Variance Reduction in Global Illumination with Monte Carlo Methods

Raymond Kim advised by Sanjoy Baruah

University of North Carolina at Chapel Hill

Abstract  Monte Carlo methods are a class of numerical algorithms that depend on repeated samples of random variables to obtain results; the numerical solution to the problem at hand can converge to the analytical solution by increasing the number of samples. However, as the problem becomes more complex, this procedure of reaching an accepted solution becomes inefficient: the problem may require higher orders of samples to achieve a single order of accuracy. Thus, variance reduction techniques exist to alter our method of performing the Monte Carlo method to achieve a faster convergence rate and a more accurate approximation given the same number of samples. In this paper we evaluate several variance reduction techniques of the Monte Carlo method and its application in the global illumination problem in computer graphics.

Introduction to Monte Carlo Methods

Background

The term Monte Carlo was first mentioned in the article "The Monte Carlo Method" by Metropolis and Ulam in 1949, just after the end of World War II [1]. The article considers a medium in which nuclear particles have the capability of producing other particles depending on their position and energy. This system is formulated by a set of integro-differential equations (equations involving both integrals and derivatives of the function) known as the Boltzmann equations; however, the classical methods used to deal with these equations were “extremely laborious and incomplete” such that the closed form solutions were unobtainable. Instead, a statistical approach is taken by sampling single chains of events upon which an analysis of the possible outcomes’ properties and distributions at various times can then be conducted.

The notion of solving problems through statistical means have existed long before the 20th century. A variant of Buffon’s needle experiment involving trials to estimate the value of π was published in the 18th century [2]. Even more so, the theoretical foundations of this method can be dated back to the 16th century with the formalization of the law of large numbers [3]. Since the Monte Carlo method by nature is repetitive, its popularity and practicality as a numerical technique only rose with the advent of modern computers. Note that performing the Monte Carlo method involves randomness which brings us to the topic of (pseudo)random number generators. Although it is out of this paper’s scope, it should not be disregarded and should be considered an important area of study.

Motivation

The previous section briefly describes the original motivation of the Monte Carlo method and the problem that was solved. As discussed, the sheer difficulty of the problem can be a factor in why the Monte Carlo might be preferred over other methods (in many cases, it is the only method that can provide a solution
to the problem of interest). Take for example, the evaluation of a definite integral. At one dimension, there are many different quadrature techniques to approximate the value of the integral, most of which perform better than the Monte Carlo method (which would require sampling the function at random points along the domain). These techniques can be extended to evaluate higher dimensional integrals; however, the number of function evaluations grow exponentially as the dimension increases. In this case, the Monte Carlo method can prove to be useful as its result does not depend in the dimensionality of the problem.

Another motivation of the Monte Carlo method is its applicability in different fields of study. It serves as a strong optimization technique when uncertainty is at play. We’ve seen how it can be used in mathematics and physical sciences, but this method can also be seen in engineering, finance, artificial intelligence, computer graphics, etc. The Monte Carlo method is also conceptually simple. When we want to find the expected value of a die, we can estimate this empirically by performing repeated simulations by rolling the die and combining our results to get an answer (this experiment can be found in the curriculum of many elementary schools).

In short, the Monte Carlo method is a simple and intuitive yet powerful method that can be used to solve the most challenging of problems.

Definition

There is no common consensus on the definition of a Monte Carlo method. Some have come up with classifications and distinctions while others have come up with qualifications on what the Monte Carlo method should entail. Putting these details and discussions aside, a nice general statement describes this method as “a numerical method of solving mathematical problems by the simulation of random variables.” [4] This will involve drawing samples from probability distributions. A general objective is to find the expectation $E$ of a transformation $f$ on a random variable $X$. This is equivalent in evaluating the integral as notated below:

$$ E[f(X)] = \int_{\Omega} f(x)p(x)dx $$

where $X$ falls under some probability density function $p(x)$ in some sample space $\Omega$. Since these two expressions are equivalent, we can use them interchangeably as done a few times in this paper.

Examples

In the following examples we discuss how the Monte Carlo method can be used to solve each problem. Note that we formally do not know if the estimator for each example converges to the right answer. Rather, we trust the correctness of each estimator using intuition and common sense. The next section hints at some of the underlying theory behind Monte Carlo methods, but will focus on variance reduction techniques.

