Cubature Points Based Triangular Spectral Elements: an Accuracy Study

Richard Pasquetti* and Francesca Rapetti

Lab. J. A. Dieudonné, Université Côte d'Azur, CNRS, Inria, F-06108 Nice, France.

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Abstract. We investigate the cubature points based triangular spectral element method and provide accuracy results for elliptic problems in non polygonal domains using various isoparametric mappings. The capabilities of the method are here again clearly confirmed.

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Key words: Spectral element method, simplicial meshes, cubature points, Fekete points, isoparametric mappings.

1 Introduction

Extending the capabilities of the quadrangle based spectral element method to the triangle based one, say from the QSEM to the TSEM, has received a considerable attention in the last twenty years. The relevant choice of interpolation points for the triangle has thus motivated a lot of works, see e.g. the review paper [15] and references herein. Several sets of points have indeed been proposed, some of them showing the advantage of simplicity in their generation, thus allowing an easy extension to the tetrahedron, *e.g.* the so-called warp & blend points [21], whereas some other ones are more satisfactory from the theoretical point of view, *e.g.* the celebrated Fekete points of the triangle, but much more difficult to compute [20].

With respect to the QSEM, based on the use of the tensorial product of Gauss-Lobatto-Legendre (GLL) nodes, such interpolation points are however not quadrature points, so that the so called spectral accuracy gets lost if not using for the quadratures a separate set of points, as *e.g.* done for the Fekete-Gauss approach [14]. Using two separate sets of points shows however some drawbacks, because the mass matrix is then no longer diagonal. Thus, when solving an evolution problem with an explicit time scheme, one

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^{*}Corresponding author. *Email addresses:* richard.pasquetti@unice.fr (R. Pasquetti), francesca.rapetti@unice.fr (F. Rapetti)

has at each time step to solve a mass matrix algebraic system. Also, a diagonal mass matrix can be useful to set up high order differentiation operators [12]. This is why it is of interest to design a *TSEM* that makes use of a single set of points showing both nice quadrature and interpolation properties. This was the goal of the cubature points based *TSEM*, as developed in the last few years [2,5,9,13].

Another difficulty with SEMs, and more generally with high order FEMs, is to maintain the accuracy when the computational domain is no longer polygonal. Several strategies have been proposed to handle curved elements, from transfinite mappings [6] to PDEs based transformations, *e.g.* the harmonic extension.

The present work follows two anterior studies:

- In [16], using the Fekete-Gauss *TSEM* we compared different strategies for curved triangular spectral elements, *i.e.* when isoparametric mappings are required to correctly approximate with curved spectral elements the presence of a curved boundary.
- In [17], for computational domains of polygonal shape we compared the cubature *TSEM*, as proposed in [9], and the Fekete-Gauss one. In terms of accuracy for elliptic problems, it turned out that these two *TSEMs* compare well. With non Dirichlet or non homogeneous Neumann conditions, *i.e.* as soon as boundary integrals are involved, some care is however needed if using the cubature *TSEM*.

Here the goal is to extend the results obtained in [16] to the case where the cubature *TSEM* is used. As in [16, 17] we only focus on elliptic problems, but keep in mind for future works the inherent difficulties associated to constrained operators, see e.g. [1].

2 Cubature points based TSEM

For the sake of completeness, let us recall that [17]:

If two different sets of points are used for interpolation and quadrature, then the space P_N(Î) of polynomials of maximal (total) degree N, defined on the reference triangle Î = {(r,s):r∈(-1,1),s∈(-1,-r)}, is usually used as approximation space. The cardinality of this space equals n=(N+1)(N+2)/2, that can be associated to n interpolation points if using Lagrange polynomials as basis functions. If 3 of these nodes coincide with the vertices of the element, then 3N of these n points should belong to the edges of Î and the remaining (N-1)(N-2)/2 are the inner nodes. Usually, the edge nodes proposed in the literature coincide with the GLL points. Since one does not know an explicit formulation of the Lagrange basis functions, say φ_i(r,s), 1≤i≤n, to compute their values or those of their derivatives at given point one generally makes use the orthogonal Kornwinder-Dubiner (KD) basis [4], for which explicit formula exist. Gauss points for the triangle and the corresponding quadrature formula may be found in the literature, up to degree M ≈ 20 if a

symmetric distribution of the points is desired. In practice, one may choose M=2N, so that both the stiffness and mass matrix are exactly computed, since their entries are polynomials of degree 2N and 2N-2, respectively. For details on the implementation of the Fekete-Gauss approach, see *e.g.* [14].

