Finite element modelling and PGD based model reduction for piezoelectric and magnetostrictive materials
Zhi Qin

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Finite Element Modeling and PGD Based Model Reduction for Piezoelectric & Magnetostrictive Materials

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Thèse de Doctorat en Electronique

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Abstract

The energy harvesting technology that aims to enable wireless sensor networks (WSN) to be maintenance-free, is recognized as a crucial part for the next generation technology megatrend: the Internet of Things (IoT). Piezoelectric and magnetostrictive materials, which can be used in a wide range of energy harvesting systems, have attracted more and more interests during the past few years. This thesis focuses on multiphysics finite element (FE) modeling of these two materials and performing model reduction for resultant systems, based on the Proper Generalized Decomposition (PGD).

Modeling these materials remains challenging although research in this area has been undergoing over decades. A multitude of difficulties exist, among which the following three issues are largely recognized. First, mathematically describing properties of these materials is complicated, which is particularly true for magnetostrictive materials because their properties depend on factors including temperature, stress and magnetic field. Second, coupling effects between electromagnetic, elastic, and thermal fields need to be considered, which is beyond the capability of most existing simulation tools. Third, as systems becoming highly integrated whole-scale simulations become necessary, which means three dimensional (3D) numerical models should be employed. 3D models, on the other hand, quickly turns intractable if not properly built. The work presented here provides solutions in respond to the above challenges.

A differential forms based multiphysics FE framework is first established. Within this framework quantities are discreted using appropriate Whitney elements. After discretization, the system is solved as a single block, thus avoiding iterations between different physics solutions and leading to rapid convergences. Next, the linear piezoelectric, and a free energy based nonlinear magnetostrictive constitutive model called Discrete Energy Averaged Model (DE-AM) are incorporated into the framework. Our implementation describes underlying material behaviors at reasonable numerical costs. Eventually, two novel PGD based algorithms for model reduction are proposed. With our algorithms, problem size of multiphysics models can be significantly reduced while final results of very good accuracy are obtained. Our algorithms also provide means to handle coupling and nonlinearity conveniently.

All our methodologies are demonstrated and verified via representative examples
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Chapter 1. Introduction

In this chapter, introduction to the research background and motivation are given. Specifically, the concepts of WSN and energy harvesting technologies are introduced in section 1. We briefly review properties of piezoelectric and magnetostrictive materials in section 2. A survey of existing models for these materials, especially those used in energy harvesting, is given in section 3, which also explains our motivation of 3D multiphysics FE modeling and models reduction. The chapter concludes with research objectives and outline of the thesis.

1.1 Energy harvesting in wireless sensor networks

With the ever-increasing computing power over the last decade, WSN, recognized as a key enabling technique for the IoT, has been applied in applications like environmental monitoring, object tracking and body networks. It is predicted that commercial use of WSN will be pervasive in the coming years [1].

A typical WSN scenario is depicted in Figure 1.1 where a large number of sensor nodes are deployed for data acquiring. The collected data are processed on the node before sent to a base station, which are finally accessed by end users through the Internet. Such sensor nodes gather and transfer data, thereby consuming considerable energy. While modern sensors are energeti-
cally low consuming, power supply can still be an issue. This is because some of them are ex-
pected to work for several years, during which period, maintenance of the power units is diffi-
cult, or even impossible in many situations. As a result, batteries are no more adequate in this
regard. On the contrary, energy harvesting technologies are promising as they require little even
zero human intervention.

Figure 1.2. Schematics of (a) piezoelectric, (b) magnetostrictive [2], and (c) magnetoelectric [3] harvester.

Energy harvesting is a mechanism for generating energy from the ambient to provide uninter-
rupted power supply [1]. The harvested energy may come from artificial sources or from the
nature directly. Accordingly, photovoltaic cells, turbines and hydro-generators can be catego-
rized in a group as they harvest energy from the nature. Other energy harvesters collect energy
mostly from mechanical vibrations or electromagnetic sources. Piezoelectric and magnetostric-
tive materials, thanks to their high energy density [4], are widely utilized in such devices. Figure
1.2.a depicts a cantilevered piezoelectric energy harvester that generates voltage on its surfaces
under vibration. It is reported that a device of similar configuration delivers 45 $\mu W$ under a 50
Hz vibration of 0.6 $m \cdot s^{-2}$ [5]. Magnetostrictive energy harvesters usually come with coils and
magnets, as shown in Figure 1.2.b. They can also be used to transfer mechanical vibration into electrical energy, while they are less advantageous than piezoelectric harvesters in terms of miniaturization of volume, due to the existence of coils and magnets. Piezoelectric and magnetostrictive materials can be combined and utilized as magnetoelectric composites (see Figure 1.2.c). In such devices, the magnetostrictive layer generates strain in respond to a varying magnetic field. Due to inter-layer elastic bonding, the strain passes to the piezoelectric layer that eventually generates electrical charges. It can also harvester energy from mechanical vibration.

1.2 Piezoelectric and magnetostrictive materials

1.2.1 Piezoelectric materials and piezoelectric effects

Piezoelectric materials refer to non-conducting ferroelectric materials that produce electric charges on the surface when mechanical stress is applied, and conversely, produce mechanical strain when electric field is applied, as represented in Figure 1.3 where $P$ is the polarization. The stress to charge conversion is called the direct piezoelectric effect, while the other is called the inverse piezoelectric effect.

![Figure 1.3. The (a) direct and (b) inverse piezoelectric effects](image)

Piezoelectric materials can be categorized into inorganic, organic and composite types [6]. Piezoelectric monocristalline materials and piezoelectric ceramics fall into the first category. Piezoelectric single crystals (e.g. quartz) are commonly seen in e.g. high-selectivity filters and high-temperature ultrasonic transducers because of their high mechanical quality factor and excellent stability. Their applicability is limited due to low piezoelectric coefficients and low machining properties. Piezoelectric ceramics (e.g. lead zirconate titanate, or PZT), on the other hand, possess strong piezoelectricity and can be easily formed into various shapes, although
they suffer from low mechanical quality factor and large electric loss. This enables them to be used in high-power transducers and wide-band filters. In the organic piezoelectric group (also referred to as piezoelectric polymers), the polyvinylidene fluoride, or PVDF is probably the most famous. These materials are flexible, low-weight and having small impedance. They are widely used for underwater ultrasonic measuring, pressure sensing, etc. For piezoelectric composites, in which piezoelectric ceramics and polymers are bonded together, properties of individual materials are enhanced. For instance, they can have both strong coupling coefficients and outstanding machining properties, which makes them ideal to be fabricated into large area films or other sophisticated applications [6]. Application of piezoelectric materials in energy harvesting systems can be found in [7].

Figure 1.4. Domain structures of piezoelectric ceramics during poling process [6]: (a) before polarization, domains are randomly oriented, (b) domains rearrange along the electric field direction during polarization, and (c) remnant polarization presents after the poling field is removed.

Piezoelectric effects can be explained from a microscopic point of view. Roughly speaking, they are structurally asymmetric in crystalline, making the electrical domains (see Figure 1.4), which are separated by walls along spontaneous polarization directions, randomly distributed at the original state. Applying increasingly larger electric fields makes these electric domains gradually align with the field, which also results in macroscopic change in shape due to the movements of domains. This procedure is called polarization. Meanwhile, applying mechanical stress causes reallocation of electrical walls and as a result, generating macroscopic polarization, or voltage on the surface. Although piezoelectric coupling is determined by several factors such as temperatures, stresses and fields, the piezoelectric effect can be regarded as linear over a wide range of conditions. We only consider linear piezoelectric behaviors in the thesis.

1.2.2 Magnetostrictive materials and magnetostrictive effects

Magnetostrictive materials refer to those exhibiting magnetostrictive effects that are found in ferromagnetic materials. Magnetostrictive effects consist of two mechanisms: the Joules effect


and the Villari effect [8]. The former implies that rotation of moments to align with an applied field generates strains, while the latter implies that applied stresses cause magnetic moments to rotate thus changing the magnetization. There has been various research on magnetostrictive effects and materials (see e.g. [9]), while we restrict ourselves to giant magnetostrictive materials whose magnetostrictive effects are more significant.

In 1971 the US Naval Ordnance Laboratory invented the first well-known giant magnetostrictive material, called Terfenol-D. Its maximum magnetostriction (i.e. the strain generated during the paramagnetic-ferromagnetic phase transition or in response to an applied field [10]) is >1000 ppm at room temperatures, being two orders of magnitude larger than common magnetostrictive materials. Another famous giant magnetostrictive material is Galfenol. It has better mechanical properties and lower temperature dependencies than Terfenol-D, although its maximum magnetostriction is smaller [10]. Giant magnetostrictive materials are wildly utilized in sonar systems, transducers and energy harvesting devices. Figure 1.5 shows a Terfenol-D based transducer in which mechanical parts are employed to provide pre-stress to the rod. In the following, we review some fundamentals of magnetostrictive materials, based on [10].

Indeed, magnetostrictive materials inherit properties from ferromagnetic materials whose nonlinear relations come from domain wall movements and domain rotations. Inside a ferromagnetic domain, magnetic moments are aligned and exhibit a spontaneous magnetization. When demagnetized, domains are randomly oriented, yielding zero net magnetization (see Figure 1.6.a). When fields are applied, domains have a trend to align themselves with the field, producing a bulk magnetization. Figure 1.6.b to Figure 1.6.d demonstrate the magnetization states in monolithic Terfenol-D under weak, median, and strong fields. When the field is weak, changes in magnetization are mainly due to reversible domain wall movement and moment rotation. As the field increases, irreversible mechanisms arise. Domain wall displacements favor the growth of domains having components in the field direction, while moments rotate to the easy axis closest
to the field direction. Eventually, moments rotate from the easy axis to the direction of the applied field, the material is hence saturated.

![Figure 1.6. Magnetization process in monolythic Terfenol-D [10]](image)

On the other hand, stress dependency of material properties is due to couplings between properties of elasticity and quantum mechanical phenomena of magnetism [11]. In simple terms, magnetism results from imbalanced spin and orbital motion of electrons located in inner electron shells. Such imbalance can cause the atom to have non-spherical electron charge cloud whose orientation is coupled to the magnetic moment, enabling magnetic and elastic energy transfer.

![Figure 1.7. Development of magnetostriction [10]](image)

It also worth noting that magnetostriction only appears below the Curie temperature $T_c$. Otherwise, material turns to be paramagnetic, and its inner structure becomes isotropic. As being cooled through $T_c$, it turns ferromagnetic. Strains are developed inside, which is called spontaneous magnetostriction (denoted by $\lambda_s$ in Figure 1.7.b). With strong fields applied, domains coherently orient to the direction of the field. Then, strain reaches its maximum, called the saturation magnetostriction (denoted by $\lambda_s$ in Figure 1.7.c).
1.3 Review of existing models

1.3.1 Existing models for piezoelectric materials

1.3.1.1 Piezoelectric constitutive models

Although hysteresis and nonlinearity are intrinsic for piezoelectric constitutive relations, a couple of mechanisms (e.g., keeping the material in low or moderate operating regimes to linearize the response) can be employed to reduce hysteretic and nonlinear effects. More importantly, polarization and fields generated with the direct piezoelectric effect, which is of main concern in our context, exhibit an almost linear dependence on the stress for low input levels [10]. All in all, the linear piezoelectric constitutive model, which relates dielectric and elastic behaviors of piezoelectric materials, provides sufficient accuracy in most cases. Therefore, we only use the linear model. More details of this model will be presented in the next chapter when we develop piezoelectric FE models.

1.3.1.2 Piezoelectric system models

Generally, piezoelectric system models can be divided into lumped-parameter type and FE type. For the former, lumped parameters for the electric domains can be easily obtained due to the inherent capacitance of piezoelectric materials and resistances from the external load. For the mechanical domain, a single mechanical degree of freedom is usually employed for the prediction of system dynamics. As long as the mechanical domain lumped-parameters are available, mechanical and electric equilibrium equations can be coupled via piezoelectric constitutive equations, leading to transformer relations (see e.g. [12] for details). While such methods provide initial insights into the coupled system through solving computationally cheap equations, the solutions lack some important aspects, such as the dynamic mode shapes, accurate strain distribution, and effects of the latter two on the electrical side [13].

FE models (see e.g. [14]), on the other hand, are more advantageous as they are more flexible to model complicated configurations and capable to obtain full field numerical solutions. With this method, weak forms are generated based on equilibrium and constitutive equations. The resulting infinite dimensional problem is then projected onto finite dimensions. As such, the problem amounts to solving a discrete system. At the same time, FE methods are also computationally more intensive, especially for large 3D models. Therefore, developing model order reduction techniques that preserve versatilities of the FE method while alleviate the computational burden, seems like a prospective approach.
1.3.2 Existing models for magnetostrictive materials

1.3.2.1 Magnetostrictive constitutive models

As summarized in [15], early magnetostrictive constitutive models consist in adapting ferromagnetic models to incorporate magnetoelastic couplings. Examples are the modified Preisach model and Jiles-Atherton model. In the latter, for instance, the effect of stress on magnetization is incorporated through adding a stress-equivalent field term into the modified Langevin equation. These models can be used to simulate e.g. variation of magnetization with stress. However, they are, to some extent, purely mathematical tools that do not address the underlying physics with sufficient accuracy. There are also a class of free-energy models for uniaxial cases [15]. In these models, the Gibbs energy for a given applied field is expressed in function of magnetization, and additional terms that were created to incorporate stress effects. This class of models is reported to be more suited to explain the physics of hysteresis rather than realistic simulations.

In [16] a new energy-based model called the modified Armstrong model is introduced. The idea is to construct magnetocrystalline, magnetoelastic and magnetic field energy terms. Sum of the latter terms corresponding to magnetization of different orientations is evaluated. The probability of a certain magnetization orientation is determined with respect to (w.r.t.) the total energy of that orientation, the lower its total energy is the larger that probability will be. Macroscopic properties of the material can be obtained as averages of all possible orientations. Therefore, accurate evaluation of macroscopic properties requires considering a large number of such orientations. The modified Armstrong model is based on 98 such orientations. A further improvement is proposed in [17] where the number of orientation is reduced to six. This model is named the Discrete Energy-Averaged Model, or DEAM. It has been validated for Galfenol [18] and Terfenol-D [19]. Compared with other magnetostrictive constitutive models, DEAM is advantageous due to its accuracy and ease to be elaborated into a FE framework. Although there are other magnetostrictive constitutive models, they are generally one dimensional or, more or less, do not properly address the underlying physics.

1.3.2.2 Magnetostrictive system models

As introduced above, magnetostrictive materials are normally inhomogeneous. Consequently, modeling magnetostrictive devices involves a hierarchy of structures on different levels – constitutive modeling on the microscopic level and system modeling on the macroscopic level. Here constitutive modeling refers to the calculation of material constants, whereas system modeling refers to the calculation of state variables (e.g. magnetic fields and stresses). As with the
piezoelectric case, popular magnetostrictive system modeling approaches can be categorized into the lumped-parameter type and the FE type. However, with the former type it is difficult, or impossible, to account for material inhomogeneity. Therefore, only FE models are reviewed.

Among the reported magnetostrictive models, a considerable amount of them are implemented using commercial packages e.g., COMSOL Multiphysics. Take the latter for example; single field interfaces are predefined while couplings need to be established via e.g. initial stress and remanant flux density features. Values of initial stresses and remanant flux densities are specified as functions of respectively, magnetic field and mechanical strain [20]. With this approach, incorporating material constitutive relations or any other un-predefined features can be cumbersome, if ever possible. Other ones, which do not rely on commercial packages, consist in enforcing the coupling weakly. For example, in [21] magnetic and elastic problems are resolved individually, while couplings are enforced through magnetic or mechanical excitations. The shortcoming with this approach is evident; data needs to be transferred between physics; iterations are also needed to enforce equilibrium, which deteriorates the simulation speed. In [22] a strongly coupled magnetostrictive model is presented. In this model, magnetic and elastic problems are solved simultaneously, using the same mesh, which eventually forms a block of discrete equations. Within the block, diagonal parts correspond to individual physics, whereas off-diagonal parts correspond to couplings. Unknowns are mechanical displacements and vector magnetic potentials $A$. However, in [22] nodal elements are used to discretize $A$. This can be improved through discretizing $A$ using Whitney elements, or vector basis functions [23]. Another issue with the model in [22] is that the electrical potential $\phi$ is not considered, while it is proved that the $A$-$\phi$ formulation is more appropriate regarding stability and convergence rate, especially for magneto-dynamic problems [24].

Magnetostrictive FE models are also commonly found in literatures of two-phase magnetolectric laminated composites. This is because in the latter, magnetostrictive and piezoelectric materials are combined via strain, due to which magnetolectric modeling is, in fact, combination of individual magnetostrictive and piezoelectric modeling. For example, in [25] a nonlinear but two dimensional (2D) FE magnetostrictive model is presented. In this model magnetostriction is assumed to be a parabolic function of the magnetization, while a Langevin-type equation is employed to consider relations between magnetization and magnetic field. In our own research group, 2D nonlinear magnetostrictive models are also developed, in the context of magnetolectric modeling, where the effect of load impedances in magnetolectric devices can be accounted for [26]. As a first attempt, the Brauer model describing the initial magnetization curve is used.
Chapter 1. Introduction

This model gives good predictions for devices of various volume ratios, epoxy bonding materials, and bias fields. Recently, this approach is extended to model Rosen-type devices [27]. While these models are effective in certain cases, there is no doubt that a fully coupled 3D magnetostrictive model which better incorporates material nonlinearities and multiphysics couplings is indispensable.

1.4 Research objectives and outline

Our ultimate objective of the thesis is to build a 3D FE framework, in which piezoelectric and magnetostrictive materials involved multiphysics problems can be modeled and simulated. To this end, several challenges, as mentioned previously, shall be addressed.

- The first one lies in constitutive modeling. For piezoelectric materials, constitutive equations based on linear piezoelectricity can be utilized. Elaborating the latter into FE systems is also well explained in numerous piezoelectric modeling literatures. For magnetostrictive materials, the state-of-the-art constitutive models are energy-based. These models are favorable because they better describe underlying physics of the material, and are applicable for a larger range of cases. However, elaborating them into FE systems can be also very computationally expensive, due to the involvement of a hierarchy of multi-level modeling. More precisely, the magnetostrictive constitutive model is fed with state variables (usually including stress and magnetic field strength) at a specific location, and gives as outputs material constants for that location. State variables can be solved with FE models on the macroscopic level, while solving the constitutive model takes place on microscopic level. Therefore, for a given profile of state variables the microscopic problem needs to be solved a lot of times (in fact, as many as the number of microscopic volumes consisted in the system), not to mention possible involvements of nonlinearity (resulted from the recursive dependency of state variables in the FE system). As such, the objective is to implement constitutive models in an efficient way, so that solutions on the microscopic level do not deteriorate the simulation speed while integration of constitutive models and FE models is convenient.

- Another challenge comes from involvements of multiphysics fields including electromagnetic, elastic, thermal, and electric circuits. Without doubt, for problems with single fields dedicated FE methodologies are vastly available. By contrast, the availability becomes questionable when multiple physics and coupling in between need to be considered. We note that in a differential forms based FE framework, the problem can be easily resolved. In
such a framework, quantities of different type are discretized using appropriate types of elements, called Whitney elements. Properties on the continuous level can be preserved after discretization. In addition, problems of different fields can be modeled in a unified fashion; boundary conditions that are normally tedious to handle with conventional FEs can also be handled straightforwardly. Hence, we define our second objective to be establishing a differential forms based FE framework, in order to elaborate piezoelectric and magnetostrictive problems into the 3D numerical system. Meanwhile, thermal fields and electric circuit related models shall be incorporated in this framework, so that we can investigate, for example, effects of thermal losses and load impedances in a piezoelectric/magnetostrictive materials based system.

- While 3D FE models are superiors compared with their counterparts, in terms of providing full-field solutions, dealing with complicated geometries and boundary conditions, etc., they can also be problematic because of their large model size. For this reason, it is vital to develop corresponding model reduction techniques, with which advantages of FE models can be preserved while at affordable computing costs. We noticed that PGD is adequate to this end. Indeed, high computational costs of 3D multiphysics FE simulation frequently come in two flavors. One consists in solving the large system repeatedly in a similar setting, such as in the case of transient analysis where the number of time steps is huge. The other consists in solving the large system for a large number of times in different settings, for example, in the case of parametric analysis where the number of parameters is large. PGD tackles these problems through decomposition, with which the original large-size problem can be converted to a series of smaller-size sub-problems, thereby significantly reducing the problem complexity. Our third objective with PGD is to develop efficient algorithms so that transient and parametric simulations using our multiphysics FE models can be performed as efficiently as possible.

With these objectives in mind, we have accomplished a couple of contributions that are presented in the remainder of the thesis.

In Chapter 2, we describe our implementation of the multiphysics framework and constitutive models. Some preliminaries of FE modeling including the principle procedures of an FE analysis, and Whitney elements, are first provided. Introduction of equilibrium equations of electromagnetic, elastic and thermal fields, which are incorporated in our framework, comes aftermath. In § 2.3, we present our piezoelectric model. The model is based on linear piezoelectricity, with electrodes and electric loads considered. Unknowns of the model are mechanical displacements
and elastic potentials, which are interpolated using the nodal Whitney elements. Simulations of a piezoelectric bimorph and a surface acoustic wave (SAW) device are presented. In § 2.4, we introduce our magnetostrictive model. More precisely, we review some fundamentals of DEAM and describe our implementation of it. After that, FE models of magnetostrictive materials, as well as elaboration of DEAM into the latter are discussed. In our model, the $A - \phi$ formulation for magnetodynamics is used. Various elements including the nodal, edge, and facet Whitney elements are employed for discretization of involved quantities. Simulations of both linear and nonlinear magnetostrictive problems are presented. FE formulations of these models for linear tetrahedron elements are provided.

In Chapter 3, model reduction via separated representations is first reviewed. Basics of PGD and the method closely related to it – Proper Orthogonal Decomposition (POD) are briefly revisited. Our two PGD-based novel algorithms are then presented. The first one is dedicated to nonlinear magneto-thermal transient analysis, in which time constants of different fields are orders of magnitude different, and thus resulting in a large number of time steps. Our proposition is to decompose quantities into space and time associated components, with which the original problem of solving a large space system at sequential time steps is converted to solving space system and time system alternatively, thereby reducing the problem complexity. The second algorithm is dedicated to parametric electro-mechanical analysis. For such problems, the number of parameters is large, due to which the large-size FE system have to be solved for as many times, being extremely time-consuming. Our solution is to decompose ensemble of the problem into components associated with different parameters, after which we solve a series of smaller-size problems to obtain approximate solutions. At the end of the chapter, we also discuss extension of the above algorithms to other applications.

In the last chapter, conclusions and prospective of the thesis are provided.
Chapter 2. Multiphysics Finite Element Modeling

In this chapter, we present our multiphysics FE models for piezoelectric and magnetostrictive materials. Preliminaries on FEM are provided in section 1, with emphasis on Whitney elements. Next, equilibrium equations used for electromagnetic, elastic and thermal fields are revisited in section 2. The piezoelectric model is presented in section 3. Applications of the model on investigation of a bimorph and a SAW device are also described. Section 4 deals with the magnetostrictive model. Introduction to DEAM (as well as the implementation for it) and FE formulations are presented separately. Elaborating DEAM into the FE system is explained through a nonlinear magnetostrictive problem. Applications of the magnetostrictive model for source current involved problems and harmonic analysis are also presented.

2.1 Preliminaries on finite element modeling

2.1.1 The finite element method

FEM is a numerical technique for finding approximate solutions to boundary value problems (BVPs) that are defined in terms of partial differential Equations (PDEs). FEM was first developed in the 1950s and is nowadays utilized in various domains, thanks to their flexibility in modeling complicated geometry and capability in obtaining full field numerical solutions. Generally, a FE analysis can be divided into the following steps:

- Construct PDEs governing the problem of interest, based on physical laws; identify boundary and initial conditions of the problem.
- Build weak forms, using either the variational method or weighted residue methods, through which an error function associated with the approximated solutions is minimized.
- Triangulate the domain into simplex that are called finite elements; over each element, interpolate quantities with degrees of freedoms and FE basis functions.
- Sum the interpolants over the triangulation, which leads to a system of matrix energetically equivalent to the weak form; resolve the discrete system.
- Post-process, analyze the results, etc.

In the remainder of the thesis, we apply FEM to specific problems, for which the convention
of notation are as follows. We use *(in the majority of the cases)* lower case letters *e.g.* \(a\) to denote scalars, bold lower case letters *e.g.* \(a\) and plain capital letters *e.g.* \(A\) to denote vectors, bold capital letters *e.g.* \(A\) to denote matrices, and calligraphic letters *e.g.* \(\mathcal{A}\) to denote tensors of order higher than two (that shall appear in Chapter 3).

### 2.1.2 Whitney elements

For certain multivariable calculus problems, particularly electromagnetics, it is more direct to solve with differential forms than vector calculus. Because with the former, field properties such as curl-free or divergence-free, and appropriate continuity across material interfaces are preserved [28]. More importantly, when the system is discretized with differential forms based elements, which are called Whitney elements [29], the favorable characteristics can be inherited from the continuous to discrete level. In this part, differential forms and Whitney elements, particularly linear tetrahedral elements, are briefly introduced. More comprehensive treatment on these topics can be found in, *e.g.* [30], [31] and [32]. Here, we follow [28].

#### 2.1.2.1 Differential forms

In differential forms calculus, four types of entries called \(p\)-forms are utilized in 3D applications. Among them the 0-form and 3-form are scalar quantities whilst the 1-form and 2-form are vector quantities. Roughly speaking, a \(p\)-form takes a \(p\)-dimensional vector and gives a number. Each type of \(p\)-form belongs to a specific functional space, called Hilbert space. Meanwhile, various operators, corresponding to those in vector calculus, are defined on \(p\)-forms. For instance, the exterior derivative, a metric free operator, corresponds to the gradient, curl and divergence in vector calculus. The wedge operator \(\wedge\), also called exterior product, corresponds to the dot and cross products. Other frequently used operators including the Hodge star operator \(\star\), the pullback and push-forward operators are mainly used for coordinate transformations [28]. Properties of \(p\)-forms are summarized as follow.

\[
\omega^3 := \beta(y) dx_1 \wedge dx_2 \wedge dx_3 \tag{2.1}
\]

The 3-form is integrated over a volume and constant in the volume. It is best suited to represent quantities like scalar densities. A 3-form is shown in Equation (2.1) where \((x_1, x_2, x_3)\) is the basis of a manifold in \(\mathbb{R}^3\), \((dx_1, dx_2, dx_3)\) the basis of the cotangent space of the manifold, \(y\) a point in \(\mathbb{R}^3\), and \(\beta\) the value at \(y\).
2.1. Preliminaries on finite element modeling

\[
L^2(\Omega) := \left\{ u; \int_\Omega u^2 d\Omega < \infty \right\}
\]

\[
\|u\|_{L^2} := (\|u\|^2)^{1/2}
\] (2.2)

The Hilbert space where \(\omega^3\) lives in, is denoted by \(L^2\) whose definition and associated norm are shown in Equation (2.2).

