A NEW ADI METHOD FOR THE POISSON-BOLTZMANN EQUATION WITH $\hbox{A TWO-COMPONENT REGULARIZATION}$

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ABSTRACT

The Poisson Boltzmann equation (PBE) is a well-established implicit solvent continuum model for the electrostatic analysis of solvated biomolecules. The solution for the nonlinear PBE is still a challenge due to its strong singularity by the source terms, dielectrically distinct regions, and exponential nonlinear terms. In this paper, a new alternating direction implicit method (ADI) is proposed for solving the nonlinear PBE using a two-component regularization. This scheme inherits all the advantages of the two-component regularization and the time-dependent PBE with ADI method while possess a novel approach to combine them. A modified version of the one-dimensional ghost fluid method (GFM) has been introduced to incorporate the nonzero jump condition into a new ADI method. The proposed scheme produced better accuracy compared to the previous ADI methods for a benchmark problem and simpler to implement by circumventing the work necessary to apply the MIB method with the regularization for a 3D problem. Though this scheme can use larger time increments than the previous ADI method, it still blows up for large time increments. Later to address this issue with the stability, Locally One Dimensional (LOD) method has been used to replace the ADI method as the operator splitting part. Finally, to test the ability of this newly proposed ADI-GFM and LOD-GFM method, we have evaluated the solvation free energy for a collection of 24 proteins with various sizes and the salt effect on protein-protein binding energy of the complex 1beb.

DEDICATION

INDEX

 $The \ superscript \ ^* \ denotes \ dimensional \ variables.$

ACKNOWLEDGMENTS

CONTENTS

A	BSTRACT	ii
D	DEDICATION	iii
L	IST OF ABBREVIATIONS AND SYMBOLS	iii
A	CKNOWLEDGMENTS	vi
L	IST OF FIGURES	X
1	INTRODUCTION	1
2	PROTEIN DATA FILE PREPARATION	5
3	CLEANING THE OLD ADIMSMS CODING PACKAGE FOR THE NEW SIMULATIONS	7
4	POISSON-BOLTZMANN EQUATION (PBE)	8
	4.1 Two-component regularization for singular sources	9
	4.2 Solvation energy from Electrostatic potential	11
	4.3 Pseudo-transient continuation approach for the PBE	12
5	OPERATOR SPLITTING SCHEMES	14

	5.1	Implicit-Euler method with Alternating Direction Implicit scheme (GFM-ADI)	14					
	5.2	Crank-Nicolson method with Locally One Dimensional Scheme(GFM-LODCN	16					
	5.3	Implicit-Euler method with Locally One Dimensional Scheme(GFM-LODIE)	18					
6	ΑN	NEW DERIVATION FOR GHOST FLUID METHOD	19					
	6.1	One dimension	19					
	6.2	Two dimensions	21					
	6.3	Three dimensions	23					
	6.4	Corner point for the new GFM method	25					
7	NU	MERICAL VALIDATION	27					
		7.0.1 Krikwood Sphere with analytical solution	27					
	7.1 Biological applications							
		7.1.1 Protein Crambin	30					
		7.1.2 Solvation energy of 24 proteins	33					
		7.1.3 Binding energy of 2a9x	34					
		7.1.4 Salt effect on the binding affinity	35					
8	СО	NCLUSIONS	39					
REFERENCES								
Δ	PPE	NDIX	/ 13					

LIST OF FIGURES

6.1	1D GFM	20
6.2	2D GFM	22
7.1	Temporal convergence	30
7.2	Electrostatic potential for 1cbn using $dt = 0.05$ and $h = 0.5$	32
7.3	Difference in solvation energy(left) and CPU time (right) in solving non-linear PB eqation for 24 protein.	34
7.4	Electrostatic potential for 2a9x using GFM-ADI for $\Delta t = 0.05$ and $h = 0.5$	36
7.5	The salt dependence of the binding affinities	38

INTRODUCTION

Solvated biomolecules and the electrostatic interaction with its solvent is critical to study various important biological process such as protein-drug binding site analysis, DNA recognition, protein folding and protein ligand bonding. In the past few decades with the development of numerical methods and computational techniques analyzing the functions and dynamics of bimolecular solvation is more practical and effective. However, imitating these interactions are still computationally expensive with biological significance. Here we are considering the Poisson-Boltzmann Equation (PBE) to describe the electrostatic potential generated by a low dielectric medium inside a protein molecule with embedded atomic charges solvated in a high dielectric medium with dissolved ions. The analytical solution of PBE is only available for some simple geometry such as a sphere or a cylinder. Efficiency and accuracy becomes a critical issue to solve PBE for biophysical models with complex geometry and a macro-molecule containing tens of thousands to millions of atoms (having partial charges).

In our mathematical model for the electrostatic analysis, PBE is a nonlinear elliptic equation on multiple domains with discontinuous dielectric coefficients separated by the solute-solvent interface or molecular surface. The difficulties with solving PBE arises from non-linearity, discontinuous dielectric coefficients, non-smoothness of the solution and singularities in the source term due to the atomic charges. Effects of non-linearity becomes significant with strong ionic presence [32].

For the non-linearity, two different approaches have been developed in the literature. The usual approach is to discretize the nonlinear PBE into an algebraic system using finite difference/finite element methods and then solve it by a nonlinear algebraic system such as (1) nonlinear relaxation method [16], [25], (2) nonlinear conjugate gradient method [23] or (3) inexact Newton method [15]. The other approach has been introduced recently and its based on the pseudo-transient continuation approach [26, 29, 38]. This approach converts time independent nonlinear PBE into a time dependent form by introducing pseudo-time derivative. The solution to the original boundary problem is then retrieved from the steady state solution of the time dependent PBE. The main advantage of introducing pseudo-time derivative is to be able to split time dependent PBE into linear and nonlinear subsystems to circumvent the blow up and overflow problem due to exponentially large term involving hyperbolic sine function.

To convert this three dimensional (3D) PBE into a set of multiple independent one-dimensional (1D) systems, the alternating direction implicit (ADI) methods in [5–7, 10] the locally one dimensional method (LOD) in [32] have been used in the literature. Specially in [10] Douglas-Rachford ADI scheme has been used to split the linear subsystem with the 3D laplacian operator into three more sub systems with one dimensional 2nd order derivatives. Altogether this method remains 2nd order of accuracy in time but conditionally stable. Later in [32] the LOD method appeared as an unconditionally stable method with reduced accuracy compared to ADI methods in [10]. Even though these 1D subsystems produced by these ADI and LOD methods are tridiagonal and can be efficiently solved by using the Thomas Algorithm [30], it lacks the treatment of the jump conditions at the interface which reduces the accuracy near the interface. Also the numerical error for these pseudo transient methods has been observed often to be dominated by the the singularity at the center of the atoms.

Even though the MIB solvers were pretty efficient to overcome the difficulty due to the jump conditions at the interface, the efficient treatment of charge singularity still remained challenging. It motivated many authors to develop different regularization methods in [1,2,9,11,13,33,42] to reduce the loss of accuracy due to the singularity. In these methods, the potential function has decomposed into a singular component plus

one or two other components to break down the PBE into a system of partial differential equations (PDEs) containing a poisson equation with the singular term plus one or two other equations. Thus the singular component can be handled separately using the analytical solution for the poisson equation as Coulomb potentials and or Green's functions. So far these type of regularization methods have never been used with the pseudo transient methods like ADI or LOD.

In an attempt to maintain the efficiency and stability of the ADI methods while restoring the accuracy to the second order near the interface, several interface schemes has been developed in [18–21]. Then as a continuation of this approach recently matched ADI (mADI) method was developed in [41] and [18] to combine the MIB method (for interface treatment) with ADI. But it was mainly focused on the parabolic equations and only 2D condition was considered for the numerical validation.

In this paper our goal was to develop a new approach to solve the PBE combining the regularization, the pseudo-transient continuation and the interface treatment so that both nonlinearity and singularity are treated. For the regularization we have chosen the two component regularization used in [11]. But it changes the jump condition to be non zero which introduces the necessity of interface treatments. Otherwise the standard central difference becomes divergent. Then motivated by the mADI method we have introduced a modified version of the Ghost Fluid Method (GFM) in [4] as the interface treatment. Compared to mADI, GFM is simpler to apply in a pseudo-continuation approach. Altogether GFM-ADI method improved the accuracy and efficiency of the ADI method to solve the non-liner PB equation. Generally it more robust than ADI method but still fails to converge in some special cases. Then to continue the search of a more stable method for our regularized pseudo continuation approach we replaced ADI scheme by LOD method to propose GFM-LODCN and GFM-LODIE method. These other two methods are combining LOD with Crank-Nicolson (CN) and Implicit Euler (IE) to discretize the pseudo time derivative. All of the three methods produced more

accurate results efficiently than their predecessors. GFM-LODIE has found to be most robust while GFM-ADI to be most accurate.

PROTEIN DATA FILE PREPARATION

To generate the molecular surface as the interface in our computational domain we had to use the 3D structural data for proteins and nucleic acids available in the Protein Data Bank(PDB). PDB contains freely accessible data on the internet submitted by biologists and biochemists from around the world. These data are usually acquired by X-ray crystallography, NMR spectroscopy, or, cryo-electron microscopy. Each molecule is represented by 4 letter .Among several types of data file available from the PDB we focused on .pdb type files available from the website of RCSB [24], one of the member organization of the PDB.