• If using a single set of points, as before 3N of the interpolations points must belong to the triangle boundary with 3 of them at the vertices. As demonstrated in [7,22], such a strong constraint forbids the possibility of finding a set of cubature points providing a sufficiently accurate cubature formula, if looking for basis functions that span the space \mathbb{P}_N . To overcome this difficulty, as first developed in [2], the idea is then to enrich the space \mathbb{P}_N by polynomial bubble functions of degree N' > N. This indeed allows to include new cubature points inside the triangle while keeping the element boundary nodes number equal to 3*N*. In the reference triangle \hat{T} , this may be achieved by introducing the polynomial space $\mathbb{P}_N \cup b \times \mathbb{P}_{N'-3}$, where b is the (unique) bubble function of $\mathbb{P}_3(\hat{T})$, namely b(r,s) = (r+1)(s+1)(r+s). The cardinality of this space then equals n'=3N+(N'-1)(N'-2)/2. Now, to compute the Lagrange polynomials at a given points one should use an extended KD basis, composed of the usual KD basis of \mathbb{P}_N completed by those KD polynomials of $\mathbb{P}_{N'-3}$ which once multiplied by the bubble function *b* are of degree strictly greater than N. Of course, N' should be chosen as small as possible to avoid a useless increase of the inner nodes number: In [2] and posterior works, N' is chosen such that it exists a cubature rule exact for polynomials of degree N+N'-2. The determination of N', together with the cubature points and weights gives rise to a difficult optimization problem, see [9] for details. It turns out that N' - N increases monotonically with N: N' = N+1, for 1 < N < 5, N' = N+2 for N = 5 and N' = N+3 for 5 < N < 10. It should be noticed that with the cubature TSEM, neither the mass matrix nor the stiffness matrix are exactly computed, since their entries are of degree 2N' and 2N'-2, respectively.

Our in house Fekete-Gauss and cubature codes make use of the condensation technique, *i.e.* one first computes the unknowns associated to the edges of the elements and then reconstructs the numerical solution inside locally. Thus, the algebraic systems that result from the two *TSEM* approximations are exactly of same size, and so the computational times compare well. They are simply solved by using a standard conjugate gradient method, with Jacobi preconditioner in a matrix free implementation [8].

A difficulty however arises when the boundary conditions are not of Dirichlet or homogeneous Neumann type, *i.e.* when the weak formulation involves boundary integrals. In case of the Fekete-Gauss *TSEM*, such boundary integrals can be easily computed since the edge Fekete nodes coincide with the GLL points. On the contrary, the edge cubature points are not of Gauss type. In [17],

• it is pointed out that for the Robin condition $\partial_n u + \alpha u = g$, being *g* a given function and α a coefficient possibly zero, using a quadrature rule based on the boundary cubature points is not satisfactory,

• it is suggested to approximate these integrals by using on each element edge of the boundary a Gauss quadrature rule, *e.g.* the one based on the GLL points.

Since the restrictions of the Lagrange polynomials φ_i at the element edges of the boundary are polynomials of degree N, one can span this polynomial space with the Legendre polynomials, say $L_i(r)$ with $r \in [-1,1]$, for the reference edge, and $0 \le i \le N$. Then, one can set up the Vandermonde matrices of size $(N+1) \times (N+1)$ based on the cubature and GLL points, say V_{Cub} and V_{GLL} . One has $e.g.(V_{Cub})_{ij} = L_j(r_i)$, with $r_i \in [-1,1]$ for the edge cubature points. Then, the matrix $V_{GLL}V_{Cub}^{-1}$ allows to compute at the GLL points quantities known at the cubature points. Especially, each column of this matrix provides the values of the edge Lagrange polynomials at the GLL points.

