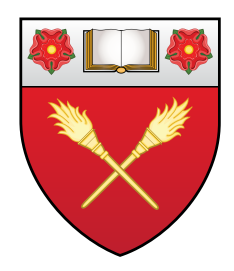
Dislocation Based Modelling of Fusion Relevant Materials.





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Dedication

"Ohana significa familia, y tu familia nunca te abandona, ni te olvida."

"Ohana means family, family means nobody gets left behind, or forgotten."

— Stitch, Lilo & Stitch

Malle santa, cuando te hablé desde el trabajo para darte la noticia me dijiste que siempre lo supiste. Recuerdo que lloraste en el teléfono, se te cerró la garganta y a mí también. A diferencia de otras veces, no pudimos platicar mucho porque nos quedamos sin aliento. Así que le hablaste a abuelita Raquel mientras le hablé a papá. Madre, David y yo te debemos tanto que no podemos repagar en mil vidas, pero ten por seguro que eres nuestro ejemplo a seguir. Nuestra madre chingona y chambeadora, solo hay una y como ella no hay ninguna. Sin tí el mundo sería un lugar más cruel y pobre, no merece un ángel tan grande y puro como tú.

Dad, "Igualito que tu jefe, wey." con tu risa característica fue lo primero que dijiste cuando te avisé "Sí dad, igualito que mi jefe." entre risas respondí. Como siempre, no platicamos mucho pero esta vez porque le querías avisar a abuelita Teté y a mis tíos, y yo le quería avisar a David. Pensé que seguirías aquí para carcajearte al verme en la túnica ridícula, así como lo hicimos nosotros cuando te vimos a tí en la tuya. Escribo esta dedicatoria antes de acabar porque una promesa es una promesa y esta madre la voy a acabar. No te tendré para pedirte consejos y contarte mis avances y tropiezos. Pero a veces pretendo que me escuchas mientras intento comprender o arreglar algo. Te queremos y te extrañamos muchísimo. Buenas noches, dad.

David, "Te mamaste we." me dijiste cuando me abrazaste al llegar a casa después del trabajo el día que me aceptaron. Me has hecho un chingo de falta ojete, te extraño mucho we. Perdón por no hablar tan seguido, pero me duele colgar we. No estoy tan chisqueado como mamá pero siento feo cuando terminamos de platicar. Me gusta mucho ver tus streams porque me recuerda un poco a sentarme a tu lado a verte jugar... o a cuando veíamos Twitch juntos y veíamos a TB missing legal.

Esto es para mi Ohana por sangre y por elección. Ustedes creyeron en mí cuando yo no lo hacía. Me empujaron a ser mejor. Me extendieron la mano cuando nadie más lo hizo. Me hicieron reir cuando solo sabía llorar. Gracias por hacerme quien soy.

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"You can always judge a man by the quality of his enemies."

— Oscar Wilde

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I guess I'll content myself with being judged unfavourably by the world.

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Outline

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Chapter 0

Preface

0.1 Notation

For the sake of clarity the following notation conventions have been used:

- Tensors are denoted by sans serif bold italics, T.
- Matrices are denoted by serif bold roman, M.
- Vectors are denoted by serif bold italics, V.
- Scalars are denoted by serif italics, S.
- Host variables are denoted by $^{\rm h}a$.
- Global device variables are denoted by $^{\rm d}a$.
- Thread variables are denoted by ^ta.

0.2 Conventions

All angles are given in radians unless stated otherwise. All pseudo-code is C-style and assuming C conventions (row-order, 0-start indexing, memory allocation).

Elements, be they Surface Elements (SEs), Finite Elements (FEs), dislocation line segments or else, are denoted by e and their total as E. Similarly nodes are denoted by n and their total N. Parallel indices are denoted by idx to distinguish them from regular indices/counters i, j, k. For disambiguation purposes the exponential function will de denoted as $\exp(x)$.

0.3 Typesetting

This document was typeset using a custom-made LaTeX document class created by the author, compiled with XaLaTeX, and the bibliography was produced with BibTeX. The custom document class can be found in https://github.com/dcelisgarza/latex_temp

0.4 Diagrams

All diagrams are vector graphics drawn using the open-source image editing software InkScape for its ease of use, elegant simplicity, high quality outputs, and "Draw Freely" philosophy.

Chapter 1

Introduction

1.1 Materials Science and History

Metals and alloys are of such importance to humankind that entire eras of our history have been defined by and named after discoveries and advances in metal-lurgy [1, 2]. A civilisation's ability to gain mastery of the properties and usage of materials is often strongly correlated with its success in history [3]. The historical influence of metals and alloys has touched all areas of human existence, from fashion to warfare [4].

It should come as no surprise that a material's use is often linked to its properties. When a civilisation learned to harness the properties of a novel material, its influence often grew as a result [5, 6]. Whether building a woodcutter's axe or a fusion reactor, the mechanical behaviour and deformation of materials—metals and alloys in particular—are still among their most important properties for engineering applications.

Traditionally such properties have been largely studied via empirical methods such as tensile and compressive tests, beam bending, and indentation [7–9]. Such tests have mainly focused on macroscopic scales, thus offering bulk-averaged properties valuable for engineers. Unfortunately such tests only describe observed behaviours without truly elucidating the fundamental mechanisms behind them [10]. This has recently started to change with the advent of micro-scale testing, where tests can be performed on single crystals or individual crystal boundaries, and can even probe specific slip systems to see how they behave under various loading scenarios [11, 12]. The increased resolution and more thorough parameter control goes a long way in providing a window through which more fundamental behaviours can be observed and hopefully explained; thus providing a way of deconvoluting macroscopic observations into their constituent phenomena [13, 14].

However, understanding the underlying mechanisms behind empirical results

through experimental means has proven difficult at every scale [15, 16]. Fortunately modelling and simulation can be powerful allies in this task. In particular, the study of dislocations is experimentally challenging [17–19]. By virtue of being such an important player in the mechanical properties of materials, dislocations demand careful study from both empirical and in-sillico avenues.

The need for a more detailed picture of material properties is especially relevant in highly engineered alloys subjected to extreme conditions, such as those found in fusion reactors. The time-dependent shift in material properties and composition is of the utmost concern such extreme environments; radiation damage, large temperature gradients, gas diffusion and plasma instabilities provide the means for properties to change drastically over a components' lifetime [20–22]. It is therefore imperative that we find ways of predicting and accounting for these changes. This requires a deeper understanding of the process driving the mechanical behaviour. Virtual experiments allow analysis and interpretation of micromechanical tests and can help provide new fundamental insight.

1.2 Fusion Energy Production

There currently exist two major branches of research for fusion energy production [23]: 1) magnetic confinement fusion (MCF) [24] which confines the plasma using magnetic fields and 2) inertial confinement fusion (ICF) [25] which uses a frozen fuel pellet which is either directly or indirectly compressed by arrays of powerful lasers. The physics and engineering challenges vary greatly between both approaches but the fundamental materials problems remain largely the same, with few a exceptions such as divertors [26, 27].

1.2.1 Operating environment

One constant feature of nuclear energy production is ionising and non-ionising radiation. In the case of fission, this is mostly in the form of low energy neutrons and residual radiation of fission products; fusion on the other hand, deals with the sparsely explored 14 MeV neutron spectrum [28]. The lack of appropriate sources of suitably energetic neutrons [29] has meant that modelling, and searching for experimental analogues of damage cascades have become a crucial part of materials research [30].

Operating environments change vastly between ICF and MCF as described by table 1.1 [31]. The table should be read with a measure of skepticism as the numbers and conditions are approximate because they are not yet fully known, especially for ICF. However it is still, a fairly reasonable first approximation into

Location	Radiation Type	MCF (ITER)	ICF (LMJ)
1 st Wall	Neutron flux Neutron fluence* γ -ray dose rate Energetic ion/atom flux	$3 \times 10^{18} \mathrm{m}^{-2} \mathrm{s}^{-1}$ $3 \times 10^{25} \mathrm{m}^{-2}$ $2 \times 10^{3} \mathrm{Gy} \mathrm{s}^{-1}$ $5 \times 10^{19} \mathrm{m}^{-2} \mathrm{s}^{1}$	$1.5 \times 10^{25} \mathrm{m}^{-2} \mathrm{s}^{-1}$ $3 \times 10^{18} \mathrm{m}^{-2}$ $\sim 1 \times 10^{10} \mathrm{Gy} \mathrm{s}^{-1}$
1 st Diagnostic	Neutron flux Neutron damage rate Neutron fluence* Neutron damage γ -ray dose rate Energetic ion/atom flux Nuclear heating Operating temperature Atomosphere	$\begin{array}{c} 1\times 10^{17}\mathrm{m^{-2}s^{-1}} \\ 6\times 10^{-9}\mathrm{dpas^{-1}} \\ 2\times 10^{24}\mathrm{m^{-2}} \\ 1\times 10^{-1}\mathrm{dpa} \\ \sim 1\times 10^{2}\mathrm{Gys^{-1}} \\ \sim 1\times 10^{18}\mathrm{m^{-2}} \\ 1\mathrm{MWm^{-3}} \\ 520\mathrm{K} \\ \mathrm{Vacuum} \end{array}$	$1 \times 10^{26} \mathrm{m}^{-2} \mathrm{s}^{-1}$ negligible $\sim 1 \times 10^{19} \mathrm{m}^{-2}$ negligible $\sim 1 \times 10^{10} \mathrm{Gy s}^{-1}$ 0 293 K Air
Other	EM pulse Shrapnel		10 to 500 kV m ⁻¹ @ 1 GHz 1 to 10 km s ⁻¹ @ \sim 30 μm

Table 1.1: Estimated operating environment comparison between MCF (ITER) and ICF (LMJ). Reproduced from [31]. Note these numbers are unrepresentative of actual fusion power plants as both ITER and the LMJ are experiments—it is likely power plants will require much harsher operating conditions.

* End of life.

the materials requirements for both "mainstream" types of fusion energy production. That said, the components needed for energy harvesting—a divertor in the case of MCF, and an open problem for ICF—are conspicuous by their absence.

Table 1.1 shows that dosages and dosage rates vary wildly between approaches. For the most part, MCF receives higher doses of neutrons and ions in both the first wall and diagnostic equipment. This seems to indicate that the material requirements for MCF are stricter than for ICF. However, shrapnel production is a strong possibility in ICF, especially in indirect drive reactors; where a specially made container called a "hohlraum" holds the fuel pellet and is subsequently obliterated when the pellet undergoes fusion (see fig. 1.1 [32]). This is a huge challenge as such high energy shrapnel would be capable of destroying diagnostic equipment and damaging the first wall [33–35].

Overall, the table is unrepresentative of potential operating environments in large-scale power stations, but sets lower limits on the demands of the materials involved in both "mainstream" proposals for fusion energy production.

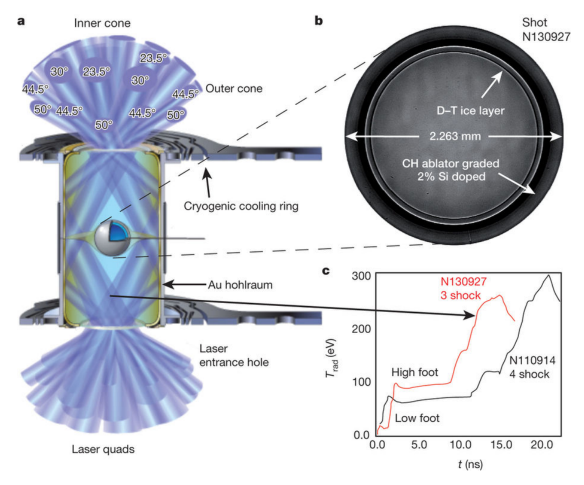


Figure 1.1: **a**, Cross section of the National Ignition Facility's ICF target design showing the gold hohlraum and plastic capsule, fuel pellet, and incident laser bundles. **b**, X-ray image of the actual capsule for N130927 with DT fuel layer and surrounding CH (carbon-hydrogen) plastic ablator. **c**, X-ray radiation drive temperature as a function of time for the National Ignition Campaign (NIC) low-foot implosion and the post-NIC high-foot implosion. Image taken from [32].

1.2.2 Effects of Radiation on Reactor Materials

1.2.2.1 Radiation Damage

One of the bigger problems in radiation damage research is the lack of a truly standardised way of measuring damage on materials [36]. The most common unit is displacements per atom (dpa) [37]. It is defined as the average number of displacements undergone by each atom in a material as a result of being irradiated. The "fundamental" unit of measurement for this is the number of displacements per unit volume per unit time, R

$$R = N \int_{E_{\min}}^{E_{\max}} \int_{T_{\min}}^{T_{\max}} \phi(E) \, \sigma(E, T) \, \upsilon(T) \, dT \, dE , \qquad (1.1)$$

where N is the atom number density (no. of atoms per unit volume); E the incoming particle's energy; T the energy transferred in a collision of a particle of energy E and a lattice atom; $\phi(E)$ the energy dependent particle flux; $\sigma(E,T)$ the cross section for the collision of a particle with energy E resulting in a transfer of energy T to the struck atom; and v(T) the number of displacements per primary knock on atom as a function of transferred energy T. DPA can be calculated by naïvely multiplying E by the sample volume and total exposure time (or we could use fluence rather than flux). This ignores the fact that $\sigma(E,T)$ and v(T) will be locally perturbed in the neighbourhood of damage cascades, since the bulk volume is much greater than that of the damage cascades', we assume the functions remain globally unchanged.

In principle, this is a rather good measure of damage [37]. The catch is that generalised, analytical expressions for $\sigma(E,T)$ and v(T) depend on a slew of parameters and are therefore incredibly hard if not impossible to derive. This does not mean they cannot be discretised and roughly approximated via Monte Carlo (MC) approaches [36–38]. However, damage cascade modelling falls squarely in the realm of pico- to nanosecond timescales and as such only feasable with Molecular Dynamics (MD) and Kinetic Monte Carlo (KMC) [39, 40] approaches. At the end of such cascades, we are often left with dislocation sources or prismatic dislocation loops [41] which can be used as inputs by a Dislocation Dynamics (DD) simulation to study larger temporal and spatial scales [30].

1.2.2.2 Transmutation

Transmutation products are one of the biggest sources of problems for materials in fusion applications [20, 21]. Not only do they tend to embrittle materials, they often also reduce their thermal conductivity [42]. The former presents significant challenges for structural materials [43]; the latter is especially egregious for energy extraction by limiting the divertor's ability to conduct heat thus lowering the reactor's efficiency and causing thermal stresses due to the generation of hot spots that cannot be easily dissipated. As a result, understanding the mechanical and thermal behaviour of transmutation alloys is crucial for moving forward [44–46] but doing so requires knowledge of the time evolution of a reactor's components. Because the composition changes in a a non-trivial way, so do the mechanical properties. Worse still are the long timescales over which this happens—on the order of years or tens of years [47]. Even if we were able to experimentally irradiate fusion-relevant materials with appropriately energetic neutrons, it would take years before we could characterise their behaviour, and doing so would be problematic due to radioactive decay.

The way we go about addressing the time-dependent compositional change of a material is to model it. Culham Centre for Fusion Energy's (CCFE) FISPACT [48] software takes an MC approach at calculating transmutation and decay products of a sample given certain conditions. The code utilises external data provided by the European Activation File which provides cross sections and decay rates for a wide range of isotopes [49]. Unfortunately there is a lack cross section data for certain neutron energies that contribute in a non-negligible manner to a fusion environment. Interpolating to fill the gaps would not be appropriate as the data is non-smooth and subject to resonance peaks, so the solution is imperfect.

