Preprocessing of Configuration Space for Improved Sampling Based Path Planning

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Abstract: Sampling based planners have been successful in path planning of robots with many degrees of freedom, but still remains ineffective when the configuration space has a narrow passage. This paper presents two new techniques of preprocessing the configuration space. The first technique called a Random Walk to Surface (RWS), uses a random walk strategy to generate samples in narrow regions quickly, thus improving efficiency of Probabilistic Roadmap (PRM) based planners. The algorithm substantially reduces instances of collision checking and thereby decreases computational time. The method is powerful even for cases where the structure of the narrow passage is not known a priori, thus giving significant improvement over other known methods. The second method, by preprocessing the configuration space, improves the efficiency of Rapidly Exploring Random Tree (RRT) like planners by identifying key regions of the configuration space to search for a solution path. The Experiments show a significant improvement in efficiency for both PRM and RRT like planners.

Key words: Robot Motion Planning, Randomized Algorithm, PRM, RRT

1. INTRODUCTION

Several areas beyond classical robotics, are benefiting from advances in algorithmic motion planning. Examples include structural studies in biology, computer games, path planning of unmanned aerial vehicles among the others. The success of these algorithms on different instances is because of availability of an abstract formulation of the model of many different real world applications, and ability to solve difficult problems in a reasonable time. Modern motion planning begins with the introduction of the notion of configuration space by Lozano-Perez and Wesley [1], [2]. In the configuration space the robot is reduced to a point; hence the motion planning problem becomes that of finding a path from an initial point to a goal point in the configuration space. However, the explicit mapping from workspace obstacles to configuration space is in general difficult. Early studies showed that the basic version of this problem is PSPACE-complete [3], [4] [5] and the best exact deterministic algorithm known is exponential in the dimension of the configuration space. On the other hand, real world problems generate instances with high dimensional configuration spaces.

Since the mid nineties, in order to break this "curse of dimensionality", sampling based approaches were introduced. The approach is based on the generation of samples to acquire information about the problem instance being solved. The first generation of these algorithms heavily relied on random samples. In contrast, there has been a recent trend to use deterministic sampling schemas [6]. For motion planning problems, samples are stored in a data structure which represents an approximation of the configuration space, as opposed to its exact combinatorial representation. The data structure is usually composed of nodes, that is, samples in the configuration space, and links, that is, valid paths connecting samples. Nodes and links can be stored in the form of graphs or trees. The entire continuum of configuration space is then approximated to a network of nodes. The implementation of these algorithms is usually quite simple. The price to pay is completeness.

Traditional combinatorial motion planning algorithms are complete, that is, they will find a solution if one exists, and will report failure otherwise. Algorithms based on randomly generated samples aims for probabilistic completeness. This means that if a solution exists, the probability to find it converges to 1 when the computation time approaches infinity [7]. Despite probabilistic completeness, randomized algorithms such as the Randomized Potential Field Planner (RPP) [8], the Probabilistic Road-map (PRM) family [9], Rapidly-
Exploring Random Trees (RRTs) [10], and others, have dominated the field of motion planning for over a decades.

Despite the success of sampling based path planners, motion planning in high dimensional configuration space is difficult. Several instances of the problem have been even proven to be undecidable. It is unlikely that sampling, can overcome such difficulty entirely. For PRM like planners one such difficulty arises when configuration space posses narrow passages. Narrow passages are those critical regions in the configuration space that require a number of sample points in order to capture the free space connectivity. This difficulty posed by narrow passages and its importance were noted in early work on PRM planners [9]. Several sophisticated sampling strategies can remove this difficulty to a large extent, but a satisfactory answer remains elusive. Likewise, the issue of exploration vs. biasing issues in case of RRT poses a significant challenge. The 'rapid' nature of RRT is due to the fact that the samples pulls the tree towards the unexplored areas of configuration space. This leads to a creation of large number of unnecessary nodes before obtaining a final solution. The exploration weightage and biasing weightage towards the goal is a critical design issue.

