

# Wavelets on irregular point sets

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In this article we review techniques for building and analysing wavelets on irregular point sets in one and two dimensions. We discuss current results on both the practical and theoretical sides. In particular, we focus on subdivision schemes and commutation rules. Several examples are included.

Keywords: multiresolution; wavelets; irregular grid; subdivision; mesh; commutation

## 1. Introduction

Wavelets are a versatile tool for representing general functions and datasets, and they enjoy widespread use in areas as diverse as signal processing, image compression, finite-element methods and statistical analysis (among many others). In essence we may think of wavelets as *building blocks* with which to represent data and functions. The particular appeal of wavelets derives from their representational and computational *efficiency*: most datasets exhibit correlation both in time (space) and frequency, as well as other types of structure. These can be modelled with high accuracy through sparse combinations of wavelets. Wavelet representations can also be computed *fast*, because they can be built using multiresolution analysis and subdivision.

Traditionally, wavelet functions  $\psi_{j,m}$  are defined as translates and dilates of one particular function, the mother wavelet  $\psi$ . We refer to these as first-generation wavelets. This paper is concerned with a more general setting in which wavelets need not—and, in fact, cannot—be translates and dilates of one or a few templates. Generalizations of this type were called second-generation wavelets in Sweldens (1997); they make it possible to reap the benefit of wavelet algorithms in settings with irregularly spaced samples, or on 2-manifolds which cannot be globally parametrized to the plane. In generalizing wavelet analysis to these more general settings one would like to preserve many of the properties enjoyed by first-generation wavelets. In particular, they should still be associated with fast algorithms and have appropriate smoothness and localization properties. In addition, they should be able to characterize various functional spaces of interest. In this paper we shall be mostly

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Figure 1. Subdivision is used to generate a smooth curve starting from a coarse description.

concerned with fast algorithms, localization and smoothness; we will not address function-space characterizations. Note though that the smoothness of the wavelets is related to their ability to form unconditional bases for certain function spaces (Dahmen 1996; Donoho 1992).

The key to generalizing wavelet constructions to these non-traditional settings is the use of generalized *subdivision schemes*. The first-generation setting is already connected with subdivision schemes, but they become even more important in the construction of second-generation wavelets. Subdivision schemes provide fast algorithms, create a natural multiresolution structure and yield the underlying scaling functions and wavelets we seek.

Subdivision is a technique originally intended for building smooth functions starting from a coarse description. In this setting there is no need for irregular grids, as one is free to choose the finer grid to be regular. However, we intend to use subdivision as part of an entire multiresolution analysis which starts from a finest irregular grid. This finest grid is gradually 'coarsified'; subdivision then gives an approximation of the original data by extrapolating the reduced data on the coarser grid back to the original finest grid. In such a setting the geometry of the grids is fixed by the finest irregular grid and the coarsification procedure; thus subdivision on irregular grids is called for.

**Remark 1.1.** Another approach would be to resample the original finest level data on a regular grid and use first-generation wavelets. Resampling, however, can be costly, introduce artefacts and is generally impossible in the surface setting. Therefore we choose to work on the original grid.

#### (a) One-dimensional subdivision

The main idea behind subdivision is the iteration of upsampling and local averaging to build functions and intricate geometrical shapes. Originally, such schemes were studied in computer-aided geometric design in the context of corner cutting (de Rham 1956; Chaikin 1974) and the construction of piecewise polynomial curves, e.g. the de Casteljau algorithm for Bernstein–Bézier curves (de Casteljau 1959) and algorithms for the iterative generation of splines (Lane & Riesenfield 1980; De Boor 1978). Later subdivision was studied independently of spline functions (Dyn *et al.* 1987; Dubuc 1986; Deslauriers & Dubuc 1987; Cavaretta *et al.* 1991; Cavaretta & Micchelli 1987, 1989) and the connection to wavelets was made (Mallat 1989; Daubechies 1988).

For example, figure 1 demonstrates the application of the four-point scheme. New points are defined as local averages of two old points to the left and two old points to the right with weights  $\frac{1}{16}(-1,9,9,-1)$ .

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Figure 2. Regular, semiregular and irregular grid hierarchies in one dimension.

In the case of spline functions, smoothness follows from simple algebraic conditions on the polynomial segments at the knots. However, in the general setting convergence and smoothness of the limit function are harder to prove. Various approaches have been explored to find the Hölder exponent of the limit function, or to determine its Sobolev class. Early references in this context are Dubuc (1986), Dyn *et al.* (1987, 1990b), Deslauriers & Dubuc (1987), Micchelli & Prautzsch (1987), Daubechies & Lagarias (1991), Cavaretta *et al.* (1991), Rioul (1992), Villemoes (1994) and Eirola (1992). These studies and their results all rely on regular, i.e. equispaced, grids. The analysis uses tools such as the Fourier transform, spectral analysis and the commutation formula.

