A PARALLEL COMPUTING TEST BED FOR PERFORMING
AN UNSUPERVISED FLUOROSCOPIC ANALYSIS
OF KNEE JOINT KINEMATICS

by

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dedicated to my husband, Prashant and my parents, Geetha and Ramanatha
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ABSTRACT

Fluoroscopic analysis of knee joint kinematics involves accurately determining the position and orientation of bones in the knee joint. This data can be derived using the static 3-D CT scan images and 2-D video fluoroscopy images together. This involves generating hypothetical digitally reconstructed radiographs (DRR) from the CT scan image with known position and orientation and comparing them to the original fluoroscopic frame. This represents a search problem in which, among all the DRRs possible from a CT image, the image that most closely matches the target fluoroscopy frame of the knee joint has to be found.

Each image in the search space differs from another by the set of position and orientation values. Position is defined by the $x$, $y$ and $z$ co-ordinates in the Cartesian coordinate system. Orientation is defined by the values of azimuth, elevation and rotation. Therefore, this constitutes a six dimensional search problem, and using a brute force method to search for the target image can take a tremendous amount of time. The fact that it is difficult to order or categorize the set of six values adds the complexity of the search. Further, previous research conducted by Scott et al. [1] suggests that even good sequential search algorithms such as the sequential Monte Carlo method can be very time-consuming. Therefore, a better search solution suitable for this kind of 6-D search that provides a significant speedup has to be found. This thesis explores using Swarm Intelligence (SI) techniques in a parallel computing environment to create a test bed for fluoroscopic analysis and increase the speed of the process. Parallel programs are developed using two SI techniques: Bees Algorithm and Particle Swarm Optimization technique.
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LIST OF ABBREVIATIONS

CT – Computer Tomography

**DRR** – Digital Reconstructed Radiograph

SI – Swarm Intelligence

**PSO** – Particle Swarm Optimization

**ACO** – Ant Colony Optimization

**SDS** – Stochastic Diffusion Search
CHAPTER 1

INTRODUCTION

1.1 Fluoroscopic Analysis

A good understanding of knee joint kinematics is essential for evaluating and researching the problems associated with the joint. To understand knee joint kinematics, determination of the position of the bones in the joint during motion in three dimensions is required. Fluoroscopy is a medical imaging technique that produces real-time, two dimensional images of an object, whereas, computer tomography (CT) is an imaging technique that produces static three dimensional image of the scanned object. The two dimensional fluoroscopy frames and the three dimensional CT scan images are not individually sufficient to provide complete information about the position of the bones in the joint in motion as a function of time. However, three dimensional position data over time can be derived using the static CT scan images and real-time information from video fluoroscopy images together.

A digitally reconstructed radiograph (DRR) is a simulation of a conventional 2-D X-ray image or a fluoroscopy frame created from CT data. Once the CT scan and a fluoroscopic video of the joint are generated, a DRR image can be generated using the CT scan image. This requires a simulation of the fluoroscope with the X-ray source or camera and the X-ray film or screen placed at equal radial distances on opposite sides of the origin. The CT cube is placed at the origin in between the X-ray source and the film. The distance between the X-ray source and the film is fixed and known. The cube in the center can translate in three
dimensions: $x$, $y$, and $z$. The film can rotate while the X-ray source can move in azimuth and elevation dimensions. This simulation is illustrated in Figure 1.1. The projection of the X-rays from the simulated source onto the film through the CT cube produces a DRR image. Different values of the X-ray source’s *azimuth* and *elevation*, the film’s *rotation* and the cube’s $x$, $y$, and $z$ co-ordinates generate different DRR images. If a generated DRR image matches the fluoroscopy frame, the position of the cube, and the orientation of the X-ray source and the film are known. Using this information, a three dimensional rendering of the joint in motion can be constructed.

Figure 1.1: Geometric configuration of the simulated fluoroscope [1]

In order to accomplish this, several algorithms are used for different operations includ-
ing generating DRR images, extracting the edges of DRR images, and comparing the edge images. These algorithms are based on the work of several researchers [2, 3]. Previous experiments used a Monte Carlo search technique to search for the matching DRR image, given an X-ray fluoroscopy image [1]. These experiments were conducted to search for azimuth and elevation of the camera and the rotation of the film. They have demonstrated that this can be achieved to an accuracy of 0.5 degree rotation, azimuth and elevation. The process, however, is very time consuming and calls for advanced search algorithms to reduce the time taken for finding the orientation of the X-ray source, the X-ray film and the position of the bones. This thesis uses swarm intelligence techniques in a parallel computing environment to achieve faster results.

1.2 Swarm Intelligence in a Parallel Computing Environment

Swarm intelligence is a branch of artificial intelligence based on the behavior of a group of entities working together to solve a common problem. “It is based on the collective behavior of decentralized, self-organized systems” [15, 26].” Swarms move through the search space and their collective intelligence leads the to the solution. Ant colony optimization (ACO), bees algorithm, particle swarm optimization (PSO), and stochastic diffusion search (SDS) are some of the techniques that exhibit swarm intelligence [5, 9, 11, 13]. These techniques have been used in many optimization problems like the traveling salesman problem, telecommunications routing, factory scheduling, etc. Swarm intelligence techniques are known to be robust and easy to implement.

Based on a review of some of the swarm intelligence techniques, bees algorithm and particle swarm optimization have been chosen to solve the search problem in fluoroscopic analysis of knee joint kinematics [5, 9]. The bees algorithm is a “biologically inspired”
technique based on the food foraging behavior of a swarm of honey bees [29, 30]. The bees in the group are divided into scout bees and worker bees. Scout bees go on a random search for flower patches and come back to the bee hive to report their findings. The quality and distance of each of the flower patches found is evaluated. Finally, the best ones are chosen and worker bees are sent to them to collect nectar. The second algorithm used in this thesis, the particle swarm optimization technique, is influenced by the flocking behavior in birds [33]. It consists of several particles organized into close knit groups or neighborhoods. The particles move in the search space searching for the destination by interacting with one another and they collectively arrive at the destination. It is based on the communication and interaction among particles which re-calculate their position and velocity values accordingly and move through the search space to find the solution [32].

Bees algorithm and particle swarm optimization are suitable methods to solve the six dimensional search problem because the entities in the system can begin the search independently and with periodic communication, they can guide each other to go in the right direction toward the destination. They can be easily mapped to the search problem discussed in Section 1.1. It is also easy to implement them in parallel. Each entity, a bee or a particle, can map to a process in the parallel computing system. Both these methods depend on random number generators just like the Monte Carlo method. But they are more easily parallelizable than the Monte Carlo method which assumes a very sequential approach. This thesis entails implementing the solution using both these techniques in a parallel computing environment.
1.3 Prior Research

This thesis is an extension to the research done by Scott et al. [1]. The goal of the research conducted by Scott et al. was to develop a method for collecting accurate, real time three dimensional kinematic data of bones and joints using two dimensional X-ray images resulting from a video fluoroscopy technique combined with the static three dimensional Computer Tomography data [1]. The process used in their research involved creating virtual X-rays from the CT image through the digitally reconstructed radiograph projections. Traditional methods used to achieve kinematic data of the joints require human intervention to initialize the starting position. The method they proposed was minimally invasive and used a Monte Carlo search technique with a variable search range. The search range decreased as matching between the target image and the image created with known position values improved. This was an optimization search. This is a very efficient method for small ranges of position values.

Considering the extent of the allowed movement of the knee joint within the fluoroscope and the allowed width of the angular movements of the X-ray source and the X-ray film, the potential range of the position values is enormous. Therefore, this calls for an efficient search algorithm that is very fast in finding the values required. The purpose of the research in this undertaking is to find suitable parallelized search mechanisms to speed up the process of determining the three dimensional position data of the bones and joints using the same image processing techniques that were used by Scott et al.

1.4 Thesis Statement

The objective of this thesis is to create a test bed or a framework for fluoroscopic analysis process. The framework will include implementation for all the components of the
fluoroscopic analysis in a parallel computing environment. This implementation is aimed at increasing the speed of the search process involved in fluoroscopic analysis. Swarm intelligence algorithms, namely, the bees algorithm and the particle swarm optimization are used for implementing the search component. The test bed is implemented in such a way that it is easy to use alternative algorithms for each component and also to test the framework with different input images.
In this chapter, the meaning of fluoroscopic analysis and the algorithms used in the analysis are discussed.

2.1 Fluoroscopy

Fluoroscopy is an imaging technique that uses a fluoroscope to obtain real-time X-ray images of a moving body part such as bones, muscles, and organs such as the heart. “A fluoroscope consists of an X-ray source and a fluorescent screen between which a patient is placed” [21]. Figure 2.1 shows a modern day fluoroscope. “Modern fluoroscopes couple the screen to an x-ray image intensifier and video camera allowing the images to be recorded and played on a monitor” [21]. Thus it is possible to view an X-ray movie of the body part being analyzed. The X-ray video is displayed using a continuous series of images produced in quick succession, similar to a television video transmission. The fluoroscopic technique has been used in several medical procedures such as diagnosing problems in the organs in the human body, analyzing functioning of the heart and other moving organs and guiding several kinds of surgeries, procedures and placements of artificial implants.
Figure 2.1: Fluoroscope [16]
2.2 Computer Tomography

Computer Tomography (CT) is a medical imaging technique that uses X-rays. It is based on the concept of tomography. “The word ‘tomography’ is derived from the Greek tomos (slice) and graphein (to write)” [22]. This means a series of slices or cross sectional images are stacked together to construct a three dimensional model of the object being scanned. These images are processed using a computer and several individual images for each slice are created. After completing all the required processing of these images, a detailed three dimensional picture of the internal structure of the body part that was scanned can be obtained. Figure 2.2 shows a modern CT scanning equipment. The patient is placed on the horizontal surface and the scanner takes cross sectional images and computes a CT scan image of the required body part [17].

A CT scan can be used to study many organs and parts of the human body [18]. A CT scan image of knee joint can be helpful in diagnosing the problems in the kinematics of the knee joint and can be useful in the placement of an artificial implant. An example of a CT scan is shown in Figure 2.3, which is a CT scan image of a knee joint.

2.3 Knee Joint Kinematics

A knee joint connects the thigh with the leg. It joins the two bones: femur, the thigh bone, and tibia, the leg bone. Figure 2.4 shows the anatomy of a knee [19]. A knee joint is one of the largest and most complicated joint in the human body. The knee joint allows different kinds of movements between the bones. The types of movements that are possible with the knee are flexion, extension, medial and lateral rotation. The knee joint is the key to the movement of human beings from one place to another.
Figure 2.2: CT Scanner [17]
Knee disorders or injuries can make the movement of the knee very difficult and painful. Surgery or any kind of knee treatments require accurate diagnosis and measurements. X-rays and CT scans are used to study the structure of the knee and take appropriate measurements. However, sometimes, diagnosis requires more than just static images and measurement of the joint structure. The different kinds of movements possible with the knee joint, the friction between bones connected during motion, the impact of the surrounding ligaments and the knee cap and other biological factors increases the complexity of diagnosis. Therefore, the structure of the knee joint has to be studied in vivo. Study of the knee joint kinematics is crucial to diagnosing knee problems. The positions of the bones and the joint in motion have to be determined. In addition, to diagnosing knee joint conditions, this kind of information can assist in the diagnosis of other joint problems.
A CT scan and an X-ray are sufficient to determine the position of the bones in a stationary position. However, doing the same while the joint is in motion is a very difficult task. In the past, invasive position capture techniques were used. Research conducted by Scott et al. developed a method to determine three dimensional real-time kinematic data of the knee joint without using invasive methods [1]. The subsequent sections in this chapter will describe how this was achieved. The focus of the research involved in this thesis is to decrease the time taken to conduct this operation.
2.4 Fluoroscopic Analysis

The process of determining three dimensional position data over time can be achieved using 3-D CT data and 2-D fluoroscopy images. This process is called fluoroscopic analysis. There are several steps required in this process. Figure 2.5 provides a block diagram of the major steps involved.

![Fluoroscopic Analysis Diagram](image)

**Figure 2.5: Procedure for fluoroscopic analysis**

Consider a fluoroscopy environment in which the knee joint is placed. Refer to Figure 1.1 in Chapter 1. The knee joint is placed in between the X-ray source and the X-ray film. The position of the knee joint is affected by the lateral movements of the joint in any of the three directions (x, y, and z) about the origin. Secondly, the position is influenced by
the azimuth and elevation of the X-ray source and thirdly, by the rotation of the X-ray film about a central axis. The knee joint can also change its azimuth, elevation and rotation values, but in this experiment, those values for the joint are kept constant, while the film and the X-ray source are rotated instead. Thus, values of the six dimensions have to be measured over time in order to obtain the real-time positional information of the knee joint.

Using the static CT scan image as input, the 3-D real time position data of the knee joint is constructed. In the first step, virtual X-ray images of the joint at different positions are created using the CT data. Then, the essential features of the image which are useful to determine the position are extracted. The essential features of the target X-ray frame from the fluoroscopy video images are also extracted. In the next step, the virtual image is compared with the target image. This process continues until there is a match. A match indicates the position of the joint for the first target fluoroscopy image. The whole process must be repeated for each fluoroscopy image in the video.

This process provides a rendering of the 3-D positional data over time. Sections 2.4.1 to 2.4.4 provided details for each of the above steps.

2.4.1 DRR Image Creation

A Digitally Reconstructed Radiograph (DRR) is a simulation of a 2D X-ray image, created from CT data. Ray tracing involves constructing a DRR from a CT image by calculating the radiological path through a three-dimensional CT array. Siddon’s algorithm [2] is used for this purpose. This method considers the CT data as consisting of the intersection volumes of three orthogonal sets of equally spaced, parallel planes. The intersection of the ray with the volume elements (voxels) are obtained as a subset of the intersections of the ray with the planes. For each voxel intersection length, the corresponding voxel indices are obtained and the products of the intersected length and voxel density summed over all
the intersections yields the radiological path and attenuation. The ray tracing mechanism employed here is similar to the ray tracing used in computer graphics to generate 3-D images. But in computer graphics, the reflection or the path of the light rays off the pixels is used to generate the image, whereas, in fluoroscopic analysis, the transmittance of the X-rays to the film is used to generate the DRR image.

2.4.2 Edge Detection

Edge detection of an image is used to “identify points in the image at which the image brightness changes sharply or more formally has discontinuities” [23]. By applying an edge detector to an image, the amount of information contained in the image can be significantly reduced by filtering out the less relevant information and retaining only the structural properties of the object in the image. Using edges for comparison increases the probability of finding a match between a DRR image and a fluoroscopy image, as opposed to the comparison of the images themselves whose X-ray intensities may not match. This also decreases the amount of data to be processed and thus significantly simplifies the computations that involve interpreting the information contained in the image. In our implementation, Sobel’s edge detection method is used [4, 24]. This method finds the edges in an image using the gradient filter method.

