A Unified Approach to Evaluation Algorithms for Multivariate Polynomials

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ABSTRACT

We present a *unified* framework for most of the known and a few new evaluation algorithms for multivariate polynomials expressed in a wide variety of bases including the Bézier, multinomial (or Taylor), Lagrange and Newton bases. This unification is achieved by considering evaluation algorithms for multivariate polynomials expressed in terms of L-bases, a class of bases that include the Bézier, multinomial, and a rich subclass of Lagrange and Newton bases. All of the known evaluation algorithms can be generated either by considering recursive evaluation algorithms for L-bases or by examining change of basis algorithms for L-bases.

Keywords: algorithms, Bézier, change of basis, duality, evaluation, Lagrange, multivariate, Newton, polynomials, recurrence, Taylor.

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1. Introduction

The Bézier, multinomial (or Taylor), Lagrange and Newton bases are some of the most popular and useful representations for expressing polynomials of degree n in s variables. Several algorithms for evaluating multivariate polynomials, represented in these bases, have been proposed. The de Casteljau algorithm with computational complexity $O(n^{s+1})$ is well-known for evaluating multivariate Bézier polynomials [dC85, Far86]. Several algorithms for evaluating multivariate polynomials in Bézier or multinomial (Taylor) form with computational complexity $O(n^s)$ have been described [CG90a, dBR92, SV86]. Generalizations with computational complexity $O(n^{s+1})$ of the Aitken-Neville algorithm for evaluating univariate Lagrange polynomials to certain subclasses of multivariate Lagrange polynomials have been proposed [CG90b, Muh78, TM60]. Evaluation algorithms with computational complexity $O(n^s)$ for multivariate polynomials expressed in Newton bases have also been described [Gas90]. In addition, evaluation algorithms related to the generalizations of forward and divided difference algorithms to multivariate polynomials have also been discussed [Vol88, dB78, Vol90, LSP87, Sch81].

Most of these evaluation algorithms seem to have been discovered independently from one another and, therefore, the literature as cited above is scattered and the various algorithms appear to be unrelated. One of the major goals of this work is to demonstrate that these seemingly disparate algorithms are, in fact, closely related. Indeed, all of these algorithms can be derived from the evaluation algorithms for multivariate L-bases, a class of bases that include the Bézier, multinomial, and certain proper subclasses of Lagrange and Newton bases. This unification of these algorithms for evaluating multivariate polynomials. This understanding, in turn, has helped us to design a few new, more efficient algorithms, for evaluating multivariate polynomials.

Our work easily generalizes to arbitrary dimensions. However, for the sake of simplicity, the results are presented and derived here only for bivariate polynomials.

We begin by describing a general parallel up recurrence algorithm with computational complexity $O(n^3)$ for the evaluation of bivariate L-bases. This algorithm specializes to the standard de Casteljau algorithm for evaluation of bivariate Bézier surfaces [dC85, Far86]. The up recurrence also specializes to $O(n^2)$ algorithms for the evaluation of multinomial (Taylor) bases, that include the algorithms proposed earlier by Carnicer-Gasca [CG90a] and de Boor-Ron [dBR92]. In addition, the up recurrence specializes to an algorithm for evaluating Newton polynomials, that has been previously discussed by Carnicer and Gasca [CG90a, Gas90]. The up recurrence also yields a new generalization of the Aitken-Neville algorithm with computational complexity $O(n^3)$ for the evaluation of bivariate Lagrange L-bases. By removing redundant computations, we show that the general parallel up recurrence algorithm can be improved to a new recurrence algorithm with computational complexity $O(n^2lgn)$ for evaluation of arbitrary L-bases. Furthermore, the up recurrence algorithm can be altered into a ladder recurrence algorithm with computational complexity $O(n^2)$ by changing the structure of the parallel up recurrence diagram for the evaluation of bivariate L-bases.

Next we describe another class of algorithms for evaluating L-bases, based on certain change of basis algorithms between L-bases. This class of algorithms include a divided difference algorithm with computational complexity $O(n^2)$ per point, a forward difference algorithm with computational complexity O(1) per point, and a new Lagrange evaluation algorithm with amortized computational complexity O(n) per point. The change of basis algorithm can be specialized into a new dual nested multiplication algorithm with computational complexity $O(n^2)$ for the evaluation of bivariate L-bases. This class of algorithms include an algorithm for evaluating multivariate polynomials in multinomial (Taylor) form described earlier by Schumaker and Volk [SV86].

This paper is organized in the following manner. Section 2 reviews the definition of L-bases and presents examples of Bézier, multinomial, Lagrange and Newton bases as L-bases. Section 3 describes up recurrence algorithms – a parallel up recurrence algorithm with computational complexity $O(n^3)$, a nested multiplication evaluation algorithm with computational complexity $O(n^2lgn)$, and a ladder recurrence algorithm with computational complexity $O(n^2)$. Section 4 presents dual evaluation algorithms – a Lagrange evaluation algorithm with computational complexity O(n) per point, a divided difference algorithm with computational complexity $O(n^2)$ per point, a forward difference algorithm with computational complexity $O(n^2)$ per point, a forward difference algorithm with computational complexity $O(n^2)$. The difference between the nested multiplication and the dual nested multiplication algorithm is highlighted by presenting as an example the evaluation of a bivariate Lagrange L-basis. Section 5 presents a brief description of other evaluation algorithms including factored and hybrid algorithms, that can be obtained by combining some of the above algorithms. Finally, Section 6 concludes with some discussion of future work.

2. L-bases

Here we review the basic definitions and certain well-known examples of affine L-bases. We also provide very brief geometric interpretations for the algebraic entities associated with the L-bases in order to explain how certain bivariate Lagrange and Newton bases arise as special cases of L-bases. Complete details are provided in [LG95b].

Throughout this paper, we shall adopt the following notation. A multi-index α is a 3-tuple of non-negative integers. If $\alpha = (\alpha_1, \alpha_2, \alpha_3)$, then $|\alpha| = \alpha_1 + \alpha_2 + \alpha_3$ and $\alpha! = \alpha_1!\alpha_2!\alpha_3!$. Other multi-indices will be denoted by β and γ . A unit multi-index e_k is a 3-tuple with 1 in the k-th position and 0 everywhere else. Scalar indices will be denoted by i, j, k, l.

A collection \mathcal{L} of 3 sequences $\{L_{1,j}\}, \{L_{2,j}\}, \{L_{3,j}\}, j = 1, \dots, n$ of affine polynomials in two variables is called a *knot-net* of affine polynomials if $(L_{1,\alpha_1+1}, L_{2,\alpha_2+1}, L_{3,\alpha_3+1})$ are affinely independent polynomials for $0 \leq |\alpha| \leq n - 1$. An affine *L*-basis $\{l_{\alpha}^n, |\alpha| = n\}$ is a collection of $\binom{n+2}{2}$ bivariate polynomials defined as follows:

$$l_{\alpha}^{n} = \prod_{i=1}^{\alpha_{1}} L_{1i} \prod_{j=1}^{\alpha_{2}} L_{2j} \prod_{k=1}^{\alpha_{3}} L_{3k}.$$
 (2.1)

It is well-known that $\{l_{\alpha}^{n}, |\alpha| = n\}$ is, in fact, an affine basis for the space of affine polynomials of degree n on R^{2} [CM92].

We assign to each affine polynomial ax + by + c the corresponding line in the affine plane defined by the equation ax + by + c = 0. (The polynomial c corresponds to the line at infinity in the projective plane.) For full details, please refer to [LG95b]. Observe that this correspondence between lines and polynomials depends on the coordinate system and is unique only up to constant multiples. Nevertheless, in the following discussions, we shall identify the polynomial with the line and vice-versa, whenever the coordinate system and constant multiples are irrelevant for the context at hand. The advantage of this correspondence is to allow us to think of algebraic entities such as polynomials in terms of geometric entities such as lines.

2.1 Bézier Bases

We can easily realize Bézier bases as special cases of L-bases by choosing the knot-net of polynomials $L_{i,j} = L_i$, $1 \le j \le n$, $L_1 = a_1x + b_1y + c_1$, $L_2 = a_2x + b_2y + c_2$, $L_3 = a_3x + b_3y + c_3$, where L_1 , L_2 and L_3 are affinely independent polynomials such that no two of the associated lines are parallel. It can easily be verified that the corresponding L-basis is, up to constant multiples, the Bézier basis defined by the three intersection points of L_1 , L_2 and L_3 . In particular, $L_1 = x$, $L_2 = y$ and $L_3 = 1 - x - y$, yields the standard Bézier basis, up to constant multiples, that is, $l_{\alpha}^n = x^{\alpha_1}y^{\alpha_2}(1 - x - y)^{\alpha_3}$.

2.2 Multinomial Bases

The multinomial basis is the standard generalization of the monomial basis to the multivariate setting. For example, the basis 1, x, y, x^2 , xy and y^2 is the bivariate multinomial basis of degree 2. Sometimes the terminology Taylor basis or power basis is also used instead of monomial or multinomial basis. However, we shall refer to this basis as the multinomial basis in accordance with [GB92] and reserve the term power basis for the basis each element of which is an *n*-th power of some linear polynomial [GB92].

The standard multinomial basis is realized as a special case of an L-basis by choosing the knot-net of polynomials $L_{i,j} = L_i, 1 \leq j \leq n, L_1 = x, L_2 = y$ and $L_3 = 1$. This yields: $l_{\alpha}^n = x^{\alpha_1} y^{\alpha_2}$. More general multinomial bases can also be realized as L-bases [LG95b].