"Hit or Miss" method. This first example is simple but does a nice job of showing how the Monte Carlo method can be used to solve a basic problem. Consider an arbitrary plane figure bounded in the region $[0, 1] \times [0, 1]$ with some area $S$. We
don’t need to know anything else about the figure, other than the fact that we can distinguish between points in its interior and exterior. Our objective is to find $\mathbb{E}[S]$. The spirit of the Monte Carlo method tells us that if we take random points in our sample space enough times, then we can use

$$\bar{S} = \frac{\# \text{ points in } S}{\# \text{ total points}}$$

as an accurate estimator for $S$. For example, let us sample 40 points as shown in Figure 1, pulled from a uniform distribution. There are 13 points inside $S$ and thus we can estimate $S$ by taking the ratio of the two. This gives us an estimated area of 0.325 whereas the actual area (knowing the coordinates that were used to plot this figure) is 0.3077. This gives us a relative error of about 5.6%. Taking 100 samples results in a relative error of 0.75%.

We can see that this estimator does indeed work. Intuitively, we also understand that as we sample more points, our estimator will become more accurate. Thix example can be extended into higher dimensions and gives us an idea of how to use Monte Carlo methods to approximate the volume of high-dimensional bodies.

**Simulation of a Mass Serving System.** Consider a system of $n$ lines that service incoming requests arriving that moments

$$T_1 < T_2 < \cdots < T_k < \cdots$$

where the $k$th request comes at time $T_k$ [4]. When the first request comes in, naturally, all the lines are free. Hence, the first line will begin servicing the request for an arbitrary time $t_h$. When the $k$th request comes in, the first open line will start serving the request, but if all are busy then the $k$th request is considered rejected. Our goal is to find the expected number of requests that are satisfied and rejected. Analytically, solving this may be tedious for a large number of requests. If we draw $t_h$ from a distribution, then our problem may become nontrivial and finding a solution becomes much harder. We can apply the Monte Carlo method here to reach our objective; if we maintain a timer $t$ we can essentially iterate through all $T_i$, updating $t$ as we go and keeping count of requests that are satisfied and rejected, so that in a single simulation, we get the number of requests that are satisfied and rejected. Our Monte Carlo estimator can simply be the sample mean of such simulations; by running it multiple times and taking the average, we can reach an estimate of the expected values we are interested in.

**Evaluating a Definite Integral** On a more analytical end, let us try to find the integral of a simple function $f$ where we define $f$ to be

$$f(x, y) = \exp(\sin(8x^2 + 3x) + \sin(11y)).$$

We want to find

$$I = \int_0^1 \int_0^1 f(x, y) \, dx \, dy = \int_0^1 \int_0^1 \exp(\sin(8x^2 + 3x) + \sin(11y)) \, dx \, dy.$$

In a more graphical sense, we want to find the volume created by this function over the unit square. Intuitively, we can define our estimator to be

$$\bar{I} = \frac{1}{N} \sum_{i=1}^{N} f(X_i, Y_i).$$
where $X_i$ and $Y_i$ are random variables drawn from the uniform distribution $U(0, 1)$. In this sense, we can estimate $I$ by repeated sampling the function at different points throughout the domain. Here we see a

**Figure 2.** Heatmap of $f$ (left) and 100 samples of $f$ (right)

heatmap of $f$ in the figure to the left and the result of 100 samples in the figure to the right. Although we do not provide a complete numerical analysis, we see that we calculate $\tilde{I}$ by taking the average of all 100 sample values.

**Variance Reduction Techniques in Monte Carlo Methods**

**Premise**

There are two theorems that are fundamental to the Monte Carlo method, and the reason why we can expect it to work. The first is stated as follows [5]:

**Strong Law of Large Numbers.** Let $X_1, X_2, \ldots$ be pairwise independent identically distributed random variables with $E[X_i] < \infty$. Let $E[X_i] = \mu$ and $Y_n = X_1 + \cdots + X_n$. Then, $Y_n/n \to \mu$ almost surely as $n \to \infty$.