In Sobel’s edge detection method, a pixel location is declared an edge location if the value of the gradient exceeds some threshold. Edges will have a higher intensity gradient than uniform regions. Whenever the gradient value exceeds the threshold value, an edge is detected. Sobel’s edge detector uses two 3x3 convolution masks. The two masks are applied to each pixel, one with a horizontal trend and one with a vertical trend. The convolution mask, which is much smaller than the actual image, is slid over the image, manipulating a square of pixels at a time. The Sobel masks \( G_x \) and \( G_y \) for the two gradients
are shown in Figure 2.6. The $G_x$ mask highlights the edges in the horizontal direction and the $G_y$ mask in the vertical direction. After taking the magnitude of both, the resulting output detects the edges in both directions.

The magnitude of the gradient is calculated using

$$|G| = \sqrt{G_x^2 + G_y^2}. \quad (2.1)$$

An approximate magnitude can be calculated using

$$|G| = |G_x| + |G_y|. \quad (2.2)$$

### 2.4.3 Image Matching

The target X-ray fluoroscopic frame has to be compared with a DRR image generated using arbitrary camera and film orientation, and cube displacement values. Both the images are converted to edge images using the Sobel filter and then compared. Our implementations involve several such comparisons with images generated using different camera and film orientation and cube displacement values. A contour matching algorithm is used to evaluate the overlap of two edge images [3]. The matching score resulting from a comparison is a value between 0 and 1. A value of 1 indicates a perfect match. The matching score is
obtained by multiplying the two images together pixel by pixel and summing the result and then normalizing by the sum of the predicted image. If \( J(x, y) \) is the input contour image and \( K(x, y) \) is the predicted contour image, then contour match score (CMS) is calculated using

\[
CMS = \frac{\sum_{(x,y)} J(x, y)K(x, y)}{\sum_{(x,y)} K(x, y)}. \tag{2.3}
\]

2.4.4 Searching

The position of the knee joint is defined by six values, viz., the \( x \), \( y \), and \( z \) co-ordinate values and the azimuth, elevation and rotation values. Since each of these values has a large range, a simple brute force method of searching is highly inefficient. Each iteration of the search is a time-consuming process, because it involves creating a virtual image with known position values using ray tracing, extracting the edges and comparing it to the target image. Therefore, the goal should be to reduce the number of iterations of the search as well as the range of search with each iteration.

The measurement of the actual fluoroscope gives the range of values possible for each of the six positional parameters. The azimuth values of the X-ray source can range from -180 to +180 degrees and the elevation can be anywhere between -90 to +90 degrees. The film can rotate from -90 to +90 degrees. The knee joint or the hypothetical CT cube that is used in our experiments has a wide range of translation possible. Typically the object being scanned is placed close to the center and hence, for the tests conducted in this project, -5cm to +5cm translations from the center, on each axis, are considered.

Efficient search algorithms, like binary search, cannot be effectively applied to a six dimensional problem of this proportion. With a binary search algorithm, we can only divide the values for one of the six dimensions into half. Even though this can potentially cut down the number of permutations of the six values to half of them, the number of
remaining permutations is still dramatically large. And moreover, there is simply no way to sort the values for any of the six parameters due to the nature of the comparisons performed in the search. All search algorithms involve comparisons of the elements being searched for in one way or another. In this case, we compare two images and the matching algorithm used for this comparison produces a match score between 0 and 1 for each comparison. The values of the six positional parameters are known when the score is 1. The images are being compared and not the six values. There is no easy way to sort the images in any manner. Thus, this indicates that the usage of any search algorithm that depends on sorted elements is ruled out.

Some search algorithms that use hashing methods require that the elements are organize-able into some form of groups. The search problem presented in this project does not fit well into this category either. Monte Carlo search methods that depend on random sampling and a few swarm intelligence algorithms are useful for this kind of search problems. The details for the Monte Carlo and the swarm intelligence algorithms are presented in Chapter 3.
CHAPTER 3

SEARCH ALGORITHMS

In Chapter 2, the necessity of an efficient search algorithm in fluoroscopic analysis was explained. This chapter provides details of the algorithms adopted in this thesis. In addition to the Monte Carlo method that was used in prior research, the two SI techniques used in this thesis, the bees algorithm and particle swarm optimization, are presented here.

3.1 Monte Carlo Search Technique

The Monte Carlo search method is a non-deterministic method that depends on random sampling. The Monte Carlo method is generally used to solve problems that are difficult to solve using deterministic methods [14, 25]. This method solves problems by generating random numbers in each try. After evaluating each of these trials, it can be predicted if the solution has been reached or not, or if the system is getting closer to the solution. Monte Carlo methods have been heavily used in a wide variety of fields ranging from economics to nuclear physics.

In the research conducted by Scott et al., the Monte Carlo method is used to search for the DRR image that matches the target image [1]. In order to search, a DRR image must be created with different values of camera’s azimuth and elevation, and the film’s rotation angles in each iteration. The DRR image thus created is compared to the target frame to find a match. Initially, the camera azimuth, elevation, and film rotation angles are set to
1. Initialize the current parameter value to zero.
   // This current value is evaluated based on the expected
   // target value in each iteration.
2. While (stopping criterion not met)
3. Generate random values and add it to the initial value.
4. Evaluate the fitness of the value obtained in step 3.
5. Negate the random values and add it to the current value.
6. Evaluate the fitness of the value obtained in step 5.
7. Select the better of the two values.
   (This becomes the current value)
8. End While.

Figure 3.1: The pseudocode for the Monte Carlo search algorithm

arbitrary values generated using a random number generator. Random step values with a range of 0 to 10° are generated for each iteration of the search. Two sets of orientation values are obtained by adding and subtracting the step values to the initial values. DRR images are then generated using both these sets. A contour match score is calculated for the comparison of each of these images with the edge image of the target fluoroscopy frame. The parameter values that produce a better score is retained and the path that is more promising is chosen. Figure 3.1 gives the pseudocode for the Monte Carlo search algorithm.

The search continues by producing two new sets of random steps, which are added and subtracted to the current values obtained from the previous iteration. This cycle is repeated until a match is found and the score of the match is greater than 0.8. Thus, in each iteration of the search, the search space decreases and the search gets closer to the solution. This results in values that are approximately equal to the desired values.

Once the preliminary search is completed, a fine search is performed with smaller step values. The fine search is used to search closer to the parameter values obtained from the previous round of search. It is assumed that the first stage of the search results in a value
that is close to the solution and hence the next stage of the search is expected to obtain a more accurate solution by searching in smaller steps. After both the stages are completed, the resulting parameter values should be fairly accurate. The sequential implementation of this method has been able to generate values within $0.5^\circ$ of the desired parameter values.

3.2 Swarm Intelligence

The word *Swarm* means a group of insects that are close to one another and engaged in a common activity. In general, *swarm* is a term used to describe the behavior of a group of animals that are similar to one another. Swarming behavior can be observed in many animals such as bees, ants, other insects, fish, bacteria and even people. This phenomenon can be called by different names such as flocking in birds, schooling in fishes, societal behavior in human beings and growth of colonies in bacteria.

Swarm intelligence is a phenomenon, based on self-organization, in which the group of insects or other beings solve complex problems together. The swarm arrives at the solution collectively. Constant and consistent communication and feedback among the group, and adherence to common rules are essential to the success of the swarm. Errors and randomness are the pillars of this system. Even if a few individuals fail to achieve their objective, the group as a whole can succeed. Examples of this behavior can be observed in the food foraging behavior in ants and bees, and migration in birds and fishes. Chances of foraging food, finding mates, avoiding predators, overcoming obstacles, migrating to a better place are made faster and easier by swarm intelligence. Swarm intelligence has a strongly noticeable benefit for all the members of the swarm.

The swarm intelligence concept, when applied to computing, can help solve many computing problems. This is a branch of artificial intelligence in which interaction and
co-operation among entities in a system is the key to solving a problem. The expression
Swarm intelligence (SI) was introduced by Gerardo Beni and Jing Wang in 1989 in the
context of cellular robotic systems [15]. There are several SI techniques such as the ant
optimization algorithm, bees algorithm, particle swarm optimization, and stochastic diffu-
sion search. SI techniques can be applied in problems involving finding the shortest path,
swarm-based data analysis, robotics, financial forecasting, search, network management,
Internet server optimization, etc. SI methods generally have simple rules and are easy to
implement.

Though the Monte Carlo method is suited for the search problem in fluoroscopic
analysis, it is very sequential in nature. It is difficult to implement it in parallel and
hence not chosen for implementation in this thesis. SI techniques were explored and it
was found that some of them are either not easy to map to the search problem in the
fluoroscopic analysis process or to implement in a parallel computing environment. Ant
colony optimization algorithm [27], which is very useful in the shortest path problems,
is based on the concept of leaving behind clues in the path and the strength of the clue
determines whether the path is desirable or not. Ants leave behind a substance called
‘pheromone’ on the paths they travel, both on the way to the destination and back. If the
path is long, the pheromone evaporates and its strength reduces. If the path is short, the ant
reinforces the pheromone by depositing on the same spot again on its way back, without
too much of a time-gap and thus strengthens it in the process. When other ants visit the
same path, if the strength of the pheromone is high then they are encouraged to take that
path. If it is weak, the path is abandoned. This is not as easy to map to the search problem
in fluoroscopic analysis as some other SI algorithms. Stochastic diffusion search [28] is
another SI algorithms that is used to solve search and optimization problems. They expect
that the objective function can be broken down into independent components that can be
worked on by individual entities in the population. An entity is expected to communicate with every other entity in the system. This leads to significant communication overhead and is therefore not very suitable for parallel implementation.

The two SI techniques chosen to solve the search problem in fluoroscopic analysis are bees algorithm and particle swarm optimization. The group of entities in both methods can be easily mapped to processes in a parallel computing system. In addition to being easily parallelizable, they are also easy to implement. Sections 3.2.1 and 3.2.2 describe these two methods in depth.

### 3.2.1 Bees Algorithm

The bees algorithm [5, 29] is inspired by the behavior of a honey bees working together to find the sources of nectar, collecting it and storing it back in their hives. It is a population-based search algorithm [5]. This approach has been used to solve many optimization problems.

A honey bee colony has many worker bees with specific roles. The main functions of the colony are to identify food sources and collect and store nectar in the bee hives. Scout bees are given the task of finding nectar sources. The scout bees begin searching in random directions. When they come back to the hive, they communicate about their findings to all other bees by performing a dance known as the *Waggle Dance*. *Waggle Dance* [7, 31] is the “figure-eight dance” performed by the scout bees to share the information about the distance, direction and quality of the flower patch. While dancing the bees move in the form of a figure-eight. If a line passing through the sun and the designated dance area is considered as a reference line, the angle between that line and the direction of the waggle dance indicates the direction of the food source. It could be in the direction of the sun, to the right or left of the sun. The duration of the waggle dance indicates the distance of
the flower patch. Figure 3.2 depicts the bee waggle dance. All the scout bees perform the waggle dance after returning from the flower patches. After learning about the distance and quality of all the flower patches through the waggle dances, the flower patches are evaluated based on some criteria. A few best sites are chosen from these if their fitness value is above a certain threshold. A large number of worker bees are recruited to collect nectar from the most promising sites and the remaining worker bees are sent to the other flower patches. Sites that are very far and those that have very few flowers with nectar and pollen are discarded. The scout bees are sent again in search of new flower patches for the next round.

Figure 3.2: Waggle Dance [31]
1. Initialize the population with random solutions.
2. Evaluate fitness of the population.
3. While (stopping criterion not met)
   //Forming new population.
4. Select sites for neighborhoods search.
5. Recruit bees for selected sites
   (more bees for best e sites)
   and evaluate fitnesses.
6. Select the fittest bee from each patch.
7. Assign remaining bees to search randomly
   and evaluate their fitnesses.
8. End While.

Figure 3.3: The pseudocode for the bees algorithm in its simplest form [6]

The bees algorithm in its simplest form requires a number of parameters [6], namely:

1. number of scout bees (n)
2. number of sites selected out of n visited sites (m)
3. number of best sites out of m selected sites (e)
4. number of bees recruited for best e sites (nep)
5. number of bees recruited for the other (m-e) selected sites (nsp)
6. initial size of the patches (ngh)
7. stopping criterion

The pseudocode is presented in Figure 3.3.

The bees algorithm is illustrated using Figures 3.4 and 3.5. In Figure 3.4, the scout bees set out in random directions away from the main bee hive. Some flower patches are big while some are small. Some are very far from the hive while some are very close. There
can also be more than one way to reach a flower patch. In Figure 3.5, the returning scout bees perform the waggle dance in the bee hive and as a result of this the majority of the worker bees fly to the larger and closer sites via the shortest route. Scout bees go in search of new sites.

Figure 3.4: Food foraging in bees - scout bees are going in search of flower patches

3.2.2 Particle Swarm Optimization

“Particle swarm optimization (PSO) is a population based stochastic optimization technique developed by Dr. Eberhart and Dr. Kennedy in 1995, inspired by social behavior of bird flocking or fish schooling” [8–10, 33]. A particle swarm system is similar to a social
Figure 3.5: Food foraging in bees - worker bees are going to the best flower patches
network and depends on a communication structure in the network. Just as a person benefits from his interaction with society, a particle in the swarm enhances its position in the system due to its interaction with other particles. People in a society talk to other people to find out their experiences, and try to solve their problems using the inputs given by others. Other people’s views and attitudes might affect an individual’s own opinions and future decisions. Some people are more influential than the others. There are various factors that can affect the social dynamics such as who the neighbors are, how they influence the direction an individual takes in his journey to a destination and his own progress at any moment.

Particle swarms are used to simulate social networks as well as several engineering applications. Particle swarm optimization techniques can be used in search problems, where the particles in the swarm communicate with one another and reach the solution faster than they would have if they were on their own. This technique has also been used to address other complex optimization problems, pattern recognition, data mining and image processing.