Record Type	Columns	Data	Justification	Data Type
	4-Jan	"ATOM"		character
	7-11	Atom serial number	right	integer
	13-16	Atom name	left*	character
	17	Alternate location indicator		character
	18-20§	Residue name	right	character
	22	Chain identifier		character
	23-26	Residue sequence number	right	integer
ATOM	27	Code for insertions of residues		character
	31-38	X orthogonal Å coordinate	right	real (8.3)
	39-46	Y orthogonal Å coordinate	right	real (8.3)
	47-54	Z orthogonal Å coordinate	right	real (8.3)
	55-60	Occupancy	right	real (6.2)
	61-66	Temperature factor	right	real (6.2)
	73-76	Segment identifier¶	left	character
	77-78	Element symbol	right	character
79-80	Charge		character	
HETATM	6-Jan	"HETATM"		character
	Jul-80	same as ATOM records		
	3-Jan	"TER"		character
	7-11	Serial number	right	integer
TER	18-20§	Residue name	right	character
11711	22	Chain identifier		character
	23-26	Residue sequence number	right	integer
	27	Code for insertions of residues		character

Table 2.1: Protein Data file (.pdb type) format

CLEANING THE OLD ADIMSMS CODING PACKAGE FOR THE NEW SIMULATIONS

POISSON-BOLTZMANN EQUATION (PBE)

We are considering the Poisson-Boltzmann Equation (PBE) as the governing equation for a solute macro molecule immersed in an aqueous solvent environment illustrated in Fig 1. Our computational domain $\Omega \in \mathbb{R}^3$ is separated into two regions Ω^- and Ω^+ by the molecular surface Γ , which is an arbitrarily shaped dielectric interface. Here Ω^- is the molecule domain with dielectric constant ϵ^- and Ω^+ is the solvent domain with dielectric constant ϵ^+ . The cubic shape boundary of $\Omega = \Omega^- \cup \Omega^+$ is denoted by $\delta\Omega$. The charges inside Ω^- has been distributed as the partial charges to assign to the nearest grid points to the center of each atom inside the molecule. The charges outside in Ω^+ are mobile ions which are described by the Boltzmann distribution. Then the electrostatic interaction of this solute-solvent system for $\mathbf{r} \in \mathbb{R}^3$ is governed by the nonlinear Poisson-Boltzmann Equation (PBE) as,

$$-\nabla \cdot (\epsilon(\mathbf{r})\nabla\phi(\mathbf{r})) + \bar{\kappa}^2(\mathbf{r})\sinh(\phi(\mathbf{r})) = \rho(\mathbf{r})$$
(4.1)

with the boundary condition,

$$\phi_b(\mathbf{r}) = \frac{e_c^2}{k_B T} \sum_{i=1}^{N_c} \frac{q_i e^{-|\mathbf{r} - \mathbf{r}_i| \sqrt{\frac{\bar{\kappa}^2}{\epsilon^+}}}}{\epsilon^+ |\mathbf{r} - \mathbf{r}_i|}$$
(4.2)

where the singular source $\rho(\mathbf{r})$ term is defined as,

$$\rho(\mathbf{r}) = 4\pi \frac{e_c^2}{k_B T} \sum_{i=1}^{N_c} q_i \delta(\mathbf{r} - \mathbf{r}_i)$$
(4.3)

There are two conditions on Γ needed to be satisfied from the dielectric theory for the potential ϕ and flux density $\epsilon \phi_{\mathbf{n}}$,

$$[\phi]_{\Gamma} = 0 \text{ and } [\epsilon \phi_n]_{\Gamma} = 0$$
 (4.4)

Here $\mathbf{n} = (n_x, n_y, n_z)$ is the outer normal direction on the interface Γ and $\phi_{\mathbf{n}} = \frac{\partial \phi}{\partial \mathbf{n}}$ is the directional derivative in \mathbf{n} . The notation $[f]_{\Gamma} = f^+ - f^-$ represent the difference of the functional value across the interface Γ .

The dielectric constant ϵ is piecewise such that, $\epsilon(\mathbf{r}) = \epsilon^-$ for $\mathbf{r} \in \Omega^-$ and $\epsilon(\mathbf{r}) = \epsilon^+$ for $\mathbf{r} \in \Omega^+$. Here N_c is the total number of atoms in the solute molecule, k_B is the Boltzmann constant, e_c is the fundamental charge and q_i , in the same unit as e_c is the partial charge on the ith atom of the solute molecule located at position \mathbf{r}_j . The Debey-Huckel parameter $\bar{\kappa}^2 = \left(\frac{2N_A e_c^2}{100k_b T}\right) I_s = 8.486902807 \text{Å}^{-2} I_s$ [14] for $\mathbf{r} \in \Omega^-$ and $\bar{\kappa} = 0$ for $\mathbf{r} \in \Omega^+$. Here N_A is Avogadro's Number and I_s is the molar ionic strength. The reader can refer to REF1 and REF2 for more details about definitions and units of these coefficients.

4.1 Two-component regularization for singular sources

To avoid the difficulty due to the source term and the work for solving the Laplace equation we consider a two component regularization proposed by Cai, Wang and Zhao [1]. For this regularization the electrostatic potential ϕ has been considered as the addition of the coulmb component ϕ_c and the reaction field component ϕ_{RF} as $\phi = \phi_c + \phi_{RF}$. Here ϕ_c satisfies the following Poission's equation,

$$\begin{cases}
-\epsilon^{-} \Delta \phi_{C}(r) = \rho(r) \text{ in } \mathbb{R}^{3} \\
\phi_{C}(r) = 0. \text{ as } |r| \to \infty
\end{cases}$$
(4.5)

which gives us the analytical solution as the Green's function G for ϕ_C as,

$$G(r) = \frac{e_c^2}{k_B T} \sum_{i=1}^{N_c} \frac{q_i}{\epsilon^- |r - r_i|}$$
(4.6)

Then Luo and his collaborators [1] proposed to solve for the original solution ϕ in Ω^+ instead of the reaction component ϕ_{RF} . Now to make the required adjustments to fit this regularization approach [1] with finite difference and finite element method Zhao and Geng [11] proposed a new elliptic problem with discontinuous function flux jumps for the two-component regularization. In particular they defined the regularized potential as,

$$\tilde{\phi} = \begin{cases} \phi_{RF} \text{ in } \Omega^{-} \\ \phi_{C} + \phi_{RF} \text{ in } \Omega^{+} \end{cases}$$

$$(4.7)$$

The jump conditions for $\tilde{\phi}$ were derived from ϕ and the definition $\phi = \phi_C + \phi_{RF}$ as,

$$\phi^{+} = \phi_{RF}^{-} + \phi_{C}^{-}$$
, and $\epsilon^{+} \frac{\partial \phi^{+}}{\partial n} = \epsilon^{-} \frac{\partial \phi_{RF}^{-}}{\partial n} + \epsilon^{-} \frac{\partial \phi_{C}^{-}}{\partial n}$ on Γ (4.8)

Thus, the regularized PB equation of $\tilde{\phi}$ with corresponding interface and boundary

conditions are given as.

$$-\nabla \cdot (\epsilon^{-}\nabla \tilde{\phi}) = 0 \text{ in } \Omega^{-}$$

$$\tag{4.9}$$

$$-\nabla \cdot (\epsilon^{+} \nabla \tilde{\phi}) + \bar{\kappa}^{2} \sinh(\tilde{\phi}) = 0 \text{ in } \Omega^{+}$$

$$(4.10)$$

$$\left[\tilde{\phi}\right] = G \text{ on } \Gamma \tag{4.11}$$

$$\left[\epsilon \frac{\partial \tilde{\phi}}{\partial n}\right] = \epsilon^{-} \frac{\partial G}{\partial n} \text{ on } \Gamma$$
 (4.12)

$$\tilde{\phi} = \phi_b \text{ on } \partial\Omega$$
 (4.13)

Now from (4.10) and (4.13) we can summarize that,

$$-\nabla \cdot (\epsilon \nabla \tilde{\phi}) + \bar{\kappa}^2 \sinh(\tilde{\phi}) = 0 \text{ in } \Omega^- \cup \Omega^+$$
(4.14)

where $\tilde{\phi}$ satisfies the PB equation without the source term. Then Zhao and Geng [11] proposed to numerically solve the regularized PB interface problem (RPB) given in (4.10)-(4.13) and to finally recover the original solution as $\phi = \tilde{\phi}$ in Ω^+ and $\phi = \tilde{\phi} + G$ in Ω^- , Where the Green's function G is analytically given.

4.2 Solvation energy from Electrostatic potential

The solvation energy can be defined as the energy released when the solute in free space is dissolved in solvent and it can be expressed in terms of the electrostatic free energy $\Delta G_{\rm ele}$ and the coulomb energy $E_{\rm cou}$ as

$$\Delta G_{\rm ele} = E_{\rm cou} + E_{\rm sol} \tag{4.15}$$

$$E_{\text{cou}} = \sum_{i=1}^{N_c} \sum_{j=1}^{N_c} \frac{q_i q_j}{\epsilon^- d_{i,j}}, i \neq j$$
 (4.16)

where q_i and q_j are the charges at the center of the atoms and $d_{i,j}$'s are the distance between the *i*-th and *j*-the atom.

Then Sharp and Honig in [27] described the calculation electrostatic free energy by,

$$\Delta G_{\text{ele}} = \int_{\mathbb{R}^3} \left(\phi \rho + \Delta \Pi - \frac{1}{2} \epsilon |\mathbf{E}|^2 \right) d\mathbf{r}$$
 (4.17)

where ϕ is the electrostatic potential, ρ is the fixed charge density represented as a smeared surface charge or as a collection of point charges, $\Delta\Pi$ is the excess osmotic pressure of the mobile ion cloud, and $\frac{1}{2}\epsilon|\mathbf{E}|^2$ is the electrostatic stress.

Now for the simplification of the numerical validations of our proposed schemes in the next chapter we omitted the energy components related to the mobile ion pressure and the electrostatic stress to report the solvation energy $E_{\rm sol}$ as,

$$E_{\text{sol}} = \Delta G_{\text{ele}} - E_{\text{cou}} \approx \sum_{i=1}^{N_c} q_i \phi_{RF}(\mathbf{r}_i)$$
(4.18)

Here the contribution of the mobile ion pressure and the electrostatic stress to the whole calculation is really small and computationally more challenging. The readers can refer to [8, 12, 27] for more details.

4.3 Pseudo-transient continuation approach for the PBE

As the pseudo-transient continuation approach, an indirect method discussed in [39] [26] [28] introduced a pseudo-time derivative to solve the PB equation as,

$$\frac{\partial u}{\partial t} = \nabla \cdot (\epsilon \nabla u) - \bar{\kappa}^2 \sinh(u) \text{ in } \Omega^- \cup \Omega^+$$
(4.19)

$$[u] = G$$
, and $\left[\epsilon \frac{\partial u}{\partial n}\right] = \epsilon^{-} \frac{\partial G}{\partial n}$ on Γ (4.20)

$$u = \phi_b \text{ on } \partial\Omega \tag{4.21}$$

Here the time independent regularized PB equation in (4.14) has been converted in to a time dependent regularized PB (TDRPB) in (4.19). Our goal is to first specify the initial condition, which could be the electrostatic potential solved from a linearized PB [39] equation or trivially u = 0 and then numerically integrate (4.19) for a sufficiently long period to get the steady state solution as the solution of the original regularized PB (4.14). Here the sign on the right hand side of eq. (4.19) has been considered as the reverse of the eq.(4.14) to ensure the numerical stability.

But, there are difficulties in the numerical integration of the TDRPB equation (4.19) because of the requirement of long time integration, when explicit time stepping methods are usually not efficient [26], [28], [39], [40]. Hence we are employing a semi-implicit time splitting method [26], [28] which have been commonly used to solve the TDPB in the literature.

