

AN AUTONOMOUS METHOD FOR MEASURING 3D JOINT KINEMATICS FROM 2D
XRAY IMAGES

By

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Dedication placeholder.

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LIST OF ABBREVIATIONS

- TKA Total Knee Arthroplasty. This is the complete or partial resurfacing of the articulating surfaces in the knee.
- TSA Total Shoulder Arthroplasty. This is the complete resurfacing of the articulating surfaces in the shoulder.
- rTSA Reverse Total Shoulder Arthroplasty. This is a TSA procedure where the "ball and socket" mechanism is reversed.
- ML Machine Learning. This is the process of feeding a computer inputs and outputs in order to determine an algorithm that goes from input → output
- CNN Convolutional Neural Network. This is a type of neural network that uses convolution kernels as the operation between each of the layers
- HRNet High Resolution Convolutional Neural Network. This is a specific CNN created by ([ADD CITATION](#)) (<https://github.com/HRNet>)

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The primary function of human synovial joints is to support the dynamic motion of the musculoskeletal system. The diseases that typically affect these systems manifest during movement, with mild to severe pain arising during specific activities or during particular ranges of motion. Unsurprisingly, the financial burden of musculoskeletal diseases is roughly USD 300 billion per year in direct and indirect costs [1]. One of the most common conditions affecting human joints is osteoarthritis, which involves the progressive loss of the cartilage between the joint surfaces over time [65]. A highly effective solution for osteoarthritis is arthroplasty, which involves a partial or complete removal and resurfacing of the affected joint with polymeric and metallic components intended to relieve pain and restore a degree of natural function and motion. Despite being highly effective, roughly 20% of patients receiving total knee arthroplasty express some form of dissatisfaction, usually manifested as pain, instability or stiffness during movement ([4], [63], [10]). Surprisingly, standard clinical musculoskeletal diagnostic methods are entirely static. That is, clinicians do not have at their disposal clinically practical ways to quantify skeletal motion during weight-bearing or dynamic movement when most pain symptoms occur. Unfortunately, most of the tools used to accurately quantify 3D dynamic motion (e.g., 3D motion capture, radiostereometry, fluoroscopic model/image registration) are prohibitively expensive or impractical to use in clinical settings. Methods using single-plane fluoroscopic or flat-panel imaging with 3D-to-2D model-image registration have been used since the 1990s. They have

been shown to provide sufficient accuracy for many clinical joint assessment applications , including natural and replaced knees ([5], [8], [46], [83]), natural and replaced shoulders ([48], [50], [82], [49], [37]), and extremities ([79], [43], [16], [15], [70]). One benefit of this approach is that suitable images can be acquired with equipment commonly found in most hospitals. The main impediment for this technology to be used clinically is the time and expense of human operators to supervise the model-image registration process. If the need for human supervision for model-image registration were eliminated, then fluoroscopic imaging could provide a reliable, inexpensive, and accurate method to provide 3D dynamic joint kinematics in a clinical setting. State-of-the-art techniques for generating kinematics using model-image registration involve numerical optimization techniques that iteratively match bone or implant model projections in dynamic x-ray images ([55], [23], [72]). These methods provide accurate 3D bone or implant kinematics when given a rough initial pose estimate for numerical optimization ([23]). However, these methods still require human input for an initial pose estimate, making them impractical for clinical use. Recent advancements in computational capabilities and machine learning algorithms provide tools that are well-suited to replace human supervision for a range of time-consuming tasks including model-image registration. In particular, convolutional neural networks can be trained to provide the image segmentation and pose-estimation capabilities required to autonomously extract knee implant kinematics from single-plane video fluoroscopy. Neural networks can be trained to segment the pixels belonging to a particular knee implant (femoral or tibial), and this pixel information can be used in a numerical optimizer to generate an implant’s 3D pose. Alternatively, a neural network can be used directly for pose-regression, using image data as input values and 3D object pose as output. This latter technique relies on the network’s ability to extract latent characteristics that determine the pose, not an object-oriented cost function to minimize pose error. This regression approach will be sensitive to study conditions, including implant geometry, projection distance and image size, all of which are “lost” when viewing a single-plane image as only a collection of pixels.

CHAPTER 1

INTRODUCTION

Total Knee Arthroplasty (TKA) is a standard procedure for alleviating symptoms related to osteoarthritis in the knee. In 2018, orthopaedic surgeons performed more than 715,000 TKA operations in the United States [2]. This number is projected to increase to 3.48 million by 2030 [40] due to an aging population and increased obesity rates. While TKA largely relieves symptomatic osteoarthritis, roughly 20% of TKA patients express postoperative dissatisfaction, citing mechanical limitations, pain, and instability as the leading causes [4, 10, 63]. Standard methods of musculoskeletal diagnosis cannot quantify the dynamic state of the joint, either pre- or post-operatively; clinicians must rely on static imaging (radiography, MRI, CT) or qualitative mechanical tests to determine the condition of the affected joint, and these tests cannot easily be performed during weight-bearing or dynamic movement when most pain symptoms occur. Unfortunately, most of the tools used to quantify 3D dynamic motion are substantially affected by soft-tissue artifacts [25, 66, 42], are prohibitively time-consuming or expensive [20], or cannot be performed with equipment available at most hospitals.

Model-image registration is a process where a 3D model is aligned to match an object's projection in an image [11]. Researchers have performed model-image registration using single-plane fluoroscopic or flat-panel imaging since the 1990s. Early methods used pre-computed distance maps [41, 83], or shape libraries [5, 73, 74] to match the projection of a 3D implant model to its projection in a radiographic image. With increasing computational capabilities, methods that iteratively compared implant projections to images were possible [46, 23, 44]. Most model-image registration methods provide sufficient accuracy for clinical joint assessment applications, including natural and replaced knees [8, 7, 38, 12], natural and replaced shoulders [37, 45, 49, 67], and extremities [16, 15, 21]. One of the main benefits of this single-plane approach is that suitable images can be acquired with equipment found in most hospitals. The main impediment to implementing this approach into a standard clinical workflow is the time and expense of human operators to supervise the model-image registration process. These methods require either (1) an initial pose estimate [23, 44], (2) a pre-segmented contour of

the implant in the image [11, 41], or (3) a human operator to assist the optimization routine out of local minima [46]. Each of these requirements makes model-image registration methods impractical for clinical use. Even state-of-the-art model-image registration techniques [23] require human initialization or segmentation to perform adequately.

Machine learning algorithms automate the process of analytical model building, utilizing specific algorithms to fit a series of inputs to their respective outputs. Neural networks are a subset of machine learning algorithms that utilize artificial neurons inspired by the human brain's connections [47]. These networks have shown a great deal of success in many computer vision tasks, such as segmentation [17, 75, 59], pose estimation [78, 35], and classification [39, 56, 57]. These capabilities might remove the need for human supervision from TKA model-image registration.

1.1 Computer Vision Primer

1.1.1 Geometric Transformations

Geometric primitives, such as points, are fundamental building blocks for representing shapes and objects in computer graphics. In this section, we will discuss how points can be represented in 2D and 3D space..

1.1.1.1 2D and 3D Points

In N-dimensional space, a point is represented as a set of N scalars, each representing a magnitude in a particular direction. This can be represented mathematically as a column vector, as shown in Equation 1-1. In the case of 2D space, a point is represented as a 2D column vector, $\mathbf{x} = [x, y]^T$. Similarly, in 3D space, a point is represented as a 3D column vector, $\mathbf{x} = [x, y, z]^T$..

$$\mathbf{x} = \begin{bmatrix} x_1 \\ x_2 \\ \vdots \\ x_{N-1} \\ x_N \end{bmatrix} \in \mathbb{R}^N \quad (1-1)$$

Homogeneous coordinates provide a way to represent points with an additional scale factor, \tilde{w} . This allows us to perform rotations and translations simultaneously and successively using matrix multiplications. Homogeneous coordinates for a point in N-dimensional space are represented as a column vector with N+1 elements, as shown in Equation 1-2. The homogeneous coordinates for a 2D point and 3D point can be represented as $\tilde{\mathbf{x}} = \tilde{w}\bar{\mathbf{x}}$, where $\tilde{w}\bar{\mathbf{x}} = [\tilde{x}_1, \tilde{x}_2, \dots, \tilde{x}_N, \tilde{w}]^T$. In most model-image registration applications, the scale factor \tilde{w} is set to 1..

$$\tilde{\mathbf{x}} = \begin{bmatrix} \tilde{x}_1 \\ \tilde{x}_2 \\ \vdots \\ \tilde{x}_N \\ \tilde{w} \end{bmatrix} = \tilde{w} \begin{bmatrix} \mathbf{x} \\ 1 \end{bmatrix} = \tilde{w}\bar{\mathbf{x}} \quad (1-2)$$

1.1.1.2 2D Transformations

Transformations are operations that change the position, orientation, or shape of an object in 2D space. One of the most basic transformations is a translation, which moves an object by adding a displacement vector to its position. This can be expressed mathematically as shown in Equation 1-3:

$$\mathbf{x}' = \mathbf{x} + \begin{bmatrix} t_x \\ t_y \end{bmatrix} = \mathbf{x} + \mathbf{t}$$

Or, using homogeneous coordinates and matrix multiplication (1-3)

$$= \begin{bmatrix} \mathbf{I} & \mathbf{t} \end{bmatrix} \tilde{\mathbf{x}}$$

Where I is the 2×2 identity matrix

In this equation, \mathbf{x} is the original position of the object, \mathbf{x}' is the transformed position, and \mathbf{t} is the displacement vector that specifies the amount of translation in the x and y directions. The

identity matrix \mathbf{I} is used when expressing the transformation using matrix multiplication, as it represents the identity transformation in the x and y dimensions. Using homogeneous coordinates and matrix multiplication allows for the convenient representation of multiple transformations as a single matrix multiplication, as well as for the composition of transformations (i.e., performing multiple transformations in a specific order).

The next type of 2D transformation is a rotation, which changes the orientation of an object, but not its shape (Eq. 1-4, 1-5).

$$\mathbf{x}' = \mathbf{Rx} \quad (1-4)$$

where

$$\mathbf{R} = \begin{bmatrix} \cos\theta & -\sin\theta \\ \sin\theta & \cos\theta \end{bmatrix} \quad (1-5)$$

This will rotate an object θ in the counter clockwise direction.

It's also possible to perform a rotation and a translation at the same time, by replacing the identity matrix \mathbf{I} in Equation 1-3 with the rotation matrix \mathbf{R} from Equation 1-5. This results in the transformation shown in Equation 1-6. This transformation preserves lengths and angles.

$$\mathbf{x}' = \begin{bmatrix} \mathbf{R}_{2 \times 2} & \mathbf{t} \end{bmatrix} \bar{\mathbf{x}} \quad (1-6)$$

A scaled rotation will change the size of the object by some scalar factor, s (Eq. 1-7); this transformation preserves angles.

$$\mathbf{x}' = \begin{bmatrix} s\mathbf{R}_{2 \times 2} & \mathbf{t} \end{bmatrix} \bar{\mathbf{x}} \quad (1-7)$$

An affine transformation preserves parallelism, and is simply a pre-multiplication by an arbitrary 2×3 matrix (Eq. 1-8).

$$\mathbf{x}' = \mathbf{A}\bar{\mathbf{x}}$$

where

$$\mathbf{A} = \begin{bmatrix} a_{11} & a_{12} & a_{13} \\ a_{21} & a_{22} & a_{23} \end{bmatrix} \quad (1-8)$$

A projection matrix (or perspective transformation) is one that operates on homogeneous coordinates (Eq. 1-9).

$$\tilde{\mathbf{x}}' = \tilde{\mathbf{H}}\tilde{\mathbf{x}} \quad (1-9)$$

To obtain inhomogeneous results, the resultant $\tilde{\mathbf{x}}'$ must be normalized (Eq. 1-10). A projective transformation preserves straight lines.

$$\begin{aligned} x' &= \frac{\tilde{x}}{\tilde{w}} & y' &= \frac{\tilde{y}}{\tilde{w}} \\ &= \frac{h_{11}x + h_{12}y + h_{13}}{h_{31}x + h_{32}y + h_{33}} & &= \frac{h_{21}x + h_{22}y + h_{23}}{h_{31}x + h_{32}y + h_{33}} \end{aligned} \quad (1-10)$$

1.1.1.3 3D Transformations

3D transformations operate on 3D points (Eq. 1-1) in a similar way to 2D transformations. In general, all types of transformations, including translations, rotations, scaled rotations, and projections, can be represented by matrices in 3D space, with the only difference being that the dimensions are increased by one.

One aspect of 3D rotations that adds complexity is the possibility of multiple axes of rotation. This introduces the issue of non-commutativity, which means that the order in which rotations are applied matters. To handle this, we can use the concept of Euler angles, which represent the rotations as composite rotations about canonical axes. These rotations are represented by matrices (Eq. 1-11). Note that these matrices are post-multiplied by each other to determine the final rotation matrix for object-centered rotations. Additionally, given a rotation matrix and the corresponding Euler rotations, it is possible to decompose the matrix into its

composite rotations about the canonical axes.

$$R_x = \begin{bmatrix} 1 & 0 & 0 \\ 0 & c_x & -s_x \\ 0 & s_x & c_x \end{bmatrix} \quad R_y = \begin{bmatrix} c_y & 0 & s_y \\ 0 & 1 & 0 \\ -s_y & 0 & c_y \end{bmatrix} \quad R_z = \begin{bmatrix} c_z & -s_z & 0 \\ s_z & c_z & 0 \\ 0 & 0 & 1 \end{bmatrix} \quad (1-11)$$

3D to 2D Projections Geometric primitives, such as points, lines, and polygons, are the building blocks of computer graphics. These basic shapes can be transformed and combined in various ways to create more complex objects and scenes. In order to represent these 3D primitives and objects in 2D image space, we must apply transformations that manipulate their position, orientation, and other properties. By using these transformations, we can create the illusion of depth and spatial relationships on a flat display. Understanding how to work with primitives and transform them is crucial for creating a wide range of visual effects and graphics in computer graphics.