Here we see that essentially as the number of samples we take approaches infinity, the sample mean approaches the true expected values. In the previous examples, we used the sample mean as our estimator; by this theorem we know that our this will converge to the correct answer and our estimator is indeed accurate. The second theorem fundamental to studying Monte Carlo methods stated below:

**Central Limit Theorem (Lindeberg-Levy).** Suppose $\{X_1, X_2, \ldots\}$ is a sequence of independently and identically distributed random variables with $E[X_i] = \mu$ and $\text{Var}(X_i) = \sigma^2 < \infty$. As $n \to \infty$,

$$\sqrt{n}\left(\frac{1}{n} \sum_{i=1}^{n} X_i\right) - \mu \to \mathcal{N}(0, \sigma^2).$$

In otherwords, for $\{X_1, X_2, \ldots, X_n\}$ and $Y_n = X_1 + \cdots + X_n$, if $n$ is sufficiently large, then $E[Y_n] = n\mu$ and $\text{Var}(Y_n) = n\sigma^2$. 

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Simply put, the Central Limit Theorem states that the sum of a large number of identical and independently distributed random variables will approximately fall under a normal distribution. The two theorems are very similar in nature, but the difference between the Law of Large Numbers and the Central Limit Theorem is that the latter tells us the shape of the distribution as well, which is helpful when evaluating variance.

Since the Monte Carlo method is based upon random variables, it is natural to expect variance in our results. As mentioned earlier, the objective of the basic Monte Carlo method is to find some expectation $\mathbb{E}[X]$ of a random variable $X$ with variance $\sigma^2$; we do so by collecting $n$ samples, $X_1, \ldots, X_n$, and using the sample mean as our estimator

$$\bar{X}_n = \frac{1}{n} \sum_{i=1}^{n} X_i.$$ 

We find the variance of the basic estimator using the Central Limit theorem as follows:

$$\text{Var}(\bar{X}_n) = \text{Var} \left( \frac{1}{n} \sum_{i=1}^{n} X_i \right) = \frac{1}{n^2} \sum_{i=1}^{n} X_i = \frac{1}{n^2} n \sigma^2 = \frac{\sigma^2}{n}.$$ 

The rate of convergence that signifies how fast the error is reduced is measured through the standard deviation, which means that the rate of convergence for the basic Monte Carlo estimator becomes $O(1/\sqrt{n})$. As we can see, the most obvious way to reduce our variance is by increasing $n$. Understandably, this does not result in a very fast convergence rate and can be inefficient when comparing the cost necessary to generate our samples versus the reduction in variance. Thus, we want to find methods in which we can alter our sampling method or come up with a new estimator so that the error converges faster than our basic case.

**Antithetic Variates**

Using antithetic variates as a means of variance reduction involves introducing a negative correlation within our samples [6]. Similar to our previous examples, suppose we are trying to find $\mathbb{E}[X]$. Take $X_i$ to be a random variable among the samples we take in the Monte Carlo scheme, where $\bar{X}_n$ was used as our basic estimator. Assume the total number of samples $n$ to be even so that we let $n = 2m$ for some $m \geq 1$. We define $Y_i$ as the average between $X_{2i-1}$ and $X_{2i}$ for $i = 1, 2, \ldots, m$ and use $Y_m$ to be our new estimator. Note the following:

$$\bar{X}_n = \frac{1}{2m} \sum_{i=1}^{2m} X_i = \frac{1}{m} \sum_{j=1}^{m} Y_j = Y_m.$$ 

Here we see that the two estimators are equivalent, and thus $Y_m$ can be used to find $\mathbb{E}[X]$. We know that

$$\text{Var}(Y_i) = \text{Var} \left( \frac{X_{2i-1} + X_{2i}}{2} \right) = \frac{1}{4} \left( \text{Var}(X_{2i-1} + X_{2i}) \right) = \frac{1}{4} (\sigma^2 + \sigma^2 + 2 \text{Cov}(X_{2i-1}, X_{2i}))$$

$$= \frac{1}{2} (\sigma^2 + \text{Cov}(X_{2i-1}, X_{2i})) = \frac{1}{2} (\sigma^2 + \text{correlation}).$$ 

The coefficient is expected as there are half as many $Y_i$ as there are $X_i$. Thus, if we are able to pair up samples of $X_i$ such that the two are negatively correlated, then our covariance value becomes negative and our overall variance is reduced.