OPERATOR SPLITTING SCHEMES

Let us consider a uniform mesh with a grid spacing h in all x, y and z directions having N_x, N_y and N_z as the number of the grid points in each direction. We assume the vector $U^n = \tilde{\phi}^n_{ijk}$ for $i = 1, ..., N_x, j = 1, ..., N_y, k = 1, ..., N_z$ denote all the nodal values of $\tilde{\phi}$ at the time level t_n . We will develop several schemes for updating U^n at time level t_n to U^{n+1} at time level $t_{n+1} = t_n + \Delta t$. In this time dependent RPB we will consider only Uas time dependent, while all other functions in (4.19) i.e. ϵ and $\bar{\kappa}^2$ are time independent.

Implicit-Euler method with Alternating Direction Implicit scheme (GFM-ADI)

In this scheme at each time step from t_n to t_{n+1} , the time dependent equation TDRPB in (4.19) will be solved by a first order time splitting in two stages,

$$\frac{\partial w}{\partial t} = -\bar{\kappa}^2 \sinh(w) \text{ with } W^n = U^n \text{ and } t \in [t_n, t_{n+1}]$$
 (5.1)

$$\frac{\partial w}{\partial t} = -\bar{\kappa}^2 \sinh(w) \text{ with } W^n = U^n \text{ and } t \in [t_n, t_{n+1}]$$

$$\frac{\partial v}{\partial t} = \nabla \cdot (\epsilon \nabla v) \text{ with } V^n = W^{n+1} \text{ and } t \in [t_n, t_{n+1}]$$
(5.1)

Altogether we have $U^{n+1} = V^{n+1}$. And for $\frac{\partial w}{\partial t} = -\bar{\kappa}^2 \sinh(w)$ we have the analytical solution as,

$$W^{n+1} = \ln \left(\frac{\cosh(\frac{1}{2}\bar{\kappa}^2 \Delta t) + \exp(-W^n)\sinh(\frac{1}{2}\bar{\kappa}^2 \Delta t)}{\exp(-W^n)\cosh(\frac{1}{2}\bar{\kappa}^2 \Delta t) + \sinh(\frac{1}{2}\bar{\kappa}^2 \Delta t)} \right)$$
 (5.3)

Which will help us to avoid to the difficulty due to the non-linear term with sinh(.) in (4.1). Here the right hand side of equation (5.3) is just a function of W and Δt which allows us to rewrite the equation (5.3) as $W^{n+1} = F(W^n, \Delta t)$ to facilitate the following discussion. For the temporal discretization of the equation (5.2) we have used Backward-Euler integration in time to get,

$$v_{i,i,k}^{n+1} = v_{i,i,k}^n + \Delta t \left(\delta_x^2 + \delta_y^2 + \delta_z^2 \right) v_{i,i,k}^{n+1}$$
(5.4)

Where δ_x^2, δ_y^2 and δ_z^2 are the central finite difference operators defined in (5.5) for the x,y and z directions, respectively with $\epsilon_{i,j,k} = \begin{cases} \epsilon^- & \text{if } x_{i,j,k} \in \Omega^- \cup \Gamma \\ \epsilon^+ & \text{if } x_{i,j,k} \in \Omega^+ \end{cases}$.

$$\delta_x^2 \left(v_{i,j,k}^n \right) = \frac{\epsilon_{i,j,k}}{h^2} \left(v_{i-1,j,k}^n - 2v_{i,j,k}^n + v_{i+1,j,k}^n \right)
\delta_y^2 \left(v_{i,j,k}^n \right) = \frac{\epsilon_{i,j,k}}{h^2} \left(v_{i,j-1,k}^n - 2v_{i,j,k}^n + v_{i,j+1,k}^n \right)
\delta_z^2 \left(v_{i,j,k}^n \right) = \frac{\epsilon_{i,j,k}}{h^2} \left(v_{i,j,k-1}^n - 2v_{i,j,k}^n + v_{i,j,k+1}^n \right)$$
(5.5)

But at the points near the interface the central difference operators defined in (5.5)-(5.5) will not be applicable. These points are defined as the irregular points where at-least one of its adjacent points is on the other side of the boundary. On the irregular points one of the three point stencils for δ_x^2 , δ_y^2 and δ_z^2 will be on the other side of the interface where the information about the function v is not available. A non-standard finite difference formula is necessary on the irregular points to discretize δ_x^2 , δ_y^2 and δ_z^2 . In this regard a modified version of the Ghost Fluid Method has been proposed in Section ?? using the fictitious points (or ghost points) and the jump conditions in (4.21) rigorously.

Then a first order Douglas-Rachford type ADI scheme has been used to decompose

the diffusion equation (5.4) in x, y and z directions as,

$$\left(1 - \Delta t \delta_x^2\right) v_{i,j,k}^* = \left[1 + \Delta t \left(\delta_y^2 + \delta_z^2\right)\right] v_{i,j,k}^n \tag{5.6}$$

$$\left(1 - \Delta t \delta_y^2\right) v_{i,j,k}^{**} = v_{i,j,k}^* - \Delta t \delta_y^2 \left(v_{i,j,k}^n\right) \tag{5.7}$$

$$(1 - \Delta t \delta_z^2) v_{i,j,k}^{n+1} = v_{i,j,k}^{**} - \Delta t \delta_z^2 (v_{i,j,k}^n)$$
(5.8)

Where v^* and v^{**} are two intermediate values to create three tridiagonal one dimensional system. Here, the three dimensional linear algebraic system in equation (5.4) has been decomposed into several one dimensional linear algebraic systems in (5.6), (5.7) and (5.8). Each one these equations has a tridiagonal structure. These three tridiagonal systems are much more efficient to solve than one non-structured system in (5.4). Then by eliminating $v^*_{i,j,k}$ and $v^{**}_{i,j,k}$ and solving for $v^{n+1}_{i,j,k}$ in (5.4) we get,

$$v_{i,j,k}^{n+1} = v_{i,j,k}^{n} + \Delta t \left(\delta_x^2 + \delta_y^2 + \delta_z^2 \right) v_{i,j,k}^{n+1} - \Delta t^2 \left(\delta_x^2 \delta_y^2 + \delta_x^2 \delta_y^2 + \delta_z^2 \delta_y^2 \right) \left(v_{i,j,k}^{n+1} - v_{i,j,k}^{n} \right)$$

$$+ \Delta t^3 \delta_x^2 \delta_y^2 \delta_z^2 \left(v_{i,j,k}^{n+1} - v_{i,j,k}^{n} \right)$$

$$(5.9)$$

Hence the Douglas-Rachford scheme (5.4) is a higher order perturbation of the Implicit-Euler method. Since both (5.1) and (5.2) are first order in time this proposed ADI-IE-GFM scheme is of first order accuracy in time. As the boundary conditions the same Dirichlet boundary boundary values are assumed for v, v^* and v^{**} for u. The entire time integration here is fully implicit.

5.2 Crank-Nicolson method with Locally One Dimensional Scheme(GFM-LODCN)

In this scheme at each time step from t_n to t_{n+1} , the time dependent non-linear equation TDPBE will be solved by a second order time splitting methods in three stages

[36],

$$\frac{\partial w}{\partial t} = -\frac{1}{2}\bar{\kappa}^2 \sinh(w) \text{ with } W^n = U^n \text{ and } t \in [t_n, t_{n+1}]$$
 (5.10)

$$\frac{\partial v}{\partial t} = \nabla \cdot (\epsilon \nabla v) \text{ with } V^n = W^{n+1} \text{ and } t \in [t_n, t_{n+1}]$$
(5.11)

$$\frac{\partial \tilde{w}}{\partial t} = -\frac{1}{2}\bar{\kappa}^2 \sinh(\tilde{w}) \text{ with } \tilde{W}^n = V^{n+1} \text{ and } t \in [t_n, t_{n+1}]$$
 (5.12)

We then have $U^{n+1} = \tilde{W}^{n+1}$. As in the subsection 5.1 we will use an analytical integration in the first and the last stage of this scheme. Symbolically, we have $W^{n+1} = F(W^n, \frac{\Delta t}{2})$ and $\tilde{W}^{n+1} = F(\tilde{W}^n, \frac{\Delta t}{2})$, where F is defined as in Eq.(5.3).

Then we have proposed another multiplicative operator splitting scheme called as Locally One Dimensional (LOD) scheme to solve the diffusion equation (5.11). This types of fractional step methods were first developed by Russian mathematicians in [3], [34], [35]. The discretization of (5.11) using Crank-Nicolson integration in time and central differencing in space results in,

$$\left(1 - \frac{\Delta t}{2}(\delta_x^2 + \delta_y^2 + \delta_z^2)\right)v_{i,j,k}^{n+1} = \left(1 + \frac{\Delta t}{2}(\delta_x^2 + \delta_y^2 + \delta_z^2)\right)v_{i,j,k}^n$$
(5.13)

Which can be decomposed in to x, y and z directions to give a LOD scheme for (5.11),

$$\left(1 - \frac{\Delta t}{2} \delta_x^2\right) v_{i,j,k}^* = \left(1 + \frac{\Delta t}{2} \delta_x^2\right) v_{i,j,k}^n$$

$$\left(1 - \frac{\Delta t}{2} \delta_y^2\right) v_{i,j,k}^{**} = \left(1 + \frac{\Delta t}{2} \delta_y^2\right) v_{i,j,k}^*$$

$$\left(1 - \frac{\Delta t}{2} \delta_z^2\right) v_{i,j,k}^{n+1} = \left(1 + \frac{\Delta t}{2} \delta_z^2\right) v_{i,j,k}^{**}$$
(5.14)

Similarly as the GFM-ADI scheme described in subsection 5.1 the tridiagonal systems in (5.14) can be efficiently solved by the Thomas algorithm.

5.3 Implicit-Euler method with Locally One Dimensional Scheme(GFM-LODIE)

For this scheme at each time step from t_n to t_{n+1} , the time dependent equation TDRPB in (4.19) will be solved by the first order time splitting similar to the GFM-ADI scheme described in the subsection 5.1 in two stages. To follow the same steps in the first stage Eq. (5.1) will be solved analytically by the function $W^{n+1} = F(W^n, \Delta t)$ in Eq. (5.3) and in the second stage for the temporal discretization of the Eq. (5.2) we have used Implicit Euler method to get Eq.(5.4).

Then we have applied the LOD scheme described in the subsection 5.2 to decompose Eq. (5.4) in to x, y and z direction as,

$$(1 - \Delta t \delta_x^2) v_{i,j,k}^* = v_{i,j,k}^n$$

$$(1 - \Delta t \delta_y^2) v_{i,j,k}^{**} = v_{i,j,k}^*$$

$$(1 - \Delta t \delta_z^2) v_{i,j,k}^{n+1} = v_{i,j,k}^{**}$$
(5.15)

Which will solved individually by the Thomas Algorithm like the previous subsections.