In computer graphics, one of the most basic methods of projecting three-dimensional objects onto a two-dimensional image plane is an orthographic projection. This projection simply drops the depth component of the object and flattens it onto the image plane (Eq. 1-12). This can be thought of as mimicking a camera with a long focal length, or when the depth of the object is shallow compared to its distance from the camera (also known as a weak perspective projection). In this equation, \mathbf{p} represents a point in 3D space and \mathbf{x} represents the projected point in 2D image space.

$$\tilde{\mathbf{x}} = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 \end{bmatrix} \tilde{\mathbf{p}} \quad (1-12)$$

Perspective projection The most common 3D-2D projection is *perspective projection*, which mimics how humans normally see by taking depth-perception into account. This can be done by scaling each point by its z position relative to the camera (Eq. 1-13). We can also

perform this using homogeneous coordinates (Eq. 1-14).

$$\bar{\mathbf{x}} = \mathcal{P}_z(\mathbf{p}) = \begin{bmatrix} x/z \\ y/z \\ 1 \end{bmatrix} \quad (1-13)$$

$$\tilde{\mathbf{x}} = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \end{bmatrix} \tilde{\mathbf{p}} \quad (1-14)$$

The perspective projection is the cornerstone of model-image registration. If a matrix can be created that mimics the fluoroscopic imaging setup in the clinic, then this offers a strong possibility of re-creating a virtual scene with the same geometry as the actual scene.

1.1.2 Image Formation and Camera Properties

Using our knowledge of geometric transformations and projective geometries, we can build up a model of a camera step by step. We standardize our reference frame by having the z direction along the focal direction of the camera, the x direction to the right, and the y direction such that the right-hand rule is maintained. The origin is at the center of the camera.

We will describe the object of interest as a collection of points,

$\mathbf{p} = [x_i, y_i, z_i, 1]$ for $i = 1, 2, \dots, N$. For a complete picture, any given operation will be performed on all points. For simplicity, the following equations will demonstrate the process on a single point of the object.

First, we need to describe the location and orientation of the object with respect to the camera. This can be done with a 3D homogeneous transformation matrix (translation and rotation) (Eq. 1-15).

$$\tilde{\mathbf{p}}' = \begin{bmatrix} R_{3 \times 3} & \mathbf{t}_{3 \times 1} \\ \mathbf{0}_{1 \times 3} & 1 \end{bmatrix} \tilde{\mathbf{p}} \quad (1-15)$$

Then, we use a projective transformation that determines the perspective scaling between

the objects actual location and the image place at the focal distance. Geometrically, this relationship can be visualized with similar triangles (Fig. 1-1), and quantified using the ratio of lengths (Eq. 1-16). In these equations, f' is the focal distance in units of length.

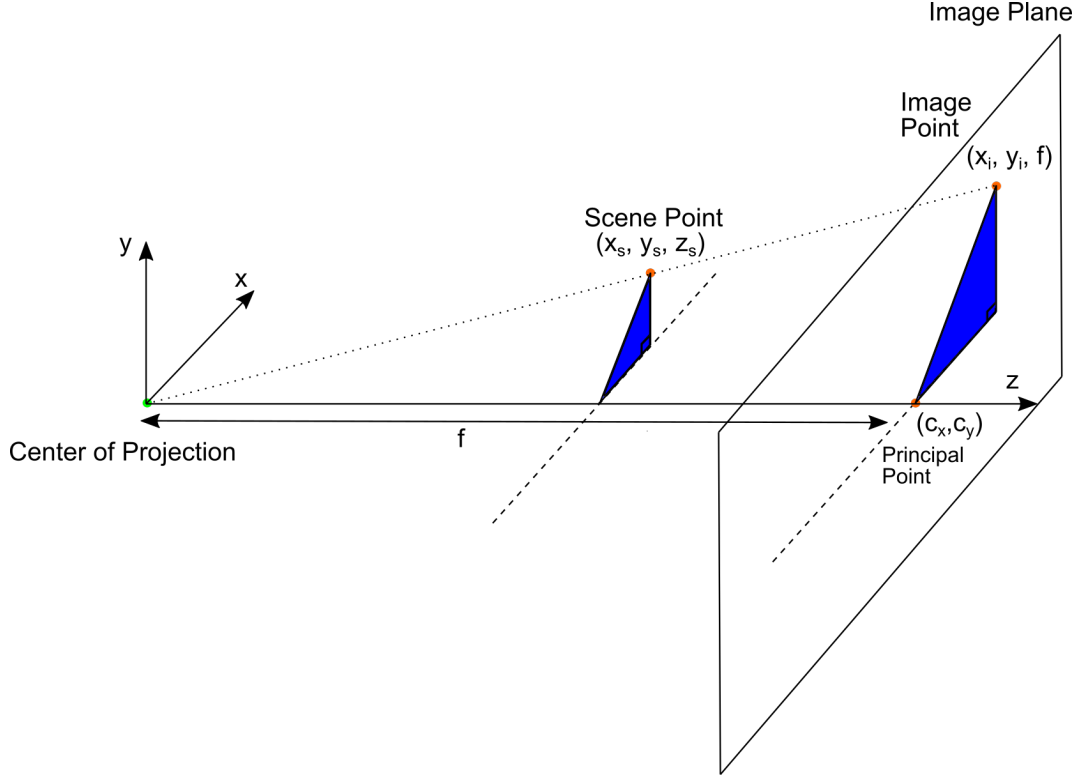


Figure 1-1. The geometry of perspective projection can be visualized by using similar triangles. The overall scaling of the image is based on the ratio of the focal length to the depth of the object.

$$\tilde{\mathbf{x}}_i = \begin{bmatrix} f' & 0 & 0 \\ 0 & f' & 0 \\ 0 & 0 & 1 \end{bmatrix} \tilde{\mathbf{p}}' \quad (1-16)$$

where

$$x_i = p'_x \frac{f'}{p'_z}$$

$$y_i = p'_y \frac{f'}{p'_z}$$

In computer graphics, the principal point, (c_x, c_y) is a key element in the mathematical

model of a camera. It is the point where the optical axis of the camera intersects the image plane, and is typically near the center of the image. The principal point is important because it determines the perspective of the image: objects that are farther away from the principal point will appear smaller in the image, while objects that are closer to the principal point will appear larger. When dealing with the shift between the reference frame embedded in the object and the reference frame embedded in the image plane, the principal point acts as a translation (Eq. 1-17).

$$\tilde{\mathbf{x}}_i = \begin{bmatrix} f' & 0 & c_x \\ 0 & f' & c_y \\ 0 & 0 & 1 \end{bmatrix} \tilde{\mathbf{p}}' \quad (1-17)$$

where

$$c_x \approx \frac{W_{image}}{2}$$

$$c_y \approx \frac{H_{image}}{2}$$

Finally, we need to convert the image coordinates, $\tilde{\mathbf{x}}_i$, to pixel coordinates using the pixel scale factor (Eq. 1-18). The pixel scale factor is defined by the parameters k_x and k_y , which represent the number of pixels per unit distance in the x and y directions, respectively. By multiplying the perspective projection matrix by the pixel scale factor, we can obtain a new matrix that maps 3D points in world coordinates directly to pixel coordinates on the image.

$$\begin{aligned}\tilde{\mathbf{x}}_{pix} &= \begin{bmatrix} k_x & 0 & 0 \\ 0 & k_y & 0 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} f' & 0 & c_x \\ 0 & f' & c_y \\ 0 & 0 & 1 \end{bmatrix} \\ &= \begin{bmatrix} f_x & 0 & c_x \\ 0 & f_y & c_y \\ 0 & 0 & 1 \end{bmatrix}\end{aligned}\tag{1-18}$$

where

$$f_x = k_x f' \text{ and } f_y = k_y f'$$

Are focal distances in units of pixels

1.1.3 Image Processing

Digital image processing is a field of computer vision that deals with the manipulation, analysis, and interpretation of digital images. It focuses on algorithms and techniques that extract meaningful information from images and to enhance visual quality. In fluoroscopy, image processing can be used to enhance the digital image, or determine important information about it such that a the model-image registration task can be performed efficiently and autonomously.

1.1.3.1 Filtering and Convolution

We have already seen how image formation yields a collection of 2D points, \mathbf{x}_{pix} . We can write the intensity values at each pixel locations as a function, $f(\mathbf{x}_{pix}) = f(i, j)$, where (i, j) represent locations in the image, and the function returns the intensity of the image at that particular pixel location. This allows us to treat images as functions, and perform similar functional operations and analysis to extract meaningful information from them.

The most widely used filter is a linear filter [68], where the output is some linear operation on the neighboring pixels (Eq. 1-19), also known as a *convolution*. In a convolution, the kernel, h is shifted along the input image, f , and the resultant image, g , is the dot product of those two matrices at that specific location.

$$\begin{aligned}
g(i, j) &= \sum_{k,l} f(i-k, j-l)h(k, l) \\
&= \sum_{k,l} f(k, l)h(i-k, j-l)
\end{aligned} \tag{1-19}$$

Where we use the following notation

$$g = f * h$$

The convolution operation is *linear shift invariant*, which means that it obeys the superposition principle (Eq. 1-20) and the shift invariance principle (Eq. 1-21). This is a powerful property, because it will behave the same everywhere on the input signal/image, which is useful for different types of feature extraction and filtering operations.

$$h * (f + g) = h * f + h * g \tag{1-20}$$

$$g(i, j) = f(i+k, j+l) \iff (h * g)(i, j) = (h * f)(i+k, j+l) \tag{1-21}$$

A common filter applied to images is the Gaussian kernel (Eq. 1-22). This kernel is shaped as a 2D discrete Gaussian, and has the effect of blurring an image and removing noise.

$$\text{Gaussian filter} = \frac{1}{256} \begin{bmatrix} 1 & 4 & 6 & 4 & 1 \\ 4 & 16 & 24 & 16 & 4 \\ 6 & 24 & 36 & 24 & 6 \\ 4 & 16 & 24 & 16 & 4 \\ 1 & 4 & 6 & 4 & 1 \end{bmatrix} \tag{1-22}$$

Another is the box kernel, which averages the value of the nearest K pixels (Eq. 1-23).

$$\text{Box filter} = \frac{1}{K^2} \begin{bmatrix} 1 & 1 & \dots & 1 \\ 1 & 1 & \dots & 1 \\ \vdots & \vdots & 1 & \vdots \\ 1 & 1 & \dots & 1 \end{bmatrix} \quad (1-23)$$

Edge filters can also be created to detect vertical (Eq. 1-24), horizontal (Eq. 1-25), or diagonal edges (Eq. 1-26). As each of the filters moves across the feature it is designed for, that region of the output will be more highly activated than other regions, extracting out the desired components. The orientation of each of these filters can be hand-selected to find desirable attributes in images.

$$\text{vertical edge filter} = \begin{bmatrix} 0 & 1 & 0 \\ 0 & 1 & 0 \\ 0 & 1 & 0 \end{bmatrix} \quad (1-24)$$

$$\text{horizontal edge filter} = \begin{bmatrix} 0 & 0 & 0 \\ 1 & 1 & 1 \\ 0 & 0 & 0 \end{bmatrix} \quad (1-25)$$

$$\text{diagonal edge filters} = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix} \text{ and } \begin{bmatrix} 0 & 0 & 1 \\ 0 & 1 & 0 \\ 1 & 0 & 0 \end{bmatrix} \quad (1-26)$$

Lastly, we can use a corner filter to find corners in images (Eq. 1-27).

$$\text{Corner filter} = \frac{1}{4} \begin{bmatrix} 1 & -2 & 1 \\ -2 & 4 & -2 \\ 1 & -2 & 1 \end{bmatrix} \quad (1-27)$$

Entire subfields of computer vision and image analysis are devoted to creating increasingly

useful and complex filters, or collections of filters. These include steerable filters, which determine information occurring in arbitrary directions [24], recursive filtering [52], and non-linear filtering [71].

1.1.3.2 Binary Image Processing

A binary image is a digital image that consists only of black and white pixels. It is often used for labeling or masking an underlying image, where the values of 1 and 0 represent the presence or absence of a particular feature or object. Binary images are often used in computer vision and image processing applications because they can be easily processed and analyzed using algorithms. They are also useful for storing and transmitting large amounts of data, as the use of only two values reduces the amount of information that needs to be stored and transmitted.

The primary form of processing on binary images is morphological, that is, changing the shape of the “blob” in order to extract useful information from it.

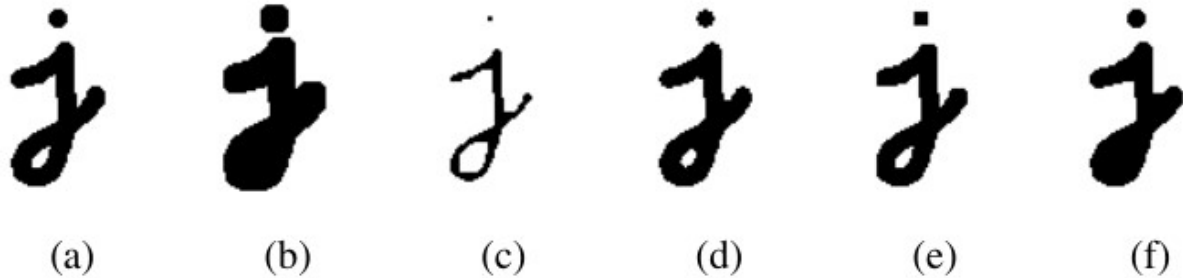


Figure 1-2. A collection of morphological operations on a binary image: (a) original image; (b) dilation; (c) erosion; (d) majority; (e) opening; (f) closing. Image from [68]

Dilation and erosion are the two main operations that are used in model-image registration (1-28). These functions are ach two-fold: first, a convolution operation is applied to the existing binary image, then a threshold is applied to the convolution output to determine if the central pixel is a zero or 1. If f is the input image, s is the convolution kernel of 1s, and $c = f \otimes s$ is the number of 1s in the convolution output, then dilation and erosion can be expressed in the following way.

$$\begin{aligned} \text{dilate}(f, s) &= \theta(c, 1) \\ \text{erode}(f, s) &= \theta(c, S) \end{aligned} \tag{1-28}$$

Where θ represents a thresholding function.

$$\theta(f, t) = \begin{cases} 1 & \text{if } f \geq t \\ 0 & \text{else} \end{cases} \quad (1-29)$$

1.1.4 Image Similarity Metrics

One of the key components in model image registration is image similarity. Fundamentally, this is the method of determining how well the user's synthetic image matches with the actual fluoroscopic image. The choice of similarity metric is going to be determined by many key factors such as the a-priori availability of implant/bone geometry and knowledge of the image quality and contrast. Broadly, there are two classes of image similarity when performing model-image registration: intensity-based and feature-based.