Another difficulty arises with curved spectral elements. Indeed, for straight triangular elements, the edge Jacobian determinant is constant on each edge and proportional to its length. On the contrary, when considering curved triangular elements, some care is needed for a relevant computation of the edge Jacobian determinants at the GLL points from those at the cubature points. Thus, once the entries of the Jacobian matrix have been computed at the edge cubature points, one can

- compute the edge Jacobian determinant and then use the polynomial interpolation to get the Jacobian at the GLL points,
- use the polynomial interpolation to get the Jacobian matrix entries at the GLL points and then compute the Jacobian determinant.

Because the edge Jacobian determinant is generally a polynomial of degree greater than N, the results of these two approaches slightly differ. There is indeed no interpolation error for the Jacobian matrix entries but there is such an error for the Jacobian matrix determinant. This is why in our cubature points based solver, it is the second approach which is implemented, even if a little more expensive than the first one.

3 Isoparametric mappings

For the cubature *TSEM*, hereafter we make use of the isoparametric mappings that we discussed in [16]:

- The bending procedure that we introduced in [8];
- The transfinite interpolation discussed for the triangles in [18];
- The harmonic extension;
- The linear elasticity approach, see e.g. [19].

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Moreover, local mappings are used, *i.e.* we simply distort the elements at the boundary.

As in [16], we assume to have at hand a parametric description of the boundary of Ω , *i.e.*, Γ is defined by a vector function, say $\mathbf{x}_{\Gamma}(t) = (\mathbf{x}_{\Gamma}(t), \mathbf{y}_{\Gamma}(t))$, where $t \in [t_{\min}, t_{\max}]$. Let $A_1A_2A_3$ be a triangular element at the boundary, with A_1 the inner node and A_2 and A_3 on the boundary. Then, for all the four approaches it is assumed that the boundary nodes, *i.e.* the interpolation points of Γ , are at the intersection of the lines joining the inner vertex A_1 and the interpolation points of the straight line (A_2A_3) , *i.e.* the edge of the element, say \tilde{T} , provided by the piecewise linear \mathbb{P}_1 mapping. This requires one to solve, in general using a simple numerical procedure, N-1 (number of nodes of an edge without the end points) generally non-linear equations. Then, it remains to define n'_p inner interpolation points, with $n'_p = n' - 3N = (N' - 1)(N' - 2)/2$, for the cubature *TSEM*.

For the paper to be self contained, let us briefly describe the previously mentioned isoparametric mappings [16]:

• The bending procedure: Assume that \tilde{F} is the interpolation point obtained with the affine \mathbb{P}_1 mapping. Let \tilde{G} and G be the points at the intersections of $(A_1\tilde{F})$ with the line (A_2A_3) and the boundary Γ , respectively. Then, the bending procedure [8] consists of stating that F is homothetic to \tilde{F} by the homothety of center A_1 and of ratio $A_1G/A_1\tilde{G}$, so that:

$$A_1F = \frac{A_1G}{A_1\tilde{G}}A_1\tilde{F}$$
 and $d \equiv \tilde{F}F = \left(\frac{A_1G}{A_1\tilde{G}} - 1\right)A_1\tilde{F}$

where *d* stands for displacement. One notices that for this bending procedure, only the point *G* of Γ influences the location of the interpolation point *F*. In order to define the inner interpolation points, one has numerically to solve n'_v equations.

