Diffraction Shader

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Inhaltsverzeichnis

1	Intr	$\mathbf{roduction}$	on					4			
	1.1	Motiva	ation					4			
	1.2	Goals						4			
	1.3	Previo	us work					5			
	1.4										
2	Theoretical Background 6										
	2.1	Definit	${f cions}$					6			
		2.1.1	Diffraction					6			
		2.1.2	Electromagnetic radiation					7			
		2.1.3	Radiant energy					8			
		2.1.4	Radiance					8			
		2.1.5	Signal					9			
		2.1.6	Fourier Transformation					9			
		2.1.7	Convolution					10			
		2.1.8	BRDF					10			
		2.1.9	Spectral Rendering					11			
		2.1.10	Taylor Series					11			
	2.2		Basis: J.Stam's Paper about Diffraction Shader					11			
	2.3	Deriva						14			
		2.3.1	BRDF formulation					14			
		2.3.2	Relative BRDF					15			
		2.3.3	Taylour approximation for BRDF					18			
		2.3.4	Sampling: Gaussian Window					21			
		2.3.5	Aplitude smooting					$\frac{1}{22}$			
		2.3.6	Final Expression					26			
3	Implementation 28										
	3.1	Setup						29			
	3.2	-	nputations in Matlab					29			
	3.3		amework					29			
	3.4		LSL Diffraction Shader								
4	Dat	a Acqu	uisition and Evaluation					32			
	4.1	-	etion Grating					32			
	4 2		Skin Parameters					33			

5	Res	ults	34
6	Con	nclusion	35
	6.1	Further Work	35
		6.1.1 Sources	35

1 Introduction

rendering diffraction colors caused by coherent (as opposed to statistical) surface micro-geometry. tables containing the result of the spectral integration predicted by the model of Stam. The theory is applied to render diffraction caused by acquired geometric models of the microstructure of snake skin, so as to reproduce the irridescent effects displayed by these animals.

1.1 Motivation

In Nature, coloring mostly comes from the inherent colors of materials but sometimes colorization has a pure physical origin such as the effect diffraction or interference of light. Both phenomenon are causing the so called structural coloration, which is the production of color through the interaction of visible light with microscopically structured surfaces. Color production is die to wave interference with quasiperiodic structures whose periodicity leads to interaction with visible light. Therefore we perceive color when the different wavelengths composing white light are selectively interfered with by matter (absorbed, reflected, refracted, scattered, or diffracted) on their way to our eyes, or when a non-white distribution of light has been emitted. In animals, such as feathers of birds and the scales of butterflies, interference is created by a range of photonic mechanisms, including diffraction grating, selective mirrors, photonic crystals. The connection between microscopic structures and coloration has been observed by Robert Hooke in the early seventeenth centrury. The discovery of the wave nature of light led to the conclusion that the cause for the coloration lies in wave interference.

In the field of computer graphics, many researchers have been attempted rendering of structural colors but most of the techniques so far, however, are either too slow for interactive rendering or rely on simplofying assumption, like modeling light as rays, to achieve real-time performance.

1.2 Goals

The purpose of this thesis is to render realtime structural colors caused by the effect of diffraction on differentr biological structures. We focus on structural colors generated by diffraction grating, in particular our approach applies to surfaces with quasiperiodic structures at the nanometer scale that can be represented as heighfields. such structures are found on the sends of snkaes,

wings of butterflies or the bodies of various insects. we focus on sknake skins and we acquirednanoscale heightfields of different snake sheds using atomic force microscopy.

In oder to achieve this rendering purpose we will rely J. Stam's formulation of a BRDF which basically describes the effect of diffraction on a given surface assuming one knows the hightfield on this surface and will further extend it. In our case, those heightfields are small patches of the microstructured surface (in nano-scale) of AFM taken snake skin patches provided by GENEVA.

In his Paper, J. Stam assuming distribution on his heightfields whereas we require a an explicit provided hightfield of the surface or at least a small patch. Therefore, this work can be considered as an extension of J. Stam's derivations for the case one is provided by a explicit height field on a quasiperiodic structure. Since one goal of this work is to render in realtime, we have to perform also precomputations which will require us to slightly modify Stam's main derivation.

1.3 Previous work

stam, hooke, see our paper, see stams paper, see own research.

Robert Hooke = observed connection between microscopic structures and colorisation wave nature of light led to conclusion that the cause for the coloration lies in wave interference.

1.4 Overview

outline chapters of thesis and show what is about.

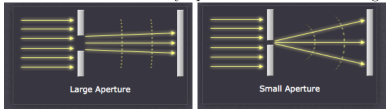
2 Theoretical Background

2.1 Definitions

2.1.1 Diffraction

Diffraction is a purely wave-like phenomenon which cannot be modeled using the standard ray theory of light. Interesting diffraction phenomena, however, occur mostly when the surface detail is highly anisotropic, viz. non-isotropic Interference produces colorful effects due to the phase differences caused by a wave traversing thin media of different indices of refraction Diffraction occurs when the surface detail is comparable to the wave- length of light

Light rays passing through a small aperture will begin to diverge and interfere with one another. This becomes more significant as the size of the aperture decreases relative to the wavelength of light passing through, but occurs to some extent for any aperture or concentrated light source.



Since the divergent rays now travel different distances, some move out of phase and begin to interfere with each other — adding in some places and partially or completely canceling out in others. This interference produces a diffraction pattern with peak intensities where the amplitude of the light waves add, and less light where they subtract. If one were to measure the intensity of light reaching each position on a line, the measurements would appear as bands similar to those shown below.



Diffraction refers to various phenomena which occur when a wave encounters an obstacle. In classical physics, the diffraction phenomenon is described as the apparent bending of waves around small obstacles and the spreading out of waves past small openings.

While diffraction occurs whenever propagating waves encounter such changes, its effects are generally most pronounced for waves whose wavelength is roughly similar to the dimensions of the diffracting objects. If the obstructing object provides multiple, closely spaced openings, a complex pattern of varying intensity can result. This is due to the superposition, or interference, of different parts of a wave that travels to the observer by different paths (see diffraction grating).

The effects of diffraction are often seen in everyday life. The most striking examples of diffraction are those that involve light; for example, the closely spaced tracks on a CD or DVD act as a diffraction grating to form the familiar rainbow pattern seen when looking at a disk

In optics, a diffraction grating is an optical component with a periodic structure, which splits and diffracts light into several beams travelling in different directions. The directions of these beams depend on the spacing of the grating and the wavelength of the light so that the grating acts as the dispersive element.

The relationship between the grating spacing and the angles of the incident and diffracted beams of light is known as the grating equation.

Fresnel and Frauenhofer diffraction frauenhofer diffraction = infinite observation distance. In multiple slit patterns each slit produces a diffraction pattern. Hence, multiple slit interference pattern is superimposed over single slit diffraction pattern.

2.1.2 Electromagnetic radiation

WIKI: http://en.wikipedia.org/wiki/Electromagnetic_radiation

Electromagnetic radiation (EM radiation or EMR) is one of the fundamental phenomena of electromagnetism, behaving as waves propagating through space, and also as photon particles traveling through space, carrying radiant energy.

EMR is characterized by the frequency or wavelength of its wave. The electromagnetic spectrum, in order of increasing frequency and decreasing wavelength, consists of radio waves, microwaves, infrared radiation, visible light, ultraviolet radiation, X-rays and gamma rays.

The eyes of various organisms sense a somewhat variable but relatively small range of frequencies of EMR called the visible spectrum or light. Higher frequencies correspond to proportionately more energy carried by each photon; for instance, a single gamma ray photon carries far more energy than a single photon of visible light.