1.1.4.1 Intensity Based

Intensity based measures are those that utilize specific pixel information in order to determine the difference between two images. This can be either a global image similarity metric, or measure the specific regions of interest in the given image.

A canonical difference between two images would be the p-norm separating them (Eq. 1-30), which iterates through each pixel of the two images and finds the p-norm difference each intensity for the pixel pair. Common p-norms are the L_1 norm (*absolute intensity differences* or *mean absolute difference*) [33] ($p = 1$) and the L_2 , or Euclidean, norm (*squared intensity differences* or *mean squared difference*) [27] ($p = 2$).

Intensity-based measures use the pixel values of the images to determine their similarity. These measures can be global, meaning they consider the entire image, or they can focus on specific regions of interest. A common intensity-based measure is the p-norm (Eq. 1-30), which calculates the difference between the intensity of corresponding pixels in the two images. The L_1 norm, also known as the absolute intensity differences or mean absolute difference, uses $p = 1$ in the equation [33], while the L_2 norm, also known as the squared intensity differences or mean squared difference, uses $p = 2$ [27].

$$\|A - B\|_p = \left(\sum_{x=0}^w \sum_{y=0}^h |a_{xy} - b_{xy}|^p \right)^{\frac{1}{p}} \quad (1-30)$$

where A and B are the two images being compared, w and h are the width and height of the images, and a_{xy} and b_{xy} are the intensity values at pixel (x,y) in the two images, respectively.

While conceptually easy to use, the main limitation of p-norm measures is their lack of spatial information. For example, an image that has been shifted by a linear transformation would not score well using a p-norm, despite the two images containing only a minor shift, scale, or rotation. One method for overcoming this limitation is to use the cross-correlation, or sliding dot product, between images [9, 27] (Eq. 1-31). When used in conjunction with projective geometry, this can help locate regions of interest for a model-based registration pipeline. The cross-correlation is calculated using the following equation:

$$\begin{aligned} (A \star B)[x, y] &= E[A_{xy} \cdot B_{x+\tau_x, y+\tau_y}] \\ &= \sum_{\tau_x=-\infty}^{\infty} \sum_{\tau_y=-\infty}^{\infty} a_{xy} b_{x+\tau_x, y+\tau_y} \end{aligned} \quad (1-31)$$

This will have the effect of determining the regions of each image that are similar, causing the correlation function to “light up” at those areas in a similar way to the convolutional operation between two images. The normalized cross-correlation can also be used (Eq. 1-32), which removes noise coming from each of the original images.

$$\text{normalized cross correlation}(A, B) = \frac{A \star B}{(A \star A)(B \star B)} \quad (1-32)$$

1.1.4.2 Feature Based

Feature based image similarity metrics involve some method of determining key features in images, and using those notable features for measuring the differences between two images. These types of methods almost always involve some type of feature-extraction step, where the various features of interest are calculated and determined for subsequent use. The two main classes of features are *keypoints* and *edges*. The simplest method of keypoint detection is using a

similar method to intensity-based matching, but having one of the “images” as a patch of the desired feature. With keypoints detected in the input image, one could determine the error of the current pose estimate by taking the Euclidean distance between all image keypoints and all projected keypoints: [12] (Eq. 1-33). With a-priori information about the keypoints, one could attach a weight to every keypoint in order to emphasize specific regions on the image and the model (Eq. 1-34)

$$\text{Keypoint Error} = \left(\sum_{i=0}^N (KP_{image,i} - KP_{proj,i})^2 \right)^{\frac{1}{2}} \quad (1-33)$$

$$\text{Weighted Keypoint Error} = \left(\sum_{i=0}^N w_i (KP_{image,i} - KP_{proj,i})^2 \right)^{\frac{1}{2}} \quad (1-34)$$

Keypoints are particularly useful when there are invariant features in images and 3D models that will always be present. However, if these features will not, or cannot always be detected, then other measures must be utilized.

Finding Edges

Edge- and contour-based matching algorithms make use of the edges that are present in the image, and aligning that with the projected edges of the 3D model. However, we must first consider the determination of edges in an image. For a human operator, it can be rather easy to find edges of interest, but how much this be incorporated computationally? The first approach might be in viewing an image topologically, with regions of different colors and intensity represented by different “heights”. Then, an edge simply becomes an area with a steep gradient (Eq. 1-35).

$$\mathbf{J}(\mathbf{x}) = \nabla I(\mathbf{x}) = \left(\frac{\partial I}{\partial x}, \frac{\partial I}{\partial y} \right)(\mathbf{x}) \quad (1-35)$$

Finding the direction of the steepest ascent/descent at any given location will give use the normal to the local edge at that point. However, the derivative operator will accentuate and amplify high frequencies in the image, causing noise to overpower the signal. Removing the

high-frequency information (a low-pass filter) in the image results in gradient detection that is much more aligned with the salient edges of the image. The Gaussian kernel is a good option for an isotropic low-pass filter on a 2D signal (image) (Eq. 1-36)

$$\begin{aligned}\mathbf{J}_\sigma(\mathbf{x}) &= \nabla[G_\sigma(\mathbf{x} * I(\mathbf{x}))] \\ &= \nabla G_\sigma(\mathbf{x}) * I(\mathbf{x})\end{aligned}\tag{1-36}$$

where

$$\nabla G_\sigma(\mathbf{x}) = \left(\frac{\partial G_\sigma}{\partial x}, \frac{\partial G_\sigma}{\partial y} \right) = [-x -y] \frac{1}{\sigma^2} \exp\left(-\frac{(x^2 + y^2)}{2\sigma^2}\right)$$

The ubiquitous edge detection algorithm was proposed by John Canny in 1986 [14], which utilizes a five-step process. First, a Gaussian kernel is applied as a low-pass filter (Eq. 1-36), second, directional filters are used to find the gradients in each direction of the image, third, a gradient magnitude threshold is applied to remove noise, fourth, a double threshold is applied to remove both strong and weak edges, and lastly, edges are determined from hysteresis.

Using Edges for Image Similarity

In model-image registration, the similarity of two contours is used as a heuristic for the correct pose. When the projected model's contour aligns accurately with the edges in the fluoroscopic image, one can say that the model is *properly registered* to the image. The main question becomes: how can we computationally determine when two contours are aligned?

As always, the simplest approach is to take the p-norm between the model and image contours (Eq. 1-30), where instead of taking the difference between the two original images, one is taking the difference of the edges of the images. This function will be minimized when there is complete overlap between image and model contours. The primary issue with this formulation is the sensitivity to slight perturbations in the model. This is due to the width of the contour being a single pixel, which would render an extremely high error if the model is shifted just a single pixel in any direction. Because the edge-detected images are binary (0-no edge, 1-edge), we can take advantage of binary morphological operations to change the images to better suit the model-image registration pipeline. The primary operation is dilation (Eq. 1-28), which is useful

because it decreases the sensitivity of the p-norm metric for image similarity, allowing for a smoother curve for optimization routines to find a global minima.

1.1.4.3 Symmetry Traps

Objects with rotational or mirror symmetry cause pathological solutions to many of the image similarity metrics when used for optimizing the pose of the object relative to the image. The simplest example of a symmetry trap can be posed as follows: given the shadow of a basketball, which direction was the logo facing? It is quickly apparent that this is an impossible question to answer with just the information given by the image and the 3D model. This problem arises when performing optimizing for the post of mediolaterally symmetric tibial implants.

Additional information must be used to find the correct pose of the implant.

However, with the knowledge of the direction of symmetry, it is possible to determine the “dual pose” of the current orientation, that is, the pose that produces indistinguishable projective geometry.

Algorithm for Determining the Dual-Pose of a Symmetric Object

1. Determine the viewing ray from camera → object (Eq. 1-37).
2. Determine the axis-angle (m, θ) rotation between the viewing ray and the symmetric axis of the object (Eq. 1-38, 1-39).
3. Rotate the object -2θ about the same axis, reflecting the rotation about the viewing ray (Eq. 1-40).
4. The final orientation of the object is exactly the “dual pose”, producing indistinguishable projective geometry (Eq. 1-41).

If T is the homogenous transformation matrix describing the object

$$\vec{v}' = T_{1:3,4} \quad (1-37)$$

$$\vec{v} = \frac{\vec{v}'}{\|\vec{v}'\|}$$

We can use trigonometry to determine the angle (Eq. 1-38) and perpendicular axis (Eq. 1-39) between two vectors. For our example, we use the normalized viewing ray and the z-axis (symmetric axis) as the two vectors.

$$\cos(\theta) = \vec{v} \cdot \vec{z} \rightarrow \theta = \arccos(\vec{v} \cdot \vec{z}) \quad (1-38)$$

$$\vec{m} = \frac{\vec{v} \times \vec{z}}{\|\vec{v} \times \vec{z}\|} \quad (1-39)$$

Then, we can build a rotation matrix using an axis and an angle [19], (Eq. 1-40).

$$c = \cos(-2\theta)$$

$$s = \sin(-2\theta)$$

$$q = -\cos(-2\theta) \quad (1-40)$$

$$R_{3 \times 3} = \begin{pmatrix} m_x^2 v + c & m_x m_y v - m_z s & m_x m_z v - m_y s \\ m_x m_y v + m_z s & m_y^2 v + c & m_y m_z v - m_x s \\ m_x m_z v - m_y s & m_y m_z v + m_x s & m_z^2 v + c \end{pmatrix}$$

Then, we obtain the final transformation matrix describing the dual pose of the object by a post-multiplication of this rotation matrix.

$$T_{dual} = T_{orig} * \begin{pmatrix} R_{3 \times 3} & \vec{0}_{3 \times 1} \\ \vec{0}_{1 \times 3} & 1 \end{pmatrix} \quad (1-41)$$

Given these two matrices, further exploration can determine which is the correct pose, though this will have to be done using information not directly present in the image contours.

1.2 Deep Learning for Image Processing in Orthopaedics

[39] Alex Krizhevsky sparked renewed interest in deep learning in 2012 by utilizing a convolutional neural network to win the ImageNet challenge [61] by more than 10 points over second place. Since then, neural networks have found their way into many different computer vision tasks, including many in the medical field. Neural networks have paved the way for improved image processing and analysis in orthopaedics utilizing a wide array of medical imaging modalities (CT, X-Ray, MRI) to segment and classify different bones and pathologies that are present in the images. This can improve the speed and accuracy of quantifying information present in images, and in some cases, it can completely remove the need for a human operator to perform common tasks.

Broadly, a neural network is an algorithm

1.2.1 The Telos of Neural Networks for Image Processing

Neural networks for image processing are an attempt to recreate the visual system in animals, creating algorithmic analogies to the various physical neuronal pathways that are present in the image. Typically, there are many different parts of our visual system that we take for granted, such as the ability to see upside down and extrapolate from prior information. While seemingly intuitive, these are difficult feats to determine algorithmically in a mathematical function.

Put images here with the visual system
represented and the zebra picture

1.2.2 Neural Network Structure

Neural networks generally have the same constitutive elements, mixed and matched based on the desired performance and complexity of the model that you are trying to build.

1.2.2.1 Neural Network Building Blocks

Generally, neural networks are formed by collections of foundational units, which can generate increasingly complex architectures and yield incredible performance. However, it is always important to start with the fundamental “atoms” of the neural network.

The most basic unit of a neural network is the perceptron (or neuron), which is composed of a summation of inputs multiplied by weights, a bias term, and a (typically non-linear) activation function (Fig. 1-3, Eq. 1-42).

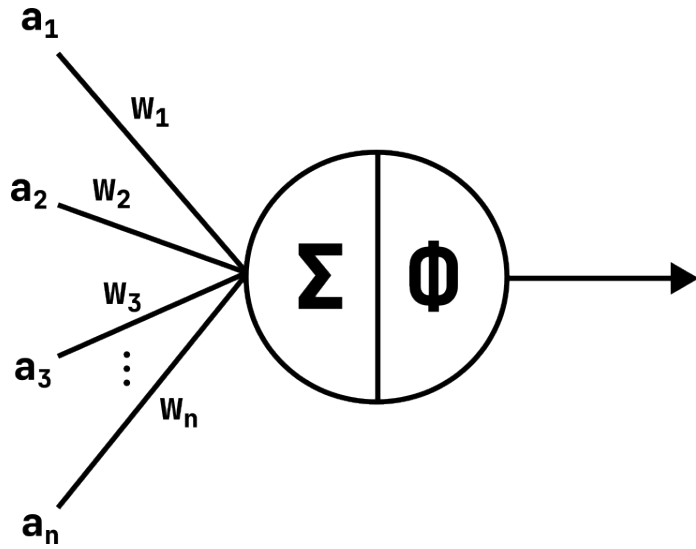


Figure 1-3. A schematic representing a single neuron that receives n inputs and applies ϕ as an activation function.

$$y = \phi\left(\sum_{i=1}^n a_i w_i + b\right) \quad (1-42)$$

Activation functions are a crucial component in neural networks as they allow for the introduction of non-linearity, which is essential for the network’s ability to learn complex representations of the input data. Without activation functions, a neural network would only be able to learn linear relationships between the input and output. However, many real-world problems involve non-linear relationships that cannot be captured by a linear model alone. Activation functions provide a way to move beyond a linear relationship, allowing the neural

network to learn more nuanced mappings between the input and output. The choice of activation function depends on the specific problem you are trying to solve and the architecture of your network. Some activation functions introduce more non-linearity than others and some trade-off between non-linearity and computational efficiency. Experimenting with different activation functions and observing the impact on the network's performance can be a useful technique for optimizing the performance of a neural network. A list of common activation functions and their equations is shown in Table 1-1.

Table 1-1. A list of activation functions and their corresponding mathematical formula

Activation Function	Equation
Linear	$\phi(x) = x$
Sigmoid	$\phi(x) = \frac{e^x}{1+e^x}$
Hyperbolic Tangent	$\phi(x) = \frac{e^x - e^{-x}}{e^x + e^{-x}}$
Rectified Linear Unit (ReLU)	$\phi(x) = \max(0, x)$
Leaky ReLU	$\phi(x) = \max(0.1x, x)$

1.2.2.2 Fully Connected Network

The fully connected network, also known as the multi-layer perceptron, is a basic type of neural network that utilizes the neurons previously discussed as building blocks. Its schematic representation is a familiar image to many when considering neural networks. By stacking the summations and multiplications of each neuron, we can derive the equation for a single layer of a fully connected network, which is simply a matrix multiplication, as seen in Eq. 1-43. In this equation, W represents the collection of weights for each neuron, a represents the input, b represents the bias, and ϕ represents the activation function.