Transmutation products will always be problematic, but they are a fact of working with fusion-relevant materials. Fortunately there are ways in which inclusions may be modelled via DD (section 1.5.4.2). DD may also be used to model dislocation movement and interaction within heterogenous media (section 1.5.4) as is the case for oxide-dispersion-strengthened (ODS) steels; and transmutation alloys of Tungsten divertors, where Osmium, Rhenium and Tantalum clusters which prove highly problematic for its temperature conductivity and structural integrity [50–56].

1.3 Parallel Computing

Processing chips are made up of millions or even billions of transistors acting as switches in logic gates. Each time a logic gate fires, the capacitors inside them charge and discharge at the chip's frequency or clock speed [57]. Consider the power of any electronic component,

$$P(t) = I(t)V(t), (1.2)$$

where I is current and V is voltage both as a function of time. The current, I, of a capacitor and the definition of power, P, as functions of time, t, are given by,

$$I(t) = C \frac{\mathrm{d}V(t)}{\mathrm{d}t}, \qquad P(t) = \frac{\mathrm{d}E(t)}{\mathrm{d}t},$$
 (1.3)

where C is capacitance and E the energy stored in the capacitor. Substituting eq. (1.3) into eq. (1.2) and integrating twice, we obtain the expression for the energy stored in a capacitor,

$$E_{\rm c} = \frac{CV_{\rm c}^2}{2},\tag{1.4}$$

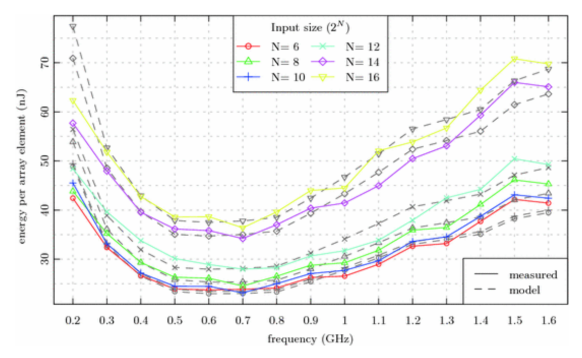


Figure 1.2: Energy required by a Samsung Galaxy S2 CPU at 37 °C to complete the Gold-Rader implementation of the *bit-reverse* algorithm. The dashed lines denote a theoretical model, image and model found in [60].

where V_c and E_c are the voltage and energy stored in the capacitor. Recalling that in a processing chip, the capacitors are charged and discharged at the chip's clock speed f, we arrive at the expression for power consumption, P_c , of a capacitor charging and discharging at frequency f [58],

$$P_{\rm c} = E_{\rm c} f \propto C V_{\rm c}^2 f \,. \tag{1.5}$$

Computer chips are rather more complicated, but their power consumption (also known as power dissipation) is described by a simple addition of terms [59],

$$P = P_{\rm dyn} + P_{\rm sc} + P_{\rm leak}, \qquad (1.6)$$

where $P_{\rm dyn}$ is the dynamic power dissipation given by eq. (1.5). The two other dissipation mechanisms are: 1) short-circuit, sc, which depends on frequency and occurs when a direct path from transistor to ground is made as a result of multiple transistors conducting simultaneously; and 2) leakage, leak, which depends on the voltage and is due to micro-currents between doped parts a transistor. Furthermore, higher voltages and frequencies result in higher temperatures, which in turn mean decreased transistor performance and increased capacitance. The overall result is an energy expenditure curve similar to fig. 1.2 [60] for every processor unit.

This set of optimum conditions is the reason behind the multicore and multi-

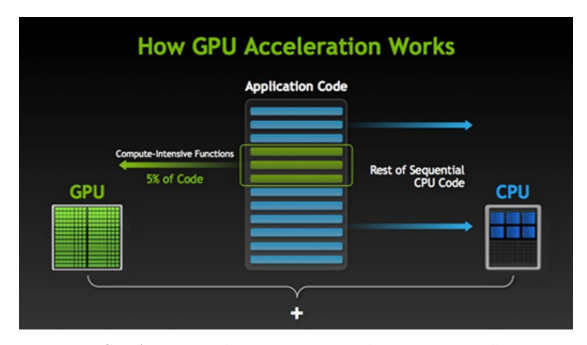


Figure 1.3: CUDA runtime schematic. Repetitive but computationally intensive tasks can be offloaded to the GPU. Both GPU and CPU are completely independent from each other and can work on different parts of the code at the same time, care must be taken to ensure proper syncronisation. Image taken from [61].

threaded design of modern Central Processing Units (CPUs) and Graphics Processing Units (GPUs). It is also why in recent years there has been such a massive push for parallelism in all computing markets. It is simply not feasable to continually increase clock speeds and voltages because cooling solutions would struggle to remove heat fast enough, and power consumption would skyrocket.

1.4 Computation on Graphics Processing Units

Central Processing Units (CPUs) are designed not only to perform mathematical operations but logical ones that control program flow. They are tailored to perform the wide variety of operations required by an operating system. These include program scheduling (load balancing), instantiation (program loading, unloading, loops, recusion instances) & branching (if/case statements, go to's), memory operations (fetching, storing, allocation), input/output (IO), and program monitoring (program counters, recursion counters). Modern CPUs have a degree of parallelism that allows them to increase their total throughput while keeping their operation within near optimum conditions as previously mentioned. They are commonly divided into cores and threads, though certain high-end chips have an additional layer named hyperthreading [62].

Given the limited scope of the first computers, the general purpose of CPUs was enough to cover their needs. With the advent of personal computers, the demands

on CPUs drastically increased. For industrial users these revolved around data acquisition, filtering and preprocessing [63–65]. On the other hand, the domestic market demanded ever increasing levels of abstraction and usability in the form of Graphical User Interfaces (GUIs) such as windows, cursors and their accompanying sound effects. Manufacturers identified this and moved to provide specialised modules that freed CPU resources and improved user experience by accelerating different processes through hardware means [66–69]. These modules include Sound Cards (SCs), Field Programmable Gate Arrays (FPGAs), Graphics Processing Units (GPUs), Application Specific Integrated Circuits (ASICs), Cryptographic Accelerators, Regular Rxpression (RegEx) Accelerators, among others. They are collectively dubbed "hardware accelerators".

Graphics Processing Units were originally intended to offload the very dataintensive but computationally simple operations needed for 3D gaming and rendering [66–68]. Graphics processing was seen as a prime candidate for hardware
acceleration because the operations on each pixel are largely the same across the
screen. Due to their original purpose as gaming and rendering accelerators, they
were never designed to operate on higher than single precision data. In fact, single
precision (32-bit precision) is still good enough to encode 32-bit colour depth (8bit channel per RGB colour + 8-bit alpha channel), which only the most high-end
monitors support [70]. It is also worth noting that because the same operations
apply to different pieces of (mostly) independent data, they can all be performed
at the same time, often trivially reducing the order of polynomial complexity
algorithms.

As previously mentioned, GPU parallelism frees up enormous amounts of CPU processing power that can be put to good use running other programs or performing complementary serial processes, while the GPU works concurrently on its dataset. Parallelism has been used by the scientific community for years, but the focus has mainly been on CPU parallelism [71]. Consequently, its scope was lagerly limited to the use of computational clusters. The two main reasons for this were the fact that GPUs lacked support for higher precision arithmetic, and the very limited to non-existent support for scientific computing in languages such as OpenGL [72].

It wasn't until the development of the OpenCL and OpenACC standards that GPUs caught the attanetion of the scientific community as a viable way of exploiting parallelisation without access to a computing cluster.

OpenCL allows one to work with heteragoenous systems and is similar to C in that it's very low level. It works on a wide range of hardware accelerators and is therefore useful for many scientific and engineering applications, but it's also relatively hard to use. One can make use of libraries written in OpenCL to

facilitate development, but it is still a fully fledged C-type language [73].

OpenACC is similar to OpenMP in that they both use pragmas¹ and they both work in shared memory environments—same GPU and same CPU respectively. This is no coincidence as OpenACC was designed as an extension of OpenMP for developing parallel applications on hardware accelerators. Unfortunately, being pragma based, the standard requires significant work by compiler manufacturers, so its adoption has therefore been slow [74]. Furthermore, despite simplifying development and minimising the barrier to entry, the use of pragmas limits the flexibility and adaptability of the framework compared to OpenCL.

Recently however, a third option has become increasingly viable. NVidia's Computer Unified Device Architecture (CUDA) framework provides the best aspects of both OpenCL and OpenACC. The tradeoff is that it only works on NVidia GPUs and is a closed source product. However, the accessibility and flexibility of CUDA provides anyone familiar with C/C++ the means to develop a GPU application with little issue. NVidia is also strongly backing scientific research by adding double precision support on their GPUs. They have also worked to provide parallel equivalents of well known serial libraries—such as cuBLAS & cuFFT—for scientific computing. They have additionally developed a wide range of specialist graphics cards tailor-made for scientific purposes. As such, they are the leaders in GPU computing in scientific communities [61].

1.4.1 Hurdles for Parallelisation

The difficulty in parallelisation varies tremendously from problem to problem. Problems where data is uncorrelated and independent—such as sampling well-behaved probability distributions—are almost trivially parallelisable. Problems where data is correlated or strongly dependent on its neighbours—such as Dislocation Dynamics—require a more careful approach [75].

The largest hurdle when implementing parallel algorithms is often the efficient use of data. In order to obtain good parallel performance, a lot of thought has to be placed on data access patterns, data read/write conflicts, and memory allocation and transfer [61]. For best performance, all this must be analysed on a case by case basis. If done incorrectly, the performance of a parallel application may be

¹Pragmas are especially formated comments that specific compilers recognise and turn into special instructions (often for moving memory to the appropriate hardware) at compile time. They are not part of the programming language's official standard, but rather compiler-specific extensions. Pragmas are usually programmed in C or Assembly and must be implemented by the compiler manufacturer. Nothing prevents application developers from writing the would-be pragma's code directly into their application, but this can prove a lengthy and difficult process. The rigid nature of pragmas, and the compiler manufacturer's priorities limit their scope and usability.

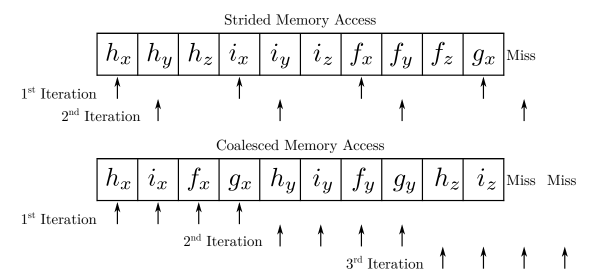


Figure 1.4: Arrows represent fetch requests by single threads in a GPU. Data should be arranged in such a way that all threads in a warp (collections of 32 threads) access contiguous memory locations to reduce time-consuming memory fetch operations. This is often unintuitive but extremely important, especially on scientific computing cards which are optimised for long computational times and low memory fetch frequency.

lower than the serial version. One must consider a wide range of parameters to successfully parallelise a problem. Among these are GPU architecture, problem size, computational and memory complexity, code branching, required arithmetic precision, and error tolerances [61, 76].

Efficient parallelisation of many problems requires "coalesced memory access" as shown in fig. 1.4, which means we have to be extremely careful when mapping CPU memory to global GPU (device) memory. The fact that threads work "simultaneously" means that in order to obtain good performance, data which is to be "simultaneously" loaded into each thread must be contiguous. This maximises cache memory use and therefore reduces slow memory fetch operations to global or shared memory.

Special cases, such as having a parallel dislocation line segment to a surface as discussed in section 1.5.3, must be treated carefully due to the way code branching works in GPUs. There are various ways of doing so: 1) if the special case is inexpensive, it can be treated within the same GPU function; 2) if the special case is expensive and always known (certain boundary conditions in FEM), it can be placed in its own GPU function that treats it separately; 3) if the special case is expessive and found at runtime it may be asynchronously treated by the CPU or buffered into its own GPU function to be executed at a later time.

One of the advantages of GPU-CPU independence is that both can work con-

²Not quite but essentially simultaneously. See [61] for details.

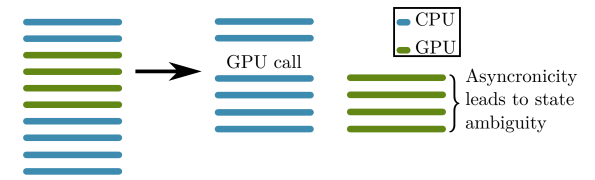


Figure 1.5: GPU and CPU code run independent of each other. Leads to state ambiguities if both sides need to talk to each other [61].

```
if (y == 0){
    x = a+b;
}
else if (y == 1){
    x = a*b;
}
else{
    x = a/b;
}
p = (y == 0);
p: x = a+b;
p: x = a+b;
p: x = a*b;
!p: x = a/b;
}
```

- (a) CPU code will only execute if the condition is met. This is called code branching.
- (b) GPU code executes every line but only stores results if the the flag before the colon is true.

Figure 1.6: The NVidia CUDA compiler replaces if and case statements with logical flags p. Every line is executed, but the data is only stored if the flag prior to the colon is true [61]. This means that having rare but computationally expensive special cases will tank parallel performance and must therefore be dealt with separately.

currently on the problem. If both systems have to talk to each other, then one must tread carefully, ensuring proper syncronisation and data mapping before moving (see fig. 1.5).

The reason why code branching is bad for GPU parallelisation is down to the fact that they work like software customisable vector machines. Where collections of threads all carry out the same operation on different pieces of data at the same time. This means that regardless of whether a condition is true or false, the code will execute. It is only data storage that depends on the condition, as illustrated in fig. 1.6.

Dislocation Dynamics (DD)—in particular 3D Discrete Dislocation Dynamics (3D DDD)—can greatly benefit from parallelisation, especially when coupling to Finite Element Methods (FEM). It is worth noting that there are potential issues arising from the very computationally expensive functions and special cases that

often arise from analytical solutions in 3D DDD. There are also potential issues with data redundancy—which are non-limiting in the short term—that may eventually require a more data-efficient approach as the computational capabilities and data capabilities of GPUs converge. Section 1.5.5 expands on these issues in the context of DD.

1.5 3D Dislocation Dynamics Modelling

The plastic deformation of materials is generally governed by the generation and motion of line defects known as dislocations through the crystal lattice. Microstructural features such as grain boundaries, precipitates and inclusions impede dislocation motion causing strengthening but often limiting ductility [77]. Understanding the behaviour of the dislocation ensemble is highly complicated even when ignoring dislocation-microstructure interactions. However, if we want to truly comprehend their real-world behaviour, we cannot limit ourselves to idealised scenarios.

One of the most often used parametrisations of DD is Discrete Dislocation Dynamics (DDD). Where dislocations are parametrised as a series of nodes linked by straight line segments. This reduces computational requirements and allows for analytic solutions to be obtained. At present, neither DDD or Finite Element (FE) models can truly handle all of the complexities of real-world alloys [78–81]. For one, DDD relies on assuming a linear-elastic isotropic solid domain with periodic boundary conditions, while FE relies on assuming continuum properties inside its finite domain.

Crystal Plasticity Finite Element Methods (CPFEM) use constitutive equations to calculate dislocation motion and generation on a set of slip systems. These are given as dislocation densities and thus are still "bulk" models because they don't interact with each other [82]. CPFEM can handle finite strains and anisotropy but not strain localisation/slip bands, etc.

