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DAMAGE AND MATERIAL IDENTIFICATION USING INVERSE ANALYSIS

by

Li Li

A Dissertation
Submitted to the Faculty of Graduate Studies
through Department of Civil and Environmental Engineering
in Partial Fulfillment of the Requirements for
the Degree of Doctor of Philosophy
at the University of Windsor

Windsor, Ontario, Canada

2012

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DAMAGE AND MATERIAL IDENTIFICATION USING INVERSE ANALYSIS

BY

Li Li

APPROVED BY:

__________________________
R. Ben Mrad, External Examiner
University of Toronto

__________________________
N. Zamani
Department of Mechanical, Automotive & Materials Engineering

__________________________
B. Budkowska
Department of Civil & Environmental Engineering

__________________________
S. Cheng
Department of Civil and Environmental Engineering

__________________________
F. Ghrib, Advisor
Department of Civil and Environmental Engineering

__________________________
J. Albanese, Chair of Defense
Department of Sociology, Anthropology & Criminology

September 11, 2012
DECLARATION OF ORIGINALITY

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ABSTRACT

In this thesis, we formulate novel solutions to two inverse problems using optical measurements as input data: i) local level damage identification of beams, and ii) material constitutive parameter identification using digital image correlation measurement of surface strain/displacements.

A novel photogrammetric procedure based on edge-detection was devised to measure the quasi-continuous deflection of beams under given loading. This method is based on the close-range photogrammetry technique made possible through recent developments of image processing algorithms and modern digital cameras.

Two computational procedures to reconstruct the stiffness distribution and to detect damage in Euler-Bernoulli beams are developed in this thesis. The first formulation is based on the principle of the equilibrium gap along with a finite element discretization. The solution is obtained by minimizing a regularized functional using a Tikhonov Total Variation (TTV) scheme. The second proposed formulation is a minimization of a data discrepancy functional between measured and model-based deflections. The optimal solution is obtained using a gradient-based minimization algorithm and the adjoint method to calculate the Jacobian. The proposed identification methodology is validated using experimental data. The proposed methodology has the potential to be used for long term health monitoring and damage assessment of civil engineering structures.

The identification of material plasticity parameters is carried out by minimizing a least-square functional measuring the gap between inhomogeneous displacement fields obtained from measurements and finite element simulations. The material parameters are
identified simultaneously by means of direct, derivative-free optimization methods where the finite element simulation is treated as a black-box procedure. Methods verifying and validating the identified results are given. Particular interest is given to the identifiability issue in deterministic and statistical sense. The validation procedure intends to detect false positive results (type-II errors). The performance of the computational procedures is illustrated by numerical and experimental examples. The proposed approach avoids using the gradient of the cost function in the identification process; it has the benefit of allowing the use of any finite element code as a black box to solve the direct problem.
ACKNOWLEDGEMENTS

First, I would like to express my sincere gratitude to my advisor Dr. F. Ghrib for his patient instructions, insightful comments and financial assistance during the research program. I also would like to thank my committee members, Dr. Zamani, Dr. Budkowska, and Dr. Cheng, for their valuable suggestions to improve this thesis. I owe much thanks to Dr. W. Altenhof, Dr. H.K. Kwan, for their teaching during my training at the University of Windsor. My deep appreciation and gratitude goes to Dr. Green of the department of Mechanical Engineering for providing me the ARAMIS optical testing system. I would like to thank the technical staff in the Civil and Mechanical Engineering Laboratories, to their assistance during the test programs. I would also like to recognize individually, friends and staff in the University of Windsor: Catherine Wilson, Lucian Pop, Pat Seguin, Andrew Jenner, Matt St. Louis, Wafa Polies; their help and support during the test programs are greatly appreciated.
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<tbody>
<tr>
<td>CONDOR</td>
<td>CONstrained, Non-linear, Direct, parallel Optimization using trust Region method for high-computing load function</td>
</tr>
<tr>
<td>DFO</td>
<td>Derivative-Free Optimization</td>
</tr>
<tr>
<td>DIC</td>
<td>Digital Image Correlation</td>
</tr>
<tr>
<td>DOF</td>
<td>Degree Of Freedom</td>
</tr>
<tr>
<td>FEM</td>
<td>Finite Element Method</td>
</tr>
<tr>
<td>FRF</td>
<td>Frequency Response Function</td>
</tr>
<tr>
<td>LCA</td>
<td>Linearized Covariance Analysis</td>
</tr>
<tr>
<td>LS</td>
<td>Least-Squares</td>
</tr>
<tr>
<td>RSM</td>
<td>Response Surface Methodology</td>
</tr>
<tr>
<td>TV</td>
<td>Tikhonov-Total Variation</td>
</tr>
<tr>
<td>VFM</td>
<td>Virtual Fields Method</td>
</tr>
<tr>
<td>V&amp;V</td>
<td>Verification and Validation</td>
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<table>
<thead>
<tr>
<th>Symbols</th>
<th>Definitions</th>
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<tbody>
<tr>
<td>B</td>
<td>The left Cauchy-Green tensor</td>
</tr>
<tr>
<td>C</td>
<td>The right Cauchy-Green tensor</td>
</tr>
<tr>
<td>d</td>
<td>Test data set</td>
</tr>
<tr>
<td>E</td>
<td>Modulus of elasticity</td>
</tr>
<tr>
<td>f</td>
<td>Force vector in FEA</td>
</tr>
<tr>
<td>f(x)</td>
<td>A function</td>
</tr>
<tr>
<td>F</td>
<td>Deformation gradient</td>
</tr>
<tr>
<td>G</td>
<td>System matrix in the equilibrium gap method</td>
</tr>
<tr>
<td>I_M</td>
<td>The relative offset measure</td>
</tr>
<tr>
<td>J</td>
<td>Objective function to be minimized</td>
</tr>
<tr>
<td>J_δf(x)</td>
<td>Mollifier of function f(x)</td>
</tr>
<tr>
<td>J(θ)</td>
<td>Jacobian evaluated at θ</td>
</tr>
<tr>
<td>K, k_i</td>
<td>Stiffness matrix and element stiffness matrix in FEA</td>
</tr>
<tr>
<td>P</td>
<td>Covariance matrix</td>
</tr>
<tr>
<td>T</td>
<td>The first Piola-Kirchhoff stress tensor</td>
</tr>
<tr>
<td>u</td>
<td>Displacement vector</td>
</tr>
<tr>
<td>v</td>
<td>Deflection of a beam</td>
</tr>
<tr>
<td>W</td>
<td>The strain energy function</td>
</tr>
<tr>
<td>x</td>
<td>Coordinates</td>
</tr>
<tr>
<td>α</td>
<td>Tikhonov regularization parameter</td>
</tr>
<tr>
<td>δ</td>
<td>The mollifier’s radius</td>
</tr>
<tr>
<td>Φ(θ)</td>
<td>The regularization function</td>
</tr>
<tr>
<td>λ</td>
<td>Lagrangian multipliers</td>
</tr>
<tr>
<td>Symbol</td>
<td>Description</td>
</tr>
<tr>
<td>--------</td>
<td>-------------------------------------------------</td>
</tr>
<tr>
<td>$\theta$</td>
<td>Rotation angle of a beam</td>
</tr>
<tr>
<td>$\theta, \theta_i$</td>
<td>Parameter vector and its components</td>
</tr>
<tr>
<td>$\Omega$</td>
<td>A region or a subset</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>Cauchy stress</td>
</tr>
<tr>
<td>$\Lambda$</td>
<td>Finite element assembly operator</td>
</tr>
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CHAPTER ONE: INTRODUCTION

“The mere formulation of a problem is far more essential than its solution, which may be merely a matter of mathematical or experimental skills. To raise new questions, new possibilities, to regard old problems from a new angle require creative imagination and marks real advances in science.”

Albert Einstein

1.1 General

Traditional techniques in experimental mechanics rely on displacement or strain transducers carefully placed in a small number of positions on the surface of the tested specimen. The data usually consist of series of test data correlated with the applied load and the measured field (usually a displacement or strain component). The recent development of new technologies combining low-cost CCD cameras and computer vision led to novel experimental methods to assess solids and structures. Non-contact measurement techniques are now becoming affordable for research and development purposes in laboratories as well as for on-site monitoring of structures.

Digital image correlation (DIC), moiré and speckle interferometry, and grid methods are among the many new technologies that can be utilized in the field of experimental mechanics. The fundamental shift in the possibilities offered by these new technologies is related to the spatial properties of the collected data. For example, a typical DIC test system commonly has the capability of collecting up to 10,000 independent displacement measurements from the surface of the specimen. The large amount of information available to the experimentalist opens new horizons that would
not be possible using traditional transducers. For example, in the context of constitutive laws identification, the geometry and the boundary conditions must be very simple when traditional transducers are used, so that an analytical solution of the stress/strain field is possible; the stress and strain fields must be homogeneous in order to determine the material properties. Consequently, the identification of constitutive laws requires more than one test, since only the mechanical properties associated with the mechanisms activated during the test can be identified. When non-contact measurement techniques are used along with the finite element method (FEM), however, the need for simplified loading and boundary conditions is relaxed. Non-contact measurement techniques with a single test set-up can replace the large number of test set-ups involving various combinations of experimental parameters that are necessary when local transducers are used.

Full-field displacement measurements allow much greater flexibility as they provide very rich experimental data and allow the use of tests under non-homogeneous conditions. However, the large amount of data produced by full-field measurement techniques requires suitable computational methods to extract the information encapsulated within the non-homogeneous displacement and strain fields. The effort to integrate new measurement technologies in experimental mechanics has introduced a completely new research area, referred to as inverse problems in solid mechanics.

Full-field displacement measurements can also be integrated with numerical simulation techniques such as FEM to build health monitoring systems for structures in service. The methodology consists of incorporating the measured displacement or strain field parameters with the mathematical model of the structure in order to estimate the
location and severity of damage. The formulation of a health monitoring problem as an inverse problem has become a very important subject that has attracted a great deal of interest and research activity over the last decade.

This thesis concerns two aspects of research based on the use of full-field measurement techniques and finite element simulations for the health monitoring of structures and for constitutive law identification. In the first part of this work, we formulate the problem of reconstruction of the stiffness of beams through the measurement of displacements along the beam. In the second part of this thesis, we study the implementation of this material identification problem using DIC to measure the displacement and strain on the surface of a specimen.

Each of these issues is of the utmost importance in practice. Currently, there is an increasing demand for effective methods to identify complex nonlinear material models. The competitive industrial environment leads engineers to develop better, but also more complex models of complicated systems. These models are generally difficult to define, and the work presented here is a step towards the development of a systematic approach for material characterisation. Furthermore, the problem of reconstruction of a beam’s stiffness can be considered as one of the fundamental problems in the health monitoring of these important structural elements.

From a mathematical point of view, the problems mentioned above can be considered as two examples of inverse problems. However, the stiffness reconstruction of beams is typically classified as an ill-posed functional identification problem, whereas the identification of material properties is considered a well-posed parameter identification problem. Traditional problems in structural mechanics consist of
evaluating the response of the studied structure to a known force or displacement where the basic parameters, such as the geometry of the structure and the mechanical properties of the materials, are known. Inverse problems, on the other hand, seek to determine unknown geometrical and/or mechanical parameters from the known measured responses of the system.

A direct procedure can rarely compute material and geometric properties from measurable responses, such as displacements or strains. Inverse methods usually formulate the problem as a minimization of error function between measured and computed responses; this is also called an estimation function. These inverse procedures usually involve finite-element (FE) model-updating techniques.

1.2 Optical full-field measurements using digital cameras

The damage identification methods developed in this thesis require a technique to measure the displacements during static loading tests. In optical measurements, objects are measured without being touched. A novel procedure based on edge-detection was devised to measure the quasi-continuous deflection of beams under given loading, and is presented in Chapter 2. This method is based on the close-range photogrammetry technique made possible through recent developments of image processing algorithms and modern digital cameras. These studies demonstrate that modern consumer cameras can be used in optical measurement procedures, offering an additional advantage in terms of cost.

The studies of material parameter identification presented in this thesis employ DIC to provide full-field displacement/strain measurements for the surface of complex
3D components as input for the inverse analysis. DIC is an image analysis technique based on grey-value digital images that can determine the contour and the displacements of an object under load in three dimensions with sub-pixel precision. This technique is already mature and commercialized; Chapter 2 provides a brief description of the DIC technique, and the commercial DIC system employed in the current work.

1.3 Damage identification based on static tests

The problem of stiffness reconstruction is presented in Chapter 3. In this chapter, we are particularly interested in formulating the identification of damage from static measurement. Static tests are the most direct way of evaluating the load-bearing capacity of structures. All structures throughout their service undergo a continual and ongoing accumulation of damage and decreased loading capacity. Consequently, damage identification is an important aspect of safety and functionality, and has attracted intensive research efforts over the past twenty years.

Two computational procedures using static deflection measurements to reconstruct the stiffness distribution, and to detect regions containing damages in Euler-Bernoulli beams, are presented and compared in Chapter 3. The first proposed formulation is based on the principle of the equilibrium gap along with a FE discretization, and the mathematical problem leads to an over-determinate linear system. The solution is obtained by minimizing a regularized function with a Tikhonov Total Variation (TTV) regularization scheme. The second proposed formulation is defined as a minimization of a data discrepancy functional between measured and model-based deflections. The optimal solution is obtained using a gradient-based minimization
algorithm and the adjoint method to calculate the Jacobian. The two proposed methodologies are validated using experimental data.

1.4 Material parameter identification using full-field measurements

Modern design and performance evaluation requires realistic simulations of structural and material behaviour, and nonlinear FE simulation has become a fundamental engineering tool. FE approaches are increasingly popular among engineers from different industries. Constitutive parameters associated with nonlinear models are not always available in standard material databases; therefore, engineers need to identify them experimentally.

Although the identification of plasticity material parameters is the focus of the current work (presented in Chapter 4), this study aims to provide a unified identification methodology. In light of recent developments in direct optimization and regression analysis, particularly the verification and validation of the material parameter identification using statistical tools, this thesis proposes novel procedures for the identification of material parameters in any given model.