One of the key strengths of this type of network is the utilization of non-linear activation functions. A well-chosen activation function can greatly impact the network's performance and ability to achieve specific tasks. For example, in a binary classification task, the sigmoid activation function can be used at the output layer, constraining the output between 0 (false) and 1 (true). The output can then represent the probability of the given input being classified as 'true'.

The ReLU activation function is another popular choice, it is computationally efficient and often used in hidden layers of deep networks. Additionally, the hyperbolic tangent (\tanh) activation function is used in the context of a model where data follows a Gaussian distribution.

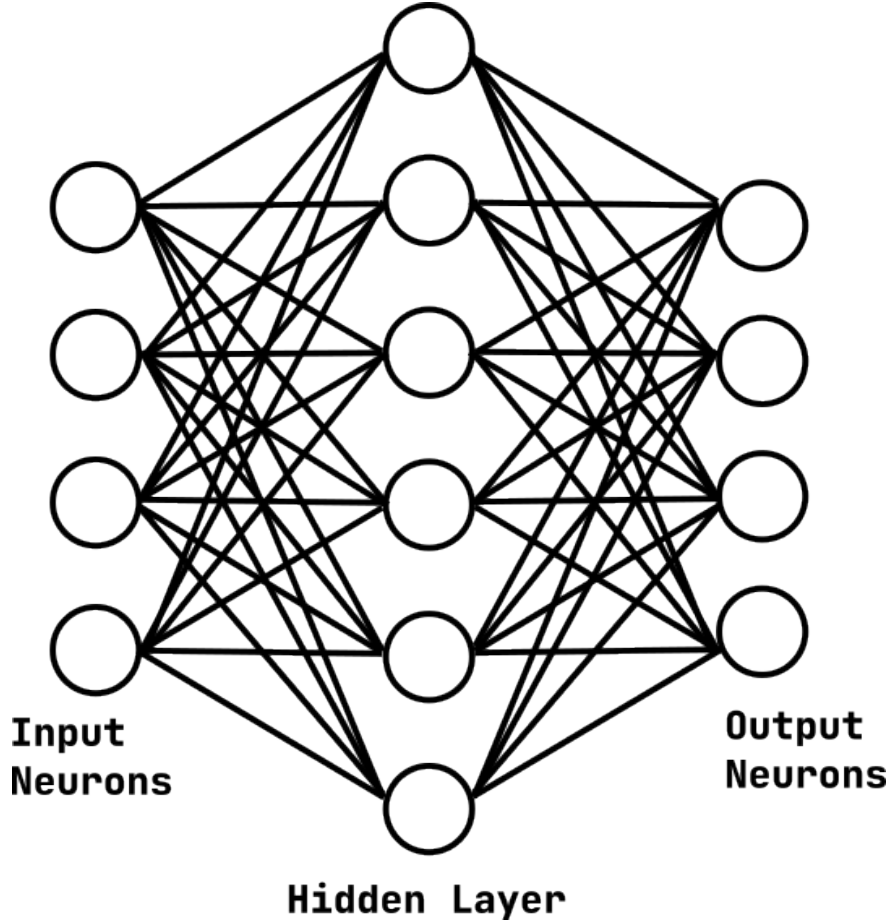


Figure 1-4. A basic fully connected network with a single hidden layer. Each of the neurons are exactly the same as the neurons shown in figure 1-3

$$y = \phi(W^T a + b) \quad (1-43)$$

Additional layers can be added by taking the output of one layer and sending that into the input of the next. Increasingly complex mappings can be generated by increasing either the size or the number of hidden layers in a network.

$$y = \phi_2(W_2^T (\phi_1(W_1^T a + b_1) + b_2)) \quad (1-44)$$

One of the main limitations of fully-connected networks is exponential increase in computational complexity as your input grows. For a standard 1024×1024 image, you have roughly 1 million input nodes, which can lead to hundreds of millions of parameters that need to be learned depending on the depth of the network. We can use standard image processing techniques to overcome some of these limitations as we explore different network architectures.

1.2.2.3 Convolutional Neural Networks

Convolutional neural networks (CNN) utilize the convolution operation (Eq. 1-19) in order to both reduce the size and complexity of the network and capture spatial information present in images. In the same way that the matrix W in the fully connected network is a learnable parameter in a FCN, the individual kernels are the learnable parameters in a convolution operation. In practice, with the correct cost function and optimization routine, this allows each of the kernels to learn the latent structure in the image and make connections between those structures. At a high level, these kernels often represent feature extractors for edges, lines, corners, curves, and other geometric primitives. However, as one traverses deeper into the network, the combinations of these features are often incomprehensible to a human (Fig. 1-5).

Typically, these networks will downsample the image to capture the most salient information, then upsample to regenerate the features from the underlying latent representations. This network architecture has been popularized by the U-Net, which is known as a standard autoencoder (Fig. 1-6). Each of the layers is composed of a collection of convolution kernels, and the downsampling occurs based on the stride or size of the kernel.

CNN architecture can be altered by changing the performance of the underlying kernels, either size, stride, shape, or behavior. Stride controls how many discrete steps the kernel takes as it is moving across the input image. Stride can be used to downsample images. An atrous convolution involves internally padding the convolution with zeros so that each of the entries are separated by a layer of zeros. This also has the effect of more aggressively downsizing an image and capturing a larger region around the center pixel.

A convolutional neural network can have an additional fully connected network appended

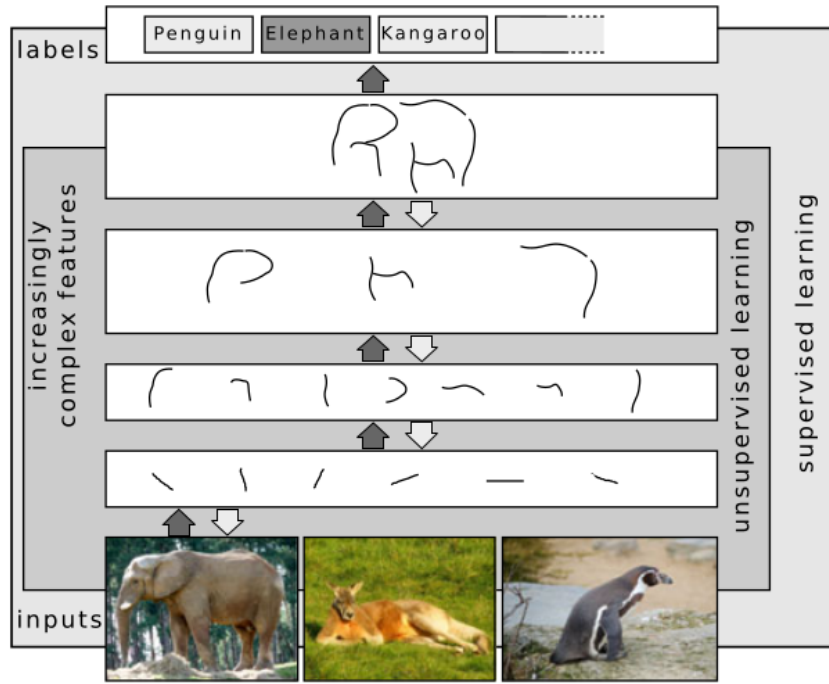


Figure 1-5. An example of extracted features from a convolutional neural network. As shown, the deeper into the network one explores, the combination of core features start to create higher level features that might represent the shape of a specific animal [62].

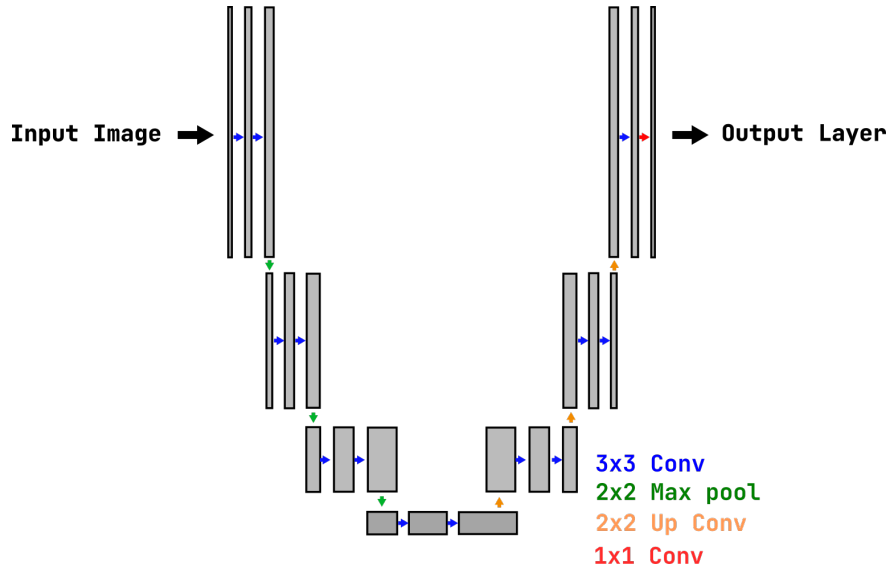


Figure 1-6. A standard U-Net architecture for a convolutional neural network. This architecture is the most common form of network used for biomedical image analysis and processing.

to the output in order to learn a mapping that involves a linear combination of the extracted features from the convolution operations. This is often used when a CNN is applied to a

classification task, where the total number of outputs of the final FCN is the number of classes.

1.2.3 Training Neural Networks

The power of neural networks is that they are able to “learn” incredibly complex mappings from a set of training data. But how *exactly* do they learn?

1.2.3.1 Neural Network Cost Functions

The first step in determine how a machine learns is knowing how correct an output is. The **cost function** is exactly the error metric that determines the wrongness of the neural network’s output. It’s this function that the neural network is attempting to minimize as it updates its weights.

The choice of cost function is dependent on the task at hand. For a regression problem, a standard mean-squared-error is typically employed

PUT TABLE WITH SOME COST FUNCTIONS HERE

1.2.3.2 Optimizing and Updating Weights

For convex problems, there often exists a closed-form solution for the parameters that minimize a given objective function. However, neural networks are highly non-linear and non-convex, forcing researcher to employ different methods of updating parameter values in a direction of minimizing the objective function. Mathematically, this means that we are determining the local gradient of the cost function with respect to each of the parameters, and updating the parameters in the direction of the steepest negative gradient.

If w is the collection of all learnable parameters, then we are trying to minimize the error between the $y = \text{model}(x)$ and our true value, t . In equation 1-45, $F(\cdot)$ is our objective function, y is our model output, and t is the true label.

$$J(\text{model}) = F(y, t) \quad (1-45)$$

The local gradient of the objective space with respect to the model weights can be used to determine the direction of movement. We also want to add a learning rate (η) to control how much each of the weights is updated in the desired direction (Eq. 1-46). This is known as gradient descent.

$$w^{(j+1)} = w^{(j)} + \Delta w \quad \text{where} \quad (1-46)$$

$$\Delta w = -\eta \nabla J(w^{(j)})$$

In order to determine the gradient at a specific weight, we must consider all the operations that occur between the weight and the final output. This can be shown visually quite simply (Fig. XYZ). Calculating the gradient at a specific weight involves a simple chain-rule operation (Eq. 1-47). The chain rule here is based on determining the local gradient at a weight only passing through a single layer. This process is known as backpropagation, and it ubiquitous for training neural networks [60].

GRGAPH NN FOR GRAD DESCENT

$$\frac{\partial J}{\partial w} = \frac{\partial J}{\partial e} \frac{\partial e}{\partial \phi} \frac{\partial \phi}{\partial v} \frac{\partial v}{\partial w} \quad (1-47)$$

One of the main limitations of gradient descent is the need to tune the learning rate. To small a value will cause a network to train extremely slowly and almost always get stuck in local minima. Too large a learning rate can cause the network to “bounce out” of the global minima due to the rate of change being too large. Hyperparameter tuning can be a difficult process without a defined methodology beyond trial and error. Some groups have proposed different methods of incorporating dynamically changing update rules in order to incorporate physical properties into the network training stage. The most common is Adaptive Moment Estimation, which incorporates Root Mean Squared Propagation [29] and momentum learning.

1.3 Model-Image Registration for Measuring Kinematics from Fluoroscopic Images

1.4 Automating Measurements of Kinematics from Fluoroscopic Images

The proposed method overcomes the various limitations of previous attempts to autonomously measure kinematics from single plane fluoroscopy. The key feature of all these limitations is that they prevent the adoption of this technique in a clinical setting, due to the labor overhead or equipment required in order to generate an accurate kinematic report. The proposed combination of methods will alleviate these requirements, and the extensibility of the software will allow this technology to be used in a clinical setting without disrupting the standard clinical workflow.

1.4.1 Autonomous Implant Detection

Determining the location of an object in an image is a historically intractable problem. These are one of those tasks that are often relegated to the corner of “easy for humans, extremely difficult for computers”, especially when there is very little a-priori information available. However, as discussed in Section 1.2, deep learning has paved the way for computer vision programs to be able to perform tasks that were once only possible by humans. Two convolutional neural networks were trained to segment the tibial and femoral components from the single plane fluoroscopic images. The network architecture used was the High-Resolution Net [75], which leverages low-level features with higher resolution parallel processing in order to better determine the spatial characteristics of the image and produce a better output. At the time of writing, this network sets the state-of-the-art standard for performance on the COCO and ImageNet datasets, demonstrating robustness for use in many different arenas.

put some pictures here of the segmentation performance of the
neural network

Many of the historic methods of determining kinematics were limited by the researchers ability to quickly and reliably determine the location in the implant. The contours were either

hand-segmented [5, 83], or a normal edge detector was used, which introduces extra tuning parameters for any given study due to variations in image quality.

1.4.1.1 Neural Network Robustness

One of the main problems with neural networks is overfitting. With millions of parameters to tune, it can be extremely easy to “overfit” on your training set, leading to the network’s inability to perform well on any image that was not directly in the training set. When dealing with fluoroscopic images, this might look like a neural network that can perform extremely well on high-quality, high frame rate, low blur images from a hospital that has a budget to support such a machine, while failing to segment images from a machine more than a decade old. We overcame this challenge through a mixture of additional image augmentations [13] and using a wide range of training data. The neural networks were trained on roughly 8000 images from 7 different human-supervised total knee kinematics studies spanning the last two decades. The image qualities range from extremely clear and high quality to nearly indiscernible without intense human supervision. The goal of this two-pronged approach was to have both artificial and real “low quality” images for the network to train on so that any hospital or researcher, regardless of the available equipment, might be able to leverage this technology in their practice. The authors hope that this approach provides equal access to this informative measurement.