Furthermore, dislocations often accumulate in the vicinity of microstructural features. The interaction of dislocations with microstructure can lead to further increased hardening and local hot spots in stress that can lead to failure initiation [83, 84]. Hardware accleration, i.e. using Graphics Processing Units (GPUs), has been shown to be very effective in DDD [85], and has the potential to enable the simulation of much larger numbers of dislocations for longer timescales. In order to study these phenomena we must find a way of coupling DDD and FEM into a single multiscale model that can simulate micromechanical tests more accurately than CPFEM. With a model such as this we may potentially be able to predict and observe emergent phenomena/properties, explain michromechanical behaviours,

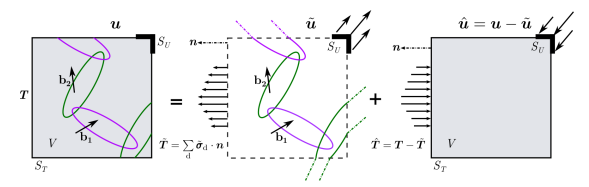


Figure 1.7: Decomposition of the boundary value problem. The volume V is bounded by surface S contains a dislocation ensemble and is subjected to initial traction conditions \mathbf{T} on S_T and \mathbf{u} on S_u . First, the traction, $\tilde{\mathbf{T}}$, and displacement, $\tilde{\mathbf{u}}$, fields due to the dislocations in the infinite domain (DDD) are evaluated on the boundaries S_T and S_u respectively. Then a traditional FE method is used to calculate the stress field satisfying the boundary conditions $\hat{\mathbf{T}} = \mathbf{T} - \tilde{\mathbf{T}}$ and $\hat{\mathbf{u}} = \mathbf{u} - \tilde{\mathbf{u}}$. Image edited from [86].

and even predict and explain experimental results based on underlying dislocation mechanisms.

1.5.1 Coupling Dislocation Dynamics to Finite Element Methods

Coupling dislocation dynamics (DD) to FEM is important to properly simulate micromechanical tests because DD provides us with a more precise set of inputs and greater granularity for solving the FE problem. There are at present two methods with which to do so, the Superposition Model and the Discrete-Continuum Model. The most accurate implementation of dislocation dynamics is discrete dislocation dynamics, where the dislocation is broken into nodes and connecting segments. This is computationally efficient and allows for analytical solutions. However, as discussed in section 1.5.1.3 explicit discretisation is not the only way to tackle the problem.

1.5.1.1 Superposition Model

The Superposition Model (SM) works by decomposing the problem into separate DDD and FE problems (fig. 2.1). It is assumed that a linear-elastic body V bounded by a surface S is subject to traction boundary conditions, T, on S_T and displacement boundary conditions, u, on S_u . The formulation proposed in [87] to impose traction-displacement boundary conditions on DDD problems in finite domains states that the total displacement and stress fields can be written as a

superposition of displacement and stress fields obtained from DDD and FE,

$$\boldsymbol{u} = \tilde{\boldsymbol{u}} + \hat{\boldsymbol{u}}, \tag{1.7a}$$

$$\boldsymbol{\sigma} = \tilde{\boldsymbol{\sigma}} + \hat{\boldsymbol{\sigma}} \,. \tag{1.7b}$$

The (~) fields are those associated with the dislocation in an infinite medium and are obtained by evaluating analytic fields in a DDD simulation. While the corrective (^) fields are those which must be superimposed to ensure the boundary conditions are met. This means that the image fields can be obtained by running a FE simulation with the "corrected" displacement and traction fields,

$$\hat{\boldsymbol{u}} = \boldsymbol{u} - \tilde{\boldsymbol{u}} \,, \tag{1.8a}$$

$$\hat{\boldsymbol{\sigma}} \cdot \boldsymbol{n} = \hat{\boldsymbol{T}} = \boldsymbol{T} - \underbrace{\tilde{\boldsymbol{T}}}_{\tilde{\boldsymbol{\sigma}} \cdot \boldsymbol{n}}, \tag{1.8b}$$

where n is the outer unit normal vector to S. As a dislocation segment moves closer to the surface, its (\sim) field diverges and starts causing numerical problems [88]. A further problem with this method is that modelling elastic inclusions not only requires the calculation of forces induced by dislocations on the inclusion's surface, but also demands the calculation of so-called polarisation stresses due to differences in the inclusion's and matrix properties [87–89].

The relative simplicity of the superposition method has made it a popular choice [86, 90, 91] for coupling DDD and FEM because all it requires is the calculation of forces and displacements on the boundaries (see section 1.5.3).

1.5.1.2 Discrete-Continuum Model

The Discrete-Continuum Model (DCM) takes an alternative approach to solving the same problem. The DCM only treats short-range dislocation-dislocation interactions analytically while all other interactions are numerically calculated via FEM [92]. It is based on the regularisation of the atomic displacement jump across the slip plane into a plastic strain inclusion according to eigenstrain theory [93]. Like the Superposition Model, the DCM also assumes the simulated volume to be linear-elastic.

The eigenstrain formalism assumes that material defects can be represented as stress-free strain distributions dubbed eigenstrains [93]. For example, a dislocation loop of any shape may be approximately represented, thin, coherent, plate-like inclusion with the same contour as the loop and a characteristic thickness h as shown in fig. 1.8 [92]. The eigenstrain tensor, ϵ^{p} , can then be defined as

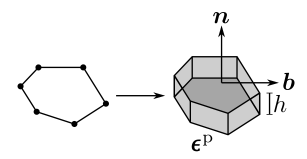


Figure 1.8: The eigenstrain formalism as defined in [93]. The dislocation is being approximated by a thin coherent inclusion of thickness h, whose strain-free stress tensor σ^{p} as defined by eq. (1.9). The vectors \boldsymbol{n} and \boldsymbol{b} are the normal vector to the slip plane and the dislocation line's Burgers vector.

a symmetric dyadic product of the Burgers vector, \boldsymbol{b} , and slip plane normal \boldsymbol{n} ,

$$\epsilon^{\mathrm{p}} \equiv \frac{1}{2h} (\boldsymbol{b} \otimes \boldsymbol{n} + \boldsymbol{n} \otimes \boldsymbol{b}).$$
 (1.9)

eq. (1.9) can be used to calculate approximate elastic fields from the stress-free eigenstrain distribution. The approximation is accurate far from the dislocation core [94]. As we move closer but still outside of the core region, the approximation tends toward the exact discrete dislocation solution as $h \to 0$. Due to linear elasticity, the total plastic strain is the sum of the individual plastic strains due to each dislocation segment.

The formalism then lets us solve the boundary value problem by finding the stress tensor σ , elastic strain ϵ^{e} and displacement u in mechanical equilibrium with the boundary conditions and plastic strain distribution ϵ^{p} from the "inclusions".

It is worth noting that in order to calculate the eigenstrains, there must be sufficient FE nodes inside the thin plate. Therefore smaller values of h necessitate finer FE meshes. Furthermore, the original DCM experienced numerical blow up as dislocation-dislocation distances approached h [95], but this has since been addressed. Assuming linear-elasticity, the revised model [92] yields,

$$\nabla \cdot \boldsymbol{\sigma} + \boldsymbol{f} = \boldsymbol{0} \quad \in V \setminus \{A\}, \tag{1.10a}$$

$$\sigma = \mathbf{E} : \epsilon \in V \setminus \{A\},$$
 (1.10b)

$$[[\boldsymbol{u}]]$$
 across $\{A\}$, (1.10c)

$$\boldsymbol{u} = \boldsymbol{u_0} \in S_u,$$
 (1.10d)

$$\boldsymbol{\sigma} \cdot \boldsymbol{n} = \boldsymbol{T} \quad \in S_T \,. \tag{1.10e}$$

At time t, $\{A\}$ denotes the area swept by the dislocation loops since the start of the simulation. [[u]] denotes displacement jumps tangent to $\{A\}$ due to dislocation glide; its magnitude and direction depend on the Burgers vector \boldsymbol{b} . \boldsymbol{E} is the 4^{th} -

order elasticity tensor, σ the small strain tensor, f are the body forces and, :, is the double dot product defined as, $(\mathbf{E}: \boldsymbol{\epsilon})_{ij} = E_{ijkl}\epsilon_{kl}$, for a rank 4 tensor \mathbf{E} and rank 2 tensor ϵ . The operator, \, is the set difference defined as, $A \setminus B = \{x \in A | x \notin B\}$. We use the same notation for the volume, surface normal and boundary conditions as in section 1.5.1.1. eq. (1.10) can then be linearly decomposed three parts which are solved via FEM or DDD and coupled.

It is worth noting that the DCM is substantially more complicated than the SM. By requiring the finite elements be small enough for the eigenstate formalism to work, it strongly couples DDD to the FE model and software. It shifts the brunt of the computational workload from DDD to FEM. Essentially trading computationally intensive long range interactions which scale on the order of $\mathcal{O}(N^2)$, where N is the number of dislocation line segments; for computationally intensive tasks scaling on the order of $\mathcal{O}(M^3)$ where M is the number of finite elements in a cubic mesh with N elements per side. Furthermore, it cannot be used with BE methods as the eigenstrain formalism demands the use of internal elements rather than simply requiring a surface mesh.

That said, the DCM allows for reductions in the computational complexity of certain parts of the problem which would otherwise have to be done via DDD [92]; namely long-range dislocation-dislocation or dislocation-surface interactions via an interaction distance cutoff that is only possible with the eigenstrain formulation. Compared to the SM, gains in computational efficiency grow as dislocation density increases with respect to the number of finite elements. Since dislocationdislocation and dislocation-surface interactions are among the most computationally expensive aspects of DDD, the DCM can reduce the overall computational cost of simulations with large enough dislocation densities.

Level Set Method 1.5.1.3

Even though the level set method has not strictly been used to couple DD to FEM, it has been used to model inclusions [96] as described in section 1.5.4.2. This type of dislocation dynamics is fundamentally distinct from DDD because Level Set DD uses arbitrary functions rather than a discretisation approach to represent dislocations lines. The idea is that in 3D, a dislocation $\gamma(t)$ can be represented by the intersection of two zero levels of two level set functions (see fig. 1.9),

$$\phi(x(t), y(t), z(t), t) = 0, \qquad \psi(x(t), y(t), z(t), t) = 0, \qquad (1.11a)$$

$$\phi(x(t), y(t), z(t), t) = 0, \qquad \psi(x(t), y(t), z(t), t) = 0, \qquad (1.11a)$$

$$\frac{d\phi}{dt} = \partial_t \phi + \boldsymbol{v} \cdot \nabla \phi = 0, \qquad \frac{d\psi}{dt} = \partial_t \psi + \boldsymbol{v} \cdot \nabla \psi = 0, \qquad (1.11b)$$

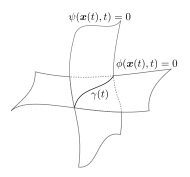


Figure 1.9: Diagram of a continuous dislocation $\gamma(t)$ as the intersection of the zero level of two level set functions $\phi(\boldsymbol{x}(t),t),\ \psi(\boldsymbol{x}(t),t)$ as posed by eq. (1.11). Image edited from [97].

where v is the dislocation velocity. This definition uses the material derivative because it is assumed that the function and its spatial coordinates are all functions of time, t. The material derivative also comes up when deriving the Navier-Stokes equations of fluid dynamics.

The level set method is significantly more computationally more expensive than DDD [97]. It must solve two coupled quasi-linear partial differential equations whose character is in general undefined. It therefore requires the use of very robust numerical solvers such as higher order Total Variation Deminishing (TVD) Runge-Kutta methods for time discretisation (TVD-RK4 or higher) and high order ENO (Essentially Nonoscillatory $\mathcal{O}(N^5)$ or higher) or WENO (Weighted Essentially Non-Oscilatory) interpolation methods for spatial discretisation [98] for the solutions to converge satisfactorily. The method also makes it impossible to obtain many of the useful analytical solutions one can find by discretising dislocations into straight line segments. Another problem is that this method has never been truly coupled to the finite element method. However, if one were to do so there are two naïve ways of going about it: 1) numerically integrating along the dislocation lines and finding a way to numerically calculate the forces on FE nodes, or 2) discretising the level set curves and using either the DCM or SM to find forces and displacements. The former would be very computationally expensive. Furthermore, no one has found a way to compute the forces or displacements exerted by generally shaped, non-discretised dislocations, numerical solutions are possible as they are needed to find self-interaction energies and dislocation-dislocation forces [97] but general, closed-form analytical ones are most likely impossible to find. The latter defeats the purpose of the level set method by abandoning the continuum framework. As a result of this, the level set method is not nearly as popular as DDD utilising either the SM or DCM.

1.5.2 Non-Singular Continuum Theory of Dislocations

The Peach-Koehler formula describes the fundamental interactions of dislocations [99] according to the force, \boldsymbol{f} , that a local stress $\boldsymbol{\sigma}$ exerts on a dislocation line with Burgers vector, \boldsymbol{b} , and line direction, $\boldsymbol{\xi}$,

$$\mathbf{f} = (\boldsymbol{\sigma} \cdot \boldsymbol{b}) \times \boldsymbol{\xi} \,. \tag{1.12}$$

According to [100], the internal stress field of a dislocation loop in a homogenous, infinite, linear-elastic medium is given by the contour integral around a loop L,

$$\sigma_{ij}^{\infty}(\boldsymbol{x}) = C_{ijkl} \oint_{L} \epsilon_{lnh} C_{pqmn} \frac{\partial G_{kp}(\boldsymbol{x} - \boldsymbol{x}')}{\partial x_{q}} b_{m} \, \mathrm{d}x'_{h}, \qquad (1.13)$$

where C_{ijkl} is the elastic stiffness tensor, ϵ_{lnh} the permutation operator, \boldsymbol{b} the Burgers vector, and $G_{kp}(\boldsymbol{x}-\boldsymbol{x}')$ is Green's function of elasticity [100]. $G_{kp}(\boldsymbol{x}-\boldsymbol{x}')$ is defined as the displacement component in the x_k direction at point \boldsymbol{x} in response to a unit point force applied in the x_p direction at point \boldsymbol{x}' [101]. In an isotropic elastic solid, $G(\boldsymbol{x}-\boldsymbol{x}')$, takes the form,

$$G_{ij}(\boldsymbol{x} - \boldsymbol{x}') = \frac{1}{8\pi\mu} \left[\delta_{ij} \partial_{pp} - \frac{1}{2(1-\nu)} \partial_{ij} \right] R(\boldsymbol{x} - \boldsymbol{x}'), \qquad (1.14)$$

where μ , ν are the isotropic shear modulus and Poisson's ratio respectively, δ_{ij} is the Kronecker Delta, $\partial_{x_1...x_n} \equiv \frac{\partial^n}{\partial x_1...\partial x_n}$, and $R = \|\boldsymbol{x} - \boldsymbol{x}'\|$. However considering that $\boldsymbol{x} \to \boldsymbol{x}' \Rightarrow R \to 0 \Rightarrow \partial_i R \to \infty$, some (or all) components of the stress field diverge. Furthermore, the total elastic energy also diverges,

$$E = \frac{1}{2} \int S_{ijkl} \sigma_{ij}(\boldsymbol{x}) \sigma_{kl}(\boldsymbol{x}) d^3 \boldsymbol{x}, \qquad (1.15)$$

where $S = C^{-1}$ is the elastic compliance tensor. These divergent properties often prove problematic when numerically computing dislocation forces and energies. Equation (1.15) can also be expressed as a double line integral [102, 103],

$$E = -\frac{\mu}{8\pi} \iint_{L} \partial_{k} \partial_{k} R(\boldsymbol{x} - \boldsymbol{x}') b_{i} b'_{j} \, dx_{i} \, dx'_{j}$$

$$-\frac{\mu}{4\pi(1-\nu)} \iint_{L} \partial_{i} \partial_{j} R(\boldsymbol{x} - \boldsymbol{x}') b_{i} b'_{j} \, dx_{k} \, dx'_{k}$$

$$+\frac{\mu}{4\pi(1-\nu)} \iint_{L} \partial_{k} \partial_{k} R(\boldsymbol{x} - \boldsymbol{x}') b_{i} b'_{i} \, dx_{j} \, dx'_{j}$$

$$-\nu \iint_{L} \partial_{k} \partial_{k} R(\boldsymbol{x} - \boldsymbol{x}') b_{i} b'_{j} \, dx_{j} \, dx'_{i}, \qquad (1.16)$$

which is important when describing how Cai et al. [101] derived their non-singular expression.