The problem of material parameter identification is formulated as a nonlinear regression problem using DIC measurement data as the input. The study presented in Chapter 4 includes the following topics: definition of the inverse problem; a brief description of the derivative-free optimization scheme; discussion of the identifiability issues related to the inverse approach; and the application of statistical inference with a non-dimensional measure of response fit.
Solution of the material parameter identification problem was performed by minimizing a least-squares (LS) function measuring the gap between inhomogeneous displacement fields obtained from DIC measurements and FE simulations. Specifically, a LS formulation of a cost function consisting of a norm for the discrepancies between the experimental data and the simulated data was minimized; the simulated data was obtained by the FE method using commercial software. Direct, derivative-free optimization methods, which treat the FE simulation as a black-box procedure, identified the material parameters simultaneously. Particular attention was paid to the identifiability and numerical stability issues of this approach, and methods for verifying and validating the identified results are discussed. In particular, consideration is given to the identifiability issue in the deterministic and the statistical sense. Sampling-based statistical inference derived from nonlinear regression theory was adopted to quantify the quality of the identification procedure, the rationale being that DIC provides a large amount of data, and thus allows the use of statistical inference. Sampling-based inference and sensitivity analysis were used to check the adequacy of the identification solution, thus avoiding Type-2 errors (i.e., the acceptance of incorrect results). Several recommended numerical approaches for validating the identification results are presented. Linearized covariance analysis (LCA) was adopted to check the fit of the parameters. An index derived from the response surface geometry was used to check the response fit, and relative curvature measures proposed to check the adequacy of LCA. The proposed validation procedure is primarily intended to detect false positives (i.e., Type-II errors).

The proposed approach avoids using the gradient of the cost function in the identification process, and has the benefit of allowing the use of any FE code as a black
box to solve the direct problem. The quantification of the quality of the result is of paramount importance to the practical application of any identification methodology.

1.5 Hyperelasticity model parameter identification for rubber and rubber-like solids

The mechanical behaviour of rubber-like materials is usually characterized by a strain energy density function $W$; the parameters of the $W$ function may be considered as material parameters. Traditional laboratory techniques require several tests — using different homogeneous deformation modes and standard cut-out samples — to determine the appropriate strain energy form and parameters.

The general approach for material parameter identification (described in Chapter 4) was used to determine the parameters of strain energy functions ($W$) with DIC test data collected for the original structural components. In this approach, whole field displacement/strain on the surface of components is measured by a DIC-based technique, which provides massive amounts of experimental data in a single test. The DIC-based methodology for parameter identification in hyperelasticity models (presented in Chapter 5) replaces the performance of multiple tests on simple-shape specimens with supposed uniform state variable distributions that are required by traditional techniques.
CHAPTER TWO: OPTICAL MEASUREMENT USING DIGITAL CAMERAS: DIGITAL IMAGE CORRELATION AND CLOSE-RANGE PHOTOGRAMMETRY

“It is necessary to measure everything that can be measured and to try making measurable what isn’t as yet.” Galileo

2.1 Introduction

The technique of obtaining information from photographs is called photogrammetry. The most important feature of photogrammetry is that objects are measured without being touched. The development of photogrammetry has a long history, particularly the branch related to aerial photogrammetry. Currently, with the development of computer and digital photography, photogrammetric technology has changed dramatically from purely optical equipment to fully digital workflow (i.e. without any type of film or plotter), and a new branch — close-range photogrammetry, also termed vision metrology or videogrammetry, is developing rapidly. Furthermore, inexpensive digital consumer cameras have reached a high technical standard with good geometric resolution, and can replace expensive metric cameras for close-range photogrammetry as long as the accuracy/precision required for the measurement is not too high (Linder, 2009).

While close-range photogrammetry is used primarily for field measurements, the application of optical measurements in an experimental setting has aroused interest. For example, holographic interferometry, moiré and moiré interferometry, and speckle methods are among the optical techniques available for strain measurements. An
innovation, digital image correlation (DIC), developed in the 1980s is now commercialized for use in industrial processes.

Close-range photogrammetry and DIC developed independently from each other. Nevertheless, DIC can be regarded as a branch of close-range photogrammetry. The research presented in this thesis uses digital cameras to optically measure beam deflection profiles and full-field displacements/strains on the surface of specimens and components for inverse identification processes. This chapter briefly reviews some of the concepts and procedures involved in close-range photogrammetry and its application to civil engineering. Two novel techniques proposed for the measurement of beam deflection profiles in a laboratory setting are presented: 1) an edge-based technique using edge-detection suitable for beams with a clean surface and smooth edges; and 2) a surface-based technique using image correlation. Finally, DIC methods are briefly described.

The novel methods are presented in the following sections: Section 2 describes the new approach: edge-based deflection measurement of beams is introduced with example. The data of this example is served as the input to the new damage identification method to be introduced in Chapter 3. In Section 3, the fundamental theory and characteristics of DIC measurement is described. Beam deflection measurement using image correlation is presented in Section 5, with example of a concrete beam.

2.2 Close-range photogrammetry and applications in civil engineering

2.2.1 Overview

The primary purpose of a photogrammetric measurement, both aerial and terrestrial, is the reconstruction of a three-dimensional object in digital form (coordinates
and derived geometric elements). For every image point, values in the form of radiometric data (intensity, grey value, and color value) and geometric data (position in image) can be obtained.

Close-range photogrammetry differs from traditional far-field photogrammetry and image-based measurement systems as the primary task in close-range photogrammetry is to measure the three-dimensional coordinates of targets placed on all areas of interest on a structure or system, whereas the primary tasks in traditional photogrammetry and image-based measurements are feature extraction, object identification, and metrological measurement at relatively lower accuracy (Heijden, 1994). Thus, the key to the technique of close-range photogrammetry is the precise calculation of the positions of each of the targets in the field of view; see (Luhmann, Robson et al., 2006) for an overview of traditional methods and models in close-range photogrammetry.

Laser-scanning measurement is an alternative to photogrammetry. The advantage of laser-scanning methods is that the object can be low-textured while photogrammetric techniques often require highly textured objects. On the other hand, laser-scanning techniques are time-consuming and usually very expensive.

Visible patterns are a very important feature of photogrammetry, particularly patterns that can be identified using pixel or color information. Patterns targeted in close-range photogrammetry are usually simply attached metallic stickers. Popular types of surface patterns targeted in photogrammetry are illustrated in Figure 1 and may include: 1) black and white sprays on the surface; 2) patterns etched onto the raw material; 3)
metallic targets attached to the surface; and 4) small, color printed circles attached to the surface.

![Image](image.png)

(a)

(b)

Figure 2.1: Examples of patterns targeted in close-range photogrammetry. a) Targets attached to the surface and used for measuring displacements of concrete beams (Niederöst and Maas, 1997); and b) a dense grid of circles printed on the surface of testing structures (Cardenas-Garcia, Wu et al., 1997) (Hegger, Sherif et al., 2004) (Franke, Franke et al., 2007).

Traditional close-range photogrammetry is applied to objects ranging from 1 m to 200 m in size; the level of accuracy depends on the distance between the camera and the object, and the size of the area to be measured. Typically, accuracy ranges from <0.1mm for smaller areas to 1 cm for larger areas (Luhmann, Robson et al., 2006; Linder, 2009). Digital image processing (DIP) techniques such as image correlation, image registration, and edge detection are used to extract the required information from digital images.
General information related to: a) basic procedures; b) digital images; c) image coordinate systems and orientation; and d) reference or control points are briefly summarized in the following pages.

**Close-range photogrammetric measurement procedures**

The fundamental procedures of a close-range photogrammetric measurement include the preparation and recording of images, pre-processing, orientation calculations, measurement, and image analysis. The steps in the general protocol for close-range photogrammetry can be summarized as follows:

1. Preparation and recording of images
   a) Application of surface targets.
   b) Determination of control points or scaling lengths.
   c) Image recording.
2. Pre-processing
   a) Computation: calculate reference point coordinates and/or distances from survey observations.
3. Orientation and measurement calculation
   a) Measurement of image points: identification and measurement of reference and scale points, including tie points (points observed in multiple images).
   b) Bundle adjustment: simultaneously calculate both interior and exterior orientation parameters as well as the object point coordinates required for subsequent analysis.
   c) Removal of outliers.
4. Measurement and analysis
   
a) Single point measurement: identify the pixel coordinates of each point to be measured in the image, and transform to physical coordinates.
   
b) Calculate displacements.
   
c) Field measurement: with interpolation and smoothing, measured points can be connected and interpolated to make a field measurement.

Specific experimental procedures for the research presented in this thesis are described in later sections. Some general concepts related to the composition of digital images, image coordinate systems and the use of reference points to establish their relationship to object coordinates (orientation), and digital imaging systems are introduced in the following pages.

**Digital images:**

The essential advantage in using digital images over traditional photos is the ease in image-processing, such as image enhancement, deblurring, edge-sharpening and denoising, thus attains the maximal utilization of information contained in images. The digital image is actually a rectangular array composed of picture elements called pixels. Each pixel is assigned an intensity value meant to characterize the color of a small rectangular segment of the scene. A high-resolution picture can contain 5 to 10 million pixels, while a low-resolution picture small picture may contain comparatively few pixels (e.g. $256 \times 256 = 65536$ pixels). Digital image processing is thus notorious for its intensive computation.

Grey-scale images are typically recorded by means of a charge-coupled device (CCD), an array of tiny detectors arranged in a rectangular grid, which is able to record
the amount, or intensity, of the light that hits each detector. Thus, we can think of a grey-scale image as a rectangular \( m \times n \) array with entries that represent light intensities.

A digital image may be considered as a matrix made up of elements that are positive integers representing surface brightness. Each of these elements is called a pixel, which has a specific value in a range that depends on the digital camera and the image acquisition electronics (e.g. for an 8-bit system, there are 256 grey values ranging from 0-255). Measurements, such as the 2D in-plane incremental displacement field, can be assessed by comparing successive digital images. Generally, grey-scale images of this type are used in image processing for edge detection, image registration, and image correlation.

Metric cameras are monochrome; using a monochrome camera has several advantages over a digital camera with Red–Green–Blue (RGB) color sensor. A monochrome sensor has no color filter in front of the sensor. This increases the International Organization for Standardization (ISO) rating of the sensor to 200 as compared to ISO 80 for the color version of the same sensor. All pixels are sensitive to the same spectrum of light. The information at each pixel location is not interpolated on a monochrome sensor. Pixels of a typical RGB color sensor are arranged in a one layer matrix of which 50%, 25%, and 25% are masked green, red, and blue, respectively. During post-processing this single layer of pixels is interpolated to a triplet of layers, meaning that 50%, 75%, and 75% of the pixels representing the green, red, and blue channels of the image, respectively, must be interpolated from pixels with a different color. This interpolation can lead to artefacts in the images, reducing the geometric quality of the image. Saving the three layers in separate color channels triples the size of
one image to 18 MB for a sensor with 6 million pixels and 8 Bit color depth, as compared to a monochrome image, while no information is added. Image correlation software typically employs only one color channel of an image for analysis (ERDAS, 2002). This implies that 100% of the original information of a monochrome sensor can be utilized by the software while a single layer of a colour image will carry at most only 50% of the original resolution of an RGB sensor. At the same time, image file size is reduced and light sensitivity of the sensor is increased when using a monochrome camera. An image size of 6MB allowed analysis to be performed on uncompressed Tagged Image File Format (TIFF) images. The image compression in JPEG format may bring more blurring and loss of information (Li, Yuan et al., 2002).

However, consumer cameras typically use color sensors. Color images in the RGB format can be separated into three grey-scale images or matrices; one of these can then be used for image processing. Alternatively, one can calculate a mixed monochrome image matrix using the following well-known formula (Linder, 2009):

\[
\text{grey value} = 0.3 \times \text{red}(R) + 0.11 \times \text{green}(G) + 0.59 \times \text{blue}(B)
\]

Generally, this calculation can be easily performed with image-processing software by activating the option “mixed image” when importing images (Linder, 2009; Umbaugh, 2011).

**World and camera coordinates**

In measuring the position and orientation of objects with computer vision methods, we have to couple the coordinates of the camera system (coordinates of the image plane) to some reference coordinates (world coordinates) in the physical space.
The coordinates of the camera system are denoted \( \mathbf{x} = [x, y, z]^T \). Usually the \( z \) axis is aligned with the optical axis orthogonal to the image plane. The world coordinates are denoted as \( \mathbf{X} = [X, Y, Z] \). The two coordinates are coupled by two linear transformations: a translation and a rotation. The translation is a shift of origin and can be described with a vector \( \mathbf{t} \). If the two origins coincide, the remaining differences can be neutralized by rotations. Here, we have also three degrees of freedom: \( \phi, \psi, \) and \( \theta \). Mathematically, rotation corresponds to multiplication of the coordinate vector with a 3 \times 3 orthogonal matrix \( \mathbf{R} \). Clearly, the matrix depends nonlinearily on the three rotation parameters. As a whole, the coupling between world coordinates and camera coordinates is given by the following expression: \( \mathbf{x} = \mathbf{R}(\mathbf{X} - \mathbf{t}) \), i.e. the ideal imaging process is a linear transformation between world coordinates/object points and sensor/image locations. However, there are potential for distortions when using this ideal pinhole model to predict image locations.

**Pin-hole camera model**

Perspective projection is a simple model describing the image formation with a lens system; it is equivalent to a pinhole camera model (Heijden, 1994). Such a model consists of a non-transparent plane with a small hole. Parallel to this plane, at a distance \( d \), the image plane is located. Light emitted from the surfaces of objects in the scene passes through the hole and illuminates the image plane. If the pinhole is small enough, an infinitesimal small surface patch of an object is mapped onto a small spot of light at the image plane. The collection of all surface patches will give rise to an irradiance called image. In the pinhole model, each point in the image plane corresponds exactly to one surface patch in the scene.
The pinhole camera model is based on the principle of collinearity, where each point in the object space is projected by a straight line through the projection center into the image plane. Usually, the pinhole model is a basis that is extended with some corrections for the systematically distorted image coordinates. The most commonly used correction is for the radial lens distortion that causes the actual image point to be displaced radially in the image plane.

**Image coordinate system (pixel coordinate system) and orientation**

In principle, the one-to-one correspondence between the physical and image coordinates of the object has to be established via a camera/lens calibration procedure. The image coordinate system described above is defined by a two-dimensional image-based reference system of rectangular Cartesian coordinates. Its physical relationship to the camera is defined by reference points.

Camera calibration in the context of three-dimensional machine vision is the process of determining the internal camera geometric and optical characteristics (intrinsic parameters) and/or the 3-D position and orientation of the camera frame relative to a certain world coordinate system. In geometrical camera calibration the objective is to determine a set of camera parameters that describe the mapping between 3-D reference coordinates and 2-D image coordinates. The whole calibration procedure may include control point extraction from images, model fitting, image correction, and an additional step to compensate for radial and tangential distortions of the lens. Correction of other error sources in feature extraction, like changes in the illumination, may also be required, especially in field measurements. Various methods for camera calibration can be found from the literature.
Physical camera parameters are commonly divided into extrinsic and intrinsic parameters. Extrinsic parameters are needed to transform object coordinates to a camera centered coordinate frame. In multi-camera systems, the extrinsic parameters also describe the relationship between the cameras. The intrinsic camera parameters usually include the effective focal length, scale factor, and the image center also called the principal point. These coefficients can be typically obtained from the data sheets of the camera and frame-grabber.