A 0-form, as in Equation (2.3), is defined on a point, giving a scalar of the function at that point. It is utilized to represent potential variables as it is continuous along all orientations.

\[
\omega^0 := \beta(y)
\] (2.3)

Concurrently a continuous scalar function represented by \(\omega^0\), belongs to the functional space \(H(\text{grad})\) whose definition and norm are shown in Equation (2.4).

\[
H(\text{grad}, \Omega) := \left\{ u; u \in L^2(\Omega); \text{grad}(u) \in \left(L^2(\Omega)\right)^3 \right\}
\]

\[
\|u\|_{H(\text{grad},\Omega)} := (\|u\|^2 + \|\text{grad}(u)\|^2)^{1/2}
\] (2.4)

The 1-form, as shown in Equation (2.5), is integrated over a line. It has tangential continuity which makes it suited for the representation of field quantities such as electric field.

\[
\omega^1 := \beta_1(y)dx_1 + \beta_2(y)dx_2 + \beta_3(y)dx_3
\] (2.5)

Functions expressed as 1-forms are defined in the functional space \(H(\text{curl})\) whose definition and associated norm are presented in Equation (2.6).

\[
H(\text{curl}, \Omega) := \left\{ U; U \in \left(L^2(\Omega)\right)^3; \text{curl}(U) \in \left(L^2(\Omega)\right)^3 \right\}
\]

\[
\|U\|_{H(\text{curl},\Omega)} := (\|U\|^2 + \|\text{curl}(U)\|^2)^{1/2}
\] (2.6)
Chapter 2. Multiphysics Finite Element Modeling

The 2-form, integrated over surfaces, has the form of Equation (2.7). It is continuous in the normal direction, making it suitable for representing flux quantities.

\[
\omega^2 := \beta_1(y)dx_2 \wedge dx_3 + \beta_2(y)dx_3 \wedge dx_1 + \beta_3(y)dx_1 \wedge dx_2 \tag{2.7}
\]

Objects expressed as a 2-form is defined in the functional space \( \mathcal{H}(\text{div}) \), as depicted in (2.8).

\[
\mathcal{H}(\text{div}, \Omega) := \{ U: U \in (L^2(\Omega))^3; \text{div}(U) \in L^2(\Omega) \}
\]

\[
\|U\|_{\mathcal{H}(\text{div}, \Omega)} := (\|U\|^2 + \|\text{div}(U)\|^2)^{1/2}
\]

Between differential forms the exterior derivative is defined, which linearly maps a \( p \)-form to a \( (p+1) \)-form, as expressed in Equation (2.9).

\[
d: \Lambda^p \rightarrow \Lambda^{p+1}; \quad \omega \mapsto d\omega \quad (p = 0,1,2)
\]

An exact sequence of \( p \)-forms can be generated with the exterior derivative operator, as depicted in Equation (2.10) where corresponding derivative operators in vector calculus are shown in blue. The sequence suggests that the exterior derivative of a \( p \)-form belongs to a subspace of the space of the \( (p+1) \)-form.

\[
\omega^0 \overset{d}{\rightarrow} \omega^1 \overset{d}{\rightarrow} \omega^2 \overset{d}{\rightarrow} \omega^3 \tag{2.10}
\]

Moreover, the generalized Stokes law can be expressed with the exterior derivative operator

\[
\int_{\Omega} d\omega^p = \int_{\partial\Omega} \omega^p \tag{2.11}
\]

The equation implies that integrating the exterior derivative of an \( p \)-form over the whole manifold \( \Omega \) equals to the integral of that \( p \)-form over the oriented boundary of the manifold.
Namely, the boundary operator and exterior derivative are dual to each other. On the other hand, successively applying the exterior derivative results in the trivial form \( i.e. d(d\omega^p) = 0 \). Physically, this correlates to, for example, the curl-free property of the electric field and the divergence-free property of the magnetic flux density.

### 2.1.2.2 Whitney elements

The edge elements were initialized by Nédélec [33] while the link between differential forms and FEs was advanced by Bossavit who also created the term Whitney elements [29]. An exhausted history review is outside the scope here, but it can be found in e.g. [30]. Here we show the involvement of Whitney elements in discretization via an abstract problem.

Consider a given Hilbert space \( V \), a bilinear continuous form \( a(\cdot, \cdot) \) defined on \( V \times V \), and a continuous linear form \( l(\cdot) \) defined on \( V \). The abstract problem can be defined in the weak form: find \( u \in V \), such that \( a(u, v) = l(v) \), for all \( v \in V \). \( V \) is the test space of \( p \)-forms with appropriate boundary conditions. Using the Galerkin method, the problem is projected onto a finite dimensional space \( V_h \subseteq V \), associated with a triangulation of the domain with \( h \) characterizing the triangulation. The finite dimensional weak form reads: find \( u_h \in V_h \), such that \( a(u_h, \vartheta_h) = l(\vartheta_h) \), for all \( \vartheta_h \in V_h \). Denoting the error between the exact solution \( u \) and the approximation \( u_h \) as \( \epsilon_h \), the Galerkin orthogonality can be deduced, as in (2.12).

\[
\tilde{a}(\epsilon_h, v) := a(u_h, \vartheta_h) - l(\vartheta_h) = 0 \tag{2.12}
\]

The equation implies that the residual, which is obtained by inserting \( u_h \) into the original equation and taking the difference between both sides, is orthogonal to the test functional space. In general, bases for \( V_h \) and \( V_h \) are identical polynomial functions. These functions have compact supports, meaning that each basis function is only nonzero over a specific subdomain – an element \( K \). The solution \( u \) over element \( K \) is interpolated with \( N \) bases functions on the element and degrees of freedom (DoFs) \( \alpha_i \), as presented in Equation (2.13). Briefly, a FE is the combination of the polynomial space \( P \), the compact support \( K \) and DoFs.

\[
I_K(u) := \sum_{i=1}^{N} \alpha_i (u) v_i \tag{2.13}
\]

Over the triangulation \( T \) containing all the elements \( K_i \), the solution \( u \) is interpolated as the
sum of local interpolants over each $\mathcal{K}_i$, see Equation (2.14).

$$I_T(u)|_{\mathcal{K}_i} := I_{\mathcal{K}_i}(u), \quad \forall \mathcal{K}_i \in \mathcal{T} \tag{2.14}$$

In this thesis, tetrahedrons are primly utilized for triangulation. Hence, $\mathcal{K}$ in what follows represent a tetrahedron if not otherwise specified. Obviously, four polynomial spaces $\mathcal{P}$ and four types of degrees of freedom $\mathcal{A}$ are needed for the discretization of $p$-forms. Such spaces are listed in Equation (2.15) where $\mathcal{P}_0, \mathcal{P}_1, \mathcal{P}_2$ and $\mathcal{P}_3$ are respectively, of dimension, 4, 6, 4, and 1, relating to the number of node, edge, facet and volume elements (see Figure 2.1).

$$\mathcal{P}_0 := \{u: u \in \mathcal{P}_1; u = a_0 + a_1 x_1 + a_2 x_2 + a_3 x_3; a_i \in \mathbb{R}\}$$

$$\mathcal{P}_1 := \{U: U = (u_1, u_2, u_3) \in (\mathcal{P}_1)^3; u_{i,j} = 0; u_{i,j} + u_{j,i} = 0, i \neq j\}$$

$$\mathcal{P}_2 := \{U: U \in ((\mathcal{P}_1)^3 + \mathcal{P}_0 \cdot r); U = (a_0 + a_3 x_1, a_4 + a_3 x_2, a_5 + a_3 x_3); a_i \in \mathbb{R}\}$$

$$\mathcal{P}_3 := \{u: u \in \mathcal{P}_0; u = a_0; a_0 \in \mathbb{R}\}$$

![Figure 2.1. Numbering of Whitney elements on a tetrahedron](image)

**Figure 2.1.** Numbering of Whitney elements on a tetrahedron

node elements: $\omega_{\mathcal{P}}^{\mathcal{P}} := \lambda_i$

edge elements: $\omega_{\mathcal{A}}^{\mathcal{E}} := \lambda_i \text{grad} \lambda_j - \lambda_j \text{grad} \lambda_i$

facets elements: $\omega_{\mathcal{A}}^{\mathcal{F}} := 2 \left( \lambda_i \text{grad} \lambda_j \times \text{grad} \lambda_k + \lambda_j \text{grad} \lambda_k \times \text{grad} \lambda_i + \lambda_k \text{grad} \lambda_i \times \text{grad} \lambda_j \right)$

volume elements: $\omega_{\mathcal{V}}^{\mathcal{V}} := 1/|\mathcal{K}_i|$
2.2 Equilibrium equations

Basis functions of the four types of elements are summarized in Equation (2.16) where \( \lambda_i \) indicates the barycentric coordinate associated with the i-th vertex \( n_i \). The value of \( \lambda_i \) ranges between 0 and 1, reaching its maximum 1 at \( n_i \) and linearly decreasing to 0 on the facet opposing to \( n_i \). \( |K_i| \) in the volume element basis function represents the volume of \( K_i \). Formulations utilized to calculate barycentric coordinates are listed in § A.1.

Meanwhile, DoFs represent discrete \( p \)-form coefficients. For node elements, they are values of the quantity on the vertices; For edge elements, they are circulations of the field over edges \( \int_{\Omega_e} U \cdot t \), where \( t \) is denotes the unit tangential vector. For facet element, they are flux through facets \( \int_{\Omega_f} U \cdot n \), where \( n \) is a unit outward normal to the boundary of the facet. For volume elements, they are volume integral of the distribution.

It is noted that when interpolants are substituted into the bilinear form in Equation (2.12), the unified single operator – exterior derivative in Equation (2.10) leads to three discrete operators: \( G \), \( C \) and \( D \), called the incident matrices, corresponding to respectively, the gradient, curl and divergence operator in vector calculus. Entries in these discrete operators are 1, 0 or -1, decided by the orientations and incident relations of the involved geometric entries [34].

2.2 Equilibrium equations

Next, we briefly revisit equilibrium equations for multiphysics fields. They are the Maxwell’s Equations for electromagnetic fields, the heat conduction equation for thermal fields, and balance of linear momentum for elastic fields. While differential forms are employed in the previous section, equations here are presented in vector calculus because we use nodal Whitney elements, (also called Lagrange elements), for both elastic and thermal quantities in the implementation (other \( p \)-forms based elements are used for electromagnetic quantities), for which vector calculus representations seem to be more common. That being said, differential forms are employed for introduction of Whitney elements, rather than continuous field equations. Nevertheless, thorough differential forms based treatments on the involved problems can be found in literatures. See e.g. [34] on magnetics, [35], [36] on elastics and [37] on thermal problems.

2.2.1 Maxwell’s Equations

Maxwell’s Equations are shown in Equation (2.17). The first is the Maxwell-Ampère’s law;
second the Faraday’s law; third and fourth respectively, Gauss’ magnetic and electric law. Quantities in the equations can be grouped into three categories – field intensities including electric field \( E(V \cdot m^{-1}) \) and magnetic field \( H(A \cdot m^{-1}) \), flux densities including electric displacement \( D(C \cdot m^{-2}) \), magnetic induction \( B(T) \), and current density \( J(A \cdot m^2) \), and volume density - the electric charge density \( \rho_e (C \cdot m^{-3}) \). On the discrete side, they are represented with respectively, edge, facet and volume elements. In practice potentials are employed to express field variables. We use electric potential \( \varphi(V) \) and magnetic vector potential \( A(V \cdot s \cdot m^{-1}) \). They are represented by node and edge elements, respectively.

\[
\begin{align*}
\text{curl} H &= J + \partial D / \partial t \\
\text{curl} E &= - \partial B / \partial t \\
\text{div} B &= 0 \\
\text{div} D &= \rho_e \\
D &= \varepsilon_0 E + P \\
B &= \mu_0 (H + M) \\
J &= \sigma_e E
\end{align*}
\]

Field and flux quantities are related by constitutive equations. See Equation (2.18) in which \( \varepsilon_0 \) and \( \mu_0 \) are the permittivity and permeability of free space, equal to, \( 8.854 \cdot 10^{-12} (F \cdot m) \) and \( 4\pi \cdot 10^{-7} (H \cdot m^{-1}) \), respectively. \( \sigma_e (S \cdot m^{-1}) \) denotes the electric conductivity.

2.2.2 Heat conduction equation

Heat conduction is governed by

\[
\rho c_p \partial \theta / \partial t + \text{div} \varphi - s = 0
\]

in which \( \rho(kg \cdot m^{-3}) \) represents the mass density per unit volume, \( c_p(J \cdot K^{-1}) \) specific heat capacity at constant pressure, \( \theta(K) \) temperature, \( \varphi(W \cdot m^{-2}) \) heat flux, and \( s(W \cdot m^3) \) the rate of heat generation per unit volume. According to the Fourier’s theorem, \( \varphi \) can be expressed using temperature gradients, as expressed in (2.20) where the minus sign implies flux
flows from higher to lower temperatures.

\[ \phi = -\kappa \cdot \text{grad} \theta \quad (2.20) \]

Meanwhile, \( \kappa (W \cdot m^{-1} \cdot K^{-1}) \) in the equation represents the thermal conductivity. Substituting the Fourier’s theorem into Equation (2.19) yields Equation (2.21) that, in conjunction with proper initial and boundary conditions, is employed to describe thermal fields.

\[ \rho c_p \frac{\partial \theta}{\partial t} - \text{div}(\kappa \cdot \text{grad} \theta) - s = 0 \quad (2.21) \]

2.2.3 Balance of linear momentum

The balance of momentum describes movements and deformations of solid matter in spatial coordinates. In general, the basis of the spatial coordinates is denoted by minuscule \( x(x, y, z) \). A reference coordinate is also needed in order to describe material properties before solution. The basis of the reference coordinate system is denoted by capitals \( X(X, Y, Z) \).

\[ x(X, t) = X + u(X, t) \quad (2.22) \]

The reference coordinates are constant over solution whereas the spatial coordinates changes. The relation between these two coordinate systems is shown in Equation (2.22) where \( u(u, v, w) \) denotes the displacement vector. In this thesis geometric linearity is assumed, which means no distinction between them is made. In other words, equations are formulated w.r.t. the undeformed state. This assumption is viable, as long as deformations are small enough, so that errors introduced by ignoring the deformation are negligible. Accordingly, the Engineering strain, Cauchy stress and linear elastic material models [38] are employed, as addressed below. Otherwise, geometric nonlinearity needs to be considered, which means the reference configuration is updated with deformation after each solution. The Green-Lagrange strain and the Second Piola-Kirchoff stress [38] shall be utilized in the latter case.

To relate displacements with forces, we first introduce the engineering strain tensor \( \varepsilon \), as shown in (2.24). Normal strain entries are defined as \( \varepsilon_{ii} = \frac{\partial u_i}{\partial x_i} \), while shear strain entries, which are denoted by \( \varepsilon_{ij} \) satisfying \( \varepsilon_{ij} = \varepsilon_{ji} \) for \( i \neq j \). \( u_i \) and \( x_i \) are components of \( u \) and \( x \) in Equation (2.22), with \( i, j = 1, 2, 3 \). The symmetric-gradient operator (which is denot-
ed by $\overrightarrow{\text{grad}}$) is applied to the displacement, to obtain the strain tensor (see the expression in compact form in (2.23) where the superscript ‘$t$’ implies transpose).

$$\overrightarrow{\text{grad}} \boldsymbol{u} := \frac{1}{2} (\text{grad} \boldsymbol{u} + (\text{grad} \boldsymbol{u})^t). \quad (2.23)$$

The strain tensor $\boldsymbol{\varepsilon}$ has six independent entries that can be denoted by $[\varepsilon_1, \varepsilon_2, \varepsilon_3, \varepsilon_4, \varepsilon_5, \varepsilon_6]^t$. The vector corresponds to $[\varepsilon_{11}, \varepsilon_{22}, \varepsilon_{33}, 2\varepsilon_{12}, 2\varepsilon_{23}, 2\varepsilon_{13}]^t$ in the Standard form, or $[\varepsilon_{11}, \varepsilon_{22}, \varepsilon_{33}, 2\varepsilon_{13}, 2\varepsilon_{12}, 2\varepsilon_{13}]^t$ in the Voigt form. In § A.2, FE formulations for expressing $\boldsymbol{\varepsilon}$ in terms of $\boldsymbol{u}$ are presented.

$$\boldsymbol{\varepsilon} := \begin{bmatrix} \varepsilon_1 & \varepsilon_{12} & \varepsilon_{13} \\ \varepsilon_{12} & \varepsilon_2 & \varepsilon_{23} \\ \varepsilon_{13} & \varepsilon_{23} & \varepsilon_3 \end{bmatrix} \quad (2.24)$$

Also utilized in linear elasticity is the Cauchy stress, which refers to the quantity of the force divided by area, considering deformation in the current configuration. The Cauchy stress tensor $\boldsymbol{\sigma}$ is also symmetric, sharing similar representations as the engineering strain tensor. In the compact form, it is usually denoted by $\boldsymbol{\sigma} = [\sigma_1, \sigma_2, \sigma_3, \sigma_4, \sigma_5, \sigma_6]^t$. Interpretation of subscripts for entries of the latter vector depends on whether the Standard or Voigt form is used.

**Table 2.1. Representation of the elasticity tensor for different materials**

<table>
<thead>
<tr>
<th>isotropic</th>
<th>orthotropic</th>
<th>anisotropic</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\lambda + 2\mu$</td>
<td>$\lambda$</td>
<td>$\lambda$</td>
</tr>
<tr>
<td>$\lambda$</td>
<td>$\lambda + 2\mu$</td>
<td>$\lambda$</td>
</tr>
<tr>
<td>$\lambda$</td>
<td>$\lambda + 2\mu$</td>
<td>$\lambda$</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>$\mu$</td>
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<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>$c_{11}$</td>
<td>$c_{12}$</td>
<td>$c_{13}$</td>
</tr>
<tr>
<td>$c_{12}$</td>
<td>$c_{22}$</td>
<td>$c_{23}$</td>
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<tr>
<td>$c_{13}$</td>
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<tr>
<td>$c_{14}$</td>
<td>$c_{24}$</td>
<td>$c_{34}$</td>
</tr>
<tr>
<td>$c_{15}$</td>
<td>$c_{25}$</td>
<td>$c_{35}$</td>
</tr>
<tr>
<td>$c_{16}$</td>
<td>$c_{26}$</td>
<td>$c_{36}$</td>
</tr>
</tbody>
</table>

The stress tensor can be obtained using the Hooke’s law, as shown in Equation (2.25) where $\boldsymbol{\sigma}_{\text{ex}}$ indicates stresses from external sources (those unrelated to strain e.g. the residual stress after heat treatment). $\boldsymbol{\varepsilon}_{\text{inel}}$ represents the sum of all strains (e.g. that due to temperature change, piezoelectric or magnetostrictive coupling effects) except for the Engineering strain. $\boldsymbol{c}$ is the fourth order elasticity tensor, whereas the operator “.” represents the double-dot tensor product. $\boldsymbol{c}$ is symmetric, having 21 independent entries in the general case.
2.3 Piezoelectric model

\[
\sigma = \sigma_{\text{ex}} + c : (\varepsilon - \varepsilon_{\text{inel}}) \tag{2.25}
\]

Depending on the property of the material, the elasticity tensor can be presented in different forms, as summarized in Table 2.1. For isotropic materials, it can be represented using the Lamé parameters, or a set of two other equivalent variables; for orthotropic materials, nine independent parameters are needed; for general anisotropic materials the number is 21. Eventually, the balance of linear momentum can be expressed as in Equation (2.26).

\[
\rho \frac{\partial^2 \mathbf{u}}{\partial t^2} + \text{div}\sigma - \mathbf{f}_v = 0 \tag{2.26}
\]

where \(\rho\) indicates the mass density per unit volume, \(\mathbf{f}_v\) the volume force vector. Forces on the boundary are incorporated into the system after enforcing boundary conditions. In the following, we present FE models, for which different conventions are employed. Constitutive equations are presented in Einstein summation convention. FE systems are presented in matrix convention, which facilitates implementation.

2.3 Piezoelectric model

In the Chapter 1, microscopic mechanisms and existing modeling methods of the piezoelectric material are briefly introduced. Here we present a piezoelectric FE model based on linear piezoelectricity. In the FE model, dependent variables are mechanical displacements and electric potentials which are both approximated using the 0-form based Whitney elements. Elaborating electric loads and electrodes into the FE model is also discussed. For demonstration, the model is utilized for simulations of a bimorph energy harvester and a SAW device.

2.3.1 Linear theory of piezoelectricity

Under the assumption of linear piezoelectricity, the equations of linear elasticity, i.e. Equation (2.26), and the equation of electrostatics are coupled through piezoelectric coefficients. To avoid notation confusions between mechanical and electrical quantities, the strain tensor is denoted by \(\mathcal{S}\) in lieu of \(\varepsilon\), the stress tensor is denoted by \(T\) in lieu of \(\sigma\), as recommended by the Standard [39]. Other notations introduced in the section of equilibrium equations remain unchanged, if not otherwise specified.
2.3.1.1 Constitutive equations

Denote the stored energy density for a piezoelectric continuum by $U$. By the conservation of energy and the first law of thermodynamics, the time derivative of $U$ can be divided into elastic and electrical parts, as shown in Equation (2.27) where $i, j = 1, 2, 3$. Note that summation convention is employed in equations of this section, which is recommended in the *IEEE Standard on Piezoelectricity* [39].

\[
\dot{U} = T_{ij} \dot{S}_{ij} - E_i \dot{D}_i
\]  
(2.27)

In the meanwhile, the electric enthalpy $\bar{H}$ is defined by

\[
\bar{H} := U - E_i D_i
\]  
(2.28)

Taking derivative on both sides of Equation (2.28) and substituting Equation (2.27) into the resultant equation yields Equation (2.29).

\[
\dot{\bar{H}} = T_{ij} \dot{S}_{ij} - D_i \dot{E}_i
\]  
(2.29)

The partial derivatives in (2.30) can be obtained based on (2.29).

\[
T_{ij} = \frac{\partial \bar{H}}{\partial S_{ij}|_E}, \quad D_i = -\frac{\partial \bar{H}}{\partial E_i|_S}
\]  
(2.30)

On the other hand, $\bar{H}$ takes the form of Equation (2.31) for linear piezoelectricity.

\[
\bar{H} = \frac{1}{2} c_{ijkl}^E S_{ij} S_{kl} - e_{klj} E_k S_{ij} - \frac{1}{2} \varepsilon_{ij}^E E_i E_j
\]  
(2.31)

where $c_{ijkl}^E (Pa)$, $e_{klj} (C \cdot m^{-2})$ and $\varepsilon_{ij}^E (F \cdot m^{-1})$ are the elastic, piezoelectric and dielectric constants, respectively. Superscripts $E$ and $S$ suggest values under constant electric field and strain, respectively. As a result, constitutive equations can be yield as (2.32).
2.3. Piezoelectric model

\[ T_{ij} = c_{ijkl}^{E} S_{kl} - e_{klj} E_k \]  
\[ D_i = e_{kl}^{S} S_{kl} + \varepsilon_{ij}^{S} E_k \]  

(2.32)

In practice, strain and electric field are expressed in terms of displacements and electric potential, respectively (see (2.33)), so that (2.34) holds for \( i, j = 1,2,3 \).

\[ S_{ij} = \frac{1}{2} (u_{ij} + u_{ji}) \quad , \quad E_i = \varphi_i \]  

(2.33)

Next, substituting (2.32) into equilibrium equations (2.26) and (2.17) yields (2.34). Note that the electric field is in fact time dependent, as it is coupled with dynamic elastic fields. Nevertheless, full electromagnetic equations are not necessary because phase velocities of elastic waves are several orders of magnitude less than velocities of electromagnetic waves [39].

\[ \rho \ddot{u}_j - e_{ijkl}^{E} u_{k,li} - e_{kj} e_{kl} \varphi_{kl} - f_j = 0 \]  
\[ e_{klj} u_{i,kl} - \varepsilon_{ij}^{S} \varphi_{ij} = 0 \]  

(2.34)

where a comma subscript followed by an orientation index denotes a derivation w.r.t. the corresponding orientation. For conciseness, \( u, v \) and \( w \) are denoted by \( u_i \) for \( i = 1,2,3 \), which are components of the mechanical displacement. In practice, contracted subscripts (i.e. the Standard or Voigt form as introduced in § 2.2.3) are usually utilized, so that the number of indices in the subscripts of \( c \) and \( e \) are reduced to two.

2.3.1.2 Elasto-piezo-dielectric material constants

When applying constitutive equations to particular piezoelectric materials, it is important to identify three coordinate systems: the crystallographic coordinate system (denoted by \( abc \)), the reference coordinate system (\( X_1X_2X_3 \)), and the spatial coordinate system (\( xyz \)). The crystallographic system, provided by the crystal itself, refers to the natural coordinate system, in terms of which properties of the crystal are described. The reference coordinate system, on the other hand, is a right-handed system, used to define material constants. The rules for determining positive sense of this system is as follow [39]. For the first axis, the first non-zero in the following list: \( e_{33}, e_{11}, e_{22}, e_{36} \) and \( e_{34} \), shall be positive. The same applies to the sec-
ond axis except that the second non-zero in the list should be positive. The last axis forms a right-handed system with the previous two axes. As a result, $e$ constants have unambiguous meanings. For instance, a positive $e_{33}$ implies that tensile stress $\varepsilon$ (by convention, the tensile stress is chosen to be positive), parallel to $X_3$ leads to an electric tension whose positive terminal is on the $+X_3$ face. Lastly, the spatial coordinate system $xyz$ is used for specifying boundary conditions and excitations. Coordinate system transforms are needed when spatial and reference systems do not coincide. Rotating matrices defined in [40] can be used in the latter case.