A NEW DERIVATION FOR GHOST FLUID METHOD

Ghost Fluid Method(GFM) is a sharpe interface technique introduced in [4] to treat the two-face contact discontinuities in the Euler equations. It extends values across the interface into an artificial fluid (ghost fluid), inducing the jump conditions at the interface. This GFM method was later extended in [22] to solve elliptic equations with variable coefficients. But in contrast to their method [22], the jump conditions are incorporated into the numerical discretization in such a way that the symmetry of the finite difference stencil is preserved. Which makes it compatible with most standard solvers. The flux jump has been decomposed in each axis direction treating the problem dimension by dimension. As a result this extended GFM becomes only first order accurate.

6.1 One dimension

In order to overcome the difficulty due to the jump-condition on the interface Γ , a modified version of Ghost Fluid Method(GFM) has been used. For a simple 1D model when we try to evaluate by the finite difference operator δ_{xx} at irregular points where the interface is at x_{Γ} and between x_i and x_{i+1} as the following figure.

Here u is the unknown function defined as solution of the PBE as $u = \begin{cases} u_{in} \text{ in } \Omega^- \\ u_{out} \text{ in } \Omega^+ \end{cases}$

Then we define $u^+ = u_{out}(x_{\Gamma}), u^- = u_{in}(x_{\Gamma}), h = x_{i+1} - x_i$ and $\lambda = \frac{x_{\Gamma} - x_i}{h}$ to get following relations,

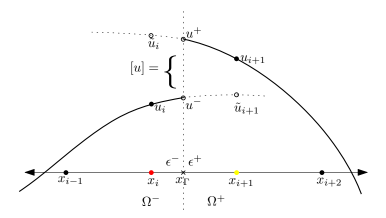


Figure 6.1: 1D GFM

$$x_{\Gamma} = x_i + \lambda h$$

$$u^- = u_i (1 - \lambda) + \tilde{u}_{i+1} \lambda \text{ and } u^+ = \tilde{u}_i (1 - \lambda) + u_{i+1} \lambda$$

$$[u] = u^+ - u^- \text{ and } \left[\epsilon \frac{\partial u}{\partial x} \right] = \epsilon^+ u_x^+ - \epsilon^- u_x^-$$

$$(6.1)$$

Where $\tilde{u_i}$ and $\tilde{u_{i+1}}$ are the fictitious values(or ghost values) of u at the irregular points extending u_{in} or u_{out} on the other side of the interface. In particular for figure 6.1, we have $\tilde{u}_i = u_{out}(x_i)$ and $\tilde{u}_{i+1} = u_{in}(x_{i+1})$. Here u_{out} has been extended to the point x_i for \tilde{u}_i and u_{in} has been extended to the point x_{i+1} for \tilde{u}_i .

Our purpose here is to apply the non-standard finite difference operator δ_x^2 at x_i and x_{i+1} to get,

$$\delta_x^2(u_i) = \frac{\epsilon^-}{h^2} (u_{i-1} - 2u_i + \tilde{u}_{i+1}) \text{ and } \delta_x^2(u_{i+1}) = \frac{\epsilon^+}{h^2} (\tilde{u}_i - 2u_{i+1} + u_{i+2})$$
 (6.2)

Now solving above equations in (6.1) for \tilde{u}_i and \tilde{u}_{i+1} we get,

$$\tilde{u}_{i} = \frac{\epsilon^{-}}{\epsilon^{+}\lambda + \epsilon^{-}(1-\lambda)} u_{i} + \frac{\lambda(\epsilon^{+} - \epsilon^{-})}{\epsilon^{+}\lambda + \epsilon^{-}(1-\lambda)} u_{i+1}
+ \frac{\epsilon^{-}}{\epsilon^{+}\lambda + \epsilon^{-}(1-\lambda)} [u] - \frac{h\lambda}{\epsilon^{+}\lambda + \epsilon^{-}(1-\lambda)} \left[\epsilon \frac{\partial u}{\partial x} \right]
\tilde{u}_{i+1} = \frac{(\epsilon^{-} - \epsilon^{+})(1-\lambda)}{\epsilon^{+}\lambda + \epsilon^{-}(1-\lambda)} u_{i} + \frac{\epsilon^{+}}{\epsilon^{+}\lambda + \epsilon^{-}(1-\lambda)} u_{i+1}
- \frac{\epsilon^{+}}{\epsilon^{+}\lambda + \epsilon^{-}(1-\lambda)} [u] - \frac{h(1-\lambda)}{\epsilon^{+}\lambda + \epsilon^{-}(1-\lambda)} \left[\epsilon \frac{\partial u}{\partial x} \right]$$
(6.3)

Then substituting equation (6.3) into equation (6.2) we get,

$$\delta_x^2(u_i) = \frac{1}{h^2} \left(a_1 u_{i-1} + b_1 u_i + c_1 u_{i+1} \right) + \frac{\epsilon^-}{h^2} \left(e_1 \cdot [u] + f_1 \cdot \left[\epsilon \frac{\partial u}{\partial x} \right] \right)$$

$$\delta_x^2(u_{i+1}) = \frac{1}{h^2} \left(a_2 u_i - b_2 u_{i+1} + c_2 u_{i+2} \right) + \frac{\epsilon^+}{h^2} \left(e_2 \cdot [u] + f_2 \cdot \left[\epsilon \frac{\partial u}{\partial x} \right] \right)$$
(6.4)

Where,

$$d = \epsilon^{+}\lambda + \epsilon^{-}(1 - \lambda)$$

$$a_{1} = \epsilon^{-}, b_{1} = -\epsilon^{-}\left(1 + \frac{\epsilon^{+}}{d}\right), c_{1} = \frac{\epsilon^{-}\epsilon^{+}}{d}, e_{1} = \frac{\epsilon^{-}}{d}, f_{1} = \frac{h\lambda}{d}$$

$$a_{2} = \frac{\epsilon^{+}\epsilon^{-}}{d}, b_{2} = -\epsilon^{+}\left(1 + \frac{\epsilon^{-}}{d}\right), c_{2} = \epsilon^{+}, e_{2} = -\frac{\epsilon^{+}}{d}, f_{2} = \frac{h(1 - \lambda)}{d}$$

$$(6.5)$$

Here the 2nd terms of the equations in (6.4) are considered to be known. Only the coefficients in the 1st terms of (6.4) contribute to the finite difference operator matrix to make it diagonally dominant(as $|b_1| - a_1 - c_1 = 0$ and $|b_2| - a_2 - c_2 = 0$) and symmetric (as $a_2 = c_1$).

6.2 Two dimensions

For two dimensional PB model we need to evaluate δ_{xx} and δ_{yy} , where the similar derivation of equations in (6.4) can be used to calculate δ_{yy} in irregular points. But in

this case, $\left[\epsilon \frac{\partial u}{\partial x}\right]$ and $\left[\epsilon \frac{\partial u}{\partial y}\right]$ is not known, while $\left[\epsilon \frac{\partial u}{\partial n}\right]$ is given. Now to get $\left[\epsilon \frac{\partial u}{\partial x}\right]$ and $\left[\epsilon \frac{\partial u}{\partial y}\right]$ in terms of $\left[\epsilon \frac{\partial u}{\partial n}\right]$ and $\left[\epsilon \frac{\partial u}{\partial \tau}\right]$ we have the following relations,

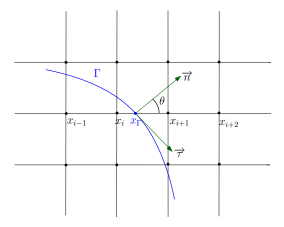


Figure 6.2: 2D GFM

$$\begin{bmatrix} \epsilon \frac{\partial u}{\partial n} \end{bmatrix} = \cos \theta \left[\epsilon \frac{\partial u}{\partial x} \right] + \sin \theta \left[\epsilon \frac{\partial u}{\partial y} \right]
\left[\epsilon \frac{\partial u}{\partial \tau} \right] = \sin \theta \left[\epsilon \frac{\partial u}{\partial x} \right] - \cos \theta \left[\epsilon \frac{\partial u}{\partial y} \right]$$
(6.6)

Which can be solved for $\left[\epsilon \frac{\partial u}{\partial x}\right]$ and $\left[\epsilon \frac{\partial u}{\partial y}\right]$ as,

$$\begin{bmatrix} \epsilon \frac{\partial u}{\partial x} \end{bmatrix} = \cos \theta \left[\epsilon \frac{\partial u}{\partial n} \right] + \sin \theta \left[\epsilon \frac{\partial u}{\partial \tau} \right]
\begin{bmatrix} \epsilon \frac{\partial u}{\partial y} \end{bmatrix} = \sin \theta \left[\epsilon \frac{\partial u}{\partial n} \right] - \cos \theta \left[\epsilon \frac{\partial u}{\partial \tau} \right]$$
(6.7)

To simplify the jump-conditions Liu, Fedkiw and Kang [22] smeared out the tangential derivative considering $\left[\epsilon \frac{\partial u}{\partial \tau}\right] = 0$ to get the following equations,

$$\begin{bmatrix} \epsilon \frac{\partial u}{\partial x} \end{bmatrix} = \cos \theta \left[\epsilon \frac{\partial u}{\partial n} \right] \\
\left[\epsilon \frac{\partial u}{\partial y} \right] = \sin \theta \left[\epsilon \frac{\partial u}{\partial n} \right]$$
(6.8)

Which are in general not true but $\left[\epsilon \frac{\partial u}{\partial x}\right]$ and $\left[\epsilon \frac{\partial u}{\partial y}\right]$ in (6.8) and (6.8) satisfies (6.6) for $\left[\epsilon \frac{\partial u}{\partial n}\right]$. It allows us to replace the unknown quantities as $\left[\epsilon \frac{\partial u}{\partial x}\right]$ and $\left[\epsilon \frac{\partial u}{\partial y}\right]$ by a known quantity $\left[\epsilon \frac{\partial u}{\partial n}\right]$ and $\left[\epsilon \frac{\partial u}{\partial \tau}\right]$. The process used to simplify the jump condition in [22] still requires the normal direction θ . In our proposed GFM scheme we have considered $\frac{\partial u^+}{\partial \tau} = 0$ to get the following derivation for $\left[\epsilon \frac{\partial u}{\partial x}\right]$ as,

$$\begin{split} \left[\epsilon \frac{\partial u}{\partial x}\right] &= \cos \theta \left[\epsilon \frac{\partial u}{\partial n}\right] + \sin \theta \left[\epsilon \frac{\partial u}{\partial \tau}\right] \\ &= \cos \theta \left[\epsilon \frac{\partial u}{\partial n}\right] + \sin \theta \left(\epsilon^{+} \frac{\partial u^{+}}{\partial \tau} - \epsilon^{-} \frac{\partial u^{-}}{\partial \tau}\right) \\ &= \cos \theta \left[\epsilon \frac{\partial u}{\partial n}\right] + \sin \theta \left(\epsilon^{+} \frac{\partial u^{+}}{\partial \tau} - \epsilon^{-} \frac{\partial u^{+}}{\partial \tau} + \epsilon^{-} \frac{\partial u^{+}}{\partial \tau} - \epsilon^{-} \frac{\partial u^{-}}{\partial \tau}\right) \\ &= \cos \theta \left[\epsilon \frac{\partial u}{\partial n}\right] + \sin \theta \left(\epsilon^{-} \left(\frac{\partial u^{+}}{\partial \tau} - \frac{\partial u^{-}}{\partial \tau}\right) + (\epsilon^{+} - \epsilon^{-}) \frac{\partial u^{+}}{\partial \tau}\right) \\ &= \cos \theta \left[\epsilon \frac{\partial u}{\partial n}\right] + \sin \theta \left(\epsilon^{-} \left[\frac{\partial u}{\partial \tau}\right] + (\epsilon^{+} - \epsilon^{-}) \frac{\partial u^{+}}{\partial \tau}\right) \end{split}$$

Now by the jump conditions we have $\left[\epsilon \frac{\partial u}{\partial n}\right] = \epsilon^{-} \frac{\partial G}{\partial n}$ and $\left[\frac{\partial u}{\partial \tau}\right] = \frac{\partial G}{\partial \tau}$.