Methods where the camera model is based on physical parameters, like focal length and principal point, are called explicit methods. In most cases, the values for these parameters are in themselves useless, because only the relationship between 3-D reference coordinates and 2-D image coordinates is required. In implicit camera calibration, the physical parameters are replaced by a set of non-physical implicit parameters that are used to interpolate between some known reference points.

In traditional aerial and terrestrial analogue photogrammetry, orientation is performed with the use of fiducial marks superimposed on the images and their nominal coordinates in the camera calibration certificate. Measuring (digitizing) the fiducial marks can set up the transformation between camera and pixel coordinates. Exterior orientation must be carried out for each image independently. It can be manually performed by measuring control points (reference points) or automatically using a method called bundle triangulation. The following section describes the use of reference points in establishing exterior orientation.

In these experiments, interior orientation was ignored, thus no lens calibration for lens distortion or check of camera quality was conducted. Measurements presented in
this thesis were taken in a laboratory setting, the camera used was new, and the distance between the object and the camera was short. Our findings show that the accuracy of the measurements was sufficient without accounting for interior orientation. Exterior orientation was determined using marks (reference points) with known (pre-measured) coordinates.

Determining the exterior orientation of an image in an automated fashion is considerably more difficult to implement than line following or driveback. The orientation of an image is typically determined by identifying four or more points of known approximate XYZ coordinates. Once these have been identified, the camera exterior orientation can be computed using a closed-form space resection. To automate the space resection procedure it is necessary to use exterior orientation devices and/or coded targets. Examples of these are shown in Figures 2.1a and 2.1b. If either an exterior orientation device or coded targets are seen in any image they are identified and decoded, and if enough object points with approximately known 3D coordinates are available the exterior orientation can be completed (and even automated with the so-called intelligent camera devices).

**Reference points (control points)**

Ultimately, the objective of determining the orientation of known coordinates is to calculate the relationship between all image and object coordinates. In order to determine the orientation, several control points printed on the surface of the object must be measured (coordinated). A control point is a point on an object that is represented in the image and for which the three-dimensional object coordinates \((x, y, z)\) are known. After obtaining suitable grey-scale or mixed monochrome digital images of the object, we
identify these control points in the image, and determine the coordinates of the control point in the array of image coordinates.

Plane affine transformation can be used to obtain a two-dimensional transformation of the image based on the relationship between the object and image coordinates of the control points (Heijden, 1994). Over-determination of object reference points is required for plane affine transformation; at least three control points are necessary. In general, in order to obtain a stable over-determination, the more control points used, the better the outcome. The research presented in this thesis required at least five well-distributed control points (where three well-distributed control points would form a triangle, not a line).

Optimal accuracy is achieved in areas contained within the control points. Consequently, if multiple cameras are needed to photograph a single object, it is beneficial to include as many identical reference points as possible in neighbouring images. Two types of reference or control points can be used: 1) signalized (or targeted) points that have been applied to the surface of the object; and 2) the object’s natural features. Control points need to be highly visible in the images captured for photogrammetry by digital imaging systems. The following section provides a brief overview of the camera features needed for photogrammetric measurement.

**Digital imaging systems**

Traditional photogrammetry uses metric cameras to acquire images for analysis. A metric camera is characterized by stability rather than flexibility, and has the following features: a known and stable interior orientation, a fixed focal length (i.e., no zoom capability), good lens correction, and a central shutter. Recently, with the rapid
development of digital photography, consumer-grade digital cameras have become commonplace; there is significant potential for their use in photogrammetric applications when calibrated (Cronk, Fraser et al., 2006). In general, the differences between metric and consumer cameras are due to the quality and stability of the camera body and the lens. Consumer cameras often have a zoom lens with large distortions that are inconsistent (e.g. can vary with focal length); thus, it is difficult to correct these distortions using calibration procedures (Linder, 2009).

Linder notes that a consumer-grade camera suitable for use in photogrammetric measurement should have the following properties (Linder, 2009):

- It should be possible to set the parameters for focal length, focus, exposure time and f-number manually.
- The resolution needs to be sufficient for photogrammetry. Generally, the higher the number of pixels, the better the resolution; however, small chips with a large number of pixels have a very small pixel size, and are not very light sensitive. The signal-to-noise ratio is poor, particularly for high ISO values (>200) and in dark parts of the image. In general, lighting requirements are more demanding when using consumer-grade cameras.
- It should be possible to deactivate the auto focus, and manually set the distance parameters.
- The digital images should be saved in a standard format (e.g. JPEG or TIFF). The image compression rate must be selectable; the best option is to turn off compression to minimize possible loss of quality.
• Accessories to reduce unwanted movement and optimize lighting should include a tripod, a remote release control, and an adapter for an external flash.

Note that the focal length or pixel size is required for digital image processing and analysis; pixel size can be obtained from the calibration process. In order to obtain accurate measurements using consumer-grade cameras, usually both interior orientation and exterior orientation determinations are required.

Captured image may be corrupted by noise due to low lighting conditions, which affect the sensors, or due to the noise generated by the electronic circuitry of the imaging hardware. Impulse noise is also commonly referred to as salt and pepper noise. Blurring is caused by a relative motion between camera and object or out of focusing or due to corruption by noise. Blurring is typically modeled by linear operation on the image. Hence restoration is also known as inverse filtering or deconvolution. Pre-processing including deblurring or restoration may be required in photogrammetry.

**Illumination**

The particular type of image formation we discuss is the formation based on radiant energy, reflection at the surface of the objects, and perspective projection. The information of the scene is found in the contrasts (local differences in irradiance). Carefully choosing the illumination of the objects is important in enhancing the contrasts for accurate measurements. The purpose of front illumination is to illuminate the objects such that the reflectance distribution of the surface becomes the defining features in the image. Many applications require this type of illumination: detection of flaws, scratches and other damages on the surface of material, DIC, etc. In outdoor applications, specular illumination or diffuse illumination may be considered.
2.2.2 Applications of close-range photogrammetry in civil engineering

Measurements in civil and mechanical engineering, particularly structural engineering applications have been made without physical contact by using interferometry, moiré technology, holographic and laser speckle interferometry, and theodolite measurement systems (Durelli and Parks, 1970) (Vest, 1979) (Ransom, Sutton et al., 1987) (Post, Han et al., 1994). Recent technological improvements in image acquisition and image analysis opened new possibilities in many fields of engineering and science; the use of close-range photogrammetry in civil engineering is the focus of recent research. Early applications of this technique at large civil engineering sites included measurements of excavation sites and damage assessment after an earthquake (Teimouri, Delavar et al., 2008). Optical methods and image analysis were applied to the observation of cracks in mortar and concrete; the analyses included RGB combination, image filtering, binarization, and shape and fractal analysis of crack patterns (Ringot and Bascoul, 2001).

Numerous laboratory and field studies have been conducted. These include a pilot study of beam deformation measurement using digital close-range terrestrial photogrammetry. Jauregui et al reported the photogrammetric measurement of global deflected shape of structures, which is not practical using traditional instruments (Jauregui, White et al., 2002) (Jauregui, White et al., 2003); in this exercise, the initial camber and dead-load deflection of pre-stressed concrete bridge girders were measured photogrammetrically and compared with level rod and total station readings, and also with dead-load deflection diagrams. Niederost et al. reported using a digital still video camera to measure deformations occurring during the dehydration process of concrete
parts over several months where displacement vectors were obtained through tracking targets (Niederöst and Maas, 1997). Similarly, Maas made photogrammetric measurements of the 3-D coordinates of signalized targets on a large water reservoir wall in Switzerland (Maas, 1998). Measurement of concrete cracks using digitized close-range photographs was reported by several researchers (Barazzetti and Scaioni, 2010) (Chen, Jan et al., 2001); in these works, edge-detection was used to identify the cracks; researchers inspected localized changes in the width of cracks. Franke et al. conducted strain analysis of solid wood and glued laminated timber constructions using close-range photogrammetry; measurements were made of the progression of deformations, cracks and deterioration when loading and relieving the specimens (Franke, Franke et al., 2007).

Hegger et al. studied the crack-opening process in pre-stressed concrete beams using a photogrammetric technique called grid-method (Hegger, Sherif et al., 2004). Whiteman et al. described the use of digital photogrammetry for measurement of deflections in concrete beams (Whiteman, Lichti et al., 2002); a precision of 0.25 mm was achieved for deflections and comparisons were made with linear variable differential transformer (LVDT) deflection measurements. Other works of metrological digital photogrammetry include testing of concrete and column (Woodhouse, 1999), thermal deflection of steel beams (Fraser, 2000), pavement deformation under rolling load (Mills, 2001), and deformation of a coal dredger (Fraser, 1995). In these tests, retro-reflective targets were usually attached to and around the beams. Both stable targets on walls and deforming targets on beams were used.

Consumer-grade digital camera was used for measuring soil erosion, generating digital elevation models from soil surfaces for a planimetric area of $16m^2$ with high
spatial and temporal resolution (Rieke-Zapp and Nearing, 2005). A consumer-grade
digital camera was used for image acquisition. The camera was calibrated using BLUH
software (Jacobsen, 2000). The program system BLUH is a commercial bundle block
adjustment program that allows camera calibration of the interior orientation including
parameters to account for radial symmetric lens distortion as well as other systematic
deviations of the camera geometry from the frame camera model. Homologous points in
overlapping images were identified with least squares matching.

Photogrammetric analysis of photographs taken from a fixed viewpoint at
different times during the loading process has been applied to soil mechanics experiments
to capture non-homogeneous deformation throughout a test (Desrues and Viggiani,
2004). The photographed surface was textured, the scale of the photograph was
determined from six or more reference marks placed on the specimen side.

2.3 Close-range photogrammetry for beam deflection measurement

2.3.1 Overview

Although various close-range photogrammetric techniques have been proposed
and applied to structural engineering problems, human knowledge, experience and skill
still play a significant role in current photogrammetric measurement applications. In
particular, these experiential factors determine the extent to which the reconstructed
model corresponds to the physical object or fulfils the task’s objectives. On the other
hand, digital image processing (DIP) has achieved a level of maturity, and MATLAB and
public domain DIP programs are widely available (Thyagarajan, 2006) (Semmlow, 2004)
(Umbaugh, 2011).
In this thesis, we examined two simple methods for measuring a quasi-continuous deflection profile of beams in the laboratory by processing camera photographs, and compared the results with dial-gauge readings and the predictions of elastic beam theory. The data obtained with these methods can be used as input for inverse damage identification. In this research, we used consumer-grade cameras obtained in the marketplace, and the widely available MATLAB image-processing toolbox to process the images for photogrammetric analysis.

The first method is an edge-based approach; the procedures can be summarized as:

a) An ordinary consumer-grade camera (Nikon D300S single lens reflex (SLR) digital camera, with an array of 2848 × 4288 pixels in each RGB image) was used to take colored photos of the beam with and without loading. The images were saved as a RGB matrix with the following characteristics:

<table>
<thead>
<tr>
<th>Size</th>
<th>Bytes</th>
<th>Data Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>2848×4288×3</td>
<td>36636672</td>
<td>Unit8 array</td>
</tr>
</tbody>
</table>

b) The contrast between structure and background was enhanced using contrast enhancement in Adobe Photoshop. This step is not necessary, but was shown to be useful.

c) Since the cameras were not fixed to the ground, there were always small movements of the camera during photo-capture. The detrimental effects of camera motion were reduced using image-registration; the image-registration technique detected camera movement, which was deleted from the calculated displacements.

d) Image-segmentation and edge-detection were used to identify the continuous boundary between structure and background. This was the key step in this procedure. Edge-detection can be done manually, at a very high cost of time and patience; however, advanced automatic edge-detection techniques have
been developed that can do this job quickly (Semmlow, 2004) (Louban, 2009). Our research found that the performance of automated edge-detection algorithms depends on the level of contrast between the structure and the background and automated edge-detection tools varied in their sensitivity to contrast. For example, in preliminary tests we did not attend to the contrast between the background and the beam; the MATLAB toolbox methods failed to reliably detect edges in these images. Another application, the GROWCUT freeware program (a cellular automata algorithm for edge-detection [www.shawnlankton.com]), worked well with these images, however. Consequently, attention should be paid to lighting and background screening to improve the appearance of the targets in the photographs and enhance the effectiveness of automatic edge-detection techniques.

e) The detected edges were saved as a black-and-white binary image, which is a binary matrix in MATLAB. An averaging process called mollification was used to calculate displacement in pixel coordinates.

f) The displacement solutions in the form of pixel coordinates were transformed to physical coordinates via homogeneous transformation.

**Image-registration**

Image registration is the alignment of two or more images so that they are optimally superimposed. In order to achieve the best alignment, it may be necessary to transform the images using affine transformations (displacement and rotation). Image registration can be assisted or unassisted; we need to perform an unassisted image registration, which relies on an optimization technique that maximizes the correlation between the images.

Unaided image registration usually involves the application of an optimization algorithm to maximize the correlation, or some measure of similarity, between the
images. The structure in the image and in the loading device moved during the tests presented in this thesis, but the supporting frame and the wall in the background remained immobile; therefore, these parts of the fixed background were cut from the whole picture. The appropriate transformation was applied to one of the images, the deformed image, and a comparison was made between this transformed image and the reference image (also termed the base image). The optimization routine seeks to vary the transformation until the correlation is optimal. In the research presented here, image registration was used to detect movement of the camera mounted on a tripod.

**Segmentation and edge-detection**

Image segmentation is the identification and isolation of components of an image into regions that correspond to structural units. Segmentation is used to isolate the structure of interest in both the deformed and the reference images, and thus enable the identification of displacements of the structure. The problems associated with segmentation have been well-studied and a large number of approaches have been developed, many specific to a particular type of image (Heijden, 1994). General approaches to segmentation can be grouped into three classes: pixel-based methods, regional methods, and edge-based methods. Pixel-based methods are the easiest to understand and to implement, but are also the least powerful and, since they operate on one element at a time, are particularly susceptible to noise. Continuity-based and edge-based methods approach the segmentation problem from different angles: edge-based methods search for differences while continuity-based methods search for similarities in the image. The second method is a surface-based approach, in which image correlation is used to find the displacement of a subset of pixels.
2.4 Photogrammetric measurement of beam deflection using an edge-based approach

The DIP methods described in Section 2.3 are useful tools in making a digital photogrammetric measurement. These methods have been extensively developed, but their wise use and commercialization to make industrial photogrammetric measurement is still a problem for the future. In this section, we present a procedure for measuring pseudo-continuous deflection profile of a beam which is based on edge-detection and other DIP techniques described in Section 2.3.

The experiments presented in this thesis used consumer-grade digital cameras (rather than expensive metric cameras) for edge-based measurement of beam deflections via close-range photogrammetry. A photogrammetric technique based on edge-detection was used in this research. The upper and lower edges of the beam were detected in digital images taken at different loading stages. The displacements of the two edges were calculated by comparison to reference images (i.e. digital images recorded when the beam was unloaded). The average displacement of the beam’s top and lower edges is considered representative of the loaded beam’s deflection.

