Towards High-order Volume Potential Evaluation for Unstructured Meshes

J. Bevan, UIUC

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Introduction: Volume Potentials are Everywhere



What is an Integral Equation?

► Integral equations one way of solving PDEs, involve operations like solution of integral equations:

$$\int_{\Omega} K(\mathbf{x}, \mathbf{y}) \, \sigma(\mathbf{y}) \, d\mathbf{y} = f(\mathbf{x}) \tag{1}$$

(K is a kernel function and $\sigma(\mathbf{y})$ unknown)

Or evaluation of integrals:

$$\phi(\mathbf{x}) = \int_{\Omega} K(\mathbf{x}, \mathbf{y}) \, \sigma(\mathbf{y}) \, d\mathbf{y} \tag{2}$$

Central idea: solution of PDE sum of "fundamental solutions" G that solve PDE for point source (in the weak sense):

$$\mathcal{L}G(x,y) = \delta(y-x) \tag{3}$$

 \mathcal{L} is linear differential operator associated with PDE, G the Green's function, δ Dirac delta.

How are They Useful?

Example: Poisson equation:

$$\nabla^2 \phi = f \tag{4}$$

Solution given by evaluating:

$$\phi(\mathbf{x}) = \int_{\Omega} G(\mathbf{x}, \mathbf{y}) f(\mathbf{y}) d\mathbf{y} \qquad G(\mathbf{x}, \mathbf{y}) = \frac{1}{4\pi |\mathbf{x} - \mathbf{y}|}$$
 (5)

- ► Integral equation methods evaluated with FMM competitive efficiency-wise with other approaches², and have excellent conditioning
- Evaluation of the integrals can pose challenges

Numerical Integral Evaluation

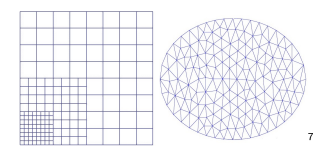
- Issue: Gaussian quadrature assumes integrand approximated well by polynomials
- ightharpoonup Poor assumption since K is usually singular or near singular
- Consider common singularity, quadrature converges³ slowly:

$$y \in [-1, 1], \ I(y) = \int_{-1}^{1} |x - y|^{-\delta} dx, \ |I - I_n| < \mathcal{O}(n^{-1+\delta})$$
 (6)

- ▶ Approaches include: product integration rules⁴; compute offline, weights that exactly integrate up to a polynomial order
- ▶ Transformation of the integral⁵(e.g. to a semi-infinite domain), causes singularity to vanish
- Adaptive quadrature; adaptively divide domain based on error estimate, compute on each subdomain

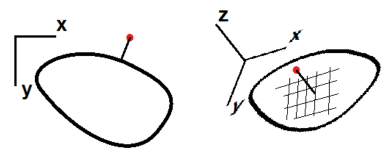
More General Scheme for Unstructured Meshes

- Mentioned approaches lack generality: tied to specific integrand and/or for unmapped/structured meshes
- We want a high-order general purpose quadrature method that works for unstructured meshes and complex geometries
- ► For inspiration look to QBX (Quadrature by Expansion)⁶: high-order, geometric flexibility



Methodology: QBX for Volume Potentials

- ▶ Volume potential share similarities to layer potentials
- Same main challenge: devising quadrature to handle singularity
- ► Take same approach: QBX
- But where do we put our expansion center, fictitious dimension?
- Off-surface: layer potential physically defined, off-volume has no requirements



Trial Scheme

- Absent any compelling choice for off-volume potential, choose obvious one:
- ▶ Consider 3D Poisson scheme: approximate G=1/r kernel with $\hat{G}=1/\sqrt{r^2+a^2}$
- Effectively a parameter is the distance from expansion center to eval point in the fictitious dimension, and kernel is no longer singular
- Choose a "good" a so the kernel is smooth and take QBX approach of evaluating Taylor expansion of de-singularized kernel back at desired eval point

Is trial scheme high-order?

- No, in fact seems to be limited to second order regardless of expansion order.
- Consider example results in figure below for 6th order expansion.
- ▶ Why only second order?