There are several drawback to this method. First, we need to come up with a method of generating good antithetic samples. Sadly, there isn’t a general method of doing so and therefore the extent of the effectiveness
of this method is based upon an entirely different issue. There are problems in which generating antithetic pairs are simple and straightforward; however, in most cases it will take some effort in doing so. Another drawback is that if the sample pairs are poorly picked, then the correlation value becomes positive and our variance increases. This is natural since picking similar values will lower the amount of ‘information’ we get compared to the ‘information’ from two independently drawn samples. Thus, we need to be careful in how we choose our antithetic samples when using this as a variance reduction technique.

Importance Sampling

Importance sampling alters the way we choose our samples. In all of our previous scenarios, we assume that the samples are drawn from a uniform distribution. The key idea behind importance sampling is choosing samples from a probability density function that reflects the "importance" of a sample’s contribution [4]. Assume that we want to evaluate the integral of some function $f(x)$ in some space $A$. If we let our probability density function be denoted by $p(x)$, then we want $p$ to be as close to $f$ as possible. To see how this works, we redefine the problem to be the following:

$$I = \int_{x \in A} f(x) \, dx = \int_{x \in A} \frac{f(x)p(x)}{p(x)} \, dx = \int_{x \in A} \frac{f(x)}{p(x)} p(X) \, dx = E \left[ \frac{f(X)}{p(X)} \right].$$

To find this value, we simply define an estimator to be

$$\tilde{f}_n^p(X) = \frac{1}{n} \sum_{i=1}^n \frac{f(X_i)}{p(X_i)}$$

where $X_i$ is drawn from $p(x)$. It has been shown that the variance is reduced most when $p(x) \propto |f(x)|$ [4]. This can be easily checked by allowing $p(x) = c f(x)$ for a constant $c$; taking the variance of the estimator results in evaluating the variance of a constant $1/c$ which is 0. When solving for $c$, we solve the integral

$$\int_{x \in A} p(x) \, dx = \int_{x \in A} c f(x) \, dx = 1$$

which becomes the problem we are trying to solve. Finding the best probability density function is extremely difficult, as the problems we use Monte Carlo methods to solve involve complex or black-box functions. It’s often impractical to find the "perfect" $p$, so we usually settle for finding a $p$ that resembles $f$. Similar to the previous variance reduction technique, if $p$ is chosen poorly, this can lead to an increase in variance (consider the case where $p(x) = 1/f(x)$).

Stratified Sampling

As the name may imply, stratified sampling involves partitioning the sample space into different strata [7]. We find the expectation of each local strata, then combine the results to estimate our overall expectation. Again, let us try to estimate $\mu = E[X]$. Instead of taking $n$ samples, we take $l$ samples from $m$ different strata so that $n = lm$ and let $X_{ij}$ be the value of the $i$th sample from the $j$th strata [9]. We define the estimator in the $j$th strata and our overall estimator to be

$$\bar{X}_j = \frac{1}{l} \sum_{i=1}^l X_{ij}, \quad \bar{X}_m = \frac{1}{m} \sum_{j=1}^m \bar{X}_j$$
Allow the expectation and variance of the random variables per strata be denoted as $\mu_j$ and $\sigma_j^2$ respectively. Using these definitions, we evaluate the variance of our estimator to be

$$\text{Var} \left( \bar{X}_m \right) = \text{Var} \left( \frac{1}{m} \sum_{j=1}^{m} \bar{X}_j \right) = \frac{1}{m^2} \sum_{j=1}^{m} \text{Var} \left( \bar{X}_j \right) = \frac{1}{m^2} \frac{1}{l} \sum_{j=1}^{m} \sigma_j^2 = \frac{1}{n} \frac{1}{m} \sum_{j=1}^{m} \sigma_j^2.$$ 

We similarly evaluate the variance for the basic estimator $X$ in terms of the $j$ strata:

$$\text{Var} \left( X \right) = E \left[ X^2 \right] - \mu^2 = \frac{1}{m} \sum_{j=1}^{m} E \left[ X^2 \mid X \text{ from strata } j \right] - \mu^2 = \frac{1}{m} \sum_{j=1}^{m} (\mu_j^2 + \sigma_j^2) - \mu^2 = \frac{1}{m} \sum_{j=1}^{m} (\mu_j - \mu)^2 + \sigma_j^2 \geq \frac{1}{m} \sum_{j=1}^{m} \sigma_j^2.$$ 

It is clear to see that the variance of the stratified estimator is less than the variance of the basic estimator. Intuitively, through stratified sampling we are reducing the variance that can exist between each strata. This specific type of variance can be a result of an issue known as sample clumping where randomly generated samples are clustered in one area; by partitioning the sample space we guaranteed some measure of spread about our space. This is visualized in the figure below.