Thus
$$\left[\epsilon \frac{\partial u}{\partial x}\right] = \cos \theta (\epsilon^{-} \frac{\partial G}{\partial n}) + \sin \theta . \epsilon^{-} \frac{\partial G}{\partial \tau} + \sin \theta (\epsilon^{+} - \epsilon^{-}) \frac{\partial u^{+}}{\partial \tau}$$

 $\approx \epsilon^{-} (\cos \theta \frac{\partial G}{\partial n} + \sin \theta \frac{\partial G}{\partial \tau}) = \epsilon^{-} \frac{\partial G}{\partial x}$, consedering $\frac{\partial u^{+}}{\partial \tau} = 0$
 $\therefore \left[\epsilon \frac{\partial u}{\partial x}\right] \approx \epsilon^{-} \frac{\partial G}{\partial x}$

Similarly it can be shown that $\left[\epsilon \frac{\partial u}{\partial y}\right] \approx \epsilon^{-} \frac{\partial G}{\partial y}$.

6.3 Three dimensions

Consider that the interface Γ intersects the grid line in the x direction at a point (i_{Γ}, j, k) which is located between (i, j, k) and (i+1, j, k), We therefore have two irregular grid points, (i, j, k) and (i+1, j, k). The fictitious values $\tilde{u}_{i,j,k}$ and $\tilde{u}_{i+1,j,k}$ are to be

determined. To use one of the jump conditions which is defined in the normal direction of the interface point, it is convenient to introduce a local coordinates (ξ, η, ζ) such that ξ is along the normal direction and η is in the xy plane. Then the coordinate transformation can be given as,

$$\begin{bmatrix} x \\ y \\ z \end{bmatrix} = \mathbf{P} \begin{bmatrix} \xi \\ \eta \\ \zeta \end{bmatrix} \tag{6.9}$$

where \mathbf{P} is the transformation matrix

$$\mathbf{P} = \begin{bmatrix} \sin \psi \cos \theta & -\sin \theta & -\cos \psi \cos \theta \\ \sin \psi \sin \theta & \cos \theta & -\cos \psi \sin \theta \\ \cos \psi & 0 & \sin \psi \end{bmatrix}. \tag{6.10}$$

Here θ and ψ are the azimuth and zenith angles with respect to the normal direction ξ . Then from equation (6.9) and (6.10) we have,

$$\left[\epsilon \frac{\partial u}{\partial x}\right] = \sin \psi \cos \theta \left[\epsilon \frac{\partial u}{\partial \xi}\right] - \sin \theta \left[\epsilon \frac{\partial u}{\partial \eta}\right] - \cos \psi \cos \theta \left[\epsilon \frac{\partial u}{\partial \zeta}\right]$$
(6.11)

Now along η axis,

$$\begin{bmatrix} \epsilon \frac{\partial u}{\partial \eta} \end{bmatrix} = \epsilon^{+} \frac{\partial u^{+}}{\partial \eta} - \epsilon^{-} \frac{\partial u^{-}}{\partial \eta}
= \epsilon^{+} \frac{\partial u^{+}}{\partial \eta} - \epsilon^{-} \frac{\partial u^{+}}{\partial \eta} + \epsilon^{-} \frac{\partial u^{+}}{\partial \eta} - \epsilon^{-} \frac{\partial u^{-}}{\partial \eta}
= \epsilon^{-} \left(\frac{\partial u^{+}}{\partial \eta} - \frac{\partial u^{-}}{\partial \eta} \right) + (\epsilon^{+} - \epsilon^{-}) \frac{\partial u^{+}}{\partial \eta}
= \epsilon^{-} \frac{\partial G}{\partial \eta} + (\epsilon^{+} - \epsilon^{-}) \frac{\partial u^{+}}{\partial \eta}. \text{ since } \left(\frac{\partial u^{+}}{\partial \eta} - \frac{\partial u^{-}}{\partial \eta} \right) = \frac{\partial}{\partial \eta} [u] = \frac{\partial G}{\partial \eta}.$$
(6.12)

Similarly along ζ axis,

$$\left[\epsilon \frac{\partial u}{\partial \zeta}\right] = \epsilon^{-} \frac{\partial G}{\partial \zeta} + (\epsilon^{+} - \epsilon^{-}) \frac{\partial u^{+}}{\partial \zeta}$$
(6.13)

Then from equation (6.11), (6.12) and (6.13)we have,

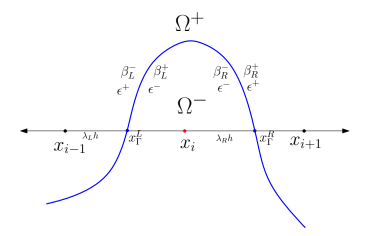
$$\left[\epsilon \frac{\partial u}{\partial x}\right] = \epsilon^{-} \left(\sin \psi \cos \theta \frac{\partial G}{\partial \xi} - \sin \theta \frac{\partial G}{\partial \eta} - \cos \psi \cos \theta \frac{\partial G}{\partial \zeta}\right)
- \sin \theta (\epsilon^{+} - \epsilon^{-}) \frac{\partial u^{+}}{\partial \eta} - \cos \phi \cos \theta (\epsilon^{+} - \epsilon^{-}) \frac{\partial u^{+}}{\partial \zeta}
= \epsilon^{-} \frac{\partial G}{\partial x} - \sin \theta (\epsilon^{+} - \epsilon^{-}) \frac{\partial u^{+}}{\partial \eta} - \cos \phi \cos \theta (\epsilon^{+} - \epsilon^{-}) \frac{\partial u^{+}}{\partial \zeta}
\approx \epsilon^{-} \frac{\partial G}{\partial x} \text{ considering } \frac{\partial u^{+}}{\partial \eta} = 0 \text{ and } \frac{\partial u^{+}}{\partial \zeta} = 0.$$
(6.14)

Similarly along x axis and y axis we have, Then from equation (6.11), (6.12) and (6.13)we have,

$$\left[\epsilon \frac{\partial u}{\partial y}\right] \approx \epsilon^{-} \frac{\partial G}{\partial y} \text{ and } \left[\epsilon \frac{\partial u}{\partial z}\right] \approx \epsilon^{-} \frac{\partial G}{\partial z}$$
 (6.15)

6.4 Corner point for the new GFM method

We have a special situation for a corner point when the interface crosses the grid line for the same axis twice around x_i where x_{i-1} and x_{i+1} are on the other side of the interface.



Corner point case for the new GFM with $G_L = -G$ and $G_R = G$. Which gives the following equations for the fictitious points \tilde{u}_{i-1} and \tilde{u}_{i+1} .

$$\tilde{u}_{i-1} = A_1 u_{i-1} + B_1 u_i + C_1 G_L - D_1 \epsilon^{-1} \frac{\partial G_L}{\partial x}$$
 (6.16)

$$\tilde{u}_{i+1} = A_2 u_{i+1} + B_2 u_i - C_2 G_R - D_2 \epsilon^{-\frac{\partial G_R}{\partial x}}$$
(6.17)

Where,

$$\beta_L^- = \epsilon^+, \beta_L^+ = \epsilon^-, \beta_R^- = \epsilon^-, \beta_R^+ = \epsilon^+$$

$$\lambda_L = \frac{x_i - x_{i-1}}{h}, \lambda_R = \frac{x_{i+1} - x_i}{h}$$

$$F_1 = \beta_L^+ \lambda_L + \beta_L^- (1 - \lambda_L), F_2 = \beta_R^+ \lambda_R + \beta_R^- (1 - \lambda_R)$$

$$A_1 = \frac{\beta_L^-}{F_1}, B_1 = \frac{\lambda_L (\beta_L^+ - \beta_L^-)}{F_1}, C_1 = \frac{\beta_L^-}{F_1}, D_1 = \frac{h\lambda_L}{F_1}$$

$$A_2 = \frac{\beta_R^+}{F_2}, B_2 = \frac{(1 - \lambda_R)(\beta_R^- - \beta_R^+)}{F_2}, C_2 = \frac{\beta_R^+}{F_2}, D_2 = \frac{h(1 - \lambda_R)}{F_2}$$

CHAPTER 7

NUMERICAL VALIDATION

To validate our proposed algorithm we are providing numerical simulations in this chapter. At first we have solved the nonlinear PB equation on a krikwood sphere and compared the numerical results with the analytical solution. Then we have considered a hypothetical protein molecule with just one atom for the PB equation and solved it to compare with the analytical result.