The particle swarm can be modeled as a population of particles in a multi-dimensional space [32]. Each particle has a position and a velocity with which it moves. All particles have a starting point and they move randomly in the search space. As the particles move from one position to another, they have to evaluate the fitness of that position based on some criteria, which determines how close they are to the solution. Navigation of the particles in the correct direction depends significantly on the accuracy of the fitness evaluation function. Periodically, a particle finds the best position it has held among the previous set of positions. It resets its position to the best position and continues the search. Each particle has a set of designated neighbors with which it should interact at regular intervals. The particles should not only keep track of the local best value, but also find out the best value in the neighborhood by communicating with other particles in the neighborhood. There is
1. Initialize the particle positions and velocities.
2. While (global best value does not match solution)
3. Move particle to a random position from the current position within a certain threshold from the current position.
4. Store the local best value.
5. Talk to neighbors and exchange neighborhood search information.
7. Store global best value.
8. Update position and velocity to the neighborhood best.
9. End While.

Figure 3.6: The pseudocode for the general PSO [32]

a particle designated in every neighborhood to keep track of the best global value. The particles adjust their positions at regular intervals based on the best local, neighborhood and global values. Thus they move towards those particles in their neighborhood, that have the best fitness values. Over time they converge toward good positions and move about in the search space in close proximity to the best global value and stop exploring the rest of search space. The rest of the search space, which has a very low likelihood of containing the solution, is discarded. The pseudocode that explains this technique is given in Figure 3.6.

Figures 3.7, 3.8 and 3.9 illustrate this concept. Figure 3.7 shows a swarm of fifteen particles organized into three groups of five particles. The particles in the three groups are denoted by circles, squares and triangles, respectively. In this figure, all the particles are initialized to some random values. In Figure 3.8, after fitness evaluation is conducted by all the particles, they are communicated to all particles in the neighborhood. These values help the particles to decide if they can continue from their current position or if they should follow a neighbor. This communication results in the movement of the particles
toward the neighbor with the best fitness value. In ideal cases, after several iterations of this exercise, the particles converge toward the best global fitness value and thus converge to the solution. This stage looks similar to what is shown in Figure 3.9. All the particles then begin to search intensively around the best value until they find the solution.

![Diagram of particle movement](image)

**Figure 3.7:** Particles in the swarm are moving randomly toward their destination

The application of the bees algorithm and the particle swarm optimization to the image search part of fluoroscopic analysis is described in detail in Chapter 4.
Figure 3.8: Particles are getting closer to the best position of the neighborhood
Figure 3.9: Particles are getting closer to the best global position
CHAPTER 4

DESIGN AND IMPLEMENTATION

The search problem involved in the fluoroscopic analysis process has been described in Chapter 1. While Chapter 2 provided an introduction to the world of fluoroscopic analysis and presented the computing problems associated with the process, Chapter 3 presented different methods to solve the search problem in fluoroscopic analysis. In this chapter, the design and implementation details as well as the pseudocode for the bees algorithm and particle swarm optimization method used for this thesis are provided.

4.1 Design Approach

This thesis consists of providing solutions to a search problem in a parallel computing environment. After finding the suitable algorithms for the search problem, a pseudocode for implementing it in the parallel environment was developed. Additionally, pseudocode for the sequential version of the algorithm was also developed. In order to measure the speed and effectiveness of the parallel implementation, comparison with its sequential counterpart is required. After a detailed analysis of the fluoroscopic analysis application, and exploration of different search strategies, two swarm intelligence algorithms, Bees and PSO were chosen. Developing pseudocode for these two algorithms for parallel and sequential implementation was the first step toward the design of this project.
The second step consisted of developing a good organization structure and an activity flow-chart, which are very essential to a good design. All the steps involved in the process were studied. Each step required an separate component. For fluoroscopic analysis, different components are required for specific tasks, such as, generating DRR images from a given position and a CT scan image, image filtering for extracting the edges of an X-ray or DRR image, image matching, and fluoroscopic searching. Figure 2.5 in Chapter 2 provides an activity flow-chart for fluoroscopic analysis that illustrates the different stages in the fluoroscopic analysis and the steps to transition from one stage to another.

The third step involved identifying the different entities in the system and designing data structures for each of them. The main entities are:

1. CT Volume - used to represent a CT scan image.
2. DRR Image - used to represent a DRR X-ray image.
3. Bee - used to represent a bee in bees algorithm.
4. Particle - used to represent a particle in PSO method.
5. Sobel filter kernels - used in the Filtering method to extract edges of an X-ray image

These data structures are presented in Appendix B.

Figure 4.1 provides the pseudocode for the general fluoroscopic analysis. Figures 4.2, 4.3 and 4.4 provide the pseudocode for the sequential bees algorithm and particle swarm optimization method respectively. The pseudocode for the parallel bees algorithm is shown in Figures 4.5 and 4.6, while Figures 4.7 and 4.8 provide the pseudocode for the parallel particle swarm optimization method.
Fluoroscopic Analysis(CTCube, TargetXrayImage)

1. TargetEdges = SobelFilter(TargetXrayImage)
2. score = 0
3. While(score < GOOD_MATCH_SCORE)
   4. x = Random()
   5. y = Random()
   6. z = Random()
   7. azimuth = Random()
   8. elevation = Random()
   9. rotation = Random()
   10. TestDRRImage = GenerateDRR(CTCube, x, y, z, azimuth, elevation, rotation)
   11. TestEdges = SobelFilter(TestDRRImage)
   12. score = ContourMatching(TestEdges, TargetEdges)
5. Match found.
   Position values are x, y, and z.
   Orientation values are azimuth, elevation, and rotation

Figure 4.1: Pseudocode for fluoroscopic analysis in the simplest form

4.2 Build Infrastructure

Build infrastructure provides the capability to add new code to the project, compile and maintain the code. The project files are organized in such a manner that it facilitates the ease of compiling and executing the application. All the code and the input files are placed in a repository controlled by a version control system called Subversion. The main makefile is placed in the main directory of the project. This makefile facilitates all the build actions required for the project. The makefile targets and the functions performed by them are given in Appendix A.

The directories are organized in such a way that it becomes convenient to maintain different libraries, input images, results, and configuration individually without affecting other parts of the project. The directory structure of the project files is provided in Appendix
Sequential_Bees(CTCube, TargetXrayImage)

n = Number of bees
m = Number of best sites
p = Number of bees chosen for the best sites (best bees)
q = n-p = Number of bees chosen as scout bees for the next iteration
r = Number of individual bee iterations

1. TargetEdges = EdgeFilter(TargetXrayImage)
2. While(BestScore < GOOD_MATCH_SCORE):
3.   Foreach Bee:
4.     For i = 0 to p:
5.       if(scout bee)
6.         //In the first iteration of #1, all bees are scout bees
7.         Generate random values for position(x,y,z) and orientation(az,el,rt)
8.       else if(worker bee)
9.         Generate random step values and add them to the previous values.
10.        TestDRRImage = GenerateDRR(x,y,z,az,el,rt, CTCube)
11.        TestEdges = EdgeFilter(TestDRRImage)
12.        BeeScore = ContourMatching(TestEdges,TargetEdges)
13. BeeScore = Calculate best of all the p scores

//Waggle dance section
14. Rank the BeeScore of all the individual bees
15. Choose the m best sites from the list based on scores.
16. Assign p bees to m sites equally (worker bees)
17. Assign q bees for random search (scout bees)
18. Save the BestScore and the values associated with it
19. [x,y,z,az,el,rt] associated with BestScore are the required position and orientation values.
20. End

Figure 4.2: Pseudocode for sequential bees algorithm
Sequential_PS0(CTCube, TargetXrayImage)

n = Number of particles
m = Number of neighborhoods
p = Number of particles in a neighborhood
r = Number of individual particle iterations

1. TargetEdges = EdgeFilter(TargetXrayImage)
2. Foreach neighborhood m:
3. NeighborhoodBest = 0.0
4. GlobalBest = 0.0
5. While(BestScore < GOOD_MATCH_SCORE):
6. Foreach Neighborhood m:
7. if NeighborhoodBest < 0.1:
   //This means, search randomly
   RANDOMSTEP = 0
8. else if NeighborhoodBest < 0.4
    //Add large random steps to previous value
    RANDOMSTEP = LARGESTEP
9. else if NeighborhoodBest < 0.6
    //Add medium random steps to previous value
    RANDOMSTEP = MEDIUMSTEP
10. else if NeighborhoodBest < 0.8
    //Add small random steps to previous value
    RANDOMSTEP = SMALLSTEP
11. end if
12. end Foreach Neighborhood m:
13. end While

Figure 4.3: Pseudocode for sequential PSO
Foreach neighbor p:
For i = 0 to r:
    if(RANDOMSTEP == 0):
        //In the 1st iteration all particles
        //search randomly
        Generate random values for position(x,y,z)
        and orientation(az,el,rt)
    else:
        Generate random step values and
        add them to the previous neighborhood best
        values.
    TestDRRImage =
    GenerateDRR(x,y,z,az,el,rt, CTCube)
    TestEdges = EdgeFilter(TestDRRImage)
    ParticleScore =
    ContourMatching(TestEdges,TargetEdges)
    ParticleBestScore =
    the best of all the r particle scores
    NeighborhoodBest =
    the best of all scores in the neighborhood
    GlobalBest = the best of all scores
End While
[x,y,z,az,el,rt] associated with GlobalBest
    are the required position and orientation values.
End

Figure 4.4: Pseudocode for sequential PSO, ctd
Parallel_Bees(CTCube, TargetXrayImage)

n = Number of bees/processes
m = Number of best sites
p = Number of bees chosen for the best sites (best bees)
q = n-p = Number of bees chosen as scout bees for the next iteration
r = Number of individual bee iterations

1. TargetEdges = EdgeFilter(TargetXrayImage)
2. Barrier() //Wait for all processes to arrive at this point
3. While(BestScore < GOOD_MATCH_SCORE):
4.   Foreach Bee:
5.     For i = 0 to p:
6.        if(scout bee)
7.           //In the first iteration of #1, all bees are scout bees
8.           Generate random values for position(x,y,z) and orientation(az,el,rt)
9.        else if(worker bee)
10.           Generate random step values and add them to the previous values.
11.     TestDRRImage = GenerateDRR(x,y,z,az,el,rt,CTCube)
12.     TestEdges = EdgeFilter(TestDRRImage)
13.     BeeScore = ContourMatching(TestEdges,TargetEdges)
14.     BeeScore = Calculate best of all the p scores

Figure 4.5: Pseudocode for parallel bees algorithm
//Waggle Dance section
14. if( process_id != 0):
   //All processes other than process 0,
   //send their scores and 6 value to process 0
   Send(BeeScore,x,y,z,az,el,rt) to Process 0
15. else if(process_id == 0):
   for(i =1 to n):
      Recv(BeeScore,x,y,z,az,el,rt)
      from Process i
   19. Rank the BeeScore of all the individual bees
20. Choose the m best sites from the list based on scores.
21. Save the BestScore and the values associated with it
22. for(i=1 to n):
   //Send information about best sites to the
   //best bees. Make some bees random.
   Send(BeeScore, x,y,z,az,el,rt,ScoutOrWorkerFlag)
   to Process i.
23. if(process_id != 0)
24.    Recv(BeeScore, x,y,z,az,el,rt,ScoutOrWorkerFlag)
25.    from Process 0
26. End While
27. [x,y,z,az,el,rt] associated with BestScore
   are the required position and orientation values.
28. End

Figure 4.6: Pseudocode for parallel bees algorithm, Ctd
Parallel_PSO(CTCube, TargetXrayImage)

n = Number of particles/processes
   Every particle in the system has a particle_id
m = Number of neighborhoods/process groups
p = Number of particles in a neighborhood
   Every neighbor in a neighborhood has a neighbor_id
r = Number of individual particle iterations

1. TargetEdges = EdgeFilter(TargetXrayImage)
2. Create m ProcessGroups //Neighborhoods
3. Initialize all neighbors in a group/neighborhood
4. GlobalBest = 0.0
5. Barrier() //Wait for all particles to get initialized
6. While(BestScore < GOOD_MATCH_SCORE):
      //This is done by all neighborhoods.
    7. if NeighborhoodBest < 0.1:
       8. RANDOMSTEP = 0 //This means, search randomly
    9. else if NeighborhoodBest < 0.4
       10. RANDOMSTEP = LARGESTEP
           //Add large random steps to previous value
    11. else if NeighborhoodBest < 0.6
       12. RANDOMSTEP = MEDIUMSTEP
           //Add medium random steps to previous value
    13. else if NeighborhoodBest < 0.8
       14. RANDOMSTEP = SMALLSTEP
           //Add small random steps to previous value
    15. For i = 0 to r:
      16. if(RANDOMSTEP == 0):
           //In the 1st iteration all particles search randomly
          17. Generate random values for position(x,y,z)
             and orientation(az,el,rt)
      18. else:
             Generate random step values and
             add them to the previous neighborhood best values
      19. TestDRRImage = GenerateDRR(x,y,z,az,el,rt, CTCube)
      20. TestEdges = EdgeFilter(TestDRRImage)
      21. ParticleScore= ContourMatching(TestEdges,TargetEdges)
      22. ParticleBestScore= the best of all the r particle scores

Figure 4.7: Pseudocode for parallel PSO
if(neighbor_id != 0):
    //All neighbors send their score
    //and values to neighbor with id=0
    Send(ParticleBestScore, x,y,z,az,el,rt)
to neighbor 0
else:
    //neighbor 0 receives all the scores
    for( i=1 to p):
        Recv(ParticleBestScore, x,y,z,az,el,rt)
        from neighbor i
    Store the NeighborhoodBest score
    //Send the neighborhood best score and
    //values back to others
    for( i=1 to p):
        Send(NeighborhoodBest, x,y,z,az,el,rt)
to neighbor i

if(neighbor_id !=0):
    Recv(NeighborhoodBest, x,y,z,az,el,rt)
    from neighbor 0

if(particle_id != 0):
    Send(NeighborhoodBest, x,y,z,az,el,rt)
to particle 0
else:
    for( i=1 to n):
        Recv(NeighborhoodBest, x,y,z,az,el,rt)
        from particle i
    Store the GlobalBest score
End While
[x,y,z,az,el,rt] associated with GlobalBest
are the required position and orientation values.
End

Figure 4.8: Pseudocode for parallel PSO, Ctd
4.3 Implementation Details

Implementation of this application consists of several files of C code that implement all the algorithms required for each step in fluoroscopic analysis. The inputs to this application are DRR images of pre-determined positions and orientation or an actual 2-D X-ray fluoroscopic frame as the target images, CT scan image, and configuration files that describe the parameters required to define the algorithms. The program outputs the position and orientation of the best matching score. By configuring the makefile to generate the debug version of the program, the program can be made to print out more information about each iteration of the search.