**Figure 3.** Uniformly distributed sampling (left) vs. stratified sampling (right)
The Quasi Monte Carlo Method

This variance reduction technique is a bit different from the rest, as we do not depend on taking random samples from a distribution. Instead, we use low discrepancy sequences to generate our samples. A low discrepancy sequence is a deterministic sequence that share properties of random variables so that each element is, as the term quasi is defined, seemingly random. These sequences are generated using prime numbers and different number bases; a few famous ones are the Halton sequence, Hammersley set, and the Sobol sequence. The rate of convergence for error using the Quasi Monte Carlo [8] is calculated to be

\[ O\left(\frac{(\log n)^s}{n}\right) \] for some dimension \( s \).

When compared to the convergence for the basic Monte Carlo method, we can see an improvement when \( n \) is high and \( s \) is relatively low.

**Figure 4.** Uniformly distributed points (left), 2D Halton sequence (middle), equidistant points (right)

The Global Illumination Problem

In global illumination, our objective is to accurately capture the light within an environment to generate realistic images. From a light source, light rays will intersect with the scene, bounce around, and eventually reach the camera where the image can then be formed. Global illumination involves the two types of illumination: direct and indirect illumination. Direct illumination occurs when a position in the environment is directly exposed to the light source whereas indirect illumination occurs when a position in the environment is exposed to light that has been reflected from other surfaces. This problem has been formalized into what is known as the rendering equation [10,11]:

\[
L_o(x, \omega_o, \lambda, t) = L_e(x, \omega_o, \lambda, t) + \int_{\Omega} f_r(x, \omega_i, \omega_o, \lambda, t) \ L_i(x, \omega_i, \lambda, t) \ (\omega_i \cdot n) \ d\omega_i
\]

where \( x \) is the location in space, \( \omega_o \) is the direction of the outgoing light, \( \lambda \) is the wavelength of light, \( t \) is time, \( \omega_i \) is the negative direction of the incoming light, and \( n \) is the surface normal at \( x \). The BRDF stands for the bidirectional reflectance distribution function which describes how the light behaves when it
hits the surface. The domain of of the integral is the hemisphere oriented about \(\mathbf{n}\) and we take the integral with respect to all incoming light rays.

Our interest lies in evaluating the integral in the right hand side of the rendering equation. Doing so analytically is impractical as the scene can be arbitrarily complex, so we use the Monte Carlo method to obtain an approximation. This requires a method of sampling different \(\omega_i\) about \(\Omega\), taking multiple samples, performing the appropriate function evaluations, and calculating the sample mean to use as our estimator. We now discuss two common methods used to do so: ray tracing and path tracing. Ray tracing establishes the foundations of how the rendering will work, then path tracing extends upon that idea to solve the rendering equation.

**Ray Tracing**

The term "ray tracing" has evolved into a very broad term, but we will referring to the basic method proposed by Whitted in 1979 [12]. In ray tracing, we start from the camera (the cone in Figure 5) and intersect viewing rays through the image plane and into the scene. This image plane is a figurative plane that the scene will project onto (and will be our resulting image on a pixel by pixel basis). When the ray intersects the scene at some position \(\mathbf{x}\), we trace the ray from \(\mathbf{x}\) to the light source(s) (cube in Figure 5) in the scene - if the ray is uninterrupted, then we include that light source when determining the amount of light gathered at \(\mathbf{x}\) (which determines the color for the pixel in the image). If the ray is interrupted (e.g. if the viewing ray intersects the darkened area on the bottom plane of Figure 5), we ignore the light source when calculating the light at that point which results in a shadow effect. A ray tracer can be extended to be recursive: in doing so,