In this paper we propose a new methods which is suitable to tackle problem of narrow passages in high dimensions for PRM. We also present a sequential sampling scheme which can guide an RRT effectively toward the goal state. In Section 2 we formulate the general motion planning problem. In Section 3 we briefly explain PRM method, problem of narrow passage and related work. Section 4 describes our algorithm for narrow passage sampling. Section 5 discuss the implementation issues and result. Section 6 reports RRT algorithm and related exploration vs. biasing issues. Section 7 gives the details of our proposed methodology to improved performance of RRT, while in subsequent sections we discuss and summarize the implementation issues and results.

2. PROBLEM FORMULATION

The configuration of a robot with \( n \) DOFs can be represented as a point in an \( n \)-dimensional space, called the configuration space \( C \), which is locally like the \( d \)-Dimensional Euclidian Space \( \mathbb{R}^d \). A configuration \( q \) is free if the robot placed at \( q \) does not collide with the obstacles or with itself. We define the free space \( C_{\text{free}} \) to be the set of all free configurations in \( C \). There are a finite number of obstacles in the configuration space, which are closed bounded sets \( O_i, i = 1, 2, \cdots, m \) and we can fairly assume that they are pairwise disjoint. Let the starting configuration be \( x_{\text{start}} \in C_{\text{free}} \) and the final configuration be \( x_{\text{goal}} \in C_{\text{free}} \). For convergence issues we define a rather general goal subset \( X_{\text{goal}} \), than a specific point. Clearly \( x_{\text{free}} \in X_{\text{goal}} \). The motion planning problem is to find a path connecting \( x_{\text{start}} \) with \( X_{\text{goal}} \), that is, a continuous function \( f : [0, 1] \rightarrow C_{\text{free}} \) such that \( f(0) = x_{\text{start}} \) and \( f(1) \in X_{\text{goal}} \). We assume an appropriate collision checker function does exist for instance of the problem. In related literature the configuration space \( C \) is frequently substituted with the state space \( X \), which includes both the degrees of freedom and their derivatives. The constraints can be non-holonomic and differential.

3. PRM AND NARROW PASSAGE PROBLEM

Probabilistic road-map methods solves motion planning problems that do not involve dynamics of the robot or have negligible dynamics. A classic multi-query PRM planner proceeds in two stages. In the first stage, it randomly chooses samples from \( C_{\text{free}} \), called milestones according to a sampling scheme (which in general uniform). It then uses these milestones as nodes to construct a graph, called a road-map, by adding an edge between every \( k \) pair of milestones that can be connected via a simple collision-free path, typically, a straight-line segment. After the road-map has been constructed, multiple queries can be answered quickly in the second stage. Each query consists of an initial configuration and a goal configuration, and asks for a collision-free path connecting \( x_{\text{start}} \) and \( x_{\text{goal}} \). The planner first finds two milestones in the road-map, such that \( x_{\text{start}} \) and \( x_{\text{goal}} \) can be connected to these nodes, respectively. The rest of the job is to search for a path (Or may be search for the shortest path).

If the configuration space possess a narrow passage, then to capture the connectivity of \( C_{\text{free}} \), it is essential to sample milestones in narrow passages. This, however, is difficult, because of small volumes of narrow passages. Any volume-based sampling distribution is likely to fail. Uniform distribution may not work well when the dispersion of the samples is higher than the narrow passage volumes. Furthermore, when dealing with multiple degree of freedom robots, we do not have an explicit representation of \( C_{\text{free}} \) and cannot locate narrow passages directly by processing the global geometry of \( C_{\text{free}} \). Intuitively, one can sample more densely near obstacle boundaries because points in narrow passages lie close to obstacles. This method called Gaussian sampler [11] is a simple and efficient algorithm that uses this idea. However, in some cases, many points near the obstacle boundaries lie far away from narrow passages and do not help in improving the connectivity of road-maps. So, despite the improvements, sampling near obstacle boundaries may generate many samples in uninteresting regions. Other geometric approaches [12], [13], [14] also exist but those are expensive to implement in high-dimensional configuration spaces.