In this paper we focus on irregular point sets. To describe the settings we are interested in, we distinguish three types of refinement grids: regular, semiregular and irregular (see figure 2). A *regular* grid has equidistant points on each level, and, each time a new point is inserted, it is placed exactly between two old points. For example, the curve shown in figure 1 is parametrized over a regular grid. A *semiregular* grid (middle, figure 2) starts with an irregular coarse grid and adds new points at parameter locations midway between successive old points. Thus the finer grids are locally regular except around the original coarsest level points. In *irregular* grids (right) parameter locations of new points need not be midway between successive old points. Note that regular grids are translation and dilation invariant, while semiregular grids are locally dilation invariant around coarsest level vertices and irregular grids generally possess no invariance property.

Similarly the weights used in subdivision schemes come in three categories: uniform, semi-uniform and non-uniform. Uniform schemes like the four-point scheme of figure 1 correspond to first-generation wavelets and use the same subdivision weights within a level and across all levels; they are typically used on regular grids or grids which can be smoothly remapped to a regular grid. Semi-uniform schemes are used on semiregular grids; they vary the weights within each level (special weights are used in the neighbourhood of the coarsest level points), but the same weights are used across levels. Such schemes are sometimes referred to as stationary. Wavelets and subdivision schemes on an interval also fall in this category. Non-uniform schemes use varying weights within and across levels and correspond to the second-generation setting.

Almost all work on smoothness for non-regular grids concerns the semiregular grids with semi-uniform subdivision schemes. Because translation-invariance is lost, Fourier-transform-based arguments can no longer be used. However, since the same weights are used on successive levels, one has dilation invariance around coarsest level points and can reduce the smoothness analysis to the study of spectral properties of certain fixed matrices. In Warren (1995, unpublished research<sup>†</sup>) it is shown that the semi-uniform version of the four-point scheme on a semiregular grid yields a  $C^1$  limit function.

† Available from www.cs.rice.edu/~jwarren.



Figure 3. An example why non-uniform subdivision is needed. The limit function with uniform (left), non-uniform (right) subdivision. The same irregular grid is used in both figures.



Figure 4. Two-dimensional loop subdivision is used to generate smooth surfaces from a coarse description.

In the irregular case the subdivision scheme must become non-uniform to account for the irregularity of the associated parameter locations. This is illustrated in figure 3, which shows the limit functions of the uniform four-point rule (left) and non-uniform four-point rule (Sweldens & Schröder 1996) (right); both use the same irregular grid.

The study of irregular subdivision is not only theoretically interesting, but also of great importance in practical applications. For example, in the semiregular setting, one can use adapted weights to better control the shape of a curve (Kobbelt & Schröder 1997) or surface (Zorin *et al.* 1996). More importantly, in many practical set-ups we start with samples associated to a very fine, but irregular, grid. Now the main task for subdivision is not further refinement, but rather aid in a multiresolution analysis on *coarser* grids. The wavelet and scaling functions from the coarsest level are generated with a subdivision scheme with new points which are no longer parametric midpoints, but are dictated by the finest level grid on which the data were originally sampled. Even though the actual number of levels is always finite for any concrete application of these methods, the asymptotic behaviour of irregular subdivision is still relevant as the finest and coarsest level could be arbitrarily far apart.

In these settings smoothness results become much harder to obtain. Because the subdivision weights vary within a level, the Fourier transform can no longer be used, and because they vary across levels, even spectral analysis cannot help. In this paper we discuss some tools that can be used to analyse smoothness; in particular, we demonstrate that the commutation formula still holds and becomes a critical tool for smoothness analysis.

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Figure 5. Sections of regular, semiregular and irregular triangle grids in two dimensions.

#### (b) Two-dimensional subdivision

The two-dimensional setting appears in the context of generating smooth surfaces (see figure 4). Here regular grids are too restrictive. For example, tensor product settings are only applicable for surfaces homeomorphic to a plane, cylinder or torus due to the Euler characteristic. Historically, this challenge was addressed by generalizing traditional spline patch methods to the semiregular biquadratic (Doo & Sabin 1978), bicubic (Catmull & Clark 1978) and quartic-box-spline settings (Loop 1987). Similar to the one-dimensional setting, researchers also developed interpolating constructions (Dyn et al. 1990a; Zorin et al. 1996; Kobbelt 1996). All these settings (and others since; for an overview see Schröder & Zorin (1998)) proceed by applying quadrisection to an initial mesh consisting of either quadrilaterals or triangles and thus belong to the semiregular setting. The weights used in the subdivision scheme are semi-uniform since they take into account the local neighbourhood structure of a vertex, i.e. how many edge neighbours a given vertex has. As in the one-dimensional semiregular setting, spectral analysis is the key to understanding the smoothness of these constructions. We refer to Reif (1995), Warren (unpublished research), Zorin (1996) and Schweitzer (1996) for more details.