2.3.2 Finite element formulations

To describe the piezoelectric problem, consider a 3D domain $\Omega_\varphi \in \mathbb{R}^3$ with Lipschitz boundary $\Gamma_\varphi := \Gamma_{\varphi_D} \cup \Gamma_{\varphi_N}$ where $\Gamma_{\varphi_D}$ and $\Gamma_{\varphi_N}$ are, respectively, Dirichlet and Neumann electrical boundaries. The mechanical domain $\Omega_u$, which is a subdomain of $\Omega_\varphi$ ($\Omega_u \subset \Omega_\varphi$), has its Lipschitz boundary $\Gamma_u := \Gamma_{u_D} \cup \Gamma_{u_N}$, with $\Gamma_{u_D}$ and $\Gamma_{u_N}$ being Dirichlet and Neumann mechanical boundary, respectively.

![Figure 2.2. Electric and mechanical domains of the multiphysics problem](image)

A common practice in modeling piezoelectric materials is to choose $\Omega_\varphi$ coinciding with $\Omega_u$, which, however, does not appropriately account for fringing effects, from the electrical modeling point of view. We choose the domains so that $\Omega_\varphi / \Omega_u \neq \emptyset$, as shown in Figure 2.2, where $\Omega_\varphi / \Omega_u$ represents an air domain. Strong forms of the coupled problem consists in finding displacements and electrical potentials subjecting to certain boundary conditions, as shown in the coupled mechanical equation (2.35)

$$\begin{align*}
c^E \nabla \left( \nabla \cdot \mathbf{u} \right) + \mathbf{e}^T \nabla \left( \nabla \varphi \right) + \mathbf{f} &= \mathbf{\rho} \ddot{\mathbf{u}} \quad \text{in} \; \Omega_u \\
\mathbf{u} &= \mathbf{u}_0 \quad \text{on} \; \Gamma_{u_D} \\
\frac{\partial T}{\partial \mathbf{n}} &= T_0 \quad \text{on} \; \Gamma_{u_N}
\end{align*}$$ (2.35)
2.3. Piezoelectric model

and the coupled electrostatic equation (2.36)

\[ e \, \text{grad} \left( \text{grad} \, u \right) - \varepsilon_s \, \text{grad} \left( \text{grad} \, \varphi \right) = 0 \quad \text{in } \Omega_u \]

\[ - \varepsilon_s \, \text{grad} \left( \text{grad} \, \varphi \right) = 0 \quad \text{in } \Omega_\varphi / \Omega_u \]

\[ \varphi = \varphi_0 \quad \text{on } \Gamma_{\varphi_D} \]

\[ \partial \varphi / \partial n = 0 \quad \text{on } \Gamma_{\varphi_N} \]

(2.36)

where \( u_0, T_0, \) and \( \varphi_0 \) are predefined values for respectively, displacements, surface tractions, and electrical potentials. \( n \) is the unit outward normal to the boundary. The symbol ‘\( \cdot \)’ indicates second order time derivative.

Meanwhile, electrodes, which are deposited on piezoelectric units, impose equipotential electrical conditions. Mechanically, their influences are negligible as they are very thin. Hence, electrodes can be considered as surfaces, involving only equipotential conditions. We address modeling for electrodes and electrical loads after setting up elementary FE systems.

2.3.2.1 Finite element approximations

First, we define some integrals, as presented in Equation (2.37).

\[ B_{UU}(u', u)_{\Omega_u} := \int_{\Omega_u} u' \left( e \, \text{grad} \left( \text{grad} \, u \right) - \rho \ddot{u} \right) d\Omega \]

\[ B_{U\Phi}(u', \varphi)_{\Omega_u} := \int_{\Omega_u} u' \, e \, \text{grad} \left( \text{grad} \, \varphi \right) d\Omega \]

\[ B_{\Phi U}(\varphi', u)_{\Omega_u} := \int_{\Omega_u} \varphi' \, e \, \text{grad} \left( \text{grad} \, u \right) d\Omega \]  \hspace{1cm} (2.37)

\[ B_{\Phi \Phi}(\varphi', \varphi)_{\Omega_\varphi} := \int_{\Omega_\varphi} \varphi' \, e \, \text{grad} \left( \text{grad} \, \varphi \right) d\Omega \]

\[ (u', f)_{\Omega_u} := \int_{\Omega_u} u' \, f \, d\Omega, \quad (u', T_0)_{\Gamma_{uN}} := \int_{\Gamma_{uN}} u' \, T_0 \, d\Gamma \]
Then consider the weak form: find \( \mathbf{u} \in (\mathcal{U})^3 \) and \( \varphi \in \mathcal{V} \) such that

\[
\begin{align*}
B_{UU}(\mathbf{u}', \mathbf{u})_{\Omega_u} + B_{U\Phi}(\mathbf{u}', \varphi)_{\Omega_u} &= -(\mathbf{u}', \mathbf{f})_{\Omega_u} + (\mathbf{u}', \mathbf{T}_0)_{\Gamma_{\text{ny}}} \quad \forall \mathbf{u}' \in (\hat{\mathcal{U}})^3 \\
B_{\Phi\Phi}(\varphi', \varphi)_{\Omega_\varphi} + B_{\Phi U}(\varphi', \mathbf{u})_{\Omega_u} &= 0 \quad \forall \varphi' \in \hat{\mathcal{V}}
\end{align*}
\] (2.38)

In (2.38), \( \mathcal{U} \) and \( \mathcal{V} \) are Hilbert spaces - \( \mathcal{H}(\text{grad}, \Omega) \); \( \hat{\mathcal{U}} \) and \( \hat{\mathcal{V}} \) are, respectively, subspaces of \( \mathcal{U} \) and \( \mathcal{V} \) subject to boundary conditions. Discrete counterparts of Equation (2.38) read: find \( \mathbf{u}_h \in (\mathcal{U}_h)^3 \) and \( \varphi_h \in \mathcal{V}_h \) so that equations in (2.39) hold. In the equation, \( \mathcal{U}_h \subset \mathcal{U}, \mathcal{V}_h \subset \mathcal{V}, \mathcal{U}_h \subset \mathcal{U}, \mathcal{V}_h \subset \mathcal{V} \), and \( \hat{\mathcal{V}}_h \subset \hat{\mathcal{V}} \) are finite dimensional subspaces associated with the triangulation, with \( h \) characterizing resolution of the latter.

\[
\begin{align*}
B_{UU}(\mathbf{u}_h', \mathbf{u}_h)_{\Omega_u} + B_{U\Phi}(\mathbf{u}_h', \varphi_h)_{\Omega_u} &= -(\mathbf{u}_h', \mathbf{f})_{\Omega_u} + (\mathbf{u}_h', \mathbf{T}_0)_{\Gamma_{\text{ny}}} \quad \forall \mathbf{u}_h' \in (\hat{\mathcal{U}})^3 \\
B_{\Phi\Phi}(\varphi_h', \varphi_h)_{\Omega_\varphi} + B_{\Phi U}(\varphi_h', \mathbf{u}_h)_{\Omega_u} &= 0 \quad \forall \varphi_h' \in \hat{\mathcal{V}}_h
\end{align*}
\] (2.39)

Within an element \( \mathcal{K}_d \), the weak form can be build using the Galerkin method, with the trial functions expressed as in Equation (2.40) where \( n_\text{n} \) is the number of vertices in \( \mathcal{K}_d \). \( \omega^e_{al} \) are 0-form based nodal basis functions. \( u^e_j \) and \( \varphi^e_j \) are nodal values associated with vertices. Practically, it is more convenient to arrange unknowns in vectors. In this case, entries of \( \mathbf{u}_h \) \( i.e. u^e_j \)'s, \( j = 1,2,3 \) can be written as \( \mathbf{U}_h = [u_1, v_1, w_1] \) in which \( u \), \( v \) and \( w \) are recovered from \( u_i \) for \( i = 1,2,3 \).

\[
\begin{align*}
\mathbf{u}_h^{(\alpha)} &= \sum_{i=1}^{n_\text{n}} \omega^e_{al} u^e_j \mathbf{j}^{(\alpha)} = \sum_{i=1}^{n_\text{n}} \omega^e_{al} \varphi^e_j \\
\end{align*}
\] (2.40)

Re-writing (2.40) in compact form gives (2.41).

\[
\begin{align*}
\mathbf{U}_h^{(\alpha)} := W^{n_u}_a U, \quad \varphi_h^{(\alpha)} := W^{n_\Phi}_a \Phi
\end{align*}
\] (2.41)

where \( U := [u_1^e \ v_1^e \ w_1^e \ \cdots \ w_{n_\text{n}}^e]^T \), \( \Phi := [\varphi_1^e \ \varphi_2^e \ \cdots \ \varphi_{n_\Phi}^e] \), \( W^{n_u}_a := [\omega^e_{al} I_3 \ \omega^e_{al} I_3 \ \cdots \ \omega^e_{al} n_3] \), \( W^{n_\Phi}_a := [\omega^e_{al} \ \omega^e_{al} \ \cdots \ \omega^e_{al} n_3] \), \( I_3 \) is the third order identity matrix.
2.3. Piezoelectric model

\[ M_u^{(\alpha)} \ddot{u}_\alpha + K_u^{(\alpha)} u_\alpha + K_{u\varphi}^{(\alpha)} \phi_\alpha = F_u^{(\alpha)} \quad \text{for } K_\varphi \in \Omega_u \]

\[ K_{u\varphi}^{(\alpha)^T} u_\alpha - K_{\varphi}^{(\alpha)} \phi_\alpha = 0 \quad \text{(2.42)} \]

\[ -K_{\varphi}^{(\beta)} \phi_\beta = 0 \quad \text{for } K_{\varphi} \in \Omega_\varphi / \Omega_u \]

Substituting (2.41) into (2.39) leads to elementary systems in (2.42) where \( u_\alpha, \phi_\alpha \) and \( \phi_\beta \) represent, respectively nodal values in the element \( \phi_\alpha \) (associated with piezoelectric materials) and element \( K_\varphi \) (associated with the air domain). Without losing generality, assume linear tetrahedral elements are used for discretization (see e.g. [41] for discussion on other types of elements). The mass matrix \( M_u^{(\alpha)} \), stiffness matrices \( K_u^{(\alpha)} \) and \( K_{\varphi}^{(\alpha)} \), coupling matrix \( K_{u\varphi}^{(\alpha)} \) and load vector \( F_u^{(\alpha)} \) are calculated as in (2.43). Superscripts \( \alpha \) and \( \beta \) correspond, respectively, to \( K_\alpha \) and \( K_\beta \) whereas \( t \) denotes tensor transpose. \( B_u^{(\alpha)} \) and \( B_{\varphi}^{(\alpha)} \) are defined as in (2.44).

\[ M_u^{(\alpha)} = \int_{K_\alpha} W_n^{n_{u\alpha}} \rho W_n^{n_{u\alpha}} d\Omega, \quad K_u^{(\alpha)} = \int_{K_\alpha} B_u^{(\alpha)^T} E B_u^{(\alpha)} d\Omega, \]

\[ K_{u\varphi}^{(\alpha)} = \int_{K_\alpha} B_u^{(\alpha)^T} E B_{\varphi}^{(\alpha)} d\Omega, \quad K_{\varphi}^{(\alpha/\beta)} = \int_{K_{\alpha/\beta}} B_{\varphi}^{(\alpha/\beta)^T} E B_{\varphi}^{(\alpha/\beta)} d\Omega, \quad \text{(2.43)} \]

\[ F_u^{(\alpha)} = \int_{K_\alpha} W_n^{n_{u\alpha}} \rho f d\Omega + \int_{\Gamma_{\alpha/\beta} \partial K_\alpha} W_n^{n_{u\alpha}} \rho \bar{t} d\Gamma \]

\[ B_u^{(\alpha)} := [B_u^{(\alpha)} B_u^{(\alpha)} B_u^{(\alpha)} B_u^{(\alpha)}] \]

\[ B_{\varphi}^{(\alpha)} := \begin{bmatrix} \frac{\partial \omega_n^{n_{u\alpha}}}{\partial x} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial x} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial x} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial x} \\ \frac{\partial \omega_n^{n_{u\alpha}}}{\partial y} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial y} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial y} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial y} \\ \frac{\partial \omega_n^{n_{u\alpha}}}{\partial z} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial z} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial z} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial z} \end{bmatrix} \quad \text{(2.44)} \]

\[ B_u^{(\alpha)} := \begin{bmatrix} \frac{\partial \omega_n^{n_{u\alpha}}}{\partial x} & 0 & 0 & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial x} \\ 0 & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial y} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial y} & 0 \\ 0 & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial z} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial z} & \frac{\partial \omega_n^{n_{u\alpha}}}{\partial z} \end{bmatrix} \]
To model structural damping, Rayleigh damping can be considered. It introduces the damping matrix as a fraction of the mass and stiffness matrices using $\alpha_{dM}$ and $\beta_{dK}$, as in (2.45). Here, $\xi$ denotes the damping factor at resonant frequency $\omega_0$. It can be measured at the first two resonant frequencies with which $\alpha_{dM}$ and $\beta_{dK}$ are calculated and used for the whole frequency range, approximately [42].

$$C_u^{(\alpha)} := \alpha_{dM}M_u^{(\alpha)} + \beta_{dK}K_u^{(\alpha)}, \quad \xi = \frac{1}{2}\left(\frac{\alpha_{dM}}{\omega_0} + \beta_{dK}\omega_0\right) \tag{2.45}$$

Adding Equation (2.45) into the first equation of (2.42) yields the elementary system

$$M_u^{(\alpha)}\ddot{U}_\alpha + C_u^{(\alpha)}\dot{U}_\alpha + K_u^{(\alpha)}U_\alpha + K_{u\phi}\phi_\alpha = F_u^{(\alpha)} \tag{2.46}$$

It is noted that damping also prevents singularity in the frequency regime when the problem is solved near frequencies of resonance. Before assembling elementary matrices into the global system, modeling electrodes and electrical loads are discussed.

2.3.2.2 Modeling the electrodes and electrical loads

Denote the surface of the $k$-th electrode as $S^k_e$. The equipotential condition means $\varphi(X) = \varphi_{\text{const}}$ for $X \in S^k_e$. When electric potentials are prescribed, this can be realized via Dirichlet boundary conditions. Otherwise, extra treatments are required. Suppose there are $n^k_e$ vertices on $S^k_e$. Then, the number of electrical DoFs is reduced to one for all the $n^k_e$ nodes. Without losing generality, assume all $n^k_e$ electrical DoFs are assigned with the numbering $k$. If there are $m^k_e$ electrodes, the total number of electrical DoFs not on electrodes is $n_\varphi - \sum_{k=1}^{m^k_e} n^k_e$. These DoFs can be numbered from $m_e + 1$ to $m_e + n_\varphi - \sum_{k=1}^{m^k_e} n^k_e - n_\phi$, with $n_\phi$ denoting the number of nodes on Dirichlet boundaries $\Gamma_{\varphi_D}$. For elastic the field, the $j$-th component of DoFs associated with the $l$-th vertex is numbered as $3 \times l + j - 1$ ($j = 1,2,3$). Hence, the total number of mechanical and electrical DoFs (denoted by $n^u_X$ and $n^\varphi_X$) are $3n_u - n_\varphi$ and $m_e + n_\varphi - n_\phi - \sum_{k=1}^{m^k_e} n^k_e$ ($n_\varphi$ number of nodes on $\Gamma_{\varphi_D}$), respectively.

$$M_u\ddot{U} + C_u\dot{U} + K_uU + K_{u\phi}\Phi = F_u \tag{2.47}$$

$$K_{u\phi}U - K_{\phi}\Phi = 0$$
After assembling, global systems can be obtained as in (2.47) where \( M_u, C_u, K_u \in \mathbb{R}^{n_2 \times n_2} \), \( K_u \varphi \in \mathbb{R}^{n_2 \times n_2}, K_\varphi \in \mathbb{R}^{n_2 \times n_2}, \) and \( F_X \in \mathbb{R}^{n_2} \). Equations in (2.47) can be transformed from the time regime to the frequency regime. For a time-harmonic excitation at angular frequency \( \omega \), the transformed system reads

\[
(-\omega^2 M_u + j \omega C_u + K_u)U + K_{u \varphi} \Phi = F_u
\]

\[
K_{u \varphi}^t U - K_\varphi \Phi = 0
\]

(2.48)

Incorporating electric circuits can be achieved by using equivalent admittance matrix [43] and modifying the dielectric matrix in the second equation of (2.48). To be more specific, \( K_\varphi \) is replaced with \( K_\varphi + \tilde{K}_\varphi \) in which the effective capacitance matrix \( \tilde{K}_\varphi \) is initially null. For each circuit element \( \tilde{x} \) connected between the \( p \)-th and \( q \)-th electrode, define a vector \( V_{\tilde{x}} \) that is of size \( n_\varphi \times 1 \) with the \( p \)-th and \( q \)-th entry being 1 or -1, and others being zero. For conciseness, We denote the product \( V_{\tilde{x}} V_{\tilde{x}}^T \) by \( K_{\varphi \tilde{x}} \). Eventually, \(-j/\omega \tilde{Z}_{\tilde{x}} \tilde{K}_{\varphi \tilde{x}} \) is added to \( \tilde{K}_\varphi \):

\[
\tilde{K}_\varphi = \sum_{\tilde{x}} \frac{j}{\omega \tilde{Z}_{\tilde{x}}} K_{\varphi \tilde{x}}.
\]

(2.49)

where \( \tilde{Z}_{\tilde{x}} = -Z_{\tilde{x}} \) when \( \tilde{x} \) denotes a capacitor, \( \tilde{Z}_{\tilde{x}} = Z_{\tilde{x}} \) when \( \tilde{x} \) denotes a resistor or inductor. \( Z_{\tilde{x}} \) is the impedance of \( \tilde{x} \). The new system can be expressed as in Equation (2.50).

\[
(-\omega^2 M_u + j \omega C_u + K_u)U + K_{u \varphi} \Phi = F_u
\]

\[
K_{u \varphi}^t U - (K_\varphi + \tilde{K}_\varphi) \Phi = 0
\]

(2.50)

In this way, a FE system that considers both electrodes and electrical loads is built. Note that while the final FE formulations are expressed in frequency domain, they can be used for transient analysis (after transferring back to time domain).

2.3.3  Numerical examples

In this part, we present simulations based on the developed model. The first one consists in modal analysis and parametric study of a cantilevered piezoelectric bimorph. The second consists in assessment of thermos-elastic attenuation in SAW device [44].
2.3.3.1 Cantilevered piezoelectric bimorph

As shown in Figure 2.3, the bimorph is fixed onto a base. In practice, acceleration is applied on the base. For modeling, it is more convenient to consider displacements w.r.t. the base, which means the system is fixed on the base and subject to an acceleration $a(f)$.

![Figure 2.3. The cantilevered bimorph. (a) model configuration, (b) triangulation](image)

Dimensions of the bimorph are set, so that piezoelectric layers are of $27 \times 3.2 \times 0.258 \, mm^3$ whereas the substrate in sandwiched by piezoelectric layers is of $27 \times 3.2 \times 0.115 \, mm^3$. Meanwhile, we suppose that electrodes are connected in series (though little work is required for the case of a parallel connection) [45]. Piezoelectric layers are made of PZT-5H, while the substrate is made of brass. Material constants are summarized in Table 2.2.

To perform modal analysis, (2.50) is re-formulated as

\[
(-\omega^2 M + j\omega C + K)X = F
\]

(2.51)


The problem amounts to setting $F$ to zero and solving the Eigen problem for resonant frequencies $\omega$ and vibration mode shapes $X$. It is noted that instead of solving a purely structural
### 2.3. Piezoelectric model

Table 2.2. Constants for piezoelectric and substrate layers

<table>
<thead>
<tr>
<th>Substrate (brass)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Mass density (kg · m⁻³)</td>
<td>9000</td>
</tr>
<tr>
<td>Young’s Modulus (N · m⁻²)</td>
<td>105 × 10⁹</td>
</tr>
<tr>
<td>Poisson’s ratio</td>
<td>0.23</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Piezoelectric (PZT-5H)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Elastic constants (× 10¹⁰ N · m⁻²)</td>
<td></td>
</tr>
<tr>
<td>$c_{11}$ = 12.72</td>
<td>$c_{12}$ = 8.02</td>
</tr>
<tr>
<td>$c_{13}$ = 8.47</td>
<td>$c_{33}$ = 11.74</td>
</tr>
<tr>
<td>$c_{44}$ = 2.30</td>
<td></td>
</tr>
<tr>
<td>Piezoelectric constants (C · m⁻²)</td>
<td></td>
</tr>
<tr>
<td>$e_{31}$ = −6.62</td>
<td>$e_{33}$ = 23.24</td>
</tr>
<tr>
<td>$e_{31}$ = −6.62</td>
<td></td>
</tr>
<tr>
<td>Relative permittivity</td>
<td></td>
</tr>
<tr>
<td>$\varepsilon_{r11}$ = 1704.40</td>
<td>$\varepsilon_{r33}$ = 1433.61</td>
</tr>
</tbody>
</table>

problem, electrodes and the electric loads can be taken into consideration, using our approach. In this example, electrodes are arranged so that the output voltage is read between the upper and lower surface of the bimorph while the substrate (which is conductor), is grounded. Loads are taken to be infinite i.e. open circuit. However, other circumstances such as short circuit and finite impedances can also be conveniently simulated.

In Figure 2.4, resonant frequencies and mode shapes of the first six modes are depicted. Real parts and imaginary parts are represented separately. As can be seen, presence of the damping matrix $\mathbf{C}$, which is related to mechanical losses in reality, is reflected in the complex valued frequencies where the imaginary part implies attenuation. In the meanwhile, results of the

---

1 This is the so-called quadratic Eigen problem (QEP) as apart from $\mathbf{M}$ and $\mathbf{K}$ the damping matrix $\mathbf{C}$ is also involved. Without $\mathbf{C}$ the problem is called the general Eigen problem (GEP), i.e. solving $(-\omega^2 \mathbf{M} + \mathbf{K}) \mathbf{X} = 0$. The QEP can be transformed into the GEP by re-writing Equation (2.51) in the generalized Schur form

$$
\begin{pmatrix}
-\omega & \mathbf{M} \\
\mathbf{0} & \mathbf{I}_n
\end{pmatrix} +
\begin{pmatrix}
\mathbf{C} \\
\mathbf{0}
\end{pmatrix}
\begin{pmatrix}
\mathbf{K} \\
\mathbf{0}
\end{pmatrix}
\begin{pmatrix}
\omega \mathbf{X} \\
\mathbf{X}
\end{pmatrix} = 0
$$

from which $\omega$ and $\mathbf{X}$ are solved for. In the Schur form $\mathbf{I}_n$ represents the $n_{th}$ order identity matrix.
mode shapes suggest that the first two modes are suited for low frequency (e.g. < 3k Hz) applications. Specifically, when the system vibrates at the first resonant frequency, electrical output reaches the maximum output voltage of \(~4V\), which is ideal for the power supply of sensor nodes in WSN.  

![Image of vibration mode shapes](image)

Figure 2.4. Real and imaginary parts of the first six vibration mode shapes. Displacements are represented by (scaled) deviations from original positions whereas potentials by colored regions (in V).

To investigate performance of the device under various excitations and load conditions, excitation frequency and electrical load are taken as parameters. A set of six resistors (100Ω, 1 kΩ, 10 kΩ, 91 kΩ, 500 kΩ and 910 kΩ) are chosen. Simulations are carried out between 0 to 3k Hz. In Figure 2.5, output voltages and power on different loads are shown. Unsurprising

Express the vibration mode shapes in exponential form: 

\[ X = X_\theta e^{j(\omega_r + j\omega_i)t} = X_\theta e^{j\omega_t} \cdot e^{-\omega_i t} \]

in which \(\omega_r\) and \(\omega_i\) represents the real and imaginary part of the angular frequency \(\omega\), respectively. It is evident that the term \(e^{-\omega_i t}\) implies attenuation of the magnitude of \(X\) over time.
Figure 2.5. Output voltage (left) and power (right) of the bimorph versus frequency for different loads. Interestingly, peak values occur near the resonant frequencies that are found in the modal analysis. In addition, it is evident that among the listed loads that of 910 kΩ delivers the optimal output. Given the simple fact that efficiency is maximized when inner and load impedances of the system are balanced, an alternative way to estimate the optimal load is to calculate the inner impedance (typically capacitance) of the piezoelectric system. See [46] for detailed explanation and examples. The above simulation results compare favorably with experimental measurements reported in [42].

2.3.3.2 Surface acoustic wave device

Figure 2.6. A delay-line structured surface acoustic wave transducer. (a) top view and (b) lateral view

A typical configuration of SAW devices is depicted in Figure 2.6. It consists of a piezoelectric substrate and two pairs of interdigital transducers (IDTs) - one for transmitting and the other for receiving electrical signals. Films with sensing abilities can be deposited on the part between IDTs, which is referred to as the delay-line area. For SAW devices, most of the energy...
(or, in other words, acoustic wave propagation) is concentrated near the surface. Consequently, even minor variations occurred in the films has significant impacts on wave propagations in the substrate, which in turn, can be quantified from outputs on receiving IDTs. Their outstanding sensitivity makes them ideal for sensing tasks in WSN.

At the same time, repeated compressive and tensile stresses in regions within the piezoelectric substrate lead to cyclic heat flow, which dissipates energy. This phenomenon is called thermos-elastic damping which should be accurately quantified when designing SAW devices. For such assessments, numerical models are more suited as more and more irregular structures are involved in modern structures, for which existing analytical models are no longer applicable [44]. To build numerical models, we note that under our multiphysics framework, thermal fields and corresponding couplings can be easily added into the piezoelectric model.

\[
\begin{align*}
T_{ij} &= c_{ijkl}S_{kl} - e_{ijkl}^pE_k - \alpha_{ij}\theta \\
D_i &= e_{ijkl}^pS_{kl} + \varepsilon_{ij}^E E_k + \gamma_i\theta \\
\rho\eta &= \alpha_{kl}S_{kl} + \gamma_k E_k + c_p\theta
\end{align*}
\]

(2.52)

In this case, the coupled thermos-piezoelectric problem is governed by Equation (2.52), in which \(\alpha_{ij}\) and \(\gamma_i\) are thermal stress coefficients and pyroelectric coefficients, respectively. Recall that \(T\) and \(S\) indicates stress and strain, \(E\) and \(D\) indicates electric field and electric displacement, \(\rho, \eta, c_p,\) and \(\theta\) indicates mass density, entropy density, specific heat per unit volume at constant deformation, and temperature, \(c_{ijkl}^E, e_{ijkl}^p,\) and \(\varepsilon_{ij}^E\) indicate values of elastic, piezoelectric coupling, and permittivity, respectively.