1.4.2 Initial Pose Estimation

Hand-in-hand with implant detection is the initial pose estimate that very often needs to be input into the optimization routine. Once the contour of the implant was determined, many methods required a human operator to place the implant in the “capture region” of their optimizer in order for it to eventually find the correct solution. This takes human supervision to get correct, thus adding another impediment toward getting this technology into the clinic.

To determine an initial estimate, we must rely on those methods that can leverage information present in the camera projection (Section 1.1.2) and the 2D CNN output (Section 1.4.1). The primary method that takes advantage of this information utilizing normalized fourier descriptor shape libraries, and extracting each of the 6 degrees-of-freedom from either the

normalized coefficients or matching with the closest entry in the library [73, 74, 6, 5]. The key feature that makes this method tractable for autonomous measurements is the availability of the implant pixels from the convolutional neural network.

1.4.2.1 Generating Normalized Fourier Descriptors

The Fourier Transform is one that takes in a continuous and repeating function and represents it as a sum of sinusoidal signals. Because the implant contour is self intersection, we can view it as a continuous function that has period 2π radians, as it will loop back onto itself and start over. This allows us to take advantage of the Fast Fourier Transform (FFT) [18], which operates on a discrete set of points rather than a continuous function. First, the contour of the segmented implant is taken then resampled into 128 equi-spaced points. The choice of 2^n points allows the FFT algorithm to perform much more quickly than another choice of points. Each contour point on the image (x, y) is then represented as a single complex point, $x = jy$, such that the 1D FFT algorithm can be used. If $s(n)$ represents the complex sequence of equi-spaced contour points in the implant, and $S(i)$ represents the frequency-domain representation of those points after the FFT is applied, then we can represent these functions as shown in Eq. 1-48.

$$\begin{aligned} S(i) &= \sum_{n=-\frac{N}{2}+1}^{\frac{N}{2}} s(n) e^{-j(2\pi in)/N} \\ &\text{for } -\frac{N}{2} + 1 \leq i \leq \frac{N}{2} \\ s(n) &= \frac{1}{N} \sum_{i=\frac{N}{2}+1}^{\frac{N}{2}} S(i) e^{j(2\pi in)/N} \\ &\text{for } -\frac{N}{2} + 1 \leq n \leq \frac{N}{2} \end{aligned} \tag{1-48}$$

As discussed in Section 1.1.4, spatial information in images is difficult for computers to understand without explicit calculation. So, we use the properties of the FFT to normalize each of the shapes based on location, rotation, and size in order to accurately compare the segmented mask to a generated library.

Normalize Position By the properties of the FFT, we know that $S(0)$ is the geometric centroid of the contour. Thus, we can set this to zero for all contours to give a consistent reference point that is independent of the location of the input contour. We can save this value as the “position normalization coefficient” to determine the (x,y) location of the implant later in the process. And, because of our usage of the 1D FFT, we know that the real portion of this coefficient is the x-value and the imaginary portion is the y-value.

$$\begin{aligned} \text{Position Coefficient} &\leftarrow S(0) \\ S(0) &= 0 \end{aligned} \tag{1-49}$$

Normalize Size Because the implant is not self-intersecting, we know that $S(1)$ is the coefficient with the largest size, and it also represent the scale of the shape of the implant. So, we can normalize each shape by dividing each coefficient by the overall size of the contour, shown below.

$$\begin{aligned} \text{Size Coefficient} &\leftarrow |S(1)| \\ S(i) &= \frac{S(i)}{|S(1)|} \\ \text{for } -\frac{N}{2} + 1 < i < \frac{N}{2} \end{aligned} \tag{1-50}$$

Normalize In-Plane Rotation and Contour Starting Point One of the most difficult parts of measuring the similarity between two contours, especially when they are composed of a set of discrete points, is that any distance measurement necessarily takes into account those discrete points in the order that they are presented. To illustrate this example, imagine two squares, each defined by the location of the corners A_i, B_i, C_i, D_i . If these two squares are perfectly overlapping, one might imagine that the Euclidean distance between each of the points, $(\sum_{P \in A,B,C,D} (P_1 - P_2)^2)^{\frac{1}{2}}$ would be equal to zero. This would only be true if the starting point of the contour was the same for each square (e.g. Corner A was always the top left corner). If each square had A starting in different locations, then even when the contours are perfectly aligned, the distance metric would be non-zero and uninformative.

We can use properties of the FFT to normalize the starting position of the contour in each of the segmentations as well as the general orientation of the contour. We can leverage the starting point shift property of the fourier transform (Eq. 1-51) and the rotation property (Eq. 1-52) in order to normalize both of these factors and ensure that similar shapes have the same orientation and starting point.

$$s(n - T) \xleftrightarrow{DFT} S(i)e^{-jiT} \quad (1-51)$$

$$s(n)e^{j\theta} \xleftrightarrow{DFT} S(i)e^{j\theta} \quad (1-52)$$

To normalize the starting point and rotation, we find k such that $S(k)$ is the coefficient of second largest magnitude. We then apply a mixture of the starting point shift and the rotation property of the FFT to orient each contour (Eq. 1-53). We also want to find the “rotation normalization coefficient”, which the the angle through which the contour needs to rotate to reach the normalized orientation. This is done by determining the phase of the normalization equation at $i = 0$, which controls the overall orientation of the contour. If u is the phase of $S(1)$, and v is the phase of $S(k)$, then we can find the normalized orientation.

$$\begin{aligned} \text{Rotation Normalization Coefficient} &\leftarrow \frac{v - ku}{k - 1} \\ S(i)_{norm} &= S(i)e^{j\frac{(i-k)u + (1-i)v}{k-1}} \\ \text{for } -\frac{N}{2} + 1 &\leq i \leq \frac{N}{2} \end{aligned} \quad (1-53)$$

Once the in-plane rotations have been normalized, the contour has been completely normalized for x, y, z translations and z rotations. And, by storing these values, we are able to utilize them in determining an initial pose estimate. Then, we can use a library made up of known x, y rotations, and compare the segmentation to this library to determine all 6 degrees-of-freedom.

1.4.2.2 Shape Library

A shape library is created using a flat panel projection (Section 1.1.2) of the implant at known x and y rotations, while holding all positions and orientations constant, then applying the normalization protocol described above to determine the normalized coefficients of each library entry. If $s_{x,y}(n)$ is the flat-panel projection of the implant with x and y rotations, then we can generate a library using the following procedure, where FFT is the fast fourier transform (Eq. 1-48) and NFD is the process of normalizing the contour and extracting the relevant coefficients (Section 1.4.2.1).

$$S_{x,y}^{lib}(i) = NFD(FFT(s_{x,y}(n))) \quad (1-54)$$

Once the shape library is generated for a specific implant, we can start to determine the pose estimates for each degree of freedom.

1.4.2.3 Generating a Pose Estimate

First, we compare the Euclidean distance of the normalized segmentation contour to each value of the shape library; the x and y rotations that minimize this function are taken as the x and y rotations of the implant (Eq. 1-55).

$$(\theta_{x,est}, \theta_{y,est}) = \operatorname{argmin}_{x,y} \left(\sum_{i=-\frac{N}{2}+1}^{\frac{N}{2}} (S^{seg}(i) - S_{x,y}^{lib}(i))^2 \right)^{\frac{1}{2}} \quad (1-55)$$

Then, we can determine the z rotation estimate by comparing the values of the normalized θ values that were needed in Eq. 1-53. This process is shown in Eq. 1-56.

$$\theta_{z,est} = \theta^{seg} - \theta_{x,y}^{lib} \quad (1-56)$$

where $\theta^{seg,lib} \equiv$ Rotation Normalized Coefficient

We can then use the principals of projective geometry (Fig. 1-1) and similar triangles to determine the out-of-plane translation of the implant, given that the library was projected at a known depth. Using similar triangles, we are able to determine that the depth is inversely

proportional to the overall magnitude of the projection, captured by the “Size Coefficient” (Eq. 1-57). We also make the assumption that we have a weak perspective projection, meaning that the out-of-plane translations are small compared to the focal length of the fluoroscopy imaging setup.

$$\begin{aligned} \frac{M^{seg}}{f} &= \frac{h}{z_{est}} \\ \frac{M^{lib}}{f} &= \frac{h}{z_{lib}} \\ \rightarrow \\ z_{est} &= \frac{M^{lib}}{M^{seg}} z_{lib} \end{aligned} \tag{1-57}$$

where

$M^{seg,lib} \equiv$ Size Coefficients

$h \equiv$ Implant Size

The x and y translations can be determined using the value of the z translation estimate, along with the location of the centroid, saved as the “Position Coefficient”. This is then refined further to account for the rotation of the implant and the distance of the implant’s centroid to its origin. We can express this with a single matrix multiplication multiplied by a scale factor (Eq. 1-58).

$$\begin{pmatrix} x_{est} \\ y_{est} \end{pmatrix} = \begin{pmatrix} Re(S^{seg}(0)) \\ Imag(S^{seg}(0)) \end{pmatrix} - \begin{pmatrix} \cos(\theta_z) & -\sin(\theta_z) \\ \sin(\theta_z) & \cos(\theta_z) \end{pmatrix} \begin{pmatrix} Re(S_{x,y}^{lib}(0)) \\ Imag(S_{x,y}^{lib}(0)) \end{pmatrix} \times \left(\frac{z_{est}}{z_{lib}} \right) \tag{1-58}$$

Once this step is complete, we have accounted for x and y rotations (Eq. 1-55), z rotations (Eq. 1-56), x and y translations (Eq. 1-58), and z translations (Eq. 1-57). This provides a robust initial estimate when the only information available is the implant geometry and the segmentation output from the neural network. Furthermore, this can be done without any human supervision, providing a fully autonomous initialization for any pose refinement strategy, so long as the

estimate is within the convergence region.

1.4.3 Objective Function

In a perfect situation, our objective function would directly measure the error between our 3D model's current pose and the true pose of the object. However, if we had a-priori access to the true pose of the object, then this entire pipeline would be worthless. Thus, we must find an objective function that can act as a heuristic for the difference between true pose of our model and the current pose of our model. Our access to the segmentation output from the CNN and the ability to project the silhouette of our model quickly makes contour comparison a natural choice for an objective function. The only assumptions that we make are (1) our projective algorithm and camera definition are the same as the camera that was used to take the original fluoroscopic image and (2) our 3D model is the same 3D model that is present in the image. If these two assumptions are correct, then the alignment of the image contour and the projected contour necessarily means that our pose is correct (SYM TRAP EXCEPTION).

First, we apply a Canny edge detector ([14]) to extract the edges from our segmentation contour, S , and our projected 3D model, P , where edges are 1, and every other pixel is 0. We can then iterate over each pixel and take the absolute values of the L_1 norm between our segmented and projected contours(Eq. 1-59).

$$J = \sum_i^{Height} \sum_j^{Width} |S_{ij} - P_{ij}| \quad (1-59)$$

Unfortunately, the contours of the projected model are extremely sensitive to pose, especially when representing angles using Euler decomposition. This results in a chaotic similarity function that has an extensive amount of local minima. Past methods have overcome this by dilating the contour of the projected image (Eq. 1-28) and performing the same L_1 optimization routine. However, a lack of an isolated contour for the fluoroscopic image still lead to a slightly noisy objective function. Our proposed method takes advantage of the segmentation output for the neural network and dilates both the segmentation contour and the projected contour for a much smoother objective function allowing for a wider search range. As our objective

function is minimized, we can decrease the level of dilation to return the metric back into its original form, which most accurately describes the difference between the projection and image.

1.4.4 Optimization Routine

Broadly, optimization is the process of minimizing or maximizing an objective function, $f(\mathbb{R}^n) \rightarrow \mathbb{R}$, potentially subject to some constraints (e.g. $x \in \Omega$) [3]. We formalize this below (Eq. 1-60). The simplest optimization problems have an analytic form of the gradient of f that can be solved directly, typically by setting the first derivative to zero (e.g. least squares).

$$\operatorname{argmin}_x \{f(x) : x \in \Omega\} \quad (1-60)$$

A drawback to our pipeline is that there is no analytic form of the objective function between each segmentation and projected contour; they must be resampled over a specified range in order to approximate objective function values and gradients. This defines a *black box* optimization routine, which is well studied in the literature [3]. In this type of function, it is not possible to use gradient-based methods to determine a minimum value for the objective function, one must use heuristics or ad-hoc methods to find the minimum location. Lipschitzian optimization offers an appealing black box optimization approach because it satisfies our need for a global search algorithm with provable convergence. First, we start with the definition of Lipschitz continuous, which places bounds on the rate of change of a function specified by some constant, called the Lipschitz constant. With a known Lipschitz constant, it is possible to find the value for the global minimum of optimization function [22].

Definition 1-1 (Lipschitz Continuous). *The function g is said to be Lipschitz Continuous on the set $\mathbf{X} \in \mathbb{R}^n$ if and only if there exists a scalar $K > 0$ for which*

$$\|g(x) - g(y)\| \leq K\|x - y\| \quad \text{for all } x, y \in \mathbf{X}$$

The scalar K is called the **Lipschitz Constant** of g relative to the set \mathbf{X} .

We can illustrate this convergence with a simple example: consider the function $f(x) \rightarrow \mathbb{R}, x \in [a, b]$. If we know that the function is Lipschitz continuous, the following conditions are true on the domain $x \in [a, b]$. This corresponds to the positive and negative slopes, K , applied to the extrema of the domain, and the intersection, x is selected as the choice for subdivision. This process is repeated, where the region is further subdivided based on the lowest value of $f(x_i)$. The iterative process is stopped once the difference between successive domain splitting is below a pre-specified global tolerance.

$$\begin{aligned} f(x) &\geq f(a) - K(x - a) \\ f(x) &\geq f(b) + K(x - b) \end{aligned} \tag{1-61}$$

While powerful, a-priori knowledge of the Lipschitz constant is needed for determining the global minima. Without it, there is no way of determining intersection points, and no way of selected new regions for subdivision and sampling. shubert's algorithm also has slow convergence, due to the inability to define parameters for when to explore local vs. global search. The Lipschitz constant, K , acts as a weight that places larger emphasis on global serach when high, and local search when low.