Perhaps the most appealing scheme to answer the problem of narrow passage is the Bridge test [15]. Here when a sample lies within an obstacle, one uses this information to build a bridge whose two end points lie in the obstacle while the mid point lies in \( C_{\text{free}} \). This method, although it requires a high computational time (because of more number of collision checking), can be very effective. In the following, we give the standard bridge test algorithm (Randomized Bridge Builder).
Algorithm: Randomized Bridge Builder (RBB)

For i ← 1 to K

q₁ ← Random Configuration(Uniform Distribution)
If Clearance(q₁) returns False
then q₂ ← Random Configuration(λₐ)
If Clearance(q₂) returns False
then q₃ ← mid point of q₁,q₂
If Clearance(q₃) returns True
then Insert q₃ to List G as a new milestone

Return(G)

Here "Clearance" is a collision checker which returns true if the sampled configuration is in $C_{free}$ and $λₐ$ is a multi-dimensional Gaussian distribution with a pre-specified variance $σ$. The performance of this algorithm depends heavily on the choice of $σ$. If one chooses $σ$ to be very small it takes a very long time and numerous collision checking before to come up with a point in $C_{free}$. Also a large $σ$ results in large number of redundant configurations. The computation time also increases with the increase in number of obstacles.

4. RANDOM WALK TOWARD SURFACE

Next we present our algorithm, which generates the sample points in the surface of the configuration space obstacle. The idea is, since in high dimension one does not know the suitable value of $σ$, and there is no a priori knowledge about narrow passage geometry, therefore the most idealistic approach will be to generate points on the surface of the obstacle. Note that in the Gaussian sampling scheme, one tries to generate points on the surface of the obstacles. But again an unsuitable value of $σ$ for a particular problem instance may take a very long time to generate a point in the narrow passage. Our algorithm is based on a simple philosophy, such as, once a sample point is generated within the obstacle one can perform a discrete random walk with a fixed length size $a$, until the particle comes out of the obstacle.

Algorithm: Random Walk Towards Surface (RWS)

For i ← 1 to K

q ← Random Configuration
If $q \in C_{free}$ then add to List L
else Flag ← 1
while Flag = 1
  $DAQ$ Select Random Direction
  $q'$ ← Random Walk $(q, p, a)$
  If $q' \in C_{free}$ then add $q'$ to the list L
  Flag ← 0
else $q \leftarrow q'$

Return L

The most important aspect of the algorithm is the reduction in collision checking computations. Both in RBB and Gaussian sampling strategy, to generate a point inside the obstacle, one has to check if the sampled random configuration belongs to either of $m, O'ₙ$'s. This increases with increase in number of obstacles. In our algorithm, once a point is generated inside obstacle $O'ᵢ$, it has to only check if the point belongs to only that obstacle $O'ᵢ$ or not. This reduces computation time drastically, as we shown on Table 1.

As in Gaussian sampling strategy, our algorithm generates unnecessary points on the surface of the obstacle which may be far away from critical regions. We accept this as a disadvantage of the algorithm, and propose a discrimination method to eliminate unnecessary points.

We define a heuristic distance $d(x, y) \propto \frac{1}{N}$ as threshold, where $N$ is the number of points. Once all the points are generated, for every point $p \in N$ a neighbor list is created which contains points that lie within $d(x, y)$. If all the neighbors originate from the same obstacle then $p$ is deleted. Thus following algorithm describes the process.