The algorithms are implemented as library plugins that can be dynamically linked to the main program. The main program performs all the main steps of extracting the CT volume information from the input CT image, and making dynamic calls to the required libraries for each of the other functions such as searching, generating DRR, etc. By implementing the algorithms as dynamically linked plugins, the flexibility for using any algorithm for a specific task is provided. This makes it possible to implement a new algorithm for any step and compile it independently. No changes are needed to any other part of the project to use the new plugin. The new plugin has to only be specified as a parameter while executing the application. This is done to allow further research and experimentation for the fluoroscopic analysis and to make it easy to explore different possibilities of solving this problem.

The environment details required to develop and execute this project is provided in Table 4.1. The instructions to execute this project are provided in Figure 4.9.
<table>
<thead>
<tr>
<th>Operating System</th>
<th>Linux - CentOS release 5.2</th>
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<tbody>
<tr>
<td>Version Control System</td>
<td>Subversion 1.4.2</td>
</tr>
<tr>
<td>Programming Language</td>
<td>C</td>
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<tr>
<td>Libraries</td>
<td>mpich2, nrutils</td>
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<td>Sequential Compiler</td>
<td>gcc</td>
</tr>
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<td>Parallel Compiler</td>
<td>mpicc</td>
</tr>
<tr>
<td>Parallel Computing platform</td>
<td>Beowulf cluster</td>
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<tr>
<td>Scripting Language</td>
<td>Korn Shell</td>
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<tr>
<td>Debugger</td>
<td>gdb</td>
</tr>
<tr>
<td>Memory Profiler</td>
<td>Valgrind</td>
</tr>
</tbody>
</table>

Table 4.1: Environment details

Execution of sequential program:

```plaintext
fluoro_s
<path to search plugin(.so lib file)>
<path to the input config file>
```

Execution of parallel program:

```plaintext
mpiexec -l -n <num of processes> fluoro_p
<path to search plugin(.so lib file)>
<path to the input config file>
```

Figure 4.9: Program execution

An example config files for the bees algorithm and the particle swarm optimization method is provided in Figures 4.10 and 4.11, respectively.

### 4.4 Test Infrastructure

The configuration driven program execution adopted in this project makes it very easy to set up tests for this project. In this project, the specification of a particular algorithm and a set of parameters constitute a test case. By varying the choice of algorithm and the parameters required to define it, plenty of test cases can be generated. To generate a test case, the creation of a config file with the required details is necessary. This can be given
# Input CT image
CT_IMAGE=<path>/images/really_small_cube.txt

# Target DRR with known position values
DRR_IMAGE=<path>/images/drr_image10_10_10_8_8_8.txt
GENERATE_DRR_LIB=<path>/lib/libp_generate_drr_raytracing.so
FILTER_LIB=<path>/lib/libp_filter_sobel_bw.so
MATCH_LIB=<path>/lib/libp_match_contour_bw.so

MAX_ITER=3
BEE_ITER=1
NUM_BEES=8
NUM_BEST_BEES=6
NUM_BEST_SITES=3
RANDOM_SEED=123
AZIMUTH_RANGE=180 # In degrees
ELEVATION_RANGE=90 # In degrees
ROTATION_RANGE=90 # In degrees
TRANSLATION_RANGE=20 # In pixels
TRANSLATION_FINE_STEP=3 # In pixels
ORIENTATION_FINE_STEP=5 # In degrees

Figure 4.10: Config file for bees algorithm
# Input CT image
CT_IMAGE=<path>/images/really_small_cube.txt

# Target DRR with known position values
DRR_IMAGE=<path>/images/drr_image10_10_10_8_8_8.txt
GENERATE_DRR_LIB=<path>/lib/libp_generate_drr_raytracing.so
FILTER_LIB=<path>/lib/libp_filter_sobel_bw.so
MATCH_LIB=<path>/lib/libp_match_contour_bw.so
MAX_ITER=3
PARTICLE_ITER=1
NUM_PARTICLES=8
NUM_NEIGHBORHOODS=4
RANDOM_SEED=123
AZIMUTH_RANGE=180  # In degrees
ELEVATION_RANGE=90  # In degrees
ROTATION_RANGE=90  # In degrees
TRANSLATION_RANGE=20  # In pixels
TRANSLATION_FINE_STEP_LEVEL1=10  # In pixels
TRANSLATION_FINE_STEP_LEVEL2=5  # In pixels
TRANSLATION_FINE_STEP_LEVEL3=3  # In pixels
ORIENTATION_FINE_STEP_LEVEL1=10  # In degrees
ORIENTATION_FINE_STEP_LEVEL2=5  # In degrees
ORIENTATION_FINE_STEP_LEVEL3=1  # In degrees

Figure 4.11: Config file for PSO
as an input to the program executable to run the test case. This structure makes it easy to write different configuration files and write a script to run the application with each config file and thus easily automate the test execution process. The output from each application can be redirected to a results folder and used to compare with subsequent test executions for the same test case in regression testing.
CHAPTER 5

TESTING AND RESULTS

The implementation details for the fluoroscopic analysis process and the test infrastructure developed are provided in Chapter 4. This chapter lists the efforts to test this implementation.

5.1 Test Cases

Software implementations require thorough testing for verifying the code and also validating the results obtained for the given inputs. Testing the implementation of fluoroscopic analysis process is required for the following purposes:

- To check the correctness of code by running it against various inputs and checking if the program runs to completion successfully.
- To verify if the parallel implementation does generate the required output in shorter time compared to the sequential implementation.
- To find the better of the two SI methods, bees algorithm and particle swarm optimization, for solving the search problem in fluoroscopic analysis.

The implementation for both the sequential and parallel versions provide the time taken for the search operation. This helps in comparing the differences in the time taken for
the two versions. Several test cases are designed to test the implementation to enable the verification of all the points mentioned above.

The implementation for the bees algorithm and the particle swarm optimization algorithm require several parameters to be set to successfully initialize the system. The bees algorithm requires values for the number of bees, number of best sites to choose, the number of bees to allocate to each best site, etc. In the case of particle swarm optimization, the set of parameters include the number of particles, number of neighborhoods and number of members in a neighborhood. These parameter values are placed in a configuration file and given as input to the main program. Examples of the configuration files for the Bees and particle swarm optimization algorithms are provided in Figures 4.10 and 4.11 respectively. In this project, different test cases are derived by specifying different parameters for each algorithm. Therefore, by creating several configuration files, one for each set of test configurations, many test cases are constructed.

An attempt has been made to diversify the parameters in such a way that the program will be tested for different sizes of the groups (bees or particles), different sizes of the sub-groups, different position and orientation values of the target image, and so on. This would help in the determination of an optimized combination of parameters. For example, a particle population consisting of 36 particles grouped into 6 neighborhoods of 6 members each, might give faster results than a population grouped into 12 neighborhoods of 3 members each. In the case of bees algorithm, if 3 best sites are chosen and 10 bees are selected for each of the 3 sites from a population of 36 bees, while allowing the remaining 6 bees to search randomly, the grouping might provide better results than any other grouping of the bees.

Bees algorithm and particle swarm optimization method are both swarm intelligence algorithms that use different mechanisms of grouping and sharing information among one
another within their colony. Which method is more suitable for fluoroscopic analysis? How many entities are required for the search to provide optimal results without costing any severe communication overhead? What is an ideal grouping of the entities involved? Do smaller number of groups with large sizes work better than larger numbers of groups with small sizes? What kind of target images result in good solutions? Does the implementation provide good results only for a small subset of the target images or does it hold for any range? The execution of tests attempts to answer these question that help determine the limitations and strengths of each algorithm by configuring several permutations and combinations of the parameter values. Tables 5.1 and 5.2 provide some of the test cases for the bees algorithm and the particle swarm optimization method respectively. Additionally, these combinations are executed multiple times with different target DRR images and for several number of maximum iterations.

<table>
<thead>
<tr>
<th>Configuration/ Testcase</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of Bees</td>
<td>12</td>
<td>24</td>
<td>24</td>
<td>24</td>
<td>36</td>
<td>36</td>
</tr>
<tr>
<td>Number of Best Bees</td>
<td>9</td>
<td>20</td>
<td>18</td>
<td>16</td>
<td>27</td>
<td>30</td>
</tr>
<tr>
<td>Number of Scout Bees</td>
<td>3</td>
<td>4</td>
<td>6</td>
<td>8</td>
<td>9</td>
<td>6</td>
</tr>
<tr>
<td>Number of Best Sites</td>
<td>3</td>
<td>2</td>
<td>6</td>
<td>4</td>
<td>9</td>
<td>6</td>
</tr>
</tbody>
</table>

Table 5.1: Test Cases for bees algorithm

<table>
<thead>
<tr>
<th>Configuration/ Testcase</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of Particles</td>
<td>12</td>
<td>24</td>
<td>24</td>
<td>24</td>
<td>36</td>
<td>36</td>
</tr>
<tr>
<td>Number of Neighbors</td>
<td>4</td>
<td>6</td>
<td>4</td>
<td>3</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>Number of Neighborhoods</td>
<td>3</td>
<td>4</td>
<td>6</td>
<td>8</td>
<td>9</td>
<td>6</td>
</tr>
</tbody>
</table>

Table 5.2: Test Cases for PSO

In order to study the communication overhead caused in the parallel implementations of the two SI algorithms, a snapshot of the inter-process communication has been taken by adding extra logging to the program and Jumpshot 4.0 software has been used to
visualize the communication in terms of send and receive operations between the processes by reading those logs. Figure 5.1 provides the visualization of the inter-process communication during the execution of bees algorithm. The left pane of the picture lists the 12 processes with id 0 to 11. The timeline is seen at the bottom pane. It can be noted that for each iteration, all the bees or processes send their values to the master process (id = 0) and the master process sends a value after comparison and evaluation of all sent values, back to all the bees. This cycle repeats for every iteration of the search. The inter-process communication for particle swarm optimization algorithm is quite different from this. Figure 5.2 provides the visualization for this communication. In this, it can be seen that in each iteration, subgroups communicate among themselves first, where all the subgroup members send message to and receive from the subgroup leader. The leaders then communicate with the master process. The communication overhead load on the master process is much lesser in the case of particle swarm optimization than bees algorithm. Considering the time taken for each iteration in generating a DRR image and comparing that to a target DRR image, the communication overhead caused by sending and receiving matching score values at the end of each iteration is not very significant.

5.2 Results

Search algorithm implementations require several iterations of DRR image generation and their comparison to target image before finding the required position and orientation values. In sequential execution, each iteration of the search is conducted sequentially and hence takes roughly \( n \times x \) amount of time for \( n \) iterations, if \( x \) is the amount of time taken for one operation of DRR image generation and comparison. In both the parallel implementations, the parallel processes share the number of iterations of the search. In ideal cases, if \( p \) is
Figure 5.1: Communication visualization for the parallel bees algorithm
Figure 5.2: Communication visualization for the parallel PSO method
the number of processes, then the time taken for $n$ iterations reduces by a factor of $n/p$. Of course, the actual value will be less than this if we account for the process communication overhead.

<table>
<thead>
<tr>
<th>Mode of Execution</th>
<th>Algorithm</th>
<th>Max Iteration</th>
<th>Number of Entities</th>
<th>Time Taken (in secs)</th>
<th>Speed Up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sequential</td>
<td>Bees</td>
<td>100</td>
<td>12</td>
<td>1161.554</td>
<td></td>
</tr>
<tr>
<td>Parallel</td>
<td>Bees</td>
<td>100</td>
<td>12</td>
<td>104.893</td>
<td>11.07</td>
</tr>
<tr>
<td>Sequential</td>
<td>PSO</td>
<td>100</td>
<td>12</td>
<td>1183.173</td>
<td></td>
</tr>
<tr>
<td>Parallel</td>
<td>PSO</td>
<td>100</td>
<td>12</td>
<td>104.777</td>
<td>11.29</td>
</tr>
</tbody>
</table>

Table 5.3: Timing results

All the experiments confirmed that the the time taken by the parallel programs were less than the time taken by their sequential counterparts by a significant amount. One such instance is provided in Table 5.3.

These results help in verifying the first test objective and part of the second objective. The sequential and parallel programs have run without any crashes or execution errors for all the configuration values for large number of iterations. In addition, for the same number of iterations, the parallel programs are considerably faster than the sequential programs.

In order to get the results, each component of the fluoroscopic analysis process have to function accurately. One of the most crucial roles in affecting the results is played by the DRR X-ray image matching algorithm. The image matching algorithm chosen in this project provided unexpected matching score values in some cases. This has made it very difficult to obtain the expected position and orientation values from the search operation. Hence, this prevents us from effectively comparing the sequential and parallel implementations and also comparing different methods of searching to find out which method takes us to the result in the shortest amount of time.
Owing to the difficulties in extracting the essential information from the complex X-ray images, and inaccuracy of the matching operation due to the presence of unsuitable information given by prior steps, the second and the third test objectives cannot be fully analyzed. The other reason could also be that the random position and orientation values used to generate the reference DRR images might not represent valid X-ray images. Due to this, all the subsequent processes of filtering and matching are bound to go wrong. There is scope for further research to improve the results and this is discussed in Chapter 6.
CHAPTER 6

CONCLUSIONS

6.1 Swarm Intelligence for Solving the Search Problem

Fluoroscopic analysis of knee joint kinematics consists of determining the position and orientation of the bones in the knee joint. In the experiments conducted in this thesis, a hypothetical fluoroscope with a CT cube placed between an X-ray camera and an X-ray film, is used. The objective of the thesis is to find the position of the CT cube and azimuth and elevation angles of the camera and the rotation of the film for a given CT scan image and a target fluoroscopy X-ray frame.

The focus of this thesis has been to develop faster search algorithms for finding the position and orientation information. This is done by implementing the two swarm intelligence techniques, the bees algorithm and the particle swarm optimization method, for sequential and parallel execution. Both these methods are biologically inspired algorithms. While the bees algorithm mimics the food foraging behavior of the bees, particle swarm optimization method is inspired by the social behavior of a flock of birds or a school of fish. By adopting the communication and collaboration among a population of entities to solve a common problem, it is easier to reach a better solution faster.

Parallel computing is a suitable method for solving problems that are enormously time consuming. Fluoroscopic analysis is one such problem and the usage of parallel computing for solving this is well warranted. The bees algorithm and the particle swarm optimization
method are easily parallelizable and are easy to apply to a problem of this nature. The experiments conducted to test the implementation provide sufficient information to indicate that parallel computing can significantly decrease the time required to search the position and orientation values in the fluoroscopic process. Several configurations of the bees and the particle swarm optimization algorithms have been successfully executed.