The singularity in R is the result of unreasonably and unphysically assuming a dislocation loop's Burgers vector distribution is a Delta function. The assumption was made to allow for closed-form and relatively simple expressions. As noted in [104], other distributions may be used but they either result in significantly more complicated expressions or destroy the analytical nature of the classical formulation.

There have been many attempts at removing this singularity, including finite-strain elasticity [105], non-local and gradient elasticity [106, 107], interaction cut-off radius [99], average stress at two points on opposite sides of the dislocation line [108, 109], and spreading the Burgers vector distribution out over a finite width [104, 110, 111]. Unfortunately all of these approaches failed in one way or another [101]. Depending on the approach, the following undesirable qualities may present themselves include inconsistencies, impractical implementation, lack of closed-form solutions for non-straight finite dislocations, lack of self-consistency, and the possibility for multiple expressions and solutions for the line integral of the dislocation line energy.

Cai et al. [101] took it upon themselves to define and justify a different Burgers vector distribution that maintains the mathematical convenience of the classical formulation that also eliminates such an unphysical assumption. They did so by introducing a Burgers vector density function,

$$\boldsymbol{b} = \int \boldsymbol{g}(\boldsymbol{x}) \, \mathrm{d}^3 \boldsymbol{x} \,, \tag{1.17a}$$

$$g(x) = b\tilde{w}(x) = b\tilde{w}(r), \qquad (1.17b)$$

where $r \equiv ||x||$. When substituting eq. (1.17a) into eqs. (1.16) and (3.2) the result of multiplying R with components of \boldsymbol{b} results in the following integrals,

$$R(\boldsymbol{x} - \boldsymbol{x}')b_m = \int R(\boldsymbol{x} - \boldsymbol{x}'')g_m(\boldsymbol{x}'' - \boldsymbol{x}') d^3\boldsymbol{x}'', \qquad (1.18a)$$

$$R(\boldsymbol{x} - \boldsymbol{x}')b_m b'_n = \iint R(\boldsymbol{x}'' - \boldsymbol{x}''')g_m(\boldsymbol{x} - \boldsymbol{x}'')g_n(\boldsymbol{x}''' - \boldsymbol{x}') d^3\boldsymbol{x}'' d^3\boldsymbol{x}'''. \quad (1.18b)$$

Using eqs. (1.17) and (1.18) as guidelines, they defined a the following convolutions,

$$w(\boldsymbol{x}) \equiv \tilde{w}(\boldsymbol{x}) * \tilde{w}(\boldsymbol{x}) = \int \tilde{w}(\boldsymbol{x} - \boldsymbol{x}') \tilde{w}(\boldsymbol{x}') d^3 \boldsymbol{x}',$$
 (1.19a)

$$R_a \equiv R(\boldsymbol{x}) * w(\boldsymbol{x}) = \int R(\boldsymbol{x} - \boldsymbol{x}') w(\boldsymbol{x}') \, d^3 \boldsymbol{x}'. \qquad (1.19b)$$

At which point they assumed there exists an integrable function w(x) such that

$$R_a = \sqrt{R(\mathbf{x})^2 + a^2} = \sqrt{x^2 + y^2 + z^2 + a^2},$$
 (1.20)

where a is an arbitrary constant meant to represent the dislocation core radius whose value may be estimated from atomistic simulations. This is essentially a definition that can be used to replace R in the classical equations and eliminate the singularity by including a free parameter. However, in order to ensure this is mathematically sound, there must indeed exist a function which yields R_a as Cai et al. [101] defined it. This can be done by making use of the following property for the convolution of two suitably differentiable functions, $\partial_i(f*g) = \partial_i f*g = f*\partial_i g$. We may take the Laplacian twice,

$$\nabla^2[\nabla^2\{R(\boldsymbol{x}) * w(\boldsymbol{x})\}] = \nabla^2[\nabla^2 R_a(\boldsymbol{x})], \qquad (1.21a)$$

$$\nabla^2[\nabla^2\{R(\boldsymbol{x})\}] * w(\boldsymbol{x}) = \nabla^2[\nabla^2 R_a(\boldsymbol{x})], \qquad (1.21b)$$

$$\nabla^2[\nabla^2\{R(\boldsymbol{x})\}] = \nabla^2\left[\frac{2}{R}\right] = -8\pi\delta^3(\boldsymbol{x}), \qquad (1.21c)$$

$$\nabla^2 [\nabla^2 R_a(\mathbf{x})] = \nabla^2 \left[\frac{2}{R_a} + \frac{a^2}{R_a^3} \right] = -\frac{15a^4}{R_a^7}, \quad (1.21d)$$

$$w(\mathbf{x}) = \frac{15a^4}{8\pi R_a^7}. (1.21e)$$

Equation (1.21c) is a very brave, handway statement given that,

$$\nabla^2 \left[R^{-1} \right] = \nabla^2 \left[(x^2 + y^2 + z^2)^{-1/2} \right] = 0, \tag{1.22}$$

but may be physically "justified" by pretending $\nabla^2 [R^{-1}]$ defines a spherical surface of radius 0. And eq. (1.21e) is a similarly handwavy statement that can be "justified" by noting that,

$$\lim_{a \to 0} w(\mathbf{x}) = \delta^3(\mathbf{x}). \tag{1.23}$$

Nevertheless, the non-singular formulation by Cai et al. [101] fixes the physically and mathematically problematic assumption that Burgers vectors follow 3D Dirac delta distributions, and proves useful in producing analytical expressions (see section 1.5.3) that are not much more complex than the singular case.

1.5.3 Analytical Forces Exerted by a Dislocation Line Segment on Surface Elements

Whether using the SM or DCM, coupling DDD to FEM requires the traction field $\sigma^{\infty}(x) \cdot n(x)$ to be distributed among the set of relevant discrete nodes of a FE or BE model. In the DCM model this applies to dislocations that are sufficiently close to the boundary; while in the SM, it applies to all dislocations. Regardless of the coupling model, the force exerted by a dislocation ensemble on a node n on element e is given by,

$$\mathbf{F}^{(n)} = \int_{S_e} \left[\boldsymbol{\sigma}^{\infty}(\boldsymbol{x}) \cdot \boldsymbol{n}(\boldsymbol{x}) \right] N_n(\boldsymbol{x}) \, dS_e \,, \tag{1.24}$$

where dS_e is the infinitesimal surface element with surface area S_e . $N_n(\boldsymbol{x})$ are so-called shape functions (interpolation functions) that distribute the traction field among the surface element's nodes.

The problematic singularity associated with the classical Volterra dislocation is avoided by using the non-singular formulation of Cai et al. [101] discussed in section 1.5.2, which changes eq. (3.3) into eq. (1.25),

$$G_{ij}(\boldsymbol{x} - \boldsymbol{x}') = \frac{1}{8\pi\mu} \left[\delta_{ij} \partial_{pp} - \frac{1}{2(1-\nu)} \partial_{ij} \right] R_a(\boldsymbol{x} - \boldsymbol{x}').$$
 (1.25)

Using the non-singular definition of $G(\mathbf{x} - \mathbf{x}')$ in eq. (3.2) we obtain the expression for the stress field of a single straight dislocation line segment bounded by two dislocation nodes (1, 2) [101],

$$\boldsymbol{\sigma}^{(12)}(\boldsymbol{x}) = -\frac{\mu}{8\pi} \int_{\boldsymbol{x_1}}^{\boldsymbol{x_2}} \left(\frac{2}{R_a^3} + \frac{3a^2}{R_a^5} \right) [(\boldsymbol{R} \times \boldsymbol{b}) \otimes d\boldsymbol{x}' + d\boldsymbol{x}' \otimes (\boldsymbol{R} \times \boldsymbol{b})]$$

$$+ \frac{\mu}{4\pi(1-\nu)} \int_{\boldsymbol{x_1}}^{\boldsymbol{x_2}} \left(\frac{1}{R_a^3} + \frac{3a^2}{R_a^5} \right) [(\boldsymbol{R} \times \boldsymbol{b}) \cdot d\boldsymbol{x}'] \mathbf{I}_2$$

$$- \frac{\mu}{4\pi(1-\nu)} \int_{\boldsymbol{x_1}}^{\boldsymbol{x_2}} \frac{1}{R_a^3} [(\boldsymbol{b} \times d\boldsymbol{x}') \otimes \boldsymbol{R} + \boldsymbol{R} \otimes (\boldsymbol{b} \times d\boldsymbol{x}')]$$

$$+ \frac{\mu}{4\pi(1-\nu)} \int_{\boldsymbol{x_1}}^{\boldsymbol{x_2}} \frac{3}{R_a^5} [(\boldsymbol{R} \times \boldsymbol{b}) \cdot d\boldsymbol{x}'] \boldsymbol{R} \otimes \boldsymbol{R}, \qquad (1.26)$$

Substituting eq. (3.5) into eq. (3.1) yields as many integrals as there are surface nodes. Their exact form depends on the shape functions used. The specifics are out of the scope of this work. For a rectangular surface element, the vector formulation can be converted into a set of local scalar coordinates via a series of

scalar projections of the set of basis vectors via eq. (1.27),

$$R = x - x' = yt + rp + sq, R_a = \sqrt{R \cdot R + a^2},$$

$$n = p \times q, y = \frac{R \cdot n}{t \cdot n}, (1.27)$$

$$r = \frac{R \cdot (q \times t)}{p \cdot (q \times t)}, s = \frac{R \cdot (p \times t)}{q \cdot (p \times t)},$$

$$dx' = -t dy,$$

where n, p, q, t are basis vectors³ that respectively correspond to the outside surface normal of the surface element, the two orthogonal vectors that define the sides of the rectangular surface element, and the dislocation line direction. The integral bounds can then be set up accordingly.

Noting that $\mathbf{n} \equiv \mathbf{p} \times \mathbf{q}$ is normal to the surface element, if \mathbf{t} is parallel to the surface element i.e. a linear combination of \mathbf{p} , \mathbf{q} then the product $\mathbf{t} \cdot \mathbf{n} = 0$, so this formulation is problematic when a dislocation line segment is parallel to a surface. An easy workaround is to slightly around the axis defined by $\mathbf{t} \times \mathbf{n}$ using the line's midpoint as a fulcrum. This works well in serial code but is problematic for parallelisation due to warp divergence (see section 1.5.5).

Figure 3.1 diagramatically shows the definition of the four basis vectors and summarises the line integrals needed to find an analytical expression of the nodal force. It is worth noting that over 40 triple integrals arise from eqs. (1.27) and (3.5). These were solved via three families of recurrence relationships found via integration by parts and six seed functions found by direct integration. A more detailed description can be found in [86].

Quadratic triangular surface elements pose a much more challenging problem due to the fact that $p \not\perp q$. This property is not the only addition that significantly complicates the integrals, the interpolation functions are quadratic which increase the number of integrals. The non-orthogonality of q, p means the integration limits in the r, s coordinates must be parametrised appropriately and the integrations should be carried out in an order suitable for the chosen parametrisation. Aside from these technical considerations, the techniques for solving the problem for quadratic rectangular surface elements are the same as the linear rectangular case [86]. It is worth noting however, that the implementation of such a procedure may require quadruple precision arithmetic.

³Unitary vectors that define a space. This space is not orthogonal because in general only $p \perp q$, $n \perp p$ and $n \perp q$.

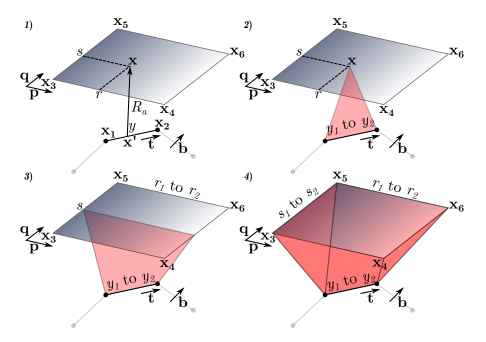


Figure 1.10: Diagram of the line integral method used to find analytical expressions for the forces exerted by dislocation on linear rectangular surface elements [86]. 1) For any given point \boldsymbol{x} on the surface element and any given point \boldsymbol{x}' on the dislocation line segment, define distance R_a . 2) Integrate from $x_1 \to x_2$ along line direction \boldsymbol{t} . 3) Integrate from $r_1 \to r_2$ along vector \boldsymbol{p} . 4) Integrate from $s_1 \to s_2$ along vector \boldsymbol{q} .

1.5.4 Multiphase Simulations

Modelling heterogeneity in DDD remains challenging. The core assumption of DDD is that we have a homogenous linear elastic space with invariant properties. However, modelling multiphase systems such as multicrystals and inclusions necessitates a weakening of such assumptions, particularly the space's homogeneity. Multiphase models are also necessary to apply DDD to more realistic scenarios.

Fusion in particular is riddled with examples of highly heterogenous materials. As previously mentioned, neutron bombardment often causes clustering of transmutation products particularly on divertor and first wall materials; such clusters often substantially change many of their mechanical and thermal properties. The whole point of Oxide-Dispersion Strengthened (ODS) steels [112] is to hamper radiation damage by distributing oxide inclusions within their matrix. If we hope to model such alloys with DDD we must find a way to relax our homogeneity criteria.

Furthermore, many potential candidates for structural and even first wall materials of fusion reactors are designed with complex microstructures in order to resist thermal creep and radiation damage. Understanding dislocation behaviours near the crystal boundaries of these microstructurally complex materials is another niche where DDD simulations can shed light into the mechanisms that give

rise to these materials' desirable properties.

1.5.4.1 Polycrystalline Materials

Polycrystalline materials have been studied using DDD, however most of these are in 2D. One of the easiest phenomena to investigate with 2D DDD are grain size effects [113, 114].

It is well known that much of the grain size strengthening effects arise from dislocation-grain boundary (GB) interactions including transmission, reflection, emission and absorption of dislocations [115, 116]; with transmission being the most typically observed.

The model depends on the grain boundary energy density, $E_{GB}(\delta\theta)$, where $\delta\theta$ is the crystallographic misorientation between neighbouring crystals; resolved shear stress, τ , on the incoming dislocation with Burgers vector $\boldsymbol{b_1}$; critical penetration stress for the GB, τ_{GB} ; and Burgers vector of the dislocation debris, $\boldsymbol{\Delta b} = \boldsymbol{b_1} - \boldsymbol{b_2}$, left behind when the incoming dislocation, $\boldsymbol{b_1}$, transmits throught the grain boundary to become a dislocation with Burgers vector, $\boldsymbol{b_2}$. The relationship may be approximated by,

$$\tau |\boldsymbol{b}_{1}|^{2} \ge \tau_{\text{pass}} |\boldsymbol{b}_{1}|^{2} = E_{\text{GB}}(\theta) |\boldsymbol{b}_{1}| + \alpha G |\boldsymbol{\Delta} \boldsymbol{b}|^{2}, \qquad (1.28)$$

where α is the material constant and G the shear modulus. The grain boundary energy density was proposed by [117] to have the simple form,

$$E_{\rm GB} = \begin{cases} k \frac{\delta \theta}{\theta_1} & 0 \le \delta \theta < \theta_1, \\ k & \theta_1 \le \delta \theta < \theta_2, \\ k \frac{\pi/2 - \delta \theta}{\pi/2 - \theta_2} & \theta_2 \le \delta \theta < \pi/2, \end{cases}$$
(1.29)

where k, θ_1 , θ_2 , are material specific.