Analogous to the piezoelectric modeling, the strong form of the problem can be obtained by combining constitutive equations (2.52) and equations of balances. Next, the discrete system can be obtained using the Galerkin method and interpolating quantities with appropriate FEs. Note that here the system is expressed in time domain, for which the Newmark method can be utilized. In this example, we solve the coupled system in the same way as previously. Namely, mechanical, electrical, and thermal DoFs are calculated simultaneously at each time step. It is noted that perfectly matched (PML) layers are added to avoid wave reflection, as shown in Figure 2.6 (b). Also, due to characteristics of SAW, components in the \(y\) direction can be decoupled, thus enabling analysis in 2D [44].
A 2D model is built for the thermos-piezoelectric problem. Simulations are carried out for both coated and uncoated conditions (that corresponds to the case with or without the presence of the film in Figure 2.6 (b), respectively). Regarding the model set-up, we use a $Y$-cut LiNbO$_3$ (see [44] for constant values of the material) substrate of $30\lambda \times 10\lambda$ (where $\lambda$ is the wavelength at a central frequency $f_0$ of 300 MHz). A total of 24 electrodes are modeled in IDTs. Meanwhile, relatively fine meshes (with maximum element length of $1/32\lambda$) and small time steps (of $0.025/f_0$) are chosen so as to ensure accuracy of the results.

In Figure 2.7 displacements in the $z$ direction at two points $A$ and $B$ (as marked in Figure 2.6) are depicted. Point $A$ is closer to transmitting IDTs, sitting four wavelength away from point $B$. Displacements at both points oscillate at constant amplitudes eventually. However, due to thermos-elastic damping slight descending in the amplitude at $B$, w.r.t. that at $A$ can be observed. Both simulated and reference results [47] in terms of wave attenuation are presented in Figure 2.8. As the figure implies, excellent correlations are obtained.
2.4 Magnetostrictive model

Magnetostrictive mechanisms and existing models of magnetostrictive materials have been presented in the Chapter 1. Here, we present a 3D magnetostrictive FE model, in which non-linear magnetostrictive behaviors are considered. As in the piezoelectric case, constitutive equations are first introduced. Next, we introduce the DEAM approach, and then FE formulations. Eventually, representative numerical examples are provided.

2.4.1 Magnetostrictive constitutive equations

The Gibbs energy $G$ of a magnetostrictive continuum [48] can be expressed as in (2.53). Recall that summation convention is employed in this section, which is recommended in the IEEE Standard on Magnetostrictive Materials [49].

$$G_b := U - T_{ij}S_{ij} - H_i B_i$$  \hspace{1cm} (2.53)

where $U$ represents the stored energy density; $i, j = 1,2,3$. Ignoring hysteresis, the material shows reversible properties. As a consequence, time derivative of $U$ can be expressed as sum of magnetic and elastic parts, see (2.54).

$$\dot{U} = T_{ij}\dot{S}_{ij} - H_i \dot{B}_i$$  \hspace{1cm} (2.54)

Taking derivative on both sides of Equation (2.53) and substituting (2.54) in yields

$$\dot{G}_b = -\dot{T}_{ij}S_{ij} - \dot{H}_i B_i$$  \hspace{1cm} (2.55)

From Equation (2.55) the following relations can be obtained

$$S_{ij} = -\left. \frac{\partial G}{\partial T_{ij}} \right|_H$$, \hspace{1cm} $$B_i = -\left. \frac{\partial G}{\partial H_i} \right|_T$$ \hspace{1cm} (2.56)

based on which, magnetostrictive constants $d^m$ can be defined as in Equation (2.57).
2.4. Magnetostrictive model

\[
d_{ij}^m = \frac{\partial S_{ij}}{\partial H_k} \bigg|_T = \frac{\partial B_i}{\partial T_{ij}} \bigg|_H
\]

(2.57)

As the equation suggests, \(d_{ij}^m\) denotes derivatives of strain (respectively, magnetic flux density) w.r.t. magnetic field (respectively, stress), under constant stress (respectively, magnetic field). Given the relations between \((S, B)\) and \((T, H)\), magnetostrictive constitutive equations can be obtained as in (2.58) where superscripts \(H\) in \(s_{ijkl}^H\) and \(T\) in \(\mu_{ij}^T\) indicates values of compliance and permeability, under constant magnetic field and stress, respectively.

\[
\begin{align*}
\delta S_{ij} &= s_{ijkl}^H \delta T_{kl} + d_{ijkl}^m \delta H_k \\
\delta B_i &= d_{ik}^m j \delta T_{kl} + \mu_{ij}^T \delta H_k
\end{align*}
\]

(2.58)

For FE modeling, it is more convenient to express \((T, H)\) in terms of \((S, B)\), as expressed in (2.59) where magnetostrictive constants \(h(N\cdot Wb^{-1} \text{ or } A\cdot m^{-1})\) instead of \(d^m(Wb \cdot N^{-1}, \text{ or } m\cdot A^{-1})\) are used; \(v^S(m \cdot H^{-1} \text{ or } 4\pi \cdot 10^{-7} \text{ Oe} \cdot G^{-1})\) indicates reluctivity under constant strain. In literature, \(B\) may be expressed in either \(G\) or \(T\) \((1T = 10^4 G)\), while \(H\) may be expressed in either \(A \cdot m^{-1}\) or \(\text{Oe}\) \((1A \cdot m^{-1} = 4\pi \cdot 10^{-3} \text{ Oe})\). Other forms of constitutive equations, as well as conversions between different sets of constants, are addressed in [49].

\[
\begin{align*}
\delta T_{ij} &= c_{ijkl}^B \delta S_{kl} - h_{ij} \delta B_k \\
\delta H_i &= -h_{ik} j \delta S_{kl} + v_{ij}^S \delta B_k
\end{align*}
\]

(2.59)

In (2.59), \(\delta\) implies quantities of perturbation. When deviations in equations in (2.59) are small enough, the linear form can be obtained, see (2.60).

\[
\begin{align*}
T_{ij} &= c_{ijkl}^B S_{kl} - h_{ij} B_k \\
H_i &= -h_{ik} j S_{kl} + v_{ij}^S B_k
\end{align*}
\]

(2.60)

2.4.2 The discrete energy averaged model

DEAM was first introduced for modeling Galfenol, and later adopted for modeling Terfenol-
D. Exceptional correlations between DEAM based simulation results and experimental measurements are reported, see [50], [17], [18], and [19]. We implemented it on a C++ platform. The program is compiled as a \textit{mex} file that can be called from MATLAB programs directly (for integration with FE programs, as the latter are implemented on MATLAB).

For DEAM, it is assumed that magnetostrictive materials are composed by non-interacting, single-domain particles whose magnetocrystalline anisotropy gives domains easy directions (denoted by \(c^k\)). When the material undergoes variation of magnetic fields \(H\) and stresses \(T\), these single-domains tend to rotate, and align themselves along the orientation \(m^k\). The bulk magnetization \(M\) and magnetostriction \(S_m\) are obtained as the weighted sum of the magnetization \(M_s m^k\) (with \(M_s\) representing the saturation magnetization) and magnetostriction \(S_m^k\) from each orientation, as expressed in Equation (2.61). Note that strain \(S\) is assumed to be decomposed into magnetostriction \(S_m\) and a purely mechanical part \(S_e\) (determined by stresses).

\[
M = \sum_{k=1}^{r} \xi_{hys}^k M_s m^k, \quad S_m = \sum_{k=1}^{r} \xi_{hys}^k S_m^k
\]  

(2.61)

where \(\xi_{hys}^k\) is the averaged hysteretic volume fractions of the particle, calculated as energy-weighted averages, see Equation (2.62).

\[
\xi_{hys}^k := \exp\left(-\frac{G_b^k}{\Omega}\right) \cdot \sum_{k=1}^{r} \exp\left(\frac{G_b^k}{\Omega}\right)
\]  

(2.62)

In(2.62) \(\Omega\) is the Armstrong smoothing factor, whereas \(G_b^k\) is the Gibbs free energy - the sum of anisotropy, coupling and the Zeeman energy, corresponding to the first, second and third term on the r.h.s. of (2.63), respectively. \(K^k\) is the anisotropy coefficient.

\[
G_b^k := \frac{1}{2} K^k \left| m^k - c^k \right|^2 - S_m^k \cdot T - \mu_0 M_s m^k \cdot H
\]  

(2.63)

Denote entries of \(m^k\) as \(m^k_i\) with \(i = 1,2,3\). For particles with cubic symmetry, magnetostriction \(S_m^k\) in \(m^k\) direction can be expressed as in Equation (2.64) where \(c_{11}\) and \(c_{12}\) are elastic moduli, while \(\lambda_{100}\) and \(\lambda_{111}\) are magnetostriction constants.
To calculate magnetic orientation \( \mathbf{m}^k \), the Gibbs energy is simplified as in (2.65).

\[
\tilde{G}_b^k := 1/2 \cdot \mathbf{m}^k \mathbf{K} \mathbf{m}^k - \mathbf{b}^{kT} \mathbf{m}^k
\]

where \( \tilde{G}_b^k \) represents some total inner energy while \( \mathbf{K} \) and \( \mathbf{b}^k \) are defined as

\[
\mathbf{K} := -3 \begin{bmatrix}
\lambda_{3100} T_1 & \lambda_{111} T_4 & \lambda_{1111} T_6 \\
\lambda_{1111} T_4 & \lambda_{100} T_2 & \lambda_{1111} T_5 \\
\lambda_{1111} T_6 & \lambda_{1111} T_5 & \lambda_{100} T_3
\end{bmatrix}, \quad \mathbf{b}^k := K^k \mathbf{c}^k + \mu_0 M_s H
\]

Next, it is easily verified that looking for a \( \mathbf{m}^k \) minimizing the l.h.s. of Equation (2.65) is equal to solving the eigenvalue problem (2.67), under the constraint that \( \| \mathbf{m}^k \|_2 = 1 \). In (2.67) \( \mathbf{I} \) is the sixth order identity matrix, whereas \( \gamma \) are unknown eigenvalues.

\[
(K - \gamma \mathbf{I}) \mathbf{m}^k = \mathbf{b}^k
\]

Finally, the problem amounts to solving a six order polynomial equation, which has six solutions for \( \mathbf{m}^k \). Among them, those complex valued or not in the vicinity of \( \mathbf{c}^k \) are discarded [50]. Once the magnetic orientations are obtained, they are substituted into equations (2.61) to (2.64). The bulk magnetization and magnetostriction are hence calculated. Meanwhile, material constants \( \mu^T, \mathbf{d}^m \) and \( \mathbf{s}^H \) in the constitutive equation (2.58) can be deduced, according to equations (2.56) and (2.57). \( \mathbf{v}^s, \mathbf{h} \) and \( \mathbf{c}^b \) in (2.59) can be obtained from \( \mu^T, \mathbf{d}^m \) and \( \mathbf{s}^H \) [49], which is needed to form the FE system.

In Figure 2.9 we depict variations of magnetization and magnetostriction under different magnetic and mechanical bias conditions. These results are obtained using our implementation, and show excellent accordance with the measurements presented in [50]. The corresponding physical scenario is that a magnetostrictive rod fixed on one end and subject to different bias conditions. Quantities in the rod are taken to be homogeneous, for which DEAM
Chapter 2. Multiphysics Finite Element Modeling

needs to be run only once for a given combination of state variables (namely, the aforementioned macroscopic and microscopic volumes refer to the same thing - the rod itself; the case of non-homogeneity i.e. the number of microscopic volumes is larger than the macroscopic one is discussed in § 2.4.4.3).

![Figure 2.9. Simulation results of nonhysteretic magnetization (right) and magnetostriction (left) for (100)Fe₈₁.₅Ga₁₈.₅ at various stress (top) and field (bottom) values](image)

2.4.3 Finite element formulations

To formulate the magnetostrictive problem, consider a domain $\Omega \in \mathbb{R}^3$ that is composed by subdomains $\Omega_D$, $\Omega_S$, and $\Omega_C$, representing the air, support of source currents and magnetostrictive materials, respectively (see Figure 2.10). Note that here we consider the effect of non-vanishing electric conductivity of magnetostrictive materials, hence $\Omega_C$ can also be interpreted as the domain of conductors. Also, we ignore eddy currents in the support of source currents. Therefore, $\Omega_S$ can be viewed as part of $\Omega_D$ except that source currents, if there are
any, are nonzero in \( \Omega_S \) while vanishing in other parts of \( \Omega_D \). For textual simplification, \( \Omega_S \) will only be mentioned for specification of source currents.

In what below low frequencies operating regimes are assumed, and thus the electric displacement term in the Maxwell-Ampère’s theorem is neglected. As a result, magnetic part of the coupled problem (that is the magneto-quasistatic problem) can be represented by (2.68), in which \( J_s \) is the imposed source current; \( \sigma_c E \) is the eddy current induced. \( \sigma_c \) is nonzero in \( \Omega_C \) and vanishing in others.

\[
\text{curl} H = \sigma_c E + J_s \\
\partial B/\partial t + \text{curl} E = 0
\]

(2.68)

We also assume \( \Omega_D \) is sufficiently large so that null boundary conditions on \( \partial \Omega_D \) are valid. Next, the magnetic vector potential \( A \) and a scalar potential \( \phi \) (which is the primitive of the aforementioned electric potential \( \varphi \)), are introduced so that

\[
B = \text{curl} A \\
E = - \partial (A + \text{grad} \phi)/\partial t
\]

(2.69)

where the negative sign in the second equation is used for symmetry of the discrete system. Substituting Equation (2.69) into (2.68) yields (2.70).

\[
\text{curl}(\nu \cdot \text{curl} A) + \partial (\sigma_c A + \sigma_c \text{grad} \phi)/\partial t = J_s \\
\text{div}(J_s - \sigma_c \partial (A + \text{grad} \phi)/\partial t) = 0
\]

(2.70)

where the second equation comes from the divergence-free property of the current density. In fact, it can be obtained by taking divergence of both sides of the first equations.

\[
\int_{\Omega} \text{curl} \mathcal{T}' \text{curl} \mathcal{T} d\Omega = \int_{\Omega} \text{curl} \mathcal{T}' J_s d\Omega
\]

(2.71)

To explicitly enforce the divergence-free property of \( J_s \), Equation (2.71) is first resolved.
Therefore, $J_s$ is expressed as $\text{curl}\mathcal{T}$ in which $\mathcal{T}$ is discretized using edge elements $\omega^e$, having boundary condition $\mathcal{T} \times n = 0$ on $\partial \Omega$. Here $n$ denotes the outward unit normal vector.

For magnetostrictive materials, a term corresponding to the magneto-elastic coupling needs to be added into (2.70). That being said, the coupled magnetic problem can be stated as: find $A \in \mathcal{H}(\text{curl}, \Omega)$ and $\phi \in \mathcal{H}(\text{grad}, \Omega)$ such that equations in (2.72) holds true for all $A' \in \mathcal{H}_0(\text{curl}, \Omega)$ and $\phi' \in \mathcal{H}_0(\text{grad}, \Omega)$. Here, $\mathcal{H}_0$ indicates Hilbert spaces with boundary conditions $A = 0$ on $\partial \Omega$ and $\phi = 0$ on $\partial \Omega$.

\begin{equation}
\int_{\Omega_0} \text{curl} A' \cdot \nu \cdot \text{curl} A \, d\Omega - \int_{\Omega_0} \text{curl} A' \cdot \mathcal{T} \, d\Omega = 0 \quad \text{in } \Omega_0
\end{equation}

\begin{equation}
\int_{\Omega} \text{curl} A' \cdot \nu \cdot \text{curl} A \, d\Omega - \int_{\Omega} \text{curl} A' \cdot h \cdot \text{grad} u \, d\Omega + \int_{\Omega} \sigma_e A' \cdot \dot{A} \, d\Omega + \int_{\Omega} \sigma_e A' \cdot \text{grad} \phi \, d\Omega - \int_{\Omega} \text{curl} A' \cdot \mathcal{T} \, d\Omega = 0 \quad \text{in } \Omega
\end{equation}

In (2.72) $u$ represents the mechanical displacement. The standard form is expressed in (2.73).

\begin{equation}
B_{AA}(A', A)_{\Omega_D} - B_{AT}(A', \mathcal{T})_{\Omega_D} = 0
\end{equation}

\begin{equation}
B_{AA}(A', A)_{\Omega_C} - B_{AU}(A', u)_{\Omega_C} + C_{AA}(A', \dot{A})_{\Omega_C} + C_{A\phi}(A', \dot{\phi})_{\Omega_C} - B_{AT}(A', \mathcal{T})_{\Omega_C} = 0 \quad \text{(2.73)}
\end{equation}

\begin{equation}
C_{\phi\phi}(\phi', \dot{\phi})_{\Omega_C} + B_{\phi\phi}(\phi', \phi)_{\Omega_C} = 0
\end{equation}

In the meanwhile, the coupled mechanical problem can be represented as in (2.74), which is treated the same way as in the piezoelectric case of previous section,

\begin{equation}
\int_{\Omega} u' \cdot \rho \cdot \ddot{u} \, d\Omega + \int_{\Omega} \text{grad} u' \cdot \mathcal{B} \cdot \text{grad} u \, d\Omega - \int_{\Omega} \text{grad} u' \cdot h' \cdot \text{curl} A \, d\Omega + \int_{\Omega} u' \cdot f_v \, d\Omega - \int_{\Gamma_{uN}} u' \cdot \bar{t} \, d\Omega = 0
\end{equation}

(2.74)
where \( u_i' \in H_0(\text{grad}, \Omega) \). Associated boundary conditions are \( u_i = u_0 \) on \( \Gamma_{uD} \) and \( n_i T_{ij} = t_0 \) on \( \Gamma_{uW} \). Appropriate initial conditions are also needed. The standard form corresponding to Equation (2.74) reads

\[
M_{UU}(u', \ddot{u})_{\Omega_C} + K_{UU}(u', u)_{\Omega_C} - K_{UA}(u', A)_{\Omega_C} + (u', f_v)_{\Omega_C} - (u', t_0)_{\Gamma_{uN}} = 0 \tag{2.75}
\]

In the above equation, a structural damping term is needed when damping effects are considered. The discrete version of the above weak formulations is presented in (2.76).

\[
B_{AA}(A'_h, A_h)_{\Omega_D} - B_{AT}(A'_h, T_h)_{\Omega_D} = 0
\]

\[
B_{AA}(A'_h, A_h)_{\Omega_C} - B_{AU}(A'_h, u_h)_{\Omega_C} + C_{AA}(A'_h, \dot{A}_h)_{\Omega_C} + C_{A\phi}(A'_h, \dot{\phi}_h)_{\Omega_C} - B_{AT}(A'_h, T_h)_{\Omega_C} = 0
\]

\[
C_{\phi\phi}(\dot{\phi}_h', \dot{\phi}_h)_{\Omega_C} + B_{\phi\phi}(\dot{\phi}_h', \dot{\phi}_h)_{\Omega_C} = 0
\]

\[
M_{UU}(u_h, \ddot{u}_h)_{\Omega_C} + M_{UU}(u'_h, u_h)_{\Omega_C} - B_{UA}(u'_h, A_h)_{\Omega_C} + (u'_h, f_v, h)_{\Omega_C} - (u'_h, t_0)_{\Gamma_{uN}} = 0
\]

Variables related to \( u \) and \( \phi \) are interpolants of nodal elements \( \omega^0_i \), whereas those related to \( T \) and \( A \) are interpolants of edge elements \( \omega^1_i \). Body forces and surface tractions, if there are any, can be treated the same way as displacements. Substituting the interpolants into (2.76) and using the Galerkin method, we can have (2.77) for elements \( K_\alpha \in \Omega_C \) and \( K_\beta \in \Omega_D \).

\[
K^{(\beta)}_\alpha \ddot{\overline{A}_\beta} = F^{(\beta)}_\alpha
\]

\[
K^{(\alpha)}_\alpha \ddot{\overline{A}_\alpha} - K^{(\alpha)}_{au} U_\alpha + C^{(\alpha)}_a \ddot{\overline{A}_\alpha} + C^{(\alpha)}_{a\phi} \ddot{\overline{\phi}_\alpha} = F^{(\alpha)}_a
\]

\[
C^{(\alpha),t}_a \ddot{\overline{A}_\alpha} + K^{(\alpha)}_{\phi} \ddot{\overline{\phi}_\alpha} = 0
\]

\[
M^{(\alpha)}_u \ddot{U}_\alpha + K^{(\alpha)}_u U_\alpha - K^{(\alpha),t}_{au} \ddot{A}_\alpha = F^{(\alpha)}_u
\]

In (2.77), \( F^{(\alpha,\beta)}_a \) and \( F^{(\alpha)}_u \) are force vectors, whereas \( \ddot{\overline{A}_\alpha}, U_\alpha \), and \( \ddot{\overline{\phi}_\alpha} \) represent discrete values for respectively, \( A, u \), and \( \phi \). The bar in \( \ddot{\overline{A}_\alpha} \) is used to suggest that values are in fact line integrals. Recall that the exterior derivative of a \( p \)-form lies in a subspace of the space of \((p + 1)\)-form, see Equation (2.10). Applying this rule to e.g. \( \text{curl} A \) and \( \text{grad} \phi \) in Equation
(2.72), the following identities can be obtained

\[
\text{curl} \mathbf{A}^{(\alpha)} = \text{curl} \left( \sum_{i=1}^{n_e} \omega_{\alpha i} \mathbf{A}_{\alpha i} \right) = \mathbf{W}_\alpha \mathbf{C}^{(\alpha)} \mathbf{\bar{A}}_{\alpha}
\]

\[
\text{grad} \phi^{(\alpha)} = \text{grad} \left( \sum_{j=1}^{n_n} \omega_{\alpha j} \phi_{\alpha j} \right) = \mathbf{W}_{\alpha}^{\phi} \mathbf{G}^{(\alpha)} \psi_{\alpha}
\]

(2.78)

Recall that \(\omega_{\alpha i}^{n}, \omega_{\alpha i}^{e},\) and \(\omega_{\alpha n}^{f}\) are nodal, edge, and facet basis functions in \(\mathcal{K}_{\alpha}\) (which reads, respectively, \(\mathbf{W}_{\alpha}^{n}, \mathbf{W}_{\alpha}^{e},\) and \(\mathbf{W}_{\alpha}^{f}\), in compact form); \(n_{n}, n_{e},\) and \(n_{f}\) are numbers of corresponding basis functions; incident matrices associated with discrete gradient, curl, and divergence are denoted by \(\mathbf{G}, \mathbf{C},\) and \(\mathbf{D},\) respectively. Implementations of incident matrices are specified in § A.3. Meanwhile \(\mathbf{K}_{\alpha}^{(\alpha/\beta)}, \mathbf{r}_{\alpha}^{(\alpha/\beta)}, \mathbf{K}_{\alpha}^{(\phi)}, \mathbf{C}_{\alpha}^{(\alpha)}, \mathbf{C}_{\alpha}^{(\phi)}\) and \(\mathbf{K}_{\phi}^{(\phi)}\) in (2.77) are calculated as in (2.79). Those associated with the elastic problem are treated as in the piezoelectric case, see Equation (2.43).

\[
\mathbf{K}_{\alpha,\beta}^{(\alpha/\beta)} := \mathbf{C}^{(\alpha/\beta),t} \left( \int_{\mathcal{K}_{\alpha,\beta}} \mathbf{W}_{\alpha,\beta}^{\tau} \cdot \mathbf{v} \cdot \mathbf{W}_{\alpha,\beta}^{\tau} \, d\Omega \right) \mathbf{C}^{(\alpha/\beta)}
\]

\[
\mathbf{K}_{\alpha}^{(\phi)} := \mathbf{G}^{(\phi)} \left( \int_{\mathcal{K}_{\alpha}} \mathbf{W}_{\alpha}^{\phi} \sigma_{\phi} \mathbf{W}_{\alpha}^{\phi} \, d\Omega \right) \mathbf{G}^{(\phi)}
\]

\[
\mathbf{C}_{\alpha}^{(\alpha)} := \int_{\mathcal{K}_{\alpha}} \mathbf{W}_{\alpha}^{\epsilon} \sigma_{\epsilon} \mathbf{W}_{\alpha}^{\epsilon} \, d\Omega
\]

\[
\mathbf{C}_{\alpha}^{(\phi)} := \left( \int_{\mathcal{K}_{\alpha}} \mathbf{W}_{\alpha}^{\phi} \sigma_{\phi} \mathbf{W}_{\alpha}^{\phi} \, d\Omega \right) \mathbf{G}^{(\phi)}
\]

(2.79)

\[
\mathbf{r}_{\alpha}^{(\alpha/\beta)} := \mathbf{C}^{(\alpha/\beta),t} \left( \int_{\mathcal{K}_{\alpha}} \mathbf{W}_{\alpha,\beta}^{\tau} \cdot \mathbf{W}_{\alpha,\beta}^{\tau} \, d\Omega \right) \mathbf{f}_{\alpha/\beta}
\]

The elementary systems are assembled according to the numbering of elements. The global system is expressed in (2.80).
2.4. Magnetostrictive model

\[ \mathbf{K}_a^{(D)} \mathbf{A}_D = F_a^{(D)} \]

\[ \mathbf{K}_a^{(C)} \mathbf{A}_C - \mathbf{K}_{au}^{(C)} U_C + \mathbf{C}_{a\phi}^{(C)} \mathbf{A}_C + \mathbf{C}_{a\phi}^{(C)} \mathbf{A}_C^T \psi_C = F_a^{(C)} \]

\[ \mathbf{C}_{a\phi}^{(C)} \mathbf{A}_C^T + \mathbf{K}_{\phi}^{(C)} \psi_C = 0 \]

\[ \mathbf{M}_{u}^{(C)} \ddot{U}_C + \mathbf{C}_{u}^{(C)} \dot{U}_C + \mathbf{K}_{u}^{(C)} U_C - \mathbf{K}_{au}^{(C)} \mathbf{A}_C = F_u^{(C)} \]  

Note that in (2.80) superscripts C and D are used to represent, respectively, domains related to magnetostrictive materials and air, similar to those in (2.77) for elementary systems. Also, the damping matrix, generated using the Rayleigh damping, is added to the fourth equation.

