

# Human Motion Database with a Binary Tree and Node Transition Graphs

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**Abstract**—Database of human motion has been widely used for recognizing human motion and synthesizing humanoid motions. In this paper, we propose a data structure for storing and extracting human motion data and demonstrate that the database can be applied to the recognition and motion synthesis problems in robotics. We develop an efficient method for building a binary tree data structure from a set of continuous, multi-dimensional motion clips. Each node of the tree represents a statistical distribution of a set of human figure states extracted from the motion clips. We also identify the valid transitions among the nodes and construct node transition graphs. Similar states in different clips may be grouped into a single node, thereby allowing transitions between different behaviors. Using databases constructed from real human motion data, we demonstrate that the proposed data structure can be used for human motion recognition, state estimation and prediction, and robot motion planning.

## I. INTRODUCTION

Using a collection of human motion data has been a popular approach for both analyzing and synthesizing human figure motions, especially thanks to recent improvements of motion capture systems. In the graphics field, motion capture data have been widely used for producing realistic animations for films and games. A body of research efforts have been directed to techniques that allow reuse and editing of existing motion capture to new characters and/or scenarios. In the robotics field, there are two major applications of such databases: building a human behavior model for robots to recognize human motions, and synthesizing humanoid robot motions.

Databases for robotics applications are required to perform at least the following two functions: First, they have to be able to categorize human motion into distinct behaviors and recognize the behavior of a newly observed motion. This is commonly achieved by constructing a mathematical model that bridges the continuous motion space and the discrete behavior space. The problem is that it is often difficult to come up with a robust learning algorithm for building the models because raw human motion data normally contain noise and error. The efficiency of search also becomes a problem as the database size increases.

Another function is to synthesize a robot motion that adapts to current situation, which is computationally demanding because of the large configuration space of humanoid robots. Motion database is a promising approach because they can reduce the search space by providing the knowledge on how

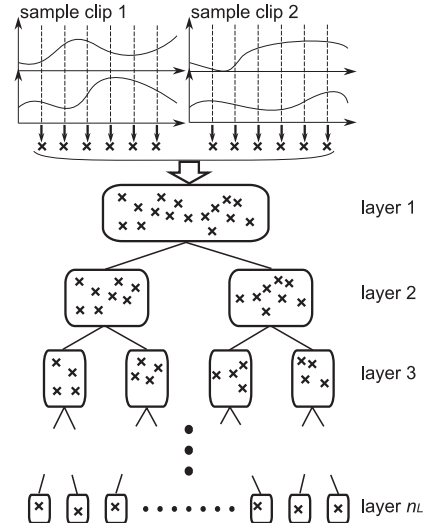


Fig. 1. A binary tree with human motion data. Each x mark represents a frame in one of the sample motion clips. The frames are iteratively split into two descendant nodes. Each frame therefore appears once in each layer. Database size can be reduced by making leaf nodes from multiple frames and keep only the statistical information of the frames included in each node.

human-like robots should move. For this purpose, however, motion capture data must be organized so that the planner can effectively extract candidates of motions and/or configurations. A database should also be able to generate high-quality motion data, which is also difficult because sample motion data are usually compressed to reduce the data size.

In this paper, we propose a highly hierarchical data structure for human motion database. We employ the binary-tree structure as shown in Fig. 1, which is a classical database structure widely used in various computer science applications because of search efficiency. However, constructing a binary-tree structure from human motion data is not a trivial problem because there is no straightforward way to split the multi-dimensional, continuous motion data into two descendant nodes. Our first contribution is a simple, efficient clustering algorithm for splitting a set of sample frames into two descendant nodes to construct a binary tree from human motion data.

We also develop algorithms for basic statistical computations based on the binary tree structure. Using these algo-

gorithms, we can recognize a newly observed motion sequence, estimate the current state and predict future motions, and plan new sequences that satisfy given constraints.

Another minor but practically important aspect of the proposed database is the ability to incorporate motion data from different sources. For example, we may want to include motion data captured with different marker sets, or include animation data from a 3D CG film. It is therefore important to choose a good motion representation to allow different marker sets and/or kinematic models. In this paper, we propose a scheme called *virtual marker set* so that motion data from different sources can be represented in a uniform way and stored in a single database.

The rest of this paper is organized as follows. In Section II, we review related work in graphics and robotics. We then present the proposed data structure and associated algorithms in Sections III and IV respectively, and provide several application examples in Section V. Section VI demonstrates experimental results using human motion capture data, followed by concluding remarks.