2.3 Lagrange Bases

Let { $\{L_{ij}\}, \{L_{2j}\}, \{L_{3j}\}, j = 1, \dots, n$ } be a knot-net of polynomials. Suppose that the polynomials $(L_{1,\alpha_1+1}, L_{2,\alpha_2+1}, L_{3,\alpha_3+1})$ are affinely dependent for $|\alpha| = n, 0 \le \alpha_k \le n-1$. It has been established in [LG95b] that, up to constant multiples, the corresponding L-basis is a Lagrange basis; that is, there exist points \mathbf{v}_{α} such that $l^n_{\alpha}(\mathbf{v}_{\beta}) = l^n_{\alpha}(\mathbf{v}_{\alpha})\delta_{\alpha\beta}$. Therefore, $\{\frac{l^n_{\alpha}}{l^n_{\alpha}(\mathbf{v}_{\alpha})}\}$ forms a Lagrange basis.

To describe the points \mathbf{v}_{α} , let us analyze the dependency conditions. Overloading the notation, let L_{ij} also denote the line in the plane defined by the equation: $L_{ij} = 0$. The linear dependency condition on the polynomials L_{i,α_i+1} means that the lines L_{i,α_i+1} are concurrent for $|\alpha| = n, 0 \leq \alpha_k \leq n - 1$. In general, these lines then meet at a point, in the affine plane if the lines intersect or at infinity in the projective plane if the lines are parallel. Let $\mathbf{v}_{\alpha} = \bigcap_{k=1}^{3} L_{k,\alpha_k+1}$ for $|\alpha| = n, 0 \leq \alpha_k \leq n - 1$. These intersections give rise to $\binom{n+2}{2} - 3$ points corresponding to $\binom{n+2}{2} - 3$ dependency conditions. To these points, we shall add three more points: $\mathbf{v}_{n00} = L_{31} \bigcap L_{21}$, $\mathbf{v}_{0n0} = L_{11} \bigcap L_{31}$, and $\mathbf{v}_{00n} = L_{11} \bigcap L_{21}$. These are precisely the $\binom{n+2}{2}$ points that give rise to Lagrange interpolation conditions.

In our earlier work, we described several interesting lattice or point-line configurations that generate Lagrange L-bases [LG95b]. Here we simply describe one interesting lattice, known as a principal lattice or geometric mesh [CY77], that admits unique interpolation by a Lagrange L-basis. Figure 2.1 is an example of a principal lattice or geometric mesh [CY77] of order n, which can be described by three sets of n lines $\{\{L_{1i}\}, \{L_{2j}\}, \{L_{3k}\}, 1 \leq i, j, k \leq n\}$ such that each set of three lines $\{L_{1,i+1}, L_{2,j+1}, L_{3,k+1}, i + j + k = n\}$ intersect at exactly one common point \mathbf{v}_{ijk} . The lines in Figure 2.1 satisfy both the the linear independence condition for $(L_{1,\alpha_1+1}, L_{2,\alpha_2+1}, L_{3,\alpha_3+1}), 0 \leq |\alpha| \leq n-1$, which is required to define a knot-net of polynomials and the linear dependence condition that is required to define a Lagrange basis. It is clear from the above construction that every geometric mesh of order n gives rise to a Lagrange L-basis.



Figure 2.1: Geometric mesh of order 3 for Lagrange L-basis

2.4 Newton Bases

By choosing the polynomials $L_{1i} = x - a_i$, $L_{2i} = y - b_i$ and $L_{3i} = 1$, we obtain the following affine Newton basis:

$$l_{\alpha}^{n} = \prod_{i=1}^{\alpha_{1}} (x - a_{i}) \prod_{j=1}^{\alpha_{2}} (y - b_{j}).$$

For example, when n = 2 this construction yields the basis functions: $1, (x - a_1), (x - a_1)(x - a_2), (y - b_1), (y - b_1)(y - b_2)$ and $(x - a_1)(y - b_1)$. A slightly more general Newton L-basis is obtained by simply choosing the polynomials $L_{1i} = a_{1i}x + b_{1i}y + c_{1i}, L_{2i} = a_{2i}x + b_{2i}y + c_{2i}$, and $L_{3i} = a_3x + b_3y + c_3$. In our earlier work [LG95b], we established that each Newton L-basis can be associated with a point and derivative interpolation problem with the following properties: (i) there exists a unique solution to the general interpolation problem expressed in this basis and (ii) the coefficients a_{α} of the interpolant $L(\mathbf{u}) = \sum_{|\alpha|=n} a_{\alpha} l_{\alpha}^n$ expressed in the Newton L-basis are the solutions of a lower triangular system of linear equations.

In [LG95b], we also exhibited a rich collection of lattices or point-line configurations that admit unique and natural solutions to an appropriate interpolation problem by means of a Newton L-basis. These lattices include the principal lattices or geometric meshes (and the associated Lagrange interpolation problems) as well as *natural lattices* of order n, which are defined by n+2distinct lines. The corresponding Newton L-bases solve the Lagrange interpolation problem at the associated $\binom{n+2}{2}$ distinct points of intersection. Figure 2.2 shows a natural lattice of order 3.



Figure 2.2: Natural lattice of order 3 for Newton L-basis

[LG95b] also presents some examples of lattices usually associated with Hermite interpolation problems [M $\acute{8}4$, NR92] that can be solved uniquely with Newton L-bases.

3. Up Recurrence Algorithms

This section describes a class of evaluation algorithms for L-bases that arise from variations on a general parallel up recurrence algorithm, which is discussed next.

3.1 Parallel Up Recurrence Algorithm

Let $L(\mathbf{u})$ be a polynomial expressed in terms of an L-basis $\{l_{\alpha}^{n}\}$ defined by the knot-net $\mathcal{L} = \{\{L_{1j}\}, \{L_{2j}\}, \{L_{3j}\}, j = 1, \dots, n\}$. Suppose we wish to evaluate $L(\mathbf{u}) = \sum_{|\alpha|=n} S_{\alpha} l_{\alpha}^{n}(\mathbf{u})$ at an arbitrary but fixed \mathbf{u} . Then we can use the up recurrence illustrated in Figure 3.1 to evaluate $L(\mathbf{u})$, where the coefficients C_{α}^{0} are normalized as $C_{\alpha}^{0} = \frac{\alpha!}{n!} S_{\alpha}$. The diagram is to be interpreted as follows: the computation starts at the base of the tetrahedron. The value at any node is computed by multiplying the value along each arrow which enters the node by the value of the node from which the arrow emerges and adding the results. Observe that the central node of the base of the tetrahedron is occluded by the apex node in Figure 3.1 and is therefore not shown. The value of $L(\mathbf{u})$ is then given by C_{000}^{n} at the apex of the tetrahedron. More formally the recurrence is described as follows:

$$C_{\alpha}^{0} = \frac{\alpha!}{n!} S_{\alpha},$$

$$C_{\alpha}^{i} = \sum_{k=1}^{3} C_{\alpha+e_{k}}^{i-1} * L_{k,\alpha_{k}+1} \text{ for } i = 1, \cdots, n. \text{ for } |\alpha| = n - i.$$

This algorithm is referred to as the *parallel* up recurrence and generalizes the parallel up recurrence algorithm for the evaluation of univariate polynomials expressed in terms of Pölya



Figure 3.1: Parallel up recurrence algorithm for L-bases



Figure 3.2: Nested multiplication evaluation algorithm for L-bases

basis functions [GB92]. The computational complexity of this algorithm is $O(n^3)$, because there are $\frac{n(n+1)(n+2)}{6}$ nodes in the tetrahedron. A careful examination of the algorithm reveals that the label along the edge from the node $\alpha + e_i$ to the node α is L_{i,α_i+1} . Due to this property of the recurrence diagram, the labels along the "parallel" edges of the tetrahedron are the same; this feature justifies the terminology *parallel*. Now observe that there are exactly $\frac{n!}{\alpha!}$ paths from the node α at the base of the tetrahedron to the node at the apex of the tetrahedron. Moreover by the parallel property the products of the labels along each path from a node α at the base to the node at the apex are identical. These products are all equal to l_{α}^n because to pass from the factors $L_{i,k}$ i = 1, 2, 3 $0 \le k \le \alpha_i$ must be encountered exactly once along each path. These observations establish the correctness of the algorithm, and are also the key to improving this algorithm as we shall soon see in the following section.

3.2 Nested Multiplication Algorithm

The nested multiplication evaluation algorithm for L-bases arises by observing that if we do not scale the coefficients in the parallel up recurrence algorithm described in Section 3.1, the same computation can be accomplished by choosing exactly *one* path from each node at the base of the tetrahedron to the node at the apex. This structure can easily be achieved by making sure that at every level of the computation there is exactly *one* arrow pointing upwards from each node. It is remarkable that it does *not* matter which set of arrows are used at each level so long as they satisfy this uniqueness condition. Therefore, this observation gives rise to a whole class of algorithms, all of which have many fewer arrows than the general up recurrence algorithm described in the previous section. A simple symmetric rule for selecting the arrows is to choose all the arrows in the topmost upright triangles, the two lower arrows on the rightmost

triangles and the leftmost arrow in all the remaining triangles as shown in Figure 3.2. However, although the number of arrows is reduced with this choice, the computational complexity of the algorithm still remains $O(n^3)$ because we have removed slightly less than $\frac{2}{3}$ of the arrows.