The irregular setting appears in two dimensions just as in the one-dimensional case when some finest irregular level is presented on input and the main task is to build a multiresolution analysis on *coarser* levels. In this case, however, we can no longer define downsampling as simply retaining every other sample. This brings us to the realm of mesh simplification; we postpone the discussion of mesh simplification and the construction of appropriate non-uniform subdivision operators to § 3.

## (c) Overview

This paper summarizes the results obtained in Daubechies *et al.* (1998, *b*), Guskov (1998) and Guskov *et al.* (1999). We start with the one-dimensional results of Daubechies *et al.* (1998, *b*). We show that even simple subdivision rules, such as cubic Lagrange interpolation, can lead to very intricate subdivision operators. To control these operators, we use commutation: because the subdivision scheme maps the space of cubic polynomial sequences to itself, we can define derived subdivision schemes for the divided difference sequences. These simpler schemes can then be used to prove growth bounds on divided differences of some order, corresponding to smoothness results for the limit function of the original scheme. The commutation formula enables us to control smoothness and is the key to the construction of wavelets associated with the subdivision scheme.

In Guskov (1998), inspiration from the one-dimensional analysis is used to tackle the much more complex two-dimensional case. Again, differences and divided differences are introduced, which can be computed from level to level with their own

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derived subdivision scheme. Control on the growth of these divided differences then leads to smoothness results. In practice, finding the right ansatz for irregular subdivision in the two-dimensional setting is much harder than in the already difficult one-dimensional case. Finally, we show how irregular subdivision schemes can be used in multiresolution pyramids for two-dimensional meshes embedded in  $\mathbb{R}^3$  and review several applications from Guskov *et al.* (1999). The 'wavelets' associated with these schemes are overcomplete and are related to frames rather than bases.

## 2. The one-dimensional case

#### (a) Multilevel grids

Consider grids  $X_j$ , which are strictly increasing sequences of points  $\{x_{j,k} \in \mathbb{R} \mid k \in \mathbb{Z}\}$ , and which are consecutive binary refinements of the initial grid  $X_0$ , i.e.  $X_j \subset X_{j+1}$  and  $x_{j+1,2k} = x_{j,k}$  for all j and k. Thus in every refinement step we insert one odd indexed point  $x_{j+1,2k+1}$  between each adjacent pair of 'even' points  $x_{j,k} = x_{j+1,2k}$  and  $x_{j,k+1} = x_{j+1,2k+2}$ , as in figure 2. We define  $d_{j,k} := x_{j,k+1} - x_{j,k}$ . We shall also use the term grid size on level j, for the quantity  $d_j := \sup_k d_{j,k}$ . As  $j \to \infty$  we want the grids to become dense, with 'no holes left'; this translates to the requirement that the  $d_j$  be summable.

#### Remark 2.1.

- (1) The above multilevel grids are called *two-nested*. One can also consider more general irregular grids such as *q*-nested grids, where we insert q-1 new points in between old points or even non-nested but 'threadable' grids. See Daubechies *et al.* (1998) for more details on this.
- (2) In case the ratio between the lengths of any two neighbouring intervals is globally bounded, we call the grid homogeneous. An example of an irregular two-nested grid that is not homogeneous is built by  $x_{j+1,2k+1} = \beta x_{j,k} + (1 \beta)x_{j,k+1}$ , where  $\beta$  is a fixed parameter satisfying  $0 < \beta < 1$ . This is an example of a dyadically balanced grid: the ratio between the lengths of two 'sibling' intervals  $d_{j,2l}$  and  $d_{j,2l+1}$  is bounded. However, the ratio between  $d_{j,-1} = \beta^j$  and  $d_{j,0} = (1 \beta)^j$  is unbounded.