## II. RELATED WORK

In the graphic field, researchers have been investigating how to reuse and edit motion capture data for new scenes and new characters. One of the popular approaches is motion graphs, where relatively large human motion data set is analyzed to build a graph of possible transitions between postures. Using the graph, it is possible to synthesize new motion sequences based on simple user inputs by employing a graph search algorithm. Kovar et al. [1] proposed the concept of Motion Graphs where similar postures in a database are automatically detected and connected to synthesize new motion sequences. They presented an application of synthesizing new locomotion sequence that follows a user-specified path. Lee et al. [2] employed a two-layered statistical model to represent a database, where the higher-level (coarse) layer is used for interacting with user inputs and the lower-level (detail) layer is used for synthesizing whole-body motions. Arikan et al. [3] also proposed a planning algorithm based on a concept similar to motion graphs. Related work in the graphics field mostly focuses on synthesizing new motions from simple user inputs using, for example, interpolation and numerical optimization [4].

In robotics, learning from human demonstration, or imitation, has been a long-term research issue [5]–[7] and a number of algorithms have been developed for storing human motion data and extracting appropriate behaviors. Because human motion varies at every instance even if the subject attempts to perform the same motion, it is necessary to model human behaviors by either statistical models [7], [8], nonlinear dynamical systems [9], [10], or a set of high-level primitives [11]. Related work relevant to this paper includes the Hidden Markov Model (HMM) representation of human behaviors [8] and the hierarchical motion database based on HMM [12]. Another hierarchical motion categorization method is also proposed using neural network models [10].

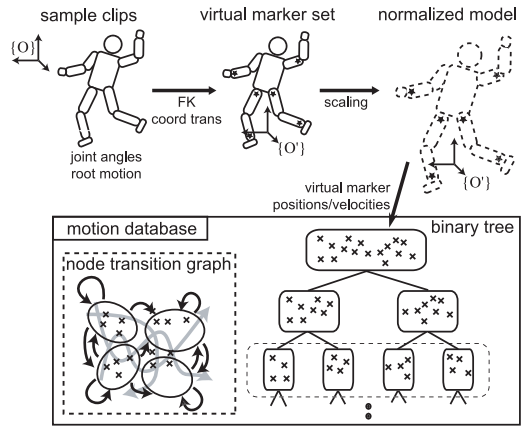


Fig. 2. Constructing the database.

However, most work in the robotics field is still focused on robust learning of human behaviors. Scalability of the motion database or synthesizing transitions between different behaviors have not been investigated well.

In the vision community, human motion database has been used to construct human motion models for human tracking in videos. Sidenbladh et al. [13] proposed a binary-tree data structure for building a probabilistic human motion model. They first performed PCA on the entire motion sequence and then projected each frame data to the principal axes to construct a binary tree.

## III. BUILDING THE DATABASE

The process for building the proposed database is summarized in Fig. 2. The user provides one or more sample motion clips represented as a pair of root motion and joint angle sequences, typically obtained by motion capture or hand animation. The joint angle data are then converted to virtual marker data through forward kinematics (FK) computation to obtain the marker positions and velocities, coordinate transformation to remove the trunk motion in the horizontal plane, and scaling to normalize the subject size. The positions and velocities of the virtual markers are used to represent the state of the human figure in each sample frame.

To construct a binary tree, we first create the root node that contains all frames in the sample motion clips. We then iteratively split the frames into two descendant nodes using the method described in Section III-B. After the tree is obtained, we count the number of transitions among the nodes in each layer to construct the node transition graphs as described in Section III-C. The binary tree and node transition graphs are the main elements of the proposed motion database.

### A. Motion Representation

There are several choices for representing the state of a frame in sample motion clips. A reasonable choice would be to use joint angles [2], [8] because they uniquely define a configuration by minimum number of parameters. However, joint angle representation strictly depends on the skeleton

model and it is difficult to map the states between different models. In addition, joint angle representation may not be consistent with visual appearance of the human figure because the contribution of each joint angle to the Cartesian positions of the links can vary.

Another possibility is to use point clouds [1]. This method is independent of underlying skeleton models and also more intuitive because it directly represent the overall shape of the figure. The problem is that it is difficult compute the distance between two poses because registration between two point clouds is required.

In our implementation, we use a much simpler approach called *virtual marker set*, where all motions are represented by the trajectory (position and velocity) of  $N_v$  virtual markers. The virtual marker set is defined by the database designer so that it well represents the motions contained in the database, and can be different from any physical marker sets. The virtual marker set approach would become similar to the point cloud method as the number of virtual markers increases, although each marker in the virtual marker set should be labeled.

If a motion is represented by joint angle trajectories of a skeleton model, it can be easily converted to the virtual marker set representation by giving the relationship between each marker in the virtual marker set and the skeleton used to represent the original motions. The relationship between a virtual marker and a skeleton is defined by specifying the link to which the marker is attached, and giving the relative position in its local frame. Although this approach requires some work on the user's side, it allows the use of multiple skeleton models with simple distance computation.