Fortunately, methods exist for utilizing the power of Lipschitzian optimization without the need for explicit knowledge of the Lipschitz constant [32]. These can both overcome the need for an a-priori knowledge of the Lipschitz constant, as well as offer some solutions of the slow convergence by providing methods for exploiting both local and global search simultaneously to find the minimum function values.

Jones et al. [32] propose a method by which the center, c , of a domain is sampled, rather than the endpoints. This produces the equations below (Eq. 1-62). The inequalities represent slopes $+K$ and $-K$, respectively, and provide a maximum value for the lower bound of the function at the endpoints, a and b . The midpoints of $[a, c]$, and $[c, b]$ are then calculated and the process is then repeated (Fig. 1-8). The power of this method is that you can determine

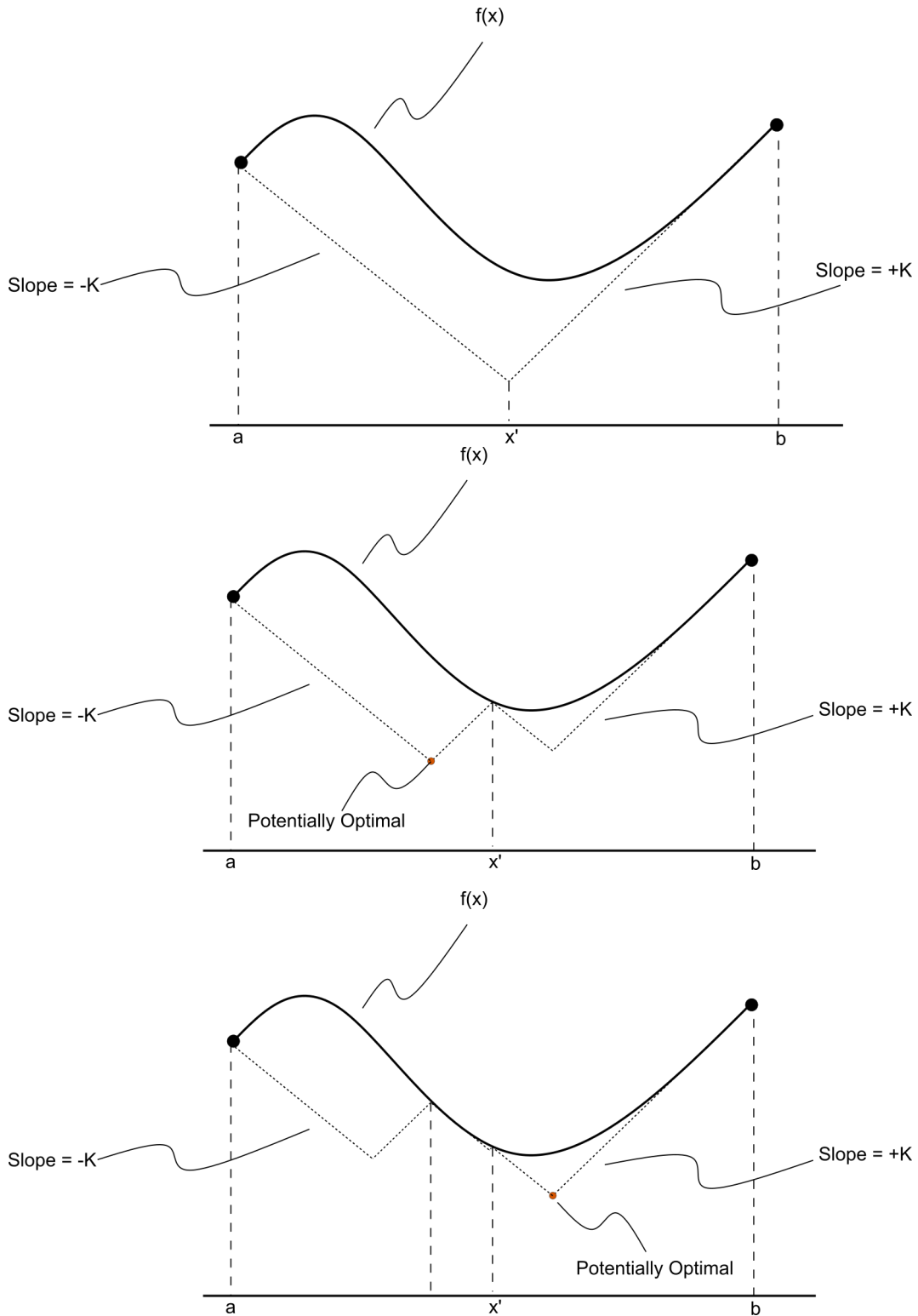


Figure 1-7. A visual representation of Shubert's algorithm, which finds the global minima iteratively through a repeated calculation of the intersection of two lines with slope $\pm K$. If K is known, it will always find the global minimum.

potentially optimal regions of the domain by choosing those points along the lower convex hull of the graph plotting sub-domain size vs center point function value. The points along this hull are those that could potentially include the function minimum, and each is chosen for further sub-sampling (Fig. 1-9). Determining the convex hull is a problem well studied in the literature (CREATE CITATION AND FIND SOME SOURCES). This elegantly mixes local vs. global search, and drastically increases the speed of convergence.

$$f(x) \geq \begin{cases} f(c) + K(x - c) & \text{if } x \leq c \\ f(c) - K(x - c) & \text{if } x \geq c \end{cases} \quad (1-62)$$

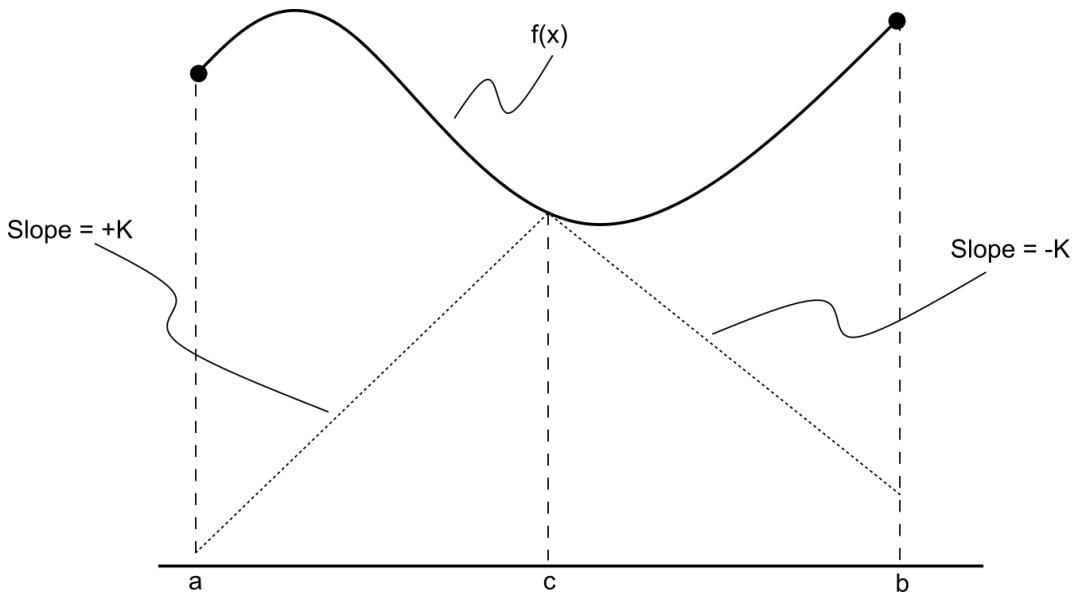


Figure 1-8. The DIviding RECTangles (DIRECT) algorithm in one dimension. It can find the global minimum of a function without a-priori knowledge of the Lipschitz constant. The value of the line with slope $\pm K$ at each of the end-points represents the theoretical minimum for the value of the function in that region. Thus, the size of the region and the value of the function at the center help the algorithm determine potentially optimal sub-regions.

This can be extended into multiple dimensions without loss of generality. First, each dimension in the domain is normalized to the range $[0, 1]$, and a hypercube is created in \mathbb{R}^D , where D is the dimension of the domain you are searching. The first iteration trisects this hypercube along an arbitrary dimension, and further iterations trisect along the largest dimension

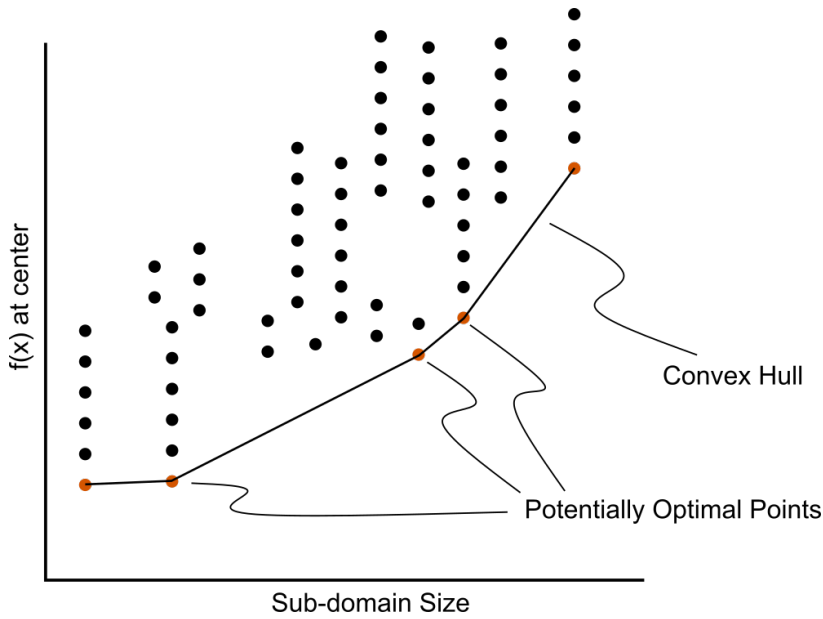


Figure 1-9. The potentially optimal regions of the DIRECT algorithm are those points that lay along the lower convex hull of the scatter plot of sub-domain size vs function value at center. This is due to those regions being the locations where the maximum possible rate of change in the function might be a minimum, without any prior knowledge of the Lipschitz constant.

of the hypercube. We select potentially optimal hypercubes by identifying points along the lower convex hull of the graph plotting hypercube size vs center point function value.

For our purposes, we construct a hypercube along each of the 6 degrees-of-freedom that describe the pose of the implant in space, using bounds set by the user. The first epoch involves minimizing the objective function with larger bounds and a larger dilation. After all iterations have been used up, the algorithm is re-started with a smaller dilation and tighter bounds. The last epoch typically has the tightest bounds and no dilation. This pyramidal scheme offers a smooth objective function when the bounds are largest, which assists in escaping local minima, and an aggressive objective when the bounds are tight, and fewer local minima are present.

1.4.5 Overcoming Single-Plane Limitations

Despite a fully autonomous solution for measuring total knee arthroplasty kinematics, there are some fundamental limitations when using monocular vision to determine the three dimensional position and orientation of an object.

CHAPTER 2
THE AIMS OF THIS DISSERTATION

2.1 Aim 1: XXX

2.2 Aim 2: YYY

2.3 Aim 3: ZZZ

CHAPTER 3

JOINT TRACK MACHINE LEARNING: AN AUTONOMOUS METHOD OF MEASURING 6-DOF TKA KINEMATICS FROM SINGLE-PLANE FLUOROSCOPIC IMAGES

3.1 Introduction

Total Knee Arthroplasty (TKA) is a standard procedure for alleviating symptoms related to osteoarthritis in the knee. In 2018, orthopaedic surgeons performed more than 715,000 TKA operations in the United States [2]. This number is projected to increase to 3.48 million by 2030 [40] due to an aging population and increased obesity rates. While TKA largely relieves symptomatic osteoarthritis, roughly 20% of TKA patients express postoperative dissatisfaction, citing mechanical limitations, pain, and instability as the leading causes [4, 10, 63]. Standard methods of musculoskeletal diagnosis cannot quantify the dynamic state of the joint, either pre- or post-operatively; clinicians must rely on static imaging (radiography, MRI, CT) or qualitative mechanical tests to determine the condition of the affected joint, and these tests cannot easily be performed during weight-bearing or dynamic movement when most pain symptoms occur. Unfortunately, most of the tools used to quantify 3D dynamic motion are substantially affected by soft-tissue artifacts [25, 66, 42], are prohibitively time-consuming or expensive [20], or cannot be performed with equipment available at most hospitals.

Model-image registration is a process where a 3D model is aligned to match an object's projection in an image [11]. Researchers have performed model-image registration using single-plane fluoroscopic or flat-panel imaging since the 1990s. Early methods used pre-computed distance maps [41, 83], or shape libraries [5, 73, 74] to match the projection of a 3D implant model to its projection in a radiographic image. With increasing computational capabilities, methods that iteratively compared implant projections to images were possible [46, 23, 44]. Most model-image registration methods provide sufficient accuracy for clinical joint assessment applications, including natural and replaced knees [8, 7, 38, 12], natural and replaced shoulders [37, 45, 49, 67], and extremities [16, 15, 21]. One of the main benefits of this single-plane approach is that suitable images can be acquired with equipment found in most hospitals. The main impediment to implementing this approach into a standard clinical workflow

is the time and expense of human operators to supervise the model-image registration process. These methods require either (1) an initial pose estimate [23, 44], (2) a pre-segmented contour of the implant in the image [11, 41], or (3) a human operator to assist the optimization routine out of local minima [46]. Each of these requirements makes model-image registration methods impractical for clinical use. Even state-of-the-art model-image registration techniques [23] require human initialization or segmentation to perform adequately.

Machine learning algorithms automate the process of analytical model building, utilizing specific algorithms to fit a series of inputs to their respective outputs. Neural networks are a subset of machine learning algorithms that utilize artificial neurons inspired by the human brain's connections [47]. These networks have shown a great deal of success in many computer vision tasks, such as segmentation [17, 75, 59], pose estimation [78, 35], and classification [39, 56, 57]. These capabilities might remove the need for human supervision from TKA model-image registration. Therefore, we propose a three-stage data analysis pipeline (Fig. 3-1) where a convolutional neural network (CNN) is used to segment, or identify, the pixels belonging to either a femoral or tibial component. Then, an initial pose estimate is generated comparing the segmented implant contour to a pre-computed shape library. Lastly, the initial pose estimate serves as the starting point for a Lipschitzian optimizer that aligns the contours of a 3D implant model to the contour of the CNN-segmented image.