This is of course a gross simplification of the real 3D problem, but this approach was used by Li et al. [113] to investigate the Hall-Petch effect, which correlates grain size with flow stress of a polycrystal,

$$\sigma = \sigma_0 + \kappa \left(\frac{d_0}{d}\right)^n \,, \tag{1.30}$$

where σ_0 is the yield stress, d_0 a reference crystal size, κ the Hall-Petch slope and n the crystal size sensitivity parameter. Li et al. [113] showed that the model reproduces the Hall-Petch effect quite successfully. As a consequence, they showed that even what might appear as an overly simplistic approach can describe a complex emergent phenomenon such as the Hall-Petch effect.

Aside from the Hall-Petch effect, the model has also been utilised by [114] to study the thickness effects of three different types of polycrystalline thin films: 1) no surface treatment, 2) surface passivation layer and 3) surface grain refinement zones. In this study, Frank-Reed sources were seeded across the domain, and the superposition principle was utilised to calculate displacement, strain and stresses in the thin films. Their results qualitatively reproduced experimental observations from expected dislocation patterns, stress distributions, and the eventual disappearance of the size effect as the films got thicker [114].

Two-dimensional models might seem useless but they have their place, particularly when thin films are concerned. However, they may also be of use when investigating the effects of dislocations on the mechanical behaviour of superconducting tape—whose applications range from medical imaging to fusion energy production. The exotic composition and crystallography of many superconductors (perovskites) would make 3D models very complex and computationally expensive, so 2D models may offer viable alternatives. On top of this, a superconducting tape's operating environment would often have it under strains which might not strictly lie in-plane, but given a small enough segment of tape, strains orthogonal to the its plane may be neglected. Thus making 2D models an acceptable first attempt at tacking the problem, at leas before computational resources and developments in 3D models make more realistic studies of such complex systems possible.

The 3D case is substantially more complicated than the 2D case. A recent study by Fan et al. [118] looked at the role of twinning on the hardening response of polycrystalline Mg using 3D DDD.

The article is a perfect example of why 3D DDD is so much more complex than 2D DDD. Admitedly, it uses a material with a HCP crystal structure which complicates matters compared to FCC or BCC.

In 3D DDD one must account for dislocation-twin boundary (TB) interactions. These may be obtained via geometric considerations, MD, and experimental evidence [119–122]. We must have information regarding how dislocations are transmitted through a TB such as 1) whether a dislocation leaves a residual dislocation on the TB, 2) the possible pre-TB and post-TB slip planes a dislocation can be transmitted to, 3) which loading conditions are conducive to which transmission behaviour and 4) which types of dislocation can be transmitted or reflected and in what ways [118]. All this obviously depends on the twinning plane, so in order to study more realistic scenarios one must have all the necessary knowledge to properly account for all dislocation-TB interactions. Furthermore, it is possible that certain dislocations may leave behind twinning dislocations that end up as

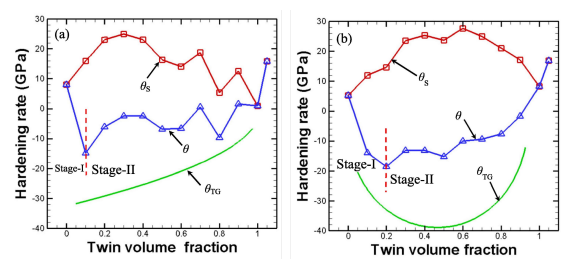


Figure 1.11: Simulated hardening rate as a function of twin volume fraction for (a) yz compressive loading and (b) tensile loading. Image edited from [118].

TB steps, whose movement can lead to TB migration and twin growth [123, 124].

Fan et al. [118] modelled four scenarios in order to deconvolute the effects that twins and grain boundaries have on the mechanical behaviour of a cubic crystal. They modelled 1) a cubic twinned polycrustal with two TBs and GBs on the edges of the cube, 2) a single crystal with no TBs or GBs, 3) a polycrystal with no TBs but GBs, 4) a twinned crystal with TBs but not GBs. It is worth noting that the effects of twin growth in plasticity were not accounted for in their DDD simulation and instead had to be numerically computed via the hardening rate θ ,

$$\theta = \frac{d\sigma}{d\epsilon} = \frac{d}{d\epsilon} \left[E \left(\epsilon - \epsilon_{\text{slip}}^{\text{p}} - \epsilon_{\text{twin}}^{\text{p}} \right) \right] = \theta_{\text{s}} + \theta_{\text{TG}}, \qquad (1.31a)$$

$$\theta_{\rm TG} = -\frac{\mathrm{d}}{\mathrm{d}\epsilon} \left[E \epsilon_{\rm twin}^{\rm p} \right] = -E \bar{m} \gamma_{\rm twin} \frac{\mathrm{d}}{\mathrm{d}\epsilon} \left[f_{\rm twin} \right] , \qquad (1.31b)$$

where θ_{TG} is the hardening rate due to twin growth, \bar{m} the mean Schmid factor of the participating twins γ_{twin} the characteristic twinning shear and $\frac{d}{d\epsilon}[f_{\text{twin}}]$ is the change in the change in the twin volume fraction with respect to applied strain. These parameters must be empirically obtained.

With their model, Fan et al. [118] managed to qualitatively reproduce experimental observations, including the concave shape of strain-stress curves that arises from the competing hardening effect of TBs restricting dislocation motion (fig. 1.11), and the softening produced by twin growth as obtained from eq. (1.31b).

Twin boundaries in 3D are analogous to grain boundaries in 2D, given that they are coherent, narrow boundaries rather than messy, often incoherent grain boundaries. Consequently, grain boundaries are much harder to treat in 3D; instead, the community focuses on multiphase models where the details of dis-

location transmission between phases are ignored or simplified as mentioned in section 1.5.4.2.

1.5.4.2 Inclusions

Inclusions have been modelled using the three methods described in section 1.5.1. However, they have gone about it differently. The SM and DCM apply their formulations heirarchically to allow for dislocation motion into and out of the inclusion. Furthermore, being coupled DDD-FEM problems, the inclusions must also be appropriately meshed, oftentimes this means inclusion geometries are limited to the mesh type. However, FEM allows for the calculation of coherency stresses which are seen by the dislocations and cause dislocation localisation and forresting in the vicinity of inclusions, just as experimentally observed [125, 126].

The level set method, having never been coupled to FEM, takes a different approach. Where the particles can be modelled as a region where dislocations experience a user-defined force. Coherency stresses have so far been neglected, but their effects may be modelled with the use of functions akin to diffuse orbital functions used in theoretical and computational quantum chemistry. Within this framework the inclusions may be arbitrarily defined to have any smooth shape, especially if one were to use piecewise parametric fourier curves.

Dislocation-permeable inclusions have been simulated via the SM [125]. As mentioned in section 1.5.1.1, there is a need to calculate polarisation stresses due to differences in the elasticity of both phases. Using the same notation as section 1.5.1.1 this can be done by breaking up the image stress into,

$$\hat{\boldsymbol{\sigma}} = {}^{\mathrm{m}}\boldsymbol{C}\hat{\boldsymbol{\epsilon}} \quad \in V_{\mathrm{m}} \tag{1.32a}$$

$$\hat{\boldsymbol{\sigma}} = {}^{\mathrm{p}}\boldsymbol{C}\hat{\boldsymbol{\epsilon}} + [{}^{\mathrm{p}}\boldsymbol{C} - {}^{\mathrm{m}}\boldsymbol{C}]\,\tilde{\boldsymbol{\epsilon}} \in V_{\mathrm{p}},$$
 (1.32b)

where m, p denote whether a variable belongs to the matrix or precipitate respectively and \boldsymbol{C} is the elastic stiffness tensor.

Yashiro et al. [125] not only modelled a cuboid inclusion, but bimetallic interfaces in 3D. They found that under the right conditions, dislocations can pass from one phase to the other. When a dislocation passes into a different phase, it leaves an antiphase boundary on the slip plane. Therefore, in order for a dislocation to move from one phase to another, it needs excess energy equal to the antiphase (APB) energy of the area it sweeps within the precipitate. This means that as a node passes from one phase to another, it feels a repulsive force F_b . The next dislocation moving along the same APB will then feel an attractive force F_b to dissolve the APB. Once both dislocations move into the new matrix, they form a superdislocation bound by the APB energy [127]. However, as the disloca-

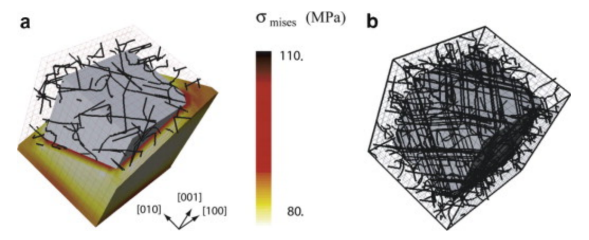


Figure 1.12: (a) Shows the initial dislocation configuration after relaxation in the presence of the calculated von Mises coherency stresses σ_{mises} . (b) Shows the dislocation forresting around the inclusion as a result of 0.2% plastic strain for the [001] case. Image obtained from [126].

tion length increases with respect to the precipitate's volume, the production of Orowan loops around it becomes more energetically favourable than moving into the new phase [125, 127]. Both of these behaviours have been observed when using the superposition method.

The eigenstrain method [128] as described in section 1.5.1.2 has been shown to be a viable solution to this problem. One can compute the stresses and displacements produced by the dislocation ensemble on the inclusions' surfaces via DCM or SM. The SM method however needs to calculate polarisation stresses [88], an operation that significantly increases the number of calculations necessary to solve the problem.

The DCM method has been used to simulate multiphase materials where the simulated dislocations reproduced experimentally observed behaviours such as the zig-zag patterns and dislocation forests produced by dislocations around precipitates in nickel superalloys [95, 126]. Reproducing such behaviours is not as trivial as simply adding inclusions to the simulation domain. In order to produce accurate results, one requires knowledge of the coherency stress between the different phases [126]. Such stresses are often due to lattice mismatches and differences in the thermal expansion coefficients. These coherency stresses localise the dislocations around the inclusions and are responsible for the observed localisation and aggregation of dislocations around precipitates as seen in fig. 1.12 [126].

The level set method has also been successfully used to model inclusions. In contrast to the aforementioned methods, they are assumed to be regions in space where dislocations experience a repelling force, instead of a region in space with distinct characteristics to the matrix. The force profile can be very simply defined to depend on the radial distance from the dislocation to a point in the "inclusion's"

volume [96]. It can also be modulated depending on the nature of the inclusion by making the force a radial function, thus avoiding many of the numerical issues facing DDD-FEM couplings (see sections 1.5.1 and 1.5.3). It is also possible to make the inclusions impermeable or semi-permeable depending on the magnitude and steepness of the force gradient as presented in [96]. Arbitrarily shaped may also be possible if one were to also use angular components to the force function, or even through the use of spherical harmonics.

Though mathematically appealing, the level set method is far from representing the reality of dislocation-inclusion interactions. More realistic scenarios require use of more commonplace DDD-FEM coupling methods. Implementing arbitrarily shaped particles is not impossible in DDD—particularly if they can be discretised. Aside from the significantly lower computational requirements, one of the biggest advantages of DDD is that one can find tractions and displacements natively through either the DCM or SM, whereas the level set method requires the force to be user-defined and displacements on the inclusion are completely ignored. Furthermore, if the inclusion is a different crystal or a twin of the same material as in [118], it is possible to use the DCM or SM to study the transmission of dislocations from one phase to the other; something that is not yet possible to do with the level set method.

1.5.5 Parallelising Discrete Dislocation Dynamics

It has been shown that DDD lends itself well to parallelisation. Ferroni et al. [85] investigated some of the more computationally intensive parts of DDD in DDLab [129], a freely available DD code for Matlab. The algorithms whose parallel performance was studied in [85] were 1. segment-segment interaction, 2. surface tractions and 3. image stresses.

The segment-segment interaction has $\mathcal{O}(N^2)$ computational complexity, where N is the number of dislocation segments. The quadratic scaling is due to the r^{-1} dependence of the dislocation stress fields, so long range interactions cannot be ignored.

Both the surface traction and image stress calculations have $\mathcal{O}(kN)$ computational where k is the total number of surface nodes and N the number of dislocation line segments.

As is usually the case in parallel computing, there is often more than one way to parallelise a problem. The optimal strategy depends on the problem itself. It was no different in [85] as shown in fig. 1.13. Ferroni et al. [85] realised that the segment-segment interaction calculation can be done via $\mathcal{O}(N^2)$ parallelism, where single pairs of dislocation segments are sent to a single thread, and then globally

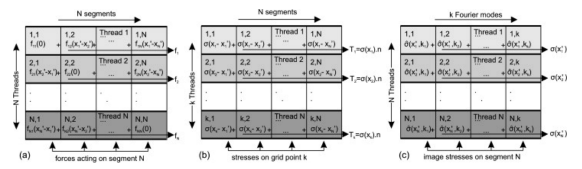


Figure 1.13: Parallelisation strategies for: (a) the N^2 segment interactions; (b) surface traction at k grid points and; (c) and image stresses on N segments. Image taken from [85].

reduced. However this approach requires $\mathcal{O}(N^2)$ memory and can lead to inefficient data reuse. So they opted instead for $\mathcal{O}(N)$ parallelisation and serialisation. In this scheme each thread is assigned a unique segment, and each thread computes the forces between its assigned segment and all the others while serially adding the force contributions. The memory access pattern they found best was that of a serial procedure, the reason is that in-thread calculations are serialised so having coalesced memory access patterns for threads would cause problems when serially accessing data for the calculation.

However, there is another approach Ferroni et al. [85] did not consider. It combines both of the aforementioned strategies, $\mathcal{O}(N^2)$ memory and $\mathcal{O}(N)$ parallelisation and serialisation. Such an approach requires two copies of the input data (one per access pattern). One pattern would allow each thread to obtain its uniquely assigned segment in a coalesced manner. The second would be the serial access pattern already present. Thus obtaining the best performance possible performance at the cost of doubling the required memory. This is a better strategy provided there is enough device memory⁴, if that is not so, the problem must be split into different GPU calls.

New NVidia architectures have made varying degrees of nested parallelisation possible, NVidia calls this "dynamic parallism". When applied to the computationally optimal solution, the $\mathcal{O}(N^2)$ memory requirement is relaxed to $\mathcal{O}(N)$ (with the same parallel-specific data structure) and $\mathcal{O}(N^2)$ computational parallelism.