After converting the motion data to the virtual marker set representation, we perform a coordinate transformation to remove the horizontal movement in the motion and scaling to normalize the subject size. Each marker position is represented in a new reference coordinate whose origin is located on the floor below the root joint,  $z$  axis is vertical, and  $x$  axis is the projection of the front direction of the subject model onto the floor. The marker velocities are also converted to the reference coordinate.

Formally, a sample motion data matrix  $\mathbf{X}$  with  $N_S$  frames is defined by

$$\mathbf{X} = (\mathbf{x}_1 \ \mathbf{x}_2 \ \dots \ \mathbf{x}_{N_S}) \quad (1)$$

where  $\mathbf{x}_i$  is the state vector of the  $i$ -th sample frame defined by

$$\mathbf{x}_i = (\mathbf{p}_{i1}^T \ \mathbf{v}_{i1}^T \ \mathbf{p}_{i2}^T \ \mathbf{v}_{i2}^T \ \dots \ \mathbf{p}_{iN_v}^T \ \mathbf{v}_{iN_v}^T)^T \quad (2)$$

and  $\mathbf{p}_{il}$  and  $\mathbf{v}_{il}$  are the position and velocity of marker  $l$  in sample frame  $i$ . If multiple motion clips are to be stored in a database, we simply concatenate all state vectors horizontally and form a single sample motion data matrix.

## B. Constructing a Binary Tree

A problem of constructing a binary tree for motion data is how to cluster the sample frames into groups with similar states. Most clustering algorithms require a large amount of computation because they check the distances between all

pairs of frames. This process can take extremely long time as the database size increases.

Here we propose an efficient clustering algorithm based on principal component analysis (PCA) and minimum-error thresholding technique. The motivation for using PCA is that it determines the axes that best characterize the sample data. In particular, projecting all samples onto the first principal axis gives a one-dimensional data set with the maximum variance, which can then be used for separating distinct samples using adaptive thresholding techniques developed for binarizing images.

The process to split node  $k$  into two descendant nodes is as follows. Assume that node  $k$  contains  $n_k$  frames whose mean state vector is  $\bar{\mathbf{x}}_k$ . Also denote the sample motion data matrix of node  $k$  by  $\mathbf{X}_k$ . We compute the zero-mean singular value decomposition of  $\mathbf{X}_k$  as

$$\mathbf{X}'_k{}^T = \mathbf{U} \mathbf{\Sigma} \mathbf{V}^T \quad (3)$$

where each column of  $\mathbf{X}'_k$  is obtained by subtracting  $\bar{\mathbf{x}}_k$  from the original state vectors,  $\mathbf{\Sigma}$  is a diagonal matrix whose elements are the singular values of  $\mathbf{X}_k$  sorted in the descending order, and  $\mathbf{U}$  and  $\mathbf{V}$  are orthogonal matrices. The columns of  $\mathbf{V}$  represents the principal axes of  $\mathbf{X}_k$ . We obtain the one-dimensional data set  $\mathbf{s}_k$  by projecting  $\mathbf{X}'_k$  onto the first principal axis by

$$\mathbf{s}_k = \mathbf{X}'_k{}^T \mathbf{v}_1 \quad (4)$$

where  $\mathbf{v}_1$  denotes the first column of  $\mathbf{V}$ .

Once the one-dimensional data is obtained, the clustering problem is equivalent to determining the threshold that minimizes classification error. We shall use a minimum-error thresholding technique [14] to determine the optimal threshold. After sorting the elements of  $\mathbf{s}_k$  and obtaining the sorted vector  $\mathbf{s}'_k$ , this method determines the index  $m$  such that the data should be divided between samples  $m$  and  $m + 1$  using the following equation:

$$m = \operatorname{argmax}_i \left\{ i \log \frac{\sigma_1}{i} + (n_k - i) \log \frac{\sigma_2}{n_k - i} \right\} \quad (5)$$

where  $\sigma_1$  and  $\sigma_2$  denote the variance of the first  $i$  and last  $n_k - i$  elements of  $\mathbf{s}'_k$ , respectively.

We obtain a binary tree by repeating this process until the division creates a node containing fewer frames than a predefined threshold. To ensure the statistical meaning of each node, we also avoid nodes with small number of sample frames by setting a threshold for minimum frame number. If Eq.(5) results in a node with fewer number of frames than the latter threshold, we do not perform the division. Therefore, a node may not be divided if it contains many similar frames.