### 6.2 Future Scope

There is scope for further research in several components of the fluoroscopic analysis process. In order to derive accurate results, it is essential to have very efficient and effective methods for each of the analysis operations such as the DRR image generation, filtering of essential information from DRR or X-ray images, matching two DRR or X-ray images as well as searching for the position and orientation values from a given CT image and a target X-ray image. While the bees and particle swarm optimization algorithms appear to be promising solutions to the search problem, more analysis has to be carried out to find the best configuration to get optimum results out of these two methods, and this would be possible with accurate results from the other steps in the fluoroscopic process.

The intent of this fluoroscopic analysis process is to provide dynamic real-time information of the knee joint to assist in several diagnostic or surgical processes. This research can be extended to include Magnetic Resonance Imaging (MRI) process to find alternate solutions. Additionally, the fluoroscopic analysis process can be applied to any joint in the human body to help diagnose and treat ailments of the joints. There is a huge potential for this process to truly help the medical community in the treatment of joint-related problems.
BIBLIOGRAPHY


APPENDIX A

README FILE

Readme file
Fluoroscopy Project

Sections:
1. Description
2. Steps Involved in Executing the Project.
3. How to Compile the Project.
4. Makefile Features.
5. How to Run the Project.
7. How to Add a Plugin.
10. Scripts.
11. Config Files.
12. Results.
13. Examples for Compiling and Executing the Project.

1. Description

This project performs an unsupervised fluoroscopic analysis of knee-joint kinematics. The operations involved include
1. reading a CT scan image,
2. generating DRR images,
3. filtering the edges from a DRR image,
4. matching the edges of two different DRR images
   and having a score associated with it,
5. and searching for the right DRR image for a CT
   image, by using different combinations of the
   film and camera orientation and CT cube offsets.

--------------------------------------------------------
2. Steps Involved in Executing the Project
--------------------------------------------------------

The main program of this project relies on the
availability of shared libraries for the following
operations: generating DRR, filtering, matching and
searching. Below are a list of steps required to execute
this project. Refer to relevant sections for details in
each step. In order to use the Makefile provided as is,
please follow the directory structure and the other
requirements as mentioned for each step. Deviation from
them would require custom makefiles or modification to
paths in the makefile or whatever is required for the
specific change.

1. Place the CT image file in the 'images' directory.
2. Place a DRR image for that CT image in the 'images'
directory. The DRR image file for a CT image,
can also be created by running the program
create_target_drr with the required camera and
film orientation and cube offsets. Refer to section
3.A and 5.A.
3. Create an input config file which lists all the
   parameters required for running the program, such
   as number of iterations to run the search, path
to the CT image, path to the DRR image, and any
   other parameters required for the algorithm chosen.
   Place this file in the 'config' directory. Refer
to section 3.E
4. Compile all the plugins required for the instance
   of the program. Each plugin should be in a
   separate directory in the 'plugins' directory.
   Place libraries (*.so files) in the lib directory
   and turn off the selinux security restrictions
   for those libraries. Refer to section 3.B and 3.C.
5. Compile the main program by specifying the compiler. Refer to section 3.D.
6. Run the program by providing the input config file. Refer to section 5.B.

--------------------------------------------------------
3. How to Compile the Project
--------------------------------------------------------

$(FLUOROSCOPY) represents the path to the fluoroscopy project source, the outermost directory of the trunk, which consists of main makefile and all other directories.

A. To compile the create_drr_image program:
   cd /$(FLUOROSCOPY)
   make clean_create_drr
   make create_drr

B. To compile all the plugins in the 'plugins' directory:
   cd /$(FLUOROSCOPY)
   make clean_plugins
   make plugins

   Note: This creates a .so file for sequential and parallel, for DOF=3 and 6, and all the permutations required.
   DOF = Degrees of Freedom

C. To compile a single plugin:
   For any plugin, you can run 'make clean' followed by 'make all' to make all libraries that are possible for that plugin. To only make a particular library, run make with required parameters.

D. To compile the main program:
   cd /$(FLUOROSCOPY)
   make clean
   make main CC=<compiler>

   NOTE: Compiler should be 'gcc' or 'mpicc'.

E. To create input config files:
Run the relevant scripts from the 'scripts'
directory or run the makefile with the right
target to create the config files.

make configure_bees #For Bees
make configure_pso #For PSO
make configure_montecarlo #For MonteCarlo

F. To create lib config file[obsolete]:
make configure_libs

NOTE: This feature is preserved for any future
use that might necessitate the need for a lib
config file.

--------------------------------------------------------
4. Makefile Features
--------------------------------------------------------
The Makefile present in the main directory of the
project trunk, facilitates many operations. Below is a
list of all the targets accepted by the makefile.

make clean_all
  executes clean, clean_create_drr, clean_libs,
  clean_plugins, and clean_swapfiles
make clean
  cleans the current directory and source directory
make clean_create_drr
  cleans files associated with create_target_drr
  program from the source directory
make clean_libs
  deletes all the libraries (*.so) files from the
  'lib' directory
make clean_plugins
  cleans all the plugin directories
make clean_swapfiles
  cleans all the swap files (*) from all the
directories in the project recursively.
make create_drr
compiles the create-target_drr program
make plugins
    compiles all the plugins in the 'plugins'
directory and copies all the libs to the lib
folder
make main CC=<compiler>
    compiles the main program. CC= <gcc|mpicc>
make help
    prints all the targets accepted by make
make configure_bees
    runs a script that creates an input config file
for the Bees algorithm by taking user inputs and
copies it to the config folder
make configure_pso
    runs a script that creates an input config file
for the PSO algorithm by taking user inputs and
copies it to the config folder
make configure_montecarlo
    runs a script that creates an input config file
for the MonteCarlo algorithm by taking user
inputs and copies it to the config folder
make configure_libs
    runs a script that creates an lib config file
and copies it to the config folder
make documentation
    compiles and creates the code documentation
and copies them to the doc folder.

--------------------------------------------------------

Makefile for the main program:
The makefile for the main program is named Makefile_main and is
present in the 'source' directory.

Makefile for the create_target_drr program:
The makefile for the create_target_drr program is named
Makefile_drr and is present in the 'source' directory.

Makefiles for plugins:
Each plugin has a directory associated with it in the 'plugins'
directory. Each plugin has its own makefile named 'Makefile'.
5. How to Run the project

A. To run the create_target_drr program:
create_target_drr
<path to generate_drr plugin(.so lib file)>
<CT image file>
<resolution>
<path to output drr file>
[<azimuth> <elevation> <rotation>]
[<x offset> <y offset> <z offset>]]

NOTE: Example for resolution is 128, 256, 512 etc.
Resolution for really_small_cube.txt is 128.

B. To run the main program
Sequential:
fluoro_s
<path to search plugin(.so lib file)>
<path to the input config file>

Parallel:
mpiexec -l -n <num of processes> fluoro_p
<path to search plugin(.so lib file)>
<path to the input config file>

6. Directory Structure of the Project

$(FLUOROSCOPY) represents the main truck directory of the project.

Create an environment variable for this as follows:
export FLUOROSCOPY=<fluoroscopy trunk absolute path>

$(FLUOROSCOPY)
Place the files in the relevant directories. The directory names are self-explanatory and clearly indicate what kind of files are expected to be present within them.

NOTE: The Makefile in the trunk directory does not compile the source in 'matlab' directory. 'matlab' directory contains the source for fluoroscopy in matlab language. All the other folders and files are associated with the C implementation for fluoroscopy.
7. How to Add a Plugin

Following are the steps to create a new plugin:

1. Create a folder in the 'plugins' directory. Follow the naming conventions provided in section 9.
2. Place all the source files in the created directory.
3. If the plugin depends on other libraries or other set of source files, or different sort techniques etc, make separate directories for each of them within the directory for the plugin. For example, generate_drr_raytracing has 3 directories for the sort function. One of them is used to compile the library.
4. Create a makefile for the plugin.
5. In the makefile, create a target 'all' which generates multiple libraries files for all the required combinations. Refer to other plugin makefiles as examples.
6. Create an input config file, as required for your libraries and execute the main program with the appropriate parameters.

8. Code Documentation

The open source tool Doxygen is used for generating the code reference manual. The configuration file Doxyfile is present in the main truck directory. Currently, it is configured to provide html, latex-pdf and perlmod-pdf documentations. Edit this file to customize the documentation to your needs. Please refer to the Doxygen tutorial for details on installing, configuring and running the tool. Some basic steps are:

To generate documentation, run the commands:
   make documentation

To view the documentation, run any of the following
commands:
  firefox docs/index.html
  acroread docs/CodeDocumentation.pdf

Alternate way to generate documentation:
doxygen Doxyfile

Alternate way to view documentation:
cd html; firefox index.html
  cd latex; make; acroread refman.pdf
  cd perlmod; make; acroread doxylatex.pdf

Currently, the documentation includes all the plugins. If desired, separate documentation can be created for each plugin, by providing a Doxygen config file for each plugin and following the steps similar to the above instructions.

--------------------------------------------------------

9. Naming Conventions
--------------------------------------------------------

The names chosen for files or directories should be self-explanatory and easy to understand.

The names of the plugin directories should have a first name that indicates the operation performed or the interface implemented, e.g., filter, match, search etc. The second name should indicate if it is parallel or sequential if applicable. The next name should indicate the algorithm used, e.g., sobel, contour, bees etc. Any other variations should be next, e.g., bw for black and white, or gs for grayscale, etc. Separate the names by underscores.

The lib names should begin with 'lib', followed by 's' or 'p' to indicate if they are sequential or parallel. After this, separate each name with an underscore and follow a pattern as mentioned above.

--------------------------------------------------------
10. Scripts

------------------------------------------------------------------

Below is the list of existing script files and its functions:

configure_libs.sh
   Used to create a lib config file [obsolete]
configure_input_bees.sh
   Used to create an input config file for Bees algorithm
configure_input_montecarlo.sh
   Used to create an input config file for Monte Carlo algorithm
configure_input_pso.sh
   Used to create an input config file for PSO algorithm
run_s_sanity.sh
   Used to run all the sequential programs for sanity checking. Basically to check if the program does not crash for the minimum number of iterations.
run_s_limited.sh
   Used to run all sequential programs for testing the limited range for search parameters
run_s_full.sh
   Used to run all sequential programs for testing the full range for search parameters
run_p_sanity.pbs
   Used to run all the parallel programs for sanity checking. Basically to check if the program does not crash for the minimum number of iterations.
run_p_limited.pbs
   Used to run all parallel programs for testing the limited range for search parameters
run_p_full.pbs
   Used to run all parallel programs for testing the full range for search parameters
run_s.sh
   Used by the run_s_* scripts internally.
run_p.sh
   Used by the run_p_* scripts internally.
helper_s.sh
    Used by the run_s.sh script internally.

helper_p.sh
    Used by the run_p.sh script internally.

NOTE: Check to see if any paths need to be changed in the scripts. In some places, the path is set to the current user’s working directory.

11. Config Files

Input configuration files can be created using the steps specified in 3E. The main program requires a config file as an input. It consists of all the algorithm parameters and also the target DRR image. In this project, a naming convention is used to name the config files for ease of use.

$(FLUOROSCOPY)/config directory consists of all the config files. The config files required for sequential tests are placed in 's_tests' folder, and files required for parallel tests are placed in 'p_tests' folder. Within these two folders, there can be many folders with separate config files, one for each test case.

Note: Naming convention for config and result directories:

Directory used for tests to just check if the programs run without error:
Name = sanity

Directory used for tests that limit the search space:
Underscore delimited name contains:
1. "limited"
2. num of processors
3. [num of neighbors]-[num of neighborhoods] (for PSO)
   Use the second number for [num of best sites] (for Bees)
5. num_bee_iter (for Bees) or num_particle_iter (for PSO)
6. drr_image: "img" followed by az, el, rt orientation
   and x, y, z translation values

Directory used for tests that search the entire search space also follow the above naming convention except that the name starts with "full" instead of "limited".

--------------------------------------------------------
12. Results
--------------------------------------------------------

$(FLUOROSCOPY)/results directory is created to contain all the output redirected from test executions. The result files required for sequential tests are placed in 's_tests' folder, and files required for parallel tests are placed in 'p_tests' folder. Within these two folders, there can be many folders, one for each test case.

The same naming convention used for naming config files may also be used for result files for easier maintenance and management of tests.

--------------------------------------------------------
13. Examples for Compiling and Executing the Project
--------------------------------------------------------

A. Steps for the sequential program:
1. cd $(FLUOROSCOPY)
2. make clean_all
3. make plugins
4. make create_drr
5. cd $(FLUOROSCOPY)/source
6. ./create_target_drr
   $(FLUOROSCOPY)/lib/libs_generate_drr_raytracing.so
   $(FLUOROSCOPY)/images/really_small_cube.txt
   128
   $(FLUOROSCOPY)/images/drr_image.txt
   60 30 45 8 17 22
7. cd $(FLUOROSCOPY)
8. make clean_create_drr
9. make main CC=gcc
10. fluoro_s $(FLUOROSCOPY)/lib/libs_search_bees_6d.so $(FLUOROSCOPY)/config/<config file>

B. Steps for the parallel program:
1. make clean_all
2. make plugins
3. make create_drr
4. cd $(FLUOROSCOPY)/source
5. ./create_target_drr
   $(FLUOROSCOPY)/lib/libp_generate_drr_raytracing.so $(FLUOROSCOPY)/images/really_small_cube.txt 128
   $(FLUOROSCOPY)/images/drr_image.txt 60 30 45 8 17 22
6. cd $(FLUOROSCOPY)
7. make clean_create_drr
8. make main CC=mpicc
9. fluoro_p $(FLUOROSCOPY)/lib/libp_search_bees_6d.so $(FLUOROSCOPY)/config/<config file>

Note: The main differences between sequential and parallel executions are in the .so library names and the executable names.

----------------------------------------
APPENDIX B

CODE DOCUMENTATION

This appendix chapter consists of the source code documentation for the essential components of the code. This documentation is extracted from the code with the help of a tool called Doxygen.

B.1 Data Structure Documentation

The following sections from Section B.2 to B.9 provide the main data structures used in the implementation of this thesis. Sections B.2 to B.6 are used in the fluoroscopic analysis process, Section B.7 is used in the configuration file reader program, and Section B.8 and B.9 are used in the SI search algorithms namely Bees algorithm and PSO method respectively.
B.2 _ctVolume Struct Reference

B.2.1 Detailed Description

This struct represents a 3-D Computer Tomography volume information used in the experimental fluoroscope.