7.0.1 Krikwood Sphere with analytical solution

The analytical solutions of the PB equation are available only for simple geometries such as spheres. So we choose the case where we have a charge at the center of a sphere known as the krikwood sphere with the analytical solution based on the reference [10].

$$\phi(\mathbf{r}) = \begin{cases} \frac{1}{\varepsilon R} - \frac{1}{R} + \frac{1}{||\mathbf{r}||} & ||\mathbf{r}|| < R, \\ \frac{1}{\varepsilon ||\mathbf{r}||} & ||\mathbf{r}|| > R. \end{cases}$$

$$\rho(\mathbf{r}) = \begin{cases} 4\pi \epsilon^{-} \delta(\mathbf{r}) & ||\mathbf{r}|| < R, \\ \bar{\kappa}^{2} \sinh(\frac{1}{\varepsilon ||\mathbf{r}||}) & ||\mathbf{r}|| > R, \end{cases}$$

$$(7.1)$$

$$\rho(\mathbf{r}) = \begin{cases} 4\pi \epsilon^{-} \delta(\mathbf{r}) & ||\mathbf{r}|| < R, \\ \bar{\kappa}^{2} \sinh(\frac{1}{\varepsilon ||\mathbf{r}||}) & ||\mathbf{r}|| > R, \end{cases}$$
 (7.2)

where $\varepsilon = \epsilon^+/\epsilon^-$ and R = 2 Å is the radius of the sphere and $\bar{\kappa} = 1$. We have a unit charge $1e_c$ located at the center of the sphere. Our dielectric constants are $\epsilon^+=80$ and $\epsilon^- = 1$ It can be shown that this analytical solution in (7.1) with the source term defined in (7.2) will satisfy the non-linear PB equation in (4.1) together with the jump

condition defined in (4.4). Both the singularity and the non-smoothness are present in this analytical solution. The singularity is from the source term in (7.2)for $||\mathbf{r}|| > R$ and the non-smoothness comes from the jump condition (4.4). Both of these difficulties gives similar kind of challenge present in the non-linear PB equation to reduce the accuracy of the spatial discretization numerically. To use this benchmark problem to test the stability and the convergence in time and space we computed the L_2 error and L_{∞} error using the following measures.

$$L_{\infty} = \max_{i,j,k} |\phi_{\text{exact}} - \phi_{\text{num}}|, L_2 = \sqrt{\frac{\sum_{i,j,k} |\phi_{\text{exact}} - \phi_{\text{num}}|^2}{N}}$$

Where ϕ_{true} is the analytical solution and ϕ_{num} is the numerical solution representing the electrostatic potential for the non-linear PB equation. For the L_2 error we have used $N = N_x \times N_y \times N_z$ as the total number of unknowns on the grid points.

Stability test: At first for the stability for the non-linear PB equation to calculate the potential $\phi(\mathbf{r})$ for the krikwood sphere we considered the domain to be [-3,3] for x,y and z direction with the spherical radius to be R=1 for the interface. In this study we used finer grid as h=0.125 to avoid the difficulty due to the larger grid spacing and focused on the effect on the stability due to the changes in time increment Δt at each time step. We have found all three of our methods to be stable for $\Delta t = [0.001, 5]$. To illustrate this we considered the sampling for Δt as $\Delta t = \{0.001, 0.002, 0.005, 0.01, 0.02, 0.05, 0.1, 0.2, 0.5, 1, 2, 5\}$ and the stopping time T as T=100 so that enough accumulations are experienced. We observed both L_2 and L_∞ error to be finite. For larger values of Δt as $\Delta t=5$ the numerical errors might feel meaningless but as long as this errors remains to be finite, this demonstrates the stability of the underlying time integration.

Spacial Convergence: In this study, we investigated the order of accuracy for the spacial convergence in Table 7.1 for the krikwood sphere. The time increment Δt

\overline{h}	L_2	Order	L_{∞}	Order	$E_{\rm sol}$			
ADI								
2	6.45E-03	N/A	3.82E-02	N/A	-92.699927			
1	4.88E-03	0.40	7.61E-02	-0.99	-83.683388			
1/2	3.43E-02	-2.81	$1.30\mathrm{E}{+00}$	-4.09	-85.921222			
1/4	4.63E-02	-0.43	$3.44\mathrm{E}{+00}$	-1.41	-83.279725			
1/8	5.11E-02	-0.14	$7.49\mathrm{E}{+00}$	-1.12	-82.680633			
		(GFM-ADI					
2	3.14E-04	N/A	1.82E-03	N/A	-81.742795			
1	1.18E-04	1.41	8.28E-04	1.13	-82.132181			
1/2	2.79E-05	2.08	3.10E-04	1.42	-82.063724			
1/4	8.51E-06	1.71	1.23E-04	1.34	-82.051117			
1/8	1.49E-06	2.51	4.47E-05	1.46	-82.046462			
		GF	M-LODCN					
2	3.14E-04	N/A	1.82E-03	N/A	-81.742788			
1	1.18E-04	1.41	8.89E-04	1.03	-82.132148			
1/2	2.87E-05	2.04	3.87E-04	1.20	-82.063684			
1/4	1.10E-05	1.39	2.13E-04	0.86	-82.051064			
1/8	7.16E-06	0.62	1.41E-04	0.60	-82.046402			
	GFM-LODIE							
2	3.16E-04	N/A	1.84E-03	N/A	-81.738399			
1	1.17E-04	1.43	8.85E-04	1.06	-82.123319			
1/2	2.84E-05	2.05	3.89E-04	1.19	-82.055388			
1/4	1.18E-05	1.27	2.18E-04	0.83	-82.043011			
1/8	9.40E-06	0.33	1.48E-04	0.56	-82.046402			

Table 7.1: Solving the nonlinear PB equation for the krikwood sphere with $\epsilon^+ = 80$, $\epsilon^- = 1$, $\Delta t = 0.001$, T = 10, $I_s = 0.01$ and $\kappa = 1$. The centered charge of unit $1e_c$ is located at (0, 0, 0).

has been kept fixed to 0.001 while reducing the grid spacing h from 2 to 1/8 in this process. Within this range of h we have noticed the accuracy of GFM-ADI to be nearly 2nd order while gradually reducing for GFM-LODCN and GFM-LODIE. Then we have computed the solvation energy $E_{\rm sol}$ using the non-linear PB equation in (4.1) with the source term defined in (4.3) for the similar setup for the krikwood sphere in 7.1. The solvation energy for this setup can also be computed analytically as -81.97820845. For all three of our proposed methods the solvation energies $E_{\rm sol}$ found to be very close to this analytical value as reported in Table 7.1.

Temporal Convergence:

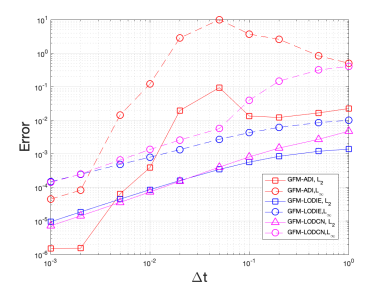


Figure 7.1: Temporal convergence

7.1 Biological applications

In this section, we focus on exploring the stability and accuracy of GFM-ADI, GFM-LODCN and GFM-LODIE schemes by considering the solvation analysis of real proteins. Even though all of our proposed schemes were found to be stable for the one atom case (Krikwood sphere) for $\Delta t = [0.001, 5]$, it is of great interest to see if these schemes are stable for real protein systems. We will compare all three methods in detail for a particular protein system and try to identify the best choice for the time increment Δt and the grid spacing h to get the most accurate result within a reasonable amount of time. The optimum Δt and h has been used later to calculate the solvation energy of 24 proteins.

7.1.1 Protein Crambin

To validate our proposed schemes with a real protein we have studied the hydrophobic protein Crambin (PDB ID : 1cbn). It is a 46 residue protein homologous to a membrane-active plant toxins [31]. It is found in the seeds of *Crambe abyssinicia* and have local anesthetic activity in a lobster walking leg axon (J. Marquis [31]). We used the MSMS package to generate the molecular surface for this protein using the crystallo-

graphic data recored at 130K as reported in [31]. At this step we used the probe radius as 1.4 the density as 10 for the MSMS package.

Δt	ADI	GFM-ADI	GFM-LODCN	GFM-LODIE
0.001	-459.5742719854	-303.00657886	-303.00154088	-302.80556740
0.002	-458.1104685049	-302.99808443	-302.98279503	-302.61375402
0.005	-452.1116748705	-302.90138868	-302.83729044	-302.03341621
0.01	NaN	-302.85075766	-302.68957418	-301.31096348
0.02	NaN	-302.69227062	-302.33466174	-300.06953533
0.05	NaN	-302.04538862	-301.08386563	-297.04896758
0.1	NaN	-300.84648128	-298.86787256	-293.05472908
0.2	NaN	-298.63624081	-294.81110682	-286.88648904
0.5	NaN	-293.21633440	-284.83429816	-274.64462149
0.7	NaN	NaN	-279.04495504	-269.01031327
1	NaN	NaN	-271.29161057	-262.50478312
2	NaN	NaN	-251.54655749	-248.89416804
5	NaN	NaN	-218.29937773	-230.91124478

Table 7.2: Solvation Energy (kcal/mol) of 1cbn for h=0.5 and the ionic strength $I_s=0.15$

For this study at first we reported the solvation energy of 1cbn calculated by all three of our proposed schemes in Table 7.2. After calculating electrostatic potential ϕ , equation (4.18) has been discretized further to calculate the solvation energy as,

$$E_{\text{sol}} = \sum_{i} \sum_{j} \sum_{k} Q(x_i, y_j, z_k) \phi_{RF}(x_i, y_j, z_k)$$
 (7.3)

where Q is the trilinear interpolation of the singular charges q_i at the center of the atoms. The potential values are obtained by scaling our calculated dimensionless potentials with the constant 0.596163438 corresponding for the room temperature (300K). In all cases a uniform mesh size h=0.5 and a large stopping time T will be used to ensure that the steady state solution is reached. For the dielectric constant we have used $\epsilon^+=80$ for water as the solvent and $\epsilon^-=1$ the region in side the protein. The ionic strength has been used as $I_s=0.15$.

Here we tried to identify the value of Δt as large as possible without losing

the adequate amount accuracy for a real protein like 1cbn. But in Table 7.2 as we have observed for $\Delta t > 0.5$, GFM-ADI diverges totally while other two schemes loses significant amount of accuracy. If we consider the solvation energy (around -302 kcal/mol) for $\Delta t = 0.005$ as the most accurate one and compare all other solvation energies in Table 7.2, it can be observed that with the increase of Δt all three proposed schemes loses accuracy but at a different rate. GFM-ADI scheme is usually more accurate while GFM-LODIE schemes more robust to the larger values of Δt . The performance of GFM-LODCN is roughly in between the other two schemes in terms of accuracy and stability. To be uniform with the studies for the other proteins in this paper we have used the optimum value for $\Delta t = 0.05$ and h = 0.5.