Carnicer and Gasca [CG90a] describe a class of evaluation algorithms for those multivariate polynomials that are expressed as the sum of a constant plus some other polynomials, each of which can be written as a product of a linear polynomial with a polynomial of degree strictly lower than the degree of the original polynomial. Polynomials represented in terms of L-bases clearly satisfy this property. Therefore, the evaluation algorithms of Carnicer and Gasca can be applied to L-bases. However their algorithm and the resulting computational complexity depend upon the way the polynomial is represented. In general, the computational complexity of their algorithm for bivariate polynomial bases including bivariate Lagrange bases [CG90b] is $O(n^3)$. We shall refer to this class of algorithms, including up recurrence algorithms and algorithms described by Carnicer and Gasca, as nested multiplication algorithms because these algorithms can clearly be viewed as nested multiplication. Both Carnicer and Gasca [CG90a] and de Boor and Ron [dBR92] describe this class of nested multiplication algorithms as the generalization of Horner's nested multiplication algorithm for the evaluation of univariate polynomials. However, as we shall see later in Section 4, there is another *different* class of nested multiplication algorithms for the evaluation of multivariate polynomials that also qualify as generalizations of Horner's algorithm. Therefore, it is not clear which of these algorithms can claim to be the generalization of Horner's evaluation algorithm, although all of them are a form of nested multiplication.

We can reduce the computational complexity of the our nested multiplication algorithm from $O(n^3)$ to $O(n^2lgn)$ by observing that it is *not* necessary to have an arrow emerging from a node if there are no arrows entering that node. Such a node will be referred to as a *dead* node. Other nodes will be referred to as *active* nodes. To reduce the computational complexity of the algorithm, we desire to increase the number of dead nodes as much as possible. One way to achieve this end is to point as many arrows as possible towards the corners and towards the sides rather than towards the center.

We now briefly illustrate how this idea can be used to reduce the computational complexity of the analogous evaluation algorithm for univariate polynomials from $O(n^2)$ to O(nlgn). This problem is purely combinatorial; one simply needs to choose enough arrows so that there is exactly one path from every node at the base to the node at the apex. The left diagram of Figure 3.3 shows how this can be achieved for univariate polynomials when n = 3. This approach can be generalized to polynomials of arbitrary degree n by a divide and conquer strategy. Here a problem of size n is broken down into two subproblems of size $\lfloor \frac{n}{2} \rfloor$ at the base followed by at most $\lceil \frac{n}{2} \rceil$ arrows on both sides of the triangle. An example for n = 7 is shown in Figure 3.4, where evaluation of a univariate polynomial of degree 7 is broken down into two subproblems of evaluation for univariate polynomials of degree 3 at the base. This approach yields the following recurrence equation for the computational complexity T(n):

$$T(n) = 2T(\lfloor \frac{n}{2} \rfloor) + O(n).$$



Figure 3.3: Nested multiplication evaluation algorithm for L-bases



Figure 3.4: Nested multiplication evaluation algorithm for univariate L-bases of degree 7

Solving this recurrence, we obtain T(n) = nlgn, which establishes the claim for univariate polynomials.

To generalize this technique to the bivariate case, we run the univariate algorithm layer by layer as illustrated in the right diagram of Figure 3.3. For the nodes in the frontmost layer of the base of the tetrahedron, we run an evaluation algorithm for a univariate L-basis of degree n. For the layer behind it at the base of the tetrahedron, we run the evaluation algorithm for a



Figure 3.5: Efficient evaluation algorithm for multinomial bases

univariate L-basis of degree n - 1 and so on. Thus we run evaluation algorithms for univariate L-bases of degrees all the way from n down to 1. This approach yields values at all the nodes along one of the edges of the tetrahedron, namely the back edge. We now simply put arrows along this back edge to compute the value at the apex of the tetrahedron. These new arrows require an additional O(n) computations. This method yields the following recurrence equation for the computational complexity T(n):

$$T(n) = \sum_{k=1}^{n} O(klgk) + O(n).$$

Therefore, the total computational complexity of this algorithm is $O(n^2lgn)$. Although this algorithm seems to give the best possible computational complexity that one can obtain by removing arrows from a parallel up recurrence algorithm, it does not give the best possible constant because one can easily figure out other ways of removing arrows which actually yield fewer arrows than the algorithm described above. However we found that this description was the easiest way to provide a simple proof that by removing arrows one can obtain an evaluation algorithm with computational complexity $O(n^2lgn)$.

3.3 Examples

This section describes how the parallel up recurrence algorithm and the nested multiplication algorithm for evaluation of bivariate L-bases can be specialized to obtain evaluation algorithms for bivariate Bézier bases, multinomial bases, Lagrange L-bases and Newton L-bases.

3.3.1 Multinomial Evaluation



Figure 3.6: de Boor-Ron's evaluation algorithm for multinomial bases

Since one of the knot-nets in the multinomial basis is always 1, considerable simplifications take place in both the parallel up recurrence algorithm and the nested multiplication algorithm for evaluation with respect to the multinomial bases. In this case it pays to choose arrows with the label 1 as often as possible. The left diagram of Figure 3.5 shows such a choice, where the arrows pointing downward have the label 1. Since no multiplication needs to be done for arrows with the label 1, one can "pull up" these nodes as shown in the right diagram of Figure 3.5. The value at any node is now computed by adding the value at that node to the value computed as before by multiplying the label along each arrow which enters the node by the value of the node from which the arrow emerges and adding the results. The right diagram of Figure 3.5 shows the same computation as in the left diagram, but now arranged in a triangular format. Since the triangular format has only n^2 nodes, this figure shows that the computational complexity of this evaluation algorithm is $O(n^2)$. Although Carnicer and Gasca [CG90a] do not explicitly discuss the evaluation of multivariate polynomials expressed in the multinomial bases, the right diagram of Figure 3.5 is very similar to the diagram in [CG90b], which appears in an analogous but slightly different context. Therefore, this evaluation algorithm can be construed to be equivalent to the one proposed by Carnicer and Gasca [CG90a, CG90b].

Interestingly, de Boor and Ron [dBR92] describe an evaluation algorithm for multinomial bases that is closely related to these algorithms, although not quite the same. In fact, their algorithm is a hybrid of the parallel up recurrence and the efficient parallel up recurrence algorithm, where some duplicate paths have not been removed. They take advantage of the arrows with label 1 and "pull up" the nodes, but they do not remove redundant paths. Therefore, their starting coefficients are slightly different from both C_{α} and S_{α} . In fact, their starting coefficients are $R_{\alpha} = \frac{\alpha_1!\alpha_2!}{(\alpha_1+\alpha_2)!}S_{\alpha}$. The diagram of Figure 3.6 shows the evaluation recurrence of de Boor and Ron for the evaluation of multinomial bases. This algorithm also has computational



Figure 3.7: Evaluation algorithm for L-bases with coefficients R_{α}



Figure 3.8: Another diagram for the evaluation algorithm for L-bases with coefficients S_{α}

complexity of $O(n^2)$, although with a larger constant.

In fact, following this strategy of using the coefficients R_{α} instead of the coefficients C_{α} or S_{α} , one can write down yet another form of nested multiplication algorithm for the evaluation of multivariate polynomials expressed in terms of L-bases. This algorithm is illustrated in Figure 3.7. Observe that the tetrahedral structure of the parallel up recurrence diagram in Figure 3.1 can be shown conveniently as in Figure 3.7, where the different layers (a layer being defined as



Figure 3.9: Tensor product evaluation algorithm for Bézier polynomials

nodes α with same α_3 value) of the tetrahedral structure are shown side-by-side. Since there is exactly one edge between different layers of this diagram, there are no "cross-edges" as there are in Figure 3.1. Therefore, the side-by-side diagram appears uncluttered. The algorithm in Figure 3.7 utilizes R_{α} as coefficients because some of the duplicate paths have not been removed. This algorithm has computational complexity $\sum_{k=1}^{n} O(k^2) = O(n^3)$. By removing all the duplicate paths and using the coefficients S_{α} , one can redraw the right diagram of Figure 3.3 as Figure 3.8. This algorithm has computational complexity $\sum_{k=1}^{n} O(klgk) = O(n^2lgn)$.

3.3.2 Bézier Evaluation

The parallel up recurrence algorithm specializes to the de Casteljau evaluation algorithm for Bézier surfaces with computational complexity $O(n^3)$ [dC85, Far86]. In this case, observe that $C^0_{\alpha} = \frac{\alpha!}{n!}S_{\alpha}$ are indeed the coefficients of the multivariate polynomials in Bézier form.

There is another interesting way of evaluating polynomials of total degree n expressed in terms of Bézier bases by viewing the polynomial as a tensor product of bi-degree $n \times n$ [War92]. This tensor product evaluation algorithm can be derived by modifying the evaluation algorithm



Figure 3.10: Evaluation algorithm for Newton L-bases

discussed in the previous section and described in Figure 3.7. First, replace the labels L_{1i} by x, L_{2i} by y and L_{3i} by 1 - x - y in Figure 3.7 to reflect the fact that we are working with the standard Bézier basis. Now reorganize the diagonal edge in Figure 3.7 with labels 1-x-y into the univariate de Casteljau evaluation algorithm as shown in Figure 3.9 and compensate for the extra factors of x + y by dividing each label of the lower triangles by x + y. Introducing the parameters t = 1 - x - y and $s = \frac{y}{x+y}$, we obtain the tensor product evaluation algorithm presented in Figure 3.9. Observe that the coefficients C_{α} in Figure 3.9 are the usual coefficients for a polynomial expressed in terms of a Bézier basis. This tensor product algorithm has $\frac{n(n+1)(n+5)}{3}$ arrows for the evaluation of polynomials of degree n. Therefore this algorithm is still $O(n^3)$. However observe that the standard bivariate de Casteljau algorithm for the evaluation of polynomials of total degree n has $\frac{n(n+1)(n+2)}{2}$ arrows, as shown in Figure 3.1. Therefore the tensor product algorithm for the evaluation of polynomials of polynomials of total degree greater than or equal to 4.