2.4.4 Numerical examples

Now, we present three numerical examples that are carried out using our magnetostrictive model. For the first one, magnetic fields are calculated for given source currents, while for the others source currents are set to zero and magnetic fields are obtained for given boundary conditions. The well-known TEAM (Testing Electromagnetic Analysis Methods) Problem 7 is utilized in the first example; the other two examples are concerned with investigation on a cantilevered magnetostrictive unimorph. More precisely, the second example deals with a linear problem whereas the third with a nonlinear one in which we also introduce elaborating DEAM into the FE model.

2.4.4.1 The TEAM Problem 7 (asymmetrical conductor with a hole)

![Figure 2.11. Configuration of the TEAM Problem 7](image)

The problem (as shown in Figure 2.11) consist in calculating non-uniform magnetic fields (that are stimulated by a harmonically time varying source current in the coil), and eddy current in a conductor plate that is placed eccentrically under the coil.
It consists of three major steps. The first is assign values of source currents to all the facet elements within the coil. While for this example analytical expressions can be obtained because of the regular shape of the coil, we adopt a more general approach, with which source
currents are calculated based on the results of an electrostatic problem. The second step is to project the source current (as in Figure 2.12.a) in $\mathcal{H}(\text{curl})$ through solving Equation (2.71). As such, results of the previous step are divergence-free (see also [51] and [52]). The final step is to calculate potentials $A$ and $\phi$ by solving the first three equations (and neglecting the mechanical terms) of Equation (2.80). Here, simulations are carried out for frequencies of 50 and 200 Hz. We compare our results with the reference and find very good accordance. Magnetic fields (around the coil) and eddy currents are depicted in Figure 2.12.b and Figure 2.12.c, respectively.

2.4.4.2 Linear harmonic magnetostrictive analysis

For better performance, magnetostrictive devices normally operate under properly biased conditions. Mechanical bias refers to pre-stresses that can be applied as in Figure 1.5 (using spring washers and compressing bolds); magnetic bias refers to fields imposed by e.g. permanent magnets. When excitations to the system are of small magnitudes, material constants can be viewed invariant, which are obtained under bias conditions. Consequently, the underlying problem is linear. In this example, we consider a linear case; additionally, we also assume materials are homogeneous (namely, material constants identical over all locations).

Table 2.3. Constants for the magnetostrictive material

\begin{equation*}
\rho = 9.2 \times 10^3 \text{ kg} \cdot \text{m}^{-3}, \quad \sigma_c = 1 \times 10^7 \text{ S} \cdot \text{m}^{-1}
\end{equation*}

\begin{equation*}
\nu^5 = \begin{bmatrix} 1.85 & 0 & 0 \\ 0 & 1.85 & 0 \\ 0 & 0 & 1.85 \end{bmatrix} \times 10^5 \text{ m} \cdot \text{H}^{-1}
\end{equation*}

\begin{equation*}
h = \begin{bmatrix} 2.90 & -1.13 & -1.13 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 2.00 & 0 \\ 0 & 0 & 0 & 2.00 & 0 \end{bmatrix} \times 10^7 \text{ A} \cdot \text{m}^{-1}
\end{equation*}

\begin{equation*}
e'' = \begin{bmatrix} 3.11 & 1.52 & 1.52 & 0 & 0 & 0 \\ 1.52 & 3.56 & 1.52 & 0 & 0 & 0 \\ 1.52 & 1.52 & 3.56 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1.57 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1.36 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1.36 \end{bmatrix} \times 10^{10} \text{ Pa}
\end{equation*}

The model consists of a cantilevered magnetostrictive unimorph sitting inside a uniform magnetic field (of 795.77 A \cdot m^{-1}, or 1 mT) along its length direction. Configuration is adopted
from [53], in which dimensions are $5 \times 1 \times 0.5 \text{ mm}^3$, the excitation frequency is $1 \text{ kHz}$ (harmonic excitations assumed), material constants are presented in Table 2.2. In addition, we add a sufficiently large air domain around the unimorph, as depicted in Figure 2.13.b.

Figure 2.13. Results of the linear magnetostrictive problem (*coordinates along the length direction are scaled for representation purpose*). (a) edges on which nonzero Dirichlet boundary conditions for $A$ are applied; (b) solution of the problem in which arrows depict the magnetic flux distribution whereas the rainbow colored region depicts mechanical deformation; (c) distribution of the magnetic flux density.

For magnetic boundary condition, we see that integrating both side of the first equation in (2.69) leads to the identity between $\int_S B$ and $\sum_i \bar{A}_i$. The latter represents the sum of DoFs associated with $A$, over any sets of edges enclosing the section. A simple way to this end is to assign zero to all $\bar{A}_i$ on the boundary surface, except for those crossing a randomly built path (that go from one end of the domain to the other), formed by a layer of elements. These edges
2.4. Magnetostrictive model

are shown in Figure 2.13.a (in red and green, for which different colors imply different edge orientations (see A.3); accordingly, DoFs associated with these edges are assigned values of opposite signs). The Biconjugate gradient stabilized (BiCGSTAB) method is employed to solve the resultant system, with diagonal preconditioners.

Simulation results are depicted in Figure 2.13.b, in which yellow arrows represent the magnetic flux, whereas the multicolored regions represent displacements. As can be seen from the figures, magnetic fields near both ends of the unimorph are distorted (with more details in Figure 2.13.c), which is due to the fact that permeability of the magnetostrictive material is relatively larger than that of the free space. Moreover, we note that deformation occurs mainly along the direction of the field: maximum displacement along the length direction is around 45 nm, while along the width and height direction it is about 2.2 and 0.7 nm, respectively.

2.4.4.3 Nonlinear static magnetostrictive analysis

This example is based on the previous one but considering material non-homogeneity and nonlinearity. Non-homogeneity results from the fact that material constants at a given location are determined by state variables on that location, while the latter (i.e. stress and magnetic field levels) are normally different from location to location. Therefore, material constants also vary spatially. As for nonlinearity, it is due to the recursive dependency between material constants and state variables. For example, to obtain material constants we need state variables, to solve for which we need to have material constants.

Figure 2.14. Modeling hierarchy involving FEM on macroscopic, and DEAM on microscopic structure.

As we have mentioned previously, the underlying problem involves a hierarchy of structures: material constants are solved on the microscopic structure (using DEAM), while state variables are solved on the macroscopic structure (using FEM). Such microscopic structures are sometimes referred to as microscopic representative volumes (MRV) [54]. Given that our FE implementation is based on linear tetrahedral elements, the concerned state variables are element-wise constant. Thus, it is natural to choose FEs as MRVs for calculations with DEAM. The results (i.e. material constants) are used to form FE systems. The schema in Figure 2.14 summarizes integration of DEAM with FEM on the hierarchy.
To solve the nonlinear problem, Newton methods are usually utilized. With these methods, we can start with an initial guess of state variables, and calculate material constants that are then used to form a new FE system, based on which new solves of state variables can be obtained. Next, previous solves are updated based on the errors, using the Jacobian. This process is repeated until reaching convergence. In our case here, however, Newton methods are not suited because calculating the Jacobian will be too much involved and expensive (recall that DEAM involves a lot of operations, even for the calculation on a single MRV). For this reason, we adopt the piecewise linear approach [55], which seems to be more feasible. This approach is based on the thought that for a nonlinear relation over a range, when the latter is divided into many sufficiently small sub-ranges, the relation over each sub-range can be approximated using linear expressions. Moreover, at the initial state where pre-stress and magnetic fields are all zero, values of stress and fields in all elements vanish over all FEs. Note that here we only deal with stationary analysis, as we assume that the magnetostrictive device operates under static biased fields supper-positioned with small dynamic excitations, for which nonlinearity involves only when solving for the biased conditions (i.e. calculating material constants over all MRVs, or FEs), which is a stationary problem.

To describe the approach, we denote the number of pieces by $n$. Say, we need to solve for material constants over all elements for under a biased condition of magnetic field $H_0^p$ and pre-stress $T_0^p$. We take $n$ values of magnetic field and pre-stress in the range between $0$ and the biased condition, denoted by $H_0^{p1}, H_0^{p2}, \cdots, H_0^{pn}$ and $T_0^{p1}, T_0^{p2}, \cdots, T_0^{pn}$. From the previous linear example, we see that the magnetic bias condition is associated with boundary conditions in the FE system. For mechanical bias, or pre-stress, we need to add an additional term (that is related to $\sigma_{ex}$ in Equation (2.25)) to the r.h.s. of the FE system (2.80). For an element $\mathcal{K}_\alpha$, this term reads $|\mathcal{K}_\alpha| \cdot B_\alpha^{w,e} \cdot T_0^{pk}$ in which $|\mathcal{K}_\alpha|$ is the volume of element $\mathcal{K}_\alpha$. For each piece, we first solve for state variables over all elements (in which magnetostrictive materials are assigned), using FEM (at the $k$-th piece, $H_0^{pk}$ and $T_0^{pk}$ are used). After that, we extract variations of stress and magnetic field from FE solutions. These variations are added to corresponding initial values, based on which the current FE problem is solved. The results are then utilized as inputs for DEAM to calculate materials constants over each element, which are, in turn, used to form the FE system for the next stage. The procedure is repeated until the final piece, for which the obtained material constants can be stored and employed for subsequent analysis. The piecewise linear approach is employed for the nonlinear problem, based on the linear FE model as described in § 2.4.4.2. Material parameters used for DEAM are adopted from [50]. The bias field is of $50 \text{ kA} \cdot \text{m}^{-1}$ while the pre-stress is zero. Note that
a relatively large field is chosen so that saturation can be observed. A series of simulation consisting of 2000 and 3000 pieces are carried out; results in terms of the maximum displacement in the length direction divided by the length are depicted in Figure 2.15. With reference to the figure, we see that results tend to converge as the number of pieces increase. In addition, the saturation phenomenon can be observed when $H_{0}^{pk}$ passes certain level. We note that the maximum strain still increases but at a very small rate. This is due to the fact that the turn point before 'saturation' in fact corresponds to the moment where materials in most elements are saturated, while the rest goes saturated afterward (due to non-homogeneous distribution of stresses and fields), gradually. Note that during the solution for $H_{0}^{pn}$, results for biased conditions $H_{0}^{p1}, H_{0}^{p2}, \ldots, H_{0}^{p,n-1}$ are also obtained.

Figure 2.15. Convergence study of the piecewise linear strategy.

2.5 Chapter summary

In this chapter, we presented our multiphysics framework for modeling piezoelectric and magnetostrictive materials. Whitney elements are employed for discretization of variables of electromagnetic, elastic and thermal fields. We’ve also seen that electrodes and electrical loads can be conveniently elaborated into the framework, which is important for investigation of performances of piezoelectric energy harvesters. For magnetostrictive modeling, it is essential to integrate DEAM and FEM when material nonlinearity needs to be considered. The underlying nonlinear problem can be solved using the piecewise linear approach.
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Chapter 3. PGD Based Model Reduction of Multiphysics Systems

In this chapter, we first review some basics of separated representation based model reduction. In particular, the PGD, as well as POD method, are introduced. Then, two PGD based model reduction algorithms dedicated to respectively, nonlinear transient and multi-parametric problems that might arise in our multiphysics modeling work, are presented in the second and third section. It is concluded in the fourth section.

3.1 Separated representation based model reduction

Although computational resource has become increasingly available today, there are still cases where the solution of some large-size problems can take enormous amount of time, or simply intractable, even using the most advanced computing facilities. This phenomenon is usually termed as ‘the curse of dimensionality’, which may frequently arise in multiphysics modeling. For instance, for transient analysis of strongly coupled problems, time constant of one physics may be orders of magnitude different than the other, which may lead to too many time steps (thus very long runtime) when using conventional time integration algorithms. Parametric problems are another important category in this regard, since ‘multiple physics’ implies more factors or, more parameters that can affect the system. On the other hand, problem size grows exponentially with the number of parameters, which can quickly become too large to be solved for large 3D FE systems.

Fortunately, model reduction techniques have been developed to alleviate this issue [56]. Such techniques consist in projecting a large system of equations onto a much smaller subspace, thereby significantly reducing the problem size while preserving underlying properties of the original problem. While such projections can be accomplished in several means, in this thesis we focus on those based on separated representation. For illustration, let’s resume the abstract problem that has been used in § 2.1.2.2: find $u \in V$, such that

$$a(u, v) = l(v), \quad \text{for all } v \in V.$$  \hspace{1cm} (3.1)

The bilinear form $a(\cdot, \cdot)$ and linear form $l(\cdot)$ is defined as it was, whereas the solution $u$
depends on parameters $x_1, x_2, \ldots, x_d$ that are associated with e.g. spatial coordinates, material constants, time, etc. When the problem in (3.1) is solved using FEM, for example, the numerical solution to a given combination of parameters reads

$$U(x_1, x_2, \ldots, x_d) = \sum_{i_1=1}^{n_1} \cdots \sum_{i_d=1}^{n_d} a_{x_{i_1} \ldots x_{i_d}} \prod_{k=1}^{d} h_{i_k}(x_k). \quad (3.2)$$

In the equation $n_i$ represents the number of DoFs related to $x_i$, $h_{i_k}(x_k)$ represents some FE basis function. The total number of unknowns is therefore $n_1 \times n_2 \times \cdots \times n_d$.

Using a rank-$J$ separated representation, the function $u(x_1, \ldots, x_d)$ can be approximated with

$$u(x_1, x_2, \ldots, x_d) = \sum_{j=1}^{J} F_j^1(x_1) \times \cdots \times F_j^d(x_d). \quad (3.3)$$

Hopefully, the approximation approaches the true solution if $J \to \infty$. Discretizing the functions $F_j^1(x_1), \ldots, F_j^d(x_d)$ for $j = 1, \ldots, J$ in the equation, yields the numerical approximation

$$U(x_1, x_2, \ldots, x_d) = \sum_{j=1}^{J} \sum_{i_1=1}^{n_1} \cdots \sum_{i_d=1}^{n_d} \prod_{k=1}^{d} a_{j, i_k} h_{i_k}(x_k). \quad (3.4)$$

Now, the total number of unknowns becomes $J \times (n_1 + n_2 + \cdots + n_d)$ which can be much smaller than in the previous case. Therefore, separated representation can bring huge computational saving [57]. The representation in (3.4) can be constructed either a posteriori or a priori, leading to e.g. the POD and PGD approach, respectively. POD has been employed in various scientific computational applications over decades [58], whereas PGD is more of an ongoing research subject [57] dating back to 2006 [59]. Note that although other similar approaches exist, in this section we restrict our review to PGD, based on which our algorithms are developed, and POD, which is closely relevant to PGD and provides possibilities of extensions to our work. Complete surveys on model reduction including other approaches can be found in e.g. [56], [60], and references therein.
3.1. Separated representation based model reduction

3.1.1 The Proper Orthogonal Decomposition

POD is a method for constructing low-dimensional approximation representations of a subspace in Hilbert space [61]. POD are normally utilized for problems with two (types of) variables [57], e.g. space and time; in this case, \( d \) in (3.4) equals 2. Also, as we are dealing with numerical problems, we are more concerned with POD in finite dimensional settings.

The finite dimensional problem of (3.1) can be stated as: find \( u_h \in V_h \), such that

\[
a(u_h, v_h) = l(v_h), \quad \text{for all } v_h \in \tilde{V}_h.
\] (3.5)

In the equation variables with subscript \( h \) stand for the same objects as in § 2.1.2.2, \( u_h \) and \( v_h \) are function of \( x_1 \) and \( x_2 \). Applying POD to this problem consists of three steps. First, solve (3.5) for a list of selected samples of the parameter (say, \( x_2 \)). Solutions are called snapshots that are utilized to compute a reduced basis with Singular Value Decomposition (SVD) in the second step. The final step is to project the discrete system in (3.5) onto the obtained POD basis, using e.g. the Galerkin method [62]. After that, we solve a much smaller-size problem before recovering solutions to the original problem from those of the reduced one.

Obviously, the selection of snapshots is critical since the quality of approximation (i.e. reduced solutions) depends totally on it. There is an approach called Reduced Basis (RB) [63], which provides means for optimal selections. However, it is reported that the increased computational cost coming with RB can be high [64]. Here, we suppose selections are properly made, and snapshots are available. In addition, we denote the finite dimensions of \( x_1 \) and \( x_2 \) by \( n_1 \) and \( n_2 \), respectively, the number of snapshots by \( m_2 \), with \( m_2 \ll n_2 \). Let \( U_{m_2} = (U_1, ..., U_{m_2}) \) be ensemble of snapshots, in which \( U_i \in \mathbb{R}^{n_1} \), \( i = 1, ..., m_2 \). Then, the second step can be seen as extracting ordered orthonormal basis vectors \( \Phi = (\Phi_1, ..., \Phi_{m_2}) \) from \( U_{m_2} \) such that the following relation between mean square errors holds [58].

\[
E[\|U - U(r)\|^2] \leq E\left\{\|U - \bar{U}(r)\|^2\right\} \quad \text{(3.6)}
\]

In (3.6) \( U \) represents an arbitrary vector in \( V_h \), \( U(r) \) is an approximation of \( U \) using the leading \( r \) vectors in \( \Phi_r := (\Phi_1, ..., \Phi_r) \), whereas \( \bar{U}(r) \) is an approximation using arbitrary \( r \) basis vectors in \( V_h \). Next, it can be easily verified that performing SVD on \( U_{m_2} \) yields (3.7).
In the equation, \( M \) and \( N \) are \( n_1 \) by \( n_1 \) and \( m_2 \) by \( m_2 \) orthogonal matrices, respectively. \( \Sigma \) is an \( n_1 \) by \( m_2 \) matrix with zero entries everywhere except on the diagonal (whose entries \( \Sigma_{i,i} \) are denoted by \( \sigma_i \)). The latter are called singular values of \( U_{m_2} \) and are non-negative numbers arranged in decreasing order: \( \sigma_1 < \sigma_2 < \cdots < \sigma_{\hat{r}} \) where \( \hat{r} \) is the rank of \( U_{m_2} \).

\[
U_{m_2} = M \Sigma N^T
\]  

(3.7)

It turns out that \( M \) is, in fact, the basis \( \Phi \) we are looking for. Substituting \( M \), the minimum error in (3.6) can be expressed as \( \sum_{i=r+1}^{\hat{r}} \sigma_i^2 \), see [58]. Hence, \( U \) can be approximated with

\[
U = \sum_{i=1}^{r} u_i^T \Phi_i = \Phi_r^T U_r
\]

(3.8)

where \( U_r \in \mathbb{R}^r \). As the last operation at the second step, we pre-multiply both sides of the discrete system of (3.5) with the POD basis \( \Phi_r \) to form a reduced system, which is called *Galerkin projection* [62]. At the final step, we solve the reduced system for all samples of the variable \( x_2 \) to obtain reduced solutions of size \( r \) by \( n_2 \), which, after being pre-multiplied with \( \Phi_r \), gives the approximated solution of size \( n_1 \) by \( n_2 \) to \( u_h \) in (3.5).

Note that when \( n_1 \) is too large, the SVD operation can be expensive, which can be circumvented by the fact that the singular vectors are also eigenvectors of the product of \( U_{m_2} \) and its transpose, which is \( r \) by \( r \), and cheaper to calculate [65]. Also, when nonlinearities are involved, there are terms to be evaluated *e.g.* \( f(U) \), for which repeated matrix-vector products, *i.e.* \( \Phi_r^T f(\Phi_r U) \), are required. The complexity of the latter can be very high which costs enormous runtime. In this case, the *Discrete Empirical Interpolation Model (DEIM)* can be employed to mitigate the situation. See examples in *e.g.* [65] and [66].

POD is effective in certain cases. Nonetheless, it has two shortcomings [57]. Firstly, it is only defined for separated representations in two variables. Although higher dimensional extensions have been studied, the optimality as in two variables is not guaranteed. Secondly, *a priori* knowledge of the solution is needed to form the reduced basis, which in some occasions can be prohibitive. PGD, as its name suggests, is a more generalized model reduction method that addresses both issues of POD.
3.1. Separated representation based model reduction

3.1.2 The Proper Generalized Decomposition

PGD is also used for construction of low-dimensional approximated representations. In contrast to POD, PGD does not require a priori calculation on selected samples. Moreover, it is theoretically unlimited in terms of the number of variables. See e.g. [67] in which applying PGD in problems in 100 variables is presented. Since its creation [59], PGD has been widely utilized in various domains, especially on mechanics, see e.g. [68] and [69] among others. For electromagnetic and electromagnetics related multiphysics modeling, examples can be found in e.g. [45], [70], and [71].

The essential assumption of PGD is that the numerical solution can be expressed as low rank tensors, or sum of rank-one tensors [72]. Each term in the sum is called a mode while the number of terms is called the rank [73]. From a numerical perspective, each mode is a multidimensional array in which every dimension is related to some physical quantity. In practice, modes are calculated successively, which is referred to as the enrichment process. During enrichment, simultaneously determining all vectors in the array leads to a nonlinear problem that can be effectively solved by employing the alternating direction method (ADM) which is also called the fixed point iteration procedure [74]. The advantage of PGD is twofold. First, instead of solving a high-dimensional problem through looping over each sample of the parameter (i.e. as with the conventional approach), it solves the problem with all samples of parameter at the same time, thus avoiding a large number of loops and reducing runtime. Second, the number of unknowns to be determined is generally dramatically reduced since PGD solutions actually form reduced basis of the full solution (though this basis is not necessarily orthonormal, as opposed to the case in POD). Numerical experience suggests that only moderate number of enrichments and iterations (within enrichments) are required. As a result, less solution data needs to be stored which is important for real-time simulations [75].

On the other hand, PGD relies on the separability of the problem. For a discrete system, separating the r.h.s. is usually feasible, because excitations and boundary conditions based on which the r.h.s. is formed are normally known, for which techniques such as SVD can be used (there exist high-dimensional versions of SVD for problems in more than two variables, see e.g. [76]). By contrast, separating a general coefficient matrix is still an open problem, albeit some literatures on this topic are now available. For instance, in [77] operator separating in mechanical and thermal problems are discussed, in [78] the same issue for high-dimensional Helmholtz problems are discussed. After separating, various PGD formulations can be built depending on underlying properties of the problem [79]. We also note that, convergence rates
and optimality of the solution (i.e. attaining the same level of accuracy at the smallest rank) are important during the PGD procedure, see [79]. In the following, we review the process of PGD construction through the discrete abstract problem in (3.5), in which \( u_h \) now might be in more than two variables (namely, \( d \geq 2 \) for the equation in (3.4)).

Since separated representation in PGD are realized in the form of low-rank tensors, it is necessary to introduce formats of the latter. In [72], five types of such formats are summarized. They are called the canonical, \( \alpha \)-rank, Tucker, tree-based Tucker, and Tensor-train format, respectively. The canonical format is simple and very easy to build. For this reason we focus on this format in the thesis. While other formats have their own advantages, discussion about them is beyond the scope of this text. More information in this regard can be found in [72]. In canonical format, ensemble of solutions to (3.5) for all \( x_1, x_2, \ldots, x_d \) (among which time in transient analysis or frequency in harmonic analysis might be included) can be expressed as

\[
\mathbf{U}^{(n)} := \sum_{i=1}^{n} \mathbf{U}_i = \sum_{i=1}^{n} \bigotimes_{k=1}^{d} U_k^{(i)} := \sum_{i=1}^{n} U_1^{(i)} \otimes \cdots \otimes U_d^{(i)}
\]  

(3.9)

where \( \mathbf{U} \) is \( U \) in calligraphic form (representing tensors of order higher than two), \( n \) is the rank, whereas \( \mathbf{U}_i \) is the \( i \)-th mode, ‘\( \otimes \)’ denotes the Kronecker product. Since \( \mathbf{U} \) belongs to \( V_h \) that is subspace of \( \mathbb{C}^{n_1 \times \cdots \times n_d} \) (\( \mathbb{R}^{n_1 \times \cdots \times n_d} \) if only real values appear in the entries), the expression in (3.9) admits that \( V_h = \tilde{V}_h \times \cdots \times \tilde{V}_h \) in which \( U_k^{(i)} \in \tilde{V}_k \), \( k = 1, \ldots, d \), \( \tilde{V}_k \subseteq \mathbb{C}^{n_k} \). Hence, the relation between norms of \( \mathbf{U}_i \) and \( U_k^{(i)} \) can be expressed as in (3.10) where \( \langle \cdot, \cdot \rangle_{V_h} \) and \( \langle \cdot, \cdot \rangle_{\tilde{V}_k} \) denote Euclidean inner products, the superscript ‘\( \dagger \)’ denotes conjugate transpose.

\[
\| \mathbf{U}_i \|_Z := \langle \mathbf{U}_i, \mathbf{U}_i \rangle_{V_h}^{\frac{1}{2}} = \prod_{k=1}^{d} \langle U_k^{(i)}, U_k^{(i)} \rangle_{\tilde{V}_k}^{\frac{1}{2}} = \prod_{k=1}^{d} \langle U_k^{(i)} U_k^{(i)} \rangle_{V_h}^{\frac{1}{2}} = \prod_{k=1}^{d} \| U_k^{(i)} \|_Z
\]  

(3.10)

In addition, suppose that a linear system is derived from the discrete weak form (3.5), after applying e.g. the Galerkin method to the latter. We denote the coefficient matrix and r.h.s. in this system by, respectively, \( A \) and \( b \) whose entries are also in variables \( x_1, x_2, \ldots, x_d \). Then, we express ensemble of samples for \( A \) and \( b \) in corresponding tensor forms: \( A \in \mathbb{C}^{(n_1 \times n_2 \times \cdots \times n_d)^2} \) and \( b \in \mathbb{C}^{n_1 \times n_2 \times \cdots \times n_d} \). The key point is to decompose them into lower-rank tensor form. We assume this can be achieved, and separations can be expressed as
3.1. Separated representation based model reduction

\[ \mathcal{U} = \bigotimes_{k=1}^{d} A_k := A_1 \otimes A_2 \otimes \cdots \otimes A_k \quad , \quad \mathbf{b}r = \sum_{j=1}^{n_b} \bigotimes_{k=1}^{d} B_k^j \]  

(3.11)

where \( A_k \) is determined by discretization of \( x_k \), \( B_k^j \) can be obtained using a posteriori techniques like SVD, \( n_b \) is the number of modes for \( \mathbf{b}r \). As such, ensemble of linear systems for all parameters can be written in a compact form, as in (3.12).

\[ \mathcal{U} \cdot \mathbf{U} = \mathbf{b}r \]

(3.12)

In the equation, \( \mathbf{U} \) denotes the exact solution to the discrete system. Note that the system in (3.12) can be extremely large and impractical to solve directly. Using PGD, we turn it into a series of small systems to obtain an approximation of \( \mathbf{U} \), denoted by \( \mathbf{U}^{(n)} \). To this end, we can define the problem in the error norm \( i.e. \) the first equation in (3.13), which is optimal [79], or in the residual norm \( i.e. \) the second equation in (3.13). \( \mathbf{U}^* \) represents an arbitrary vector in the space \( V_h \). Since we do not solve (3.12) \( a \ priori \), and have no information about \( \mathbf{U} \), it seems more practical to choose the residual norm.

\[ \mathbf{U}^{(n)} = \arg \min_{\mathbf{U} \in V_h} \| \mathbf{U} - \mathbf{U}^* \|_2 \]

(3.13)

\[ \mathbf{U}^{(n)} = \arg \min_{\mathbf{U} \in V_h} \| \mathbf{b}r - \mathcal{U} \cdot \mathbf{U}^* \|_2 \]

The error norm can be simplified to \( ( \mathbf{b}r - \mathcal{U} \cdot \mathbf{U}^*)^\dagger \cdot ( \mathbf{b}r - \mathcal{U} \cdot \mathbf{U}^*) \), which, after expansion and dropping \( \mathbf{b}r^\dagger \mathbf{b}r \) (as it is unrelated to \( \mathbf{U}^* \)), leads to the function to be minimized in (3.14).

\[ f (\mathbf{U}^*) = \mathbf{U}^*^\dagger \mathcal{U}^\dagger \mathbf{U}^* - \mathbf{b}r^\dagger \mathbf{U}^* - \mathbf{U}^*^\dagger \mathbf{U}^\dagger \mathbf{b}r \]

(3.14)

Finally, using vector calculus (namely, taking derivative of \( f (\mathbf{U}^*) \) w.r.t. \( \mathbf{U}^* \) and setting the result to zero) \( \mathbf{U}^* \) can be solved, as in (3.15).

\[ \mathcal{U}^\dagger \mathbf{U}^* - \mathcal{U}^\dagger \mathbf{b}r = 0 \]

(3.15)
When $\mathcal{A}$ is Hermitian and defining a norm, $\mathcal{A}^\dagger$ can be cancelled from (3.15), yielding

$$\mathcal{A} \mathcal{U}^* - \mathcal{E} = 0 \quad (3.16)$$

which leads to the Galerkin-PGD. Otherwise, (3.15) is used, which leads to the Minimal Residual-PGD. But it is advised not to use the latter, because of its slow convergence rate [79]. Apart from the above two, there is another called Minimax-PGD in [79], which can be interpreted as the Petrov-Galerkin method based on (3.15). It delivers better convergence rates than Minimal Residual-PGD in certain cases. A more recent type of PGD, based on the Ideal Minimal Residual (IMR) method is presented in [80]. IMR-PGD can allegedly enhance convergence rate when others fail to. The idea is to use an energy norm that is equivalent to the error norm in (3.13), while requires no information about $\mathcal{U}$.