Some branches may be shorter than others because we extend each branch as much as possible and some of them may hit the thresholds earlier. In such cases, we simply extend shorter branches by attaching a copy of the leaf node so that the length of all branches become the same. Each node therefore can have 0 (leaf nodes), 1 or 2 descendant nodes.

### C. Node Transition Graphs

After constructing the binary tree, we then build the node transition graphs based on the transitions observed between nodes in each layer. Because we know the set of frames included in each node, we can easily determine the transition probabilities by dividing the number of transitions to a specific node by the total number of frames in the node.

We build two kinds of node transition graphs at each layer. The *global transition graph* describes the average node transition probabilities observed in all sample clips. The transition probability from node  $m$  to node  $n$  in the same layer is computed by

$$p_{m,n} = \frac{t_{m,n}}{\sum_i t_{m,i}} \quad (6)$$

where  $t_{k,l}$  denotes the total number of transitions from nodes  $k$  to  $l$  observed in all sample clips. A *clip transition graph* describes the node transition probabilities observed in a specific sample clip. We can use the same equation (6) to compute the transition probabilities, except that  $t_{k,l}$  only considers the transitions within the same sample clip.

The global transition graph at each layer is similar to motion graphs [1] in the sense that all possible transitions between nodes are included. However, the way we construct the graph is different from existing motion graph techniques, resulting in a more efficient database construction. Our method generally requires  $O(N \log N)$  for a database with  $N$  sample frames because the depth of the tree is typically  $O(\log N)$  and splitting the frames at each layer requires  $O(N)$  computations, while most motion graph and other clustering techniques require  $O(N^2)$  computations because they usually compute the distance between each pair of frames in the database.

The clip transition graphs are similar to human behavior models using HMMs [8]. In most HMM-based approaches, a simple left-to-right model or single-chain cyclic model with fixed number of nodes is assumed because it is difficult to train an HMM with arbitrary length or arbitrarily interconnected nodes. In our method, we do not assume the structure of the node transition or the number of nodes used to represent a sample clip. If a sample clip includes a cyclic pattern, for example, our method automatically models the cycle by producing a single transition loop, while a left-to-right model tries to encode the whole sequence within a given number of nodes.

## IV. ALGORITHMS

For a given tree and node transition graph, we should be able to perform the following two basic operations:

- find the optimal node transition to generate a newly observed motion clip, and
- compute the probability that a newly observed motion clip is generated by a node transition graph.

In the rest of the section, we shall denote the newly observed motion comprising  $M$  frames by  $\hat{\mathbf{X}} = (\hat{\mathbf{x}}_1 \hat{\mathbf{x}}_2 \dots \hat{\mathbf{x}}_M)$  where  $\hat{\mathbf{x}}_i$  is the state vector at frame  $i$ . Here we assume that both positions and velocities of virtual markers are given.

### A. Optimal Node Transition

The probability that the observed motion  $\hat{\mathbf{X}}$  was produced by a node transition  $\mathcal{N} = \{n_1, n_2, \dots, n_M\}$  is given by

$$P(\mathcal{N}|\hat{\mathbf{X}}) = \prod_{i=1}^M P_t(n_{i-1}, n_i) P_s(n_i|\hat{\mathbf{x}}_i) \quad (7)$$

where  $P_t(k, l)$  is the transition probability from node  $k$  to  $l$  ( $P(n_0, n_1) = 1$ ) and  $P_s(k|\mathbf{x})$  is the probability that the state was at node  $k$  when the observed state was  $\mathbf{x}$ .  $P_s(k|\mathbf{x})$  is obtained by the Bayesian inference:

$$P_s(k|\mathbf{x}) = \frac{P_o(\mathbf{x}|k)P(k)}{P(\mathbf{x})} = \frac{P_o(\mathbf{x}|k)P(k)}{\sum_i P_o(\mathbf{x}|i)} \quad (8)$$

where  $P_o(\mathbf{x}|k)$  is the likelihood that state vector  $\mathbf{x}$  is output from node  $k$  and  $P(k)$  is the *a priori* probability that the state is at node  $k$ .

The actual form of  $P_o(\mathbf{x}|k)$  depends on the probability distribution used for each node. In this paper, we assume a simple Gaussian distribution with mean  $\bar{\mathbf{x}}_k$  and covariance  $\Sigma_k$ , in which case  $P_o(\mathbf{x}|k)$  can be computed by

$$P_o(\mathbf{x}|k) = \frac{1}{(\sqrt{2\pi})^N |\Sigma_k|} \exp\left(-\frac{1}{2}(\mathbf{x} - \bar{\mathbf{x}}_k)^T \Sigma_k^{-1} (\mathbf{x} - \bar{\mathbf{x}}_k)\right).$$

$P(k)$  can be either a uniform distribution among the nodes, or weighted according to the number of sample frames included in the nodes. In our implementation, we use a uniform distribution for the global transition graph. We also use a uniform distribution for each clip transition graph, excluding the nodes that did not appear in the sample clip.