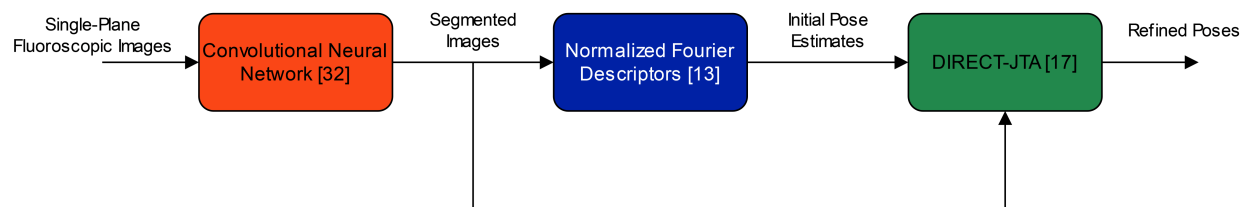


Figure 3-1. An overview of the pipeline for autonomous measurements of total knee arthroplasty kinematics. First, the data is processed through a convolutional neural network to locate the pixels belonging to the femoral and tibial implants [75], then, Normalized Fourier Descriptor shape libraries are used to determine and initial pose estimate [5], and lastly, DIRECT-JTA [23] is run on those segmented images using the NFD estimates as initializations for pose.

This paper seeks to answer the following three questions: (1) How well does a convolutional

neural network segment the femoral and tibial implants from fluoroscopic and flat-panel images?

(2) Can a Fourier descriptor-based pose estimation method produce useful initial guesses of 3D implant pose from the CNN-segmented images? (3) Can the Lipschitzian optimizer, given reasonable initial guesses, replicate human-supervised TKA kinematic measurements?

3.2 Methods

Data from seven previously reported TKA kinematics studies were used for this study [34, 54, 53, 76, 31, 77, 64]. These studies utilized single-plane fluoroscopy or flat-panel imaging to measure tibiofemoral implant kinematics during lunge, squat, kneel, and stair climbing movements from 8248 images in 71 patients with implants from 7 manufacturers, including 36 distinct implants. From each of these studies, the following information was collected: (1) deidentified radiographic images, (2) x-ray calibration files, (3) manufacturer-supplied tibial and femoral implant surface geometry files (STL format), and (4) human supervised kinematics for the tibial and femoral components in each of the images. CNNs were trained with images from six of the studies using a transfer-learning paradigm with an open-source network [75]. CNN performance was tested using two image collections: a standard test set including images from the six studies used for training and a wholly naïve test set using images from the seventh study, where the imaging equipment and implants were different from anything used in training (Fig. 3-2). We used both test image sets to compare human-supervised kinematics with autonomously measured kinematics. Separately, two independent groups utilized our software to assess the accuracy of TKA kinematics measurements compared to their previously reported reference standard systems using RSA [69] or motion capture [20].

3.2.1 Image Segmentation

Images were resized and padded to 1024x1024 pixels. Images containing bilateral implants had the contralateral knee cropped from the image. Segmentation labels were created by taking the human-supervised kinematics for each implant and generating a flat-shaded ground-truth projection image (Fig. 3-3). Two neural networks [75] were trained to segment the tibial and femoral implants, respectively, from the x-ray images. Each network was trained using a random

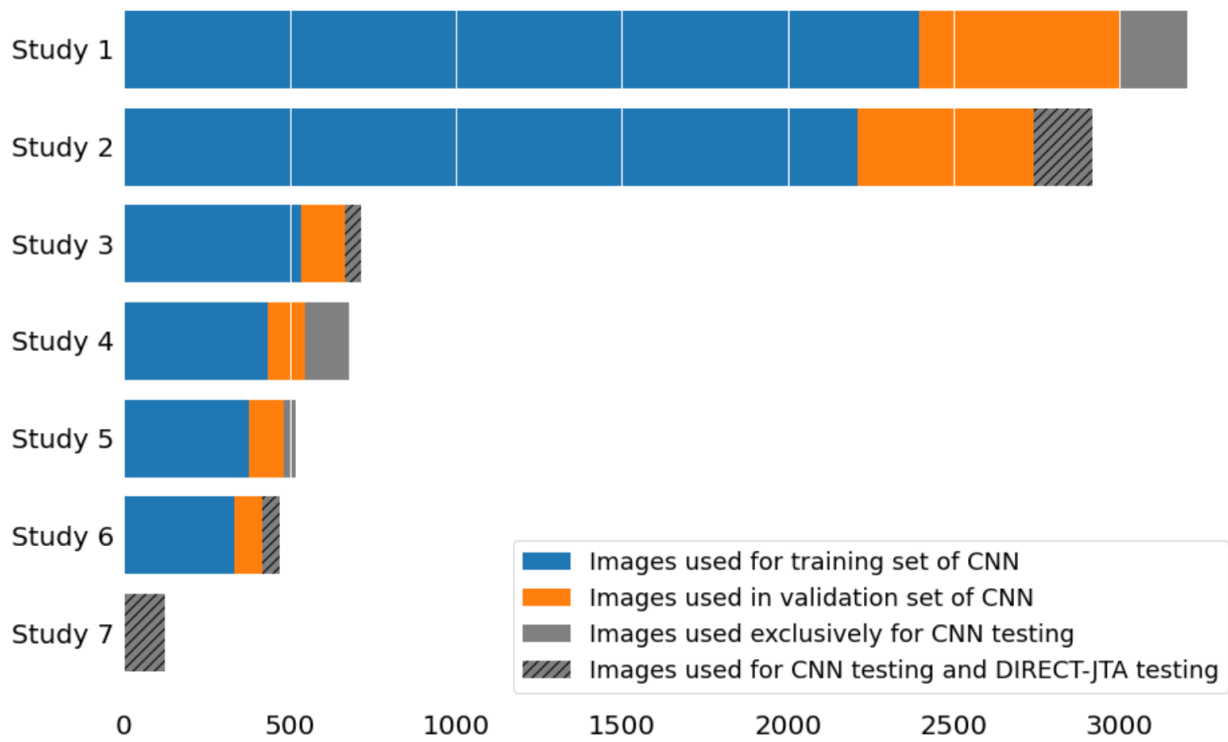


Figure 3-2. Data from seven studies were used to train and test the TKA kinematics measurement pipeline. Color coding in the figure identifies how many images were used for the training, validation, and testing functions. Images from the seventh study were used exclusively for testing the measurement pipeline that was trained using images from the other six studies.

6284/1572 (80/20) training/validation split. Augmentations were introduced in the training pipeline to improve the network's generalization to new implants and implant types [13]. Each neural network was trained on an NVIDIA A100 GPU for 30 epochs. The performance of the segmentation networks was measured using the Jaccard Index [30]. This calculates the intersection between the estimated and ground-truth pixels over the union of both sets of pixels. The ideal Jaccard index is 1.

3.2.2 Initial Pose Estimates

Initial pose estimates were generated from bounding contours of the CNN-segmented implant regions using Normalized Fourier Descriptor (NFD) shape libraries [5, 73, 74]. Shape libraries were created by projecting 3D implant models using the corresponding x-ray calibration parameters with $\pm 30^\circ$ ranges for the out-of-plane rotations at 3° increments (Fig. 3-4). Pose

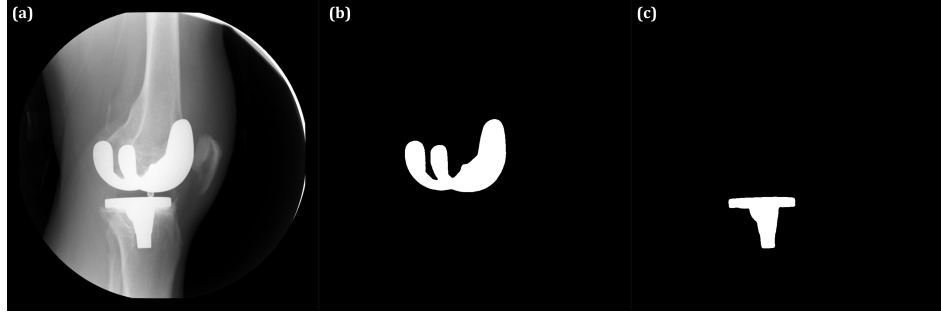


Figure 3-3. A representative fluoroscopic image is shown (a) with corresponding femoral (b) and tibial (c) ground-truth images created by flat-shaded projections of registered implant models.

estimates were determined as previously described [5] NFD-derived femoral and tibial implant poses were transformed to anatomic joint angles and translations [26] and compared to the human-supervised kinematics for the same images using RMS differences for each joint pose parameter. The performance of this method was also assessed using flat-shaded projection images with perfect segmentation as a ground-truth reference standard.

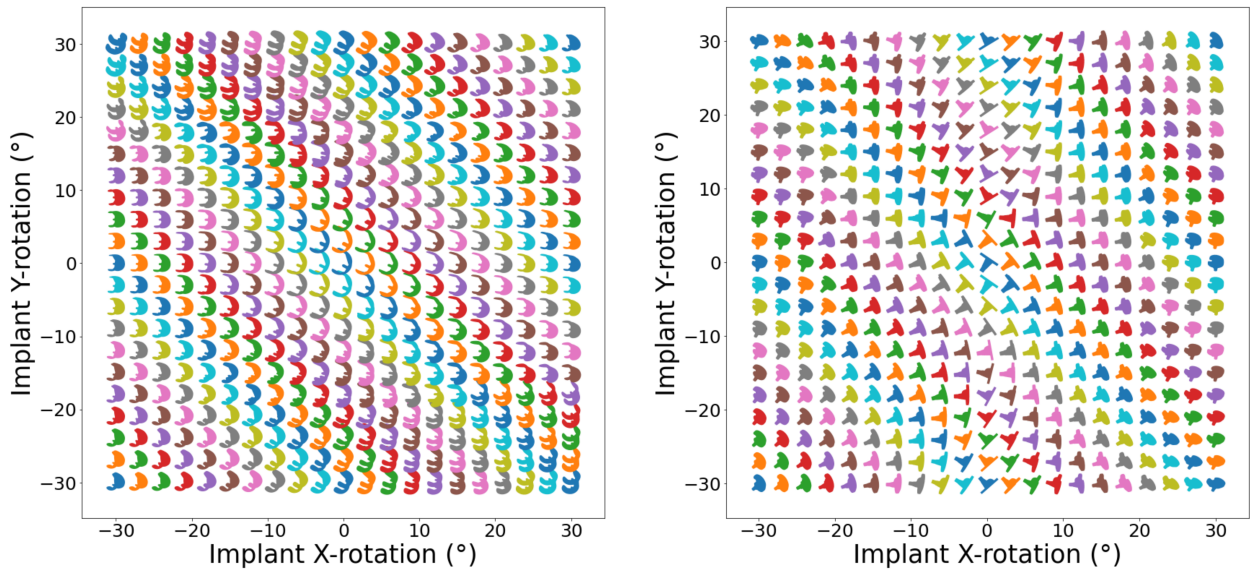


Figure 3-4. Femoral (left) and tibial (right) NFD shape libraries were generated to capture the variation in projection silhouette geometry with out-of-plane rotation [5]. Initial pose estimates were generated by comparing the NFD contour from the x-ray image to the shape library.

3.2.3 Pose Refinement

A modified Dividing Rectangles (DIRECT) algorithm called DIRECT-JTA [23] generated the final pose estimates. This method of Lipschitzian optimization divides the search into three stages, the “trunk,” “branch,” and “leaf.” Each of the three stages was assigned distinct cost function parameters and search regions. The cost function used a computationally efficient L1-norm between the dilated contour from the segmentation label and the projected implant. Successively decreasing the dilation coefficient allowed the optimization routine to escape local minima, and the leaf branch served to find the optimal out-of-plane translation. Transversely symmetric tibial implants posed problems during registration because two distinct poses produced roughly identical projections [36]. Because of this pose ambiguity, the tibial implant was always optimized after the non-symmetric femoral implant. In addition to the dilation metric, the tibial mediolateral translation and varus/valgus rotations relative to the femur were penalized. Final implant poses were transformed into knee joint rotations and translations [26] and compared to the human-supervised kinematics for the same images using RMS differences for each joint pose parameter. Squared differences between data sets were compared using one-way MANOVA with post-hoc multiple pair-wise comparisons using the Games-Howell test (R v4.2.0 using R Studio, rstatix, and stats).

3.2.4 Pose Ambiguities and Registration Blunders

A blunder was defined as an image frame with the squared sum of rotation differences greater than 5° between autonomous and human-supervised measures. These blunder frames contain errors considerably larger than would be clinically acceptable and warrant further exploration. Blunders were analyzed with respect to the tibial implant’s apparent varus/valgus rotation relative to the viewing ray (Fig. 3-5). A probability density function and cumulative density function were calculated for the blunder likelihood. Due to the high likelihood of blunders in this region, an ambiguous zone was defined for all apparent tibial varus/valgus-rotation less than 3.6 degrees, which is the mean + 1std of the blunder distribution (Fig. 3-5). Squared measurement differences between images inside and outside the ambiguous

zone were also compared using one-way MANOVA with post-hoc multiple pair-wise comparisons using the Games-Howell test.