In this thesis, we only utilize Galerkin-PGD, which is straightforward to implement and efficient over a large range of applications [68]. To proceed with (3.16), we first need to re-write the tensor system into explicit matrix-vector form, for which properties of Kronecker product in (3.17) can be used [81].

\[
\begin{align*}
\text{Transpose:} & \quad (B \otimes C)^\dagger = B^\dagger \otimes C^\dagger \\
\text{Inverse:} & \quad (B \otimes C)^{-1} = B^{-1} \otimes C^{-1} \\
\text{Product:} & \quad (B \otimes C) \cdot (D \otimes F) = B \cdot D \otimes C \cdot F \\
\text{Associativity:} & \quad B \otimes (C \otimes D) = (B \otimes C) \otimes D
\end{align*}
\quad (3.17)
\]

Further, we replace $\mathcal{U}^*$ with $\mathcal{U}^{(n)}$, which is the solution we eventually obtain. Substituting (3.9) and (3.11) into (3.16) and applying properties in (3.17), yields (3.18).

\[
\sum_{i=1}^{n} \otimes_{k=1}^{d} A_{k} U_{k}^{i} = \sum_{j=1}^{n} \otimes_{k=1}^{d} B_{k}^{j} \quad (3.18)
\]

As mentioned above, a PGD process consists of multiple enrichments in which sequences of iterations using ADM are enclosed inside. Take the $m$-th iteration for instance (with $1 \leq m \leq n$, the maximum enrichment number). Modes obtained in previous enrichments (zeros if $m$ is 1) are put onto the r.h.s., as presented in (3.19).
3.2 A non-incremental algorithm for transient magneto-thermal problems

\[
\bigotimes_{k=1}^{d} A_k U_k^m = \sum_{j=1}^{n_b} \bigotimes_{k=1}^{d} B_k^j - \sum_{i=1}^{m-1} \bigotimes_{k=1}^{d} A_k U_k^i
\]  \hspace{1cm} (3.19)

Using ADM, both sides of (3.19) are pre-multiplied with \((U_{m}^{m'} \prod_{\alpha=1}^{k} U_{\alpha}^{m})^T\), in which \(l\) takes value from 1 to \(d\) in order during iterations, \(U_{m}^{m'}\) denotes a test vector whereas \(U_{\alpha}^{m}\) are known vectors which can be assigned with arbitrary values in the very first iteration (namely, the first ADM iteration for \(m = 1\)) and with results obtained in previous iterations otherwise. Multiplication of \(U_{m}^{m}\) with \(A_{\alpha} U_{\alpha}^{\beta}\) (\(\beta\) stands for \(m\) and \(i\)) and \(B_{\alpha}^{j}\) in (3.19) results in scalars. As such, a system for \(U_{m}^{m}\) can be obtained, as expressed in (3.20), where \(\hat{B}_i\) is obtained from the aforementioned scalars and other terms in the r.h.s. This is repeated during enrichments.

\[
A_i U_i^m = \hat{B}_i
\]  \hspace{1cm} (3.20)

Meanwhile, enrichments and iterations can be controlled with prescribed convergence criterions and maximum counters. For faster convergence, solutions obtained from (3.20) are normalized and updated in practice. See [44] for examples on these topics. In this fashion, the original large-size problem is turned into a series of smaller-size problems. Since the total number of ADM iterations is normally moderate, considerable runtime can be saved. In addition, space for storing solutions can also be significantly spared. Based on what has been introduced above, we present in the following two novel algorithms that are dedicated for multiphysics problems within our multiphysics framework. The first is designed for solving non-linear transient problems, in which electromagnetics and thermal fields are strongly coupled, through non-incremental approaches, with which solutions are separated into space and time components. The second algorithm is dedicated to parametric problems involving piezoelectric materials. The algorithm is introduced directly into the discrete piezoelectric system, which has been presented in § 2.3, after which solutions are decomposed into components associated with different parameters.

3.2 A non-incremental algorithm for transient magneto-thermal problems

In this section, we present a PGD based algorithm [70] for transient magneto-thermal problems, in which magnetodynamic and thermodynamic fields are strongly coupled, as electric
material properties depend on temperature, while the latter evolves due to the accumulated Joule heat. For our applications, such problems arise when, for example, eddy currents generated in magnetostrictive materials need to be considered. Solving the problem using conventional time integration algorithms are impractical, due to significant differences between electromagnetic and thermal time constants, which can lead to enormous amount of time steps, rendering the problem too large to be solved. We propose a novel algorithm circumventing this issue through decomposing variables into the space and time components, and solving linearized systems in space and time iteratively. We also show that material nonlinearities in the problem can be handled in a very convenient way.

3.2.1 The coupled nonlinear dynamic problem

The underlying method utilized is referred to as the space-time separated representation method [67]. It has been applied in e.g. problems of rheology [82], fluid flow [83], and electrodynamics [80] on mechanics, as well as magnetoquasistatic problems [84] on electromagnetics. The work presented in this section appears to be the first one, which applies the method on nonlinear magneto-thermal problems. The latter category of problems is important in cases where losses due to eddy currents are involved. Available solutions to these problems are mostly based on the harmonic balance method, which provides a way to trading off between costs and accuracy. However, according to [85] the computational burden may not be significantly decreased if high accuracies of the solution are expected. For the presentation of our proposition, mathematical formulations governing the coupled problem are first present. To keep focusing on the algorithm, we do not take into account elastic field in the magnetostrictive system in (2.80). Also, we present our formulations in cylindrical coordinates, so that the current work can be directly applied to many realistic applications e.g. inductive heating. Elaborating this algorithm for magnetostrictive materials involved simulations (like those introduced in § 2.4.3) are discussed at the end of the section.

There are numerous references dealing with magnetodynamic fields involved multiphysics problem, such as [86], [87] and [88]. We recall that equations governing the magnetodynamics are introduced in (2.17) and (2.18). Additionally, we assume that the electric conductivity \( \sigma_e \) in (2.18) depends on temperature, which is denoted by \( g(\theta) \) for the moment. We still employ potential variables in (2.69) and start from (2.70), while differences here is that for \( A \) and \( J_0 \), only components in the \( \varphi \) axis are non-vanishing (denote them by \( A^\varphi \) and \( J_0^\varphi \)); further, the scalar potential \( \phi \) in (2.70) disappears. Gauging conditions are automatically satisfied if Dirichlet conditions are applied to \( A^\varphi \). Therefore, the governing equation can be simplified as
3.2. A non-incremental algorithm for transient magneto-thermal problems

\[
\sigma_c \frac{\partial A^\varphi}{\partial t} + \frac{\partial}{\partial r} \left( 1 \frac{\partial (rA^\varphi)}{\partial r} \right) + \frac{\partial}{\partial z} \left( 1 \frac{\partial A^\varphi}{\partial z} \right) = J^\varphi_s. \tag{3.21}
\]

where \(r\) and \(z\) represent axis of the cylindrical coordinates. Introducing new variables

\[
\hat{A} = rA^\varphi, \quad \hat{J} = J^\varphi_s. \tag{3.22}
\]

and substituting them into (3.21) leads to

\[
\frac{\sigma_c}{r} \frac{\partial \hat{A}}{\partial t} + \frac{\partial}{\partial r} \left( 1 \frac{\partial \hat{A}}{\partial r} \right) + \frac{\partial}{\partial z} \left( 1 \frac{\partial \hat{A}}{\partial z} \right) = \hat{J}. \tag{3.23}
\]

Let \(\Omega_A\) denote the magnetic domain whose boundary \(\Gamma_A\) can be parted as union of Dirichlet boundaries \(\Gamma_A^D\) and Neumann boundaries \(\Gamma_A^N\), associated with boundary conditions as in the first two equations in (3.24) where \(\mathbf{n}\) denotes the outward unit vector normal to \(\Gamma_A^N\). Initial conditions can be expressed by the third one.

\[
\hat{A} = \hat{A}_D \quad \text{on} \quad \Gamma_A^D
\]

\[
\frac{\partial \hat{A}}{\partial \mathbf{n}} = \hat{A}_N \quad \text{on} \quad \Gamma_A^N \tag{3.24}
\]

\[
\hat{A}(t = 0) = \hat{A}_0
\]

The thermal problem is governed by the equation in (2.21), in which the rate of heat generation \(s\) now is related to the magnetic problem, and can be expressed by the r.h.s. of (3.25).

\[
\rho c_p \frac{\partial \theta}{\partial t} - \kappa \left( \frac{\partial^2 \theta}{\partial r^2} + \frac{\partial^2 \theta}{\partial z^2} \right) = \sigma_c \left( \frac{1}{r} \frac{\partial \hat{A}}{\partial t} \right)^2. \tag{3.25}
\]

Let \(\Omega_\theta\) denote the thermal domain whose boundary \(\Gamma_\theta\) can be parted as union of Dirichlet boundaries \(\Gamma_\theta^D\) and Neumann boundaries \(\Gamma_\theta^N\), associated with boundary conditions as in the first two equations in (3.26) where \(\mathbf{n}\) denotes the outward unit vector normal to \(\Gamma_\theta^N\). Initial conditions for \(\theta\) can be expressed by the third equation in (3.26).
\[ \theta = \theta_D \quad \text{on } \Gamma^D_\theta \]
\[ -\kappa \cdot \nabla \theta / \partial n = s_N \quad \text{on } \Gamma^N_\theta \]
\[ \theta(t = 0) = \theta_0 \]  

(3.26)

### 3.2.2 Conventional time integration approach

Weak forms of the coupled problem can be obtained by pre-multiplying equations in (3.23) and (3.25) with test functions \( \hat{A}' \) and \( \theta' \), respectively, and integrating the resultant system over corresponding domains, as expressed in (3.27).

\[
\int_{\Omega_A} \hat{A}' \left\{ \sigma_c \frac{\partial \dot{A}}{\partial t} + \frac{1}{\mu r} \left( \frac{\partial^2 \dot{A}}{\partial r^2} + \frac{\partial^2 \dot{A}}{\partial z^2} \right) - \dot{\mathbf{j}} \right\} d\Omega = 0
\]
\[
\int_{\Omega_\theta} \theta' \left\{ \rho c_p \frac{\partial \theta}{\partial t} - \kappa \left( \frac{\partial^2 \theta}{\partial r^2} + \frac{\partial^2 \theta}{\partial z^2} \right) - \sigma_c \frac{1}{r} \left( \frac{\partial \dot{A}}{\partial t} \right)^2 \right\} d\Omega = 0
\]

(3.27)

Next, we treat the problem in (3.27) the same fashion as in Chapter 2; namely, we use the same triangulations for \( \Omega_A \) and \( \Omega_\theta \), then solve magnetic and thermal unknowns simultaneously. Denote the triangulation by \( \Omega^h \), which is the union of \( n_X \) FEs. Equations in (3.27) can be projected onto finite-dimensional spaces. For simplicity, \( \sigma_c \) and \( r \) are approximated as element-wise constant (denote them as \( \sigma_{c\alpha} \) and \( r_{\alpha} \) for the element \( \mathcal{K}_\alpha \)), as expressed in (3.28).

\[
\sum_{\alpha=1}^{n_X} \int_{\mathcal{K}_\alpha} \hat{A}' \left\{ \sigma_{c\alpha} \frac{\partial \dot{A}}{\partial t} + \frac{1}{\mu r_{\alpha}} \left( \frac{\partial^2 \dot{A}}{\partial r^2} + \frac{\partial^2 \dot{A}}{\partial z^2} \right) - \dot{\mathbf{j}} \right\} d\Omega = 0
\]
\[
\sum_{\alpha=1}^{n_X} \int_{\mathcal{K}_\alpha} \theta' \left\{ \rho c_p \frac{\partial \theta}{\partial t} - \kappa \left( \frac{\partial^2 \theta}{\partial r^2} + \frac{\partial^2 \theta}{\partial z^2} \right) - \sigma_{c\alpha} \frac{1}{r_{\alpha}^2} \left( \frac{\partial \dot{A}}{\partial t} \right)^2 \right\} d\Omega = 0
\]

(3.28)

Techniques used for 2D Cartesian coordinate system based FE problems can be directly applied to (3.28). We use 2D nodal elements (defined in (2.16)) for discretization. Over \( \mathcal{K}_\alpha \), \( \dot{A} \) and \( \theta \) are interpolated with shape functions \( W^{n,2D}_\alpha \) and nodal values \( \dot{A}_\alpha \) and \( \theta_\alpha \), as depicted in (3.29) where we use superscript ‘†’ in lieu of ‘t’ to avoid confusions in the subsequent.
3.2. A non-incremental algorithm for transient magneto-thermal problems

\[
\hat{A} = W^{n,2D,t}_\alpha \hat{A}_\alpha, \quad \theta = W^{n,2D,t}_\alpha \theta_\alpha
\]  
(3.29)

Note that notation \(\hat{A}\) is used for both functions and vectors of discrete values, whose meaning are supposed to be told from the context. Eventually, the global FE system can be deduced by following the same procedure as in § 2.3.2.1, as depicted in (3.30). Definitions of terms in the equation are provided in § B.1.

\[
C_A(\theta) \frac{\partial \hat{A}}{\partial t} + K_A \hat{A} = J
\]
(3.30)

\[
C_\theta \frac{\partial \theta}{\partial t} + K_\theta \theta = S(\theta, \hat{A})
\]

For time discretization, we use the Crank-Nicholson schema, with which ordinary differential equations (ODE) in (3.30) are solved in a step by step fashion. To account for nonlinearity, both the Picard (also referred to as the fixed point method) and Newton method can be employed. However, for a general nonlinear relation \(g(\theta)\), the Newton method might not be appropriate, because calculating the Jacobian can be very involved. On the other hand, the Picard method is simpler as it linearizes the system at the current time step by substituting results from the previous time step into the nonlinear terms. Nonetheless, the Picard method may converge slowly or fail to converge, if the time step size is large. To prevent this, we choose sufficiently small time steps. The conventional approach is usually referred to as incremental, while our PGD based approach is non-incremental.

3.2.3 Non-incremental space-time separation approach

3.2.3.1 Weak form and space-time discretization

Our non-incremental approach is based on PGD, with which it is assumed that functions \(\hat{A}\) and \(\theta\) admit separation as in (3.31) where subscripts \(X\) and \(t\) implies spatial and temporal association, \(r\) denotes the maximum number of modes, or rank that depends on the prescribed accuracy and maximum counter of enrichments (as presented in the coming sections).

\[
\hat{A}^{(r)} \approx \sum_{i=1}^{r} \hat{A}_X^i \hat{A}_t^i, \quad \theta^{(r)} \approx \sum_{i=1}^{r} \theta_X^i \theta_t^i
\]
(3.31)
Meanwhile, separation of $f$ in (3.23) is trivial, as we introduced in § 3.1.2. We present the separation in (3.32), in which $r_j$ represents the number of modes (corresponding to e.g. the number of stranded inductors in reality), $f_X^j$ and $f_t^j$ represents, respectively, the unit current density in coils and the time function of electric currents.

$$f^{(r_j)} \approx \sum_{i=1}^{r_j} f_X^i f_t^i$$  \hspace{1cm} (3.32)

On the other hand, terms in (3.31) are calculated \textit{a priori}, through enrichments. With reference to (3.19), we assume that at the $m$-th enrichment, the previous $m-1$ modes are available, as depicted in (3.33), where $A_{x}^m A_t^m$ and $\theta_x^m \theta_t^m$ are new modes to be calculated.

$$A^{(m)} = A^{(m-1)} + A_{x}^m A_t^m := \sum_{i=1}^{m-1} A_{x}^i A_t^i + A_{x}^m A_t^m$$
$$\theta^{(m)} = \theta^{(m-1)} + \theta_x^m \theta_t^m := \sum_{i=1}^{m-1} \theta_x^i \theta_t^i + \theta_x^m \theta_t^m$$  \hspace{1cm} (3.33)

At the same time, nonlinear terms involved in the Joule heat source are handled so that at the $m$-th enrichment we have

$$\sigma_c = g(\theta^{(m-1)}) \quad , \quad \frac{\partial A}{\partial t} = \frac{\partial}{\partial t}(A^{(m-1)}) = \sum_{i=1}^{m-1} A_{x}^i \frac{\partial A_t^i}{\partial t}.$$  \hspace{1cm} (3.34)

Since each mode has both space and time unknowns components, test functions $\hat{A}'$ and $\theta'$ in (3.27) now correspond to $\hat{A}_{x}^{m'} \hat{A}_t^{m'} + \hat{A}_{x}^{m} \hat{A}_t^{m}$ and $\theta_x^{m'} \theta_t^{m'} + \theta_x^{m} \theta_t^{m'}$, respectively. To obtain the weak form, integrals are now over the entire \textit{space-time domain}, which can be denoted by $\Omega \otimes I$. $\Omega$ is the spatial domain, whereas $I$ is the temporal domain. The weak form is expressed in (3.35), which needs to be projected onto a finite-dimensional subspace (\textit{i.e.} discretization).

For $\Omega$, we use the same discretization as the in § 3.2.2. For $I$, we also use FE discretization. More precisely, we denote the discrete time domain by $I^h$, which consists of $t_0, t_1, \ldots, t_{n_t}$ with $t_0 < t_1 < \cdots < t_{n_t}$ ($n_t$ is the total number of time instants).
3.2. A non-incremental algorithm for transient magneto-thermal problems

\[ \int_{\Omega^h} \left( A^{n'} A'^m + \hat{A}^m A^{m'} \right) \left\{ \frac{g(\theta^{(m-1)})}{r} \frac{\partial (\hat{A}^{(m-1)} + \hat{A}^m A'^m)}{\partial r^2} + \frac{\partial t}{\partial t} \frac{\partial^2 (\hat{A}^{(m-1)} + \hat{A}^m A'^m)}{\partial z^2} \right\} d\Omega dI = 0 \]

\[ \int_{\Omega^h} \left( \theta^{m'} \theta'^m + \theta^m \theta^{m'} \right) \left\{ \frac{\rho c_p}{r} \frac{\partial (\theta^{(m-1)} + \theta^m \theta'^m)}{\partial t} - \kappa \left( \frac{\partial^2 (\theta^{(m-1)} + \theta^m \theta'^m)}{\partial r^2} + \frac{\partial^2 (\theta^{(m-1)} + \theta^m \theta'^m)}{\partial z^2} \right) \right\} d\Omega dI = 0 \]

(3.35)

The discrete space-time domain can be denoted by \( \Omega^h \otimes I^h \), and expressed as in (3.36) where \( I_j \) denotes the time element \([t_{j-1}, t_j]\).

\[ \Omega^h \otimes I^h := \sum_{\alpha=1}^{n_X} \sum_{\beta=1}^{n_T} K_{\alpha} \otimes I_{\beta} \]  

(3.36)

The discrete weak form is obtained through projecting continuous weak forms in (3.35) onto the discrete space-time domain \( \Omega^h \otimes I^h \). The resultant discrete systems are presented in § B.2. We denote solutions to the discrete system by \( \hat{A}^{X,m}, \hat{A}^{t,m}, \theta^{X,m}, \theta^{t,m} \) (which are associated with continuous variables in (3.33) \( \hat{A}^m, \theta^m \), respectively).

Further, we denote vectors containing spatial nodal values in \( K_{\alpha} \) and temporal nodal values in \( I_{\beta} \) for \( \hat{A} \) (respectively, \( \theta \)) by \( \hat{A}_{\alpha} \) and \( \hat{A}_{\beta} \) (respectively, \( \theta_{\alpha} \) and \( \theta_{\beta} \)), and the 1D nodal shape functions for time discretization as \( W_{\beta}^{n,1D} \). As such, \( \hat{A} \) and \( \theta \) over the space-time element \( K_{\alpha} \otimes I_{\beta} \) can be interpolated as

\[ \hat{A} = (W_{\alpha}^{n,2D,\dagger} \hat{A}_{\alpha}) \otimes (W_{\beta}^{n,1D} \hat{A}_{\beta}) \] \[ \theta = (W_{\alpha}^{n,2D,\dagger} \theta_{\alpha}) \otimes (W_{\beta}^{n,1D} \theta_{\beta}) \]

(3.37)

Eventually, the FE system used to solve for \( \hat{A}^{X,m}, \hat{A}^{t,m}, \theta^{X,m}, \theta^{t,m} \) can be formed, through replacing variables in the discrete weak form (in § B.2) with the interpolants in (3.37). For
conciseness, we define coefficient matrices $C_A^X$, $K_A^X$, $M_A^t$, $K_A^t$, $C_\theta^X$, $K_\theta^X$, $K_\theta^\theta$, and $M_\theta^t$, that are shape functions related to integrals, as presented in § B.3.