The parallisation strategies for the other two problems, (b) and (c) in fig. 1.13,

⁴Assuming the line direction is calculated inside the program, each dislocation line segment represents 3, 3-element vectors of type double (8-bytes per entry), totalling 72 bytes in global memory per dislocation line segment (two nodes + Burgers vector per segment). There must also be enough global memory to output the forces, which represent 1, 3-element vector of type double per dislocation node, totalling 24 bytes per node. Assuming a single dislocation loop with N nodes (therefore N segments), this brings the total global memory requirement to 96N bytes. Using two access patterns for the dislocation line segments and Burgers vector means an extra 72N bytes, giving a grand total of 168N bytes. Assuming the material parameters are stored in cosntant memory.

involved similar decompositions as the segment-segment interaction. For surface tractions Ferroni et al. [85] parallelised over k surface nodes with sequential calculation and addition of the stress from each dislocation line segment. For the image stresses, they chose the reverse strategy since it allows for the serialised addition of the image stresses on dislocation line segments.

Their findings showed very promising results for parallelising DDD simulations. Compared to C, their findings showed that depending on the specific problem and parameters used, the parallel implementation reduced computational time by a factor of ~ 110 at best and ~ 3 at worst for sufficiently large problems. This is of course, provided one uses a card designed for high performance computing, as these are tailored towards double precision arithmetic and computationally intensive procedures. However, gaming graphics cards are increasingly capable of double precision arithmetic and though not as drastic, speed ups are still substantial.

Chapter 2

Coupling Discrete Dislocation Dynamics to Finite Element Methods

2.1 Superposition Scheme

Coupling Discrete Dislocation Dynamics (DDD) to a Finite Element Method (FEM) [86] is important to properly simulate micromechanical tests because DDD provides us with a more precise set of inputs and greater granularity for solving the FE problem. This can be achieved by using a so-called superposition scheme fig. 2.1 that enables the independent solution of both problems, whilst feeding information from one to the other in a continuous feedback loop.

2.2 Extracting Surface Nodes

The FEM coupler arranges the nodes starting on the xz-plane where $y \in \{0 \rightarrow n \ dy\}$ and $n \in \mathbb{N}$. However in order to couple DDD to FEM we only require the surface nodes where displacements are not calculated. Because we're working with rectangular prisms, we can easily pick out the surface nodes using a search algorithm with a logical mask. MATLAB and Fortran provide vector intrinsics that allow one to do so. Figure 2.2a illustrates only the surface nodes according to our implementation's node arrangement—which is the xz-plane whose domain is $[y_{\min}, y_{\max}]$. However, due to the nature of the analytical solutions in chapter 3, we need all the surface nodes of the rectangular faces for which there are no displacements. This means that edge nodes are shared between 2 adjacent faces and corner nodes between 3 adjacent faces. In order to properly apply the logical mask to find only the surface nodes need to know that each FEs' nodes are numbered

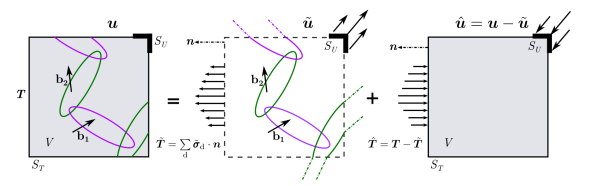
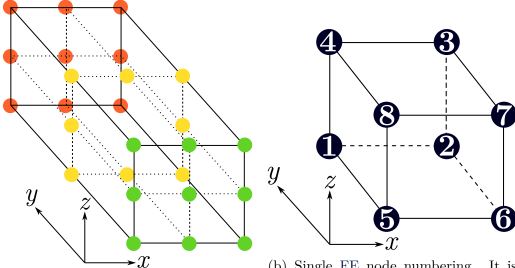


Figure 2.1: The dislocation ensemble in a volume V is bounded by surface S. First, the traction field $\sum_{\mathbf{d}} \tilde{\boldsymbol{\sigma}}_{\mathbf{d}}$ due to the dislocation ensemble is evaluated at the surface. Then, a traditional FEM or Boundary Element Method (BEM) calculates the image traction field $\hat{\boldsymbol{\sigma}} \times \boldsymbol{n}$. Which is then fed back to the DDD problem to evolve the dislocation positions and repeat the cycle. Image edited from [86].



(a) Arrangement of the surface nodes of our FE model. FE nodes are arranged in chunks of $\Delta x \Delta z$ nodes in our implementation, but we only want the surface nodes.

(b) Single FE node numbering. It is necessary to know which FE plane corresponds to which node labels. This lets us design the auxiliary matrix that selects nodes according to the planes we want to extract.

Figure 2.2: FE node arrangement for coupling to DDD.

according to fig. 2.2b.

Using fig. 2.2 one can work out which nodes are of interest to whichever surface is being extracted. The ordering of the nodes in the final array will depend on the definition of the problems in chapter 3.

Node selection remains an expensive operation and minimising array indexing is of the utmost importance for the best perfomance. Selecting nodes in the traditional sense, i.e. with code branching such as if statements or case selection is unmantainable, verbose and very prone to mistakes. The issue was solved by introducing an auxiliary matrix which defines various parameters that aid node selection and greatly reduces code size, improves readability, and eliminates the need for code branching. The matrix can be constructed utilising fig. 2.2 in order to know which nodes correspond to which FE planes. The p^{th} column of the matrix¹ corresponds to the p^{th} plane (according to an arbitrary plane numbering) and is defined as,

$$\boldsymbol{V_p}^{\mathsf{T}} = \begin{bmatrix} L_{1p} & L_{2p} & \cdots & L_{Np} & A_p & C_p \end{bmatrix}, \tag{2.1}$$

where L_{np} is the numeric label for node n as given by fig. 2.2, A_p is the area of the plane, and C_p is the numeric label of the orthogonal coordinate to the plane $C_p = 1, 2, 3$ for the x, y, z coordinates respectively. A_p lets us segment our output and transitional arrays so that the only data being modified is that which corresponds to the correct plane and C_p lets us know which coordinate we must use in our selection criteria. Using our particular node labelling scheme (with dimensions Δx , Δy , Δz respectively in the x, y, z directions), the matrix is defined as,

$$\mathbf{V} = \begin{bmatrix} 5 & 2 & 6 & 1 & 5 & 4 \\ 1 & 6 & 5 & 2 & 6 & 3 \\ 8 & 3 & 7 & 4 & 1 & 8 \\ 4 & 7 & 8 & 3 & 2 & 7 \\ \Delta y \Delta z & \Delta y \Delta z & \Delta x \Delta z & \Delta x \Delta z & \Delta x \Delta y & \Delta x \Delta y \\ 1 & 1 & 2 & 2 & 3 & 3 \end{bmatrix}.$$
 (2.2)

The information codified in eq. (2.2) lets us index and process only the necessary columns to extract the surface nodes we're interested in. The advantage of this setup over a naïve implementation is that it can be relatively easily expanded, maintained, and is general enough that it lends itself to a variety of selection criteria. The columns from left to right (1 to 6) represent: face $1 \equiv \min(x)$, yz-

 $^{^1\}mathrm{MATLAB}$ uses column-major ordering, so this gives us the best performance for vectorised code.

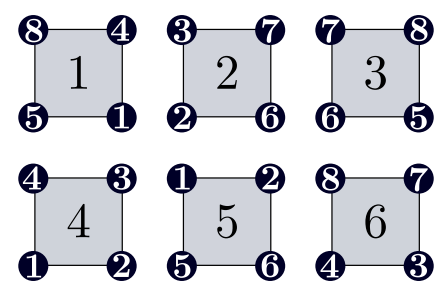


Figure 2.3: Self-consistent, chirality preserving surface planes. Plane and node numbering according to the columns of eq. (2.2) and fig. 2.2.

plane; face $2 \equiv \max(x)$, yz-plane; face $3 \equiv \min(y)$, xz-plane; face $4 \equiv \max(y)$, xz-plane; face $5 \equiv \min(z)$, xy-plane; face $6 \equiv \max(z)$, xy-plane.

It is important to make sure the nodes are extracted in a self consistent manner because the problem definitions in chapter 3 assume a specific node ordering. In particular, the labelling chirality must remain the same. This is due to the fact that the problems in chapter 3 utilise internal coordinates derived from a set of basis vectors, and scalar projections onto them. If the relationship between the vectors is kept, but the chirality is different (opposite) the calculated quantities will have the wrong sign. If however the relationship between the vectors is different to the problem formulation, the program will crash due to division by 0. Self-consistency was achieved by placing an imaginary observer inside the FE model facing the $\min(x)$, yz-plane (face 1) and rotating it to view all 6 planes according to fig. 2.3.

2.3 Mapping Forces

 \hat{F} only cares about the global node numbers, of which there are M. The columns of N_L , F_N represent the 4 nodes of a SE, the rows represent the SE they are part of. The column vector \boldsymbol{x} represents the x, y, z-coordinates of the node-element or global node corresponding to their subscript. The column vector γ_t is the list

Algorithm 2.1 If \hat{F} 's columns are arranged the same way as γ_t .

- 1: \triangleright Loop through the array containing the node labels of the relevant surface nodes.
- 2: for i = 0; $i < \text{length}(\gamma_t)$; i + + do
- 3: ▷ Save the global node label for the current iteration.
- 4: $n \leftarrow \gamma_t[i]$
- 5: Use the node label to find a vector with the linearised index of all surface nodes whose labels correspond to the node whose forces we want to pass on to the FEM coupler.
- 6: $L \leftarrow \text{find}(\mathbf{N_L} == n)$
- 7: ▷ Loop over coordinates.
- 8: **for** k = 0; k < 3; k + + do
- 9: Arrows Use global node label vector to index the force array from the analytical force calculation due to dislocations on surface nodes. Multiplied by 3 because there are three coordinates per node. We sum the forces from the analytical calculation because the same global node can be part of multiple surface elements. We add k because the x, y, z coordinates are consecutively stored in $\mathbf{F_n}$.

10:
$$\mathbf{\hat{F}}[3 \times \mathbf{L} + k] \leftarrow \mathbf{\hat{F}}[3 \times \mathbf{L} + k] + \sum \mathbf{F_n}[3 \times \mathbf{L} + k]$$
11: end for

12: end for

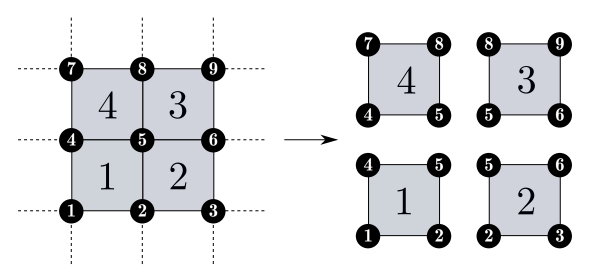


Figure 2.4: FE nodes shared by multiple SEs.

of global nodes for which the forces induced by dislocations must be calculated.

$$\mathbf{x}_{e,n}^{\mathsf{T}} \equiv \begin{bmatrix} x_{en} & y_{en} & z_{en} \end{bmatrix}, \quad \hat{\mathbf{x}}_{n}^{\mathsf{T}} \equiv \begin{bmatrix} x_{n} & y_{n} & z_{n} \end{bmatrix}$$
 (2.3)

$$\mathbf{N_{L}} = \begin{bmatrix} l_{1,1} & l_{1,2} & l_{1,4} & l_{1,5} \\ l_{2,2} & l_{2,3} & l_{2,5} & l_{2,6} \\ l_{3,5} & l_{3,6} & l_{3,8} & l_{3,9} \\ l_{4,4} & l_{4,5} & l_{4,7} & l_{4,8} \end{bmatrix}, \quad \boldsymbol{\gamma} = \begin{bmatrix} l_{1} \\ l_{2} \\ \vdots \\ l_{9} \end{bmatrix}$$

$$\mathbf{F_{e}} = \begin{bmatrix} \boldsymbol{x_{1,1}} & \boldsymbol{x_{1,2}} & \boldsymbol{x_{1,3}} & \boldsymbol{x_{1,4}} \\ \boldsymbol{x_{2,1}} & \boldsymbol{x_{2,2}} & \boldsymbol{x_{2,3}} & \boldsymbol{x_{2,4}} \\ \boldsymbol{x_{3,1}} & \boldsymbol{x_{3,2}} & \boldsymbol{x_{3,3}} & \boldsymbol{x_{3,4}} \\ \boldsymbol{x_{4,1}} & \boldsymbol{x_{4,2}} & \boldsymbol{x_{4,3}} & \boldsymbol{x_{4,4}} \end{bmatrix}, \quad \hat{\boldsymbol{F}} = \begin{bmatrix} \hat{\boldsymbol{x}_{1}} \\ \hat{\boldsymbol{x}_{2}} \\ \hat{\boldsymbol{x}_{3}} \\ \hat{\boldsymbol{x}_{4}} \end{bmatrix}. \quad (2.5)$$

$$\mathbf{F}_{\mathbf{e}} = \begin{bmatrix} x_{1,1} & x_{1,2} & x_{1,3} & x_{1,4} \\ x_{2,1} & x_{2,2} & x_{2,3} & x_{2,4} \\ x_{3,1} & x_{3,2} & x_{3,3} & x_{3,4} \\ x_{4,1} & x_{4,2} & x_{4,3} & x_{4,4} \end{bmatrix}, \quad \hat{\mathbf{F}} = \begin{bmatrix} \hat{x}_1 \\ \hat{x}_2 \\ \hat{x}_3 \\ \hat{x}_4 \end{bmatrix}. \tag{2.5}$$

Chapter 3

Analytical Forces Induced by Dislocations on Linear Rectangular Surface Elements

3.1 Forces Exerted by a Dislocation Line Segment on Linear Rectangular Surface Elements

Coupling DDD to FEM requires the traction field $\sigma^{\infty}(x) \cdot n(x)$ to be distributed among the set of relevant discrete nodes of a FE or Boundary Element (BE) model. This is usually achieved as follows,

$$\mathbf{F}^{(m)} = \int_{S^e} \left[\boldsymbol{\sigma}^{\infty}(\mathbf{x}) \cdot \mathbf{n}(\mathbf{x}) \right] N_m(\mathbf{x}) \, dS_e \,, \tag{3.1}$$

where dS_e is the infinitesimal surface element with surface area S_e . $N_m(\boldsymbol{x})$ are so-called shape functions (interpolation functions) that distribute the traction field among the surface element's nodes.

The problematic singularity associated with the classical Volterra dislocation is avoided by using the non-singular formulation of Cai et al. [101]. The stress field of a dislocation in a homogenous infinite linear elastic domain can be calculated as a contour integral along the loop [100],

$$\sigma_{ij}^{\infty}(\boldsymbol{x}) = C_{ijkl} \oint \epsilon_{lnh} C_{pqmn} \frac{\partial G_{kp}(\boldsymbol{x} - \boldsymbol{x}')}{\partial x_q} b_m dx'_h, \qquad (3.2)$$

where C_{ijkl} is the elastic stiffness matrix, ϵ_{lnh} the permutation operator, \boldsymbol{b} the Burgers vector, \boldsymbol{x}' the coordinate that spans the dislocation, and $G_{kp}(\boldsymbol{x}-\boldsymbol{x}')$ is Green's function of elasticity [100]. $G_{kp}(\boldsymbol{x}-\boldsymbol{x}')$ is defined as the displacement component in the x_k direction at point \boldsymbol{x} due to a force applied in the x_p direction

at point \mathbf{x}' . The traditional singularity comes from taking the Burgers vector distribution as a delta function. Cai et al. [101] proposed an alternative definition of $G_{kp}(\mathbf{x} - \mathbf{x}')$ which has a wider isotropic spread mainly localised in a radius a around the dislocation core,

$$G_{ij}(\boldsymbol{x} - \boldsymbol{x}') = \frac{1}{8\pi\mu} \left[\delta_{ij} \partial_{pp} - \frac{1}{2(1-\nu)} \partial_{ij} \right] R_a, \qquad (3.3)$$

where μ , ν are the isotropic shear modulus and Poisson's ratio respectively, δ_{ij} is the Kronecker Delta, $\partial_{x_1...x_n} \equiv \frac{\partial^n}{\partial x_1...\partial x_n}$. R_a is the defined as,

$$R_a(\mathbf{x}) = R(\mathbf{x}) * w(\mathbf{x}) = \int R(\mathbf{x} - \mathbf{x}') w(\mathbf{x}') d^3 \mathbf{x}'$$
$$= \sqrt{R^2 + a^2}$$
(3.4a)

$$w(\mathbf{x}) = \frac{15a^4}{8\pi (R^2 + a^2)^{7/2}},$$
(3.4b)

where $w(\mathbf{x})$ is the isotropic Burgers vector distribution derived in the appendix of [101], $\mathbf{x} = (x, y, z)$ and $R(\mathbf{x}) = \sqrt{x^2 + y^2 + z^2}$.