#### (b) Subdivision schemes

Subdivision starts with a set of initial function values  $f_0 = \{f_{0,k}\}$ , which live on the coarsest grid  $X_0$ . The subdivision scheme S is a sequence of linear operators  $S_j, j \ge 0$ , which iteratively computes values  $f_j = \{f_{j,k}\}$  on the finer grids via the formula  $f_{j+1} = S_j f_j$ , or

$$f_{j+1,l} = \sum_{k} S_{j,l,k} f_{j,k}.$$

We consider only *local* schemes in the sense that the above summation has a globally bounded number of terms centred around k = 2l. Subdivision gives us values defined on the grid points  $x_{j,k}$ . By connecting these points we can define a piecewise linear function  $f_j(x)$  (see figure 6). Our ambition is to synthesize a continuous limit function



Figure 6. On the left an interpolating scheme is applied to function values; the new function values are shown as open circles. On the right, arrows show the dependencies in the computation of those values; the vertical dashed arrows indicate that function values which were already assigned are kept unchanged in the subdivision, because this is an interpolating scheme.

 $\phi(x)$  as the pointwise limit for  $j \to \infty$  of  $f_j(x)$ . We are interested in the existence and smoothness of  $\phi(x)$ .

The subdivision coefficients  $S_{j,k,l}$  will depend on the application one has in mind. We pointed out in the introduction that one cannot simply stick with the coefficients from the regular case; typically the coefficients need to be spatially varying, and will be linked to the spatial variation of the grid.

One such subdivision scheme which allows for a spatial interpretation is Lagrangian interpolating subdivision (Dubuc 1986; Dyn *et al.* 1987; Deslauriers & Dubuc 1987, 1989). Here, the value  $f_{j+1,2k+1}$  at a new point is found by defining a polynomial which interpolates the points  $(x_{j,l}, f_{j,l})$  for l in the neighbourhood of k, and evaluating this polynomial at  $x_{j+1,2k+1}$  (see figure 7). In the regular cubic case, this corresponds to the standard four-point scheme, with

$$S_{j,2k+1,k} = S_{j,2k+1,k+1} = \frac{9}{16}$$
 and  $S_{j,2k+1,k-1} = S_{j,2k+1,k+2} = -\frac{1}{16}$ .

In the irregular setting the coefficients are a non-trivial quotient of cubic polynomials in the  $x_{j,k}$  (see Daubechies *et al.* 1999).

Lagrangian subdivision is *interpolating* in the sense that in each subdivision step the values at the even grid points are kept, i.e.  $f_{j+1,2k} = f_{j,k}$ , and the limiting function thus interpolates the original data  $\phi(x_{0,k}) = f_{0,k}$ . For non-interpolating or *approximating* schemes, the  $f_{j+1,2k}$  can differ from  $f_{j,k}$  (see figure 8).

## (c) Smoothness results

To derive smoothness estimates, we use Lemarié's *commutation formula* idea, generalized to the present irregular setting. (Note that this is similar to Rioul (1992) and Dyn *et al.* (1991), who studied the regular case.) For the cubic Lagrange interpolation example, this amounts to introducing divided difference sequences,

$$f_{j,k}^{[1]} := \frac{f_{j,k+1} - f_{j,k}}{x_{j,k+1} - x_{j,k}},$$

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Figure 7. Cubic Lagrangian interpolation. The value  $f_{j+1,2k+1}$  at the odd grid point  $x_{j+1,2k+1}$  is obtained by evaluating a cubic polynomial P(x) interpolating values at four neighbouring even grid points  $x_{j+1,2k-2} = x_{j,k-1}, \ldots, x_{j+1,2k+4} = x_{j,k+2}$ .



Figure 8. The top row shows the limit function and weights of the cubic Lagrange interpolation scheme; the bottom row illustrates the non-interpolating subdivision scheme producing cubic B-splines.

and observing that the  $f_{j,k}^{[1]}$  are also related by local subdivision, i.e. there exists a local subdivision scheme with entries  $S_{j,l,k}^{[1]}$  so that

$$f_{j+1,l}^{[1]} = \sum_{k} S_{j,l,k}^{[1]} f_{j,k}^{[1]}.$$

The existence of the  $S_j^{[1]}$  follows from the fact that every  $S_j^{[0]} := S_j$  maps a constant sequence to itself (see Daubechies *et al.* 1999). (It is clear that if  $S^{[0]}$  does not leave constants invariant, then  $S^{[1]}$  cannot exist, since it would need to map the zero sequence to a non-zero result.) Moreover, one can show that if the  $f_j^{[1]}$  'converge' to a continuous function  $\phi_1$ , then the  $f_j^{[0]} := f_j$  also converge, to a continuously differentiable function  $\phi$ , and that  $\phi' = \phi_1$ . (For details, see Daubechies *et al.* (1998, *b*).) This is the essence of the commutation idea:

$$\begin{array}{ccc} f_j & \xrightarrow{\text{divided difference}} & f_j^{[1]} \\ & & \downarrow \text{limit} & & \downarrow \text{limit} \\ \phi & \xrightarrow{\text{differentiation}} & \phi_1 = \phi' \end{array}$$