The consumer-grade camera (Nikon D300S single lens reflex (SLR) digital camera) used in the present work has the following features; the sensor size of the camera can be up to 12 megapixels, with an array of $4288 \times 2848$ pixels in each RGB image. More than one camera can be used to measure longer beams, with the field of view for each camera overlapping slightly to ensure image alignment. This technique is becoming standard procedure in state-of-the-art digital image processing; multiple cameras must be
triggered simultaneously. In the current experiments, RGB images (typically JPEGs) were taken for post-processing at each load step and after the deformation had stabilized. The schematic in Figure 2.2 outlines the procedural steps for measuring the beam deflection using close-range photogrammetry.

**Figure 2.2: Procedural steps for beam deflection measurement using photogrammetry based on edge-detection.**

**Edge-detection**: The beam’s edges in the images taken before and after loading were defined using edge-detection image analysis techniques (Semmlow, 2004) (Louban, 2009). The pixel coordinates of detected edges were then transformed to physical coordinates and the displacements between images taken at different load levels were computed. The edge-detection was performed on a sub-image that included both the region of interest on the beam and the background; this increased the efficiency of the
algorithm. The edge detection algorithm operated on the original RGB image; the portion of the image containing the detected edge was transferred to a grey-scale image, and a threshold in pixel values was set to separate the edge from the background. The criteria would either be set manually by observing the grey-scale values of the image or through the histogram of pixel values. To enhance the edge-detection operation, a black screen was placed behind the beam to enhance the contrast. The image was then transformed to a binary grey-scale format. The experimental set-up is shown in Figure 2.3; and one set of detected edges is shown in Figure 2.4. The edge-based photogrammetric measurement technique was shown to be feasible in laboratory settings.

Figure 2.3. The laboratory set-up for tests of a step-cut beam.
Figure 2.4. The detected upper-edge of a cantilever beam: a) a cut-out RGB image; b) the detected edge in a grey-scale image; and c) a binary image of the edge generated using a pixel threshold.

Perspective projection is a simple model describing the image formation with a lens system. It is equivalent to a pinhole camera model. Homogeneous coordinates are a powerful concept to describe the transformations. Translation, rotation, scaling and perspective projection can be handled mathematically with a single matrix multiplication. The transformations can be expressed as $\mathbf{x} = M \mathbf{X}$. In principle, the one-to-one correspondence between the physical and image coordinates of the object has to be established via a camera/lens calibration procedure.

**Coordinate transformation**

Mechanical devices, such as dial gauges, require a stable base for mounting; photogrammetric measurements require the selection of stable reference points on a stable background to calibrate coordinate transformation from pixel coordinates to global physical coordinates. In this study, several stable points in the background and geometric
feature points on the beam area were used as reference points. In the initial state, these points were assigned physical coordinates and these control points were then used to define the homogeneous transformation between pixel and physical coordinates. A homogeneous coordinate system was used to define this transformation which is defined by a $4 \times 4$ transformation matrix ($T$) with a 4th row of $[0 \ 0 \ 0 \ 1]$. The homogenous transformation equation is (Luhmann, Robson et al., 2006):

$$
\begin{bmatrix}
\alpha_{1,1} & \alpha_{1,2} & \cdots & \alpha_{1,r} \\
\alpha_{2,1} & \alpha_{2,2} & \cdots & \alpha_{2,r} \\
\alpha_{3,1} & \alpha_{3,2} & \cdots & \alpha_{3,r} \\
1 & 1 & 1 & 1
\end{bmatrix}
\begin{bmatrix}
t_{1,1} & t_{1,2} & t_{1,3} & t_{1,4} \\
t_{2,1} & t_{2,2} & t_{2,3} & t_{2,4} \\
t_{3,1} & t_{3,2} & t_{3,3} & t_{3,4} \\
0 & 0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
\beta_{1,1} & \beta_{1,2} & \cdots & \beta_{1,r} \\
\beta_{2,1} & \beta_{2,2} & \cdots & \beta_{2,r} \\
\beta_{3,1} & \beta_{3,2} & \cdots & \beta_{3,r} \\
1 & 1 & 1 & 1
\end{bmatrix}
$$

or equivalently

$$
A = TB
$$

(2.1)

(2.2)

where $A$ is the physical coordinate of the reference point, $B$ is the corresponding coordinate of the reference point in the image coordinate system, and $T$ is the transformation matrix to be defined. At least four pairs of non-collinear points are required for computing the matrix $T$, however, as stated before, this transformation is usually optimized (in terms of least squares) with more than four reference points.

**Image registration**

Though mounted on a tripod, temporary movements of the camera cannot be avoided, especially during shooting. This motion may cause changes in image orientation, and induce inaccurate photogrammetric measurements. In order to detect potential detrimental camera movements, image registration of stable background regions
in the image sequence was used to detect any changes in camera orientation; when detected, the change in image orientation was corrected (Semmlow, 2004).

**Mollification**

Model updating and damage identification are often performed with FE models; however, the number of points measured along the beam (>4000 in this research) is much higher than the number of nodes in a FE model. A mollification-based procedure was used for noise-filtering and data reduction. The mollification method is an inverse-problem technique, and has multiple applications when implemented as a filtering algorithm, including the reduction of data through filtering and numerical differentiation (Murio, 1993) (Murio, Mejia et al., 1998). Furthermore, although displacements can be obtained directly through close-range photogrammetry, the rotation angles are indirectly accessible, and the mollification technique allows their approximation (Murio, Mejia et al., 1998).

To illustrate the mollification technique, let us consider the locally integrable two-dimensional function $f(x)$ in a domain $\Omega$ in $\mathbb{R}^2$. The mollifier of $f$, denoted as $J_\delta f(x)$, is defined as:

$$J_\delta f(x) = \int_{\mathbb{R}^2} w_\delta(x-y)f(y)dy \quad (2.3)$$

The mollifier’s radius is $\delta$. For any number $\delta > 0$, we define the family of functions $w_\delta(x)$ as:

$$w_\delta(x) = \frac{1}{\delta^2} w\left(\frac{x}{\delta}\right) \quad (2.4)$$

The weight function $w(x)$ is characterized as $C^\infty$, is non-negative, has a total integral of 1, and vanishes outside the unit ball centered at 0: $B(0,1) = \{x \in \mathbb{R} : \|x\| < 1\}$. The mollifier $J_\delta f(x)$ is a smoothed version of the original function $f(x)$, and it can be used to filter out noise and reduce the data.
1\}. Under such conditions, it is clear that \( w_\delta \in C_0^\infty(B(0, \delta)) \), and moreover, that

\[
\int_{\mathbb{R}} w_\delta(x) \, dx = 1.
\]

In the present study, we selected the following weight function:

\[
w(x) = \begin{cases} 
  C \exp \left( - \frac{1}{1 - \|x\|^2} \right) & \text{for } \|x\| < 1 \\
  0 & \text{for } \|x\| \geq 1
\end{cases}
\]  

(2.5)

where \( C \) represents a constant normalizing the kernel function. It is evident that the corresponding discretized version of the mollification can be computed using a numerical convolution. The numerical convolution and the optimal selection of the mollifier’s radius using cross-validation (Woodbury, 2003) was used to filter the data and numerically compute the derivatives in the current research.

Within the Euler-Bernoulli beam theory, the rotation angle is the derivative of the deflection, \( \theta = dv/dx \). It is well known that direct numerical differentiation of the raw data is an ill-posed problem. The mollification approach to numerical differentiation can be used to substitute the original ill-posed problem of finding the derivative of the deflection, \( v' \), by a new problem of finding the derivative of its mollifier \((J_\delta v)'\). It is demonstrated that this numerical technique is consistent and stable (Murio, Mejia et al., 1998). This means that the reconstruction of the derivative of the mollified data function is stable with respect to the noise existing in the measurements. The central difference scheme is applied to evaluate the first derivative. The rotation at a given node \( x_i \) is obtained as follows:

\[
\theta(x) = \theta_{x_i} = D_0(J_\delta v) = \frac{v(x_i + \Delta x) - v(x_i - \Delta x)}{2\Delta x}
\]  

(2.6)

where \( \Delta x \) is the distance between two successive nodes.
The following example illustrates the implementation of the measurement technique described above; a cantilever beam subjected to a concentrated load was tested. The experimental set-up is shown in Figure 2.5. The camera-to-object distance was approximately 0.5 m, which is sufficient for one-camera measurement of a 1 m-long beam depending on the range of view. The deflection profile and rotation angle that was obtained using the described edge-detection based method are reported in Figure 2.6 and 2.7. These results were compared with the predicted theoretical deflections, and the photogrammetric measurements were found to be accurate.
Figure 2.5: The experimental set-up for close-range photogrammetric measurement of the displacement of a loaded cantilever beam with a high-contrast stable background using consumer-grade optical equipment.
Figure 2.6: Beam deflection: Experimentally-obtained measurements using edge-detection methods and close-range photogrammetry technique.

Figure 2.7: Beam Deflection: Theoretical deflections with $E = 66$ GPa and $E = 72$ GPa, and correspondence of the predicted values to the experimentally-measured deflection profile shown in the previous figure. Predictions were based on elastic beam theory. The elasticity modulus of the aluminum alloy material was calibrated at $70.14 \text{ GPa}$. Measurements were obtained using edge-detection methods and close-range photogrammetry techniques.
2.5 Photogrammetric measurement of beam deflection using a surface-based approach

A surface-based approach to photogrammetry employs image correlation to obtain a measure of displacement. Digital Image Correlation (DIC) effectively tracks the movement of natural surface features, or reference patterns applied to the surface (Fig. 2.1), when a beam is displaced during the test or experiment. The displacement of these surface patterns within discretized subsets or facet elements of the whole image is analyzed. The maximum correlation in each window corresponds to the displacement, and gives the vector length and direction for each window.

2.6 Digital image correlation

The commercially most successful image processing-based measurement technique is the digital image correlation (DIC) method. In recent years, DIC has become a popular tool in the experimental mechanics community for full-field
displacement measurement and experimental strain analysis. A wide range of applications demonstrate the versatility of this technique.

Digital Image Correlation is a full-field image analysis method, based on grey value digital images that can determine the contour and the displacements of an object under load in three dimensions. A recent review on 2-dimensional DIC for in-plane displacement measurement is given in (Pan, Qian et al., 2009); a more comprehensive review of DIC, including both 2D and 3D measurements, and also a comprehensive presentation is given in the book (Sutton, Orteu et al., 2009).

Peters and Ranson (1982) proposed and implemented a DIC technique that is applicable to the computation of surface strains and displacements (Peters and Ranson, 1982) (Chu, Ranson et al., 1985) (Sutton, Turner et al., 1991). This technique relies on the analysis of intensity patterns in digital images of the object under consideration (reference/undeformed and deformed), with a random pattern of white speckles on its surface. (Digitizers were used before the invention of digital CCD cameras). The digital image is actually a rectangular array made up of picture elements called pixels. The determination of local disparities existing between pairs of images, or the problem of image registration, is a basic requirement of computer vision. In image registration, cross-correlation is frequently used to detect local similarities between two images. A common correlation function is the least-squares correlation function (sum of squared differences):

\[ C = \frac{\sum_s (f-g)^2}{\sum_s f^2} \]  

(2.7)
It is non-negative and approaches 0 when the values for f and g are similar. Different alternative versions of the sum of squares exist: normalized sum of squared differences and zero-normalized sum of squared differences; the latter was shown to be the most reliable and robust criterion in case of image blurring and variations in lighting and exposure conditions while taking reference and deformed photos (Tong, 2005). The image correlation algorithm is actually the pattern matching techniques used to compute image motion from a sequence of two or more images (Giachetti, 2000),

Another option is the normalized cross-correlation coefficient (Giachetti, 2000):

\[ C = \frac{\sum (f \times g)}{(\sum (f^2 + g^2))^{\frac{1}{2}}} \] (2.8)

Unlike the least-squares correlation function (which should be minimized in the search for matched pairs of pixel subsets), the cross-correlation function should be maximized.

The DIC technique for the research presented in this thesis consisted of the following: a) Consider two images of the same object before and after deformation. b) Define a subset \( \Omega \) in the undeformed image, centered at point \( A(i, j) \), as an \( (n \times n) \) pixels reference area. c) Define a bigger subset \( \Omega^* \) in the deformed image, centered at the same position, as an \( (m \times m) \) pixel area \( (m > n) \).

The normalized cross-correlation coefficient as a function of position coordinate \( (x, y) \) is written as:

\[ C(x, y) = \frac{\sum_{i=1}^{n} \sum_{j=1}^{n} l(i, j) l^*(i+x, j+y)}{\sqrt{\sum_{i=1}^{n} \sum_{j=1}^{n} l^2(i, j) \sum_{i=1}^{n} \sum_{j=1}^{n} l^*(i+x, j+y)}} \] (2.9)
where \( I(i,j) \) is the grey value of the subset \( \Omega \) at the point \((i,j)\), and \( I^*(i+x,j+y) \) is the grey value of the subset \( \Omega^* \) at the point \((i+x,j+y)\). For the DIC analysis using the ARAMIS system presented in this thesis, the \( n \) value was chosen to be 15.

In order to achieve sub-pixel accuracy (so that the maximum correlation coefficient value can be located between two pixels, rather than at a discrete pixel position), a continuous correlation distribution is constructed by fitting the discrete correlation coefficients to a 2D curved surface. From this theoretical continuous distribution, the maximum correlation coefficient value can be determined using an optimum search technique, thus achieving sub-pixel accuracy.

Before evaluating the similarity between reference and deformed subsets using the correlation criterion, the intensity of these points with subpixel locations must be provided. Thus, a certain subpixel interpolation scheme should be utilized. In the literature, various sub-pixel interpolation schemes including bilinear interpolation, bicubic interpolation, bicubic B-spline interpolation, biquintic B-spline interpolation and bicubic spline interpolation have been used. A high-order interpolation scheme (e.g. bicubic spline interpolation or biquintic spline interpolation) is recommended by Schreier et al. (Schreier, Braasch et al., 2000) since they provide higher registration accuracy and better convergence character of the algorithm than the simple interpolation schemes do.

**Basic principles of DIC measurement**

DIC is an optical full-field measurement technique that measures the deformation of an object’s surface. The image correlation system tracks random grey-value patterns in small areas (image subsets) of images taken during deformation. Two cameras (placed according to triangulation principle) can be used along with stereo vision models to
obtain 3D deformation measurements. The corresponding surface strains are calculated from surface deformations or calculated simultaneously with displacement, depending on the method and representation used.

Several commercial DIC systems have already been developed and commercialized, including: ARAMIS, Correlated Solutions, and LaVision’s StrainMaster DIC system (LaVision). The work presented in this thesis was conducted using the ARAMIS DIC system (GOM and Trillion Quality Systems).