For two dislocation nodes (1, 2) connected by straight line segments eq. (3.2) becomes,

$$-\frac{\mu}{8\pi} \int_{x_1}^{x_2} \left(\frac{2}{R_a^3} + \frac{3a^2}{R_a^5} \right) [(\mathbf{R} \times \mathbf{b}) \otimes d\mathbf{x}' + d\mathbf{x}' \otimes (\mathbf{R} \times \mathbf{b})]$$

$$+ \frac{\mu}{4\pi (1 - \nu)} \int_{x_1}^{x_2} \left(\frac{1}{R_a^3} + \frac{3a^2}{R_a^5} \right) [(\mathbf{R} \times \mathbf{b}) \cdot d\mathbf{x}'] \mathbf{I}_2$$

$$- \frac{\mu}{4\pi (1 - \nu)} \int_{x_1}^{x_2} \frac{1}{R_a^3} [(\mathbf{b} \times d\mathbf{x}') \otimes \mathbf{R} + \mathbf{R} \otimes (\mathbf{b} \times d\mathbf{x}')]$$

$$+ \frac{\mu}{4\pi (1 - \nu)} \int_{x_1}^{x_2} \frac{3}{R_a^5} [(\mathbf{R} \times \mathbf{b}) \cdot d\mathbf{x}'] \mathbf{R} \otimes \mathbf{R}$$

$$(3.5)$$

3.1.1 Resolving Singularities when Dislocation Line Segments are Parallel to Surface Elements

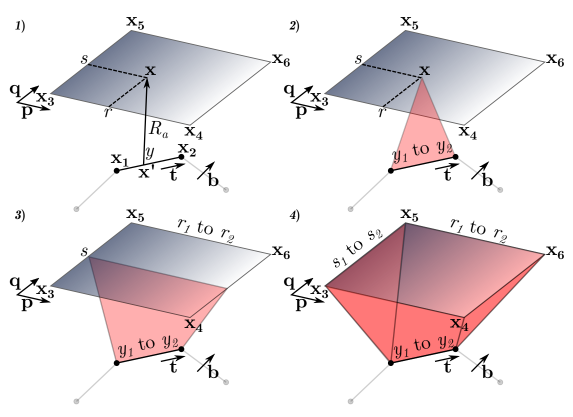


Figure 3.1: Diagram of the line integral method used to find analytical expressions for the forces exerted by dislocations on linear rectangular SEs [86]. 1) For any given point \boldsymbol{x} on the SE and any given point \boldsymbol{x}' on the dislocation line segment, define distance R_a . 2) Integrate from $x_1 \to x_2$ along line direction \boldsymbol{t} . 3) Integrate from $r_1 \to r_2$ along vector \boldsymbol{p} . 4) Integrate from $s_1 \to s_2$ along vector \boldsymbol{q} .

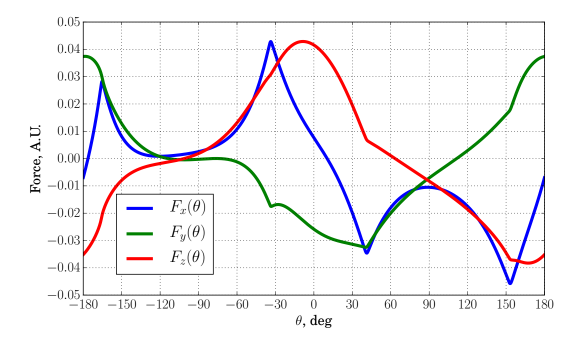


Figure 3.2: Effects of rotating a single dislocation line segment on the forces exerted by it on a linear rectangular SE. The specific values of this function are not known a priori, all that is known is that it must be periodic $(T=2\pi)$ and have finite maximum and minimum values. The singularity is avoided by perturbing the angle $\theta=0 \to \theta=\pm\epsilon$, $\epsilon\gtrsim 0$.

Chapter 4

Parallelisation of the Analytical Forces Induced by Dislocations on Surface Elements

4.1 Parallelisation on Graphics Processing Units

Efficient parallelisation requires coalesced memory access, which means we have to be extremely careful when mapping CPU memory to device memory. The fact that threads work "simultaneously" means that in order to obtain good performance, data which is to be "simultaneously" loaded into each thread must be contiguous. This maximises cache memory use and therefore reduces slow memory fetch operations to global or shared memory.

The parallelisation was done only over the SEs in order to avoid the undesirable and inefficient GPU branching that would occur under other schemes.

The most natural form of parallelisation is to have blocks of 4n threads where $n \leq 8 \in \mathbb{N}$. This also fits nicely into the 32 thread per warp paradigm.

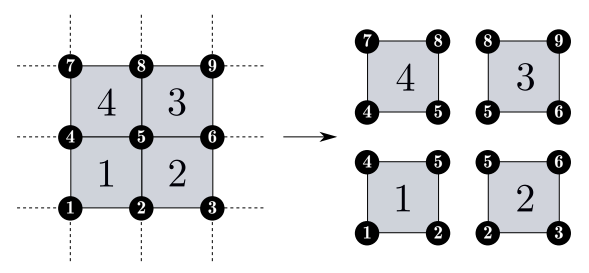


Figure 4.1: Each linear rectangular SE is mapped to one thread.

Algorithm 4.1 Elements in host \mapsto device.

```
1: function ELEMENT_HOST_DEVICE_MAP({}^{\text{h}}\boldsymbol{X}[N][3 \times E])
                                         ▶ Input, temporary, and output indices set to zero.
 2:
         i, j, k \leftarrow 0
 3:
               \triangleright <sup>d</sup>X accommodates all 3 coordinates of all N nodes in all E elements.
 4:
         {}^{\mathrm{d}}\boldsymbol{X} \leftarrow \mathrm{malloc}(3 \times E \times N)
 5:
                                                                     ▷ Loop over nodes in element.
         for (n = 0; n < N; n + +) do
 6:
                \triangleright Set output index to point at the n^{\text{th}} node of the first coordinate of
 7:
     the first element.
 8:
              k \leftarrow j
              for (c = 0; c < 3; c + +) do
                                                                             ▷ Loop over coordinates.
 9:
                   \triangleright Set input index to point at the c^{\text{th}} coordinate of the first element
10:
    of the n^{\text{th}} node.
                   i \leftarrow c
11:
                   for (e = 0; e < E; e + +) do
                                                                                 ▶ Loop over elements.
12:
                        {}^{\mathrm{d}}\boldsymbol{X}[k+e] \leftarrow^{\mathrm{h}} \boldsymbol{X}[n][i]
13:
                              \triangleright Advance output index to point at the n^{\rm th} node of the c^{\rm th}
14:
     coordinate of the first element.
                        i \leftarrow i + 3
15:
16:
                   end for
                              \triangleright Advance output index to point at the n^{\mathrm{th}} node of the c^{\mathrm{th}}
17:
     coordinate of the first element.
                   k \leftarrow k + E \times N
18:
19:
               \triangleright Advance temporary index to point at the (n+1)^{\text{th}} node of the first
20:
     coordinate of the first element.
21:
              j \leftarrow j + E
         end for
22:
         return {}^{\mathrm{d}}X
23:
24: end function
```

Algorithm 4.2 Elements in device \mapsto thread.

```
function ELEMENT_DEVICE_THREAD_MAP({}^{d}X[3 \times E \times N],
 1: GPU
    {}^{\rm t}X[N][3]
                                                                             ▶ Thread, input indices.
 2:
         idx, i \leftarrow \text{threadIdx.} x + \text{blockIdx.} x \times \text{blockDim.} x
 3:
 4:
         for (c = 0; c < 3; c + +) do
                                                                                ▶ Loop over elements.
              for (n = 0; n < N; n + +) do
                                                                     ▶ Loop over nodes in element.
 5:
                   {}^{\mathrm{t}}\boldsymbol{X}[n][c] \leftarrow^{\mathrm{d}} \boldsymbol{X}[i+n \times E]
 6:
 7:
              i \leftarrow i + E \times N \triangleright \text{Advance input index to the } (n+1)^{\text{th}} \text{ node of element}
 8:
    i.
 9:
         end for
         return {}^{\mathrm{t}}X[n][c]
10:
11: end GPU function
```

Algorithm 4.3 Force in thread $\stackrel{+}{\mapsto}$ device.

```
1: GPU function ADD_FORCE_THREAD_DEVICE({}^{\mathrm{t}}F_n[N][3], {}^{\mathrm{t}}F_e[3], {}^{\mathrm{d}}F_n[3 \times
    E \times N,
    {}^{\mathrm{d}}\mathbf{F}_{e}[3\times E], \mathrm{idx})
 2:
                                                                                         ▶ Nodal force.
                 ▷ Set output index to whatever parallelisation index is given to the
3:
    function. In this case the same index we use for the main parallelisation.
         i \leftarrow idx
 4:
         for (n = 0; n < N; n + +) do
                                                                                  ▶ Loop over nodes.
5:
             for (c = 0; c < 3; c + +) do
6:
                                                                           ▶ Loop over coordinates.
                    ▶ Ensure nodal forces are correctly added and mapped from local
 7:
    thread memory to global device memory.
                  atomicAdd({}^{\mathrm{d}}\boldsymbol{F_n}[i+c\times E\times N], {}^{\mathrm{t}}\boldsymbol{F_n}[n][c])
8:
9:
             end for
                         ▷ Displace output index to point at the first coordinate of the
10:
     (n+1)^{\text{th}} node of the idx<sup>th</sup> SE.
             i \leftarrow i + E
11:
         end for
12:
                                                                                          ▶ Total force.
13:
         i \leftarrow idx
                                                                               ▷ Reset output index.
14:
         for (c = 0; c < 3; c + +) do
                                                                           ▶ Loop over coordinates.
15:
             atomicAdd({}^{\mathrm{d}}\boldsymbol{F_e}[i], {}^{\mathrm{t}}\boldsymbol{F_e}[c])
16:
             \triangleright Advance the output index to point at the (i+1)^{\text{th}} coordinate of the
    idx<sup>th</sup> SE.
             i \leftarrow i + E
18:
19:
         end for
         return {}^{\mathrm{d}}F_n, {}^{\mathrm{d}}F_e
20:
21: end GPU function
```

```
Algorithm 4.4 Force (parallelise over dislocations) in thread \stackrel{+}{\mapsto} device.
```

```
1: GPU function DLN_ADD_FORCE_THREAD_DEVICE({}^{\mathrm{t}}F_{n}[N][3], {}^{\mathrm{t}}F_{e}[3],
    {}^{\mathrm{d}}\boldsymbol{F_n}[3 \times E \times N], {}^{\mathrm{d}}\boldsymbol{F_e}[3 \times E], k)
                                                                                     ▷ Nodal force.
 2:
                           ⊳ Set output index to correspond to whatever SE we're on.
 3:
    By parallelising over dislocation lines, we must loop through SEs in the main
    code, this is the value of k we provide. Multiply by 3 and N because each SE
    has 3 coordinates and N nodes.
 4:
         i \leftarrow 3 \times N \times k
         j \leftarrow 0
 5:
                                                                                for (n = 0; n < N; n + +) do
                                                                               ▶ Loop over nodes.
 6:
             for (c = 0; c < 3; c + +) do
                                                                       ▶ Loop over coordinates.
 7:
                   ▶ Ensure nodal forces are correctly added and mapped from local
    thread memory to global device memory.
                  atomicAdd({}^{\mathrm{d}}\boldsymbol{F_n}[i+j+c], {}^{\mathrm{t}}\boldsymbol{F_n}[n][c])
 9:
10:
             end for
                     ▶ Displace auxiliary index to point at the first coordinate of the
11:
    (n+1)^{\text{th}} node of the k^{\text{th}} SE.
             j \leftarrow j + 3
12:
         end for
13:
                                                                                      ▶ Total force.
14:
                 \triangleright Set auxiliary index to point at the first coordinate of the k^{\text{th}} SE.
15:
         j \leftarrow 3 \times k
16:
         for (c = 0; c < 3; c + +) do
17:
                                                                       ▶ Loop over coordinates.
             atomicAdd({}^{d}\boldsymbol{F_e}[j+c], {}^{t}\boldsymbol{F_e}[c])
18:
         end for
19:
         return {}^{\mathrm{d}}F_n, {}^{\mathrm{d}}F_e
20:
21: end GPU function
```

Algorithm 4.5 Nodal force in device \mapsto host.

```
1: function fx_device_host_map({}^{d}\boldsymbol{F_n}[3 \times E \times N], {}^{h}\boldsymbol{F_n}[N][3 \times E])
                                                     ▶ Set input and output indices to zero.
2:
        i, j \leftarrow 0
        for n = 0; n < N; n + + do
3:
                                                                              ▶ Loop over nodes.
                                      ▶ Reset output index to point at the first element.
 4:
             for e = 0; e < E; e + + do
                                                                          ▶ Loop over elements.
 5:
                 for c = 0; c < 3; c + + do
                                                                       ▶ Loop over coordinates.
6:
                      {}^{\mathrm{h}}\mathbf{F}_{n}[n][j+c] = {}^{\mathrm{d}}\mathbf{F}_{n}[i+e+c\times E\times N]
 7:
8:
                       ▶ Advance output index to point at the first coordinate of the
9:
    (e+1)^{\text{th}} element.
                 j \leftarrow j + 3
10:
             end for
11:
             \triangleright Advance input index to point at the first coordinate of the (n+1)^{\text{th}}
12:
    node of the first element.
             i \leftarrow i + E
13:
        end for
14:
        return {}^{\rm h}F_n
15:
16: end function
```

4.1.1 Data Mapping

- 4.1.1.1 Elements: Host \mapsto Device
- 4.1.1.2 Elements: Device \mapsto Thread
- 4.1.1.3 Force: Thread $\stackrel{+}{\mapsto}$ Device
- 4.1.1.4 Force (Parallelise over Dislocations): Thread $\stackrel{+}{\mapsto}$ Device
- 4.1.1.5 Nodal Force: Device \mapsto Host
- 4.1.1.6 Total Force: Device \mapsto Host

$$\boldsymbol{X}_{en} \coloneqq [x_{en}, y_{en}, z_{en}] \tag{4.1a}$$

 $\boldsymbol{X}_{(1 \to E)n} \mapsto \boldsymbol{X}_n$

$$X_n := [x_{1n}, y_{1n}, z_{1n}, \dots, x_{En}, y_{En}, z_{En}]$$
 (4.1b)

 $oldsymbol{X}_{1 o N}\mapsto oldsymbol{X}^{ ext{SE}}$

$$[x_{11}, \dots, x_{E1}, x_{12}, \dots, x_{E2}, \dots, x_{1N}, \dots, x_{EN},$$

$$\boldsymbol{X}^{\text{SE}} := y_{11}, \dots, y_{E1}, y_{12}, \dots, y_{E2}, \dots, y_{1N}, \dots, y_{EN},$$

$$z_{11}, \dots, z_{E1}, z_{12}, \dots, z_{E2}, \dots, z_{1N}, \dots, z_{EN}]$$

$$(4.1c)$$

where e is the SE label, n the node label, E the number of SEs in the scope. N

¹Not quite but essentially simultaneously.