Obtaining the optimal node transition  $\mathcal{N}^*$  is the problem of finding the node sequence that maximizes Eq.(7). A common method for this purpose is to perform forward-backward algorithm or dynamic programming. However, such algorithms can be computationally expensive for long sequences or densely connected graphs. We could omit nodes far enough from each observed frame using a threshold, but it is difficult to determine the threshold so that enough number of candidates are left for the search.

Instead of searching the entire node sequence at a single layer, we utilize the binary tree data structure by starting from the top layer. Because the top layer only contains the single root node  $r$ , the trivial optimal sequence at the top layer,  $\mathcal{N}_1^*$ , is to visit the root node  $M$  times, i.e.,  $\mathcal{N}_1^* = \{r, r, \dots, r\}$ . Starting from this initial sequence, we perform a dynamic programming to find the best way to trace the descendants of the nodes in the sequence all the way down to the bottom layer. We could also terminate at an intermediate layer if we do not need precise results, in which case the result would be obtained faster.

### B. Motion Generation Probability

Motion generation probability is defined as the probability that a node transition graph generates the observed motion. This probability can be used for identifying the type of behavior. We can compute the motion generation probability

by summing up the probability of generating the motion by all possible node transitions. However, there may be huge number of possible node transitions for long motions or large transition graphs.

An alternative used in this paper is to use the dynamic programming described in the previous subsection to find multiple node sequences. Because the algorithm returns node sequences in the descending order of probability, we can approximate the total motion generation probability by using the top few node sequences.

## V. APPLICATIONS

### A. State Estimation and Prediction

Estimating the current state is accomplished by taking the last node in the most likely node transition in the global node transition graph. Once the node transition is estimated with high probability, we can then predict the next action by tracing the node transition graph. By combining the probability of the node transition and the probability of the future transition candidate, we can also obtain the confidence of the prediction.

### B. Motion Recognition

Motion recognition is the process to find a motion clip in the database that best matches a newly observed motion sequence. This is accomplished by comparing the motion generation probability from the clip transition graphs.

### C. Motion Planning

The global transition graph can also be used for planning a new motion sequence subject to kinematic constraints. In addition, because the tree has multiple layers that model the sample clips at different granularities, motion planning can also be performed at different precision levels. The only issue is how to compute the motion of the root in the horizontal plane because those information are removed from the database.

Our solution is to use the marker velocity data to obtain the root velocity, and then integrate the velocity to obtain root trajectory. The velocity can be obtained by employing a numerical inverse kinematics algorithm, which essentially solves the following linear equation:

$$\dot{\theta} = \mathbf{J}v \quad (9)$$

where  $\dot{\theta}$  is the joint velocities including the root linear and angular velocities,  $v$  is the vector of all marker velocities extracted from the mean vector of a node, and  $\mathbf{J}$  is the Jacobian matrix of the marker positions with respect to joint angles. This is usually an overconstrained IK problem because the virtual marker set contains more markers than necessary to determine the joint motion. We therefore apply the singularity-robust (SR) inverse [15] of  $\mathbf{J}$  to obtain  $\dot{\theta}$ .

## VI. RESULTS

### A. Sample Data Set

We use two different sets of data to demonstrate the generality of our approach. The minimum number of sample

TABLE I  
PROPERTIES OF THE TWO DATABASES.

	<i>Database 1</i>	<i>Database 2</i>
# of frames	5539	11456
# of layers	17	16
# of nodes	1372	1527
# of nodes in bottom layer	167	205

frames in a node is set to 16 in both databases. The properties of the databases are summarized in Table I.

The first set (*Database 1*) consists of 19 motion clips including a variety of jog, kick, jump, and walk motions, all captured separately in the motion capture studio at University of Tokyo. The motions were captured using our original marker set consisting of 34 markers. This marker set is also used as the virtual marker set to construct all the databases in the experiments. *Database 1* is used to demonstrate the analysis experiments.

The second set (*Database 2*) is generated from selected clips in a publicly available motion library [16] and will be used for the planning experiment. The database includes two long locomotion sequences with forward, backward and sideward walk of the same subject. Although *Database 2* contains twice as many sample frames as *Database 1*, it resulted in relatively small number of layers and nodes probably because they consist of similar motions and hit the minimum frame number threshold earlier.

Because the virtual marker set consists of 34 markers, the dimension of the state vector is 204.