Figure 2.9: the ARAMIS system of the GOM Company

Figure 2.8 shows a typical experimental setup using the ARAMIS DIC system. A random speckle pattern using black and white spray paints is applied to the surface of the specimen. Note that two cameras are positioned on a special tripod with a stable mounting base and with support bars that allow flexibility in positioning the cameras.
External light was used to provide optimal exposure. The lighting is especially important and the contrast between the object and the background affects the quality of the measurements. The stability of the camera set-up is particularly important when capturing images; camera movement and vibration causes errors in calibration and deformation calculations. Trials were conducted to determine optimal conditions for a) camera placement and b) random speckle pattern.

High zoom lengths can be used in the ARAMIS system along with a 1-inch² (25.4 mm) calibration block to measure a small region. A calibration block is a target with a dedicated pattern with known pattern dimensions, and is used to calibrate the cameras. The calibration block must be produced with special care to ensure the accuracy of the pattern’s dimensions. The ARAMIS system comes with pre-manufactured calibration blocks. Random speckle patterns were created and optimized using trial and error.

2.7 Summary

In this chapter we developed one photogrammetric measurement for beam deflection and presented the optical method – DIC, for obtaining strain/displacements on the surface of a specimen. The edge-detection-based photogrammetric measurement of beam deflection profiles provide input data to drive a damage identification procedure which is to be developed in the next chapter. The results were comparable to the predictions of calculated dead-load deflection diagrams. The DIC measurement provides input data to drive the material parameter identification procedures which will be presented in chapters 4 and 5.
“To perform an effective analysis is an art (K.J. Bathe)”, to perform an effective damage identification is even a finer art.

3.1 Introduction and background

3.1.1 An introduction to damage identification problem

Structural systems undergo degradation processes and become aged through the course of time, or may deteriorate suddenly for a variety of reasons, such as unexpected loading or natural causes. As a result, damage occurs, generally in the form of localized stiffness decrease in structural members. In general, early detection of damage is vital in order to monitor the continuous degradation of the structure. Without monitoring, continuous damage can ultimately lead to catastrophic failure of the system.

Recent events show that structures are not immune from disastrous collapse. From some of these events we list: The Silver Bridge failure in West Virginia in 1967 caused 46 deaths; the collapse of the Mianus Bridge in 1983; the de la Concorde overpass collapse in Quebec crushed five people to death; and, on August 1, 2007, the I-35W St. Anthony Falls Bridge over the Mississippi River in Minneapolis, Minnesota collapsed in the middle of rush hour. The collapse of the Minneapolis Bridge in particular has raised a public safety issue concerning the 73,784 bridges in U.S.A that are rated “structurally deficient” by the U.S.A Department of Transportation (Arnoldy, 2007). In response to disasters, there is an ever-increasing demand for assessment of the integrity and
reliability of structures in service. As such, the identification of structural damage has attracted intensive research efforts in the past thirty years.

Structural damage identification technique involves the localization and detection of damages that occur in a structure from measurements of its dynamic and static responses. The process of implementing a damage identification strategy for aerospace, civil and mechanical engineering infrastructure is referred to as Structural Health Monitoring (SHM) (Farrar and Worden, 2007) (Worden, Farrar et al., 2007). This area involved a full research specialization in the last two decades. For example, when replacing the I-35W St. Anthony Falls Bridge, vibrating-wire strain gauges were positioned inside piers and shafts to monitor the new bridge. SHM has become a fundamental tool in maintaining safety and integrity of structures and avoiding loss of life and property, as can be testified by the dedicated academic journal. Since 2002, the journal “Structural Health Monitoring” is fully dedicated to this entire area of research (http://shm.sagepub.com/). In the following, we give a brief presentation of the most used approaches for damage identification.

3.1.2 Material-level methods

Some visual and localized experimental techniques are widely used for monitoring structural behaviours, these techniques include acoustic emission tests, ultrasonic methods, magnetic field methods, radiographs, and eddy-current methods (Phares, Rolander et al., 2001) (Kundu, 2004). These techniques have been called local methods (Doebling, Farrar et al., 1996), but they should more appropriately be referred to as material-based methods, as they detect defects on the material level. The results of these techniques do not directly translate to stiffness changes on the structural level.
3.1.3 System identification

Many investigators have used system identification techniques for the non-destructive assessment of structural integrity. System identification refers to the determination of a mathematical model through observation of the relationships between a system’s inputs and the corresponding outputs (Juang, 1994) (Ljung, 1999). System identification approaches are commonly used for the dynamic systems employed in electrical and mechanical engineering. In civil engineering, the application of system identification focuses on the estimation of modal properties: modal frequencies, damping, modal shapes, etc. Standard techniques exist for selecting the mathematical model structure from a set of model candidates (Ljung, 1999), including physical and non-physical classes of models, such as ARMA models or state-space models. Popular methods for model determination include least squares and recursive least square methods, maximum likelihood methods, and Bayesian methods. System identification relates strongly to theory of optimal controls.

Unfortunately, mathematical models such as state space models and ARMA models are generally not directly applicable to structural damage identification because damage needs to be associated to physical changes; in particular, through the use of parameters that define the structural properties, such as the reduction in stiffness or material modulus, or the size and location of cracks. In contrast, system identification techniques usually provide a phenomenological mathematical model, which requires additional effort to translate changes in the model to damage states in the structure. For example, ARMA models were used for SHM (Peeters, 2000), where modal frequencies
and modal damping could be obtained, the change in these two parameters was used to indicate possible damage.

In structural damage assessment, damage is usually related to changes in the internal material structure for ductile behaviour such as steel materials and/or the appearance of cracks for geomaterials. Damage accumulation is associated with a reduction of the structural members stiffness, and is generally expressed as a “damage variable” comparing the remaining stiffness with a reference value (Bicanic and Chen, 1997) (Kokot and Zambaty, 2009a).

3.1.4 Model updating

Damage identification can be formulated as a parameter-based model updating problem. Model updating focuses on the improvement of a mathematical model using experimental data. Usually a mathematical model is defined to translate parameters describing structural damage to measurable responses or data characteristic of the structure, such as modal characteristics, static deflections, strains, derived characteristics, and dynamic time response history. An optimization problem is usually formulated to minimize a measure of the difference between the experimental and computed outputs. Model updating is solved as an estimation of predefined uncertain structural parameters involving least-square output matching.

As defined, model updating is often solved as a parameter identification problem. Formally, parameter identification is the process used to inversely determine or update a set of unknown parameters from a mathematical model through the examination of measured responses and derived data to a given input. This process is usually called parameter estimation, and statistical analyses are often associated with the estimation
process to determine a confidence region for the estimated parameters. Within structural health monitoring literature, terms such as model identification, structural identification, or damage identification are often used inter-changeably. Thus, these terms are used somewhat arbitrarily.

The model-updating problem is also considered as an inverse problem; depending on the definition of the cost function, the type of data, and the amount of data available, the mathematical problems need to be solved can be well-conditioned or ill-conditioned.

3.1.5 Finite element model updating

Since the FE model has been proven to be the most appropriate tool for modeling structures, model updating for structures usually refers to FE model updating. The purpose of damage identification is to update the parameters of the FE model to match the results with measured data.

Research in the area of FE model-updating methods were initiated in the 1990’s to identify the state of structures from given measurements. Generally, in damage identification problems, a finite-element model of the studied structure, with a known geometry and topology, is constructed with parameterized constitutive models at the element level. Identifying these parameters provides the location and severity of structural damage; understanding of the damage state and prediction of future performance can also be achieved with an updated FE model (Brownjohn, Xia et al., 2001).

3.1.6 The four levels of damage assessment

A system of classification for damage-identification methods, as presented by (Doebling, Farrar et al., 1996), who defined four levels of damage identification:
- Level 1 (Existence): Determination that damage is present in the structure.
- Level 2 (Location): Determination of the geometric location of the damage.
- Level 3 (Severity): Quantification of the severity of the damage.
- Level 4 (Diagnosis & prognosis): Prediction of the remaining service life of the structure.

Levels 1 to 3 pertain to the problem of damage identification. Different methods are used to solve the various levels of a damage identification problem with various degrees of precision. Level 4 refers specially to structural health monitoring after damage is located and quantified or when an updated model is developed to precisely describe the structure’s behaviour. The identified structural parameters or member properties (e.g. $EA$ for trusses or $EI$ for beams) can be used for damage assessment, and for load rating in the management of structural systems (e.g. bridge management systems). Model-based updating and detection techniques are necessary for the assessment of damage severity and the prognosis of the future behaviour of the structure.

3.1.7 Classification of damage identification methods

Nowadays, an extensive literature for damage identification in structures is available. The majority of existing approaches can often be classified into two major categories: i) dynamic damage identification methods using dynamic testing and dynamic test data (modal properties, time histories, etc.); and ii) static damage identification methods using static test data (displacements, strains, etc.).

We can also adopt another classification for damage identification methods to distinguish between model-based and non-model-based methods. As stated before, the
model-based methods are usually FE model updating techniques for damage identification; these methods can be further divided into two classes: i) matrix updating that are non-parametric methods based on property matrix updating; and ii) parametric updating that use a parameterized representation of the model. Most FE model updating methods for damage identification are parametric-based techniques. Non-model-based methods include matching response contours, checking irregularity (often called damage index in the response data and/or derived data from responses), and detection of nonlinearity and pattern recognition paradigms using neural networks and time series analysis models such as AR models, ARMA models (Sohn, Worden et al., 2002) (Omenzetter and Brownjohn, 2006).

The classification of damage identification methods can be based on the targeted application to define two classes: 1) local level damage identification; and 2) global level damage identification. The local level methods focus on the details related to the damage in a given structural member. Usually for a beam member or a plate/shell member, the problem is to locate and quantify localized damage in the form of discrete cracks, or to reconstruct a spatially distributed damage field. A great deal of work has been done to address the problem of crack localization in a beam, using dynamic or static test data. Recently, the problem of identification of stiffness variation in a beam has also attracted interests among researchers. Local level methods reveal details about damage to a structural member. By exhaustive application to an entire structure, these local level methods could provide a complete picture of the current damage state, or health state, of the structure. For large and complex structures, these methods are best used to periodically monitor specific parts of the structure.
A structural model consisting of an assembly of many members is considered when applying global level methods. Parameters are defined as the stiffness of a cross section area/inertia of each member, i.e. damages are defined as the “average” reduction of stiffness for a member, which is usually modeled as one element or a substructure in the FE model. After the localization of a possible damage in a structural member, usually a truss or beam, a direct examination of that member is thereafter performed to determine its condition in detail.

Table 3.1: A synopsis for the classification of damage identification methods

<table>
<thead>
<tr>
<th>Model-based methods</th>
<th>Matrix updating</th>
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<tr>
<td></td>
<td>Parametric updating</td>
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<td></td>
<td>Non-model-based methods, i.e. response-based methods</td>
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</table>

3.1.8 Overview of this chapter

In Section 3.2, the state-of-the-art of damage identification methodologies is reviewed with an emphasis on the static response-based methods. We will also discuss some related issues including measurement of responses, quantification of damage, identifiability, and regularization of ill-posed inverse problems in Sections 3.2 to 3.5. Next, we propose a new methodology for damage identification in Euler-Bernoulli beams using measurements of static deflections. Two formulation techniques are discussed: (i) The first method uses the equilibrium gap concept; and (ii) the second technique employs the adjoint optimization technique to minimize a data discrepancy functionally expressed as a misfit between the measured and model-based deflections. To overcome the ill-posedness inherent in these types of formulations, a regularization technique based on the
Tikhonov-Total Variation (TTV) is used. The proposed methodology is validated using synthetic data for beams with known damage locations and levels (i.e., the distribution of stiffness is known). In the second phase of validation, a series of experiments are presented; four beams with different types of predefined damage were tested and used for validation.

3.2 Literature review

3.2.1 Dynamic damage identifications

The dynamic identification techniques are by large account more extensively studied than the static-based methods, and the corresponding literature is quite extensive. The main assumption is that damages alter the dynamic response of the structure; difference between the predicted and measured dynamic characteristics can be used to retrieve damage information.

Engineers and researchers in the aerospace and offshore oil industries began to study vibration-based damage detection during the late 1970s and early 1980s. Early works used correlation changes in the modal properties to the changes in structural properties (Silva and Maia, 1999). Afterwards different types of dynamic characteristics have been employed, including natural frequencies (Bicanic and Chen, 1997) (Salawu, 1997), modal shapes (Doebling, Farrar et al., 1996) (Kim, Ryu et al., 2003), modal shape derivatives (Maeck and Roeck, 1999) (Ndambi, Vantomme et al., 2002), frequency response function shapes (Liu, Lieven et al., 2009), response time histories, wavelet analysis of dynamic signals (Kim and Melhelm, 2004), and harmonic responses (Kokot and Zambaty, 2009a) (Liu and Chen, 2002) (Kokot and Zambaty, 2009b), impulse responses (Sophia and Karolos, 1997) (Mangal, Idichandy et al., 2001).
The literature on vibration-based damage identification methods is extensive, as testified by the number of reviews published. We cite few review papers in this area: Natke reviewed the FE model updating techniques in frequency domain (Natke, 1988). Friswell and Mottershead gave an extensive review of FE model updating using dynamic data (Friswell and Mottershead, 1995), a special issue of Mechanical Systems and Signal Processing is devoted to this topic (Mottershead and Friswell, 1998), and a recent overview of dynamic based methods is available in (Friswell, 2007).

Literature review with a focus on the statistical pattern recognition viewpoint, and emphasized data fusion, cleansing, outlier analysis was given in (Doebling, Farrar et al., 1996), later extended in (Sohn, Farrar et al., 2003), the authors referred to these techniques as statistical models. Other important reviews include (Lynch and Loh, 2006) (Carden and Fanning, 2004) (Doebling, Farrar et al., 1998) (Doebling and Farrar, 1997).

A recent literature review that includes the basic approaches of dynamic monitoring, and guidelines for sensor selection, data collection is available in (Hsieh, Halling et al., 2006). In this review, the experimental approaches were separated into ambient vibration, forced vibration (usually harmonic), and free vibration monitoring. Ambient vibration tests are easiest to conduct; however, in ambient vibration methods, stationarity, whiteness, and unidirectional excitation are often assumed.

Because of the extensive literature for vibration-based damage identification and health monitoring research, an adequate classification is necessary to simplify the review. In the classical review by Doebling et al, dynamic identification methods were categorized as (Doebling, Farrar et al., 1996):

- Modal frequency Based Methods
• Modal Shape Based Methods
• Modal Shape Derivatives such as Curvature/Strain Modal Shape Based Methods
• Dynamically Measured Flexibility Based Methods
• Matrix Update Based Methods (direct, non-parametric updating method)
• Non-linear Methods (detection of nonlinearity)
• Neural Network Based Methods
• Other Methods

This categorization is mainly based on the types of dynamic characteristics employed; this classification paradigm was later followed in many other reviews, with sometimes minor modification (Carden and Fanning, 2004).