![Figure 5. Camera setup in a scene](image)

we can achieve effects like reflection or perhaps even some basic caustics. However, one of the downsides to ray tracing is that only direct lighting can be computed; this limits the features of our rendering to the ones mentioned previously. There exist shortcuts and workarounds that can be applied to our basic ray tracer in order to get more intricate details; however, this requires indepth knowledge about the scene and an artistic eye to imitate such realism. Another requirement for a ray tracer are intersection functions for each object in the scene. Whether or not the objects were created with a modeling-tool in a third party software or
defined analytically, it is required to know where each ray intersects with the object given any orientation.
As the scene becomes more complicated, this intersection function becomes costly which is a reason why ray
tracing has been mostly used for offline rendering. Other rendering methods exist in cases where latency is
an important issue (e.g. video-games, VR displays). With the rise in GPU’s and parallel processing power,
real-time raytracing has become feasible, allowing for beautiful renderings of virtual worlds.

Path Tracing

The second method, path tracing, is similar to that of ray tracing but has more of a probabilistic aspect
[10]. Before we discuss the algorithm and its implementation, we first need to understand the behavior of
light in a scene and indirect lighting. On a bright sunny day, it is obvious to see that the shadow of a tree
is not pitch black. This ambient lighting comes from what we call indirect illumination: light from a source
reaches an object and scatters around the environment, providing light to other objects in the scene. In the
example of the tree, the light ray may hit a nearby stone and get reflected into the leaves and then reach
ground where the tree’s shadow is located. If we relate back to the rendering equations, this is why we take
the integral with respect to the incoming light rays about the hemisphere.

Figure 6. Scattering of light rays at $x$

With this behavior in mind, path tracing makes an attempt at simulating the bouncing behavior of light.
Forward path tracing starts from the light source and follows the paths of light into the scene, keeping
track of all the light rays that reach our camera. However, this is really inefficient which is why we consider
backwards path tracing. Similar to ray tracing, we from the camera and shoot viewing rays into the scene.
However, instead of shooting a single ray through each pixel, we send a large number of rays through the
image plane. This number is often referred to as samples per pixel (SPP) and is a common metric for path
tracers; the number of samples can reach into the tens of thousands (resulting in a better image but slower
runtime). Next, for each ray, we find the intersection of the ray and the scene at some location $x$. In ray
tracing we checked whether or not $x$ is exposed to any light. However, in path tracing, we bounce the ray
randomly within the hemisphere about the surface normal (the domain of our integral) as depicted in Figure
6. Of course, how we determine our reflected ray depends on the material and the BRDF at $x$: if the surface
is completely reflective then we can calculate the appropriate mirrored ray. If the surface is not perfectly
reflective, then we can choose a random ray that is biased towards the general direction of the mirrored
ray. Finally if the surface was perfectly diffuse, then we send the ray in a uniform random direction about
the hemisphere. Finally, if the viewing ray intersects with the light source, we can factor this back into the
original radiance value of $x$. If the viewing ray never intersects with the light source given some upper bound
of bounces, then it is left alone.

Ideally, we want to take multiple samples per bounce and take the average to get a better estimate of the
light collected at location $x$. However, in practice, we usually only shoot a single ray off the surface because
otherwise the runtime will exponentially increase (for each of the thousands of original rays, we create \( n \) new rays). If we assume that we have the computing power to sample the hemisphere at multiple locations even at high SPP, then it is essential to use a quality Monte Carlo estimator to evaluate our integral.

### Variance in Global Illumination

We see how Monte Carlo methods are used in path tracing and in solving the rendering equation, but the problem we focus on is the variance associated with approximating the integral. In computer graphics, variance comes in the form of visual artifacts: noise and aliasing. Noise resembles the random pixel pattern from analog TV sets and is produced through the use of random numbers. In path tracing, if we use a low SPP value then there can be a high likelihood that a majority of those samples will miss the light source and return a darker radiance than the true value. This is why images with low SPP values tend to look darker in general.