EMR carries energy—sometimes called radiant energy—through space continuously away from the source

In general, EM radiation (the designation 'radiation' excludes static electric and magnetic and near fields) is classified by wavelength into radio, microwave, infrared, the visible spectrum we perceive as visible light, ultraviolet, X-rays, and gamma rays. Arbitrary electromagnetic waves can always be expressed by Fourier analysis in terms of sinusoidal monochromatic waves, which in turn can each be classified into these regions of the EMR spectrum.

The behavior of EM radiation depends on its frequency. Lower frequencies have longer wavelengths, and higher frequencies have shorter wavelengths, and are associated with photons of higher energy. There is no fundamental limit known to these wavelengths or energies, at either end of the spectrum, although photons with energies near the Planck energy or exceeding it (far too high to have ever been observed) will require new physical theories to describe.

2.1.3 Radiant energy

 $WIKI: http://en.wikipedia.org/wiki/Radiant_energy$ Radiant energy is the energy of electromagnetic waves.[1] The quantity of radiant energy may be calculated by integrating radiant flux (or power) with respect to time and, like all forms of energy, its SI unit is the joule. The term is used particularly when radiation is emitted by a source into the surrounding environment. Radiant energy may be visible or invisible to the human eye.

The term "radiantenergy" is most commonly used in the fields of radiometry, solar energy, heating and lighting,

2.1.4 Radiance

WIKI: http://en.wikipedia.org/wiki/Radiance Radiance and spectral radiance are measures of the quantity of radiation that passes through or is emitted from a surface and falls within a given solid angle in a specified direction. They are used in radiometry to characterize diffuse emission and reflection of electromagnetic radiation. In astrophysics, radiance is also used to quantify emission of neutrinos and other particles. The SI unit of radiance is watts per steradian per square metre (W·sr-1·m-2), while that of

spectral radiance is $W \cdot sr-1 \cdot m-2 \cdot Hz-1$ or $W \cdot sr-1 \cdot m-3$ depending on whether the spectrum is a function of frequency or of wavelength.

Radiance characterizes total emission or reflection. Radiance is useful because it indicates how much of the power emitted by an emitting or reflecting surface will be received by an optical system looking at the surface from some angle of view. In this case, the solid angle of interest is the solid angle subtended by the optical system's entrance pupil. Since the eye is an optical system, radiance and its cousin luminance are good indicators of how bright an object will appear. For this reason, radiance and luminance are both sometimes called "brightness". This usage is now discouraged – see Brightness for a discussion. The nonstandard usage of "brightness" for "radiance" persists in some fields, notably laser physics

Def
$$L = \frac{d^2\Phi}{dAd\Omega cos(\theta)} \approx \frac{\Phi}{\Omega Acos(\theta)}$$

2.1.5 Signal

A signal is a function that conveys information about the behavior or attributes of some phenomenon. In the physical world, any quantity exhibiting variation in time or variation in space (such as an image) is potentially a signal that might provide information on the status of a physical system, or convey a message between observers

2.1.6 Fourier Transformation

The Fourier-Transform is a mathematical tool which allows to transform a given function or rather a given signal from defined over a time- (or spatial-) domain into its corresponding frequency-domain.

Let f an measurable function over \mathcal{R}^n . Then, the coninious Fourier Transformation, denoted as FT, $\mathcal{F}\{t\}$ of f is defined as, ignoring all constant factors in the formula:

$$\mathcal{F}\{w\}_{FT} = \int_{\mathcal{R}^n} f(x)e^{-iwt}dt \tag{1}$$

whereas its inverse transform is defined like the following which allows us to obtain back the original signal:

$$\mathcal{F}\{w\}_{FT}^{-1} = \int_{\mathbb{R}} \mathcal{F}\{w\} e^{iwt} dt \tag{2}$$

By using fourier analysis, which is the approach to approximate any function by sums of simpler trigonometric functions, we gain the so called discrete time fourier transform (in short DTFT). The DTFT operates on a discrete function. Usually, such an input function is often created by diitally sampling a continius function. The DTFT itself is operation on a discretized signal on a continious, periodic frequency domain and looks like the following:

$$\mathcal{F}\{w\}_{DFT} = \sum_{-\infty}^{\infty} f(x)e^{(-iwk)}$$
(3)

we can further discretize the frequency domain and will get then the discrete fourier transformation (in short DFT) of the input signal:

$$\mathcal{F}\{w\}_{DFT} = \sum_{n=0}^{N-1} f(x)e^{(-iw_n k)}$$
 (4)

Where the angular frequency w_n is defined like the following $w_n = \frac{2\pi n}{N}$ and N is the number of samples within an equidistant periode sampling.

2.1.7 Convolution

$$(f * g)(t) = \int_{\mathcal{R}^n} f(t)g(t - x)dx \tag{5}$$

Note that the Fourier transform of the convolution of two functions is the product of their Fourier transforms. This is equivalent to the fact that Convolution in spatial domain is equivalent to multiplication in frequency domain. Therefore, the inverse Fourier transform of the product of two Fourier transforms is the convolution of the two inverse Fourier transforms

2.1.8 BRDF

The bidirectional reflectance distribution function, in short BRDF, denoted as $f_r(w_i, w_r)$ is a four dimensional function that defines how light is reflected at an opaque surface. The function takes a negative incoming light direction, ω_i , and outgoing direction, ω_r , both defined with respect to the surface normal \mathbf{n} and returns the ratio of reflected radiance exiting along ω_r to the irradiance incident on the surface from direction ω_i

$$f_r(w_i, w_r) = \frac{dL_r(w_r)}{dE_i(w_i)} \tag{6}$$

$$= \frac{dL_r(w_r)}{L_i(w_i)cos(\theta_i)dw_i} \tag{7}$$

where L is radiance, or power per unit solid-angle-in-the-direction-of-a-ray per unit projected-area-perpendicular-to-the-ray, E is irradiance, or power per unit surface area, and θ_i is the angle between ω_i and the surface normal, \mathbf{n} . The index i indicates incident light, whereas the index r indicates reflected light.

The reason the function is defined as a quotient of two differentials and not directly as a quotient between the undifferentiated quantities, is because other irradiating light than $d E_i(\omega_i)$, which are of no interest for $f_r(\omega_i, \omega_r)$, might illuminate the surface which would unintentionally affect $L_r(\omega_r)$, whereas $d L_r(\omega_r)$ is only affected by $d E_i(\omega_i)$.

2.1.9 Spectral Rendering

In Computer Graphics, spectral rendering is where a scene's light transport is modeled considering the whole span of wavelengths instead of R,G,B values (still relating on geometric optic, which ignore wave phase). The motivation is that real colors of the physical world are spectrum; trichromatic colors are only inherent to Human Visual System.

2.1.10 Taylor Series

Taylor series is a representation of a function as an infinite sum of terms that are calculated from the values of the function's derivatives at a single point.

The Taylor series of a real or complex-valued function f(x) that is infinitely differentiable at a real or complex number a is the power series:

$$\mathcal{T}(f;a)(x) = \sum_{n=0}^{\infty} \frac{f^n(a)}{n!} (x-a)^n$$
 (8)

2.2 Thesis Basis: J.Stam's Paper about Diffraction Shader

GOAL main task in the theory of diffraction is to solve this wave equation for different geometries. we are interested in computing the reflected waves from various types of surfaces

abstract: before: most reflection models empirically or based on ray-theory of light. now: new reflection model based on wave theory modeling the effect of diffraction

In his Paper Diffraction Shader, Jos Stam derives a BRDF which modeling the effect of diffraction for various analytical anistropic reflaction models using the scalar Kirchof theory and the theory of random processes. By employing the so called scalar wave theory of diffraction [source 5 in stams paper] in which a wave is assumed to be a complex valued scalar. It's noteworthy, that stam's BRDF formulation does not take into account the polarization of the light. Fortunately, light sources like sunlight and light bulbs are unpilarizaed. In our simulations we will always assume we have given i directional light source, i.e. sunlight. Hence, we can use stam's model for our derivations

A further assumption in Stam's Paper is, the emanated waves from the source are stationary, which implies the wave is a superposition of independent monochromatic waves. This implies that each wave is associated to a definite wavelangth lambda. However, sunlight once again fulfills this fact.