We had given a different synopsis in section 3.1.7: model-based methods, and non-model-based methods. Most damage identification methods are model-based approach, in which a model (usually a FE model) is updated to minimize the difference between measured and calculated properties; therefore model-based methods are normally FE model updating methods, and can be further divided into two classes: matrix updating and parametric updating.

Model-based methods - I: Matrix updating for dynamic FE model updating

The non-parametric, matrix updating methods adjusts the stiffness, mostly along with the mass, and in some cases, the damping matrices, from measured data (mostly modal testing data or dynamic FRFs). These methods tend to fit the available experimental data exactly, update the mathematical model, but generally do not provide
any direct physical meanings, and thus they are usually not suitable for damage assessment purposes, but they are useful for control design and predictive modeling. This type of methods have been called “direct updating” methods in the literature since no iterative solution process is required (Caesar and Pete, 1987), but we prefer to call them “non-parametric updating” methods, or “matrix-updating” over “direct methods” or “direct updating methods”, since the name “direct methods” is widely associated to optimization algorithms without utilizing gradient information.

Non-parametric updating methods are global level methods; although they can be performed to deal with any level of complexity, for damage identification purposes, the mathematical models need to be very simple, such as discrete spring-mass systems.

**Model-based methods - II: parametric updating methods for dynamic FE model updating**

In parametric-updating methods, the problem of damage identification is formulated as a parameter estimation problem where an error function is defined as the discrepancy between a mathematical model response and measured data. The formulated problem is in the form of an optimization problem where the unknowns are the parameters defining the mathematical model that fit the best the measured data. Parametric model updating methods can be subdivided according to the nature of the measured data they use which can be: mode frequency, mode shapes, mode shape curvatures, etc, as in the classification we have mentioned by Doebbling et al. (Doebbling, Farrar et al., 1996).
Based on the optimization scheme employed to solve the problem, a model-updating problem can be divided as iterative or non-iterative methods, in the latter direct analytical solutions are possible and the former require iterative updating. Furthermore, iterative methods include sensitivity-based and non-sensitivity-based; the former employ gradient-based optimization schemes, and the latter, direct search methods such as response surface methods, genetic algorithms, Nelder-Mead simplex method, particle swarm, cluster-based stochastic search, etc.

In the early 1990’s most of parametric model updating formulation employ gradient-based search methods. For this reason, the parameter estimation methods for damage identification have been referred to as sensitivity-based methods (Doebling, Farrar et al., 1996) (Farhat and Hemez, 1993), because they usually make use of sensitivity of the stiffness matrix to the unknown structural parameters in the gradient-based iterative updating process. However, sensitivity evaluations are not always necessary in solving optimization problems, direct derivative-free methods and evolutionary methods can be considered as an alternative to solve the parameter identification problems for damage identification as they have specific advantages such as stability and exhaustiveness of search space. Recent reviews of the alternative shows that methods based on derivative-free algorithms are becoming a competitive choice.

**Response-based Dynamic Identification Methods without reference to models**

Dynamic identification methods that do not use a specific mathematical model are called response-based. Techniques used in these methods include direct matching contours, irregularity checking in responses and derived properties, damage index from
response data or derived data. In response-based approaches, no baseline model is required, and the irregularities are detected from the measured or derived responses, such as curvature mode shapes.

Possibly the first work in this category is made in (Lifshitz and Rotem, 1969), who proposed a method for damage detection from measure frequencies. The authors related the changes in the dynamic moduli to the frequency changes in particle-filled elastomers. Most of early work of vibration-based damage identification focused on rotating blades and machinery. An early literature survey of damage detection using modal properties was due to (Richardson, 1980), which cites large amount of research work dealing with rotating machinery.

Masoud and Al-Said (2009) developed a crack localization algorithm exploring the variation in a single frequency of a beam as a function of rotor speeds (which is equivalent to varying axial loading of the beam) to detect and localize a crack. The frequencies are obtained analytically using Lagrange equations and an assumed mode method (Masoud and Al-Said, 2009); the contour lines of the cracked beam frequencies are plotted with crack location and crack depth as its axis; the identification procedure is developed utilizing the contour plots. Similar crack identification procedures in cantilever beams based on contour plots of modal frequencies in terms of crack depth and crack location were developed by (Nahvi and Jabbari, 2005) (Batabyal, Sankar et al., 2008), in which FEA are used to evaluate modal parameters. It is worth mentioning that these methods are non-model-based methods, because the analytical or FEA solutions are used to obtain the contour plots only, and the identification is performed with the contour plots without reference to any model.
Pandey, Biswas et al (1991) suggested another damage identification method based on curvature mode shape of beam, instead of natural frequencies changes, to identify the location of a crack (Pandey, Biswas et al., 1991). It is assumed that large change in curvature indicates the location of damage. This method is further improved by Ratcliffe (Ratcliffe, 1997) (Ratcliffe and Bagaria, 1998); he suggested using the third order polynomial interpolation of the second-order finite difference of modal shapes; this indicator is more sensitive than the curvature. One limitation of these methods is that only a single crack can be considered in the beam to be detected. Beams with more than one single crack cannot be analysed with this methodology.

3.2.2 Static response data-based damage identification

In comparison to dynamic-based identification methods, the literature of damage identification methods based on quasi-static responses is rather limited. The synopsis table in section 3.1.7 can also be used to group static identification methods, in which most static identification methods belong to the group of parametric updating, and the non-model based methods include only a few works using signal processing to detect irregularities in static deflection.

Early work on static-based methods started in the 1980s. Sheena et al presented a method for improving the analytical stiffness matrix from noise-free static measurements, in which they optimize the difference between the theoretical and correct stiffness subjected to measured displacements constraints (Sheena, Zalmanovitch et al., 1982b) (Sheena, Zalmanovitch et al., 1982a). The method requires the measurement of the displacements at all the active DOFs associated to a FE model. If only a limited number
of DOFs are measured, spline interpolation can be used to create the complete set of displacement data. Within this formulation, the stiffness is computed as a whole matrix, not directly related to the internal parameters controlling the properties of each element. To the author’s best knowledge, this is the only published work on matrix updating using static responses.

Most static response-based identification methods can be considered as part of parametric FE model updating group. Early research in this area was due to a group of researchers including Sanayei, Hajela, Banan, Hjelmstad. For example, Sanayei and Scampoli presented an identification procedure of plate-bending stiffness parameters for a one-third scale, reinforced-concrete pier-deck model; the formulation is based on FE model updating using static test results through minimization of the equilibrium gap functional (Sanayei and Scampoli, 1991). Sanayei and Onipede presented an analytical method for the identification of properties of structural elements using static test data (Sanayei and Onipede, 1991). In this approach, the forces are applied to a set of DOFs and the associated displacements are measured at another set of DOFs. An iterative procedure is used to minimize the difference between the measured and the model response. The sensitivity analysis proposed by Adelman and Haftka (1986) is used at the heart of this iterative method to identify directly the structural element parameters (Adelman and Haftka, 1986). The stiffness matrix properties such as element connectivity, positive definiteness, symmetry and bandedness are automatically preserved.

Hajela and Soeiro (1990) presented a review on damage detection techniques; the authors classified both static and dynamic identification techniques into three categories:
the equation error approach, the output error approach, and the minimum deviation approach (Hajela and Soeiro, 1990a). Given an equilibrium equation in the form $K(\theta)x = f$, where $x$ is the state variable, and $f$ is the excitation, $\theta$ is a set of internal variables, then the equation error, also called force error estimator is defined as: $\min_\theta K(\theta)u - f$, and the output error (displacement error) estimator is defined as $\min_\theta QK^{-1}(\theta)f - u_{measured}$, where $Q$ is a Boolean matrix extracting the measured responses from the whole set of DOFs. The advantage of the equation error approach is that the error is a linear function of the entries in $K(\theta)$. Its disadvantage lies in the necessity to measure all the DOFs in the model. The advantage of the output error approach is that the minimization can be operated on the measured DOFs and therefore no need to measure all the DOFs. The disadvantage of the output error estimator is that the error is now a highly nonlinear function of the entries in $K(\theta)$.

The equation error and output error approaches applied for static identification are studied in details by Banan et al (Banan, Banan et al., 1994a) (Banan, Banan et al., 1994b). In 1994, the authors studied the two formulations for estimating the constitutive parameters needed to calibrate finite-element model results with measured displacements from known static loading. The authors solved the problem by minimizing the error between the simulated displacements and the on-site measurements using sensitivity-based optimization which is similar to the method developed by Sanayei and Onipede in 1991 (Sanayei and Onipede, 1991). The problem was formulated as a constrained nonlinear optimization using either a force-error (output error) estimator or a displacement (equation error) estimator. The methodology was tested on a simulated 25 member bowstring truss; Monte Carlo simulations are used to study the performance.
Later, Sanayei and Saletnik conducted a similar work as that of Banan et al, but used static strain measurements instead of the displacements (Sanayei and Saletnik, 1996a) (Sanayei and Saletnik, 1996b). The internal parameters were estimated by minimizing the error between the theoretical and measured strains in the structure resulting from a series of loading cases of concentrated forces. A similar approach was used by Hjelmstad and Shin, where they proposed to group the parameter reduce the number of unknowns, especially in case of inadequate measurements (Hjelmstad and Shin, 1997).

In 1997, Liu and Chian developed a procedure for identifying the cross-sectional areas of a truss using static strain measurements resulting from a series of concentrated force loading cases (Liu and Chian, 1997). A closed-form solution was obtained for the truss. A numerical example is presented along with model test results.

Chou and Ghaboussi (Chou and Ghaboussi, 2001) used a genetic algorithm to identify damage in a truss based on measured deflections and verified with numerical examples. An optimization problem is formulated for the detection and identification of structural damage. Both the “‘output error’” defined as a measure of the difference between the measured and computed responses under static loading condition and the “‘equation error’” indicating the residual force in the system of equilibrium equations are used to formulate the objective function to be optimized. Recently, Shenton and Hu used a misfit cost function using strain measurements and a genetic based minimization algorithm to identify damage in beams subjected to static loading (Shenton and Hu, 2006) (Hu and Shenton, 2007). Shenton and Hu proposed a damage distribution formulation based on the re-distribution of the dead load strain taking place in a structure when
damage occurs; since the structure’s own weight is always present, this method eliminates the need for an external loading apparatus and is also suited for designing a permanent SHM system. In this technique, the damage is identified by minimizing the error between the measured strain changes and the model strains prediction in the damaged structure. The problem is formulated as a constrained optimization problem and solved by a real-coded genetic algorithm; the basic procedure is introduced for an example of a single-span fixed-fixed beam using a closed-form solution (Shenton and Hu, 2006) (Hu and Shenton, 2006). In a later paper, this procedure was extended to use finite elements (Hu and Shenton, 2007).

In 2005, Nejad et al. presented a method to describe the change in the static displacement of certain degrees of freedom by minimizing the difference between the load vectors of damaged and undamaged structures (Nejad, Rahai et al., 2005).

From the review of the literature, it appears that all of the static-based methods developed to date require one or more external loading cases to be applied to the structure and that the corresponding static responses being measured. The loading configuration to excite the structure needs to be designed to make sensitive results. One may note that in practice, this requirement might be feasible in bridges, but may pose a difficulty in many other types of structures.

Choi et al. presented a solution based on the conjugate beam method; it was shown that for determinate beams, the shape of the displacement variation due to damage is related to the influence line of the moment in the conjugate beam (Choi, Lee et al., 2004). Other research work proposed a formulation of locating cracks through wavelet
analysis of the static deflection profile using signal processing techniques (Rucka and Wilde, 2006).

Some authors proposed that static response data to be combined with dynamic data in the damage identification. It is believed that more information can be available by combining different type of measurements. Hajela and Soeiro proposed that static displacement information be incorporated to supplement the modal data for damage detection (Hajela and Soeiro, 1990b). Simulated static deflections and vibration mode data are used successfully for parameter estimations (Hajela and Soeiro, 1990b). Another method making use of composite data was proposed by Wang et al.; the proposed method can be considered a two-stage identification algorithm for identifying the structural damages by employing the changes in natural frequencies and measured static displacements (Wang, Hu et al., 2001).

Oh and Jung proposed a method combining both static displacements measurement and identified dynamic modes (Oh and Jung, 1998); the proposed approach allows the use of composite data consisting of a combination of static displacements and eigen-modes. In dynamic tests the curvature and slope of mode shapes are included in the formulation of the error responses. To examine the capability of the proposed damage assessment algorithm a series of tests for a predetermined damaged two-span continuous beam and a planar bowstring truss structure were performed. This work showed that the combination of the curvature or slope of mode shape and the static displacement data results a superior capabilities for the damage detection and assessment.
However, conducting dynamic and static tests complicates the application in real world structures.

The methods reviewed before are global level methods, which the damage is identified as average stiffness reduction of a structural member. Local level methods can be divided into two groups based on the way of parameterization.

Most existing damage detection methods aim at identifying localized damage zones and quantifying their intensity (Yang, 2002) (Buda, 2006) (Nikolakopoulos, 1997) (Chen, 2005). Within this approach, cracks are usually parameterized beforehand assuming accurate knowledge of the structure stiffness distribution.

For methods developed to localize and quantify concentrated cracks, it is usually assumed that an accurate knowledge of the stiffness distribution of the undamaged structure is available, and the unknown parameters define the crack location and depth. Buda and Caddemi presented a crack identification scheme for Euler-Bernoulli beams based on static displacements (Buda and Caddemi, 2007).

For methods developed to localize and quantify concentrated cracks, it is usually assumed that an accurate knowledge of the stiffness distribution of the undamaged structure is available. Earlier research showed that numerical algorithms that perform well for the discrete damage identification may be inefficient for the stiffness reconstruction problem.

Lesnic et al. presented a mathematical analysis of the identification of a heterogeneous flexural stiffness distribution of an Euler-Bernoulli beam. The authors established the conditions for the well-posedness of the associated inverse problem (Lesnic, Elliott et al., 1999).
All of the static-based methods developed to date require one or more cases of external loadings to be applied to the structure and corresponding static responses to be measured. The loading configuration to excite the structure need to be designed to make sensitive results; one may note that in practice, this requirement is simple for bridges, but may pose a difficulty in many types of structures.