3.3.3 Newton Evaluation

Since all the polynomials in one of the sequences in the knot-net of a Newton basis are 1, considerable simplifications take place as well in both the parallel up recurrence algorithm and the nested multiplication algorithm for the evaluation of Newton bases. In this case too it pays to choose arrows with labels 1 as often as possible. The left diagram of Figure 3.10 shows such a choice, where the arrows pointing downwards have the label 1. Since no multiplication needs to be done for arrows with labels 1, one can "pull up" these nodes as shown in the right diagram of Figure 3.10. The right diagram of Figure 3.10 shows the same computation as in the left diagram, but now arranged in a triangular format. Since the triangular format has only $O(n^2)$ nodes, this figure shows that the computational complexity of this evaluation algorithm is $O(n^2)$.

This algorithm is exactly the same as the evaluation algorithm described by Gasca in [Gas90].

3.3.4 Lagrange Evaluation: Generalization of the Aitken-Neville Algorithm

We now derive a variation of the parallel up recurrence diagram for the evaluation of Lagrange L-bases, which gives rise to a bivariate generalization of the univariate Aitken-Neville algorithm. In the past the Aitken-Neville algorithm has been generalized to some restricted classes of bivariate Lagrange polynomials [Ait32, Nev34, Gas90, CG90b, CG90a, Muh78, GL87, GM89, Muh74, TM60, Muh88]. This new generalization extends the class of multivariate Lagrange polynomials to which an Aitken-Neville algorithm can be applied.

Given a knot-net $\mathcal{L} = \{\{L_{1j}\}, \{L_{2j}\}, \{L_{3j}\}, j = 1, \dots, n\}$ of linear homogeneous polynomials, consider the three knot-nets $\mathcal{M}_1 = \{(\hat{L}_{11}, \dots, L_{1n}), (L_{21}, \dots, \hat{L}_{2n}), (L_{31}, \dots, \hat{L}_{3n})\}, \mathcal{M}_2 = \{(L_{11}, \dots, \hat{L}_{1n}), (\hat{L}_{21}, \dots, L_{2n}), (L_{31}, \dots, \hat{L}_{3n})\}, \text{ and } \mathcal{M}_3 = \{(L_{11}, \dots, \hat{L}_{1n}), (L_{21}, \dots, \hat{L}_{2n}), (\hat{L}_{31}, \dots, L_{3n})\}$ respectively, where \hat{L} means that the term L is missing. Observe that if \mathcal{L} satisfies the linear independence condition on the knot-net of polynomials for $0 \leq |\alpha| \leq n - 1$, then the knot-net \mathcal{M}_1 satisfies the linear independence condition for $0 \leq |\beta| \leq n - 2$. This observation follows by setting $\alpha_1 = \beta_1 + 1$, $\alpha_2 = \beta_2$ and $\alpha_3 = \beta_3$. Similarly the knot-nets \mathcal{M}_2 and \mathcal{M}_3 satisfy the linear independence condition for $0 \leq |\beta| \leq n - 2$. Moreover, if the knot-net \mathcal{L} satisfies the linear dependence condition for a Lagrange L-basis. Let $\{l_{\alpha}^n\}, \{p_{1,\alpha}^{n-1}\}, \{p_{2,\alpha}^{n-1}\}$ and $\{p_{3,\alpha}^{n-1}\}$ be the L-bases corresponding to the knot-nets $\mathcal{L}, \mathcal{M}_1, \mathcal{M}_2$ and \mathcal{M}_3 respectively.

We can associate point interpolation problems to these knot-nets as follows. Let $L(\mathbf{u})$ be the unique polynomial of degree n that interpolates the values S_{α} at the points \mathbf{v}_{α} , where the points \mathbf{v}_{α} are the points of intersection of certain lines from the collection \mathcal{L} as described in Section 2.3. Then, $L(\mathbf{u}) = \sum_{|\alpha|=n} S_{\alpha} \frac{l_{\alpha}^{n}(\mathbf{v}_{\alpha})}{l_{\alpha}^{n}(\mathbf{v}_{\alpha})}$. Similarly let $M_{1}(\mathbf{u}), M_{2}(\mathbf{u})$ and $M_{3}(\mathbf{u})$ be the unique polynomials of degree n-1 that satisfy the point interpolation conditions $M_{i} = S_{\alpha+e_{i}}\delta_{\alpha\beta}$ for $|\alpha| = n-1$. Then $M_{i}(\mathbf{u}) = \sum_{|\alpha|=n-1} S_{\alpha+e_{i}} \frac{p_{i,\alpha}^{n-1}}{p_{i,\alpha}^{n-1}(\mathbf{v}_{\alpha})}$.

Now create the following variation of the parallel up recurrence algorithm: Replace the labels L_{k,α_k} by the labels $\frac{L_{k,\alpha_k}}{L_{k,\alpha_k}(\mathbf{v}_{\alpha+le_k})}$ on the arrows pointing from the node C_{α}^l to the node $C_{\alpha-e_k}^{l+1}$. Then we can express the solution to the point interpolation problem as an affine combination of the three subproblems of point interpolation and thus generalize the Aitken-Neville algorithm for the evaluation of Lagrange polynomials from univariate to bivariate polynomials [Ait32, Nev34]. **Theorem 1:**

$$L(\mathbf{u}) = \frac{L_{11}}{L_{11}(\mathbf{v}_{n00})} M_1(\mathbf{u}) + \frac{L_{21}}{L_{21}(\mathbf{v}_{0n0})} M_2(\mathbf{u}) + \frac{L_{31}}{L_{31}(\mathbf{v}_{00n})} M_3(\mathbf{u}).$$

Proof: The proof is by induction. The case n = 1 reduces to a simple triangular point interpolation problem and in this case the linear solution $L(\mathbf{u})$ is indeed a barycentric combination of constant solutions given by

$$L(\mathbf{u}) = \frac{L_{11}}{L_{11}(\mathbf{v}_{100})} C_{100} + \frac{L_{21}}{L_{21}(\mathbf{v}_{010})} C_{010} + \frac{L_{31}}{L_{31}(\mathbf{v}_{001})} C_{001}.$$



Figure 3.11: Ladder recurrence algorithm for bivariate L-bases

The inductive hypothesis assumes that the statement of the theorem is true for n - 1. We will now prove that the statement of the theorem holds for n. First, observe that for i = 1, 2, 3

$$M_i(\mathbf{u}) = \sum_{|\alpha|=n-1} S_{\alpha+e_i} \frac{p_{i,\alpha}^{n-1}}{p_{i,\alpha}^{n-1}(\mathbf{v}_{\alpha})}.$$
(3.1)

The inductive hypothesis asserts that the values at the three nodes that contribute to the apex of the tetrahedron, which are simply the apexes of the recurrence diagrams of the three subproblems, are $M_1(\mathbf{u})$, $M_2(\mathbf{u})$ and $M_3(\mathbf{u})$. Now let

$$M(\mathbf{u}) = \frac{L_{11}}{L_{11}(\mathbf{v}_{n00})} M_1(\mathbf{u}) + \frac{L_{21}}{L_{21}(\mathbf{v}_{0n0})} M_2(\mathbf{u}) + \frac{L_{31}}{L_{31}(\mathbf{v}_{00n})} M_3(\mathbf{u}).$$
(3.2)

We will prove that $M(\mathbf{u}) = L(\mathbf{u})$. Substituting Equation 3.1 into Equation 3.2 and recalling also that

$$L_{i1}p_{i,\alpha}^{n-1} = l_{\alpha+e_i}^n,$$

we obtain

$$M(\mathbf{u}) = \sum_{|\alpha|=n} S_{\alpha} \frac{l_{\alpha}^{n}}{l_{\alpha}^{n}(\mathbf{v}_{\alpha})} I(\mathbf{v}_{\alpha}),$$

where

$$I(\mathbf{v}) = \frac{L_{11}(\mathbf{v})}{L_{11}(\mathbf{v}_{n00})} + \frac{L_{21}(\mathbf{v})}{L_{21}(\mathbf{v}_{0n0})} + \frac{L_{31}(\mathbf{v})}{L_{31}(\mathbf{v}_{00n})}.$$

To show that I = 1, we observe that I is a linear polynomial, which evaluates to 1 at three affinely independent points \mathbf{v}_{n00} , \mathbf{v}_{0n0} and \mathbf{v}_{00n} ; therefore I is identically equal to 1. This proves that $M(\mathbf{u}) = L(\mathbf{u})$ and establishes the theorem. \Box

3.4 Ladder Recurrence Algorithm

By removing redundant arrows, we were able to design in the previous section an $O(n^2 lgn)$ algorithm for the evaluation of bivariate L-bases. Can we do better? The answer is yes, although we must modify still further the structure of the up recurrence algorithm. In the previous

section, we derived an $O(n^2lgn)$ algorithm by extending a univariate evaluation algorithm with computational complexity O(nlgn). By modifying the structure of the univariate parallel up recurrence, Warren developed a ladder recurrence algorithm for the evaluation of univariate L-bases (or Pölya bases) with computational complexity O(n) [War94]. Since each triangle in Figure 3.7 has the same structure as the univariate parallel up recurrence, we can replace each triangle of height p by a ladder of height p.