### B. Properties of the Database

We first visualize *Database 1* to investigate its properties. In Fig. 3, the spheres (light-blue for readers with color figures) represent all nodes in the bottom layer and the red lines represent the transitions among the nodes. The white mesh represents the plane of first and second principal axes. The projection onto the first-second principal axes plane is also shown in Fig. 4. Figure 5 shows the nodes and global transition graphs at layers 4 and 7, with the top layer numbered layer 1.

The location of each node is computed by projecting its mean vector onto the first three principal components of the root node of the tree. The size of the node is proportional to the maximum singular value of the sample motion data matrix of the node.

Figure 6 shows the mean marker positions and velocities of the nodes marked in Fig. 4. These images clearly show the geometrical meaning of the first and second principal axis: the first axis represents the vertical velocity and the second axis represents which leg is raised. Apparently there is no verbally explainable meaning in the third axis.

Figure 7 shows the node transition graphs of two representative clips. As expected from the above observation, motions such as jump that involves mostly the vertical motion stay in the first and third axes plane, while jog, kick and walk motions stay in the second and third axes plane.

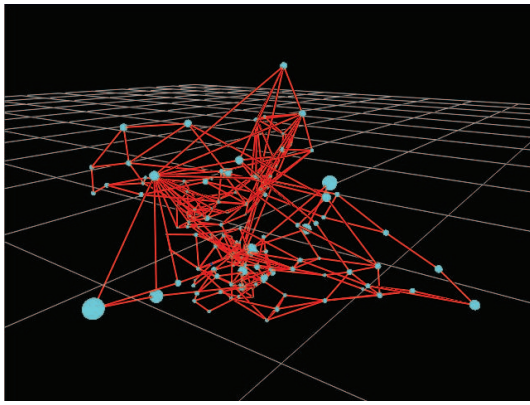


Fig. 3. Visualization of the database in the three principal axes space.

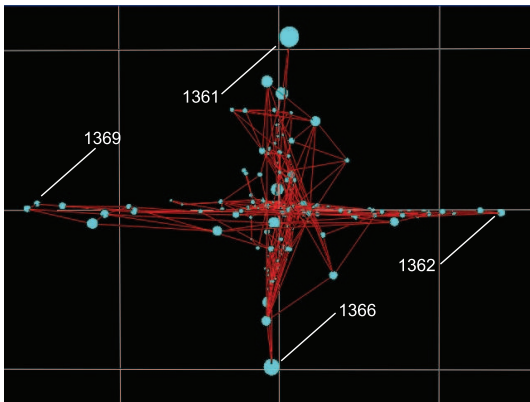


Fig. 4. Visualization of the database in the first (horizontal) and second (vertical) principal axes space.

### C. State Estimation and Prediction

We experimented the state estimation ability by providing the first 0.2 s of a novel kick motion by the same subject and a jump motion by a different subject. The results are shown in Figures 8 and 9 respectively. Note that the global node transition is used to compute the best node sequence, although only the node transition in relevant sample clip is drawn in each figure for clarity.

In both cases, the database could predict the subjects' actions before they actually occurred. In Fig. 8, all nodes on the identified node sequence (drawn as yellow spheres)

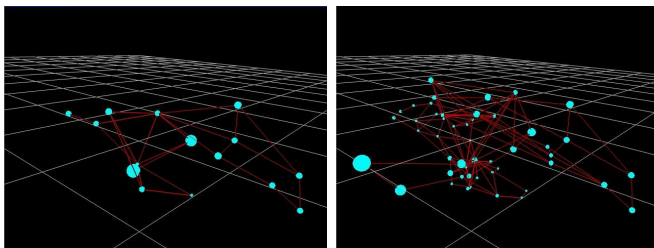


Fig. 5. Nodes and global transition graphs at layers 4 and 7.

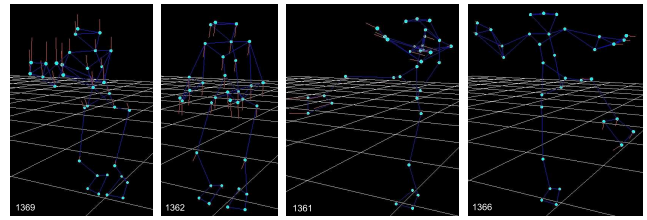


Fig. 6. Mean marker positions and velocities of selected nodes. The lines rooted at markers denote the velocities. From left to right: nodes 1369, 1362, 1361, and 1366 as marked in Fig. 4.