2nd order convergence plot of K2S6

Preliminary Error Analysis

- We would like to examine the error $\epsilon = |\mathsf{Exact}|$ potential QBX computed potential and it's dependence on a
- ▶ Call T_k the k-th order Taylor series expansion of \hat{G} about d and evaluated at a=0:

$$T_k(r,d) = \sum_{n=0}^{k} \frac{(-d)^n}{n!} \hat{G}^{(n)}(r,d)$$

So our error is:

$$\epsilon_k = \int_{\Omega} G(r)\sigma(r) dr - \int_{\Omega} T_k(r,d)\sigma(r) dr$$

► This form seems complicated to inspect, is there a way to avoid the integrals and factor out the density?

Error in Fourier Space

Consider the action of the Fourier transform on the error:

$$\mathcal{F}[\epsilon] = \mathcal{F}\left[\int G \,\sigma \,dr\right] - \mathcal{F}\left[\int T_k \,\sigma \,dr\right]$$

and by the convolution theorem:

$$= \mathcal{F}[G] \mathcal{F}[\sigma] - \mathcal{F}[T_k] \mathcal{F}[\sigma] = \mathcal{F}[\sigma] (\mathcal{F}[G] - \mathcal{F}[T_k])$$
$$\mathcal{F}[T_k] = \sum_{n=0}^k \frac{(-d)^n}{n!} \mathcal{F}[\hat{G}^{(n)}(r,d)]$$

▶ This looks more reasonable, let's examine the behavior of $\mathcal{F}[G] - \mathcal{F}[T_k]$ with respect to d.

Fourier Transform Particulars

- ▶ Need 3D Fourier transform; both *G* and *T* are radially symmetric, so simplifications can be made: transforms can be given in terms of the scalar *k* in Fourier space.
- ▶ It is known that $\mathcal{F}[1/r] = 1/\pi k^2$
- ▶ With some work one can show:

$$\mathcal{F}\left[\frac{1}{\sqrt{r^2 + a^2}}\right] = \frac{2a}{k} K_1(2\pi ak)$$

where $K_n(x)$ is the modified Bessel function of second kind

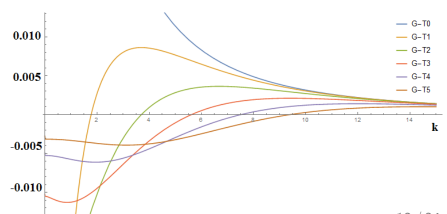
- ▶ Reduces to expected form for $\lim_{a\to 0} \frac{2a}{k} K_1(2\pi ak) = 1/\pi k^2$
- ▶ Without concerning ourselves with details, in general we find:

$$\mathcal{F}[T_k] = \sum_{n=-1}^{k} C_n \, d^{n+2} \, k^n K_n(2\pi k d)$$

Examination of Error

- k dependence tells us how well the expansion preserves low vs high modes in real space (in figure below d=0.2)
- ▶ Taylor expansion of $\mathcal{F}[T_5]$ wrt d about 0:

$$\frac{1}{\pi k^2} + \frac{\pi d^2}{10} + \frac{1}{20} \pi^3 d^4 k^2 + \mathcal{O}(d^6)$$



Cause: Approximation Error

- It seems there is some additional approximation error that limits convergence, truncation error from Taylor series not the issue
- We replaced Green's function with de-singularized approximation, what does approximate kernel correspond to?
- Remember that for our Laplace equation

$$\nabla^2 G(x, y) = \delta(y - x) \tag{7}$$

 However our de-singularized Green's function doesn't satisfy this, instead of solving for a point source it solves for a blob source: ζ

$$\nabla^2 \hat{G}(x, y) = \zeta(y - x) \tag{8}$$

Dirac Delta Approximation and Moment Conditions

- ► The quality of our Green's function approximation then depends upon the quality of our Dirac delta approximation from our choice of a blob
- ► The order of convergence of this approximation can be shown^{8,9} to depend upon the *moment conditions*, where for a s order accurate approximation we require:

$$\int \zeta(\mathbf{x}) \, d\mathbf{x} = 1 \tag{9}$$

$$\int \mathbf{x}^{\mathbf{i}} \zeta(\mathbf{x}) d\mathbf{x} = 0, |\mathbf{i}| < s - 1$$
 (10)

$$\int \mathbf{x}^s \, \zeta(\mathbf{x}) \, d\mathbf{x} < \infty \tag{11}$$

High Order De-singularized Kernels

- ▶ We can see that $1/\sqrt{r^2+a^2}$ is actually 0th order approximation (the third condition isn't totally satisfied for s=2)
- ► This approximation error in turn bounds our overall error and limits convergence rate to at best 2nd order
- Nhy better convergence than expected from kernel? Postulated QBX expansion satisfies moment conditions for s=2, remains to be verified
- Possible to construct higher-order kernels by satisfying moment conditions 10 for larger s
- For example consider the 2nd order kernel (i.e. satisfies moment conditions for s=2:

$$\zeta = \frac{15/2}{(r^2 + a^2)^{7/2}} \to \hat{G} = \frac{r^2 + \frac{3}{2}a^2}{(r^2 + a^2)^{3/2}}$$
(12)

Fourier Space Error Analysis of High Order Kernels

- Define $T_{s,k}$ to be similar to our T_k from before, but now for \hat{G}_s the sth order algebraic approximate kernel
- ► We can examine again the error in our computed integral in Fourier space, but now for our higher order kernel
- We find that in general:

$$\mathcal{F}[T_{s,k}] = \frac{2}{\left(\frac{s}{2}\right)!} d^n k^{n-2} \pi^{n-1} K_{\frac{s}{2}+1}(2\pi k d) + \sum_{n=\frac{s}{2}-1}^{k} C_n d^{n+2} k^n \pi^{n+1} K_n(2\pi k d)$$
(13)

► For example, consider the Taylor expansion of higher order kernels wrt *d* about 0:

$$\mathcal{F}[T_{2,5}] \to \frac{1}{\pi k^2} + \frac{k^2 \pi^3 d^4}{20} + \mathcal{O}(d^6) \ , \ \mathcal{F}[T_{4,7}] \to \frac{1}{\pi k^2} + \frac{k^4 \pi^5 d^6}{252} + \mathcal{O}(d^8)$$

Results: Test Setup

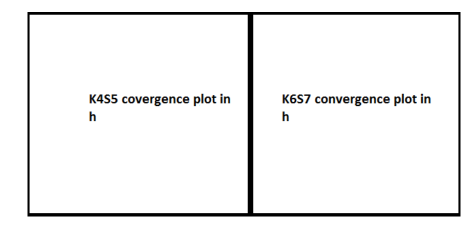
- Theoretical computed convergence rates were verified empirically
- Integral evaluated for 3D Laplace Green's function with constant density in domain $[-0.5, 0.5] \times [-0.5, 0.5] \times [-0.5, 0.5]$
- Possible to compute exact analytical expression for any target in domain
- Domain split into cube elements and tensor product Gauss-Legendre quadrature of varying order was used
- ► Computed result compared with exact result to determine error, and with *h* refinement the order of convergence

Required Minimum Quadrature Order

- \blacktriangleright Accuracy of result dependent on choice of quadrature order q and value chosen for d
- Min required quadrature order to accurately evaluate integral, smaller d less smooth kernel and higher required q
- Past min q, error dominated by truncation error in expansion

Error of result compared to q for a particular d K6S7, d=0.01

Observed Convergence



Quadrature Error Bounds

- Content dependent on how far I get with TV error estimate by Thurs night
- Explanation of bounding quadrature error¹¹in terms of
- Analytic continuability
- and Total Variation (TV)

Rough TV Bound Behavior

▶ Relate bound on required q and d, and behavior wrt h refinement

Observed Error Correctly Bounded by Estimate

► Hopefully show here that observed *quadrature error* (not *total error*) is properly bounded by the TV estimate

Conclusion

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