The transfinite interpolation: For the triangle, the method makes use of the barycentric coordinates, say (λ₁,λ₂,λ₃). Recall that λ_i=0 stands for the edge opposite to the vertex *i*, λ_i=*Constant* to a line parallel to this edge and that vertex *i* belongs to λ_i=1. General transfinite interpolation formula are given in [18] (different approaches are however possible, see *e.g.* [3]). For the triangle, if the displacement *d* vanishes at the two edges (A₁A₂) and (A₁A₃), again assuming that A₂A₃ is the curved edge, one has:

$$\boldsymbol{d}(\lambda_1,\lambda_2,\lambda_3) = \lambda_2 \boldsymbol{d}(0,1-\lambda_3,\lambda_3) + \lambda_3 \boldsymbol{d}(0,\lambda_2,1-\lambda_2).$$

This means that two points of the boundary Γ influence the location of an inner interpolation point *F*. They are those at the intersections of Γ with the lines parallel to (A_1A_2) and (A_1A_3) and passing by the point \tilde{F} . One notices that the method requires solving $2n'_p$ equations.

• Harmonic extension: The goal is here to define the inner interpolation points by solving, for each curved element *T*, the weak form of the Laplace Dirichlet problem:

$$\Delta d = 0$$
 in T , $d|_{\partial \tilde{T}} = g$

with g given on (A_2A_3) and g = 0 on the two other sides. To resolve this vector Laplace problem, that in fact yields two uncoupled scalar problems, one has to set up the differentiation matrices, say $D_{\tilde{x}}$ and $D_{\tilde{y}}$ that allows one to compute derivatives in \tilde{T} . To this end, one makes use of the differentiation matrices D_r and D_s and applies the chain rule. Since the mapping from \hat{T} to \tilde{T} is affine, the Jacobian matrix and Jacobian determinant are constant. For the same reason, one may solve directly for the interpolation point coordinates. Finally, one should invert a matrix of size $n'_p \times n'_p$, in order to compute the coordinates of the inner interpolation points. The approach is thus rather simple and in practice not costly. In [11], where global mappings are considered, it is however outlined that the harmonic extension may fail to define a mapping.

 Linear elasticity: Here the curved domain is viewed as the deformation of triangles, that is governed by the equation of linear elasticity. Introducing the Lame coefficients λ and μ, the displacement field is governed by the Navier-Cauchy equation:

$$\mu \Delta d + (\lambda + \mu) \nabla (\nabla \cdot d) = 0 \quad \text{in } \tilde{T}, \quad d|_{\partial \tilde{T}} = g.$$

There is now a coupling between the components of the displacement field. Using the ingredients previously described for the harmonic extension, the weak form of this elasticity problem yields a matrix system of size $2n'_p$. The solution of this system provides the displacements of the inner interpolation points. Again, one can also compute directly their coordinates. Because the forcing term is zero, the mapping is here parametrized by the ratio $\mu/(\lambda+\mu)=1-2\eta$, where $\eta \in (-1,0.5)$ is the so called Poisson ratio.

4 Accuracy study

First, one of the test-cases introduced in [16] has been revisited. Using the cubature *TSEM* we solve the Poisson equation, $-\Delta u = f$, with exact solution $u_{ex} = \cos(10x)\cos(10y)$, so that $f = 200u_{ex}$, and Dirichlet boundary conditions, in the computational domain and mesh (the so-called medium mesh of [16]) presented in Fig. 1 (left). For the elasticity isoparametric mapping, the Poisson ratio is taken equal to 0.2.

For the different isoparametric mappings, the max norm of the error, obtained from the errors at the cubature points, is shown in Figure 2 (left). The curve labeled "polygon" is the one obtained for the domain defined by the \mathbb{P}_1 mesh, *i.e.* no isoparametric mapping is involved in this case. Such results compare well with those obtained with the Fekete-Gauss *TSEM*, recalled in Figure 2 (right). Again, one observes that the best results are obtained with the harmonic extension or with the linear elasticity approach, *i.e.* with the isoparametric mappings based on PDEs rather than transfinite interpolations. As in [17], one observes that a loss of accuracy occurs for N = 9. It is due to a slight lack of accuracy of the cubature rule [10].



Figure 1: Computational domains and corresponding \mathbb{P}_1 meshes.



Figure 2: Max norm of the error vs the polynomial degree N for the Poisson-Dirichlet problem using the cubature TSEM (at left) and the Fekete-Gauss TSEM (at right).