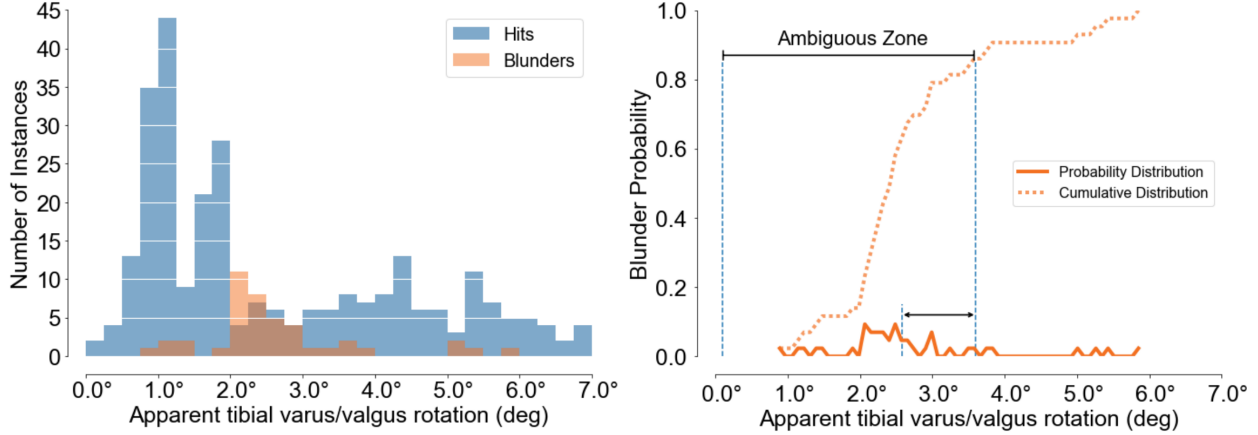


Figure 3-5. The histogram (left) shows the correctly registered frames (Hits, blue) and incorrectly registered frames (Blunders, orange) plotted as a function of the apparent tibial varus/valgus angle relative to the viewing raw. The probability plot (right) shows the distribution of blunders (solid orange) and the cumulative probability of blunders (dotted orange). The Ambiguous Zone is defined as apparent tibial varus/valgus rotations less than the mean + one standard deviation of the blunder probability distribution, capturing approximately 85 % of the blunders.

3.3 Results

CNN segmentation of standard test set images produced Jaccard indices of 0.936 for the femoral and 0.883 for the tibial components. CNN segmentation performance on the completely naïve test set was lower, 0.715 and 0.753, respectively.

The initial pose estimates were within the range of convergence for the DIRECT-JTA optimizer and offered a robust initialization for optimization (Table 1). The RMS differences for initial pose estimates on ground-truth images were smaller (better) than for CNN-segmented images, but the differences were mostly within a few millimeters or degrees. Due to poor sensitivity for measuring out-of-plane translation with monocular vision, the mediolateral translation had the largest RMS differences for both image types.

RMS differences between DIRECT-JTA optimized kinematics and human-supervised kinematics were sub-millimeters for all in-plane translations (Table II). Mediolateral translations and out-of-plane rotation differences were smaller when the pose of the tibia was outside the

Table I
RMS Differences Between NFD Initial Estimates and Human-Supervised Kinematics

Implant	Images	Translation (mm)			Rotation (deg)		
		x	y	z	z	x	y
Femoral	CNN-Segmented Images	2.37	0.71	17.59	2.54	2.45	4.75
	Ground-Truth Projections	2.06	0.57	13.53	0.85	1.42	4.00
Tibial	CNN-Segmented Images	2.06	1.49	29.93	0.94	5.59	9.47
	Ground-Truth Projections	2.05	0.87	14.60	0.55	4.73	6.23

ambiguous zone. The RMS differences for the completely naïve test set were within 0.5 mm or 0.5 deg compared to the standard test set, indicating similar performance on the entirely novel dataset.

Table II
RMS Differences Between DIRECT-JTA Optimized and Human-Supervised Kinematics

Test Set	Image Group	Number of Images	A/P (mm)	S/I (mm)	M/L (mm)	Flx/Ext (°)	I/E (°)	V/V (°)
Standard	Inside AZ	187	0.694	0.523 ^b	1.752 ^a	0.730 ^a	3.380	1.938 ^a
	Outside AZ	83	0.685	0.466 ^c	0.917	1.029	1.811	0.605
Naïve	Inside AZ	47	0.802	0.739	1.715 ^d	1.388	4.044	2.480 ^d
	Outside AZ	75	0.692	0.644	0.691	1.031	1.154	0.846

AZ = Ambiguous Zone

Superscripts denote pairwise differences ($p < 0.05$) in squared errors for:

- a. Standard Inside AZ vs Standard Outside AZ
- b. Standard Inside AZ vs Naïve Inside AZ
- c. Standard Outside AZ vs Naïve Outside AZ
- d. Naïve Inside AZ vs Naïve Outside AZ

There was one femoral blunder and 43 tibial blunders out of 392 test images. Using the definition of the ambiguous zone as apparent tibial varus/valgus rotation less than 3.6 deg, 11% of images have a tibial blunder within this zone, compared to 3.2% outside. Sixty-six percent of tibial blunders were due to symmetry ambiguities (Fig 3-6).

One-hundred thirteen image pairs from an RSA study of TKA were used to independently assess the accuracy of the autonomous kinematics measurement for single-plane lateral TKA images. RMS errors were 0.8mm for AP translation, 0.5mm for SI translation, 2.6mm for ML translation, 1.0° for flexion-extension, 1.2° for abduction-adduction, and 1.7° for internal-external rotation. At a different institution, 45 single-plane radiographic images were acquired with an instrumented sawbones phantom that was independently tracked using motion capture.

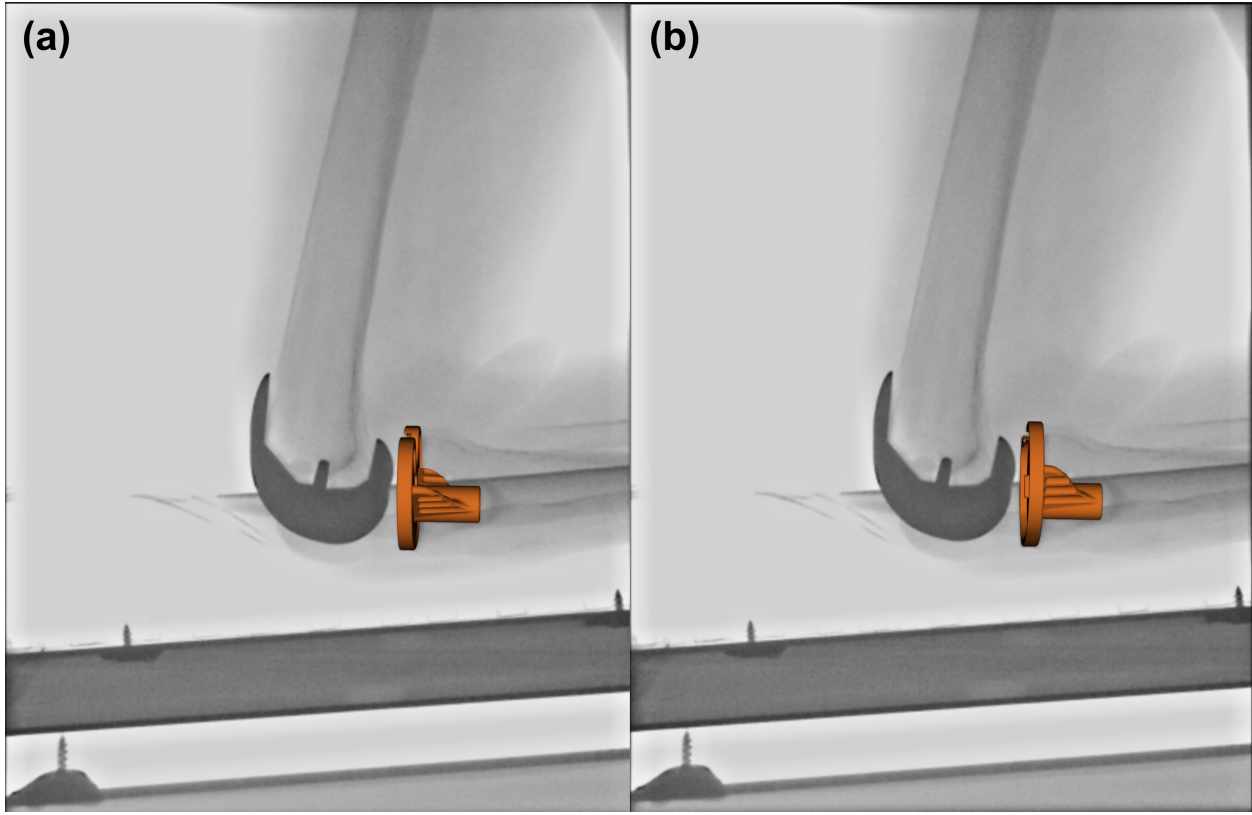


Figure 3-6. The figure shows the same radiographic image with two registered tibial implant poses: (a) shows a correctly registered tibial implant, while (b) shows an implant caught in a local cost function minimum corresponding to a nearly symmetric pose.

Comparing the motion capture and autonomously measured radiographic kinematics, the RMS errors were 0.72mm for AP translation, 0.31mm for SI translation, 1.82mm for ML translation, 0.56° for flexion-extension, 0.63° for abduction-adduction, and 0.84° for internal-external rotation.

3.4 Discussion

Dynamic radiographic measurement of 3D TKA kinematics has provided important information for implant design and surgical technique for over 30 years. Many surgeons have expressed an interest in utilizing this type of measurement in their clinical practices; however, current methods are impractical. We developed a completely autonomous TKA kinematics measurement pipeline that can potentially provide a practical method for clinical implementation. This study sought to answer three questions, (1) How well does a neural network segment TKA implants from fluoroscopic and flat-panel images? (2) How well can an NFD shape library

estimate the pose of a TKA implant given a CNN-segmented image? And (3) How well does a Lipschitzian optimization routine replicate human-supervised kinematics for TKA implants given an approximate initial guess?

CNN image segmentation of TKA implants worked well, with Jaccard indices greater than 0.88 for the standard test set, and greater than 0.71 for the naïve test set. Segmentation performance for the standard test set outperformed published examples by 0.05-0.1 Jaccard points [81, 58], with the naïve test set on par with other segmentation examples. The most notable decrease in segmentation performance occurred along the perimeter of the segmented pixel region, especially in areas where implant projections occluded each other. These imperfectly segmented perimeter regions likely affect the initial pose estimate and the DIRECT-JTA optimization solution since both methods rely heavily on the segmented implant boundary. Further improvements can be made for the perimeter segmentation results by introducing intelligent augmentations during training using generative models [28] and performing neural network bolstered contour improvement strategies [80].

Our initial pose estimates were satisfactory as an initialization for the DIRECT-JTA optimization, falling within the convergence region of $\pm 30^\circ$ [23]. However, the performance for the ground-truth projections was not as good as the cited method [5], which achieved errors of less than 1mm for in-plane translation and 2° for rotation. The cited method utilized an additional refinement step for the NFD estimation, interpolating the apparent out-of-plane angles between nearest shapes in the library. This extra step was not done because only approximate initial pose estimates were needed. In addition, the current study incorporated a vastly larger set of implant shapes (36 vs. 2) and image quality and calibration variations. Distinct implant shapes manifest unique normalization maps, where there can be discontinuities or jumps in normalization angles which affect the best-fitting library entry (Fig. 3-4) [73, 74]. These details are easily upgraded with additional code using previously reported methods but were not pursued because the initial pose results were well within the DIRECT-JTA convergence region. The initial pose estimates for the CNN-segmented images were not as good as for the ground-truth projections. This follows

directly from the fact that the perimeter of the segmented implants was not as accurately rendered, leading to poorer results with the edge-based NFD method. Finally, the out-of-plane translation estimates were relatively poor for both ground-truth projects and CNN-segmented images. This translation estimate is extremely sensitive to model projection and edge detection details and can be adjusted for better results if required.

RMS differences between human-supervised and DIRECT-JTA optimized kinematics demonstrate the two methods provide similar results. In-plane translation differences of less than 0.8mm and out-of-plane less than 1.8 mm, indicate good consistency in determining the relative locations of TKA implants. Rotation differences of 4° or less for frames within the ambiguous zone, and less than 1.7° for frames outside the ambiguous zone, indicate joint rotation measures with sufficient resolution to be clinically useful. We observed two important characteristics in the measurement comparisons that will affect future implementations and use. First, we identified an ambiguous zone of apparent tibial rotations wherein there is a higher incidence of registration errors. These errors resulted in significant differences in measurement performance for the out-of-plane translations and rotations. This phenomenon, resulting from the nearly symmetric nature of most tibial implants [41, 83, 5, 46, 23] prompts either practical modification to imaging protocols to bias the tibial view outside the ambiguous zone or modifications of the model-image registration code to enforce smooth kinematic continuity across image frames and/or to impose joint penetration/separation penalties [51]. Second, we observed similar measurement performance for the standard and naïve test sets, which differed only in the superior/inferior joint translation. This suggests that the autonomous kinematic processing pipeline can provide reliable measures for implants and imaging systems that were not part of the training set, which will be important for application in novel clinical environments.

Two independent research teams utilized our software to evaluate the accuracy of our autonomous measurement pipeline compared to their reference standard methods using implants and image detectors that were not part of our training sets. In both cases, the accuracy results were comparable to results reported for contemporary human-supervised single-plane

model-image registration methods for TKA kinematics [5, 23, 8, 7, 38]. Interestingly, the independent accuracy results appeared superior to our assessment of differences between autonomous and human-supervised measures of TKA kinematics. In both cases, the independent centers used high-resolution flat-panel detectors that provided better spatial resolution and grayscale contrast than most of the imaging systems included in our datasets. With images of similar quality, it is reasonable to expect similar measurement accuracy.

This work has several limitations. First, the image data sets resulted from previous studies in our labs, so there was no prospective design of which implant systems and image detectors should be included for a pipeline that generalizes well to other implants and detectors. Nevertheless, the naïve data set and the independent assessments, all involving implants and detectors not used for training, performed well and suggest that the method can usefully generalize to measurements of traditionally configured TKA implants. Future work is required to evaluate measurement performance with partial knee arthroplasty or revision implants. Second, many methodologic and configuration options and alternatives remain to be explored, and the current pipeline implementation should not be considered optimal. How best to disambiguate tibial poses and determine the most effective and robust optimization cost functions are areas of current effort.

We present an autonomous pipeline for measuring 3D TKA kinematics from single-plane radiographic images. Measurement reproducibility and accuracy are comparable to contemporary human-supervised methods. We believe capabilities like this will soon make it practical to perform dynamic TKA kinematic analysis in a clinical workflow, where these measures can help surgeons objectively determine the best course of treatment for their patients.

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BIOGRAPHICAL SKETCH

Andrew Jensen is a Florida native from Sarasota, Florida. He attended the University of Florida for his undergraduate degree in Mechanical Engineer, for which he received high honors. He took a brief hiatus from school to work at an orthopaedic solutions company, Exactech. The COVID-19 pandemic cut his time at Exactech short, so he joined the Gary J Miller Orthopaedic Biomechanics Laboratory as a part-time researcher during the summer leading up to his first official semester of graduate school.

Andrew enjoys being outdoors, hiking, reading, and doing different things.