Algorithm 4.6 Total force in device \mapsto host.

```
1: function ftot_device_host_map({}^{d}F_{e}[3 \times E], {}^{h}F_{e}[3 \times E])
                                                                    ▶ Set output index to zero.
 2:
 3:
         for e = 0; e < E; e + + do
                                                                           ▶ Loop over elements.
             for c = 0; c < 3; c + + do
 4:
                                                                       ▶ Loop over coordinates.
                  {}^{\mathrm{h}}\boldsymbol{F_e}[i+c] = {}^{\mathrm{d}}\boldsymbol{F_e}[e+c\times E]
 6:
                   > Advance the output index to point at the first coordinate of the
     (e+1)^{\text{th}} surface element.
             i \leftarrow i + 3
 8:
 9:
         end for
         return {}^{\rm h}F_e
10:
11: end function
```

the number of nodes in a SE.

Data-mapping according to ?? and relabelling the nodes so they go from $0 \rightarrow N-1$, the data from fig. 4.1 would be arranged in device memory like eq. (4.2),

$$\boldsymbol{X}^{\text{SE}} = \begin{bmatrix} \underbrace{b_{x}, i_{x}, f_{x}, g_{x}}_{\mathbf{x_{0}}}, \underbrace{a_{x}, b_{x}, i_{x}, h_{x}}_{\mathbf{x_{1}}}, \underbrace{i_{x}, d_{x}, e_{x}, f_{x}}_{\mathbf{x_{2}}}, \underbrace{b_{x}, c_{x}, d_{x}, i_{x}}_{\mathbf{x_{3}}}, \\ \dots y\text{-coord} \dots, \\ \dots z\text{-coord} \dots \end{bmatrix}$$

$$(4.2)$$

In the GPU, each thread will cater to one SE at a time. This means that each thread will have to extract the relevant data from the 1D array with length $3 \times E \times N$ into four 1D arrays of length 3. The purpose of CNE mapping is to provide the warp with coalesced memory access. This is achieved via ??.

Since ?? is performed in a CUDA GPU, threads in a single block execute sequentially from,

threadIdx.
$$x = 0 \rightarrow \text{threadIdx.} x = \text{blockDim.} x - 1$$
, (4.3)

while threads in different blocks execute in parallel. Coalesced memory access is ensured by having each block load a cache line whose entries are contiguously accessed by the threads in the block. Using the same notation as $\ref{eq:local_second}$, full cache line utilisation (optimal cache use) is achieved if cache lines can accommodate l entries

Figure 4.2: Minimum working complete example of the backward coodinate-node-element-map. The explicit calculations of global indices correspond to substituting the values for "idx" and "idxi" as in ??. X_a^b denotes a 1D array of length three containing the xyz-coordinates of node a of element b. Each thread concerns itself with only one element at a time. The dash-dot and dashed boxes represent cache lines for a thread block, the dotted line represents a memory fetch, and the dashed lines in X^E represent steps of E entries (the start of the data for the next node of the element type we're dealing with). The memory operations are not shown twice to minimuse redundancy.

given by eq. (4.4),

$$l = \begin{cases} a \times N \times E &, a > 0 \in \mathbb{N} \\ & \text{or} \\ \frac{1}{2^a} \times N \times E &, a \ge 0 \in \mathbb{N}, N \times E \equiv 0 \pmod{2^a} . \end{cases}$$
 (4.4)

Figure 4.2 shows an example of ?? up to the y-coordinate.

4.1.1.7 Node Coordinate Element Map

The node-coordinate-element (NCE) data mapping in eq. (4.5) is carried out by algorithm 4.7, each thread looks after a given SE.

Algorithm 4.7 NCE data mapping.

```
for all n nodes \in surface element do

for all c coordinates \in [x, y, z] do

for all e surface elements \in surface mesh section do

list.append(data of the n^{\text{th}} node with c^{\text{th}} coordinate of the e^{\text{th}} SE)

end for

end for
```

$$\boldsymbol{X}_{en} \coloneqq [x_{en}, y_{en}, z_{en}] \tag{4.5a}$$

$$X_{(1 \to E)n} \mapsto X_n$$

$$\mathbf{X}_n := [x_{1n}, \dots, x_{En}, y_{1n}, \dots, y_{En}, z_{1n}, \dots, z_{En}]$$
 (4.5b)

$$X_{1 \rightarrow N} \mapsto X^{\operatorname{SE}}$$

$$\mathbf{X}^{\text{SE}} := \begin{bmatrix} x_{11}, \dots, x_{E1}, y_{11}, \dots, y_{E1}, z_{11}, \dots, z_{E1} \\ \dots, \dots, \\ x_{1M}, \dots, x_{EN}, y_{1N}, \dots, y_{EN}, z_{1N}, \dots, z_{EN} \end{bmatrix}$$
(4.5c)

where e is the surface element, n the node, E the total number of SEs in scope, N the total number of nodes in each SE.

Data-mapping according to algorithm 4.7 and relabelling the nodes so they go from $0 \to N-1$, the data from fig. 4.1 would be arranged in device memory like eq. (4.6),

$$\underbrace{\begin{bmatrix} h_x, i_x, f_x, g_x, h_y, i_y, f_y, g_y, h_z, i_z, f_z, g_z \\ \dots & \mathbf{x_1}, xyz\text{-coords} \dots, \\ \dots & \mathbf{x_2}, xyz\text{-coords} \dots, \\ \dots & \mathbf{x_3}, xyz\text{-coords} \dots \end{bmatrix}}$$
(4.6)

4.1.1.8 Resolving Data Write Conflicts

Data write conflicts can be a problem in parallel applications where global data is changed by multiple threads within the same clock tick. This can be avoided with atomic operations.

4.1.2 Resolving Parallel Dislocation Line Segments to Surface Elements

Dealing with the special case when the dislocation line segment t is parallel to the SE is relatively trivial when in a serial program. By the mean value theorem, we

can slightly perturb t by rotating it by a small angle around the midpoint of t with respect to the axis of rotation defined by $t \times n$. In contrast to serial code, program branches can have a serious effect on parallel performance due to warp divergence. Due to the complexity of the force calculation and relative rarity of the edge case, there is no branching behaviour in the parallel code until *after* the calculation is performed. Where algorithm 4.8 is performed.

Algorithm 4.8 Resolving cases when $t \parallel n$ on GPUs.

```
for all surface elements and line segments do \dots

if t_{\mathrm{thread}} \parallel n_{\mathrm{thread}} then x_{\mathrm{buffer}} \leftarrow x_{\mathrm{thread}}
t_{\mathrm{buffer}} \leftarrow t_{\mathrm{thread}}
else F_{\mathrm{total}} + = F_{\mathrm{thread}}
end if end for return to serial code if x_{\mathrm{buffer}} \neq \mathrm{empty} then for all surface elements and line segments do perform calculation for t \parallel n end for end if
```

4.1.2.1 Implementation

The node mapping would be the same as ??. The parallelisation would be on individual packets of a SE with a slightly rotated dislocation line segment, as in a single iteration of the serial code where the dislocation line segment is rotated. The forces would be averaged with atomic operations and <code>__syncthreads()</code> before they are added to the total nodal forces.

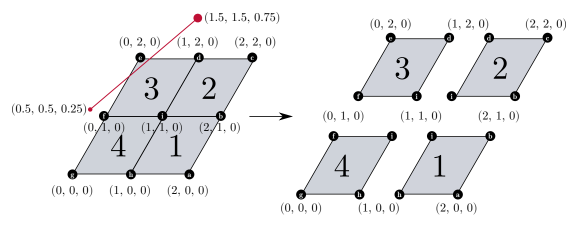


Figure 4.3: Test case for CUDA development. Dislocation line not shown in node separation as the parallelisation happens across surface elements. Not to perspective or scale.

4.1.3 Test Case

According to ?? and the node labelling scheme of fig. 3.1 the data mapping would look like:

$$\mathbf{X}^{\text{SE}} = \underbrace{\begin{bmatrix} \underbrace{x \text{-coord}} \\ 1, 1, 0, 0, \underbrace{2, 2, 1, 1}, \underbrace{1, 1, 0, 0}, \underbrace{2, 2, 1, 1}, \\ \underbrace{y \text{-coord}} \\ \underbrace{y \text{-coord}} \\ \underbrace{x_{3}} \underbrace{x_{4}} \underbrace{x_{5}} \underbrace{x_{5}} \underbrace{x_{6}}$$

$$\underbrace{x_{5}} \underbrace{x_{6}} \underbrace{x_{6}}$$

$$\underbrace{0, 0, 0, 0, \underbrace{0, 0, 0, 0}, \underbrace{0, 0, 0, 0}, \underbrace{0, 0, 0, 0}, \underbrace{0, 0, 0, 0}, \underbrace{0, 0, 0}, \underbrace{0, 0, 0}, \underbrace{0, 0, 0, 0}, \underbrace{$$

In general the number of indices in the array would be $3 \times e \times n$ where e is the number of surface elements, n the number of nodes per element and 3 the number of spatial dimensions.

4.2 Thread Block Size Optimisation

 $http://docs.nvidia.com/cuda/cuda-c-best-practices-guide/index.html\#memory-optimizations \\ http://docs.nvidia.com/cuda/cuda-c-best-practices-guide/index.html\#execution-configuration-optimizations$

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Appendix A

Best Practices

This chapter defines the best practices that will make improve the development and testing pipeline by defining rules and standards that facilitate collaboration.

A.1 Filesystem

All files must be within /project_name.

```
1. All files must be within /project name,
```

```
~/ = project_name/,
~/~/ = project_name/src/ or project_name/dev/.
```

- 2. Release code must be within ~/src/.
- 3. Development code must be within ~/dev/.
- 4. Test data must be within ~/tests/.
- 5. Documentation must be within ~/~/doc/.
- 6. Examples must be within $\sim/\sim/\exp/$.
- 7. External libraries must be within ~/lib/.
- 8. Generated images must be within ~/~/images/.
- 9. Old versions recordkeeping must be within ~/prv/va.b.c/.

A.2 Versioning

1. Use a version control system like GitHub or PasteBin.

- 2. There must be a master branch that is only changed when the code is stable and bug free.
- 3. Development branches should be exploited as seen fit without making things overly convoluted.
- 4. Commits must be as bug free and regular as possible. When to commit is left to the developer's discresion.
- 5. Commit messages should be as descriptive as possible.
- 6. Versions should be specified as va.b.c where a, b, c = integers. The three levels are a = release version (usable, bug free code), b = beta version (code that is undergoing testing), c = alpha version (code that is under active development).

A.3 Documentation

A.3.1 Commenting

Every codefile must be appropriately commented by meeting the following guidelines.

- 1. The start of each codefile must have a heading detailing the creator, date of creation and edit history (date and name of editor).
- 2. Below the heading there must be a general explanation of the code. It must state any procedures, structures, objects and how they are to be utilised. Any backward or forward dependencies must be stated.
- 3. Below the description and edit history any relevant literature must be mentioned (dois are preferred). Must be as detailed as possible, include equation numbers/ranges if necessary.
- 4. The start of every procedure has an explanation of its purpose, inputs, outputs and inputs-outputs.
- 5. Particularly complicated code blocks must have an in-depth explanation of what it does. Comment each line if necessary.
- 6. Corrections or additions must be explicitly bounded by comments at the start and end of the change. Both bounding comments must have the author's name and the date. Below the starting comment, there should be an explanation of the change. Any punctual comments can be made as normal.

A.3.2 README

Every codefile must have an associated README .tex document that documents the codefile's contents. It must meet the following guidelines as appropriate.

- 1. The name must be that of the file it documents (minus the extension of course).
- 2. Description and overall explanation of the codefile's purpose.
- 3. Overall flow chart or pseudo code describing the file's purpose.
- 4. Document the codefile's procedures. This means describing and explaining their corresponding inputs, outputs, inputs-outputs, forwards and backwards dependencies, and flow charts or pseudo codes.
- 5. Unit test designs and results for each procedure. If appropriate also include those of integral tests.

The codefile may also be appended at the end of documentation if desired (the minted package is highly recommended).

A.4 Modularisation

- 1. Code repetition must be kept to a *strict* minimum. Any piece of code that will be reused must be modularised.
- 2. Procedures must be as self-sufficient as possible *without* repeating code. If repeating code is necessary, replace it with a procedure that is to be repeatedly called instead. Minimising repeated code >>> procedure self-sufficiency.

A.5 Coding Style

All names must meet the following guidelines.

- 1. Indent appropriately. Four space tabs are a good compromise between code necking and readability.
- 2. Minimise the use of nested code blocks, use intrisics, libraries or create procedures instead.
- 3. Break up lines that are uncomfortably long, typically anything over 80–100 characters.

- 4. Names should be appropriately descriptive and human readable.
- 5. All code and names must be systematic and logical.
- 6. the use of upper cases should be reserved for parameters (const variables in C).
- 7. Delimit words with "_" not case changes.
- 8. Long and descriptive \gg short and cryptic.

A.5.1 Filenames

1. If old versions are to be kept, the old stable versions of file must have the date of last modification appended suffixed after the file extension in the _yyyymmdd format. For example, if the stable version of the release code hello_world_parallel.c was last modified on April 25, 2017 it should be archived as

hello_world_parallel.c_20170425.

A.5.2 Variables, Structures and Objects

- 1. Only counters and indices can be single letter variables.
- 2. Structure and object definitions are suffixed with _s and _o respectively.
- Inputs, outputs and input-outputs to procedures must be prefixed with i_,
 o_ and io_ respectively.

A.5.3 Procedures

1. Functions and subroutines must be *prefixed* with f_ and s_ respectively.

Appendix B

Coupling Discrete Dislocation
Dynamics to Finite Element
Methods

Appendix C

Implementation of Analytical Forces Induced by Dislocations on Linear Rectangular Surface Elements

- C.1 Serial C Code MEX File
- C.2 Parallel CUDA C Code MEX File

Appendix D

Talks

D.1 Durham July 12–14 2017

Bridging the gap between the microscopic and macroscopic world is an on-going challenge for science and technology. If we hope to understand complex emergent phenomena we need to study systems that blur the line between micro and macro. In materials science, one such area is the study of extended defects called dislocations; whose nucleation and movement mediate the permanent deformation of materials. These large, high energy defects present very complex and long ranged interactions with each other, crystal boundaries, impurities, free surfaces, and themselves. As such, their dynamics are difficult to study experimentally. So we make justified assumptions and simplifications and create models that let us study them in detail. However if our models are to prove useful in real applications, they must be continually refined and improved by weakening assumptions and removing simplifications. Unfortunately, with increased refinement comes increased computational cost and new challenges. Therefore, finding faster alternatives that do not sacrifice accuracy is of the utmost importance—better yet if the alternatives are more accurate or exact. With the advent of increasingly accessible graphics processor units (GPUs) typically used in video gaming, the power of parallel processing is no longer exclusive to researchers with access to supercomputers. In this talk we will discuss the GPU implementation of exact solutions for the forces dislocations exert on the surfaces of materials.



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