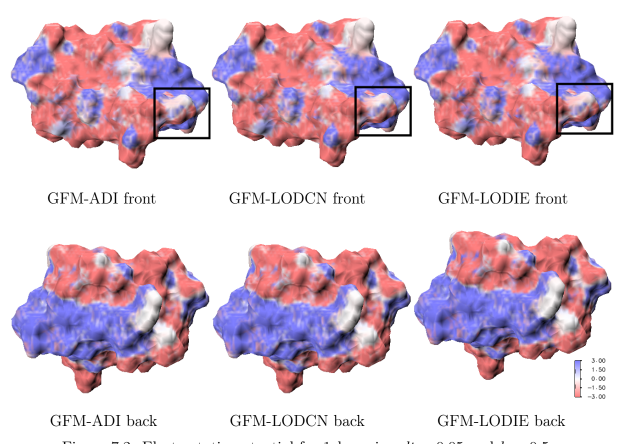


Figure 7.2: Electrostatic potential for 1cbn using dt = 0.05 and h = 0.5.

In Figure 7.2 we focus on the electrostatic potential on the surface of the protein

1cbn for all three of our prosed schemes. The difference is not that much noticeable unless we focus in the black squared region of the front side. This shows that, even though there are some differences in solvation energies calculated by the proposed three methods there is no significant difference for the electrostatic potentials for our choice for the optimum values of the parameters.

7.1.2 Solvation energy of 24 proteins

Next we solved the nonlinear PB equation and computed solvation energy for a collection of 24 proteins as in [9,11]. The dielectric constants, Ionic strength and all other parameters have been set similar to our study for the protein Crambin(1cbn).

PDB							
2erl 573 -952.36 -1165.28 -952.13 -951.03 -945.96 1cbn 648 -303.33 -459.51 -302.06 -301.09 -297.06 1vii 596 -902.31 -1154.67 -901.78 -901.36 -895.15 1fca 729 -1204.44 -1458.16 -1205.53 -1205.09 -1199.33 1bbl 576 -988.40 -1302.49 -986.15 -985.72 -979.60 2pde 667 -820.97 -1018.66 -819.17 -819.83 -813.45 1sh1 702 -753.99 -999.92 -751.69 -751.84 -745.15 1vjw 826 -1241.07 -1513.17 -1244.74 -1243.52 -1236.67 1uxc 809 -1139.25 -1478.20 -1138.50 -1135.22 -1128.39 1ptq 795 -873.32 -1170.00 -867.98 -867.40 -859.71 1bor 832 -853.47 -1102.40 -852.49 -851.24		No. of Atoms	$_{\mathrm{rMIB}}$	ADI	GFM-ADI	GFM-LODCN	GFM-LODIE
1cbn 648 -303.33 -459.51 -302.06 -301.09 -297.06 1vii 596 -902.31 -1154.67 -901.78 -901.36 -895.15 1fca 729 -1204.44 -1458.16 -1205.53 -1205.09 -1199.33 1bbl 576 -988.40 -1302.49 -986.15 -985.72 -979.60 2pde 667 -820.97 -1018.66 -819.17 -819.83 -813.45 1sh1 702 -753.99 -999.92 -751.69 -751.84 -745.15 1vjw 826 -1241.07 -1513.17 -1244.74 -1243.52 -1236.67 1uxc 809 -1139.25 -1478.20 -1138.50 -1135.22 -1128.39 1ptq 795 -873.32 -1170.00 -867.98 -867.40 -859.71 1bor 832 -853.47 -1102.40 -852.49 -851.24 -844.68 1fxd 824 -3321.39 -3653.81 -3321.68 -3321.34 <td>1ajj</td> <td>519</td> <td>-1139.48</td> <td>-1371.10</td> <td>-1139.45</td> <td>-1139.07</td> <td>-1133.06</td>	1ajj	519	-1139.48	-1371.10	-1139.45	-1139.07	-1133.06
1vii 596 -902.31 -1154.67 -901.78 -901.36 -895.15 1fca 729 -1204.44 -1458.16 -1205.53 -1205.09 -1199.33 1bbl 576 -988.40 -1302.49 -986.15 -985.72 -979.60 2pde 667 -820.97 -1018.66 -819.17 -819.83 -813.45 1sh1 702 -753.99 -999.92 -751.69 -751.84 -745.15 1vjw 826 -1241.07 -1513.17 -1244.74 -1243.52 -1236.67 1uxc 809 -1139.25 -1478.20 -1138.50 -1135.22 -1128.39 1ptq 795 -873.32 -1170.00 -867.98 -867.40 -859.71 1bor 832 -853.47 -1102.40 -852.49 -851.24 -844.68 1fxd 824 -3321.39 -3653.81 -3321.68 -3321.34 -3313.83 1r69 997 -1088.62 -1419.35 -1085.45 -1084.6	2erl	573	-952.36	-1165.28	-952.13	-951.03	-945.96
1fca 729 -1204.44 -1458.16 -1205.53 -1205.09 -1199.33 1bbl 576 -988.40 -1302.49 -986.15 -985.72 -979.60 2pde 667 -820.97 -1018.66 -819.17 -819.83 -813.45 1sh1 702 -753.99 -999.92 -751.69 -751.84 -745.15 1vjw 826 -1241.07 -1513.17 -1244.74 -1243.52 -1236.67 1uxc 809 -1139.25 -1478.20 -1138.50 -1135.22 -1128.39 1ptq 795 -873.32 -1170.00 -867.98 -867.40 -859.71 1bor 832 -853.47 -1102.40 -852.49 -851.24 -844.68 1fxd 824 -3321.39 -3653.81 -3321.68 -3321.34 -3313.83 1r69 997 -1088.62 -1419.35 -1085.45 -1084.66 -1076.32 1mbg 903 -1353.31 -1685.70 -1352.63 -13	$1 \mathrm{cbn}$	648	-303.33	-459.51	-302.06	-301.09	-297.06
1bbl 576 -988.40 -1302.49 -986.15 -985.72 -979.60 2pde 667 -820.97 -1018.66 -819.17 -819.83 -813.45 1sh1 702 -753.99 -999.92 -751.69 -751.84 -745.15 1vjw 826 -1241.07 -1513.17 -1244.74 -1243.52 -1236.67 1uxc 809 -1139.25 -1478.20 -1138.50 -1135.22 -1128.39 1ptq 795 -873.32 -1170.00 -867.98 -867.40 -859.71 1bor 832 -853.47 -1102.40 -852.49 -851.24 -844.68 1fxd 824 -3321.39 -3653.81 -3321.68 -3321.34 -3313.83 1r69 997 -1088.62 -1419.35 -1085.45 -1084.66 -1076.32 1mbg 903 -1353.31 -1685.70 -1352.63 -1351.30 -1343.68 1bpi 898 -1304.37 -1672.02 -1301.61 -12	1vii	596	-902.31	-1154.67	-901.78	-901.36	-895.15
2pde 667 -820.97 -1018.66 -819.17 -819.83 -813.45 1sh1 702 -753.99 -999.92 -751.69 -751.84 -745.15 1vjw 826 -1241.07 -1513.17 -1244.74 -1243.52 -1236.67 1uxc 809 -1139.25 -1478.20 -1138.50 -1135.22 -1128.39 1ptq 795 -873.32 -1170.00 -867.98 -867.40 -859.71 1bor 832 -853.47 -1102.40 -852.49 -851.24 -844.68 1fxd 824 -3321.39 -3653.81 -3321.68 -3321.34 -3313.83 1r69 997 -1088.62 -1419.35 -1085.45 -1084.66 -1076.32 1mbg 903 -1353.31 -1685.70 -1352.63 -1351.30 -1343.68 1bpi 898 -1304.37 -1672.02 -1301.61 -1299.86 -1291.40 1hpt 858 -812.49 -1147.42 -809.02 -	1fca	729	-1204.44	-1458.16	-1205.53	-1205.09	-1199.33
1sh1 702 -753.99 -999.92 -751.69 -751.84 -745.15 1vjw 826 -1241.07 -1513.17 -1244.74 -1243.52 -1236.67 1uxc 809 -1139.25 -1478.20 -1138.50 -1135.22 -1128.39 1ptq 795 -873.32 -1170.00 -867.98 -867.40 -859.71 1bor 832 -853.47 -1102.40 -852.49 -851.24 -844.68 1fxd 824 -3321.39 -3653.81 -3321.68 -3321.34 -3313.83 1r69 997 -1088.62 -1419.35 -1085.45 -1084.66 -1076.32 1mbg 903 -1353.31 -1685.70 -1352.63 -1351.30 -1343.68 1bpi 898 -1304.37 -1672.02 -1301.61 -1299.86 -1291.40 1hpt 858 -812.49 -1147.42 -809.02 -808.09 -799.24 451c 1216 -1027.21 -1379.27 -1023.71 <t< td=""><td>1bbl</td><td>576</td><td>-988.40</td><td>-1302.49</td><td>-986.15</td><td>-985.72</td><td>-979.60</td></t<>	1bbl	576	-988.40	-1302.49	-986.15	-985.72	-979.60
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1uxc 809 -1139.25 -1478.20 -1138.50 -1135.22 -1128.39 1ptq 795 -873.32 -1170.00 -867.98 -867.40 -859.71 1bor 832 -853.47 -1102.40 -852.49 -851.24 -844.68 1fxd 824 -3321.39 -3653.81 -3321.68 -3321.34 -3313.83 1r69 997 -1088.62 -1419.35 -1085.45 -1084.66 -1076.32 1mbg 903 -1353.31 -1685.70 -1352.63 -1351.30 -1343.68 1bpi 898 -1304.37 -1672.02 -1301.61 -1299.86 -1291.40 1hpt 858 -812.49 -1147.42 -809.02 -808.09 -799.24 451c 1216 -1027.21 -1379.27 -1023.71 -1022.61 -1012.70 1svr 1435 -1711.11 -2257.80 -1707.87 -1706.38 -1693.11 1frd 1478 -2862.50 -3376.35 -2863.69	$1\mathrm{sh}1$	702	-753.99	-999.92	-751.69	-751.84	-745.15
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1fxd 824 -3321.39 -3653.81 -3321.68 -3321.34 -3313.83 1r69 997 -1088.62 -1419.35 -1085.45 -1084.66 -1076.32 1mbg 903 -1353.31 -1685.70 -1352.63 -1351.30 -1343.68 1bpi 898 -1304.37 -1672.02 -1301.61 -1299.86 -1291.40 1hpt 858 -812.49 -1147.42 -809.02 -808.09 -799.24 451c 1216 -1027.21 -1379.27 -1023.71 -1022.61 -1012.70 1svr 1435 -1711.11 -2257.80 -1707.87 -1706.38 -1693.11 1frd 1478 -2862.50 -3376.35 -2863.69 -2863.03 -2850.42 1a2s 1272 -1921.20 -2292.15 -1919.28 -1917.70 -1907.96 1neq 1187 -1731.71 -2223.08 -1729.87 -1728.61 -1716.47 1a63 2065 -2374.41 -3149.69 -2370.80 -2369.26 -2350.42	1ptq	795	-873.32	-1170.00	-867.98	-867.40	-859.71
1r69 997 -1088.62 -1419.35 -1085.45 -1084.66 -1076.32 1mbg 903 -1353.31 -1685.70 -1352.63 -1351.30 -1343.68 1bpi 898 -1304.37 -1672.02 -1301.61 -1299.86 -1291.40 1hpt 858 -812.49 -1147.42 -809.02 -808.09 -799.24 451c 1216 -1027.21 -1379.27 -1023.71 -1022.61 -1012.70 1svr 1435 -1711.11 -2257.80 -1707.87 -1706.38 -1693.11 1frd 1478 -2862.50 -3376.35 -2863.69 -2863.03 -2850.42 1a2s 1272 -1921.20 -2292.15 -1919.28 -1917.70 -1907.96 1neq 1187 -1731.71 -2223.08 -1729.87 -1728.61 -1716.47 1a63 2065 -2374.41 -3149.69 -2370.80 -2369.26 -2350.42	1bor	832	-853.47	-1102.40	-852.49	-851.24	-844.68
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1hpt 858 -812.49 -1147.42 -809.02 -808.09 -799.24 451c 1216 -1027.21 -1379.27 -1023.71 -1022.61 -1012.70 1svr 1435 -1711.11 -2257.80 -1707.87 -1706.38 -1693.11 1frd 1478 -2862.50 -3376.35 -2863.69 -2863.03 -2850.42 1a2s 1272 -1921.20 -2292.15 -1919.28 -1917.70 -1907.96 1neq 1187 -1731.71 -2223.08 -1729.87 -1728.61 -1716.47 1a63 2065 -2374.41 -3149.69 -2370.80 -2369.26 -2350.42	1mbg	903	-1353.31	-1685.70	-1352.63	-1351.30	-1343.68
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$\begin{array}{cccccccccccccccccccccccccccccccccccc$	1 frd	1478	-2862.50	-3376.35	-2863.69	-2863.03	-2850.42
1a63 2065 -2374.41 -3149.69 -2370.80 -2369.26 -2350.42	1a2s	1272	-1921.20	-2292.15	-1919.28	-1917.70	-1907.96
	1neq	1187	-1731.71	-2223.08	-1729.87	-1728.61	-1716.47
1a7m 2809 -2160.34 -2771.41 -2155.05 -2152.48 -2135.73	1a63	2065	-2374.41	-3149.69	-2370.80	-2369.26	-2350.42
2000 2100.01 2,11.11 2100.00 2102.10 2100.10	1a7m	2809	-2160.34	-2771.41	-2155.05	-2152.48	-2135.73