Rather than the Dirichlet condition, we now use the Robin condition $\partial_n u + u = g$, with g derived from the exact solution $u_{ex} = \cos(10x)\cos(10y)$. Results obtained with the Fekete-Gauss solver and with cubature points based solver are compared in Figure 3, using the PDEs based isoparametric mappings (harmonic extension and linear elasticity). Clearly, the results obtained with the Fekete-Gauss and the cubature *T*SEM compare well. Thus, even if curved elements are involved, if the boundary data are correctly handled [16] then the cubature approach proposed in [9] is quite satisfactory (at least for $N \leq 8$).

To extend the present study we consider the computational domain and \mathbb{P}_1 -mesh shown in Figure 1 (right). The boundary of this star domain is parametrized by the polar angle θ :

$$x = \cos\theta (1 + 0.3\cos(7\theta))$$
$$y = \sin\theta (1 + 0.2\cos(7\theta)).$$



Figure 3: Max norm of the error vs the polynomial degree N for the Poisson-Robin problem using the cubature TSEM and the Fekete-Gauss TSEM.

The \mathbb{P}_1 -mesh is generated from 50 boundary nodes, defined by $\theta \propto 2\pi/50$, and the number of elements equals 174.

Isoparametric mappings based on the harmonic extension and linear elasticity approaches are used. Till now, the linear elasticity isoparametric mapping was characterized by a Poisson ratio $\eta = 0.2$. Then, it is of interest to carry out a sensitivity study to this parameter. For isotropic materials one has $\eta \in (-1,0.5)$, most materials are such that $\eta \in (0,0.5)$, and for typical engineering materials $\eta \in (0.2,0.5)$. Hereafter we make computations with the following Poisson ratio $\eta = \{0.1,0.2,0.4\}$.

For one particular element of the mesh and a polynomial approximation degree N=6, Figure 4 shows the distribution of the cubature nodes obtained from the \mathbb{P}_1 -mapping and from the isoparametric \mathbb{P}_6 -mapping, when using the harmonic extension or the linear elasticity mappings associated to the different Poisson ratio.

We make use of the cubature *TSEM* and the exact solution $u_{ex} = \cos(10x)\cos(10y)$ to set up the source and boundary terms, f and g, of the following elliptic PDE $-\Delta u + u = f$. Accuracy tests are made using Dirichlet conditions, *i.e.* u = g at the boundary nodes, and also the Robin condition $\partial_n u + u = g$. The results obtained for these Dirichlet and Robin problems are shown in Figure 5, at left and at right, respectively.

One observes that:

- For the linear elasticity isoparametric mapping, the sensitivity to the Poisson ratio η is low. With the Robin condition, all curves coincide. With the Dirichlet condition, one can discern a slight improvement for the largest value $\eta = 0.4$, the convergence curve being then the closest to the harmonic extension one. Note that this is surprising, since if $\eta \rightarrow 0.5$ then $\mu \rightarrow 0$.
- The decrease of the error for the Robin problem is exponential, whereas this is less obvious in the Dirichlet case. In fact, the convergence curves show two parts, be-



Figure 4: Distribution of the cubature nodes before and after the isoparametric mapping: Harmonic extension (top left), elasticity with $\eta = 0.1$ (top right), $\eta = 0.2$ (bottom left) and $\eta = 0.4$ (bottom right).



Figure 5: Max norm of the error vs the polynomial degree N, using the cubature TSEM for an elliptic problem in the star domain, with Dirichlet (at left) and Robin (at right) conditions.

low and beyond a critical value of the polynomial degree. Roughly speaking, for the smallest values of N the convergence rate is governed by the approximation of the solution in the bulk of the domain, whereas for the largest ones it is enforced by its approximation at the boundary. This is why, see Figure 5, for small N the convergence curves coincide with those obtained when using as computational domain the polygon obtained from the 50 boundary nodes, but differ for N larger than threshold values, say, for the present \mathbb{P}_1 discretization of the star-shaped domain, N=5 and N=3 with Dirichlet and Robin conditions, respectively.

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