Figure 3.11 presents an example of the bivariate ladder recurrence algorithm for degree 3. The recurrence starts at the bottom of Figure 3.11, at all the nodes shown as cross-hatched circles. These nodes have values 1. As before, the value at any node is computed by multiplying the value along each arrow that enters the node by the value of the node from which the arrow emerges and adding the results. The computation proceeds upwards and the value of $L(\mathbf{u})$ emerges at the apex node of the triangle, shown as a black circle in Figure 3.11. A more formal description of this algorithm for arbitrary degree n can be found in [LGW94]. Since a ladder with p steps requires O(p) computations, the computational complexity of this algorithm is $\sum_{p=1}^{n} O(p) = O(n^2)$. Observe that in contrast to other algorithms described in this work, this algorithm employs coefficients as labels along the edges.

4. Dual Evaluation Algorithms

Thus far we have described a generic $O(n^3)$, a generic $O(n^2lgn)$, and a generic $O(n^2)$ algorithm for the evaluation of bivariate L-bases. In this section we describe dual evaluation algorithms with computational complexity $O(n^2)$, O(n), and O(1) per point. This class of algorithms is based upon certain change of basis algorithms between L-bases which we discuss next.

4.1 Change of Basis Algorithms

In our earlier work [LG95a], we derived change of basis algorithms with computational complexity $O(n^3)$ between any two bivariate L-bases. Here we establish how these change of basis algorithms can be used to derive evaluation algorithms with computational complexity O(n) or O(1) per point for bivariate L-bases. This class of algorithms include a Lagrange evaluation algorithm, a divided difference algorithm, and a forward difference algorithm, each of which we shall now describe.

4.1.1 Lagrange Evaluation Algorithm

The general change of basis algorithm between any two L-bases was derived using the duality between B-bases and L-bases and is dual to the general change of basis algorithm between any two B-bases [Sei91], derived using the blossoming principle [Ram87, Ram89]. We begin with a specific example of a change of basis algorithm from a Bézier L-basis to a Lagrange L-basis to: (i) illustrate the general procedure for change of basis between any two L-bases that will be needed subsequently to derive the divided difference and the forward difference algorithm, and (ii) to describe how this algorithm can be used to evaluate any L-basis at $O(n^2)$ points with computational complexity $O(n^3)$, that is, an amortized cost of O(n) computations per point.

Suppose we are given the coefficients R_{α} of a quadratic polynomial L with respect to the Bézier L-basis $\{p_{\alpha}^n\}$ defined by a knot-net $\mathcal{L} = \{\{L_{1j}\}, \{L_{2j}\}, \{L_{3j}\}, j = 1, 2\}$, where

The corresponding Bézier L-basis is then given by $p_{200}^2 = x^2$, $p_{020}^2 = y^2$, $p_{002}^2 = (1 - x - y)^2$, $p_{110}^2 = xy$, $p_{101}^2 = x(1 - x - y)$, and $p_{011}^2 = y(1 - x - y)$.

We would like to compute the coefficients U_{α} of this quadratic polynomial L with respect to the Lagrange L-basis $\{l_{\alpha}^n\}$ defined by the knot-net $\mathcal{M} = \{\{M_{1j}\}, \{M_{2j}\}, \{M_{3j}\}, j = 1, 2\}$, where

$$M_{11} = x ; M_{12} = x - \frac{1}{2} ; M_{21} = y ; M_{22} = y - \frac{1}{2} ; M_{31} = 1 - x - y ; M_{32} = \frac{1}{2} - x - y .$$



Figure 4.1: Labeling of the tetrahedron



Figure 4.2: Change of basis from Bézier L-basis to Lagrange L-basis

The corresponding Lagrange L-basis is then given by $l_{200}^2 = x(x - \frac{1}{2}), \ l_{020}^2 = y(y - \frac{1}{2}), \ l_{002}^2 = (1 - x - y)(\frac{1}{2} - x - y), \ l_{110}^2 = xy, \ l_{101}^2 = x(1 - x - y), \ \text{and} \ l_{011}^2 = y(1 - x - y).$

To describe the change of basis algorithm, we construct three tetrahedra. We first explain the labeling scheme for these tetrahedra. For each tetrahedron, $\frac{(3-i)(4-i)}{2}$ nodes are placed at the *i*-th level of the tetrahedron for i = 0, 1, 2 and the nodes along one of the lateral faces are indexed by α for $|\alpha| = 2$. An arrow is placed pointing downward from a node α at *i*-th level to the three nodes $\alpha + e_1 - e_3$, $\alpha + e_2 - e_3$ and $\alpha - e_3$ at (i - 1)-st level directly below it. This labeling scheme for the nodes is shown in Figure 4.1. Values, referred to as *labels*, are placed along the arrows. The labels are indexed as $g_{k,\alpha}$ for k = 1, 2, 3 and $|\alpha| = 0, 1, 2$ for an arrow from a node $(\alpha_1, \alpha_2, 2 - |\alpha|)$ at the $|\alpha|$ -th level to the three nodes below it. This labeling scheme for the labels and the arrows is also shown in Figure 4.1. For the first tetrahedron the known coefficients R_{α} with $|\alpha| = 2$ are placed at the nodes along one of the lateral faces of the tetrahedron as depicted in the first diagram of Figure 4.2. The labels $g_{k,\alpha}$ are computed as follows: For $|\alpha| = 0, 1$, let $i = 2 - |\alpha|$; then

$$L_{3i} = g_{1,\alpha}L_{1,\alpha_1+1} + g_{2,\alpha}L_{2,\alpha_2+1} + g_{3,\alpha}M_{3,\alpha_3+1}.$$

Thus finding $g_{k,\alpha}$ amounts to solving a 3×3 system of linear equations. For our example, the labels are: $g_{3,001} = 2$, and $g_{1,001} = g_{2,001} = g_{3,010} = g_{3,100} = g_{3,000} = 1$. The rest of the labels are zero. These labels are shown in the first diagram of Figure 4.2. The computation is now carried out as follows. At the start all the nodes at all levels of the pyramid are empty or zero other than the nodes α with $|\alpha| = 2$, where the coefficients R_{α} are placed. The empty or zero nodes are shown as hatched circles in Figures 4.1 and 4.2. The computation starts at the apex of the tetrahedron and proceeds downwards. A value at an empty node is computed by multiplying the label along each arrow that enters the node by the value of the node from which the arrow emerges and adding the results. A value already at that node. After the computation is complete, the new coefficients $S_{\alpha+(2-|\alpha|)e_3}$ emerge at the nodes α on the base triangle. These coefficients are as follows: $S_{200} = R_{200}$, $S_{110} = R_{110}$, $S_{020} = R_{020}$, $S_{101} = R_{002} + R_{101}$, $S_{011} = R_{002} + R_{011}$, $S_{002} = 2R_{002}$. These new coefficients now express the polynomial L with respect to the L-basis defined by the knot-net $\{\{L_{1j}\}, \{L_{2j}\}, \{M_{3j}\}, j = 1, 2\}$.

We now repeat the above procedure with a second tetrahedron, where the coefficients S_{α} are placed at the nodes α with $|\alpha| = 2$ as shown in the middle diagram of Figure 4.2. The labels on the tetrahedron are permuted from (i, j, k) to (i, k, j) because now we wish to retain the polynomial M_{3j} and replace the polynomials L_{2j} by M_{2j} . The labels $g_{k,\alpha}$ are now computed as follows: For $|\alpha| = 0, 1$, let $i = 2 - |\alpha|$; then

$$L_{2i} = g_{1,\alpha}L_{1,\alpha_1+1} + g_{2,\alpha}M_{2,\alpha_2+1} + g_{3,\alpha}M_{3,\alpha_3+1}.$$

These labels are also shown in the middle diagram of Figure 4.2 and in our special case turn out to be the same as in the first tetrahedron. After the computation is complete, the new coefficients T_{α} emerge at the nodes on the base triangle. These coefficients are as follows: $T_{200} = S_{200}$, $T_{110} = S_{020} + S_{110}$, $T_{020} = 2S_{020}$, $T_{101} = S_{101}$, $T_{011} = S_{020} + S_{011}$, $T_{002} = S_{002}$. These coefficients now express the polynomial L with respect to the L-basis defined by the knot-net $\{\{L_{1j}\}, \{M_{2j}\}, \{M_{3j}\}, j = 1, 2\}$.