Mention Helmolth equation, which has the solution $k = \frac{2\pi}{\lambda}$ which is the wavenumber

Based on his these previous assumptions and applying Stam starts his derviations by applying the so called Kirchhoff integral, which is relating the reflected field to the incoming field. This equation is a formalization of Huygen's well-known principle that states that if one knows the wavefront at a given moment, the wave at a later time can be deduced by considering each point on the first wave as the source of a new disturbance, i.e. once the field $\psi_1 = e^{ik\mathbf{x}\cdot\mathbf{ss}}$ on the surface is known, the field everywhere ψ_2 else away from the surface can be computed. More precisely, we want to compute the wave ψ_2 equal to the reflection of an incoming planar monochromatic wave $\psi_1 = e^{ikk_1*x}$ traveling in the direction k_1 from a surface S. Mathematically this can be formulized the following:

$$\psi_2 = \frac{ike^{iKR}}{4\pi R} (F\mathbf{v} - \mathbf{p}) \cdot \int_S \hat{\mathbf{n}} e^{ik\mathbf{v} \cdot \mathbf{s}d\mathbf{s}}$$
 (9)

In applied optics, when dealing with scattered waves, one does use differential scattering cross-section rather than defining a BRDF which has the following identity:

$$\sigma^{0} = 4\pi \lim_{R \to \infty} R^{2} \frac{\langle |\psi_{2}|^{2} \rangle}{\langle |\psi_{1}|^{2} \rangle}$$
 (10)

Relationship between the BRDF and the scattering cross section is the follwing:

The relationship between the BRDF and the scattering cross section can be shown to be equal to $BRDF = \frac{1}{4\pi} \frac{1}{A} \frac{\sigma^0}{cos(\theta_1)cos(\theta_2)}$ Wheras $\theta_1 a$ and θ_2 are the angles that the vectors $\hat{k_1}$ and $\hat{k_2}$ make with the vertical direction.

ADD FIGURE for k1, k2

where R is the disance from the center of the patch to the receiving point x_p , $\hat{\mathbf{n}}$ is the normal of the surface at s and the vectors:

$$\mathbf{v} = \hat{\mathbf{k_1}} - \hat{\mathbf{k_1}} = (u, v, w)$$

$$p = \hat{k_1} + \hat{k_1}$$

During his derivations, Stam provides a analytical representation for the Kirchhoff integral by using his assumptions. He restricts himself to the reflaction of waves from height fields h(x,y) with the assumption that the surface is defined as an elevation over the (x,y) plane using the surface plane approximation.

Which will lead him to the following identity for the Kirchhoff integral

$$\mathbf{I}(ku,kv) = \int \int \frac{1}{ikw} (-p_x, -p_y, ikwp) \tag{11}$$

wheras stam formulates for a hightfield auxiliary function $p(x,y) = e^{iwkh(x,y)}$ where $w = -(cos(\theta_i) + cos(\theta_r))$ and θ_i and θ_r are the angles of incident and reflected directions with the surface normal (ADD picture) and the wavenumber $k = \frac{2\pi}{\lambda}$

$$p(x,y) = e^{ikwh(x,y)} \tag{12}$$

We the observation that the integral is a Fourier transform by -iku and -ikv which will lead us to his final derivation, using the identity of BRDF, and computing the limes:

$$BRDF_{\lambda}(w_i, w_r) = \frac{k^2 F^2 G}{4\pi^2 A w^2} \langle |P(ku, kv)|^2 \rangle$$
 (13)

 $BRDF_{\lambda}(w_i, w_r)$ is BRDF where wavelength λ w_i and w_r are incident and reflected normalized directions vectors, pointing away from the given surface. Which can be written, using the fourier transform (FT) P(u, v) = F(p)(u, v), as: $BRDF_{\lambda}(w_i, w_r) = \frac{F^2G}{\lambda^2 A w^2} abs(P(\frac{u}{\lambda}, \frac{v}{\lambda}))^2$ where F represents the Fresnel term, uv, v, w are derived from the incident and reflected directions as $(u, v, w) = -\omega_i - \omega_r$, abs(P) represents the expected valuess of a random variable X and A is an area of integration on the surface that is considered to contribute to diffraction, G is the geometry term which is $G = \frac{(1-\hat{\mathbf{k}}_1 \cdot \hat{\mathbf{k}}_2)^2}{cos(\theta_1)cos(\theta_2)}$

and P(x,y) is the Fourier transform (FT) of the function p(x,y) from above. This identity for the BRDF is the starting point for our derivations.

2.3 Derivations

2.3.1 BRDF formulation

EXPLAIN: Why do we want a formulation for $L_{\lambda}(w_r)$ in some words. what does it represent?

Definition of $BRDF(w_i, w_r) := f_r(w_i, w_r) = \frac{dL_r(w_r)}{dE_i(w_i)} = \frac{dL_r(w_r)}{L_i(w_i)cos(\theta_i)dw_i}$ Hence, we can dervie the following expression:

$$\begin{split} f_r(w_i,w_r) &= \frac{dL_r(w_r)}{L_i(w_i)cos(\theta_i)dw_i} \\ &=> f_r(w_i,w_r)L_i(w_i)cos(\theta_i)dw_i = dL_r(w_r) \\ &=> \int_{\Omega} f_r(w_i,w_r)L_i(w_i)cos(\theta_i)dw_i = \int_{\Omega} dL_r(w_r) \\ &=> L_r(w_r) = \int_{\Omega} f_r(w_i,w_r)L_i(w_i)cos(\theta_i)dw_i \end{split}$$

We assume, that our incident light is a directional light source like sunlight and therefore its radiance is given as $L_{\lambda}(w) = I(\lambda)\delta(w - w_i)$ where $I(\lambda)$ is the intensity of the relative spectral power for the wavelength λ . Thus we get for our the brdf formulation:

$$L_{\lambda}(w_r) = \int_{\Omega} BRDF_{\lambda}(w_i, w_r) L_{\lambda}(w_i) cos(\theta_i) dw_i$$
 (14)

$$= BRDF_{\lambda}(w_i, w_r)I(\lambda)cos(\theta_i) \tag{15}$$

where w_i is the solid angle for the incoming light, θ_i is the angle of incidence, w_r is the solid angle for the reflected light, λ wavelength, Ω is the hemisphre we of integration for the incomming light. Radiance reflected by given surface in given direction: $L_{\lambda}(w_i)$ is the incomming radiance, $L_{\lambda}(w_r)$ is the reflected radiance

For the $BRDF(w_i, w_r)$ we are going to use the formulation dervied by Stam described above which looks like this using the fact that wavenumber $k = \frac{2\pi}{\lambda}$:

$$BRDF(w_{i}, w_{r}) = \frac{k^{2}F^{2}G}{4\pi^{2}Aw^{2}} \langle |P(ku, kv)|^{2} \rangle$$

$$= \frac{k^{2}F^{2}(1 - \hat{\mathbf{k_{1}}} \cdot \hat{\mathbf{k_{2}}})}{\cos(\theta_{1})\cos(\theta_{2})4\pi^{2}Aw^{2}} \langle |P(ku, kv)|^{2} \rangle$$

$$= \frac{4\pi^{2}F^{2}(1 - \hat{\mathbf{k_{1}}} \cdot \hat{\mathbf{k_{2}}})}{\cos(\theta_{1})\cos(\theta_{2})4\pi^{2}A\lambda^{2}w^{2}} \langle |P(ku, kv)|^{2} \rangle$$

$$= \frac{F(w_{i}, w_{r})^{2}(1 - \hat{\mathbf{k_{1}}} \cdot \hat{\mathbf{k_{2}}})}{\cos(\theta_{1})\cos(\theta_{2})A\lambda^{2}w^{2}} \langle |P(ku, kv)|^{2} \rangle$$