Table 7.3: Solvation energies (kcal/mol) of 24 Proteins considering $\Delta t = 0.001$ for ADI and $\Delta t = 0.05$ for GFM-ADI, GFM-LODCN, GFM-LODIE

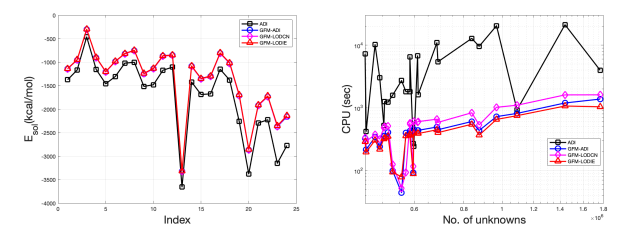


Figure 7.3: Difference in solvation energy(left) and CPU time (right) in solving nonlinear PB equation for 24 protein.

Solvation energies for all three of our proposed schemes have been compared with the rMIB and MIB schemes in Table 7.3. The results from our proposed schemes have been observed to be very close to the results from rMIB and MIB schemes while solving nonlinear PB equation instead of linear PB equation. As we have identified in previous subsections GFM-ADI appeared to be more accurate than the other two of our proposed schemes and obtained the same level of accuracy as rMIB and MIB schemes. Table 7.3 also confirms that if GFM-ADI fails to converge for any protein then GFM-LODCN or GFM-LOD can also be used since they are more stable and the results are not that much different than GFM-ADI.

7.1.3 Binding energy of 2a9x

Binding energies play an important role in viral transcription and antiviral drug designs. In particular a better accuracy of the binding energy of the BIV Tat Protein and BIV TAR RNA in HIV viral replication can significantly help in search for the new antiviral drugs that repress the replication by blocking transactivation of viral RNA transcription [17]. In this section we will demonstrate the ability of GFM-ADI scheme to compute the binding energy of the BIV Tat Protein and BIV TAR RNA.

The electrostatic binding free energy can be calculated by the following formula

based on the free energy cycle,

$$E_{\text{bind}}^{\text{AB}} = \Delta G_{\text{ele}}^{\text{AB}} - \Delta G_{\text{ele}}^{\text{A}} - \Delta G_{\text{ele}}^{\text{B}}$$

$$= [E_{\text{sol}}^{\text{AB}} + E_{\text{cou}}^{\text{AB}}] - [E_{\text{sol}}^{\text{A}} + E_{\text{cou}}^{\text{A}}] - [E_{\text{sol}}^{\text{B}} + E_{\text{cou}}^{\text{B}}]$$

$$(7.4)$$

where the free energies of the complex AB and its monomers A and B on the RHS can be calculated using the calculated solvation energies in equation (4.15).

h	$E_{\rm sol}^{\rm complex}$	$E_{\rm sol}^{ m protein}$	$E_{ m sol}^{ m RNA}$	$E_{\rm bind}^{\rm complex}$			
rMIB							
1	-5816.38	-1021.94	-8893.39	383.60			
1/2	-5821.22	-1025.86	-8898.54	387.84			
1/4	-5823.39	-1026.27	-8900.52	388.05			
GFM-ADI							
1	-5834.10	-1027.14	-8915.17	392.84			
1/2	-5824.82	-1025.98	-8905.59	391.38			
1/4	-5841.62	-1026.40	-8916.69	386.12			

Table 7.4: Binding energy of 2a9x

7.1.4 Salt effect on the binding affinity

The nonlinear PB equation is often used to describe the salt effects on the binding of ligands, peptides, and proteins to nucleic acids, membranes, and proteins. In this investigation we have tested the the performance of the proposed GFM-ADI scheme for the evaluation of the salt effect in the protein-protein binding of the complex 1beb and 1emv. Physically, the binding affinity can be quantitatively represented based on the binding-free energies, which reflect the non-specific salt dependence of the formation of macro- molecular complexes. The binding affinity is then calculated as the slope ratio of the salt-dependent binding energy at certain salt strength I_s against the natural logarithm of I_s . The electrostatic binding-free energy can be further split into $E_{\text{cou}}(I_s)$'s as the salt-dependent parts and $E_{\text{sol}}(I_s)$'s as the salt-dependent parts. The variation of the salt-dependent part of the binding-free energy $\Delta E_{\text{bind}}(I_s)$ can thus be calculated as the difference in $E_{\text{bind}}(I_s)$ for some nonzero salt strength and the zero salt concentra-

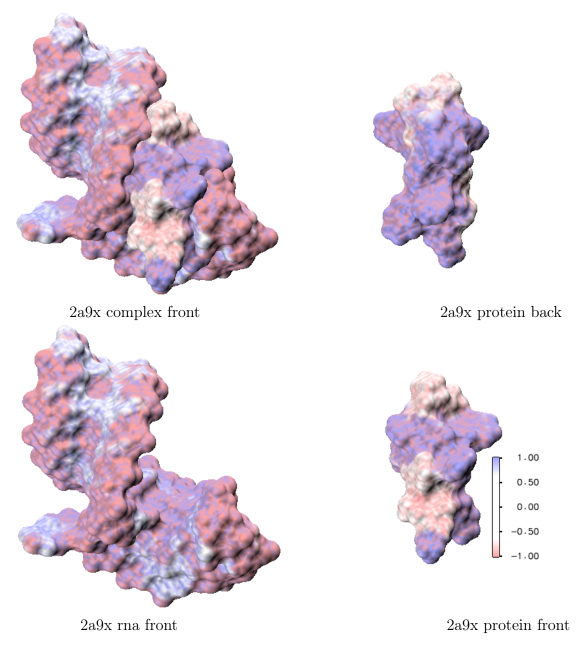


Figure 7.4: Electrostatic potential for 2a9x using GFM-ADI for $\Delta t = 0.05$ and h = 0.5

tion, because the salt independent parts gets simply cancelled. Altogether we have the following formula.

$$\Delta E_{\text{bind}}(I_s) = E_{\text{bind}}(I_s) - E_{\text{bind}}(0)$$

$$= [E_{\text{sol}}^{AB}(I_s) - E_{\text{sol}}^{AB}(0)] - [E_{\text{sol}}^{A}(I_s) - E_{\text{sol}}^{A}(0)] - [E_{\text{sol}}^{B}(I_s) - E_{\text{sol}}^{A}(0)] (7.5)$$

For this study we have used the same model parameters used earlier for 24 proteins. In Figure 7.5 we reported the calculated binding free energy with the experimental results. The slope ration or the binding affinity is calculated and reported in Table (7.5) as in [39]. The results attained by the Lagrangian formulation linearized PB (LFLPB) model [37] are also given in Table (7.5) for comparison. For 1beb the binding affinity calculated by GFM-ADI scheme sufficiently close to experimental data and better than LFLPB. For 1emv the results from GFM-ADI is not as good as those of the LFLPB model but qualitatively it agrees with the experimental observations; that is as the hetero-diemric complex, the binding-free energy increases when the ionic strength is larger. This is probably because the calculation of the binding affinity requires a physical cutoff to obtain two monomers A and B.

	Charges				Slope ratios		
Complex	PDB	AB	A	В	Experimental	GFM-ADI	LFLPB
Lactoglobulin dime(A-B)	1beb	+26	+13	+13	-1.62	-1.82	-2.02
E9Dnase-Im9(10)(B-A)	$1 \mathrm{emv}$	-3	-8	+5	2.17	0.52	2.4

Table 7.5: Comparison of binding affinities of the protein complexes 1emv and 1beb

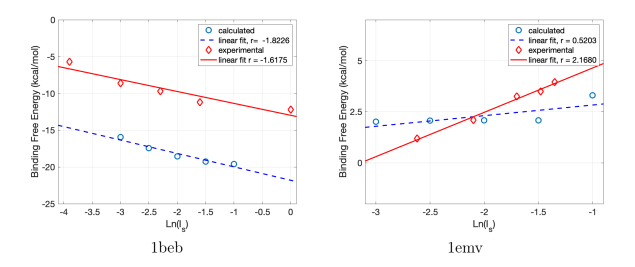


Figure 7.5: The salt dependence of the binding affinities

CHAPTER 8

CONCLUSIONS

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