Finally we repeat the above procedure with a third tetrahedron, where the coefficients T_{α} are placed at the nodes α with $|\alpha| = 2$ as shown in the rightmost diagram of Figure 4.2. The labels on this tetrahedron are permuted from (i, j, k) to (j, k, i) because now we wish to retain the polynomials M_{2j} and M_{3j} and replace the polynomials L_{1j} by M_{1j} . The labels $g_{k,\alpha}$ are computed as follows: For $|\alpha| = 0, 1$, let $i = 2 - \alpha$; then

$$L_{1i} = g_{1,\alpha} M_{1,\alpha_1+1} + g_{2,\alpha} M_{2,\alpha_2+1} + g_{3,\alpha} M_{3,\alpha_3+1}$$

Again in our special case these labels are the same as in the first tetrahedron and are shown in the rightmost diagram of Figure 4.2. After the computation is complete, the new coefficients U_{α} emerge at the nodes on the base triangle. These new coefficients are as follows: $U_{200} = 2T_{200}$, $U_{110} = T_{200} + T_{110}$, $U_{020} = T_{020}$, $U_{101} = T_{200} + T_{101}$, $U_{011} = T_{011}$, $U_{002} = T_{002}$. These coefficients express the polynomial *L* with respect to the L-basis defined by the knot-net $\mathcal{M} = \{\{M_{1j}\}, \{M_{2j}\}, \{M_{3j}\}, j = 1, 2\}$. The change of basis algorithm is now complete. In terms of the original coefficients R_{α} , the final coefficients U_{α} , are: $U_{200} = 2R_{200}$, $U_{110} = R_{200} + R_{020} + R_{110}$, $U_{020} = 2R_{020}$, $U_{101} = R_{200} + R_{002} + R_{101}$, $U_{011} = R_{020} + R_{002} + R_{011}$, $U_{002} = 2R_{002}$.

The above change of basis algorithm is the crucial step in the Lagrange evaluation algorithm that we now describe. Given the coefficients R'_{α} of the quadratic polynomial L in the quadratic Bézier basis x^2 , y^2 , $(1 - x - y)^2$, 2xy, 2x(1 - x - y), and 2y(1 - x - y), we first compute the coefficients R_{α} with respect to the Bézier L-basis by $R_{\alpha} = \frac{n!}{\alpha!}R'_{\alpha}$, where n = 2 in this case. Then we run the change of basis algorithm to compute the coefficients U_{α} of the quadratic polynomial L with respect to the Lagrange L-basis l^n_{α} . Since the Lagrange basis is given by $\{\frac{l^n_{\alpha}}{l^n_{\alpha}(\mathbf{v}_{\alpha})}\}$, the coefficients U'_{α} with respect to the Lagrange basis are computed by $U'_{\alpha} = U_{\alpha}l^n_{\alpha}(\mathbf{v}_{\alpha})$. In the current case, since $\mathbf{v}_{200} = (1,0)$, $\mathbf{v}_{020} = (0,1)$, $\mathbf{v}_{002} = (0,0)$, $\mathbf{v}_{110} = (\frac{1}{2},\frac{1}{2})$, $\mathbf{v}_{101} = (\frac{1}{2},0)$, $\mathbf{v}_{011} = (0,\frac{1}{2})$, we find $l^2_{200}(\mathbf{v}_{200}) = \frac{1}{2}$, $l^2_{020}(\mathbf{v}_{020}) = \frac{1}{2}$, $l^2_{110}(\mathbf{v}_{110}) = \frac{1}{4}$, $l^2_{101}(\mathbf{v}_{101}) = \frac{1}{4}$. The coefficients U'_{α} are the values of the quadratic polynomial L at the points \mathbf{v}_{α} ; that is, $L(\mathbf{v}_{\alpha}) = U'_{\alpha}$.

The general Lagrange evaluation algorithm is obtained similarly by first converting the coefficients in a given basis to an L-basis by multiplying if necessary by appropriate constants, running the change of basis algorithm from the given L-basis to the Lagrange L-basis, and finally multiplying the derived coefficients by appropriate constants to obtain the value of the polynomial at the points defined by the Lagrange L-basis.

A general change of basis algorithm from any L-basis to any other L-basis is obtained by following essentially the same procedure. Suppose we are given the coefficients R_{α} of a polynomial L of degree n with respect to an L-basis $\{l_{\alpha}^n\}$ defined by the knot-net $\mathcal{L} = \{\{L_{1j}\}, \{L_{2j}\}, \{L_{3j}\}, j = 1, \dots, n\}$. We would like to compute the coefficients U_{α} of this polynomial L with respect to another L-basis $\{p_{\alpha}^n\}$ defined by another knot-net $\mathcal{M} =$ $\{\{M_{1j}\}, \{M_{2j}\}, \{M_{3j}\}, j = 1, \dots, n\}$.

The general change of basis algorithm is constructed in the following manner:

1. Build three tetrahedra. For each tetrahedron, $\frac{(n+1-i)(n+2-i)}{2}$ nodes are placed at the *i*-th level of the tetrahedron for $i = 0, \dots, n$. The labels $g_{k,\alpha}$ along the edges of the first tetrahedron are computed for $|\alpha| = 0, \dots, n-1$, from

$$L_{3i} = g_{1,\alpha}L_{1,\alpha_1+1} + g_{2,\alpha}L_{2,\alpha_2+1} + g_{3,\alpha}M_{3,\alpha_3+1}, \quad i = n - |\alpha|.$$

The labels for the second and the third tetrahedron are computed in a similar fashion. We assume that the intermediate knot-nets $\{\{L_{1j}\}, \{L_{2j}\}, \{M_{3j}\}, j = 1, \dots, n\}$ are linearly independent.

2. Point the arrows on the tetrahedron downwards and place the original coefficients R_{α} along the lateral face of the pyramid. Carry out the computation and collect the new coefficients S_{α} along the base of the pyramid.

$$\begin{bmatrix} x_4, x_3, x_2, x_1 \end{bmatrix} \xrightarrow{1} \begin{bmatrix} x_5, x_4, x_3, x_2 \end{bmatrix} \xrightarrow{1} \begin{bmatrix} x_6, x_5, x_4, x_3 \end{bmatrix}$$

$$\begin{bmatrix} x_4, x_3, x_2, x_1 \end{bmatrix} \xrightarrow{1} \begin{bmatrix} x_4, x_3, x_2 \end{bmatrix} \xrightarrow{1} \begin{bmatrix} x_5, x_4, x_3 \end{bmatrix}$$

$$\begin{bmatrix} x_5, x_4, x_3 \end{bmatrix}$$

$$\begin{bmatrix} x_1, x_2, x_1 \end{bmatrix} \xrightarrow{1} \begin{bmatrix} x_3, x_2 \end{bmatrix}$$

$$\begin{bmatrix} x_1, x_2, x_1 \end{bmatrix} \xrightarrow{1} \begin{bmatrix} x_2, x_1 \end{bmatrix}$$

$$\begin{bmatrix} x_2, x_1 \end{bmatrix} \xrightarrow{1} \begin{bmatrix} x_2 \end{bmatrix}$$

$$\begin{bmatrix} x_2, x_1 \end{bmatrix}$$

$$\begin{bmatrix} x_2 \end{bmatrix}$$

$$\begin{bmatrix} x_2, x_1 \end{bmatrix}$$

$$\begin{bmatrix} x_2 \end{bmatrix}$$

$$\begin{bmatrix} x_3, x_2 \end{bmatrix}$$

$$\begin{bmatrix} x_3, x_2 \end{bmatrix}$$

$$\begin{bmatrix} x_3, x_2 \end{bmatrix}$$

$$\begin{bmatrix} x_3, x_2 \end{bmatrix}$$

$$\begin{bmatrix} x_4, x_3 \end{bmatrix}$$

$$\begin{bmatrix} x_4, x_3 \end{bmatrix}$$

$$\begin{bmatrix} x_1, 1 \end{bmatrix}$$

$$\begin{bmatrix} x_2 \end{bmatrix}$$

$$\begin{bmatrix} x_2 \end{bmatrix}$$

$$\begin{bmatrix} x_3 \end{bmatrix}$$

$$\begin{bmatrix} x_3 \end{bmatrix}$$

$$\begin{bmatrix} x_3 \end{bmatrix}$$

$$\begin{bmatrix} x_3 \end{bmatrix}$$

$$\begin{bmatrix} x_4, x_4 \end{bmatrix}$$

Figure 4.3: Univariate divided and forward difference evaluation algorithms

3. Repeat steps 1 and 2 two more times with the second and third tetrahedron using the output of the previous step as the input into the next step. After 3 steps, the coefficients at the base of the tetrahedron are the desired coefficients U_{α} .

This general change of basis algorithm is $O(n^3)$ since each tetrahedron has $O(n^3)$ nodes. Therefore, a general Lagrange evaluation algorithm takes $O(n^3)$ computations to evaluate the polynomial at $O(n^2)$ points. Since the geometric mesh or principal lattice configuration admits unique interpolation by a Lagrange L-basis, this algorithm can be used repeatedly to evaluate any bivariate L-basis on a regular rectangular lattice with an amortized cost of O(n) computations per point.

4.1.2 Divided and Forward Difference Algorithms

The divided difference algorithm is an algorithm for evaluating polynomials at several points, where successive computations take advantage of previous computations. The forward difference evaluation algorithm is a divided difference algorithm for evaluating a polynomial at several *equidistant* points. In the univariate case, these algorithms can be viewed as change of basis algorithms from one Newton basis to another Newton basis. We shall establish that the situation is analogous in the multivariate setting.