For methods dealing with the identification of distributed stiffness, the problem consists of recovering a continuous distribution of the beam’s stiffness using experimental data. Mathematically, the problem involves a minimization of a functional, and it is generally more difficult inverse problem to solve than the identification of discrete cracks. Usually, the identification of distributed stiffness requires extensive amount of information than for the case of discrete cracks. Focusing on the theoretical aspect of the problem, it is worth to mention the work Lesnic et al. who presented a mathematical analysis of the identification of a heterogeneous flexural stiffness distribution of an Euler-Bernoulli beam. The authors established the conditions for the well-posedness of the associated inverse problem (Lesnic, 1999). Earlier research showed that numerical algorithms that perform well for the discrete damage identification may be inefficient for the stiffness reconstruction problem.

Chou and Ghaboussi (Chou and Ghaboussi, 2001) used a genetic algorithm to solve an optimization problem formulated for detection and identification of structural damage. The “output error” indicating the difference between the measured and computed responses under static loading and the “equation error” indicating the residual force in the system of equilibrium equations are used to formulate the objective function to be optimized.
The response surface methodology, which is a combination of statistical and mathematical techniques, was employed by Ren et al to update a FE model based on static responses of structures (Ren, Fang et al., 2011); uniform DOE was used in building the response surface models. This method was verified in a numerical beam and an experimental box-girder bridge.

3.2.3 Advantages and disadvantaged of static and dynamic identifications

Advantages and disadvantaged of dynamic identification

Vibration testing is easier to conduct; it is generally easier to excite a large structure dynamically, particularly with harmonic loading or environmental loading, although one often cited challenging problem when applied to real structures is that it is often impractical to excite full scale structures in controlled way. It is also easier to measure and continuously monitor acceleration response.

One significant disadvantage of dynamic identification is the requirement of mass matrix and damping in dynamic model, while the required results are usually of stiffness only. Estimating damping is difficult, and although model masses can be estimated from structural drawings, it adds uncertainty in modeling errors.

Soil-structure interaction poses another difficulty in dynamic identification, but not for static identification; non-structural elements is another source of uncertainty need to be considered in dynamic identification. Dynamic-based identifications are mostly based on modal identification results. In modal identification, only lowest frequencies and modal properties can be estimated with certainty. For ordinary structures, no more than 3~5 frequencies can be accurately identified; for very large bridges, this number can
be 10. However, Damage is typically a local phenomenon. Localized damages often require higher modes to be accurately measured which are impossible.

Some modal shape-based algorithms require complete mode shapes to be measured. When only a small number of sensors are installed, one can use either system condensation techniques or modal expansion techniques. Some methods require a very dense measurement of displacement or strain mode shape (Goldfeld, 2007);

System condensation techniques reduce the DOFs defined in the analytical model to the measured DOFs. Reduction techniques often produce a condensed matrix that does not resemble the member connectivity of the original model. On the other hand, modal expansion methods generally do not produce the results that are accurate enough to provide reliable information about damaged DOFs or damaged structural members.

Up to date the most successful dynamic-based damage identification practice is the local-level frequency-based detections of rotating blades which is already a tool in routine practice for rotating machinery. Rotating blades can be regarded as cantilever beams under various axial loads induced by centrifugal forces; however this type of cantilever beams is scarce in real civil infrastructures. Furthermore in civil structures, it can be difficult to isolate local modes, generally a global mean, some local identification using modal properties, this is the major difficulty; another problem is that cracks in truss behaviour members are not revealed by frequency-change, but are dangerous. One needs to use local bending models to identify cracks, i.e. local level test need local bending modes.

In beam damage identification using dynamic responses, most of the investigations reported in the literature focused on testing single free–free beams, for
example, references: (Maeck, Wahab et al., 2000) (Ren and De Roeck, 2002) (Cerri and Vestroni, 2003) (Wahab, De Roeck et al., 1999), i.e. the global and local modes are identical and are for the beam only; this type of condition rarely occur in real practice.

Another difficulty using dynamic properties is the variability of dynamic parameters identified using system identification techniques as a result of environmental and operational conditions, such as temperature, moisture, wind and others (Cornwell, Farrar et al., 1999). It has been demonstrated that changes in modal parameters due to environmental and operational factors may well exceed those caused by even severe damage, and thus if this variability is neglected, it is very difficult to draw reliable conclusions about structural condition (Sohn, Dzwonczyk et al., 1999) (Farrar, Cornwell et al., 2000).

If measured over a fair frequency range, frequency response functions (FRF) is high density data and generally contain much more data and information than frequencies and modal shapes, however the difficulty is that FRFs are heavily influenced by damping, and it is generally difficult to model damping exactly in a FE model. One may try to assume Rayleigh damping and identify the damping ratios or Rayleigh coefficients multiplying the mass and stiffness matrices. Some modal identification methods give complex modal shapes in most types of damping; in this case a complex transformation is often required to transform a complex modal shape to a real one (Niedbal, 1984). This may introduce further uncertainties in following damage identification.

Frequency response functions (FRFs) contain a large amount of redundant information since there are many more points in a FRF than parameters in the model. It is
therefore necessary to choose which points of the FRF should be used in updating the model. On the other hand, if the sensitivity is too high, it will suffer from noise since it will be highly sensitive to noise. Another advantage of FRF is that they are not restricted to linear systems, and modal parameters in structures with high modal density are difficult to identify.

**Advantage and disadvantages of static identification:**

There is no need to assume mass, damping in static identification, and in some cases, the test is easier, especially when optical measurements are used. The measurement devices are being developed, including inclinometers, strain gauges, and especially the close-range photogrammetry, which considered by the author as one very promising development for civil engineering in future.

In global level identification, the effect of the damage may be concealed due to the limitation of load paths. For a real structure, the damaged components which have fairly little contribution to structural deformations under a certain load case will be difficult to identify. These kinds of limitations can be practically overcome by optimizing the loading scheme according to the proper pre-analysis or loading several groups of load synthetically. A static test is easy to excite the whole structure in local level. Multiple loading cases are usually required in global level identification.

Some of the most successful results of damage identification have been achieved on laboratory scale truss structures, both dynamic and static identifications, in dynamic tests, the lack of rotational degrees of freedom and ease of accessibility allow entire mode shapes to be measured (Kosmatka and Ricles, 1999); while in static tests, displacements
can be measured at joints only or strains to be measured in each single bar member (Hjelmstad and Shin, 1997) (Liu and Chian, 1997).

Successful damage identification and stiffness recovery require efficient measurement techniques of the structural response. Traditional measurement techniques using LVDTs and strain gauges allow only localized measurements along the beam and therefore provide reduced amount of information. Recent progresses in image processing techniques and digital cameras permit a quasi-continuous deflection profile measurements of beams (Jiang, 2008). This progress offers new possibilities in structural parameter identification, damage localization. Unlike localized testing sensors, digital images (Digital Image Correlation and close range digital photogrammetry) are able to provide a large number of spatially distributed measurements. The large quantity of experimental data in the form of quasi-continuous measurements of the deflection profile becomes a valuable input for the damage identification problem.

3.3 Equilibrium gap method

Claire et al. (Claire, 2004) developed the general concept of the equilibrium gap to identify damages in 2D structures. When restricted to the elasticity domain and in the absence of volumetric loading, the continuous format of the equilibrium equation is given by: \( \text{div}(\sigma(u)) = 0 \), or equivalently \( \text{div}(\sigma(E,u(x))) = 0 \), where the Cauchy stress \( \sigma \) is a function of the displacement field, \( u(x) \), and the material parameters \( E \) and \( \nu \). Using finite element discretization, the continuous equilibrium equations can be written as a set of discrete equations in the form of \( [\sigma \cdot n] = 0 \), where \([*]\) denotes the jump of the
quantity \* in the continuum. The stress jump equations can also be given as \( \sigma_i n = \sigma_j n \), where \( n \) denotes the normal of an interface between the two sides of a section. Given a measured displacement field, a piecewise-constant material parameter distribution is assumed and the distribution is evaluated through a finite element formulation.

The principle of equilibrium gap formulation can be applied specifically for beams where the equilibrium equations can be written as functions of the generalized forces at any given section:

\[
F_l + F_r = P_{lr} \tag{3.1-a}
\]

\[
M_l + M_r = M_{lr} \tag{3.1-b}
\]

where the indices \( l \) and \( r \) denote the internal force to the left and right of the section, respectively. The variables \( P_{lr} \) and \( M_{lr} \) denote the external force and moment at the cross section. Equations (3.1-a) and (3.1-b) are the equilibrium conditions associating the external and internal forces; in each material point, the sum of all internal forces arising from various adjacent elements equals the applied load (Figure 3.1).
Figure 3.1: Internal force equilibrium in Euler-Bernoulli beams subjected to flexure.

In contrast with the equilibrium of 2D solids, the external loading is directly included in the beam’s equilibrium expression since the boundary conditions are explicit. The equilibrium between the internal forces and external loads is therefore used as a condition that must be imposed as a constraint. Within the context of FE formulation of Euler-Bernoulli beams and considering isotropic damage within a given cross-section, the force-displacement relations of an Euler-Bernoulli beam are given by:
\( k_d \textbf{d} = f \)
\[
(1 - D) k_0 \textbf{d} = f
\]
\[
(1 - D) \frac{E I_0}{L^3} \begin{pmatrix}
12 & 6L & -12 & 6L \\
6L & 4L^2 & -6L & 2L^2 \\
-12 & -6L & 12 & -6L \\
6L & 2L^2 & -6L & 4L^2
\end{pmatrix} \begin{pmatrix}
v_1 \\
\theta_1 \\
v_2 \\
\theta_2
\end{pmatrix} = \begin{pmatrix}
V_1 \\
M_1 \\
V_2 \\
M_2
\end{pmatrix}
\]

(3.2)

where \( (v_1, v_2, \theta_1, \theta_2)^T \) are the nodal degrees of freedom (vertical deflection, rotation angle), and \( (V_1, M_1, V_2, M_2) \) are the dual variables (shear, moment). The stiffness matrix of the element at the damaged state is denoted \( k_d \) and is proportional to a reference value for stiffness \( k_0 \). The variable \( D \) (0 ≤ D ≤ 1) is a scalar indicating the damage level (stiffness decrease). When \( D = 0 \), the beam element is considered undamaged and its stiffness is equal to the reference value. At the other limit, the ultimate condition \( D = 1 \) represents a complete loss of the beam’s stiffness. This definition for damage is in accordance to the continuum damage mechanics (Lemaitre, 1996). When the axial deformation is neglected, the stiffness matrix depends only on the flexural stiffness \( E I \), and the beam’s element length. For each element, we define a reference value for stiffness \( E I_0 \), and note that its value can vary along the beam. We will assume that a reference distribution is \( E I_0(x) \), and take this as the initial estimate for the identification process. For example, this reference stiffness can, but not necessarily, correspond to the stiffness of the undamaged beam. From Eq. (3.2), a possible definition for the damage variable would be \( D(x) = 1 - \frac{E I(x)}{E I_0(x)} \), where \( E I(x) \) is the actual stiffness of the cross-section at a position \( x \). It is clear that when \( E I \) decreases, damage increases linearly, so
that the theoretical limit of 1 is reached when the cross-section loses its flexural resistance. Using this definition, the beam’s stiffness matrix is expressed as a function of the damage variable $D$ (with $k = (1 - D)k_0$). The parameter $(1 - D)$ corresponds to the stiffness reduction factor. When FE discretization is used, a constant damage parameter $D$ is assumed for each element, which corresponds to a piecewise-constant definition of the damage field along the beam. (this is original application of the equilibrium gap)

At present, we will assume that the deflection profile is available from non-contact measurements made using close-range photogrammetry. For each pair of adjacent elements, $e$ and $e + 1$, the following equilibrium equations can be written:

\[(1 - D_e)(k_{0e}^3)_{e} d_{e,r} + (1 - D_{e+1})(k_{0e}^1)_{e+1} d_{e+1,l} = (V_2)_l + (V_1)_r = P_{lr} \quad (3.3-a)\]

\[(1 - D_e)(k_{0e}^3)_{e} d_{e,r} + (1 - D_{e+1})(k_{0e}^2)_{e+1} d_{e+1,l} = (M_2)_l + (M_1)_r = M_{lr} \quad (3.3-b)\]

where $(k_{0e}^3)_{e}$ is the third row in the reference stiffness matrix of element $e$; and $D_e$ is the associated damage variable. Similarly, $(k_{0e}^1)_{e+1}$ is the first row of the reference stiffness matrix of element $e + 1$; $d_{e,r}$ and $d_{e+1,l}$ are the displacement vectors of the node on the right and left sides of elements $e$ and $e + 1$, respectively. The generalized forces $P_{lr}$ and $M_{lr}$ are the concentrated load and moment applied at the node connecting elements $e$ and $e + 1$. In writing these equilibrium equations for each pair of adjacent elements, one can arrange the linear system in terms of the damage variables to obtain the following algebraic equations:
\[
\begin{pmatrix}
(k_0^3)_{1}d_1 & (k_1^3)_{2}d_2 \\
(k_0^4)_{1}d_1 & (k_0^5)_{2}d_2 \\
(k_0^4)_{2}d_2 & (k_0^6)_{3}d_3 \\
(k_0^4)_{2}d_2 & (k_0^7)_{3}d_3 \\
(k_0^4)_{2}d_2 & (k_0^8)_{3}d_3 \\
\vdots
\end{pmatrix}
\begin{pmatrix}
1 - D_1 \\
1 - D_2 \\
\vdots \\
1 - D_{\text{nel}}
\end{pmatrix}
= 
\begin{pmatrix}
P_1 \\
M_1 \\
\vdots \\
M_{\text{nel}}
\end{pmatrix}
\]

Finally, the problem for identification of the damage variable is written as:

\[
\begin{cases}
\text{Find} & \boldsymbol{\theta} = (1 - D_1, 1 - D_2, \ldots, 1 - D_{\text{nel}})^T \\
\text{such that} & \mathbf{G}\theta = \mathbf{R}
\end{cases}
\]

where \(\boldsymbol{\theta}\) is the set associated with the damage variables corresponding to each element

\(\theta = (1 - D_1, 1 - D_2, \ldots, 1 - D_{\text{nel}})\), and \(\mathbf{R} = (P_1, M_1, \ldots, P_i, M_i, \ldots)^T\) is the external force

at the nodes associated with the generalised degrees of freedom (i.e. displacements) \((d_1, d_2, \ldots, d_n)\).

This formulation leads to a typical inverse problem, where the coefficient matrix \(\mathbf{G}\) defines the mathematical model. Generally, the forward problem associated with Eq. (3.5) would be to solve the system \(\mathbf{K}_d \mathbf{d} = \mathbf{R}\) given a set \(\boldsymbol{\theta}\). In the inverse problem associated with Eq. (3.5), the matrix \(\mathbf{G}\) encapsulates both the model and the data. The discretization process leads inevitably to modeling errors in forming the system, whereas measurement errors in the data affect the solution of the inverse problem. These two phenomena lead to the ill-posedness of the inverse problem for Eq. (3.5), therefore indicating the need for a regularization procedure. Eq. (3.5) is typical to problems in many engineering areas, including biomedical and seismic imaging, digital tomography and digital image reconstruction. A detailed analysis of regularization techniques and
solution methods for such inverse problems can be found in (Aster, Borchers et al., 2005).