Data Fields

- double * volume
- int size
- double width
- double height
- double length
- float cam_to_table
- double cube_adjust [3]
- double cube_offset [3]
- int origin [3]
- double far_corner [3]
- double dimension [3]

B.2.2 Field Documentation

B.2.2.1 float _ctVolume::cam_to_table

Distance between the camera and the fluoroscope table
B.2.2.2 double ctVolume::cube_adjust[3]
CT cube adjustment buffer

B.2.2.3 double ctVolume::cube_offset[3]
CT cube offset (x,y,z translations from the origin)

B.2.2.4 double ctVolume::dimension[3]
CT volume dimensions

B.2.2.5 double ctVolume::far_corner[3]
Coordinates of far corner of the CT cube

B.2.2.6 double ctVolume::height
CT volume height in cm

B.2.2.7 double ctVolume::length
CT volume length in cm

B.2.2.8 int ctVolume::origin[3]
CT cube origin coordinates(x,y,z), in pixels

B.2.2.9 int ctVolume::size
number of elements in volume
B.2.2.10  double* _ctVolume::volume

3-D CT volume data

B.2.2.11  double _ctVolume::width

CT volume width in cm

The documentation for this struct was generated from the following file:

- fluoro.h
B.3  _drrImage Struct Reference

B.3.1  Detailed Description

This struct represents the DRR image.

Data Fields

- int n
- int m
- double ** image

B.3.2  Field Documentation

B.3.2.1  double** _drrImage::image

2-D image

B.3.2.2  int _drrImage::m

Number of rows

B.3.2.3  int _drrImage::n

Number of columns

The documentation for this struct was generated from the following file:

- fluoro.h
B.4  _film Struct Reference

B.4.1  Detailed Description

This struct represents the film in the experimental fluoroscope.

Data Fields

- double size
- double resolution
- double dist
- double table_to_film
- double pixel_size
- double offset [3]
- double center [3]
- double rotation
- double azimuth
- double elevation
- double distance

B.4.2  Field Documentation

B.4.2.1  double _film::azimuth

Film azimuth
B.4.2.2  double_film::center[3]

B.4.2.3  double_film::dist

Distance

B.4.2.4  double_film::distance

Distance

B.4.2.5  double_film::elevation

Film elevation

B.4.2.6  double_film::offset[3]

Film offset (x,y,z)

B.4.2.7  double_film::pixel_size

Pixel size

B.4.2.8  double_film::resolution

Pixel resolutions

B.4.2.9  double_film::rotation

Film rotation

B.4.2.10 double_film::size

Size of the film
B.4.2.11  double *film::*table_to_film

Distance between the film and the fluoroscopic table

The documentation for this struct was generated from the following file:

- fluoro.h
B.5 _camera Struct Reference

B.5.1 Detailed Description

This struct represents the X-ray camera in the experimental fluoroscope.

Data Fields

- int _camera::camera_to_table
- double _camera::azimuth
- double _camera::elevation
- double _camera::dist
- double _camera::offset [3]
- double _camera::coord [3]

B.5.2 Field Documentation

B.5.2.1 double _camera::azimuth

Camera azimuth

B.5.2.2 int _camera::camera_to_table

Distance between the camera and the fluoroscopic table

B.5.2.3 double _camera::coord[3]

Camera coordinates
B.5.2.4  double _camera::dist

Camera distance

B.5.2.5  double _camera::elevation

Camera elevation

B.5.2.6  double _camera::offset[3]

Camera offset

The documentation for this struct was generated from the following file:

- fluoro.h
B.6 _edges Struct Reference

B.6.1 Detailed Description

This struct represents the filtered DRR image that consists of edges of the image.

Data Fields

- int n
- int m
- double ** image

B.6.2 Field Documentation

B.6.2.1 double** _edges::image

2-D image

B.6.2.2 int _edges::m

Number of rows

B.6.2.3 int _edges::n

Number of columns

The documentation for this struct was generated from the following file:

- fluoro.h
B.7 dict Struct Reference

B.7.1 Detailed Description

This struct defines a dictionary to hold the key-value pairs of a configuration item present in a config file.

Data Fields

- char key [MAX_VALUE_LEN]
- char value [MAX_VALUE_LEN]

B.7.2 Field Documentation

B.7.2.1 char dict::key[MAX_VALUE_LEN]

Configuration item key

B.7.2.2 char dict::value[MAX_VALUE_LEN]

Configuration item value

The documentation for this struct was generated from the following file:

- config_reader.h
B.8 bee Struct Reference

B.8.1 Detailed Description

This data structure represent a Bee in the Bees Algorithm. Each bee that is involved in the search for the right drr image, is associated with the values for orientation of camera and film, offset of the cube and score of the resulting DRR image, at a given time.

Data Fields

- double orientation [3]
- double offset [3]
- double score
- int finesearch
- int rank

B.8.2 Field Documentation

B.8.2.1 double bee::orientation

Represents the orientation of the camera (az and el) and the film (rt)

B.8.2.2 int bee::finesearch

Indicates whether the bee will next search around the previous values or in random. Value is either 1 or 0

B.8.2.3 double bee::offset

Represents the cube offset values
B.8.2.4  int _bee::rank

Represents the ranking of the bee based on the score value

B.8.2.5  double _bee::score

Represents the score of the resulting DRR image

The documentation for this struct was generated from the following files:

- search_parallel_bees.c
- search_sequential_bees.c
B.9 .particle Struct Reference

B.9.1 Detailed Description

This data structure represent a particle in the Particle Swarm Optimization method. Each particle that is involved in the search for the right DRR image, is associated with the values for orientation of camera and film, offset of the cube and score of the resulting DRR image, at a given time.

Data Fields

- double orientation [3]
- double offset [3]
- double score

B.9.2 Field Documentation

B.9.2.1 double .particle::orientation

Represents the orientation of the camera(az and el) and film(rt)

B.9.2.2 double .particle::offset

Represents the cube offset values

B.9.2.3 double .particle::score

Represents the score of the resulting DRR image

The documentation for this struct was generated from the following files:

- search_parallel_pso.c
• search_sequential_pso.c
B.10 Plugin Interface Documentation

The following sections from Section B.11 to B.13 provide the details of the header files used as interfaces to the plugins to the fluoroscopic project. Implementation of fluoroscopic analysis requires algorithms to be implemented using these interfaces. The algorithm implementation will define all the functions declared in the corresponding header file. This makes it possible for any algorithm that implements this interface to fit into main fluoroscopic analysis program. Provisions have been made to specify which algorithm to use during run time of the main program.
B.11 generate_drr.h File Reference

B.11.1 Detailed Description

This header file provides an interface for implementing an algorithm for generating Digitally Reconstructed Radiograph images from a given CT image.

Typedefs

- typedef void * CT_Volume_Ptr
- typedef void * Drr_Image

Functions

- void generate_drr (double camera_azimuth, double camera_elevation, double film_rotation, double cube_offset[], CT_Volume_Ptr cube, Drr_Image out_drr_image)

B.11.2 Typedef Documentation

B.11.2.1 typedef void * CT_Volume_Ptr

This typedef is used to define a CT volume.

B.11.2.2 typedef void * Drr_Image

This typedef is used to define a DRR image.
B.11.3 Function Documentation

B.11.3.1 void generate_drr (double azimuth, double elevation, double rotation,
        double offset[], void *cube, void *drr_image)

This function creates a DRR model.

Parameters:

    azimuth  Azimuth angle of the camera
    elevation Elevation angle of the camera
    rotation  Rotation angle of the film
    offset   x,y,z offsets of the CT cube
    cube     3-D CT volume data
    drr_image Pointer to the generated output DRR image [output parameter]

Returns:

    Void
B.12 filter.h File Reference

B.12.1 Detailed Description

This header file provides an interface for implementing an algorithm for filtering X-ray images to extract required information from an image.

Typedefs

- typedef void * Filter_Image

Functions

- Filter_Image filter (Filter_Image image)

B.12.2 Typedef Documentation

B.12.2.1 typedef void * Filter_Image

This typedef is used to define an image.

B.12.3 Function Documentation

B.12.3.1 Filter_Image filter (void * image)

Filters the input image and provides an image with only the requires features of the input image.

Parameters:

- image Image to be filtered

Returns:

Filtered image
B.13 match.h File Reference

B.13.1 Detailed Description

This header file provides an interface for implementing an algorithm for matching or comparing two images.

Typedefs

- typedef void * Match_Image

Functions

- double match (Match_Image ref_image, Match_Image target_image)

B.13.2 Typedef Documentation

B.13.2.1 typedef void * Match_Image

This typedef is used to define an image.

B.13.3 Function Documentation

B.13.3.1 double match (void * ref_image, void * target_image)

Computes the score of an image. This routine is intended to be used after an image has been filtered using a sobel operator for black and white images.

Parameters:

ref_image Reference Contour image

target_image Target Contour image
Returns:

Score value between 0 and 1
B.14 search.h File Reference

B.14.1 Detailed Description

This header file provides an interface for implementing a search algorithm for searching the position and orientation information with a given CT image and configuration information required for the algorithm.

Typedefs

- typedef void * Search_CT_Volume_Ptr

Functions

- void search (Search_CT_Volume_Ptr cube, FILE *search_config_filename)

B.14.2 Typedef Documentation

B.14.2.1 typedef void * Search_CT_Volume_Ptr

B.14.3 Function Documentation

B.14.3.1 void search (void * _cube, FILE * config_file)

This function attempts to search for an fluoroscopy image. The search algorithm is based on a Bees algorithm, where random camera positions are tried until a close match is found.

Parameters:

_cube  Pointer to the CT_VOLUME

config_file  Pointer to the open file containing all the configuration values required for this algorithm
Returns:

Void
B.15 Source File Documentation

The following sections from Section B.16 to B.31 provide the details of the source files included in this thesis.

Section B.16 contains information about the program that is used to create a DRR image from a given CT image along with position and orientation data. This program is kept separate from the rest of the fluoroscopic analysis code, so that, required DRR images can be created beforehand and used as input to the main fluoroscopy program. Images once created and stored, need not be recreated during each program execution.

Section B.17 contains information about the program which is the main entry point to the entire fluoroscopic analysis process. Sections B.21 to B.31 provide information about the plugins that implement the interfaces defined in Sections B.11 to B.13.
B.16 create_target_drr.c File Reference

B.16.1 Detailed Description

This program is used to generate a DRR image from the specified orientation and offset configuration inputs for a given CT image. USAGE: <ProgramName> <PathToGenerate-DrrPlugin> <PathToCTImageFile> <Resolution> <PathToOutputFile> [ <Azimuth> <Elevation> <Rotation> [ <X_Offset> <Y_Offset> <Z_Offset> ] ]

Functions

- static void print_usage (const char ∗appname)
- int main (int argc, char ∗argv[])

Variables

- void ∗handle
- void(∗) generate_drr_function (double azimuth, double elevation, double rotation, double offset[], void ∗cube, void ∗drr_image)
- const char ∗dlError

B.16.2 Function Documentation

B.16.2.1 int main (int argc, char ∗argv[])

This is the main function for creating DRR images from a given CT image. It is responsible for creating a drr image file from the specified orientation and position offset configuration inputs for a given CT image.
Parameters:

\textit{argc}  Number of command line parameters

\textit{argv}  Array of command line parameters

Returns:

0 on success, 1 on failure

B.16.2.2 \texttt{static void print\_usage (const char * appname)} [static]

Displays the command line parameters required for this program

Parameters:

\textit{appname}  Name of the program

Returns:

Void

B.16.3 Variable Documentation

B.16.3.1 \texttt{const char* dlError}

B.16.3.2 \texttt{void(*) generate\_drr\_function(double azimuth, double elevation, double rotation, double offset[], void *\_cube, void *\_drr\_image)}

B.16.3.3 \texttt{void* handle}
B.17 main.c File Reference

B.17.1 Detailed Description

This is the main entry point for the fluoroscopy program. This program is dependent on shared libraries for filtering and searching DRR images. USAGE: `<ProgramName> <PathToSearchPlugin> <PathToInputConfigFile>`.

Functions

- static void print_usage_exit (const char *appname)
- int main (int argc, char *argv[])

Variables

- void * search_handle
- void(* search_function (void *, FILE *)
- const char * dlError

B.17.2 Function Documentation

B.17.2.1 int main (int argc, char * argv[])

This is the main function for the fluoroscopy program.

- The input configuration file is parsed to extract the CT image file and the DRR image file.
- CT volume and DRR image are constructed from these two files.
- Following this the edges of the drr image are extracted.
• Using this as a target, a search for the orientation and the offset of the CT cube is begun.

• This function relies on shared libraries for filtering and searching which have to be provided while compiling this program.

Parameters:

\begin{itemize}
  \item \texttt{argc} Number of command line parameters
  \item \texttt{argv} Array of command line parameters
\end{itemize}

Returns:

0 on success, 1 on failure

\textbf{B.17.2.2} \texttt{static void print\_usage\_exit (const char \ast \ appname)} \hspace{1em} [\texttt{static}]

Displays the command line parameters required for this program

Parameters:

\begin{itemize}
  \item \texttt{appname} Name of the program
\end{itemize}

Returns:

Void

\textbf{B.17.3} Variable Documentation

\textbf{B.17.3.1} \texttt{const char\ast \ dlError}

\textbf{B.17.3.2} \texttt{void\ast \ search\_function(void \ast, FILE \ast)}

\textbf{B.17.3.3} \texttt{void\ast \ search\_handle}
B.18 write_image.c File Reference

B.18.1 Detailed Description

This file contains routines for writing matrices representing a CT cube and a DRR image to a text file.

Functions

- `void write_image (double **image, int m, int n, const char *filename)`
- `void write_array (double *array, int n, const char *filename)`
- `void write_drr (DRR_IMAGE *drr_image, const char *filename)`

B.18.2 Function Documentation

B.18.2.1 `void write_array (double * array, int n, const char * filename)`

Writes an array to a text file. Each value is delimited by a space

Parameters:

- `array` Array of doubles
- `n` Number of elements
- `filename` File name to create and write into

Returns:

Void
B.18.2.2  void write_drr (DRR_IMAGE * drr_image, const char * filename)

Writes a DRR image to a text file. First two lines contain the number of rows and number of columns respectively. The image values are inserted from the third line. Each value is delimited by a space.

Parameters:

   drr_image  DRR image

   filename  File name to create and write into

Returns:

   Void

B.18.2.3  void write_image (double ** image, int m, int n, const char * filename)

Writes an image array to a text file. Each value is delimited by a space

Parameters:

   image  Image array (output parameter)

   m  Number of rows

   n  Number of columns

   filename  File name to create and write into

Returns:

   Void
B.19  util.c File Reference

B.19.1 Detailed Description

This file contains general application utility functions.