When it comes to variance reduction techniques in global illumination, stratified sampling is one of the more popular methods to reduce noise. Importance sampling and antithetic variates requires an in-depth knowledge about the scene we are rendering. Use of antithetic variates would require us to find negatively correlated sample pairs which is difficult for a single scene. Even more so, in the film industry, the scene we are trying to render is constantly changing which makes importance sampling and antithetic variates even more difficult to use. On the other hand, stratified sampling would be used to partition the hemisphere into subregions from which samples will be taken. There is no need for knowledge about the scene we are trying to render which is extremely convenient. Quasi-Monte Carlo methods have become an area of interest, as it can have a faster convergence rate due to the low dimensionality of the problem [12]. A benefit of Quasi-Monte Carlo methods in global illumination is that it can visually produce less noise, but can give way to aliasing due to the periodic nature of low discrepancy sequences. Aliasing can be identified by unusual or out-of-place patterns in our final render.

### Sampling Technique Proposal

For the remainder of this report, I propose a proof of concept for a method using low-discrepancy sequences to generate samples in our hemisphere. Generating these samples is not a trivial task, as it is incorrect to use uniform random variables as parameters for our spherical coordinates and expect a uniform distribution about the surface of the hemisphere. This brings us to my main objective: to sample the hemisphere using low discrepancy sequences while preventing any potential aliasing artifacts that may incur by introducing a variability factor in the sampling process. I make the assumption that we are sampling a perfectly diffuse surface (and thus our samples are generated uniformly about the surface of the hemisphere). We assume the hemisphere has radius 1 and is centered at the origin with the surface normal vector oriented upwards.

We first generate a 3D and 2D low-discrepancy sequence cube and square oriented about the origin and with length values of 2 (that is, an inscribed sphere will have radius 1). Call these set of points \( A \) and \( B \). For \( n \) samples, generating these values will have a runtime of \( O(n^3) \) and \( O(n^2) \) but is only used at the beginning and is a constant factor. For each sample in \( A \), if \( |A_i| \geq 1 \), we discard the sample and continue. By doing so, we start from a point cloud in a cube and transform it into a sphere. We perform a similar process to \( B \), discarding those whose length is greater than 1. Once we are left with valid values for \( A \) and \( B \), we normalize \( A \) so that the points are projected onto the surface of the sphere. If the \( A \) lies on the lower hemisphere, we simply find its reflected value about the origin. At this moment, our samples are evenly spread out about.
the upper hemisphere. Next, we calculate a unit vector \( \mathbf{u} \) that lies in the plane orthogonal to \( A_i \). We use an arbitrary \( \mathbf{u} \) for the time being. We let \( \mathbf{v} \) be the cross product of \( \mathbf{u} \) and \( A_i \), which gives us a vector that is orthogonal to \( \mathbf{u} \) and \( A_i \). We want to adjust the lengths of \( \mathbf{u} \) and \( \mathbf{v} \) depending on the density of the samples - if samples are generated pretty closely then our \( \mathbf{u} \) and \( \mathbf{v} \) should be low to effectively "cover" the distance between each sample points. Using \( \mathbf{u} \) and \( \mathbf{v} \) as a basis of sorts, our new sample is calculated by evaluating

\[
A_i + B_{i,x} \mathbf{u} + B_{i,y} \mathbf{v}
\]

where \( B_{i,x} \) and \( B_{i,y} \) is the \( x \) and \( y \) values of our sample \( B_i \). In doing so, we a method of sampling our hemisphere using low discrepancy sequences but add in a variability factor. The runtime of this method will be similar to the speed up from using Quasi-Monte Carlo methods and should reduce the possibility of aliasing artifacts.

**Discussion**

Potential problems that may occur is the amount of normalizing we are doing in the process. Square roots are expensive operations on a low level, and performing this billions of times can add up and become a significant overhead. Another issue is being able to realistically test and analyze this method; could take possibly days to generate a single render, depending on the number of samples we take. Finally, the periodicity of low discrepancy sequences can lead to unexpected effects. The transformations we perform on each sample can unknowingly lead to patterns which will inaccurately sample the hemisphere.

**Future Work**

For future work, the most immediate step would be to implement this process and conduct numerical tests to make sure of its viability. The transformation of low discrepancy sequences can potentially be an interesting area of study, since they are ultimately deterministic despite being quasirandom. It would be nice to conduct a rigorous study of error bounds and discrepancy, providing a formal argument for the correctness of the this method.
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