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It turns out that one can also consider higher-order divided differences; for the cubic Lagrange interpolation case, one can go up to fourth-order differences because  $S_j^{[0]}$  maps cubic polynomials sampled at  $x_{j,k}$  to cubic polynomials sampled at  $x_{j+1,k}$ . These  $f_{j,k}^{[4]}$  no longer converge, but we can control their growth, and this helps us prove that  $\phi_1$  is continuous, and  $\phi$  continuously differentiable. In fact, detailed (and rather technical) estimates in Daubechies *et al.* (1999) show that, for homogeneous grids,

$$|f_{j,k}^{[4]}| \leqslant C \frac{\lambda^j}{d_{j,k}^3},$$

where  $\lambda < 1$  is determined by the bound on the ratio between neighbouring interval lengths. Once such a bound is known, a general theorem (see theorem 4 in Daubechies *et al.* (1999)) can be used to prove that  $\phi \in C^{2-\epsilon}$ . This result is optimal in the sense that even in the regular case better smoothness cannot be obtained.

#### Remark 2.2.

- (1) This result for cubic Lagrange interpolation on homogeneous grids can be extended to grids that are dyadically balanced only. The analysis becomes much more delicate.
- (2) A similar approach can be used for non-interpolating subdivision. In that case it turns out that one has to use appropriately defined divided differences, which are different from the 'standard' definition. See Daubechies *et al.* (1998) for a complete discussion of this situation.

#### (d) Wavelets

Wavelets at level j are typically used, in the regular case, as building blocks to represent any function in the multiresolution analysis that 'lives' in the (j + 1)st approximation space  $V_{j+1}$ , but not in the coarser resolution approximation space  $V_j \subset V_{j+1}$ . One can introduce similar wavelets in the present irregular setting. The scaling functions  $\phi_{j,k}$  are the limit functions obtained from starting the subdivision scheme at level j, from the 'initial' data  $f_{j,l} = \delta_{l,k}$ , and refining from there on. Under appropriate assumptions on the subdivision operators  $S_j$ , the  $\phi_{j,k}$  are independent;  $V_j$  is the function space spanned by them. Clearly,  $V_j \subset V_{j+1}$ . As in the regular case, there are many different reasonable choices for complement spaces  $W_j$  (which will be spanned by the wavelets at level j) that satisfy  $V_{j+1} = V_j \oplus W_j$ .

When the scaling functions are interpolating as in the Lagrangian case, i.e.

$$\phi_{j,k}(x_{j,k'}) = \delta_{k,k'},$$

then a simple choice for a wavelet is given by  $\psi_{j,m} = \phi_{j+1,2m+1}$ , i.e. the wavelet is simply a finer-scale scaling function at an odd location. This is sometimes called an *interpolating* wavelet. This is in general not a very good wavelet as it does not have any vanishing moments. It can be turned into a wavelet with vanishing moments using the lifting scheme (Sweldens & Schröder 1996).

Another way to select a complement space  $W_j$  is to use commutation between two biorthogonal multiresolution hierarchies,  $V_j$  and  $\tilde{V}_j$ . If both are associated to local

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subdivision schemes, then the biorthogonality of the  $\phi_{j,k}$  and  $\phi_{j,l}$  imposes consistency requirements on the  $S_j$  and  $\tilde{S}_j$ . Commutation can be used, as in the regular case, to pass from one dual pair of multiresolution analyses to another, by operations related to differentiating and integrating, respectively. For instance, the above choice of an interpolating wavelet corresponds formally to letting the dual scaling function be a Dirac. Applying the commutation rule each time reduces the order of the scaling functions, but increases the order of the dual scaling function. In particular, the Dirac will become a box and later on a general B-spline. It turns out that there is a natural definition of wavelets  $\psi_{j,k}$  and  $\tilde{\psi}_{j,k}$  corresponding to the dual multiresolution structures. It is shown in Daubechies *et al.* (1998) that, as in the regular case, the new wavelet after commutation is the derivative of the old wavelet and the new dual wavelet is the integral of the old dual wavelet. By repeatedly applying commutation starting from the Lagrangian setting, one can thus build the entire family of biorthogonal compactly supported irregular B-spline wavelets and their duals (Daubechies *et al.* 1998).