Functions

- static void set_volume_defaults (CT_VOLUME *volume)
- FILM * create_film ()
- void free_film (FILM *film)
- CAMERA * create_camera ()
- void free_camera (CAMERA *cam)
- DRR_IMAGE * alloc_drr (int m, int n)
- DRR_IMAGE * create_drr (const char *filename)
- void free_drr (DRR_IMAGE *drr)
- CT_VOLUME * create_volume (const char *filename)
- void free_volume (CT_VOLUME *volume)
- double *** alloc_dcube (int m, int n, int p)
- void free_dcube (double ***cube)
- double ** alloc_dmat (int m, int n)
- void free_dmat (double **dmat)
- void * xmalloc (size_t bytes)
- double xrandom ()
- double yrandom ()
- int zrandom (double limit)
• double d_abs (double val)

B.19.2 Function Documentation

B.19.2.1 double *** alloc_dcube (int m, int n, int p)

 Allocates memory for a 3 dimensional array of doubles as a contiguous block of memory.

Parameters:

\[ m \] Number of rows
\[ n \] Number of columns
\[ p \] Number of planes

Returns:

A 3-D array of doubles

B.19.2.2 double ** alloc_dmat (int m, int n)

 Allocates memory for a 2 dimensional array of doubles as a contiguous block of memory.

ARGS: Number of rows Number of columns

Returns:

Data array(double)

B.19.2.3 DRR_IMAGE * alloc_drr (int m, int n)

 Allocates memory for a DRR image.

Parameters:

\[ m \] Number of rows
\[ n \] Number of columns

\textbf{Returns:}

Pointer to a DRR\_IMAGE object

\textbf{B.19.2.4 \texttt{CAMERA} \textbullet{} \texttt{create\_camera}()}\n
Creates a new \texttt{CAMERA} object.

\textbf{Returns:}

Pointer to a new \texttt{CAMERA} object

\textbf{B.19.2.5 \texttt{DRR\_IMAGE} \textbullet{} \texttt{create\_drr} (const char \texttt{*filename})} \n
Creates an DRR image initialized by the data contained in the specified file.

\textbf{Parameters:}

\textit{filename} Name of the file containing the DRR\_IMAGE data

\textbf{Returns:}

Pointer to a DRR\_IMAGE object

\textbf{B.19.2.6 \texttt{FILM} \textbullet{} \texttt{create\_film}()}\n
Creates a \texttt{FILM} object.

\textbf{Returns:}

Pointer to a new \texttt{FILM} object
B.19.2.7  CT_VOLUME * create_volume (const char *filename)

Creates an volume object initialized by the data contained in the specified file.

Parameters:

filename  Name of the file containing the CT volume data

Returns:

Pointer to a CT_VOLUME object

B.19.2.8  double d_abs (double val)

Returns the absolute value.

Parameters:

val  Input value

Returns:

Absolute value

B.19.2.9  void free_camera (CAMERA * cam)

Frees up the CAMERA object.

Parameters:

cam  Pointer to a CAMERA object

Returns:

Void
B.19.2.10  void free_dcube (double *** cube)

Frees a dcube object.

Parameters:

    cube  Pointer to a dcube

Returns:

    Void

B.19.2.11  void free_dmat (double ** dmat)

Frees a dmat object.

Parameters:

    dmat  Pointer to a dmat object to free

Returns:

    Void

B.19.2.12  void free_drr (DRR_IMAGE * drr)

Frees memory allocated for a DRR_IMAGE object.

Parameters:

    drr  Pointer to a DRR_IMAGE

Returns:

    Void
B.19.2.13  void free_film (FILM *film)

Frees a FILM object.

Parameters:

  *film  Pointer to a FILM object

Returns:

void

B.19.2.14  void free_volume (CT_VOLUME *volume)

Frees memory for a CT_VOLUME object.

Parameters:

  *volume  Pointer to a CT_VOLUME object

Returns:

Void

B.19.2.15  static void set_volume_defaults (CT_VOLUME *volume)  [static]

Initializes the default values for the volume object.

Parameters:

  *volume  Pointer to the CT volume

Returns:

void
B.19.2.16  void * xmalloc (size_t bytes)

Allocates memory. If memory allocation fails, the application will abort.
Parameters:

    bytes  Number of bytes to allocate

Returns:

    Pointer to a chunk of memory

B.19.2.17  double xrandom ()

Generates a random number (double) between 0 and 1. It is always positive
Returns:

    Random number (double)

B.19.2.18  double yrandom ()

Generates a random number (double) between 0 and 1. It can be a positive or a negative
number. The positiveness or negativeness is also generated randomly.
Returns:

    Random number (double)

B.19.2.19  int zrandom (double limit)

Generates a random number (int) that lies within the specified limit. It can be a positive or
a negative number. The positiveness or negativeness is also generated randomly.
Returns:

    Random number (int)
B.20 config_reader.c File Reference

B.20.1 Detailed Description

This file contains functions to extract the configuration information from a file consisting of key-value pairs in a suitable format. Example for a configuration item: `MAX_ITER=500`, where MAX_ITER is the key and 500 is the value.

Functions

- void read_config (FILE *config)
- void print_dict ()
- int get_config_value (char *key, char **value)

B.20.2 Function Documentation

B.20.2.1 void get_config_value (char * key, char ** value)

Searches for the given key in the dictionary of configuration items and returns 1 or 0 to indicate the success of the search. It also returns the value through the output parameter.

Parameters:

- key String that denotes the key to be searched for in the configuration.
- value This will hold the value associated with the key as a result of this operation.

Returns:

1 if key-value pair is found, 0 otherwise.
B.20.2.2  void print_dict ()

Prints the configuration dictionary contents in the form of key-value pairs.

Returns:

void

B.20.2.3  void read_config (FILE * config)

Reads the configuration file and constructs a dictionary of key-value pairs for each configuration item.

Parameters:

config  FilePointer to the configuration file

Returns:

void
B.21 generate_drr.c File Reference

B.21.1 Detailed Description

This file implements the interface generate_drr.h (p. 91). This file contains utility routines for creating Digitally Reconstructed Radiograph (DRR) images. It contains the routine for creating a DRR model which includes setting up the point on a film array and uses ray tracing to create a DRR image for the given orientation and offset values.

Functions

- static void sph2cart (double theta, double phi, double rho, double cart[])
- void generate_drr_helper (CAMERA *cam, CT_VOLUME *cube, double ***film_array, int m, int n, int p, DRR_IMAGE *drr_image)
- void generate_drr (double azimuth, double elevation, double rotation, double offset[], void *cube, void *drr_image)

B.21.2 Function Documentation

B.21.2.1 void generate_drr (double azimuth, double elevation, double rotation, double offset[], void *cube, void *drr_image)

This function creates a DRR model.

Parameters:

- azimuth Azimuth angle of the camera
- elevation Elevation angle of the camera
- rotation Rotation angle of the film
- offset x,y,z offsets of the CT cube
**.cube** 3-D CT volume data

**.drr_image** Pointer to the generated output DRR image [output parameter]

**Returns:**

Void

**B.21.2.2 void generate_drr_helper (CAMERA * cam, CT_VOLUME * cube, double *** film_array, int m, int n, int p, DRR_IMAGE * drr_image)**

This is a helper function that calls the ray tracing routine to create a DRR image from the given information.

**Parameters:**

- **cam** Pointer to camera orientation
- **cube** Pointer to a CT volume
- **film_array** Pointer to a 3D array containing points on a film
- **m** Number of rows
- **n** Number of columns
- **p** Depth of the array
- **drr_image** Pointer to the DRR image [output parameter]

**Returns:**

Void

**B.21.2.3 static void sph2cart (double theta, double phi, double rho, double cart[ ])**

[static]

This function converts spherical coordinates to Cartesian coordinates.
Parameters:

\( \theta \)  Value of theta

\( \phi \)  Value of phi

\( \rho \)  Value of rho

\( \text{cart} \)  Array for 3-D Cartesian coordinates [output parameter]

Returns:

Void
B.22 ray_tracing.c File Reference

B.22.1 Detailed Description

This file implements Siddon’s ray tracing algorithm”. Ray tracing involves constructing a DRR from a CT image by calculating the radiological path through a three-dimensional CT array.

Functions

- **void trace_CT** (double *cam_pt, double *film_array, int *film_dims, double *CT_image, double *N_vox, double *CT_size, int *CT_orig, double *DRR_out-im)

Variables

- **const int * film_dims**

B.22.2 Function Documentation

B.22.2.1 **void trace_CT** (double *cam_pt, double *film_array, int *film_dims, double *CT_image, double *N_vox, double *CT_size, int *CT_orig, double *DRR_out-im)

B.22.3 Variable Documentation

B.22.3.1 **const int * film_dims**
B.23 filter_sobel_bw.c File Reference

B.23.1 Detailed Description

This file implements the interface “filter.h”. It is the plugin for filtering using Sobel algorithm for black and white values.

Functions

- static int ** conv2d (double **image, int num_rows, int num_cols, const int filter_kernal[][3], int fk_row, int fk_col)
- void * filter (void *image)

Variables

- static const int x_filt ][3]
- static const int y_filt ][3]
- static const int d1_filt ][3]
- static const int d2_filt ][3]

B.23.2 Function Documentation

B.23.2.1 static int ** conv2d (double **image, int num_rows, int num_cols, const int filter_kernal[][3], int fk_row, int fk_col) [static]

Function for two dimensional convolution: \( g(x,y) = \text{SUM}\{ \text{SUM}\{ w(s,t)f(x+s,y+t), t=-b..b \}, s=-a..a \} \)

Parameters:

- image 2-D image array
\textbf{num\_rows}  Number of rows in image array

\textbf{num\_cols}  Number of cols in image array

\textbf{filter\_kernal}  2-D filter kernal

\textbf{fk\_row}  Number of rows in filter kernal

\textbf{fk\_col}  Number of cols in filter kernal

**Returns:**

Weighted average value

\textbf{B.23.2.2 Filter\_Image filter (void * image)}

Filters the input image and provides an image with only the requires features of the input image.

**Parameters:**

\textit{image}  Image to be filtered

**Returns:**

Filtered image

\textbf{B.23.3 Variable Documentation}

\textbf{B.23.3.1 const int d1\_filt[][3] [static]}
**B.24 filter_sobel_gs.c File Reference**

**B.24.1 Detailed Description**

This file implements the interface “filter.h”. It is the plugin for filtering using Sobel algorithm for grayscale values.

**Functions**

- static int ** conv2d (double **image, int num_rows, int num_cols, const int filter_kernal[][3], int fk_row, int fk_col)
- void * filter (void *image)

**Variables**

- static const int x_filt [][]3
- static const int y_filt [][]3
- static const int d1_filt [][]3
- static const int d2_filt [][]3

**B.24.2 Function Documentation**

**B.24.2.1 static int ** conv2d (double **image, int num_rows, int num_cols, const int filter_kernal[][3], int fk_row, int fk_col)  [static]**

Function for two dimensional convolution:  
\[ g(x,y) = \sum \{ \sum \{ w(s,t)f(x+s,y+t), t=-b..b \}, s=-a..a \} \]

**Parameters:**

- image 2-D image array
\( \text{num\_rows} \) Number of rows in image array

\( \text{num\_cols} \) Number of cols in image array

\( \text{filter\_kernal} \) 2-D filter kernal

\( \text{fk\_row} \) Number of rows in filter kernal

\( \text{fk\_col} \) Number of cols in filter kernal

Returns:

Weighted average value

B.24.2.2 \( \text{void\* filter (void * image)} \)

Filters the input image and provides an image with only the requires features of the input image.

Parameters:

\( \text{image} \) Image to be filtered

Returns:

Filtered image

B.24.3 Variable Documentation

B.24.3.1 \( \text{const int d1\_filt[][3]} \) [static]
B.25 match contour bw.c File Reference

B.25.1 Detailed Description

Provides contour matching for black and white images. This file implements the interface "match.h". It uses contour matching algorithm for comparing two black and white contour images.

Functions

- double match (void *image, void *target_image)

B.25.2 Function Documentation

B.25.2.1 double match (void *image, void *target_image)

Computes the score of an image. This routine is intended to be used after an image has been filtered using a sobel operator for black and white images.

Parameters:

- image Reference Contour image

- target_image Target Contour image

Returns:

- Score value between 0 and 1
B.26  match_contour_gs.c File Reference

B.26.1  Detailed Description

Provides contour matching for grayscale images. This file implements the interface
"match.h". It uses contour matching algorithm for comparing two grayscale contour
images.

Functions

- double match (void *image, void *target_image)

B.26.2  Function Documentation

B.26.2.1  double match (void *image, void *target_image)

Computes the score of an image. This routine is intended to be used after an image has
been filtered using a sobel operator for black and white images.

Parameters:

- image  Reference Contour image
- target_image  Target Contour image

Returns:

Score value between 0 and 1
B.27 search_sequential_montecarlo.c File Reference

B.27.1 Detailed Description

This file implements the interface `search.h` using Monte Carlo search technique. This is a
sequential program that contains routines for attempting to find a fluoroscopy image frame
by deriving a matching DRR image from a CT volume. Monte Carlo method is used to
achieve this. This program depends on libraries for filtering and matching a DRR image.

Functions

- static double get_score (double cam[], double offset[], double step[], EDGES
  *t_edges, CT_VOLUME *cube)
- void search (void *cube, FILE *config_file)

Variables

- void *generate_drr_handle
- void *filter_handle
- void *match_handle
- void(*) generate_drr_function (double, double, double, double[], void *, void *)
- void (*)(*) filter_function (void *)
- double(*) match_function (void *, void *)
- const char *dlError
- double offset [3] = {0,0,0}
B.27.2 Function Documentation

B.27.2.1 static double get_score (double cam[], double offset[], double step[],
    EDGES * t_edges, CT_VOLUME * cube) [static]

Creates a DRR image for the given orientation and offset and returns its score as compared
to the target DRR image.
Parameters:

   cam  Camera coordinates for the required DRR image

   offset Cube offsets for the required DRR image

   step  Camera offset step values for the required DRR image

   t_edges  Pointer to the edges of the target DRR image

   cube  Pointer to the CT volume data

Returns:

   Image score

B.27.2.2 void search (void * _cube, FILE * config_file)

This function attempts to search for an fluoroscopy image. The search algorithm is based
on a Bees algorithm, where random camera positions are tried until a close match is found.
Parameters:

   _cube  Pointer to the CT_VOLUME

   config_file  Pointer to the open file containing all the configuration values required for
                this algorithm

Returns:

   Void
B.27.3 Variable Documentation

B.27.3.1 const char* dlError

B.27.3.2 void (*)(void *) filter_function(void *)

B.27.3.3 void * filter_handle

B.27.3.4 void(*) generate_drr_function(double, double, double, double[], void *,
void *)

B.27.3.5 void* generate_drr_handle

B.27.3.6 double(*) match_function(void *, void *)

B.27.3.7 void * match_handle

B.27.3.8 double offset[3] = {0,0,0}
B.28 search_sequential_bees.c File Reference

B.28.1 Detailed Description

This file implements the interface 'search.h' using Bees Algorithm. This is a sequential program that contains routines for attempting to find a fluoroscopy image frame by deriving a matching DRR image from a CT volume. Bees algorithm is used to achieve this. This program depends on libraries for filtering and matching a DRR image.