where $\hat{\mathbf{k_t}}$ represents a unit vector whose spherical coordinates are given by the solid angle t. Since we are going to integrate over a sphere Ω we can write the component $w = (\cos(\theta_i) + \cos(\theta_r))$ SHOW WHY WE ARE ALLOWED TO WRITE IT LIKE THIS => SPHERICAL COORDINATES DIFFERENCE (k1-k2) = (u,v,w) and so on. this our identity for $L_r(w_r)$ will lead us to the following identity using our identity:

$$\begin{split} L_{\lambda}(w_{r}) &= \frac{F(w_{i}, w_{r})^{2}(1 - \hat{\mathbf{k}_{1}} \cdot \hat{\mathbf{k}_{2}})^{2}}{A\lambda^{2}cos(\theta_{i})cos(\theta_{r})(cos(\theta_{i}) + cos(\theta_{r}))^{2}} \left\langle \left| P_{cont}(\frac{2\pi u}{\lambda}, \frac{2\pi v}{\lambda}) \right|^{2} \right\rangle cos(\theta_{i})I(\lambda) \\ &= I(\lambda) \frac{F(w_{i}, w_{r})^{2}(1 - \hat{\mathbf{k}_{1}} \cdot \hat{\mathbf{k}_{2}})^{2}}{\lambda^{2}A(cos(\theta_{i}) + cos(\theta_{r}))^{2}cos(\theta_{r})} \left\langle \left| P_{cont}(\frac{2\pi u}{\lambda}, \frac{2\pi v}{\lambda}) \right|^{2} \right\rangle \\ &= I(\lambda) \frac{F(w_{i}, w_{r})^{2}(1 - \hat{\mathbf{k}_{1}} \cdot \hat{\mathbf{k}_{2}})^{2}}{\lambda^{2}A(cos(\theta_{i}) + cos(\theta_{r}))^{2}cos(\theta_{r})} \left\langle \left| T_{0}^{2}P_{dtft}(\frac{2\pi u}{\lambda}, \frac{2\pi v}{\lambda}) \right|^{2} \right\rangle \end{split}$$

 P_{cont} is the continious inverse Fourier transform for the taylor seies of our hight-field representing the nano structure, i.e. $P(k,l) = \mathcal{F}^{-1}\{p\}(k,l)$ and

 P_{dtft} is the dicrete-time inverse Fourier Transform for the same problem domain and T_0 the sampling distance for the discretization pf p(x,y) assuming equal and uniform sampling in both dimensions x, y.

2.3.2 Relative BRDF

reason why relative brdf: In order to scale the reflactiance such that we are able to texture. convex combination reflectance with texture. Scale illumination.

Let us examine what $L_{\lambda}(w_r)$ will be for $w_r = w_0 := (0,0,*)$ i.e. specular reflection case, denoted as $L_{\lambda}^{spec}(w_0)$. When we know the expression for $L_{\lambda}^{spec}(w_0)$ we would be able to compute the relative reflected radiance for our problem by simply dividing $L_{\lambda}(w_r)$ by $L_{\lambda}^{spec}(w_0)$, denoted as

$$\rho_{\lambda}(w_i, w_r) = \frac{L_{\lambda}(w_r)}{L_{\lambda}^{spec}(w_0)}$$
(16)

But first, let us derive the following expression:

$$F(w_0, w_0)^2 (1 - \begin{pmatrix} 0 \\ 0 \\ 1 \end{pmatrix} \cdot \begin{pmatrix} 0 \\ 0 \\ -1 \end{pmatrix})^2$$

$$L_{\lambda}^{spec}(w_0) = I(\lambda) \frac{\lambda^2 A(\cos(0) + \cos(0))^2 \cos(0)}{\lambda^2 A(\cos(0) + \cos(0))^2 \cos(0)} \langle \left| T_0^2 P_{dtft}(0, 0) \right|^2 \rangle$$

$$= I(\lambda) \frac{F(w_0, w_0)^2 (1+1)^2}{\lambda^2 A (1+1)^2 1} \left| T_0^2 N_{sample} \right|^2$$

$$= I(\lambda) \frac{F(w_0, w_0)^2}{\lambda^2 A} \left| T_0^2 N_{sample} \right|^2$$

Where $N_{samples}$ is the number of samples of the dtft.

Thus, we can plug our last derived expression into the definition for the relative reflectance radiance in the direction w_r and will get:

$$\begin{split} \rho_{\lambda}(w_{i},w_{r}) &= \frac{L_{\lambda}(w_{r})}{L_{\lambda}^{spec}(w_{0})} \\ &= \frac{I(\lambda) \frac{F(w_{i},w_{r})^{2}(1-\hat{\mathbf{k_{1}}}\cdot\hat{\mathbf{k_{2}}})^{2}}{\lambda^{2}A(\cos(\theta_{i})+\cos(\theta_{r}))^{2}\cos(\theta_{r})} \langle \left| T_{0}^{2}P_{dtft}(\frac{2\pi u}{\lambda},\frac{2\pi v}{\lambda}) \right|^{2} \rangle}{I(\lambda) \frac{F(w_{0},w_{0})^{2}}{\lambda^{2}A} \left| T_{0}^{2}N_{sample} \right|^{2}} \\ &= \frac{F^{2}(w_{i},w_{r})(1-\hat{\mathbf{k_{1}}}\cdot\hat{\mathbf{k_{2}}})^{2}}{F^{2}(w_{0},w_{0})(\cos(\theta_{i})+\cos(\theta_{r}))^{2}\cos(\theta_{r})} \langle \left| \frac{P_{dtft}(\frac{2\pi u}{\lambda},\frac{2\pi v}{\lambda})}{N_{samples}} \right|^{2} \rangle \end{split}$$

for simplification and a better overview, let us introduce the following expression, the so called gain factor

$$C(w_i, w_r) = \frac{F^2(w_i, w_r)(1 - \hat{\mathbf{k_1}} \cdot \hat{\mathbf{k_2}})^2}{F^2(w_0, w_0)(\cos(\theta_i) + \cos(\theta_r))^2 \cos(\theta_r) N_{samples}^2}$$
(17)

Using this substitute, we will end up with the following expression for the relative reflectance radiance

$$\rho_{\lambda}(w_i, w_r) = C(w_i, w_r) \langle \left| P_{dtft}(\frac{2\pi u}{\lambda}, \frac{2\pi v}{\lambda}) \right|^2 \rangle$$
 (18)

using the previous definition for the relative reflectance radiance $\rho_{\lambda}(w_i, w_r) = \frac{L_{\lambda}(w_r)}{L_{\lambda}^{Spec}(w_0)}$ which we can rearrange to the expression

$$L_{\lambda}(w_r) = \rho_{\lambda}(w_i, w_r) L_{\lambda}^{spec}(w_0) \tag{19}$$

Let us choose $L_{\lambda}^{spec}(w_0) = S(\lambda)$ such that is has the same pforifle as the relative spectral power distribution of CIE Standard Illuminant D65. Further, when integration over λ for a specular surface we should get CIE_XYZ values corresponding to the white point for D65

the corresponding tristimulus values using CIE color matching functions for the CIE_XYZ values look like:

SEE HOW THIS DEFINITION DIFFERS FROM THE WIKIDEF AND HOW WE COULD END UP WITH A SIMILAR DEFINITION.

$$X = \int_{\lambda} L_{\lambda}(w_r) \overline{x}(\lambda) d\lambda \tag{20}$$

$$Y = \int_{\lambda} L_{\lambda}(w_r)\overline{y}(\lambda)d\lambda \tag{21}$$

$$Z = \int_{\lambda} L_{\lambda}(w_r)\overline{z}(\lambda)d\lambda \tag{22}$$

where \overline{x} , \overline{y} , \overline{z} are the color matching functions