Algorithm: Elimination(L)

$N \leftarrow Size(L)$
For i ← 1 to N

$Q \leftarrow Neighbors(q(i), d)$
$I \leftarrow q(i).index$
$M \leftarrow Size(Q)$
For k ← 1 to M

if $I = Q(k).index$
  then $p \leftarrow p + 1$
If $p = M$
  then delete $q(i)$
else insert $q(i)$ to List $S$
Return $S$

This eliminates unnecessary points. We also try to provide an approximate analysis of RWS in terms of mean escape time $τ_{es}$ required to come out of the obstacle. Although our proposed random walk is discrete, but we approximate this with a continuous time diffusion process. This allows us to
state a much more simple expression for mean escape time $\tau_{ex}$. Since the expression for mean exit time is available in existing literature, therefore we only state the result. For a somewhat elementary proof, see Appendix. A more general discussion and in depth analysis of diffusion process can be found in [16]. In particular, the expected exit time from a ball of radius $R$ in $d$ dimension is given by

$$\tau = \frac{R^2}{d} \mathbb{E}[x^2], \quad x \in \mathbb{R}^d$$  \quad (1)

where $x$ is the location of initial position of particle from origin. Therefore, maximum of expected exit time is $\tau_{max} = \frac{R^2}{d}$. Now the question is, how many steps will be taken up to and including time $\tau_{max}$. We assume the waiting time between successive steps is finite and constant. We denote this by $\beta$. Therefore, the expected number of collision checking required before a particle comes out of domain $\Omega$ is,

$$N_{col} \sim \frac{R^2}{d\beta}$$  \quad (2)

Interestingly this indicates that as the dimension increases it becomes less and less probable for a random walker to return to the starting point as shown by Pólya [17]. This means particles once sampled within the obstacles, are more probable to quickly appear onto the obstacle surface as the dimension goes high.

5. COMPARISON RESULTS BETWEEN RWS AND RBB

We compare the samples generated in the critical region and time to generate them for both RBB and RWS algorithms in different critical environments, Fig 3 to Fig 8. We choose several benchmark problem, tested in a Pentium dual core processor and standard MATLAB runtime environment. Configuration space is $[0,100] \times [0,100]$. In case of RWS the random walk has a fixed step size of 1 m. The comparison results in Table 1 shows the capability of RWS algorithm compared to RBB. Note that with increase of $\sigma$, RBB may generate samples not in the critical region as shown in Fig 7. The differences in the computation time with different $\sigma$ clearly indicates the importance of proper $\sigma$ selection. With increase in number of obstacles, computations become slower in case of RBB as shown in Fig 7 (which also evident from Table 1).

6. RAPIDLY EXPLORING RANDOM TREE

Rapidly Exploring Random Tree (RRT) have been shown to be very effective in solving robot motion planning problems in complex configuration space with kynodynamic constraints. RRT was introduced in [18],[10] as an efficient data structure and sampling scheme to quickly search high dimensional spaces that have algebraic constraints (arising from obstacle) and differential constraints (arising from nonholonomy and dynamics). The algorithm incrementally builds a tree whose nodes are different configurations of the robot/vehicle. The edges of the graph correspond to the feasible path between the configurations. The key idea of RRT is to bias the exploration toward unexplored portions of the space by sampling configurations, and incrementally pulling the search tree toward the unexplored portion [19]. In the following we present the basic RRT algorithm.

Algorithm: Build-RRT($q_{init}$)

1. $T \cdot init(q_{init}) \triangleright$ Initialize tree $T$
2. For $k \leftarrow 1$ to $K$ do
   1. $q_{rand} \leftarrow$ Random Configuration
   2. $Ext(T, q_{rand})$
3. Return $T$

![Table 1: Comparison between RWS and RBB](image)

Table 1: Comparison between RWS and RBB
7. PREPROCESSING THE CONFIGURATION SPACE

Many variants and modifications of the basic RRT algorithm can be found in [20], [21], [22], [23] etc, in order to address exploration vs biasing issues, nearest-neighborhood queries, and probabilistic convergence. In [24], it is mentioned that different choice of metric in $C_{free}$ can lead to a drastic difference in the performance of RRT. Apart from this issues one of the biggest problem is the issue of exploration vs biasing. While many used an artificial bias in generating samples towards the goal [25], it is certainly not the best approach because of the potential problem of local minima. One variant is to pick up milestones in Probabilistic Roadmap method and try to push RRT to sequentially follow the PRM vertices [10]. But in general, for dynamical systems, the problem of reachability may occur, and one never knows whether a desired milestone is reachable or not a priori. In [20], [21] an adaptive biasing is used. There the focus is on the selection of the best among $k$-nearest neighbors, and adaptively increase or decrease the sampling bias. But again this approach depends on too many tuning parameters that depend on configuration space criticality.