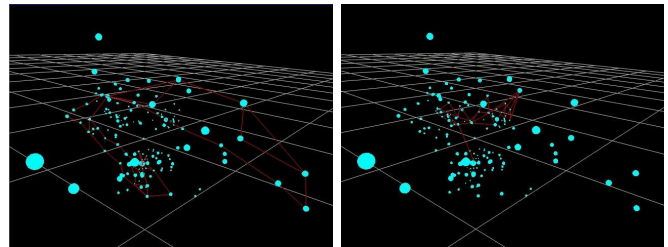


Fig. 7. Node transition graphs of selected clips. Left: jump, right: kick.

are included in the transition graph of the kick samples. The last node in the sequence is likely to transit to the dark blue node in the left figure with the probability of 0.36, which corresponds to the marker positions and velocities in the right figure. Similarly, the result of Fig. 9 indicates that the database can correctly identify that the subject is in preparation for a jump.

### D. Motion Recognition

Figures 10–11 show the results of motion recognition experiments. The three graphs in Fig. 10 show differences between layers for the same observation. We computed the motion generation probability of new motions with respect to the node transition graph of each sample motion clip. Because the new motion is 2.5 s long and computation of the probabilities takes long time, we computed the probability of generating the

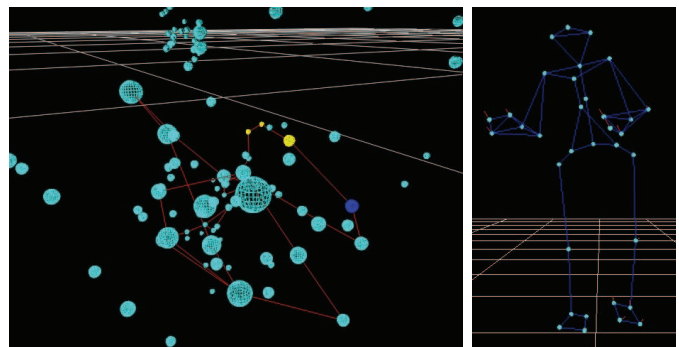


Fig. 8. Result of state estimation and prediction for a kick motion of the same subject. Left: the nodes on the identified node transition are drawn in yellow. Right: the marker positions and velocities corresponding to the blue node in the left figure.

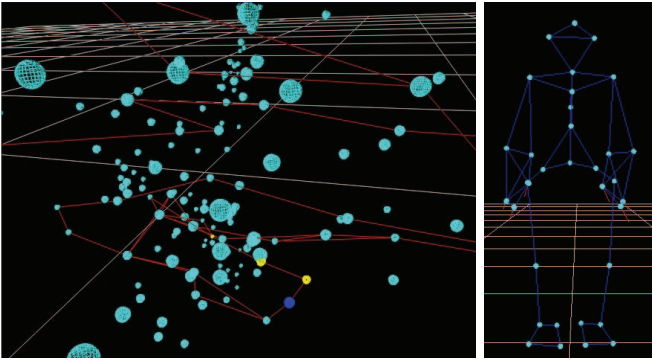


Fig. 9. Result of state estimation and prediction for a jump motion of a different subject. Left: the nodes on the identified node transition are drawn in yellow. Right: the marker positions and velocities corresponding to the blue node in the left figure.

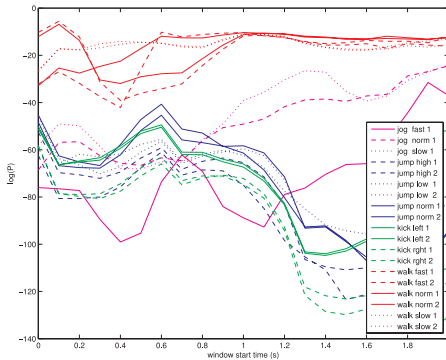


Fig. 11. Time profile of generation probability of a walk motion of a different subject (bottom layer).

sequences within a sliding window of width 0.4 s. The graphs depict the time evolution of probabilities when we moved the window from [0.0s, 0.4s] to [2.0s, 2.4s]. The lines are colored according to the type of motions. We use same line color and type for very similar sample clips for clarity of presentation.

The first three graphs show the probability of a new walk motion by the same subject as the database, while a walk motion from a different subject was used for Fig. 11. These results show that the model can successfully recognize the type of observed motion even for different subjects. It is also suggested that the statistical models at different layers have different granularity levels because the variation of probabilities is smaller for upper layers. In particular, similar motions such as walk and jog begin to merge at layer 7.

Figure 12 show the result at the bottom layer for a completely unknown motion (lifting an object from the floor). It is clear that the database cannot tell whether the motion is jump or kick, which is intuitively reasonable because these three motions start from small bending.