Using our last finding for $L_{\lambda}(w_r)$ and the definition for the tristimulus values we can actually derive an expression for computing the colors for our brdf model. Since X, Y, Z are defined similarly, it satisfies to derive an explicit expression for just one tristimulus term, for example X. The other two will look the same, except the we have to replace all X with Y or Z respectively. Therefore, we get:

$$X = \int_{\lambda} L_{\lambda}(w_{r})\overline{x}(\lambda)d\lambda$$

$$= \int_{\lambda} \rho_{\lambda}(w_{i}, w_{r})L_{\lambda}^{spec}(w_{0})\overline{x}(\lambda)d\lambda$$

$$= \int_{\lambda} \rho_{\lambda}(w_{i}, w_{r})S(\lambda)\overline{x}(\lambda)d\lambda$$

$$= \int_{\lambda} C(w_{i}, w_{r})\langle \left| P_{dtft}(\frac{2\pi u}{\lambda}, \frac{2\pi v}{\lambda}) \right|^{2} \rangle S(\lambda)\overline{x}(\lambda)d\lambda$$

$$= C(w_{i}, w_{r})\int_{\lambda} \langle \left| P_{dtft}(\frac{2\pi u}{\lambda}, \frac{2\pi v}{\lambda}) \right|^{2} \rangle S(\lambda)\overline{x}(\lambda)d\lambda$$

$$= C(w_{i}, w_{r})\int_{\lambda} \langle \left| P_{dtft}(\frac{2\pi u}{\lambda}, \frac{2\pi v}{\lambda}) \right|^{2} \rangle S_{x}(\lambda)d\lambda$$

Where we used the definition $S_x(\lambda)\overline{x}(\lambda)$ in the last step.

2.3.3 Taylour approximation for BRDF

Based on J. Stam's Paper about Diffraction shaders we will show that there is an approximation of his equation (5), $\mathbf{p}(\mathbf{x},\mathbf{y})$, for a explicitly given heightfield $\mathbf{h}(\mathbf{x},\mathbf{y})$. This approximation is achieved by using Taylor-Series and using this identity we will further be able to approximate the Fourier-Transformation of $\mathbf{p}(\mathbf{x},\mathbf{y})$, denoted as $\mathbf{P}(\mathbf{u},\mathbf{v})$. Finally we will give an error bound for this approximation. Finally, we will put our new found identity into our so far found relative BRDF representation.

Taylor Series of p Given $p(x,y) = e^{ikwh(x,y)}$ form Stam's Paper where h(x,y) is here a given heightfield. Also given the definition $e^y = 1 + y + \frac{y^2}{2!} + \frac{y^3}{3!} + \dots = \sum_{n=0}^{\infty} \frac{y^n}{n!}$ where y can be real or even complex valued - note this identity can either be derived by power series or by Taylor-Series (using the derivatives of the exp-function and developing the Taylor-Series around the point a=0). Let us now set y=ikwh(x,y) where i is the imaginary number. For simplification, let us denote h(x,y) as h. It follows by our previous stated identities: $e^y = 1 + (ikwh) + \frac{1}{2!}(ikwh)^2 + \frac{1}{3!}(ikwh)^3 + \dots = \sum_{n=0}^{\infty} \frac{(ikwh)^n}{n!}$. Hence it holds $p(x,y) = \sum_{n=0}^{\infty} \frac{(ikwh(x,y))^n}{n!}$.

Fourier Transformation of function p Let us now compute the Fourier Transformation of p(x,y) form above: $\mathcal{F}\{p\}(u,v) = \mathcal{F}\left\{\sum_{n=0}^{\infty} \frac{(ikwh(x,y))^n}{n!}.\right\} = \mathcal{F}^{lin\,Operator}$ $\sum_{n=0}^{\infty} \mathcal{F}\left\{\frac{(ikwh(x,y))^n}{n!}\right\} = \sum_{n=0}^{\infty} \frac{(ikwh)^n}{n!} \mathcal{F}\{h(x,y)^n\}. \text{ Hence it follows: } P(\alpha,\beta) = \sum_{n=0}^{\infty} \frac{(ikwh)^n}{n!} \mathcal{F}\{h^n\}(\alpha,\beta).$ $\mathbf{NB}: \mathcal{F}\{h^n\}(u,v) \text{ denotes the two dimensional Fourier Transformation}$

NB: $\mathcal{F}\{h^n\}(u,v)$ denotes the two dimensional Fourier Transformation of p(x,y) and can be numerically computed by the two dimensional **DFT** or rather by the two dimensional **FFT** over h(x,y).

Approximation of function P Next we are going to look for an $N \in \mathbb{N}$ s.t. $\sum_{n=0}^{N} \frac{(ikwh)^n}{n!} \mathcal{F}\{h^n\}(\alpha,\beta) \approx P(\alpha,\beta)$. is a good approximation. We have to prove two things:

- 1. Show that there exist such an $N \in \mathbb{N}$ s.t the approximation holds true.
- 2. Find a value for B s.t. this approximation is below a certain error bound, for example machine precision ϵ .

Proof Sketch of 1. By the **ratio test** (see [1]) we can show that the series $\sum_{n=0}^{N} \frac{(ikwh)^n}{n!} \mathcal{F}\{h^n\}(\alpha,\beta)$ converges absolutely:

Proof: Consider $\sum_{k=0}^{\infty} \frac{y^n}{n!}$ where $a_k = \frac{y^k}{k!}$. By the definition of the ratio test for series it follows: $\forall y : limsup_{k\to\infty} |\frac{a_{k+1}}{a_k}| = limsup_{k\to\infty} \frac{y}{k+1} = 0$

Thus this series converges absolutely, no matter what value we will pick for y.

Part 2: Find such an N Let $f(x) = e^x$. We can formulate its Taylor-Series, stated above. Let $P_n(x)$ denote the n-th Taylor-Polinomial, $P_n(x) = \sum_{k=0}^n \frac{f^{(k)}(a)}{k!} (x-a)^k$, where a is our developing point (here, in this case a=0). We can define the error of the n-th Taylor-Polinomial to be $E_n(x) = f(x) - P_n(x)$. That error is the actual value minus the Taylor polinomial. It holds true: $|E_n(x)| = |f(x) - P_n(x)|$. By using the Lagrangien Error Bound (see source [2]) it follows: $|E_n(x)| \leq \frac{M}{(n+1)!} |x-a|^{n+1}$ with a=0, where M is some value satisfying $|f^{(n+1)}(x)| \leq M$ on the interval I = [a,x]. Since we are interested in an upper bound of the error and since a is known, we can reformulate the interval as $I = [0, x_{max}]$, where $x_{max} = |i| * k_{max} * w_{max} * h_{max}$, since we are interested in computing an error bound for $e^{ikwh(x,y)}$. From Stam's Paper about diffraction shader we know some paramters for the length, width and height for a given sample patch, i.e. heightfield h(x,y) and when using those parameters are able to find a explicit number for x_{max} .

Facts we are using from Stam's Paper:

- Height of bump: 0.15micro meters
- Width of a bump: 0.5micro meters
- Length of a bump: 1micro meters
- $k = \frac{2\pi}{\lambda}$ is the wavenumber and $\lambda \in [\lambda_{min}, \lambda_{max}]$ its wavelength hence $k_{max} = \frac{2\pi}{\lambda_{min}}$
- w is a component of the vector $\vec{v} = \vec{k_1} \vec{k_2} = (u, v, w)$, where $\vec{k_1}$ and $\vec{k_2}$ are **normalized** direction vectors and this each component can have a value in range [-2, 2].
- for simplification, assume $[\lambda_{min}, \lambda_{max}] = [400nm, 700nm]$.