3.2.3.2 Enrichment procedure

Obviously, the obtained global system is nonlinear. As introduced in § 3.1.2, ADM can be employed. It this case, ADM consists in iteratively solving the temporal problem — giving $(\hat{A}^{X,m}, \Theta^{X,m})$ — then solving for $(\hat{A}^t, \Theta^t)$, and the spatial problem — giving $(\hat{A}^{X,t}, \Theta^{X,t})$ — then solving for $(\hat{A}^{X,m}, \Theta^{X,m})$. The temporal problem can be formulated by assigning $\hat{A}^{X,m} \hat{A}^{t,m}$ to $\hat{A}$ and $\Theta^{X,m} \Theta^{t,m}$ to $\Theta$ in (3.35), whereas the spatial problem can be formulated by assigning $\hat{A}^{X,t} \hat{A}^{t,m}$ to $\hat{A}$ and $\Theta^{X,t} \Theta^{t,m}$ to $\Theta$. Algebraic systems for the temporal and spatial problems are presented in respectively, (3.38) and (3.39). Subscripts $A$ and $\theta$ of matrices imply magnetic and thermal problems, whereas superscripts $X$ and $t$ (not for tensor transpose) imply spatial and temporal problems, respectively. Note that vectors $f^X$ and $f^t$ result from discretization of $\hat{f}_{X}^i$ and $\hat{f}_{t}^i$ in (3.32), whereas $S^t$ and $S^X$ result from separating the term related to Joule heat in (3.35).

\[
\begin{align*}
\left[(\hat{A}^{X,m} C_A^X \hat{A}^{X,m}) \right] K_A^t + \left[(\hat{A}^{X,m} K_A^X \hat{A}^{X,m}) \right] M_A^t \right] \hat{A}^{t,m} \\
= \sum_{i=1}^{m-1} \left[\left((\hat{A}^{X,m} C_A^X \hat{A}^{X,m}) \right) K_A^t + \left((\hat{A}^{X,m} K_A^X \hat{A}^{X,m}) \right) M_A^t \right] f^{t,i} \\
&= \sum_{j=1}^{r_j} \left[\left((\hat{A}^{X,m} C_A^X \hat{A}^{X,m}) \right) K_A^t + \left((\hat{A}^{X,m} K_A^X \hat{A}^{X,m}) \right) M_A^t \right] f^{t,i}
\end{align*}
\]

(3.38)

\[
\begin{align*}
\left[(\Theta^{X,m} C_\theta^X \Theta^{X,m}) \right] K_\theta^t + \left[(\Theta^{X,m} K_\theta^X \Theta^{X,m}) \right] M_\theta^t \right] \Theta^{t,m} \\
= \sum_{i=1}^{m-1} \left[\left((\Theta^{X,m} C_\theta^X \Theta^{X,m}) \right) K_\theta^t + \left((\Theta^{X,m} K_\theta^X \Theta^{X,m}) \right) M_\theta^t \right] f^{t,i} \\
&= \sum_{j=1}^{r_\theta} \left[\left((\Theta^{X,m} C_\theta^X \Theta^{X,m}) \right) K_\theta^t + \left((\Theta^{X,m} K_\theta^X \Theta^{X,m}) \right) M_\theta^t \right] f^{t,i}
\end{align*}
\]

(3.39)

Meanwhile, for equations in (3.31) it might occur that space values turn to infinitely large,
while temporal values to infinitely small (thus leading to numerical overflow). To prevent it, we can normalize \(A_{m+1}^t\) and \(\Theta_{m+1}^t\) by dividing them with their corresponding norms. Also, for efficiency considerations, the maximum iteration counter in ADM (denote it by \(n_{\text{in}}\)) can be set to lower than 10. A tolerance of \(\sim 10^{-4}\) can be used as converging criterion (denoted by \(\varepsilon_{\text{in}}\)), which can be utilized as in (3.40). The symbol \(Y\) refers to both \(A\) and \(\Theta\), which applies to (3.41) and (3.42) in the following, too.

\[
\|Y^{t,m} - Y^{t,m-1}\|_2 / \|Y^{t,m-1}\|_2 \leq \varepsilon_{\text{in}} \tag{3.40}
\]

Once \(A_{m+1}^t, \bar{A}_{m+1}^t, \Theta_{m+1}^t, \) and \(\Theta_{m+1}^t\) are calculated, we can obtain approximations for the real solutions as: \(A^{(m)}\) (i.e. \(\sum_{i=1}^{m} \bar{A}^{t,i} \otimes \bar{A}^{t,i}\)) and \(\Theta^{(m)}\) (i.e. \(\sum_{i=1}^{m} \Theta^{t,i} \otimes \Theta^{t,i}\)). To improve convergence rates, we can update all time components through solving for \(\bar{A}^{t,i}\) and \(\Theta^{t,i}\) from (3.41), in which \(\mathfrak{U}_Y\) and \(\delta_Y\) denote ensemble of the matrices and r.h.s.s corresponding to all time values, \(Y^{t,i}\) are test vectors. Spatial values \(Y^{X,i}\) are not updated, because otherwise it results in a very large system (of size \(2m \cdot n_x\)^2, as a matter of fact). In contrast, the problem size for updating \(Y^{t,i}\) is \((2m \cdot n_t)^2\), which is much smaller.

\[
\left(\sum_{i=1}^{m} Y^{X,i} \otimes Y^{t,i}\right)^\dagger \mathfrak{U}_Y \left(\sum_{i=1}^{m} Y^{X,i} \otimes Y^{t,i}\right) = \left(\sum_{i=1}^{m} Y^{X,i} \otimes Y^{t,i}\right)^\dagger \delta_Y \tag{3.41}
\]

The enrichment procedure can be terminated when reaching the maximum counter (denoted by \(n_{\text{out}}\), or the following criterion is satisfied.

\[
\|Y^m\|_2 / \|Y^1\|_2 = \|Y^{t,m}\|_2 / \|Y^{t,1}\|_2 \leq \varepsilon_{\text{out}} \tag{3.42}
\]

The algorithm is summarized and presented in Figure 3.1. We see that during enrichments, iterations for magnetic and thermal problems are actually “no longer coupled” (namely, unknowns of the magnetic and thermal problems are solved separately), which makes it possible to solve these two problems in parallel. With the novel algorithm, complexity of the

\[\text{3 The subscript ‘In’ and ‘Out’ (that is to appear in the following) can be explained from the fact that ADM iterations are inside loops that are enclosed by enrichments, which can be called as outside loops.}\]
original problem can be considerably reduced, depending on the number of enrichments and iterations required before convergence. See [70] for more detailed analysis.

\[
\sigma_c = \sigma_{co}/[1 + \gamma(\theta - \theta_0)]
\]  

(3.43)

Temperature dependence of the electric conductivity of the conductor (or workpiece), \textit{i.e.} the function \(g(\theta)\) is given in equation (3.43) where the coefficient \(\gamma\) equals \(3.9 \times 10^{-3}\), the ref-

3.2.4 Numerical application

The proposed algorithm is applied on a numerical example whose configurations, including model configuration and triangulation are provided in § B.4.
3.2. A non-incremental algorithm for transient magneto-thermal problems

erece temperature \( \theta_0 \) equals 293.15 K. Other constants are depicted in the following table.

Table 3.1. Material constants in the inductive heating model

<table>
<thead>
<tr>
<th>Property</th>
<th>work-piece</th>
<th>coil</th>
<th>air</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \sigma_{\text{el}} ) (( \Omega^{-1} \cdot \text{m}^{-1} ))</td>
<td>5.7013 \times 10^7</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>( \mu_r )</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>( \rho c_p ) (( \text{J} \cdot \text{kg}^{-1} \cdot \text{K}^{-1} ))</td>
<td>3.3495 \times 10^6</td>
<td>1.2 \times 10^3</td>
<td>1.2 \times 10^3</td>
</tr>
<tr>
<td>( \kappa ) (( \text{W} \cdot \text{m}^{-1} \cdot \text{K}^{-1} ))</td>
<td>400</td>
<td>0.02</td>
<td>0.02</td>
</tr>
</tbody>
</table>

Three sets of simulations are performed: the first set with the novel algorithm; the second set with the conventional approach as introduced in 3.2.2; the last set with the conventional approach, too, while using refined triangulations (of space and time). Results of the last set of simulations are used as references. In Figure 3.2, results for eddy currents at selected time instants during two periods, which are extracted from PGD results, are depicted in two rows. The upper row contains results of the first period, whereas those in the lower one are of the ninth period. We see that currents in the

![Figure 3.2. Snapshots of eddy currents during the first (1-9 ms) and ninth period (91-99 ms).](image)

lower row are propagating deeper (from the surface) than those in the upper row; meanwhile, peak values of the currents in the lower row are smaller. This is due to the fact that temperature rises because of the accumulated Joule heat, which in turn, reduces the electric conductivity and eventually affects the peak values.
Figure 3.3. Results from different sets of simulations for (a) the magnetic and (b) thermal problem.

Results from all three sets of simulations are presented in Figure 3.3. Excellent accordance between results of our PGD-based non-incremental approach and the conventional incremental approach can be observed. To be more precise, it is found that relative errors between the latter two are 0.36% for magnetic results, and 0.20% for thermal results. Comparison in terms of runtime is not presented here, because simulations with novel and conventional approaches were carried out on different platforms. On the other hand, the solution size of PGD simulation is nearly 75% smaller than its conventional counterpart, suggesting that the complexity has been reduced to almost a quarter thanks to the novel algorithm.

As for elaborating the algorithm for magnetostrictive simulations, since it is designed for transient analysis, elastodynamics in time domain will get involved. The latter is parabolic, and requires extra efforts to ensure good convergence (see [80]). Still, when DEAM is employed to model magnetostrictive nonlinearity, separability of the system might be problematic. In that case, an alternative way is to resort to POD, which does not have such constraints.
3.3 A fast algorithm for parametric electro-mechanical problems

In this section, we present another algorithm that can be used to reduce complexity of parametric piezoelectric problems. We present it via the bimorph energy harvester example that has been studied in Chapter 2. Parametric piezoelectric problems are important. For example, we have seen in the study in § 2.3.3.1, that outputs of the energy harvester can be significantly affected by both mechanical inputs and electric loads. On the other hand, the complexity of such problems grows exponentially with the number of parameters, which renders parametric studies of large FE models impractical. To preserve advantages of our multiphysics FE models while reducing complexity, we propose a PGD based algorithm [45]. Applying our approach on the piezoelectric bimorph example, variables are decomposed into space, frequency, and electric load related components; the large-size system is then converted to a sequence of smaller-size subsystems. In this manner, not only the problem complexity is reduced, memories for storing results can also be significantly spared.

3.3.1 Separated representation of the parametric problem

We recall that the problem being investigated consists of $n_X^u$ mechanical and $n_X^p$ electrical DoFs, in which $n_X^u$ and $n_X^p$ are usually large. Also, we intend to calculate outputs for different excitation frequencies in $[\omega_1, \omega_2, \cdots, \omega_{n_\omega}]$ and loads in $[r_1, r_2, \cdots, r_{n_R}]$, $n_\omega$ and $n_R$ denoting numbers of respective parameters. Expressing solutions to the system in (2.50) for all parameters in an ensemble leads to three-way arrays. Our goal is to find low-order tensors that approximate these arrays. As illustrated in Figure 3.4 where $\mathcal{Y}$ represents ensemble of solutions of displacements $U$ and electrical potentials $\mathcal{P}$, while $Y$ stands for both $U$ and $\Phi$ in (2.50).

$$
\mathcal{U}(X, \omega, R) \approx \sum_{i=1}^{r} U_{X1} \otimes U_{\omega i} \otimes U_{Rl}
$$

$$
\mathcal{B}(X, \omega, R) \approx \sum_{i=1}^{r} \Phi_{X1} \otimes \Phi_{\omega i} \otimes \Phi_{Rl}
$$

(3.44)

Figure 3.4. Schema of approximating solutions of three-way arrays with low-order tensors.
Chapter 3. PGD Based Model Reduction of Multiphysics Systems

The approximation can also be expressed by equations in (3.44), where \( r \) is the rank, while \( U_{Xi} \otimes U_{oi} \otimes U_{ri} \) and \( \Phi_{Xi} \otimes \Phi_{oi} \otimes \Phi_{ri} \) are, respectively, displacement and potential mode. As in (3.11), we also express ensemble of coefficient matrices and r.h.s.s of the system in (2.50) in tensor form. Regarding the r.h.s.s, we assume that identical mechanical excitations are applied in all cases of \( \omega \) and \( R \). Then, the r.h.s. of the whole system can be expressed as \( F_X \otimes F_\omega \otimes F_R \), in which entries of \( F_\omega \) and \( F_R \) are all equal to 1, are vectors of size \( n_\omega \) and \( n_R \), respectively. Regarding coefficient matrices, the expression is straightforward, thanks to our modeling of electric loads and electrodes in § 2.3.2.2, with which variables related to \( X, \omega \) and \( R \) are individual terms in products. The tensor consists of four blocks, as shown in (3.45).

\[
\mathcal{A} = \begin{bmatrix} \mathcal{U}_u & \mathcal{U}_{u\varphi} \\ \mathcal{U}_{\varphi u} & \mathcal{U}_\varphi \end{bmatrix} \tag{3.45}
\]

where blocks are defined by

\[
\begin{align*}
\mathcal{U}_{u\varphi} & := K_{u\varphi} \otimes D_\omega \otimes D_r \\
\mathcal{U}_{\varphi u} & := K_{u\varphi}^T \otimes D_\omega \otimes D_r \\
\mathcal{U}_\varphi & := K_{\varphi} \otimes D_\omega \otimes D_r - j\tilde{K}_{\varphi} \otimes G_\omega \otimes G_r \\
\mathcal{U}_u & := -M_u \otimes A_\omega \otimes D_r + jC_u \otimes B_\omega \otimes D_r + K_u \otimes D_\omega \otimes D_r
\end{align*}
\tag{3.46}
\]

For definitions in (3.46), \( D_\omega \) and \( D_r \) equal identity matrices of respectively \( n_\omega \) and \( n_R \) order. Others that are not defined in (2.50) are diagonal matrices, as shown in the following.

\[
\begin{align*}
A_\omega & := \text{diag}\{\omega_1^2, \omega_2^2, \ldots, \omega_{N_\omega}^2\} \in \mathbb{R}^{n_\omega \times n_\omega} \\
B_\omega & := \text{diag}\{\omega_1, \omega_2, \ldots, \omega_{N_\omega}\} \in \mathbb{R}^{n_\omega \times n_\omega} \\
G_r & := \text{diag}\{1/r_1, 1/r_2, \ldots, 1/r_{N_R}\} \in \mathbb{R}^{n_R \times n_R} \\
G_\omega & := \text{diag}\{1/\omega_1, 1/\omega_2, \ldots, 1/\omega_{N_\omega}\} \in \mathbb{R}^{n_\omega \times n_\omega}
\end{align*}
\tag{3.47}
\]

To keep notations neat, we introduce two calligraphic symbols \( b \) and \( x \), so that objects can be accessed via subscripts of these latter, see (3.48).
3.3. A fast algorithm for parametric electro-mechanical problems

\[ b = [b_1, b_2] = [F_X \otimes F_\omega \otimes F_R, 0] \tag{3.48} \]

\[ x^i = [x^i_1, x^i_2] = [U_{Xi} \otimes U_{\omega i} \otimes U_{Ri}, \Phi_{Xi} \otimes \Phi_{\omega i} \otimes \Phi_{Ri}] \] \tag{3.48}

Specifically, subscript 1 and 2 indicate association with mechanical and electrical variables, respectively; components associated with \( X, \omega \) and \( R \) are accessed with an additional subscript \( k \) for \( k = 1, 2 \) and 3, respectively. As examples, \( x^i_{11} \) stands for, \( u \) and \( \Phi \), while \( b_{12} \) stands for \( F_\omega \). For individual blocks in (3.45), a superscript \( k \) is added to \( \mathcal{A} \) so that \( \mathcal{A}_{ij}^k \) depicts matrices in the block \( \mathcal{A}_{ij} \) relating to the space, frequency and resistor when \( k \) equals to respectively, 1, 2 and 3 (i, j = 1, 2 refers to \( u \) and \( \varphi \)). In this way, ensemble of linear systems in (2.50) for all parameters can be expressed in a compact form

\[ \mathcal{A} \cdot x^{(r)} := \mathcal{A} \sum_{i=1}^{r} x^i \approx b. \tag{3.49} \]

The goal is to find all modes \( x^i \) in appropriate subspaces through enrichments, with components \( x^m_{\lambda k} \) inside modes calculated using e.g. ADM. Meanwhile, we define subspaces \( \mathcal{S}_{\lambda k} \) so that \( x^m_{\lambda k} \in \mathcal{S}_{\lambda k} \). Examples of such subspaces are \( \mathcal{S}_{11} \subset \mathbb{C}^{n_U} \) and \( \mathcal{S}_{23} \subset \mathbb{C}^{n_\varphi} \). With reference to (3.10), we see that \( x^m \in \mathcal{S}_{\lambda} := \mathcal{S}_{\lambda 1} \times \mathcal{S}_{\lambda 2} \times \mathcal{S}_{\lambda 3} \) while \( x^m \in \mathcal{S} := \{ (\beta_1, \beta_2)^\dagger; \beta_k \in \mathcal{S}_{\lambda}, \lambda = 1,2 \} \).

3.3.2 The enrichment process

We start from the definition based on residual norm, which is introduced in (3.13). For simplicity, we denote \( b_{\lambda} = \mathcal{A}_{\lambda k} \cdot \{ x^{(m-1)} + x^m \} \) by \( r^m_{\lambda} \). The subscript \( \lambda \) of \( \mathcal{A}_{\lambda k} \) represents the vector containing \( \lambda \)-related row (i.e. mechanical or electrical when \( \lambda \) equals 1 and 2, respectively) indicators. The residual norm based definition translates to the expression in (3.50).

\[ x^m = \arg \min_{x^m \in \mathcal{S}_{\lambda}} \| x^m(x^{m^*}) \|_2, \quad \lambda = 1,2 \tag{3.50} \]

The above problem amounts to

\[ \langle \mathcal{A}_{\lambda k} \cdot x^m, x^{m^*} \rangle_{\mathcal{S}_{\lambda}} = \langle t^{m-1}_{\lambda}, x^{m^*} \rangle_{\mathcal{S}_{\lambda}} \forall x^{m^*} \in \mathcal{S}_{\lambda} \tag{3.51} \]
where $\langle \cdot , \cdot \rangle_{\mathcal{S}_\lambda}$ is the inner product defined on $\mathcal{S}_\lambda$, it is calculated as $\prod_{k=1}^{3} \langle \cdot , \cdot \rangle_{\mathcal{S}_{\lambda_k}}$, in which $\langle \ast_k , \ast_k \rangle_{\mathcal{S}_{\lambda_k}}$ is obtained through $(\ast_k^\dagger \cdot \ast_k)$. Technically, we should follow the Minimal Residual-PGD approach that is related to (3.15), to solve the problem in (3.51), because matrices in (3.46) are symmetric and real-valued, which, when multiplied with the imaginary unit (see equations in (2.50)), become non-Hermitian. Nevertheless, we can also take the Galerkin-PGD approach that is related to (3.16). As explained in [68], the latter approach proves efficient to capture good approximations, even if in some non-Hermitian cases. As a result, the problem is equivalent to solving (3.52).

\[
\begin{align*}
\mathbf{x}^{m+1} &= \mathbf{x}^m + \mathbf{m}^{m-1} \\
\mathbf{x}^{m+1} &= \mathbf{x}^m + \mathbf{m}^{m-1}
\end{align*}
\]  

(3.52)

Now ADM can be employed to solve this nonlinear problem. We assume that components except $x^{m}_{\lambda\mu}$ in the $m$-th mode are known when calculating $x^{m}_{\lambda\mu}$, see (3.53). Vectors $x^{m}_{\lambda\mu}$ can be assigned with values as introduced before.

\[
x^{m}_{\lambda} = x^{m}_{\lambda\mu} \otimes \left( \bigotimes_{k \neq \mu} x^{m}_{\lambda k} \right)
\]  

(3.53)

Substituting (3.53) into (3.52) leads to a linear system for $x^{m}_{\lambda\mu}$. We recall that unknowns associated with $X$, $\omega$ and $R$ are calculated when $\mu$ takes value from 1 to 3, while displacements and electrical potentials are calculated when $\lambda$ takes 1 and 2. To control iterations and enrichments, proper maximum counters and convergence tolerances need to be defined. We first denote the iteration counter in ADM procedure by $i$ whose maximum value is $n_{\text{in}}$. Also, we denote the results for $x^{m}_{\lambda\mu}$ at the $i$-th iteration by $x^{m,i}_{\lambda\mu}$. Then, the convergence criterion can be expressed as in (3.54) where $\varepsilon_{\text{in}}$ is prescribed.

\[
\prod_{\mu=1}^{3} \frac{\| x^{m,i}_{\lambda\mu} - x^{m,i-1}_{\lambda\mu} \|_2}{\| x^{m,i-1}_{\lambda\mu} \|_2} < \varepsilon_{\text{in}}, \quad \lambda = 1,2
\]  

(3.54)

After iterations, we can choose any two vectors in the mode $x^{m}_{\lambda}$ to normalize, in order to prevent numerical overflow. For the enrichment process, we also define a maximum counter (denoted by $n_{\text{out}}$). The following convergence criterion can also be used to control the process. In the equation $\varepsilon_{\text{out}}$ denotes the prescribed tolerance.
3.3. A fast algorithm for parametric electro-mechanical problems

\[ \frac{\|x_{\lambda_2}^m\|_2}{\|x_{\lambda_1}^1\|_2} < \varepsilon_{\text{out}}, \quad \lambda = 1,2 \]  

(3.55)

The criterion in (3.55) implies that the newly calculated mode is sufficiently small in front of the first one. In fact, the problem resulting from (3.52) can be viewed as the Nearest Kronecker Product Problem [81], in which the first mode is usually among the most dominant ones.

Table 3.2. Algorithm for parametric electro-mechanical problems

<table>
<thead>
<tr>
<th>No.</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Initialize ( x_{\lambda_0} ) with ( \lambda = 1,2 ); ( \mu = 1,2,3 )</td>
</tr>
<tr>
<td>2</td>
<td>\textbf{for} ( i_{\text{out}} = 1 ) to ( n_{\text{out}} ) \textbf{do}</td>
</tr>
<tr>
<td>3</td>
<td>\textbf{for} ( i_{\text{in}} = 1 ) to ( n_{\text{in}} ) \textbf{do}</td>
</tr>
<tr>
<td>4</td>
<td>Compute ( x_{\lambda_i}^{\text{in}} ) from equations in (3.52)</td>
</tr>
<tr>
<td>5</td>
<td>Check convergence criterions in (3.54)</td>
</tr>
<tr>
<td>6</td>
<td>\textbf{end for}</td>
</tr>
<tr>
<td>7</td>
<td>Normalize ( x_{\lambda_i}^{\text{in}} ) (( \lambda = 1,2 ); ( \mu = 1,2 ))</td>
</tr>
<tr>
<td>8</td>
<td>Update ( x_{\lambda_i} ) using equations in (3.57)</td>
</tr>
<tr>
<td>9</td>
<td>Check convergence criterions in (3.55)</td>
</tr>
<tr>
<td>10</td>
<td>\textbf{end for}</td>
</tr>
<tr>
<td>11</td>
<td>Recover the approximated solution using (3.44)</td>
</tr>
<tr>
<td>12</td>
<td>\textbf{end} of the algorithm</td>
</tr>
</tbody>
</table>

To compare the relative error between the results obtained with conventional and PGD base approaches, we use the equation in (3.56).

\[ \varepsilon_{\text{rel}} = \max_{\lambda=1,2} \frac{\|x_{\lambda}^{\text{pgd}} - x_{\lambda}^{\text{con}}\|_2}{\|x_{\lambda}^{\text{con}}\|_2} \]  

(3.56)

3.3.3 Updating strategy

At the same time, updating plays an important role in enhancing convergence rates. In this
case, it can be achieved through enforcing that \( \mathbf{v}_m^{(\lambda)} \) obtained using the newly calculated mode \( \mathbf{x}_m \) is orthonormal to the subspace \( (\mathcal{S}_\lambda)^m := \mathcal{X}_{\mu=1,2,3}(\mathcal{S}_\lambda^\mu)^m \), see (3.57).

\[
x_\lambda^{l+} \cdot (b_\lambda - \mathcal{G}_\lambda \cdot \mathbf{x}_m^{(l)}) = 0, \quad l = 1,2,\ldots,m
\]  

(3.57)

Again, ADM can be employed to solve (3.57). It consists in solving all \( m \) components \( x_\lambda^{l+} \) while fixing other sets of components \( x_\lambda^{l+} \) for \( \kappa \neq \mu \), and take \( \mu = 1,2,3 \) in turn. However, it is found in our numerical tests that the convergence rate can be extremely slow, especially when \( m \) is large. In other words, increasing iteration numbers has insignificant effects in the current case. Consequently, we only perform iteration once in the updating. Also, when updating space related components, the problem size is square of \( m \times \left( n_X^b + n_X^p \right) \), which quickly goes large as \( m \) increases. Thus, we do not consider updating space related components. The algorithm is summarized and presented in Table 3.2.

### 3.3.4 Numerical application

The novel algorithm is applied to solve the bimorph example in § 2.3.3.1. For performance comparison, model configuration, and samples values of parameters are kept identical as with the conventional approach. Recall that the number of frequency samples is 600, while that for resistors is six. In contrast to the example in § 3.2.4, the problem of interest here is solved with conventional and PGD based approach on the same platform, using MATLAB.

#### Table 3.3. Comparison in performance of different implementations

<table>
<thead>
<tr>
<th></th>
<th>runtime</th>
<th>Solutions size</th>
<th>Relative error</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conventional (sequential)</td>
<td>11228 s</td>
<td>1144 Mb</td>
<td>reference</td>
</tr>
<tr>
<td>Conventional (parallel)</td>
<td>5125 s</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PGD (without updating)</td>
<td>3094 s</td>
<td>30.72 Mb</td>
<td>0.178</td>
</tr>
<tr>
<td>PGD (with updating)</td>
<td>2483 s</td>
<td></td>
<td>8.01 \times 10^{-6}</td>
</tr>
</tbody>
</table>

In the first set of tests, convergence criterions for iterations and enrichments are set to be \( 10^{-3} \) and \( 10^{-5} \), respectively. Corresponding maximum counters are set to 10 and 100. Meanwhile, conventional approaches in both sequential and parallel (based on the built-in
3.3. A fast algorithm for parametric electro-mechanical problems

*parallel mechanism* of MATLAB) are used. For the PGD approach, cases with and without updating are tested. For different implementations, the consumed runtime and memory required for solution storing, as well as relative errors are depicted in Table 4.2.

![Figure 3.5](image)

Figure 3.5. Convergence of enrichment process for PGD approach with and without updating.

We notice that for parallel implementation of the conventional approach, runtime is less than half of that of the serial one, due to the efficient use of processors. On the other hand, both PGD approaches take less runtime than the parallel conventional approach. Specifically, the updated version of PGD approach takes less than half of the latter (namely, less than a quarter of the sequential conventional approach). Difference in runtime can be explained from complexities of respective approaches. In fact, complexity of both versions of conventional approaches is proportional to $n_\omega \times n_R \times \left( n_X^{uv} + n_X^w \right)$, whereas that of both PGD approaches is proportional to $\sum_{i=1}^{n_{out}} \hat{R}_{in}^i \left( n_X^{uv} + n_X^w + 2n_\omega + 2n_R \right)$ in which $\hat{R}_{in}^i$ is the number of iterations undergone during the i-th enrichment. Apparently, the latter can be much smaller than the former, especially when $n_X^{uv}$ and $n_X^w$ are large. The ratio can be observed from the spaces required to store final results (it equals around 40). The time-saving due to updating for PGD approaches can be seen from Figure 3.5, which suggests that updating leads to better approximation with less enrichment.

In Figure 3.6 and Figure 3.7, components related to spatial coordinates and frequencies for the first eight modes are depicted. We see that only bending modes (none torsion nor other higher order modes) appear in the former, which is due to the fact that frequencies chosen here only cover the first two eigen frequencies of the bimorph (similar to the case of Figure 2.4). For Figure 3.7, most peak values locate near the eigen frequencies of the system. In Figure 3.8,
Figure 3.6. Components related to spatial coordinates of the first to eighth enrichment. Displacements are depicted by the (scaled) deformation from original positions, while potentials are represented using the colored region. As in Figure 2.4, area for air is not shown in the figure. Values are normalized.

Figure 3.7. Components related to frequencies of the first to eighth enrichment (Values are normalized). We depict components related to the resistor of 91 kΩ whose counterparts for other resistors show similar shapes.
3.3. A fast algorithm for parametric electro-mechanical problems

In order to investigate influence of different algorithm parameters (namely, \( n_{\text{Out}} \), \( \varepsilon_{\text{Out}} \) for controlling enrichments and \( n_{\text{In}}, \varepsilon_{\text{In}} \) for controlling iterations) in the PGD approach, additional sets of simulations are carried out. Results are listed in different groups and presented in Table 3.4 and Table 3.5, for respectively, iteration and enrichment parameters. Comparison are made in terms of: \( \#_{\text{ite}} \) the total number of iterations, \( \varepsilon_{\text{conv}} \) calculated with (3.55) when testing enrichment convergence, and \( \varepsilon_{\text{rel}} \) obtained using the expression in (3.56).