The RRT algorithm has a known exponential bound on its run length tail probabilities. Its nearest-neighbor operation implies that individual iteration require increasing time. Therefore, it is reasonable to assume that run time tail probabilities may arise that are heavier than exponential. It is shown in [26], that two independent forward search RRT, solving the same query, can reach roughly the same level of progress at drastically different run lengths.

So far there is little proof on probabilistic completeness of RRT for arbitrary kinodynamic problem. In [10], a theorem states that for both convex and non-convex state space and for holonomic path planning problem, the probability, that the dispersion of the RRT vertices will be less than an arbitrary positive threshold, will converge to 1 as the number of vertices approaches infinity.

**Theorem [10]:** Suppose $x_{init}$ and $x_{goal}$ lie in the same connected components of a convex/non-convex, bounded open $n$ dimensional state space. The probability that an RRT constructed from $x_{init}$ will find a path to $x_{goal}$ approaches one as the number of RRT vertices approaches infinity.

Based on the above theorem, we realize that, identification of necessary segments of the configuration space (state space) that is worth searching sequentially for the goal, may generate a lower dispersion value for a given number of RRT vertices, that is, the solution can be found ‘quickly’. This motivates us to pre-process the configuration space to find most suitable regions to be sequentially explored before becoming close enough to $x_{goal}$. As an analogy, imagine RRT to be a big ship approaching a harbor and requiring the guidance of a small pilot ship which knows the different channels to approach having different quality measures. An RRT used for kinodynamic problem therefore requires a set of guiding milestones representing the key segments. These guiding milestones cannot be chosen using elementary PRM, because those may not be reachable from the current configuration.

Before describing our approach, we define the following:

1. **State Space:** A topological space $X \in [0, 1]^n$.
2. **Boundary Value:** $x_{init} \in X$ and $x_{goal} \in X$.
3. **Collision Detector:** A function, $D : X \rightarrow \{true, false\}$ that determines whether global constraints are satisfied for state $x$.
4. **Inputs:** A set $U$ which specifies the complete set of controls.
5. **Incremental Simulator** $A(\eta \Delta t)$ : Given the current state $x(t)$ and inputs applied over a time interval $\eta \Delta t$, the incremental simulator computes $x(t + \eta \Delta t)$.
6. **Metric:** A real valued function $\rho : X \times X \rightarrow [0, \infty]$.
7. **Reachable set:** $R(x^{k}, \hat{X}, \eta \Delta t) = \{x \in X \mid u \in U, x = x^{k} + \int_{0}^{\Delta t} f(x, u) dt \}$
8. **Voronoi partition of set $P$** Let $P := \{p_{1}, p_{2}, ..., p_{n}\}$ be a set of $n$ distinct points in $\mathbb{R}^{n}$, these points are the sites. The Voronoi diagram of $P$ as the subdivision of the space into $n$ cells, one for each site in $P$, with the property that a point $q$ lies in the cell corresponding to a site $p_{i}$ if and only if $\rho(q, p_{i}) < \rho(q, p_{j})$ for each $p_{j} \in P$ with $j \neq i$. The Voronoi partition of set $P$ is $V(P)$, Fig(2).

We assume no a priori knowledge about the configuration space (state space). By uniform sampling scheme we choose $N$ number of points in the free portions of the state space. The goal is to generate a non-uniform partition of the space. The suitable value of $N$ of course depends on the dimension and range of the problem instance. Next, we do a Voronoi partition of the space based on these $N$ registration points (sites), $V(P)$, according to the given metric $\rho$. The goal is to find a single step transition probability of a particle from region $i$ to region $j$ of the Voronoi partition. One problem may occur that, once we partitioned the state space, connectivity of a region or cell may be lost. Since the shape of the configuration space obstacle is unknown, the lowest grid resolution required to overcome the problem is also unknown. Only we can assume that the $C-$space obstacles $O_{t}$s having large measure are more likely to create problems for a specific $N$. It is clear that points on the surface of the obstacle generated in an uniform manner can reduce the problem of disconnectedness.
substantially. We use therefore, the RWS algorithm to find points on the surface of the obstacles. We call this as enrichment of the initial Voronoi cell distribution. The degree of reduction is dependent on the number $M$ of additional points generated.