### 3. The two-dimensional case

We mentioned in the introduction that the importance of the irregular setting arises from the practical need to coarsify in settings in which the initial input is given as a function over a fine triangulation of the plane (functional setting) or as a triangulation of a 2-manifold (surface setting). In the one-dimensional setting, the downsampling operation to create a coarser level is straightforward as we can simply 'skip' every other sample. In the irregular two-dimensional setting, downsampling is much less straightforward. Before delving into the details of irregular two-dimensional subdivision, we first discuss a number of approaches which can be employed to define irregular downsampling in the surface setting. This problem has received a lot of attention in computer graphics, where it is generally referred to as polygonal simplification.

## (a) Polygonal simplification

In polygonal mesh simplification, the goal is to simplify a given (triangulated) mesh  $\mathcal{M}^L = (\mathcal{P}^L, \mathcal{K}^L)$  into successively coarser, homeomorphic meshes  $(\mathcal{P}^l, \mathcal{K}^l)$  with  $0 \leq l < L$ , where  $(\mathcal{P}^0, \mathcal{K}^0)$  is the coarsest or base mesh. Here  $\mathcal{P}^l$  is a set of l point positions, while  $\mathcal{K}^l$  encodes the topological structure of the mesh and consists of triples  $\{i, j, k\}$  (triangles), pairs  $\{i, j\}$  (edges) and singletons  $\{i\}$  (vertices). The goal now is to allow certain topological operations on  $\mathcal{K}^l$  which preserve the manifold property and genus of the mesh. These changes go hand in hand with geometric changes which are typically subject to an approximation quality criterion.

Several approaches for such mesh simplification have been proposed (the interested reader is referred to the excellent survey by Heckbert & Garland (1997) for more details). The most popular methods are the so-called 'progressive meshes' (PM). In a PM construction a sequence of edge collapses is prioritized based on the error it introduces. An edge collapse brings the endpoints of the chosen edge into coincidence, in the process removing two triangles, three edges and one vertex (in the case of interior edges). The point location of the merged vertex can be chosen so as to minimize some error criterion with respect to the original mesh. The error can be



Figure 9. If an irregular finely detailed mesh is given, the first task in building a multiresolution analysis is coarsification.

measured in various norms such as  $L^\infty$  (symmetric Haussdorff distance),  $L^2$  and Sobolev norms.

For our purposes we are using a PM construction based on half-edge collapses, i.e. the point position for the collapsed edge is one of its end points. This results in a mesh hierarchy which is interpolating in the sense that the point position sets  $\mathcal{P}^l$  are nested. There are several possible ways to define levels of a hierarchy. The most flexible way treats a single half-edge collapse operation as defining a level. In contrast to the usual wavelet setting this results in a linear, rather than logarithmic, number of levels.

Before going to the surface case, we first consider the functional setting and then treat the surface setting as three instances of a functional setting.

#### (b) Functional setting: multivariate commutation formula

Just as in the one-dimensional case, irregular multivariate subdivision schemes act on sequences whose elements are associated with irregular parameter locations. We introduce levels numbered  $0, 1, \ldots$  with level 0 corresponding to the coarsest scale. Within each level *n*, the collection of all parameter locations constitute an irregular grid  $\chi_n$ .

We can now introduce a subdivision scheme S as a sequence of linear operators  $S_n, n \ge 0$ , which iteratively compute sequences  $f_n$  defined on  $\chi_n$ , starting from some coarsest level data  $f_0$  via

$$f_{n+1} = S_n f_n.$$

In the one-dimensional setting we analysed the regularity of the functions produced by subdivision through the behaviour of properly defined divided differences. We proceed similarly for the irregular two-dimensional setting. Let  $\mathcal{D}_n^{[p]}$  denote the operator which maps the data sequence  $f_n$  into the corresponding sequence  $f_n^{[p]}$  of divided differences of order p, that is  $f_n^{[p]} = \mathcal{D}_n^{[p]} f_n$ . We say that there exists a derived subdivision scheme  $S^{[p]}$  satisfying the commutation formula if the sequences  $f_n^{[p]}$  are related via the relation  $f_{n+1}^{[p]} = S_n^{[p]} f_n^{[p]}$ , where the  $S_n^{[p]}$  constitute a local bounded subdivision scheme. Thus we can write

$$\mathcal{D}_{n+1}^{[p]}S_n = S_n^{[p]}\mathcal{D}_n^{[p]}.$$



difference of normals lies in parameter plane

Figure 10. Second differences are associated with an edge. Since they are the difference of two adjacent triangle normals (first divided differences), one can see that the second differences are orthogonal to the common edge in the parameter plane.

We then prove that the bounds on the growth of sequences  $f_n^{[p]}$  can be translated into the smoothness estimates for the functions produced by the original subdivision scheme S.

