number of regression degrees of
freedom (DF), the mean sum of squared errors (SSE) and the F
value of the goodness of fit (F).
% [B,R2,DF,SSE,F,B1SE,T] = REGRES1(X,Y) returns the standard
error of the slope B(1,1) (B1SE) and t score (T) for the slope
B(1,1) of the linear fit.
% NOTES: 1. The calculation of R2 assumes
        SS_reg+SS_err=SS_tot. While correct for linear
        regression, this is not the most general form for R2.  %
        See p. 491 in "An Introduction to Statistical Methods %
        and Data Analysis" by Lyman Ott, 3rd Ed., 1988 and:
        %
        %
        2. SSE is the mean sum of squared errors (residuals)
        (i.e. the sum of squared errors divided by the degrees
        of freedom DF).

% 25-Sep-09 * Mack Gardner-Morse

% Check for Inputs
if (nargin<2)
    error(' *** ERROR: REGRES1 requires two input arguments.');
end
%
% Check that Both Data Inputs are Vectors
[n 1] = size(x);
[n2 12] = size(y);
%
m = min([n 1]);
m2 = min([n2 l2]);

if ((m~=1) | (m2~=1))
    error(' *** ERROR: REGRES1 only works with data vectors. ')
end

x = x(:);
y = y(:);

[n 1] = size(x);
[n2 12] = size(y);

% Check that both inputs have the same number of rows
if (n~=n2)
    error(' *** ERROR: REGRES1 requires X and Y have the same length. ')
end

% Least Squares Linear Regression: y = b1*x+b0
p = 1;
b = polyfit(x,y,p);

% R^2 and t Score Statistics
if (nargout>1)
    ybar = mean(y);
yhat = polyval(b,x);
dy = y-ybar;
tss = dy'*dy;
dr = yhat-ybar;
ssr = dr'*dr;
if (tss>eps)
    r2 = ssr/tss;
    % Only valid for linear regression, assumes SS_reg+SS_err = SS_tot
else
    r2 = NaN;
end

df = n-p-1;
se = norm(y-yhat)/sqrt(df);
xbar = mean(x);
sx = norm(x-xbar);
b1se = se/sx;
if (se>eps)
    t = b(1,1)/b1se;
else
    t = 0;
end
sse = se*se;
if (sse>eps)
    f = (ssr/p)/(sse/df);
else
    f = 0;
end

return
function [indx,pidx,nidx] = slope(x,icol,iplot)
%SLOPE  Returns the indices to the points with positive and negative
% slopes in a column of the input matrix.
%    INDX = SLOPE(X,ICOL,IPLOT) returns the index INDX to the
%    points at changes in the slope in a column of the input
%    matrix.  ICOL is the column to be used (defaults is the
%    first column [=1]).  IPLOT plots the column data and the index
%    points (default is no plotting [=0]).
%    [INDX,PIDX,NIDX] = SLOPE(X,ICOL,IPLOT) returns the index to
%    points with a positive slope in PIDX and the index to points
%    with a negative slope in NIDX.
%  26-Mar-01 * Mack Gardner-Morse
%
%######################################################################
%
%% Check for Inputs
%%
if (nargin<3)
    iplot = 0;
end
if (nargin<2)
    icol = 1;
end
if (nargin<1)
    error(' *** ERROR in SLOPE:  An input argument is required.');
end
%
%% Check that Last Two Inputs are Scalar
%%
1 = length(icol);
L2 = length(iplot);
if ((1==1)||(L2==1))
    error(' *** ERROR in SLOPE:  Column and plotting arguments must be scalar.');
else
end
%
%% Set Up Index into Matrix
%%
indx = 1;
k = 2;
n = length(x);
%
%% Slope
%%
if (x(2,icol)-x(1,icol))>0
    pos = 1;
else
    neg = 1;
end

% Calculate slopes
s = (x(2:end,icol)-x(1:end-1,icol))/ones(n,1);

% Check for changes in slope
for i = 1:n-1
    if ((s(i+1))>0) & (s(i)<=0)
        indx = [indx,i+1];
    elseif ((s(i+1))<=0) & (s(i)>0)
        indx = [indx,i+1];
    end
end

% Plotting
if iplot
    plot(x,icol,hcolor)
    hold on
    plot(x(indx),icol,hcolor)
end
else
    pos = 0;
end
%
% Look for Turning Points
%
pidx = [];
nidx = [];
%
while k <= n
    j = k-1;
    if pos
        pidx = [pidx; j];
        if (x(k,icol)<x(j,icol))
            pos = 0;
            indx = [indx; j];
            nidx = [nidx; j];
        end
    else
        nidx = [nidx; j];
        if (x(k,icol)>x(j,icol))
            pos = 1;
            indx = [indx; j];
            pidx = [pidx; j];
        end
    end
%
    if (x(k,icol)==x(j,icol))
        error ([' *** ERROR in SLOPE: Input at ' num2str(j) ' and ' ... num2str(k) ' in column ' num2str(icol) ' are the same.']);
    end
    k = k+1;
end
%
indx = [indx; n];
%
if pos
    pidx = [pidx; n];
else
    nidx = [nidx; n];
end
%
% Plot Slopes
%
if (iplot)
    figure
    subplot(2,1,1)
    plot(x,'b.-');
    hold on;
    plot(indx,x(indx,icol),'go');
    plot(pidx,x(pidx,icol),'r.');
    title ('Positive Slopes');
    subplot(2,1,2)
plot(x,'b.-');
hold on;
plot(indx,x(indx,icol),'go');
plot(nidx,x(nidx,icol),'r.');
title('Negative Slopes');

pause(2);
close;

end
Appendix III: Data Processing Plots

Figure IIIa: AP Displacement vs. Time Anterior-posterior (x-direction) displacement is plotted versus time, and slope changes ("peaks") are marked.
Figure IIIb: AP Angular Peaks vs. Cycle Number Valid peak displacements are converted to angular displacement (degrees) and plotted versus cycle number. Linear and combination fits are applied to data. Note the difference in fit between models.
**Figure IIIc: AP and Lateral Displacements vs. Time** AP and lateral (z-direction) displacement are plotted versus time, and valid AP peaks are marked. Markers are also placed at corresponding times on lateral data to help visualize the degree of synchronicity between motion in AP and lateral directions.
Figure IIId: AP Displacement vs. Lateral Displacement  Pendulum displacement in the x-z plane is plotted to help visualize the motion of the pendulum. Valid peaks are marked and numbered, highlighting changes in the orientation of the pendulum’s plane of motion over time.
Figure IIIe: 2D Displacement vs. Time Overall displacement (magnitude of pendulum displacement in x-z plane) is plotted versus time, and slope changes (“peaks”) are marked as before.
Figure IIIf: 2D Angular Peaks vs. Cycle Number  Valid peak displacements are converted to angular displacement (degrees) and plotted versus cycle number, and linear and combination fits are applied to data.