We do a repartitioning using Voronoi grid of $N + M$ points. Next, using Monte-Carlo method we approximate the volume measure of the free portions of Voronoi grid cells, with a confidence level of 0.95 [27]. We generate $N'$ particles in $C_{\text{free}}$ by uniform distribution.

Once we have an approximate measure of the free volume in each cell $w_i$, we give random excitation input to the individual particle for a time period of $\eta \Delta t$. The incremental simulator gives us the position of the particle within a Voronoi cell. We then calculate the transition probability $P_{ij}$, defined as the probability that the particle at cell $i$ at time $t$, will go to cell $j$ at time $t + \eta \Delta t$. For all such grid cell we get the $\eta$ step transition probability matrix $P_{\eta}$. Assuming the time interval $T = \eta \Delta t$, we rename this $P_{\eta}$ as single step transition probability $P_{ij}$. Note that for the transition probability calculation, the underlying assumption of Markovian nature of the process exists. So

$$P_{ij} = \frac{\text{Number of particles that goes into cell } j \text{ from cell } i}{\text{Number of particles fall inside cell } i}$$

$$\sum_{i=1}^{N+M} P_{ij} = 1 \quad \forall j \in [1, 2, ..., N + M]$$

Once we have the transition probability of the entire grid regions we can calculate, such as, $P^\eta$ passage times from the Cell$_i$ containing the initial configuration to Cell$_j$ containing final configuration and most probable path $\gamma$ from Cell$_i$ to Cell$_j$. Assuming one such path exists, the most probable path $\gamma_m$ has the maximum probability over all such paths.

Next, we describe the dynamic programming procedure to find $\gamma_m$ for a given $V(P_{N+M})$.

**Algorithm for Most Probable Path:** To find the single best cell sequence $Q = \{q_1, q_2, ..., q_T\}$ ($q_1, q_2, ..., q_T$ are corre-

- $\delta_i(i) = \max_{q_1, q_2, ..., q_T} P(q_1, q_2, ..., q_T = i)$

that is, $\delta_i(i)$ is the best score along a single path at stage $t$ which ends at cell $S_i$. By induction

$$\delta_{t+1}(j) = \max_i \delta_t(i) P_{ij}$$

To retrieve the cell sequence, we need to keep track of the argument that maximized (4), for each $t$ and $j$. We do this via the array $\psi(j)$. The complete procedure for finding the best state sequence can now be stated as

1. Initialization

$$\delta_1(i) = \pi_{i \omega_i}$$

$$\psi_1(i) = 0$$

2. Recursion

$$\delta_t(j) = \max_{1 \leq i \leq N} \delta_{t-1}(i) a_{ij} w_i$$

$$\psi_t(j) = \arg \max_{1 \leq i \leq N} \delta_{t-1}(i) a_{ij} w_i$$

3. Termination

$$P^* = \max_{1 \leq i \leq N} \delta_T(i)$$

$$q_T^* = \arg \max_{1 \leq i \leq N} \delta_T(i)$$

4. Back Tracking

$$q_t^* = \psi_{t+1}(q_{t+1}^*)$$

$T = 1, T - 2, ..., 1$

It is important to choose the number of stages $T$ over which the maximization requires to be done. One can evaluate first passage characteristic from cell $i$ to cell $j$ and compute the mean first passage time. There are standard Monte Carlo based procedures that exist for finding approximate mean first passage time $T$ [27]. The interpretation of the most probable sequence cell is that these zones are to be sequentially searched by RRT. The Voronoi registration points of these cells are the guiding milestones.