To appreciate the divided difference algorithm in the multivariate setting, we first very briefly describe this algorithm in the univariate setting. Suppose a polynomial L(x) of degree n in one variable is to be evaluated at points $x_1, \dots x_m$, where m is much larger than the degree n of the polynomial. For the sake of illustration, assume that the given polynomial is of degree three. Moreover suppose that the polynomial is represented in the Newton basis $1, x - x_1, (x - x_1)(x - x_2)$ and $(x - x_1)(x - x_2)(x - x_3)$. If that is not the case, then the polynomial can always be evaluated at n+1 distinct points and then the coefficients of the polynomial L(x) with respect to the Newton basis can be computed using well-known techniques. The coefficients of the polynomial with respect to the Newton basis are the divided differences as shown below:

$$L(x) = L[x_1] + L[x_2, x_1](x - x_1) + L[x_3, x_2, x_1](x - x_1)(x - x_2) + L[x_4, x_3, x_2, x_1](x - x_1)(x - x_2)(x - x_3)$$

Given these coefficients, one can compute the new coefficients of the same polynomial with respect to the new Newton basis 1, $x - x_2$, $(x - x_2)(x - x_3)$ and $(x - x_2)(x - x_3)(x - x_4)$ by applying the change of basis algorithm from one Newton basis to another Newton basis as shown



Figure 4.4: One step of bivariate divided difference evaluation algorithm

in the left diagram of Figure 4.3. Observe that the coefficient of the polynomial with respect to the basis function 1 in the new Newton basis (designated as $[x_2]$ in Figure 4.3) is the value of the polynomial at x_2 . We now "roll" this algorithm along to compute the values of the given polynomial at successive points x_3 , x_4 and so on. This algorithm has computational complexity O(n).

The forward difference algorithm is akin to the divided difference algorithm but takes advantage of the fact that the points at which the polynomial is to be evaluated are equidistant. In this case, it is possible to get rid of all multiplications so that each successive evaluation is based purely on addition. This goal is achieved by premultiplying the divided differences by appropriate constants, which depend upon the fixed distance between successive points, and then reducing the rest of the computation to pure addition as shown on the right diagram of Figure 4.3, where h refers to the distance between successive equidistant points. Both the divided and forward difference algorithms are discussed in standard numerical analysis textbooks [CdB80]. We refer the readers to [Muh73, GLC82] for a generalization of divided differences to the multivariate setting.

We now describe how a bivariate version of the divided difference and forward difference algorithms can be obtained by specializing the change of basis algorithms between L-bases described in Section 4.1.1. Given any bivariate polynomial of degree n, we first compute the representation of the polynomial in the standard Newton L-basis defined by the knot-net $\mathcal{L} = \{\{L_{1j}\}, \{L_{2j}\}, \{L_{3j}\}, j = 1, \dots, n\}$, where $L_{1j} = 1, L_{2j} = x - a_j$ and $L_{3j} = y - b_j$. Let the coefficients of the given polynomial with respect to this basis be R_{α} . We now perform the change of basis algorithm from this Newton L-basis to the "next" Newton L-basis defined by the knot-net $\mathcal{M} = \{\{M_{1j}\}, \{M_{2j}\}, \{M_{3j}\}, j = 1, \dots, n\}$, where $M_{1j} = 1, M_{2j} = x - a_{j+1}$ and $M_{3j} = y - b_{j+1}$. This change of basis algorithm can be achieved in two steps: first, perform the change of basis



Figure 4.5: Second step of bivariate divided difference evaluation algorithm

algorithm from the given knot-net to the knot-net $\mathcal{P} = \{\{P_{1j}\}, \{P_{2j}\}, \{P_{3j}\}, j = 1, \dots, n\}$, where $P_{1j} = 1, P_{2j} = x - a_j$ and $P_{3j} = y - b_{j+1}$. Let S_{α} be the coefficients of the given polynomial with respect to this intermediate basis. In the next step perform the change of basis algorithm from the knot-net \mathcal{P} to the knot-net \mathcal{M} . The left diagram of Figure 4.4 illustrates the first step of this algorithm for the case n = 2. Since many arrows are 0 or 1, this left diagram simplifies to the right diagram of Figure 4.4. In general for degree n, there will be i + 1 arrows with label $b_{n-i} - b_1$ in the first step of this algorithm for $i = 0, \dots, n-2$. Therefore this step of the algorithm requires $O(n^2)$ computations. The second step of the algorithm yields the left diagram in Figure 4.5, which then simplifies to the right diagram in Figure 4.5. This step also requires $O(n^2)$ computations. Therefore, the computational complexity of the bivariate divided difference algorithm is $O(n^2)$ computations per point.

The bivariate forward difference algorithm takes advantages of equidistant points by premultiplying the coefficients by appropriate constants as in the univariate case, thus reducing the computations to pure additions. To be more precise, the coefficients R_{α} in the right diagram of Figure 4.4 are multiplied by $\alpha_3!k^{\alpha_3}$ and the coefficients S_{α} in the right diagram of Figure 4.5 are multiplied by $\alpha_2!h^{\alpha_2}$, where h is the distance between grid points in the x-direction and k is the distance between grid points in the y-direction. After these premultiplications, the forward difference algorithm is obtained simply by changing all the labels on the arrows on these diagrams to 1. Thus every successive evaluation can be achieved simply by addition without any multiplication, thus reducing the cost to O(1) computations per point.

4.2 Dual Nested Multiplication Evaluation Algorithm

We now describe the dual nested multiplication evaluation algorithm for an L-basis that is obtained from the dual formulation of the de Boor evaluation algorithm for a B-basis [LG95a].

To evaluate a polynomial expressed in terms of an L-basis at a point \mathbf{u} , let M and N be two polynomials that vanish at \mathbf{u} , that is, $M(\mathbf{u}) = N(\mathbf{u}) = 0$. We now peform a change of basis algorithm from the L-basis defined by the given knot-net \mathcal{L} to the L-basis defined by the knot-net $\{L_{1j}, L_{2j}, N; j = 1, \dots n\}$ and then from this basis to the L-basis defined by the knotnet $\{L_{1j}, M, N; j = 1, \dots n\}$. Since all the basis functions in the L-basis defined by this last knot-net have a factor of M or N except the basis function $l_{n00}^n = L_{11} \cdots L_{1n}$, the value of the given polynomial at \mathbf{u} can now be computed simply by multiplying the coefficient of l_{n00}^n with $l_{n00}^n(\mathbf{u})$.

We shall refer to this algorithm as the dual nested multiplication algorithm. It is shown in [LG95a] that for polynomials written in terms of the multinomial basis, this algorithm specializes to a bivariate generalization of Horner's evaluation algorithm for univariate polynomials expressed in the monomial (Taylor) basis, and agrees with the algorithm for evaluation of multivariate polynomials proposed by Schumaker and Volk [SV86]. Notice that in this algorithm, one performs computations only for two tetrahedra in contrast to computations for three tetrahedra in the general change of basis algorithm. Also, for the first tetrahedron, the computation needs to be carried out only along one of the faces of the tetrahedron, as shown in the left diagram of Figure 4.6, because only the values along one of the edges at the base of the tetrahedron are needed as input for the next tetrahedron. Therefore, the computation needs to be carried out only along one edge of the tetrahedron, as shown in the right diagram of Figure 4.6, because only the value at one corner of the base of the tetrahedron is needed to evaluate the given polynomial. The computational complexity of this step is therefore only O(n), giving an overall computational complexity of $O(n^2)$.

Observe that this dual nested multiplication algorithm can be generalized somewhat in order to further simplify the computation of the labels along the edges. In this evaluation algorithm if we are given only one value of \mathbf{u} , then we have a good deal of flexibility in choosing M and N. In fact, even more generally, we can choose any set of 2n lines $\{M_i, N_i, i = 1, \dots, n\}$ passing through \mathbf{u} and the same argument goes through as long as the intermediate knot-nets are linearly independent.

In the case of a Bézier or multinomial L-basis, $L_{1j} = L_1$, $L_{2j} = L_2$ and $L_{3j} = L_3$. Given $\mathbf{v}_0 = (x_0, y_0)$, one possible choice is $M = x - x_0$ and $N = y - y_0$. But another possible choice $M = L_3(\mathbf{v}_0)L_1 - L_1(\mathbf{v}_0)L_3$ and $N = L_2(\mathbf{v}_0)L_1 - L_1(\mathbf{v}_0)L_2$ is more symmetric and has the advantage that the labels $g_{k,\alpha}$ are much simpler, as the following identities reveal:

$$L_{3} = 0 \times L_{2} + \frac{L_{3}(\mathbf{v}_{0})}{L_{1}(\mathbf{v}_{0})}L_{1} - \frac{1}{L_{1}(\mathbf{v}_{0})}M,$$
$$L_{2} = 0 \times M + \frac{L_{2}(\mathbf{v}_{0})}{L_{1}(\mathbf{v}_{0})}L_{1} - \frac{1}{L_{1}(\mathbf{v}_{0})}N.$$



Figure 4.6: Dual nested multiplication evaluation algorithm for L-bases

We close this section by highlighting with an example the difference between the up recurrence algorithms described in Section 3 and the dual nested multiplication algorithm introduced here. Consider a polynomial $L(\mathbf{u})$ represented in terms of the Lagrange L-basis defined by the knot-net

Suppose $L(\mathbf{u}) = \sum_{|\alpha|=n} R_{\alpha} l_{\alpha}^{n}$ is to be evaluated at a point (x_{0}, y_{0}) . Then a typical up recurrence algorithm organizes the computation as follows:

$$L(\mathbf{u}) = R_{200}(x_0 - \frac{1}{2})x_0 + (R_{110}x_0 + R_{020}(y_0 - \frac{1}{2}))y_0 + (R_{101}x_0 + R_{011}y_0 + R_{002}(\frac{1}{2} - x_0 - y_0))(1 - x_0 - y_0)$$

In contrast, the dual nested multiplication algorithm organizes the computation very differently. We first need to make choices. Let us define $N = x + y - x_0 - y_0$ and $M = y - y_0$. For the first tetrahedron, one obtains the following required labels by solving the appropriate system of linear equations: $g_{1,000} = g_{2,000} = a$, $g_{1,100} = g_{2,100} = g_{1,010} = g_{2,010} = b$, where $a = \frac{1}{2(x_0+y_0)} - 1$, and $b = \frac{1}{2(x_0+y_0)-1} - 1$, as shown on the left diagram of Figure 4.6. This tetrahedron yields the computation: $S_{200} = (R_{002}a + R_{101})b + R_{200}$, $S_{110} = (R_{002}a + R_{101})b + (R_{002}a + R_{011})b + R_{110}$, and $S_{020} = (R_{002}a + R_{011})b + R_{020}$. For the second tetrahedron, the necessary labels are: $g_{1,000} = c$, and $g_{1,100} = d$, where $c = \frac{2y_0-1}{2x_0}$ and $d = \frac{2y_0}{2x_0-1}$. These labels are shown on the right diagram of Figure 4.6. This tetrahedron yields the computation: $T_{200} = (S_{020}c + S_{110})d + S_{200}$. The value of the original polynomial is then finally obtained by multiplying the value T_{200} by $l_{200}^2(x_0, y_0)$, which in this case is $x_0(x_0 - \frac{1}{2})$.