Data Structures

- struct _bee

Typedefs

- typedef _bee bee
- typedef _bee * BeePtr

Functions

- static BeePtr create_bee ()
- static void copy_bee (BeePtr from_bee, BeePtr to_bee)
- static void free_bee (BeePtr the_bee)
- static double get_score (BeePtr the_bee, EDGES *t_edges, CT_VOLUME *cube)
- static void print_info (long iteration, int bee_id, BeePtr the_bee, char *msg)
- void search (void * _cube, FILE * config_file)
Variables

- void * generate_drr_handle
- void * filter_handle
- void * match_handle
- void(*) generate_drr_function (double, double, double, double[], void *, void *)
- void(*)(*) filter_function (void *)
- double(*)(*) match_function (void *, void *)
- const char * dlError

B.28.2 Typedef Documentation

B.28.2.1 typedef struct _bee bee

B.28.2.2 typedef struct _bee* BeePtr

B.28.3 Function Documentation

B.28.3.1 static void copy_bee (BeePtr from_bee, BeePtr to_bee) [static]

Copies the values of one bee to another bee.

Parameters:

from_bee  Pointer to the bee whose values are to be copied

to_bee    Pointer to the bee whose values should be copied from the from_bee

Returns:

Void
B.28.3.2 static BeePtr create_bee () [static]

Creates a new bee and allocates memory to it. It also initializes all its members to 0.

Returns:

Pointer to a bee

B.28.3.3 static void free_bee (BeePtr the_bee) [static]

Deallocate the memory allocated to the bee.

Parameters:

the_bee Pointer to the bee

Returns:

Void

B.28.3.4 static double get_score (BeePtr the_bee, EDGES ∗ t_edges, CT_VOLUME ∗ cube) [static]

Creates a DRR image for the given orientation and offset and returns its score as compared to the target DRR image.

Parameters:

the_bee Pointer to the bee which contains the values of camera and film orientation, cube position offsets and score

t_edges Pointer to the edges of the target DRR image

cube Pointer to the CT volume data

Returns:

Image score
B.28.3.5 void print_info (long iteration, int bee_id, BeePtr the_bee, char * msg)

    [static]

This function is used to print debug information. It is not a general purpose logger, but very specific to printing orientation and offset values for a specific bee at a specific iteration.

Parameters:

    iteration  Iteration of the search

    bee_id    Id of the bee whose info is to be printed

    msg       String that displays the message associated with the information

B.28.3.6 void search (void * _cube, FILE * config_file)

This function attempts to search for an fluoroscopy image. The search algorithm is based on a Bees algorithm, where random camera and film orientation and CT cube positions are tried until a close match is found.

Parameters:

    _cube    Pointer to the CT_VOLUME

    config_file    Pointer to the open file containing all the configuration values required for this algorithm

Returns:

    Void
B.28.4 Variable Documentation

B.28.4.1 const char* dlError

B.28.4.2 void(*)(void*) filter_function(void*)

B.28.4.3 void* filter_handle

B.28.4.4 void(*) generate_drr_function(double, double, double, double[], void*, void*)

B.28.4.5 void* generate_drr_handle

B.28.4.6 double(*) match_function(void*, void*)

B.28.4.7 void* match_handle
B.29  search_sequential_pso.c File Reference

B.29.1 Detailed Description

This file implements the interface 'search.h' using Particle Swarm Optimization method. This is a sequential program that contains routines for attempting to find a fluoroscopy image frame by deriving a matching DRR image from a CT volume. Particle Swarm Optimization algorithm is used to achieve this. This program depends on libraries for filtering and matching a DRR image.

Data Structures

- struct _particle

Typedefs

- typedef _particle particle
- typedef _particle * ParticlePtr

Functions

- static ParticlePtr create_particle ()
- static void copy_particle (ParticlePtr from_particle, ParticlePtr to_particle)
- static void free_particle (ParticlePtr the_particle)
- static double get_score (ParticlePtr the_particle, EDGES *t_edges, CT_VOLUME *cube)
- static void print_info (long iteration, int neighborhood_id, int particle_id, ParticlePtr the_particle, char *msg)
- void search (void * _cube, FILE *config_file)
Variables

- `void * generate_drr_handle`
- `void * filter_handle`
- `void * match_handle`
- `void(*) generate_drr_function (double, double, double, double[], void *, void *)`
- `void (*)(*) filter_function (void *)`
- `double(*) match_function (void *, void *)`
- `const char * dlError`

B.29.2 Typedef Documentation

B.29.2.1 typedef struct _particle particle

B.29.2.2 typedef struct _particle* ParticlePtr

B.29.3 Function Documentation

B.29.3.1 static void copy_particle (ParticlePtr from_particle, ParticlePtr to_particle) [static]

Copies the values of one particle to another particle.

Parameters:

- `from_particle` Pointer to the particle whose values are to be copied
- `to_particle` Pointer to the particle whose values should be copied from the from_- particle

Returns:

Void
B.29.3.2 static ParticlePtr create_particle () [static]

Creates a new particle and allocates memory to it. It also initializes all its members to 0.
Returns:

Pointer to a particle

B.29.3.3 static void free_particle (ParticlePtr the_particle) [static]

Deallocate the memory allocated to the particle.
Parameters:

the_particle Pointer to the particle

Returns:

Void

B.29.3.4 static double get_score (ParticlePtr the_particle, EDGES * t_edges, CT_VOLUME * cube) [static]

Creates a DRR image for the given orientation and offset and returns its score as compared to the target DRR image.
Parameters:

the_particle Pointer to the particle which contains the values of camera and film orientation, cube position offsets and score

t_edges Pointer to the edges of the target DRR image

cube Pointer to the CT volume data

Returns:

Image score
B.29.3.5  void print_info (long iteration, int neighborhood_id, int particle_id,
        ParticlePtr the_particle, char * msg) [static]

This function is used to print debug information. It is not a general purpose logger, but very
specific to printing orientation and offset values for a specific bee at a specific iteration.

Parameters:

    iteration  Iteration of the search
    neighborhood_id  ID of the particle’s neighborhood
    particle_id  ID of the particle whose info is to be printed
    the_particle  Pointer to the particle
    msg  String that displays the message associated with the information

B.29.3.6  void search (void * _cube, FILE * config_file)

This function attempts to search for an fluoroscopy image. The search algorithm is based
on a Bees algorithm, where random camera and film orientation and CT cube positions are
tried until a close match is found.

Parameters:

    _cube  Pointer to the CT_VOLUME
    config_file  Pointer to the open file containing all the configuration values required for
                  this algorithm

Returns:

    Void
B.29.4 Variable Documentation

B.29.4.1 const char* dlError

B.29.4.2 void(*)(void*) filter_function(void*)

B.29.4.3 void* filter_handle

B.29.4.4 void(*) generate_drr_function(double, double, double, double[], void*, void*)

B.29.4.5 void* generate_drr_handle

B.29.4.6 double(*) match_function(void*, void*)

B.29.4.7 void* match_handle
B.30 search_parallel_bees.c File Reference

B.30.1 Detailed Description

This file implements the interface ‘search.h’ using Bees Algorithm. This is parallel program that contains routines for attempting to find a fluoroscopy image frame by deriving a matching DRR image from a CT volume. Bees algorithm is used to achieve this. This program depends on libraries for filtering and matching a DRR image.

Data Structures

- struct _bee

Typedefs

- typedef _bee bee
- typedef _bee * BeePtr

Functions

- static BeePtr create_bee ()
- static void copy_bee (BeePtr from_bee, BeePtr to_bee)
- static void free_bee (BeePtr the_bee)
- static double get_score (BeePtr the_bee, EDGES *t_edges, CT_VOLUME *cube)
- static void print_info (long iteration, int bee_id, BeePtr the_bee, char *msg)
- void search (void *_cube, FILE *config_file)
Variables

- void * generate_drr_handle
- void * filter_handle
- void * match_handle
- void(*) generate_drr_function (double, double, double, double[], void *, void *)
- void (*)(*) filter_function (void *)
- double(*) match_function (void *, void *)
- const char * dlError

B.30.2 Typedef Documentation

B.30.2.1 typedef struct _bee bee

B.30.2.2 typedef struct _bee* BeePtr

B.30.3 Function Documentation

B.30.3.1 static void copy_bee (BeePtr from_bee, BeePtr to_bee) [static]

Copies the values of one bee to another bee.

Parameters:

from_bee  Pointer to the bee whose values are to be copied

      to_bee  Pointer to the bee whose values should be copied from the from_bee

Returns:

      Void
B.30.3.2 static BeePtr create_bee () [static]

Creates a new bee and allocates memory to it. It also initializes all its members to 0.

Returns:

Pointer to a bee

B.30.3.3 static void free_bee (BeePtr the_bee) [static]

Deallocation the memory allocated to the bee.

Parameters:

the_bee Pointer to the bee

Returns:

Void

B.30.3.4 static double get_score (BeePtr the_bee, EDGES * t_edges, CT_VOLUME * cube) [static]

Creates a DRR image for the given orientation and offset and returns its score as compared to the target DRR image.

Parameters:

the_bee Pointer to the bee which contains the values of camera and film orientation, cube position offsets and score

t_edges Pointer to the edges of the target DRR image

cube Pointer to the CT volume data

Returns:

Image score
B.30.3.5  void print_info (long iteration, int bee_id, BeePtr the_bee, char * msg)  
 [static]

This function is used to print debug information. It is not a general purpose logger, but very specific to printing orientation and offset values for a specific bee at a specific iteration.

Parameters:

iteration  Iteration of the search

bee_id  Id of the bee whose info is to be printed

msg  String that displays the message associated with the information

B.30.3.6  void search (void * _cube, FILE * config_file)

This function attempts to search for an fluoroscopy image. The search algorithm is based on a Bees algorithm, where random camera and film orientation and CT cube positions are tried until a close match is found.

Parameters:

_cube  Pointer to the CT_VOLUME

config_file  Pointer to the open file containing all the configuration values required for this algorithm

Returns:

Void
B.30.4 Variable Documentation

B.30.4.1 const char* dlError

B.30.4.2 void(*)(void*) filter_function(void*)

B.30.4.3 void* filter_handle

B.30.4.4 void(*) generate_drr_function(double, double, double, double[], void*, void*)

B.30.4.5 void* generate_drr_handle

B.30.4.6 double(*) match_function(void*, void*)

B.30.4.7 void* match_handle
B.31 search_parallel_pso.c File Reference

B.31.1 Detailed Description

This file implements the interface ‘search.h’ using Particle Swarm Optimization method. This is a parallel program that contains routines for attempting to find a fluoroscopy image frame by deriving a matching DRR image from a CT volume. Particle Swarm Optimization algorithm is used to achieve this. This program depends on libraries for filtering and matching a DRR image.

Data Structures

- struct _particle

Typedefs

- typedef _particle particle
- typedef _particle * ParticlePtr

Functions

- static ParticlePtr create_particle ()
- static void copy_particle (ParticlePtr from_particle, ParticlePtr to_particle)
- static void free_particle (ParticlePtr the_particle)
- static double get_score (ParticlePtr the_particle, EDGES *t_edges, CT_VOLUME *cube)
- static void print_info (long iteration, int neighborhood_id, int particle_id, ParticlePtr the_particle, char *msg)
- void search (void *_cube, FILE *config_file)
Variables

- void * generate_drr_handle
- void * filter_handle
- void * match_handle
- void(*) generate_drr_function (double, double, double, double[], void *, void *)
- void (*)(*) filter_function (void *)
- double (*)(*) match_function (void *, void *)
- const char * dlError

B.31.2 Typedef Documentation

B.31.2.1 typedef struct _particle particle

B.31.2.2 typedef struct _particle* ParticlePtr

B.31.3 Function Documentation

B.31.3.1 static void copy_particle (ParticlePtr from_particle, ParticlePtr to_particle)

[static]

Copies the values of one particle to another particle.

Parameters:

(from_particle) Pointer to the particle whose values are to be copied

(to_particle) Pointer to the particle whose values should be copied from the from_particle

Returns:

Void
B.31.3.2 static ParticlePtr create_particle () [static]

Creates a new particle and allocates memory to it. It also initializes all its members to 0.

Returns:

Pointer to a particle

B.31.3.3 static void free_particle (ParticlePtr the_particle) [static]

Deallocate the memory allocated to the particle.

Parameters:

the_particle  Pointer to the particle

Returns:

Void

B.31.3.4 static double get_score (ParticlePtr the_particle, EDGES * t_edges,
CT_VOLUME * cube) [static]

Creates a DRR image for the given orientation and offset and returns its score as compared
to the target DRR image.

Parameters:

the_particle  Pointer to the particle which contains the values of camera and film
orientation, cube position offsets and score

t_edges  Pointer to the edges of the target DRR image

cube  Pointer to the CT volume data

Returns:

Image score
B.31.3.5  void print_info (long iteration, int neighborhood_id, int particle_id, 
        ParticlePtr the_particle, char * msg)  [static]

This function is used to print debug information. It is not a general purpose logger, but very 
specific to printing camera and film orientation and CT cube offset values for a specific bee 
at a specific iteration.

Parameters:

  iteration  Iteration of the search

  neighborhood_id  ID of the particle’s neighborhood

  particle_id  ID of the particle whose info is to be printed

  the_particle  Pointer to the particle

  msg  String that displays the message associated with the information

B.31.3.6  void search (void * cube, FILE * config_file)

This function attempts to search for an fluoroscopy image. The search algorithm is based 
on a Bees algorithm, where random camera and film orientation and CT cube positions are 
tried until a close match is found.

Parameters:

  cube  Pointer to the CT_VOLUME

  config_file  Pointer to the open file containing all the configuration values required for 
               this algorithm

Returns:

    Void
B.31.4  Variable Documentation

B.31.4.1  const char* dlError

B.31.4.2  void(*)(void*) filter_function(void*)

B.31.4.3  void * filter_handle

B.31.4.4  void(*) generate_drr_function(double, double, double, double[], void *,
                                           void *)

B.31.4.5  void* generate_drr_handle

B.31.4.6  double(*) match_function(void *, void *)

B.31.4.7  void * match_handle