8. IMPLEMENTATION ISSUES AND RESULTS

Although there is no guarantee that every time the above pre- processing of configuration space lead to the best sequence path (zone). The procedure clearly depends on

1. The non-uniform grid distribution.
2. The number of grid cells.
3. The number of points generated on the surface of the obstacles.
4. The criticality of the configuration space.
5. The reachable set $\mathcal{R}_k$.
While implementing one should carefully evaluate the transition probability matrix, as it is the heart of the whole procedure. Depending on grid distribution one may find cases where $P_{ij} = 1$ which indicates a transient region. There one may generate additional points with increased $\eta \Delta t$ and try to approximate $P_{ij}$. Also, there may be cases where $P_{ij} = 0$ because of small cell measure. In this case one can excite the corresponding cell registration point with random control input and thus approximate the value of $P_{ij}$. Depending on the configuration space criticality there may be large uncertainty in evaluating transition probability matrix. Inclusion of uncertainty in $P$ and generating a robust procedure for the method is our future concern. We have tested our method for path planning of nonholonomic mobile robots, although it can be applied to any other configuration space as well. The vehicle dynamics is defined by

$$\dot{x} = v \cos(\theta)$$

(12)

$$\dot{y} = v \sin(\theta)$$

(13)

$$\dot{\theta} = \frac{(v/L) \tan(\theta)}{2}$$

(14)

The configuration space is $\mathbb{R}^2 \times S^1$. We intentionally created 6 critical obstacle traps so that a conventional RRT requires much longer time to come to the goal region. For each of the 6 problem instances a most probable path is found by our algorithm. Once the sequential milestones are available one can guide the RRT using biased sampling towards the milestones. We used 3000-5000 samples to approximate the cell free volume and to calculate the transition probability. For problem instances A to E (Fig [9-13]) the start configuration is (50,50,0) while the goal configuration is (100,100,0). A limited position is (50,50,0) and goal configuration is (100,100,20) and problem instances A to E (Fig [9-13]) the start configuration is (0,0,0) while the goal configuration is (100,100,0). A limited number (usually 10-100, depending on the problem instance) of initial Voronoi cells ($N_{Vor}$) are created. Enrichment $N_9$ depends on the criticality of the problem instance. One can argue about the added computation required for this approach, but clearly the results shows it is worth investing in the extreme case.

9. CONCLUSIONS AND FUTURE WORK

We developed configuration space preprocessing techniques to improve the efficiency of sampling based planners like PRM and RRT in critical situation. The approaches are promising. In future our focus will be on forming a robust approach to tackle uncertainty in transition probability matrix calculation and consequently development of an improved pre-processing technique.

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REFERENCES

APPENDIX

We begin by assuming a random sample generated within an obstacle and calculate maximum of mean exit time max(τk). This is equivalent to evaluate the maximum mean exit time from a domain Ω, of a particle, which is showing brownian motion in a stationary fluid medium. If b(t, x) ∈ ℝ³ is the velocity of the fluid at the point x at time t, then a reasonable mathematical model for the position Xt of the particle at time t would be a stochastic differential equation of the form

\[ dX_t = b(t, X_t) + \sigma(t, X_t)W_t \]  

(15)

where, Wt ∈ ℝ³ is a white noise. The Itô interpretation of this equation is

\[ dX_t = b(t, X_t)dt + \sigma(t, X_t)dB_t \]  

(16)

where Bt is m-dimensional Brownian motion, Xt ∈ ℝ³, b(t, Xt) ∈ ℝ³, σ(t, Xt) ∈ ℝ³×m. We call b the drift coefficient and σ the diffusion coefficient.

We can associate a second order partial differential operator A to an Itô diffusion X. The basic connection between A and X is that A is the generator of the process X. Let \( X_t \) be a time-homogeneous Itô diffusion in ℝ³. The infinitesimal generator A of X is defined by

\[ Af(x) = \lim_{t \to 0} \frac{E^t[f(X_t)] - f(x)}{t} \]  

(17)

where, \( f: \mathbb{R}^d \to \mathbb{R} \) such that limit exists at \( x \). E^t define expectation on \( X_t \) with respect to probability measure.