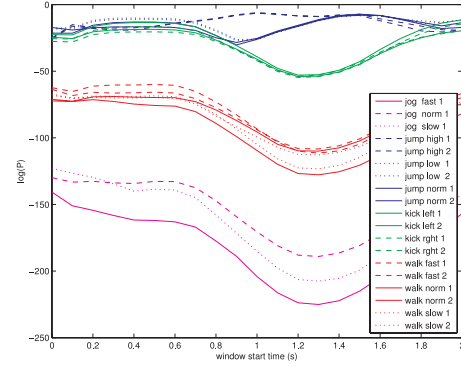


Fig. 12. Time profile of generation probability of a lifting motion (bottom layer).

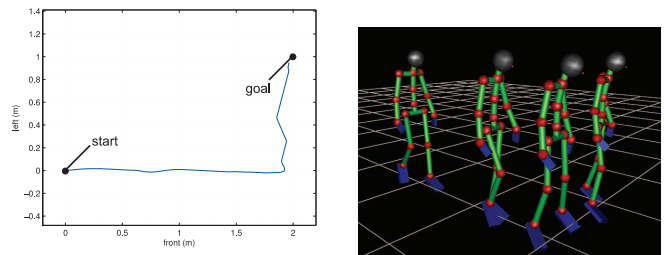


Fig. 13. Result of motion planning. Left: path of the root in horizontal plane. Right: postures at selected frames.

### E. Motion Planning

We performed a simple motion planning test with only start and goal position constraints. Figure 13 shows the planned motion when the goal position is given as 2.0 m front and 1.0 m left of the current position. The planner outputs a reasonable motion under the available samples, which is to walk forward for 2 m and do a side walk to the left for 1 m. We can observe that the feet occasionally penetrate the floor. We would need the contact status information to fix this problem.

## VII. CONCLUSION

In this paper, we proposed a new data structure for storing multi-dimensional, continuous human motion data, and demonstrated its basic functions through experiments. The main contributions of the paper are summarized as follows:

- 1) We proposed a method for constructing a binary tree data structure from human motion data. We applied PCA and a minimum-error thresholding technique to efficiently find the optimal clustering of sample frames.
- 2) We proposed to build a global node transition graph representing the node transitions in all sample clips, as well as a clip transition graph representing the node transitions in each sample motion clip.
- 3) We developed two algorithms for computing the most probable node transition and generation probability for

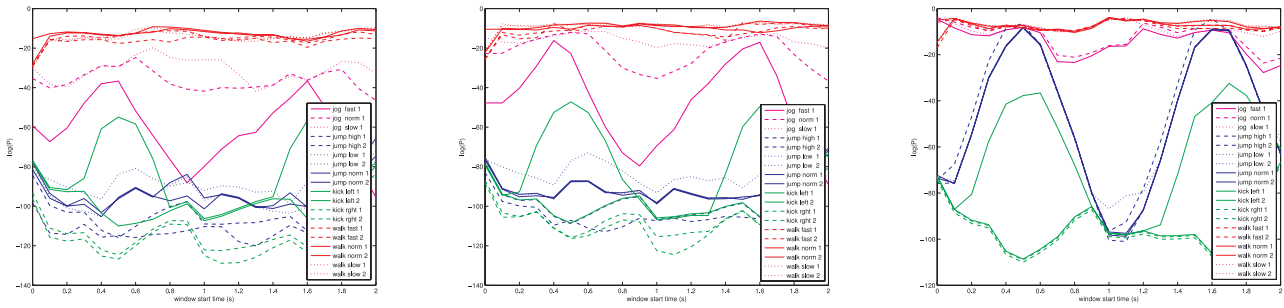


Fig. 10. Time profile of generation probability of a new walk motion of the same subject. From left to right: bottom layer, layer 10, layer 7.

a given new motion sequence, based on the binary tree data structure.

- 4) We demonstrated three applications of the proposed database through experiments using real human motion data.

There are several functions yet to be addressed by our database. We currently do not support incremental learning of new motions because all sample frames must be available to perform the PCA. If the new sample clips do not drastically change the distribution of whole samples, we could apply one of the extensions of PCA for online learning techniques [17]. Some techniques for balancing binary trees for multi-dimensional data could also be employed to reorganize the tree [18].

It should be relatively easy to add segmentation and clustering functions because the sample motion clips are abstracted by node transition graphs. We can easily detect same or similar node transitions in different motion clips, which could be used for segmentation. Clustering of sample clips can be achieved by evaluating the distance between motion clips based on their node transition graphs and applying standard clustering algorithms [19].

Our current database only contains the marker position and velocity data. It would be interesting to add other modalities such as contact status, contact forces, and muscle tensions. In particular, we would be able to solve the contact problem in our planning experiment if we have access to the contact status information. Contact force and muscle tension information would also help generating physically feasible motions for humanoid robots.

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