In order to extend this construction to the multivariate case we need to define the multivariate divided differences in such a way that the algebra of the commutation formula works. This is done in Guskov (1998) for a class of polynomial reproducing subdivision schemes. It is also shown there that, for multilevel grids satisfying some natural conditions, the bounds on the growth of these divided differences can be used to analyse the regularity of functions produced by subdivision.

### (c) Constructing a subdivision scheme

In this section we provide a particular example of a subdivision scheme which in the functional setting produces visually smooth functions on irregular triangulations. Our subdivision algorithm relies on minimizing divided differences. Consider a triangle  $\{i, j, k\}$  in the parameter plane with corners  $(x_i, y_i), (x_j, y_j)$  and  $(x_k, y_k)$ , and function values  $f_i$ ,  $f_j$  and  $f_k$ . These three function values define a plane. The gradient to this plane can be seen as a first-order divided difference corresponding to this triangle. The gradient is zero only if the plane is horizontal  $(f_i = f_j = f_k)$ .

Next, we define the second-order differences. They are computed as the difference between two normals on neighbouring triangles and can be thought of as being associated with the common edge (see figure 10, left). It is easy to see that the difference between gradients of two adjacent triangles is orthogonal to their common edge (see figure 10, right). Thus the component  $D_e^2 f$  normal to the edge e can be used for the second-order difference. It depends linearly on the four function values of these two triangles. The coefficients can be found in Guskov (1998) or Guskov *et al.* (1999). The second-order difference operator is zero only if the two triangles lie in the same plane, and one can see that its behaviour is closely related to the dihedral angle.

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Figure 11. From left to right: a portion of the fine mesh; the coarse mesh; function produced by the non-interpolating scheme.

The central ingredient in the design of our subdivision scheme is the use of a nonuniform relaxation operator which inserts new values in such a manner that secondorder differences are minimized. Define a quadratic energy, which is an instance of a discrete fairing functional (Kobbelt 1997):

$$Rf_i = \arg\min \boldsymbol{E}(f_i) = \sum_{e \in \mathcal{K}} (D_e^2 f)^2.$$

Setting  $\partial \boldsymbol{E} / \partial f_i = 0$  yields

Note that if f is a linear function, i.e. all triangles lie in one plane, the fairing functional E is zero. Consequently, linear functions are invariant under R. In particular, R preserves constants from which we deduce that the  $w_{i,j}$  sum to one.

Subdivision is computed one level at a time starting from level  $n_0$  in the PM. Reversing the PM construction back to the finest level adds one vertex  $(x_n, y_n, f_n)$  per level; the non-uniform subdivision is computed one vertex at a time. The position of each new vertex n is computed according to (3.1), using areas and lengths of the original finest level mesh. Next, the immediate neighbours of n are relaxed using (3.1) as well. The ambition of our strategy of minimizing  $D_e^2 f$  is to obtain  $C^1$  smoothness. However, there is currently no ansatz on the bounds of the divided differences to prove regularity of the limit function. Figure 11 shows an irregular grid of 20 493 triangles (left), simplified down to 86 triangles (middle). Now associate the value f = 1 with the centre vertex and 0 with all others. The figure on the right is the result of running the subdivision scheme back to the finest level. Even though the grid is irregular the resulting function appears smooth.

#### (d) Functions on surfaces

In order to build a multiresolution structure on meshes, we first need to introduce the relaxation operator acting on functions defined over triangulated surfaces in three dimensions. We shall follow the strategy of the planar case and introduce second differences for such functions. For this we need to specify some locally consistent parametrization over the support of the difference operator. Consider a triangular mesh  $\mathcal{P}$  in  $\mathbb{R}^3$ , and let  $f: \mathcal{P} \to \mathbb{R}$ . We would like to define the second difference operator  $D_e^2 f$  for an edge e from the triangulation  $\mathcal{P}$ . For this we only need a consistent



Figure 12. Burt–Adelson-style pyramid scheme.

parametrization (i.e. flattening) for two neighbouring triangles at a time. Let the edge  $e = \{i, j\}$  be adjacent to two triangles  $\{i, j, k\}$  and  $\{j, i, l\}$ . We use the 'hinge map' to build a pair of adjacent triangles in the plane. These two triangles in the parameter plane have the same angles and edge lengths as the two triangles in  $\mathbb{R}^3$ . We then define  $D_e^2 f$  as described in the previous section. Using these second differences, it is easy to extend the definition of the relaxation operator and the corresponding subdivision scheme to work with functions defined over triangulated surfaces.