Hence $x_{max} = |i| * k_{max} * w_{max} * h_{max} = k_{max} * w_{max} * h_{max} = 2 * (\frac{2\pi}{4*10^{-7}m}) * 1.5 * 10^{-7} = 1.5\pi$ and it follows for our intervall $I = [0, 1.5\pi]$. Next we are going to find the value for M. Since the exponential function is monoton growing (on the interval I) and and the derivative of the \exp function is the exp function itself, we can find such an M: $M = e^{x_{max}} = exp(1.5\pi)$ and $|f^{(n+1)}(x)| \leq M$ holds. With $|E_n(x_{max})| \leq \frac{M}{(n+1)!} |x_{max} - a|^{n+1} = \frac{exp(1.5\pi)*(1.5\pi)^{n+1}}{(n+1)!}$ we now can find a value of n for a given bound, i.e. we can find an value of $N \in \mathbb{N}$ s.t. $\frac{exp(1.5\pi)*(1.5\pi)^{N+1}}{(N+1)!} \leq \epsilon$. With Octave/Matlab we can see:

- if N=20 then $\epsilon \approx 2.9950 * 10^{-4}$
- if N=25 then $\epsilon \approx 8.8150 * 10^{-8}$
- if N=30 then $\epsilon \approx 1.0050 * 10^{-11}$

Conclusion With this approach we have that $\sum_{n=0}^{25} \frac{(ikwh)^n}{n!} \mathcal{F}\{h^n\}$ (α,β) is an approximation of P(u,v) with $error \epsilon \approx 8.8150*10^{-8}$. This means we can precompute 25 Fourier Transformations (for example via FFT2) and then sum them up in order to approximate P(u,v) and $\epsilon \approx 8.8150*10^{-8}$. This approach will allow us to speed up our shader. Furthermore we see that when we just take 5 more iterations, we will reduce the error bound to the dimension of 10^{-11} .

Using $P_{dtft} = \mathcal{F}^{-1}\{p\}(u,v)$ definied in the section of the taylor approximationwe get for the tristumulus value X, we will get:

$$X = C(w_i, w_r) \int_{\lambda} \left\langle \left| P_{dtft}(\frac{2\pi u}{\lambda}, \frac{2\pi v}{\lambda}) \right|^2 \right\rangle S_x(\lambda) d\lambda$$

$$= C(w_i, w_r) \int_{\lambda} \left| \sum_{n=0}^{N} \frac{(wk)^n}{n!} \mathcal{F}^{-1} \{i^n h^n\} (\frac{2\pi u}{\lambda}, \frac{2\pi v}{\lambda}) \right|^2 S_x(\lambda) d\lambda$$

2.3.4 Sampling: Gaussian Window

why this identity works: The DFT of a discrete heightfield patch is equivalent to the DTFT of an infinitely periodic function consisting of replicas of the same discrete patch. By windowing with a window function that is zero outside the central replica, the convolution of either the DFT or the DTFT of heightfield with the fourier transfrom of the window becomes equivalent.

Let $window_g$ denote the gaussian window with $4\sigma_s$ μm where $\sigma_f = \frac{1}{2\pi\sigma_s}$ let us further substitute $\mathbf{t}(\mathbf{x}, \mathbf{y}) = i^n h(x, y)^n$

$$\mathcal{F}_{dtft}^{-1}\{\mathbf{t}\}(u,v) = \mathcal{F}_{fft}^{-1}\{\mathbf{t}\}(u,v)window_g(\sigma_f)$$
 (23)

Therefore we can deduce the following expression from this:

$$\mathcal{F}_{dtft}^{-1}\{\mathbf{t}\}(u,v) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} F_{fft}^{-1}\{\mathbf{t}\}(w_{u},w_{v})\phi(u-w_{u},v-w_{v})dw_{u}dw_{v}
= \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \sum_{i} \sum_{j} F_{fft}^{-1}\{\mathbf{t}\}(w_{u},w_{v})\delta(w_{u}-w_{i},w_{v}-w_{j})\phi(u-w_{u},v-w_{v})dw_{u}dw_{v}
= \sum_{i} \sum_{j} \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} F_{fft}^{-1}\{\mathbf{t}\}(w_{u},w_{v})\delta(w_{u}-w_{i},w_{v}-w_{j})\phi(u-w_{u},v-w_{v})dw_{u}dw_{v}
= \sum_{i} \sum_{j} F_{fft}^{-1}\{\mathbf{t}\}(w_{u},w_{v})\phi(u-w_{u},v-w_{v})$$

where
$$\phi(x,y) = \pi e^{-\frac{x^2+y^2}{2\sigma_f^2}}$$

2.3.5 Aplitude smooting

Let us consider the so called 1-dimensional Box-function with length T which is defined as the following: ADD AN IMAGE OF BOXFUNCTION

$$Box(x) = \begin{cases} 1 & \text{if } x \le T \\ 0 & \text{if } else \end{cases}$$

We assume, that our given heighfield can be represented as a 2-dimensional box-function. Note that we can use any explicit given constrainted 2-dimensional function and will get some identities like we get from the box-function.

Further we are assuming that we can model the overall surface be assuming this heighfield being distributed in a periodic manor. Therfore, the whole surface can be represented like this $f(x) = \sum_{n=0}^{N} Box(x+nT_1,y+mT_2)$ assuming the given heighfield has the dimensions T_1 by T_2 . But let us first consider the 1-dimensional Box-function case before deriving an identity for the Fourier transform of our 2-dimensional Box-function, i.e. the fourier transform of our heighfield.

Note: A function f periodic with periode T means: $\forall x \in \mathcal{R} : Box(x) = Box(x+T)$

A so called bump can be represented by our 1-dimensional Box-function. We assume periodicity which is equaivalent to: $f(x) = \sum_{n=0}^{N} Box(x + nT)$

We are insterested in the 1-dimensional inverse Fourier transform of the 1-dimensional Box-function:

$$\mathcal{F}^{-1}{f}(w) = \int f(x)e^{iwx}dx$$

$$= \int_{-\infty}^{\infty} \sum_{n=0}^{N} Box(x+nT)e^{iwx}dx$$

$$= \sum_{n=0}^{N} \int_{-\infty}^{\infty} Box(x+nT)e^{iwx}dx$$

Next, apply the following substituation x + nT = y which will lead us to:

$$x = y - nT$$
$$dx = dy$$

Plugging this substituation back to the equation from above we will get

$$\mathcal{F}^{-1}\{f\}(w) = \int f(x)e^{iwx}dx$$

$$= \sum_{n=0}^{N} \int_{-\infty}^{\infty} Box(y)e^{iw(y-nT)}dy$$

$$= \sum_{n=0}^{N} e^{-iwnT} \int_{-\infty}^{\infty} Box(y)e^{iwy}dy$$

$$= \sum_{n=0}^{N} e^{-iwnT} \mathcal{F}\{f\}(w)$$

$$= \mathcal{F}^{-1}\{f\}(w) \sum_{n=0}^{N} e^{-iwnT}$$

We used the fact that the term e^{-iwnT} is a constant when integrating along dy and the identity for the inverse Fourier transform of the Box function. Next, let us consider $\sum_{n=0}^{N} e^{-uwnT}$ further:

$$\sum_{n=0}^{N} e^{-uwnT} = \sum_{n=0}^{N} (e^{-uwT})^n$$
$$= \frac{1 - e^{iwT(N+1)}}{1 - e^{-iwT}}$$

We recognize the geometric series identity for the left-handside of this equation. Since our series is bounded we can derive our right-handside.