Consider the stationary case of Itô diffusion when \( b = 0 \) and \( \sigma = I_d \) i.e.,

\[ dX_t = dB_t \]  

(18)

The generator of \( B_t \) is

\[ Af = \frac{1}{2} \sum \frac{\partial^2 f}{\partial x_i^2} \]  

(19)

where, \( f = f(x_1, x_2, ..., x_d) \) twice differentiable, that is, \( A = \frac{1}{2} \Delta \) where \( \Delta \) is the Laplace operator.

Now we state Dynkin’s Theorem [16]: Let f is a twice differentiable function. Suppose τ is the stopping time (escape time from a domain Ω, and \( E^\tau[f] < \infty \)) then

\[ E^\tau[f(X_\tau)] = f(x) + E^\tau[\int_0^\tau Af(X_s)ds] \]  

(20)

Now consider a Brownian motion \( B = (B_1, B_2, ..., B_d) \) starting at \( a = (a_1, a_2, ..., a_d) \) ∈ ℝ³ and assume \( |a| < R \). We define the first exit time as \( \tau_K \) of \( B \) from the ball

\[ K_R = \{ x \in \mathbb{R}^d; |x| < R \} \]  

(21)

According to the Dynkin’s theorem we choose \( f(x) = |x|^2 \) for \( |x| \leq R \), \( X = B \) and \( \tau = \sigma_k = \min(k, \tau_K) \), where \( k \) is any integer. So,

\[ E^\sigma[f(B_{\sigma_k})] = f(a) + E^\sigma[\int_0^{\sigma_k} \frac{1}{2} \Delta f(B_s)ds] \]  

(22)

\[ = |a|^2 + E^\sigma[\int_0^{\sigma_k} d \cdot ds] \]  

(23)

\[ = |a|^2 + d \cdot E^\sigma[\sigma_k] \]  

(24)

Letting \( k \to \infty \), and \( \tau_k = \lim \sigma_k < \infty \) a. s. and

\[ E^\sigma[\tau_k] = \frac{1}{d}(R^2 - |a|^2) \]  

(25)
Figure 3: Problem Instance A (a) RWS without elimination (b) RWS with elimination (c) RBB ($\sigma = 2$)

Figure 4: Problem Instance B (a) RWS without elimination (b) RWS with elimination (c) RBB ($\sigma = 2$) (d) RBB ($\sigma = 10$) (e) RBB ($\sigma = 5$) (f) RBB ($\sigma = 2$)

Figure 5: Problem Instance C (a) RWS without elimination (b) RWS with elimination (c) RBB ($\sigma = 20$) (d) RBB ($\sigma = 10$) (e) RBB ($\sigma = 5$) (f) RBB ($\sigma = 2$)

Figure 6: Problem Instance D (a) RWS without elimination (b) RBB ($\sigma = 2$)
Preprocessing of Configuration Space

Figure 7: Problem Instance D (a) RBB ($\sigma = 2$) (b) RBB ($\sigma = 10$)

Figure 8: Problem Instance E (a) RWS without elimination (b) RBB ($\sigma = 2$) (c) RBB ($\sigma = 5$)

Figure 9: Problem instance A: $N_{grid} = 20$, $N_{\eta} = 100$, 3000 samples

Figure 10: Problem instance B: $N_{grid} = 100$, $N_{\eta} = 30$, 5000 samples

Figure 11: Problem instance C: $N_{grid} = 10$, $N_{\eta} = 50$, 3000 samples

Figure 12: Problem instance D: $N_{grid} = 10$, $N_{\eta} = 100$, 5000 samples
Figure 13: Problem instance E: $N_{grid} = 10$, $N_q = 100$, 3000 samples

Figure 14: Problem instance F: $N_{grid} = 10$, $N_q = 30$, 3000 samples

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