5. Other Algorithms

In this section, we briefly review a few other algorithms that are closely related to the evaluation algorithms discussed so far.

5.1 Hybrid Algorithms

In addition to the algorithms for evaluating multivariate polynomials discussed so far, we are aware of some different algorithms for evaluating multivariate polynomials, most of which are spurred by considerations of stability in numerical computations [LSP87, Vol90, Pet94]. First, Volk [Vol88] has proposed a hybrid univariate nested multiplication and divided difference algorithm. Volk [Vol90] has also proposed a slightly different evaluation algorithm based on the forward difference algorithm for evaluation of polynomials at several points. In order to improve the stability of the forward difference evaluation algorithm, Volk reduces the level of indirection in the computation by solving an upper triangular system [Vol90]. A second method, discussed by Peters [Pet94], is again motivated by concerns of stability and efficiency. This algorithm extracts univariate polynomials using a forward difference or dual nested multiplication algorithm. We have not addressed the very important considerations of stability in numerical computations, for which we refer the reader to [FR88, Far91, FR87], because rather than stability the focus of this work has been to provide a conceptual framework for the unification of a large variety of evaluation algorithms for multivariate polynomials.

5.2 Factored Algorithms

By factored algorithms we mean algorithms where an intermediate basis is used to perform evaluation. For example, the divided and forward difference algorithms for evaluation were obtained by "factoring" through the Newton basis, while the Lagrange evaluation algorithm is obtained by "factoring" through the Lagrange basis.

We have concentrated on evaluation algorithms for multivariate polynomials that are expressed in terms of L-bases. Another important class of bases are B-bases, bases that are dual to L-bases. B-bases are blending functions for B-patches which were introduced by Seidel [Sei91] and shown to agree with certain multivariate B-splines on a special region of the parameter domain [DMS92]. Evaluation algorithms with computational complexity $O(n^{s+1})$ for multivariate polynomials of degree n in s variables expressed in terms of B-bases have been derived in [Sei91] using blossoming. Barry and Goldman [BG93b] and Sankar, Silbermann and Ferrari [SSF94] have discussed a technique for speeding up knot insertion and evaluation of univariate polynomials expressed in terms of univariate progressive or B-spline bases by factoring through the Newton dual basis, that is, the dual to the univariate Newton basis. Here, we only indicate why this technique of factoring can easily be extended to speed up the algorithms for evaluating multivariate polynomials, expressed in terms of B-bases, at several points. This factoring algorithm

can be viewed as dual to the divided difference algorithm. To derive this algorithm, one first converts the given polynomial into the special Newton dual B-basis defined by the knot-net of points $\mathbf{u}_{1i} = (a_{1i}, b_{1i})$, $\mathbf{u}_{2i} = (1, 0)$ and $\mathbf{u}_{3i} = (0, 1)$ for $i = 1, \dots, n$. The interesting and crucial property of this basis is that one can express a point $\mathbf{u} = (x, y)$ as follows:

$$\mathbf{u} = 1\mathbf{u}_{1i} + (x - a_{1i})(1, 0) + (y - b_{1i})(0, 1)$$

Therefore, the labels 1, $x - a_{i1}$, and $y - b_{1i}$ do not involve any divisions. This observation is the crucial property that is needed to minimize divisions and speed up the computations.

5.3 Derivative and Other Algorithms

Although we have focused only on algorithms for evaluating multivariate polynomials, these algorithms are closely related to and can easily be extended to procedures to compute derivatives of multivariate polynomials. This technique of unifying evaluation algorithms and derivative algorithms has been discussed for univariate polynomials by Goldman and Barry [GB92]. Here we briefly indicate how this extension from evaluation algorithms to derivative algorithms can be derived. The key observation is that a multinomial (Taylor) basis is defined by a point and two vectors and, therefore, change of basis algorithms to a multinomial basis defined by a point **p** and two vectors \mathbf{v}_1 and \mathbf{v}_2 amounts to computing the directional derivatives of the polynomial at the point \mathbf{p} in the directions \mathbf{v}_1 and \mathbf{v}_2 . In particular, given a Bézier B-basis defined by three points \mathbf{p}_1 , \mathbf{p}_2 and \mathbf{p}_3 , the change of basis algorithm to the multinomial B-basis defined by the point \mathbf{p}_1 and the unit vectors in the direction $\mathbf{p}_2\mathbf{p}_1$ and $\mathbf{p}_3\mathbf{p}_1$ is the generalization of the Boehm's derivative algorithm from curves to surfaces. To compute the directional derivatives at a point palong the vectors \mathbf{v}_1 and \mathbf{v}_2 of a polynomial $L(\mathbf{u})$ given with respect to an L-basis, perform the change of basis algorithm from the given L-basis to the uniform L-basis defined by the following three lines: the line through p along the direction \mathbf{v}_1 , line through p along the direction \mathbf{v}_2 , and the line at infinity.

Using principles of homogenization, blossoming, and duality [BGD91, BG93a, CLR80, LM86, Mar70, Boe80] univariate evaluation and differentiation algorithms have been unified with several other very well-known algorithms, formulas, and identities, including the Oslo algorithm, Boehm's knot-insertion and derivative algorithms, Marsden's identity, the binomial theorem, Ramshaw's blossoming algorithm, a two-term differentiation algorithm, and a two-term degree elevation formula. Although in this work we have focused solely on evaluation algorithms, it follows in conjunction with our earlier work [LG95a, LG95b] that the principles of blossoming, duality, and homogenization can be extended to provide a similar unification in the multivariate setting.

6. Conclusions and Future Work

We have presented a unified framework for evaluation algorithms for multivariate polynomials expressed in a wide variety of polynomial bases including the Bézier, multinomial, Lagrange, and Newton bases. Although in the past several different evaluation algorithms have been constructed by organizing the nested multiplications in different ways, the interpretation and unification of all these algorithms either as a way of reorganizing the computation in an up recurrence algorithm or as change of basis algorithms is new.

Variations of the up recurrence algorithm include a parallel up recurrence algorithm with computational complexity $O(n^3)$, a nested multiplication algorithm with computational complexity $O(n^2lgn)$, a ladder recurrence algorithm with computational complexity $O(n^2)$, and a generalization of the Aitken-Neville algorithm for Lagrange L-bases with computational complexity $O(n^3)$. Specializations of this class of algorithms include the de Casteljau algorithm for Bézier bases, the evaluation algorithms for multinomial and Newton bases proposed by Carnicer and Gasca, and the evaluation algorithms for multinomial bases proposed by de Boor and Ron.

Variations of change of basis algorithms between L-bases yielded a divided difference algorithm with computational complexity $O(n^2)$ per point, a forward difference algorithm with computational complexity O(1) per point, and a Lagrange evaluation algorithm with amortized computational complexity O(n) per point for the evaluation of polynomials at several points. This class of algorithms also include a dual nested multiplication algorithm with computational complexity $O(n^2)$ which along with the up recurrence algorithms described in the previous paragraph can be considered as generalization of Horner's algorithm for the evaluation of univariate polynomials. The evaluation algorithm for multinomial bases proposed by Schumaker and Volk is a special case of a dual nested multiplication algorithm.

New algorithms derived and discussed in this work can best be appreciated in the case of Lagrange bases, where unlike multinomial or Newton bases considerable simplifications do not occur. For multivariate polynomials expressed in Lagrange L-bases, we have described several evaluation algorithms including a nested multiplication algorithm with computational complexity $O(n^2lgn)$ generated by removing redundant arrows from the up recurrence algorithm, a generalization of the Aitken-Neville algorithm with computational complexity $O(n^2)$, and a dual nested multiplication algorithm with computational complexity $O(n^2)$. We have presented specific examples to demonstrate that these algorithms are all distinct both conceptually and in practice.

It has been very satisfying to discuss all the well-known algorithms for evaluating multivariate polynomials in a single unified framework. In fact, we are not aware of any evaluation algorithm for multivariate polynomials that cannot be derived from the techniques presented in this work. It would be satisfying to integrate into our formulation evaluation algorithms for multivariate polynomials expressed in terms of other useful bases such as multivariate Hermite bases. A generalization of the Aitken-Neville recurrence for Hermite bases defined over geometric meshes is currently under investigation by Habib and Goldman [HG95].

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