Since e^{-ix} is a complex number and every complex number can be written in its polar form, i.e. $e^{-ix} = cos(x) + isin(x)$ we can go even further, using the trigonometric idententities that cos(-x) = cos(x) and sin(-x) = -sin(x):

$$\frac{1 - e^{iwT(N+1)}}{1 - e^{-iwT}} = \frac{1 - \cos(wT(N+1)) + i\sin(wT(N+1))}{1 - \cos(wT) + i\sin(wT)}$$

Which is still a complex number (p+iq). Every complex number can be written as a fraction of two complex numbers. This means that the complex number (p+iq) can be written as $(p+iq) = \frac{(a+ib)}{(c+id)}$ for any (a+ib), (c+id) $id) \neq 0$. For our case, let us use the following substituations:

$$a := 1 - cos(wT(N+1)) \qquad b = sin(wT(N+1)) \qquad (24)$$

$$c = 1 - cos(wT) \qquad d = sin(wT) \qquad (25)$$

$$c = 1 - \cos(wT) \qquad \qquad d = \sin(wT) \tag{25}$$

hence it follows $\frac{1-e^{iwT(N+1)}}{1-e^{-iwT}} = \frac{(a+ib)}{(c+id)}$. By rearanging the terms it follows (a+ib) = (c+id)(p+iq) and multiplying the right handside out we get the following system of equations:

$$(cp - dq) = a \tag{26}$$

$$(dp + cq) = b (27)$$

Which gives lead us we some further math (trick: mult first eq. by c and 2nd by d, then adding them together using distributivity and we have the identity for p for example, similar for q) to

$$p = \frac{(ac + bd)}{c^2 + d^2} \tag{28}$$

$$q = \frac{(bc + ad)}{c^2 + d^2} \tag{29}$$

Putting our substituation for a, b, c, d back into the current representation for p and q and using some trigonometric identities, this we then get:

$$p = \frac{1}{2} + \frac{1}{2} \left(\frac{\cos(wTN) - \cos(wT(N+1))}{1 - \cos(wT)} \right)$$
 (30)

$$q = \frac{\sin(wT(N+1)) - \sin(wTN) - \sin(wT)}{2(1 - \cos(wT))}$$
(31)

Since we have seen, that $\sum_{n=0}^{N} e^{-uwnT}$ is a complex number and can be written as (p+iq) and we know now the explicit identity for those p and q we get for the 1-dimensional Fourier transform of the 1-dimensional Boxfunction the following final identity:

$$\mathcal{F}^{-1}{f}(w) = \mathcal{F}^{-1}{f}(w) \sum_{n=0}^{N} e^{-iwnT}$$
$$= (p+iq)\mathcal{F}^{-1}{Box}(w)$$

In oder to derive next a identity for the Fourier transform for our 2-dim heighfield, we can proceed similarly, the only fact which changes is, that we are now in a 2-dimensional domain, i.e. we are about to compute a two-dimensional Fourier transform: Let us again us again a Box-function, this time a 2-dimensional Box-function Box(x,y) just for the sake of convenience.

$$\mathcal{F}^{-1}\{f\}(w_{1},w_{2}) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \sum_{n_{2}=0}^{N_{1}} \sum_{n_{2}=0}^{N_{2}} Box(x_{1} + n_{1}T_{1}, x_{2} + n_{2}T_{2})e^{iw(x_{1}+x_{2})}dx_{1}dx_{2}$$

$$= \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \sum_{n_{2}=0}^{N_{1}} \sum_{n_{2}=0}^{N_{2}} Box(y_{1},y_{2})e^{iw((y_{1}-n_{1}T_{1})+(y_{2}+n_{2}T_{2}))}dx_{1}dx_{2}$$

$$= \sum_{n_{2}=0}^{N_{1}} \sum_{n_{2}=0}^{N_{2}} \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} Box(y_{1},y_{2})e^{iw(y_{1}+y_{2})}e^{-iw(n_{1}T_{1}+n_{2}T_{2})}dy_{1}dy_{2}$$

$$= \sum_{n_{2}=0}^{N_{1}} \sum_{n_{2}=0}^{N_{2}} e^{-iw(n_{1}T_{1}+n_{2}T_{2})} \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} Box(y_{1},y_{2})e^{iw(y_{1}+y_{2})}dy_{1}dy_{2}$$

$$= \left(\sum_{n_{2}=0}^{N_{1}} \sum_{n_{2}=0}^{N_{2}} e^{-iw(n_{1}T_{1}+n_{2}T_{2})}\right) \mathcal{F}^{-1}\{Box\}(w_{1},w_{2})$$

$$= \left(\sum_{n_{2}=0}^{N_{1}} e^{-iwn_{1}T_{1}}\right) \left(\sum_{n_{2}=0}^{N_{2}} e^{-iwn_{2}T_{2}}\right) \mathcal{F}^{-1}\{Box\}(w_{1},w_{2})$$

$$= (p_{1}+iq_{1})(p_{2}+iq_{2})\mathcal{F}^{-1}\{Box\}(w_{1},w_{2})$$

$$= (p_{1}p_{2}-q_{1}q_{2})+i(p_{1}p_{2}+q_{1}q_{2}))\mathcal{F}^{-1}\{Box\}(w_{1},w_{2})$$

$$= (p+iq)\mathcal{F}^{-1}\{Box\}(w_{1},w_{2})$$

Where we define $p := (p_1p_2 - q_1q_2)$ and $q := (p_1p_2 + q_1q_2)$. For this identity we used green's integration rule which allowed us to split the double integral to the product of two single integrations. Also, we used the definition of the 2-dimensional inverse Fourer transform of the Box-function. We applied the same substituation like we did in for the 1 dimensional case, but this time twice, once for each variable seperately. The last step, substituting with p and q will be useful later in the implementation. The insight should be, that the product of two complex numbers is again a complex number. We will have to compute the absolute value of $\mathcal{F}^{-1}\{f\}(w_1, w_2)$ which will then be equal $(p^2 + q^2)^{\frac{1}{2}} |\mathcal{F}^{-1}\{Box\}(w_1, w_2)|$

2.3.6 Final Expression

As the last step of our series of derivations, we plug all our findings together to one big equation in order to compute the colors in the CIE_XYZ colorspace:

For a given heigh-field h(x,y), representing a small patch of the nanostructure of our surface, the resulting CIE_XYZ caused by the effect of diffraction can be computed like the following: Let $P(u, v, \lambda) = F_{fft}^{-1} \{i^n h^n\} (\frac{2\pi u}{\lambda}, \frac{2\pi v}{\lambda})$

Let
$$P(u, v, \lambda) = F_{fft}^{-1}\{i^n h^n\}(\frac{2\pi u}{\lambda}, \frac{2\pi v}{\lambda})$$

$$\begin{pmatrix}
X \\
X \\
Z
\end{pmatrix} = C(w_i, w_r) \int_{\lambda} \sum_{n=0}^{N} \frac{(wk)^n}{n!} \sum_{r} \sum_{s} |P(u - w_r, v - w_s, \lambda)|^2$$

$$\phi(u - w_r, v - w_s) \begin{pmatrix} S_x(\lambda) \\ S_y(\lambda) \\ S_z(\lambda) \end{pmatrix} d\lambda$$
(32)

where $\phi(x,y)=\pi e^{-rac{x^2+y^2}{2\sigma_f^2}}$ is the gaussian window, where w_s and w_r are ... explain them

3 Implementation

how to discretize from final derivation to computation? what do we have to precompute, what during runtime? how does the final algorithm look like explain shaders: vertex(geometriy, precomp) - and fragment-shader(in local space-tspace) how from cie_xyz to cie_rgb how gamma correction how texturing can we do better?

TODO: explain that there is the jrtr and the scene code - what are their responsibilities.

shader

In computergraphics, we are interested in rendering a given scene containing our 3d geometries by using so called shader programs. The purpose of such programs, which run directly on the gpu hardware device, is to compute the colorization and illumination of the objects living in our scene. This computation happens in several stages and depends on the provided input parameters like the camera, light sources, objects material constants and the desired rendering effect one is interested in to model. The shader stages are also modeled as small little programs, the so called vertex-, geometry-and fragment-shaders. Those stages are applied within the rendering pipeline sequencially.