### (e) Burt-Adelson pyramid

For meshes we found it more useful to generalize an oversampled Burt-Adelsontype pyramid (Burt & Adelson 1983) than a critically sampled wavelet pyramid. Let  $(\mathcal{P}^n)$  be some fixed PM hierarchy of triangulated surfaces. We start from the function  $f_N : \mathcal{P} = \mathcal{P}^N \to \mathbb{R}$ , defined on the finest level, and compute a sequence of functions  $(f_n) (n_0 \leq n \leq N)$  as well as oversampled differences  $d_i^{(n)}$  between levels.

Like subdivision, the Burt–Adelson pyramid is computed vertex by vertex. Thus the four critical components of a BA pyramid: presmoothing, downsampling, subdivision and detail computation are done for one vertex n at a time (see figure 12). The presmoothing comes down to applying the relaxation operator to the neighbours of n. Downsampling simply removes the vertex n through a half-edge collapse. We perform subdivision as described above and compute details  $d^{(n)}$  for all neighbours of n.

In order to see the potential of a mesh pyramid in applications, it is important to understand that the details  $d^{(n)}$  can be seen as an approximate frequency spectrum of the mesh. The details  $d^{(n)}$  with large n come from edge collapses on the finer levels and thus correspond to small scales and high frequencies, while the details  $d^{(n)}$  with small n come from edge collapses on the coarser levels and thus correspond to large scales and low frequencies. Hence, the sequence of  $d^{(n)}$  for running n can be seen as an approximate frequency spectrum. Moreover, while the superscript n of an individual detail vector  $d_i^{(n)}$  corresponds to its level/frequency, the subscript i corresponds to its location. Thus we actually have a space-frequency decomposition.

It is theoretically possible to build a critically sampled wavelet transform based on the lifting scheme (Sweldens 1997). The idea is to use an interpolating subdivision scheme which only affects the new vertex and omits the relaxation of the even vertices. Consequently, only one detail per vertex is computed and the sampling is always critical. However, at this point it is not clear how to design updates that make the transform numerically stable. Additionally, interpolating subdivision schemes do not yield very smooth meshes and have unwanted undulations. Therefore, critically sampled wavelet transforms have had limited use in graphics applications.

Wavelets on irregular point sets



Figure 13. Smoothing and filtering of the Venus head. (a) Original; (b) low-pass filter; (c) stopband filter; (d) enhancement filter.

#### 4. Applications

In the surface setting we deal with a triangulated mesh  $\mathcal{P}$  of arbitrary topology and connectivity embedded in three dimensions with vertices  $p_i = (x_i, y_i, z_i)$ . It is important to separate the two capacities the mesh  $\mathcal{P}$  fulfils in our analysis. First, the original mesh and its PM representation serve as the source of local parametrization and connectivity information which determines the coefficients of our adaptive relaxation operator.

Second, if our purpose is to process the geometry of the mesh, it is crucial to treat all three coordinates x, y and z as *dependent* variables. In fact, we consider the coordinates of the mesh to be real functions on the current PM vertex set. Initially, before any changes in geometry take place, these functions can be viewed as identities. When the wanted processing operations, such as filtering or editing, are applied to the data, these functions become more meaningful.

As an example of possible application of our scheme we present various manipulations of the scanned Venus's head model. The original mesh has 50 000 vertices. After building a PM hierarchy, we use our BA pyramid scheme to build a multiresolution representation. We can use different manipulations of the detail coefficients in order to achieve various signal processing tasks. Specifically, if all the detail coefficients finer than some level are put to zero, we achieve a smoothing effect (in figure 13b all the details on the levels above 1000 were set to zero). The stopband filter effect is achieved by setting to zero some range of coefficients (in figure 13c all the details between the levels 1000 and 15 000 were set to zero). One can also enhance certain frequencies (in figure 13d all the details between the levels 1000 and 15 000 were multiplied by two).

#### 5. Conclusion

One of the current frontiers in wavelet research and applications is the generalization of multiresolution methods from the regular to the semiregular and, more recently, irregular setting. We have given a brief review of these developments, starting with

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the one-dimensional setting and moving on to the two-dimensional functional and manifold settings. While there exists an extensive set of tools for the analysis of wavelet constructions in the regular setting, such tools have only recently begun to emerge for the irregular setting. One such tool is the generalization of commutation from the regular to the irregular setting. We have applied these ideas by proposing new irregular subdivision schemes in the manifold setting which are explicitly designed to minimize certain differences. Little is as yet known about the analytic smoothness properties of the resulting constructions, but numerical evidence suggests that they are quite useful for practical applications.

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