Table 3.4. Study on parameters of iterations

<table>
<thead>
<tr>
<th>( \varepsilon_{\text{In}} )</th>
<th>( n_{\text{In}} = 5 )</th>
<th>( n_{\text{In}} = 10 )</th>
<th>( n_{\text{In}} = 20 )</th>
<th>( n_{\text{In}} = 30 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( #_{\text{ite}} )</td>
<td>( \varepsilon_{\text{conv}} ) (10^{-5})</td>
<td>( \varepsilon_{\text{rel}} ) (10^{-5})</td>
<td>( #_{\text{ite}} )</td>
<td>( \varepsilon_{\text{conv}} ) (10^{-5})</td>
</tr>
<tr>
<td>( 10^{-2} )</td>
<td>480</td>
<td>2.2</td>
<td>7.2</td>
<td>624</td>
</tr>
<tr>
<td>( 10^{-3} )</td>
<td>495</td>
<td>3.7</td>
<td>7.8</td>
<td>839</td>
</tr>
<tr>
<td>( 10^{-4} )</td>
<td>498</td>
<td>2.0</td>
<td>6.7</td>
<td>937</td>
</tr>
<tr>
<td>( 10^{-5} )</td>
<td>500</td>
<td>52</td>
<td>44</td>
<td>863</td>
</tr>
</tbody>
</table>

In Table 3.4, the maximum enrichment counter is fixed at 100 while enrichment criterion is set to \( 10^{-5} \). It is found that increasing the maximum iteration counter \( n_{\text{In}} \) does not significantly improve accuracy, while complexity increases dramatically. It is more sophisticated with \( \varepsilon_{\text{In}} \); when it is set too large, solutions for step 4 of the algorithm i.e. solving (3.52) can be far from being good approximations, which then requires more enrichments (indeed, enrichments are successive corrections); on the other hand, when it is set too small, results may
get deteriorated, because more iterations are needed for higher precision, while it can go diverging after certain number of iterations. Therefore, the solved modes are not necessarily more accurate, albeit the larger number of iterations.

<table>
<thead>
<tr>
<th>$\varepsilon_{\text{Out}}$</th>
<th>$n_{\text{Out}} = 30$</th>
<th>$n_{\text{Out}} = 50$</th>
<th>$n_{\text{Out}} = 100$</th>
<th>$n_{\text{Out}} = 200$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$10^{-2}$</td>
<td>228</td>
<td>228</td>
<td>228</td>
<td>228</td>
</tr>
<tr>
<td></td>
<td>0.72</td>
<td>7.2</td>
<td>72</td>
<td>72</td>
</tr>
<tr>
<td></td>
<td>$4.8 \times 10^{-2}$</td>
<td>$48 \times 10^{-3}$</td>
<td>$480 \times 10^{-4}$</td>
<td>$480 \times 10^{-5}$</td>
</tr>
<tr>
<td>$10^{-3}$</td>
<td>254</td>
<td>417</td>
<td>623</td>
<td>623</td>
</tr>
<tr>
<td></td>
<td>2.7</td>
<td>2.7</td>
<td>0.92</td>
<td>0.92</td>
</tr>
<tr>
<td></td>
<td>$5.4 \times 10^{-3}$</td>
<td>$1.9 \times 10^{-3}$</td>
<td>$3.6 \times 10^{-4}$</td>
<td>$3.6 \times 10^{-5}$</td>
</tr>
<tr>
<td>$10^{-4}$</td>
<td>254</td>
<td>417</td>
<td>835</td>
<td>948</td>
</tr>
<tr>
<td></td>
<td>5.7</td>
<td>2.7</td>
<td>0.17</td>
<td>0.02</td>
</tr>
<tr>
<td></td>
<td>$5.4 \times 10^{-4}$</td>
<td>$1.9 \times 10^{-4}$</td>
<td>$0.69 \times 10^{-5}$</td>
<td>$0.58 \times 10^{-6}$</td>
</tr>
</tbody>
</table>

In Table 3.5, maximum iteration number is set to 10, iteration convergence criterion is fixed at $10^{-3}$ for all cases. Influences of enrichment parameters are obvious. With more enrichment, it is more likely to arrive at better approximations. Thus, both smaller values of $\varepsilon_{\text{Out}}$ and larger values of $n_{\text{Out}}$ are in favor of accuracy. However, it is also important to keep in mind that the limit of approximation precision is also decided by factors other than parameters being investigated here, such as updating. Therefore, results are no longer improved to some limit by simply increasing the number of modes. In the present case, maximum counter of 100 and tolerance of $10^{-5}$ are found to be appropriate.

### 3.4 Chapter summary

In this chapter, we have seen the potential of complexity reducing with separated representations. Based on the latter, POD and PGD are introduced through an abstract problem. We noticed that the former provides good approximations in certain occasions, while subjecting to the number of variables and requiring calculation of the original problem at selected samples of parameter. PGD, on the other hand, can be applied in higher dimensional problems, and construct solutions approximations a priori. Nevertheless, approximations obtained with PGD are normally less optimal than with POD. Besides, effectiveness of the former depends on separability of specific problems.
Fortunately, the two algorithms that we’ve proposed, based on PGD, prove to be very satisfactory. The first algorithm can be used to significantly reduce problem size of nonlinear magneto-thermal problems. Moreover, when combined with the Picard method, we can solve the strongly-coupled problem in a ‘decoupled’ way, in parallel, thereby, nonlinearity is treated straightforwardly. The second algorithm can be used for large electro-mechanical problems with multiple parameters. In the presented numerical application, our algorithm outperforms the conventional one in terms of runtime and memory requirements, by a large margin, while still delivers highly accurate results (whose relative error is $\sim 10^{-5}$).
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Conclusions and Prospective

In this thesis, a multiphysics finite element framework for piezoelectric and magnetostrictive materials modeling has been presented. We used differential forms based Whitney elements for discretization of quantities of electromagnetic, elastic, and thermal fields. It is shown that the involved fields, as well as coupling effects between these fields, can be conveniently modeled in this framework. Elaborating electrical circuits (including electrodes) into FE systems has also been demonstrated. For constitutive modeling, we adopted the linear piezoelectricity for piezoelectric materials, and the free-energy based the discrete energy-averaged model, or DEAM, for magnetostrictive materials. Due to non-homogeneity of magnetostrictive materials, integration of DEAM with FEM requires that FEM be solved on the macroscopic structure, and DEAM solved on the microscopic structure. To this end, we have chosen FEs as microscopic representative volumes. As a result, FE results were utilized to extract inputs for DEAM, whereas DEAM results were utilized to assemble FE systems. To deal with nonlinearity involved in this hierarchy, the piecewise linear strategy was adopted. Throughout our numerical tests, we observed that FE models were versatile and able to provide more detailed insights into the simulated system than others; however, FE models also demanded more computing resources, especially in cases of transient and parametric analysis. To preserve advantages of our FE models while reduce the price in terms of runtime and memories; we proposed two novel model reduction algorithms, based on the proper generalized decomposition, or PGD method. In these algorithms, solutions of the multiphysics problem were decomposed into components associated with different parameters, through which the original large-size problem could be converted to a multitude of much smaller-size sub-problems. We presented through a couple of examples that our algorithms were able to significantly reduce problem complexity, while generating very high quality solutions at the same time. To sum up, we have achieved the objectives listed at the beginning of the thesis: 1) building a 3D FE framework to model piezoelectric and magnetostrictive materials involved multiphysics fields and coupling effects; 2) elaborating state-of-the-art constitutive models for better description of materials behaviors; and 3) performing model reduction so as to carry out 3D FE simulations efficiently.

In the meanwhile, we also realized that the work presented in this thesis could be extended in some possible orientations. For instance, we noticed that when elastic problems of thin structures involved (for both piezoelectric and magnetostrictive materials), it becomes expensive,
Conclusions and Prospective

sometimes problematic to obtain accurate results using our first order elements, due to the well-known shear locking phenomenon. Therefore, it would be helpful to add in higher order elements for such kinds of problems. Also, shell elements can also be considered in such cases. Second, although several applications for energy harvesting have been accomplished (and presented) during the thesis, extra efforts are necessary in order to leverage the potential of the developed framework for simulations of devices that are under investigation within our research group. Take the DEAM model for example. Running simulation using the latter requires materials specific parameters, which are available for Galfenol and Terfenol-D while currently not for Metglas (per the author’s knowledge). Consequently, additional measurements and some fine tuning based on the present work may be needed. Third, for uncertainty quantification it is becoming a trend to perform stochastic simulations for piezoelectric and magnetostrictive materials involved systems, so that a broader range of realistic factors can be accounted for. In this case, our multiphysics models can take the role of modeling the deterministic part of the problem, whereas our PGD based algorithms can serve to carry out model reductions in problems related to uncertainties. Last but not the least important, when nonlinearities in general cases (such as those in DEAM) involved, it becomes questionable for the separability of applying PGD. Therefore, it would be beneficial to investigate novel methodologies addressing such issue, so that our algorithms can be applied to more general nonlinear problems.
Appendix A

§ A.1 Formulations for barycentric coordinates in tetrahedron elements

Assume a tetrahedron $P_1P_2P_3P_4$, as shown in Figure A.1. A point $P$ at $(x,y,z)$ is located within the tetrahedron. $P_1P_2P_3P_4$ is denoted by $\mathcal{K}$ for simplicity. The barycentric coordinates at $P$ are denoted by $\lambda_i$ with the arrows indicating its decreasing direction (i.e. directions approaching the surfaces of the tetrahedron). Meanwhile, vectors $r_j$ on the edges are oriented as pointing from smaller numbered vertex to larger numbered vertex.

To express $\lambda_i$ in terms of the coordinates, consider the following determinants:

\[
u_0 = \begin{vmatrix} x_1 & y_1 & z_1 & 1 \\ x_2 & y_2 & z_2 & 1 \\ x_3 & y_3 & z_3 & 1 \\ x_4 & y_4 & z_4 & 1 \end{vmatrix}, \quad \nu_1 = \begin{vmatrix} x & y & z & 1 \\ x_2 & y_2 & z_2 & 1 \\ x_3 & y_3 & z_3 & 1 \\ x_4 & y_4 & z_4 & 1 \end{vmatrix}, \quad \nu_2 = \begin{vmatrix} x_1 & y_1 & z_1 & 1 \\ x_2 & y_2 & z_2 & 1 \\ x_3 & y_3 & z_3 & 1 \\ x_4 & y_4 & z_4 & 1 \end{vmatrix}, \quad \nu_3 \text{ and } \nu_4 = \begin{vmatrix} x_1 & y_1 & z_1 & 1 \\ x & y & z & 1 \\ x_2 & y_2 & z_2 & 1 \\ x_3 & y_3 & z_3 & 1 \\ x_4 & y_4 & z_4 & 1 \end{vmatrix}.
\]

Based on the linear interpolating relation, it is easy to verify that $\lambda_i$ equals $\nu_i/\nu_0$. Hence, a scalar field $u$ that takes the value of $u_k$ at vertex $P_k$ ($k = 1,2,3,4$) can be expressed as $\sum_{i=1}^{4} \lambda_i u_i$ at $P$. In fact, the first determinant $\nu_0$ is the determinant of the so-called Jacobian,
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denoted by \( J \) as in Equation (A.1), which can also be obtained from, \textit{e.g.} \((r_1 \times r_2) \cdot r_3\). The latter is six times of the volume of the tetrahedron \((v_0 = 6|\mathcal{K}|)\).

\[
J = \begin{bmatrix} x_1 & y_1 & z_1 & 1 \\ x_2 & y_2 & z_2 & 1 \\ x_3 & y_3 & z_3 & 1 \\ x_4 & y_4 & z_4 & 1 \end{bmatrix}
\]  
(A.1)

As the barycentric coordinates are first order function, their gradients w.r.t. coordinates \(x, y\), and \(z\) are constant over \(\mathcal{K}\). These constants are shown in Equation (A.2).

\[
\begin{align*}
\text{grad}(\lambda_1) &= r_4 \times r_5 / 6v_0, \\
\text{grad}(\lambda_2) &= r_2 \times r_3 / 6v_0, \\
\text{grad}(\lambda_3) &= r_3 \times r_1 / 6v_0, \\
\text{grad}(\lambda_4) &= r_1 \times r_2 / 6v_0
\end{align*}
\]  
(A.2)

Since all basis function of Whitney elements are calculated via the barycentric coordinates, the above relations allow for a convenient implementation.

\section*{§ A.2 Formulations relating the engineering strain and displacement}

By definition, the engineering strain tensor \( \varepsilon \) within an element \( \mathcal{K} \) is expressed as in Equation (A.3) where \((u, v, w)\) represents the displacement of a particle located on the point \((x, y, z)\).

\[
\varepsilon = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \varepsilon_3 \end{bmatrix} = \begin{bmatrix} \frac{\partial u}{\partial x} \\ \frac{\partial v}{\partial y} \\ \frac{\partial w}{\partial z} \end{bmatrix}
\]  
(A.3)

Denote DoFs on \(\mathcal{K}\) – displacements at the \(i_{th}\) vertex for \(i = 1, 2, 3, 4\) by \((u_i, v_i, w_i)\), and interpolate \((u, v, w)\) via \(a = \Sigma_{i=1}^{4} \lambda_i a_i\) for \(a = u, v, w\).

Then, insert the interpolants into Equation (A.3) and re-arrange DoFs in a vector as \([u_1, v_1, w_1, u_2, v_2, w_2, u_3, v_3, w_3, u_4, v_4, w_4]^T\). As such, \( \varepsilon \) can be re-written as
Note that nonzero entries in the central large matrix (denoted as $B_u$) can be divided into three groups, as marked in different color. Let $B_{u,i,:}$ and $B_{u,:,j}$ represent respectively, the $i$th line and $j$th column of $B_u$. It can be verified that $B_u^{[1,4,6],[1,4,7,10]}$, $B_u^{[4,2,5],[2,5,8,11]}$, $B_u^{[6,5,3],[3,6,9,12]}$ all refer to the same matrix:

$$
\begin{bmatrix}
\frac{\partial \lambda_1}{\partial x} & \frac{\partial \lambda_2}{\partial x} & \frac{\partial \lambda_2}{\partial y} & \frac{\partial \lambda_3}{\partial x} & \frac{\partial \lambda_3}{\partial y} & \frac{\partial \lambda_4}{\partial x} & \frac{\partial \lambda_4}{\partial y} \\
\frac{\partial \lambda_1}{\partial y} & \frac{\partial \lambda_2}{\partial y} & \frac{\partial \lambda_2}{\partial z} & \frac{\partial \lambda_3}{\partial y} & \frac{\partial \lambda_3}{\partial z} & \frac{\partial \lambda_4}{\partial y} & \frac{\partial \lambda_4}{\partial z} \\
\frac{\partial \lambda_1}{\partial z} & \frac{\partial \lambda_2}{\partial z} & \frac{\partial \lambda_2}{\partial x} & \frac{\partial \lambda_3}{\partial z} & \frac{\partial \lambda_3}{\partial x} & \frac{\partial \lambda_4}{\partial z} & \frac{\partial \lambda_4}{\partial x} \\
\end{bmatrix}
$$

This property of the elastic gradient matrix allows for rapid implementing of the elastic stiffness matrix, as the above matrix is very common for FE problems of the scalar field, and can be readily obtained with equation in § A.1. Therefore, while elastic DoFs are vector valued (i.e. three DoFs per vertex) the elastic stiffness matrix can be built based on those of scalar field problems in a straightforward manner.

§ A.3 Calculation of the incident matrices

Denote the number of tetrahedrons, triangles, edges, and nodes in the concerned triangulation by respectively, $n_t$, $n_f$, $n_e$, and $n_n$. Dimensions of the incident matrices $D$, $C$, and $G$ can thus be expressed as $n_t \times n_f$, $n_f \times n_e$, and $n_e \times n_n$, respectively. Meanwhile, $D$, $C$, and $G$ are composed of values $\pm 1$ and 0. Take $C_{ij}$ for example, $C_{ij} = 0$ means the $j$th edge $E^j$ does not locate within the $i$th facet $F^i$. Otherwise, $E^j$ is one of the constituting edges of $F^i$. Furthermore, $C_{ij} = +1$ implies the local orientation of $E^j$ is consistent with the global orientation, whereas $C_{ij} = -1$ implies inconsistent.
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Figure A.2 Illustration of incident relations for (a) tetrahedron to facets
\( T_{\alpha_1\alpha_2\alpha_3\alpha_4} \rightarrow \{ F_{\alpha_2\alpha_3\alpha_4}, F_{\alpha_1\alpha_2\alpha_3}, F_{\alpha_1\alpha_2\alpha_4}, F_{\alpha_1\alpha_3\alpha_4} \} \), (b) facet to edges \( F_{\beta_1\beta_2\beta_3} \rightarrow \{ E_{\beta_2\beta_3}, E_{\beta_3\beta_1}, E_{\beta_1\beta_2} \} \), and (c) edge to nodes \( E_{\gamma_1\gamma_2} \rightarrow \{ N_{\gamma_2}', N_{\gamma_1}' \} \).

Denote an \( n \)-dimensional geometric object using its \( n + 1 \) vertices. For instance, \( T_{\alpha_1\alpha_2\alpha_3\alpha_4} \) represents a tetrahedron (consisting vertices \( \alpha_1, \alpha_2, \alpha_3 \) and \( \alpha_4 \)); \( F_{\alpha_1\alpha_2\alpha_3} \) represents a triangle (consisting vertices \( \alpha_2, \alpha_3 \) and \( \alpha_4 \)); \( E_{\beta_2\beta_3} \) represents an edge (consisting vertices \( \beta_2 \) and \( \beta_3 \)); whereas \( N_{\gamma_1} \) represents the node (numbered by \( \gamma_1 \)). To uniquely identify an object, it is assumed that numberings of vertices are arranged in ascend order. Namely, \( \alpha_1 < \alpha_2 < \alpha_3 < \alpha_4 \) in \( T_{\alpha_1\alpha_2\alpha_3\alpha_4} \), and \( \beta_2 < \beta_3 \) in \( E_{\beta_2\beta_3} \), and so forth. This is how global identities of geometric objects are defined. The local identities, on the other hand, indicate how \( n \)-dimensional sub-objects are identified in higher dimensional objects comprising them. Here, the induced ordering is used, which is depicted in Figure A.2. Specifically, \( T_{\alpha_1\alpha_2\alpha_3\alpha_4} \) induces four facets \( F_{\alpha_2\alpha_3\alpha_4}, F_{\alpha_1\alpha_2\alpha_3}, F_{\alpha_1\alpha_2\alpha_4}, F_{\alpha_1\alpha_3\alpha_4} \); \( E_{\beta_1\beta_2\beta_3} \) induces three edges \( E_{\beta_2\beta_3}, E_{\beta_3\beta_1}, E_{\beta_1\beta_2} \); and \( E_{\gamma_1\gamma_2} \) induces two nodes \( N_{\gamma_2}', N_{\gamma_1}' \).

Suppose that the global and local identities are called consistent if the vertex numbering of the latter is an even permutation of the former. Then, the induced local identities \( F_{\alpha_2\alpha_3\alpha_4}, F_{\alpha_1\alpha_2\alpha_4}, E_{\beta_2\beta_3}, E_{\beta_1\beta_2} \) are consistent with their global counterparts, whereas \( F_{\alpha_1\alpha_4\alpha_3}, F_{\alpha_1\alpha_3\alpha_2}, E_{\beta_3\beta_1} \) not. Given an incident matrix whose entries are initially all zero, entries are changed to +1 in the case of consistency, and -1 for inconsistency. For \( G \) that links edges with
vertices, the second vertex of an edge is assigned with -1, whereas the first with +1. Note that most entries are zero in $D$, $C$, and $G$. As a result, the incident matrices are better stored as sparse matrices; namely only nonzero entries are stored.
Appendix B

§ B.1 Definitions of terms in the systems of Equation (3.30) of Chapter 3

\[
C_A(\Theta) := \sum_{\alpha=1}^{n_X} \int_{\mathcal{K}_\alpha} \frac{c_{e_A}}{r_\alpha} W_a^{n,2D} \cdot W_a^{n,2D,\dagger} \, d\Omega \\
C_\theta := \sum_{\alpha=1}^{n_X} \int_{\mathcal{K}_\alpha} \rho c_p W_a^{n,2D} \cdot W_a^{n,2D,\dagger} \, d\Omega
\]

\[
K_A := \sum_{\alpha=1}^{n_X} \int_{\mathcal{K}_\alpha} \frac{1}{m_\alpha} (\text{grad} \ W_a^{n,2D}) \cdot (\text{grad} \ W_a^{n,2D,\dagger}) \, d\Omega
\]

\[
K_\theta := \sum_{\alpha=1}^{n_X} \int_{\mathcal{K}_\alpha} \kappa (\text{grad} \ W_a^{n,2D}) \cdot (\text{grad} \ W_a^{n,2D,\dagger}) \, d\Omega
\]

\[
J := \sum_{\alpha=1}^{n_X} \left( \int_{\mathcal{K}_\alpha} W_a^{n,2D} \cdot W_a^{n,2D,\dagger} \, d\Omega \right) J_i
\]

\[
S(\Theta, \hat{A}) := \sum_{\alpha=1}^{n_X} \int_{\mathcal{K}_\alpha} \frac{\sigma_{e_A}}{r_\alpha} \left( \frac{\partial \hat{A}}{\partial t} \right)^2 \cdot (W_a^{n,2D} \cdot W_a^{n,2D,\dagger}) \, d\Omega
\]

§ B.2 Discrete weak forms associated with Equations (3.35) and (3.36)

Note that in an additional subscript ‘h’ is added to variables \( \hat{A}^{(m-1)} \) and \( \theta^{(m-1)} \), to make them read respectively, \( \hat{A}_h^{(m-1)} \) and \( \theta_h^{(m-1)} \), implying corresponding variables in (B.2) are discrete values related to triangulations (characterized by ‘h’).

\[
\sum_{\alpha=1}^{n_X} \sum_{\beta=1}^{n_T} \int_{\mathcal{K}_\alpha \otimes \mathcal{K}_\beta} \begin{pmatrix}
\frac{g(\hat{A}_h^{(m-1)})}{r_\alpha} \frac{\partial}{\partial t} (\hat{A}_h^{(m-1)} + \hat{A}_x^m \hat{A}_t^m) \\
\frac{\partial^2 (\hat{A}_h^{(m-1)} + \hat{A}_x^m \hat{A}_t^m)}{\partial r^2} \\
\frac{\partial^2 (\hat{A}_h^{(m-1)} + \hat{A}_x^m \hat{A}_t^m)}{\partial z^2} \\
- \sum_{i=1}^{r_j} f X_i j t_{ci}
\end{pmatrix} d\Omega dI = 0 \quad (B.2a)
\]
\[ \sum_{a=1}^{n_X} \sum_{\beta=1}^{n_T} \int_{J_{a}} \int_{J_{\beta}} \left( \Theta^{m,\alpha}_{h} \Theta^{m,\alpha}_{c} + \Theta^{m,\alpha}_{h} \Theta^{m,\alpha}_{c} \right) \left\{ \frac{\partial}{\partial t} \left( \Theta^{(m-1)}_{h} + \Theta^{X,m}_{h} \Theta^{m,\alpha}_{c} \right) \right\} d\Omega dI = 0 \]  

(B.2b)

§ B.3 Coefficient matrices obtained through substituting (3.37) into (B.2)

\[ C_{A}^{X} := \sum_{a=1}^{n_X} \int_{J_{a}} \frac{\sigma_{c} a}{r_{a}} W_{x}^{n,2D} \cdot W_{x}^{n,2D} d\Omega \]

\[ K_{A}^{X} := \sum_{a=1}^{n_X} \int_{J_{a}} \frac{1}{r_{a}} \left( \text{grad} W_{x}^{n,2D} \right) \cdot \left( \text{grad} W_{x}^{n,2D} \right) d\Omega \]

\[ C_{\beta}^{X} := \sum_{a=1}^{n_X} \int_{J_{a}} \rho c_{p} W_{x}^{n,2D} \cdot W_{x}^{n,2D} d\Omega \]

\[ K_{\beta}^{X} := \sum_{a=1}^{n_X} \int_{J_{a}} \kappa \left( \text{grad} W_{x}^{n,2D} \right) \cdot \left( \text{grad} W_{x}^{n,2D} \right) d\Omega \]

\[ M_{A}^{t} := \sum_{\beta=1}^{n_T} \int_{J_{\beta}} W_{\beta}^{n,1D} \cdot W_{\beta}^{n,1D} dI \]

\[ K_{A}^{t} := \sum_{\beta=1}^{n_T} \int_{J_{\beta}} \frac{\partial W_{\beta}^{n,1D}}{\partial t} dI \]

\[ M_{\beta}^{t} := \sum_{\beta=1}^{n_T} \int_{J_{\beta}} W_{\beta}^{n,1D} \cdot W_{\beta}^{n,1D} dI \]

\[ K_{\beta}^{t} := \sum_{\beta=1}^{n_T} \int_{J_{\beta}} \frac{\partial W_{\beta}^{n,1D}}{\partial t} dI \]

It is worth noting that due to the consistency of time discretization, time basis functions in \( K_{A}^{t} \) and \( K_{\beta}^{t} \) are changed from \( W_{\beta}^{n,1D} \) to \( \tilde{W}_{\beta}^{n,1D} \), which equals \( W_{\beta}^{n,1D} + 0.5|j_{\beta}| \cdot \left( \frac{\partial W_{\beta}^{n,1D}}{\partial t} \right) \) (i.e. the so-called upwind Petrov-Galerkin formulation). \(|j_{\beta}|\) is the length of time element \( J_{\beta} \).

§ B.4 Settings for the numerical model in §3.2.4 (inductive heating)

The model consists of a conductor that is axisymmetric to the z-axis, an excitation coil, and
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an air boxed (of $0.2 \, m \times 0.5 \, m$). Geometric dimensions related to the conductor and coil are depicted in Figure B.1.a. Linear triangular elements are used to discretize the space domain. Size of elements near the conductor skin is set to be relatively small, so as to account for the skin effect. A total of 5659 elements with 2900 nodes are generated.

Figure B.1 Model configuration (a) geometric dimensions, and (b) spatial triangulation
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