Our shaders are written in GLSL, developed for OpenGl. The decission for using OpenGl has been made since the underlying framework which is responsible for the precomputation of all scene date is based on a framework written in Java using JOGL in oder to communicate with the GPU and precompute all the relevant scene data. This framework, the so called jrtr framework has been developed as an exercise during the class computer graphics held by M. Zwicker which I attended in autumn 2012. The framework itself has been extended during this thesis quite a lot. Further, there are also some precomputations involved, performed in matlab. This is basically addressing all the required precomputations for the provided heigh-fields, refering to computation of the inverse two dimensional Fourier transformations which are further explained within this chapter.

It's noteworthy that all the vertices are processed within the vertexshader, whereas the fragement shader's responsibility is to perfrom pixelwise rendering, using the input from the vertex shader. Just remember, fragements are determined by a triple of vertices, hence each pixel has assigned a trilinear interpolated value of all input parameters of its spanning vertices. Usually, all necessary transformations are applied vertex-wise, considering the vertex-shader as the precomputation stage for the later rendering within the rendering pipeline, in the fragment-shader. In the geometry shader, new vertices around a considered vertex can be created, this is useful for debugging - displaying normals graphically for example.

In this section we are going to explain how to get a fragment-shader from our findings for our BRDF formultion from the last section. this fragment-shader will render the effect of diffraction on our given geometry pixelwise. Therefore, the quality of diffraction depends on the number of pixels we are going to use for the rendering process and this is directly determined by the resolution of the canvas in which the rendered images are being displayed. But, before we can start formulating our fragment-shader we first have to write our vertex shader which does all the precomputations.

By the end of the day we will end up with two different shaders, one which basically samples the whole lambda space using a gaussian window. This shader will be modeling the effect of diffraction completely but will also be rather slow. The other shader will use a gaussian window too but will just use a few wavenumber for the sampling process. Furthermore, this shader will thread specularity seperatly as a special case which will be more like an approximation.

tell how we are going to sample - uniformly along lambda - explain drawback of this approach - explain possible solutions for this issue. maybe refer to reference shader or leave this for the disscusion part.

3.1 Setup

explain geometry computation explain light(source) setup explain factories explain camera setup explain how materials are stored explain how assigned to jrtr explain how passed to glsl shader - see computer graphics slides maybe show schematically the architecture

3.2 Precomputations in Matlab

explain matlab code explain shifts explain what will be outputed

3.3 jrtr Framework

explain how this will work

3.4 GLSL Diffraction Shader

start using the final findings from chapter 2 and substitute explain how all the components are computed and why they are computed like this.

Algorithm 1 Vertex diffraction shader

for each $Vertex v \in Shape do$ end for

Algorithm 2 Fragment diffraction shader

```
foreach Pixel p \in Fragment do
    BRDF_{XYZ}, BRDF_{RGB} = vec4(0.0)
    (u, v, w) = \hat{\mathbf{k_1}} - \hat{\mathbf{k_2}}
    for (\lambda = \lambda_{min}; \lambda \leq \lambda_{max}; \lambda = \lambda + \lambda_{step}) do
        k = \frac{2\pi}{\lambda}
         (w_u, w_v) = (ku, kv)
        w_{color} = (S_x(\lambda), S_y(\lambda), S_z(\lambda))
         for (r) do
             for (s) do
                  coords = getLookUpCoord(r, s)
                  P = taylorApprox(coords, k, w)
                  w_{r,s} = gaussianWeight(dist)
                  scale_{pq} = pqFactor(w_u, w_v)
                  P* = scale_{pq}
                  P_{abs} = |P|^{2}
                  P_{abs}* = w_{r,s}
                  BRDF_{XYZ} + = vec4(P_{abs} * w_{color}, 0.0)
             end for
        end for
    end for
    BRDF_{XYZ} = BRDF_{XYZ} * C(\hat{\mathbf{k_1}}, \hat{\mathbf{k_2}}) * shadowF
    BRDF_{XYZ}.xyz = D_{65} * M_{XYZ-RGB} * BRDF_{XYZ}.xyz
    BRDF_{RGB}.xyz = D_{65} * M_{XYZ-RGB} * BRDF_{XYZ}.xyz
    BRDF_{RGB} = gammaCorrect(BRDF_{RGB})
end for
```

4 Data Acquisition and Evaluation

what is this chapter about how is evaluation perfromed our shader

4.1 Diffraction Grating

Gratings may be of the reflective or transmissive type, analogous to a mirror or lens respectively. A grating has a zero-order mode (where m=0), in which there is no diffraction and a ray of light behaves according to the laws of reflection and refraction the same as with a mirror or lens respectively.

An idealised grating is considered here which is made up of a set of slits of spacing d, that must be wider than the wavelength of interest to cause diffraction. Assuming a plane wave of wavelength with normal incidence (perpendicular to the grating), each slit in the grating acts as a quasi pointsource from which light propagates in all directions (although this is typically limited to a hemisphere). After light interacts with the grating, the diffracted light is composed of the sum of interfering wave components emanating from each slit in the grating. At any given point in space through which diffracted light may pass, the path length to each slit in the grating will vary. Since the path length varies, generally, so will the phases of the waves at that point from each of the slits, and thus will add or subtract from one another to create peaks and valleys, through the phenomenon of additive and destructive interference. When the path difference between the light from adjacent slits is equal to half the wavelength, /2, the waves will all be out of phase, and thus will cancel each other to create points of minimum intensity. Similarly, when the path difference is , the phases will add together and maxima will occur. The maxima occur at angles m, which satisfy the relationship dsin m/=|m|where m is the angle between the diffracted ray and the grating's normal vector, and d is the distance from the center of one slit to the center of the adjacent slit, and m is an integer representing the propagation-mode of interest.

Thus, when light is normally incident on the grating, the diffracted light will have maxima at angles θ_m given by:

$$dsin(\theta_m) = m\lambda$$

It is straightforward to show that if a plane wave is incident at any arbitrary angle i, the grating equation becomes:

$$d(sin(\theta_i) + sin(\theta_m)) = m\lambda$$

When solved for the diffracted angle maxima, the equation is:

$$sin(\theta_m) = \left(\frac{m\lambda}{d} - sin(\theta_i)\right)$$

The light that corresponds to direct transmission (or specular reflection in the case of a reflection grating) is called the zero order, and is denoted m = 0. The other maxima occur at angles which are represented by non-zero integers m. Note that m can be positive or negative, resulting in diffracted orders on both sides of the zero order beam.

This derivation of the grating equation is based on an idealised grating. However, the relationship between the angles of the diffracted beams, the grating spacing and the wavelength of the light apply to any regular structure of the same spacing, because the phase relationship between light scattered from adjacent elements of the grating remains the same. The detailed distribution of the diffracted light depends on the detailed structure of the grating elements as well as on the number of elements in the grating, but it will always give maxima in the directions given by the grating equation.

$$\forall \begin{pmatrix} x \\ y \\ z \end{pmatrix} \in \mathbb{R}^3 : \exists r \in [0, \infty) \exists \phi \in [0, 2\pi] \exists \theta \in [0, \pi] \text{ s.t.}$$

$$\begin{pmatrix} x \\ y \\ z \end{pmatrix} = \begin{pmatrix} rsin(\theta)cos(\phi) \\ rsin(\theta)sin(\phi) \\ rcos(\theta) \end{pmatrix}$$

4.2 Snake Skin Parameters

5 Results

differece of this shader compared to evaluation shader

6 Conclusion

explain why we did our derivations explain why our approach is a good idea explain how the straight foreward approach would behave compared to our approach, computing the fourier transformations straight away. explain what we achieved, summary say something about draw-backs and about limitations of current apporach say something about the ongoing paper

6.1 Further Work

6.1.1 Sources

- [1] http://en.wikipedia.org/wiki/Ratio_test
- [2] http://math.jasonbhill.com/courses/fall-2010-math-2300-005/lectures/taylor-polynomial-error-bounds