In this study, a Tikhonov-Total Variation (TTV) regularization scheme is used to solve Eq. (3.5) (Aster, Borchers et al., 2005). The problem is treated in the least-square sense augmented with the TTV regularization, leading to:

$$\min_{\theta} \frac{1}{2} \| G\theta - R \|^2 + \alpha \Phi(\theta)$$

(3.6)

where $\alpha$ is the Tikhonov parameter controlling the relative weight between the least squares and the regularization term $\Phi(\theta)$. In this thesis, the value of this parameter chosen is $10^{-4}$, which is found to be a sub-optimal choice in most practical applications. A better value can be chosen for this parameter is possible using selecting methods such as the L-curve method (Hansen, 1992) (Hansen, 1998). The function $\Phi(\theta)$ represents the total variation (TV) functional of the vector $\theta$. Minimization of the regularized TTV functional yields an efficient scheme to solve the discrete inverse problem. This type of regularization penalizes highly oscillatory solutions while allowing jumps-like discontinuities. Vogel (Vogel, 2002) showed that in the case of two-dimensional image deblurring, TTV regularization tends to produce qualitatively correct reconstructions of blocky images (Dobson, 1996).

The accuracy of the regularized solution depends very much on the smoothness of the true model. If $m_{\text{true}}$ is not smooth, then Tikhonov regularization solution simply will not give an accurate solution.
To obtain the numerical scheme for calculating $\Phi(\theta)$, consider the following general definition of the total variation (TV) of a function $f(x)$ on the interval $[0, 1]$: 

$$TV(f) \equiv \sup \sum_i |f(x_i) - f(x_{i-1})|$$

(3.7)

where the index, $i$, is taken over all partitions ($0 = x_0 < x_1 < \cdots < x_n = 1$) of the interval. In the one-dimensional case, the smooth form of TV functional defines the total variance function $\Phi$ as:

$$\Phi(f) = \int_0^L \sqrt{\left(\frac{df}{dx}\right)^2} \, dx$$

(3.8)

This equation is a continuous form of the discrete total variation that includes the sum of the magnitude of the jumps for $f$. However, due to the non-differentiability of the Euclidean norm at the origin, special attention is required for numerical implementation. One can select a smooth approximation to the Euclidean norm, $|x|$, such as $\sqrt{|x|^2 + \beta^2}$, where $\beta$ is a small positive parameter giving rise to the following smooth total variance functional:

$$\Phi(\theta(x)) = \int_0^L \sqrt{\left(\frac{d\theta}{dx}\right)^2 + \beta^2} \, dx$$

(3.9)

To derive the discrete form of the TTV term, let $\Delta x$ be the interval distance used to discretize $\theta(x)$ (i.e. the length of a beam element in our case), and assume that the number of elements is $nel$ (i.e. the index of $\theta$ ranging from 1 to $nel$); then, the discretized version of Eq. (3.9) can be written as:

$$\Phi(\theta) = \sum_{i=2}^{n} \psi((\Delta \theta)^2) \Delta x, \quad \text{and} \quad \psi((\Delta \theta)^2) = \sqrt{\left(\frac{\theta_i - \theta_{i-1}}{\Delta x}\right)^2 + \beta^2}$$

(3.10)
With this discretization of $\Phi(\theta)$, the regularized least-square problem shown in Eq. (3.5) can be solved using a primal-dual Newton method (Chan, Golub et al., 1996) (Vogel, 2002).

Within the present framework, both statically determinate and indeterminate beams can be analyzed with this formulation as long as the applied load and displacement measurements are available. It is also worth mentioning that the damage identification can be restricted to a given segment of the beam, and that the search area does not need to enclose the entire beam.

### 3.4 Data discrepancy-based FE model updating

Using a measured deflection of a beam, $u^m$, the problem of reconstructing the stiffness of a beam can be formulated as an optimization problem using a misfit function between the measurements and the model-based $u(\theta)$ displacements:

$$\begin{align}\min_{\theta} J(u, \theta) &= \frac{1}{2} \sum_{i=1}^{N} \left| u_i(\theta) - u^m_i \right|^2 + \alpha \Phi(\theta) \\
\text{such that. } K(\theta)u &= f \end{align}$$

(3.11)

where $J$ is the data-discrepancy functional augmented with a TTV functional similar to that in Equation (3.10). $N$ is the total number of measured points along the beam, $\theta$ is the vector collection of the unknown damage parameters, the vector $u^m$ is the measured displacement data and $u(\theta)$ is the set of simulated results corresponding to a set of distributed parameters $\theta$. The constraint equations appearing in the minimization problem expressed in Eq. (3.11) is the static finite element direct problem corresponding
to a set of parameters $\theta$ and a given load vector $f$. Solution of Eq.(16) leads to the expression of a FE model updating problem.

To solve a minimization problem efficiently, one needs to evaluate the gradient of the cost functional, $J$, which is constrained by the static equilibrium FE problem (Eq. 3.11). An adjoint formulation is used to compute the gradient vector of the cost functional to the unknown stiffness distribution parameters, $\theta$. The adjoint method allows computation of the gradient of a cost functional with a large number of input parameters. The constraint equations are the equilibrium-governing equations of the discretized beam using FE, and are written in the following residual form:

$$R(\theta, u(\theta)) = K(\theta)u - f = 0$$

(3.12)

where $u$ is the state variable (representing displacements in the present case), and the residual is an implicit function of the unknown internal variables $\theta$. Introduction of the Lagrangian multipliers, $\lambda$, converts the constrained problem into an unconstrained optimization. The augmented functional that enforces the governing equations is expressed as:

$$L(\theta, u) = J(\theta, u) - \lambda^T R(\theta, u)$$

(3.13)

Differentiating the Lagrangian with respect to $\theta_i$, gives:

$$\frac{dL(\theta, u)}{d\theta_i} = \frac{\partial J}{\partial \theta_i} + \left(\frac{\partial J}{\partial u} - \lambda^T \frac{\partial R}{\partial u}\right) \frac{du}{d\theta_i} - \lambda^T \frac{\partial R}{\partial \theta_i}$$

(3.14)

Usually, the derivative of the displacement with respect to the internal parameters, $du/d\theta_i$, is difficult to evaluate, and is not directly accessible. However, with a suitable
choice of \( \lambda \), it is possible to make the term in the bracket, \( \left( \frac{\partial J}{\partial u} - \lambda^T \frac{\partial R}{\partial u} \right) \), equal zero, thus avoiding the need to evaluate the gradient \( d\mathbf{u}/d\theta_i \). Therefore, an additional equation can be written as:

\[
\left( \frac{\partial J}{\partial u} - \lambda^T \frac{\partial R}{\partial u} \right) = 0
\]  

(3.15)

Equation (3.15) is the adjoint equation, which is solved for \( \lambda^T \); this equation reduces to the following linear system:

\[
(u - u^m) = \lambda^T K(\theta)
\]  

(3.16)

where \( u \) is the displacement vector computed from the FE model in Eq.(16), and \( u^m \) is the corresponding measured displacement vector. Therefore, Eq. (3.14) is simplified to:

\[
\frac{dL(\theta, u)}{d\theta_i} = \frac{\partial J}{\partial \theta_i} - \lambda^T \frac{\partial R}{\partial \theta_i}
\]  

(3.17)

in which the two derivative terms can be easily computed as:

\[
\frac{\partial J}{\partial \theta} = \alpha \frac{\partial \Phi}{\partial \theta} \quad \text{and} \quad \frac{\partial R}{\partial \theta_i} = \frac{\partial K(\theta)}{\partial \theta_i} u - \frac{\partial f}{\partial \theta_i}
\]  

(3.18)

The regularized TV-functional \( \Phi(\theta) \) is defined in the previous section (see Eq. 3.9 and Eq. 3.10). In order to calculate the gradient of the TV term \( J \), \( \frac{\partial J}{\partial \theta} = \alpha \frac{\partial \Phi}{\partial \theta} \), it is easy to deduce from Eq.(3.10) that

\[
\frac{\partial \Phi}{\partial \theta} = \Delta x \left( \Delta^T diag \left( \Psi'(\theta) \right) \Delta \right) \theta
\]  

(3.19)
Δx is the size of the beam element, and the term in the brackets, 
Δ^T diag\left(\Psi'(\theta)\right)Δ, is a positive, semi-definite, symmetric matrix. The Δ matrix has 
(n−1) rows and (n) columns, with the i^{th} row being Δ_i (see Eq.11). The diag\left(\Psi'(\theta)\right) matrix is a (n−1)×(n−1) diagonal matrix with the i^{th} diagonal entry equal to \Psi'((Δ,θ)^2).

The global stiffness matrix \(K\) is the classical assembly of the elements’ stiffness matrices \(k_i\):

\[ K = \bigwedge_{i=1}^{N_{el}} (1-D_i)k_i \]  

(3.20)

where \(\bigwedge\) is the FE assembly operator, and \(N_{el}\) is the number of elements in the FE model. As stated earlier, we assume that for each element, the damage is represented by a scalar parameter, \(D_i\), in the interval [0,1]. Therefore, the derivative of the stiffness matrix with respect to \(\theta\) can be computed analytically given a specific beam element formulation. For the Euler-Bernoulli bending element the following expression holds:

\[ \frac{\partial K(\theta)}{\partial \theta_i} = \bigwedge_{j=1}^{N_{el}} \frac{\partial k_j(\theta_j)}{\partial \theta_i} = \bigwedge_{j=1}^{N_{el}} k_j \]  

(3.21)

In conclusion, the adjoint method leads to an efficient computational method to calculate the gradient of the cost functional and consists of two consecutive steps: 1) solve the adjoint equation (3.17) for \(\lambda'\), then insert \(\lambda'\) into Eq. (3.18) to calculate the gradient of the Lagrangian. With the gradient calculated, any efficient gradient-based optimization technique can be used to find the optimal solution. A procedure based on the BFGS pseudo-Newton optimization technique is summarized in Table-2. The algorithm
shown in Table-1 is general, and can be applied to other discretization schemes, providing that the governing relation, \( R(\theta, u(\theta)) = K(\theta)u - f = 0 \), holds. Numerical experiments show that TTV regularization produces better results than the classical Tikhonov regularization method for this particular type of inverse problem.

Table 3.2: Damage identification algorithm using the adjoint-optimization method.

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Perform test (image acquisition and data collection);</td>
</tr>
<tr>
<td>2.</td>
<td>Choose an initial distribution of ( \theta^0(x) ), usually starting with ( D_j = 0 );</td>
</tr>
<tr>
<td>3.</td>
<td>Set ( k = 0 ), ( H_0 = I );</td>
</tr>
<tr>
<td>3.1</td>
<td>Finite element simulation of ( u(x) ) with given ( \theta^k );</td>
</tr>
<tr>
<td>3.2</td>
<td>Solve the adjoint equation (3.17) for ( \lambda^T );</td>
</tr>
<tr>
<td>3.3</td>
<td>Calculate the Lagrangian ( L ) and its gradient ( \nabla L );</td>
</tr>
<tr>
<td>3.4</td>
<td>Determine a scalar value ( \zeta ) through line-search along the direction ( -H_k \nabla L ) so that ( L(\theta^{k+1}) = L(\theta^k) + \zeta \nabla L ) and ( L(\theta^{k+1}) &lt; L(\theta^k) ).</td>
</tr>
<tr>
<td>3.5</td>
<td>Update the stiffness distribution, and compute: ( s_k = \theta^{k+1} - \theta^k ), and ( y_k = \nabla L(\theta^{k+1}) - \nabla L(\theta^k) );</td>
</tr>
<tr>
<td>3.6</td>
<td>( k = k + 1 ); and update ( H_k )</td>
</tr>
<tr>
<td></td>
<td>( H_k = H_{k-1} + \frac{(y_{k-1} - H_{k-1}s_{k-1})y_{k-1}^T}{y_{k-1}^Ts_{k-1}} - \frac{(y_{k-1} - H_{k-1}s_{k-1})^T}{(y_{k-1}^Ts_{k-1})^2} y_{k-1}y_{k-1}^T );</td>
</tr>
<tr>
<td>3.7</td>
<td>Check convergence;</td>
</tr>
<tr>
<td>4.</td>
<td>Solution = best point found.</td>
</tr>
</tbody>
</table>


3.5 Examples

3.5.1 Numerical examples

3.5.1.1 Validation of the Equilibrium Gap Method

To validate the proposed algorithms, we start with an ideal case where the “experimental” data are synthetic and obtained from a direct problem using a FE solution. The objective is to validate the proposed algorithms for locating discrete cracks and reconstructing the stiffness distribution along a beam. In this first validation attempt, we seek to validate the equilibrium gap-based method for a beam with two localized, damaged sections. A simply supported 10 m long beam with a constant stiffness is subjected to a concentrated load of 5 kN at the mid-span. The stiffness of the two elements at locations \( x_1 = 2.5 \text{ m} \) and \( x_2 = 5 \text{ m} \) are reduced by 50\% and 30\% (\( D = 0.5 \) and \( D = 0.3 \)), respectively. The deflection of the FE model is used as input for the equilibrium gap method to identify the damage. The beam is discretized with 100 elements, and therefore the stiffness distribution function is represented by 100 discrete unknown parameters. A Gaussian noise is added to the simulated response to emulate real measured signals as expressed by:

\[
noise(x) = NRND \cdot a \cdot RMS(u)
\]  

(3.22)

where \( NRND \) is a Gaussian random distribution with zero-mean and a unit standard deviation; \( a \) is the applied noise level; and \( RMS(u) \) is the root-mean-square of the measured displacement \( u(x) \). The problem is solved using a Primal-Dual method (Vogel, 2002).
Figure 3.2-(a) shows the damage pattern when no noise is added to the deflection measurements; in this case the two damaged zones (cracks) and damage levels are identified with very good accuracy. In Figures 3.2-(b) and 3.2-(c), noise levels of 2.5% and 5% are added to the measured deflection data, respectively. The numerical simulation is capable of identifying the positions and levels of the damage under these noise conditions. However, it is observed that the noise added to the synthetic measurements can mask the damage at the fine-scale level. If the noise level is known, the minimum scale of detectable damage corresponding to this level of noise can be approximately estimated by inspecting the variations of the identified damage indices. For example, at a 5% noise level, the finest detectable level of damage seems to be 10%, as it can be seen from inspection of Figure 3.2-(c).

To demonstrate the effect of noise in masking fine-scale damage, the beam is re-simulated using a different damage scenario, the beam’s stiffness is reduced by 30% and 10% at two locations, 2.5 m and 6 m from the left support, (i.e. \( D(2.5m) = 0.3 \) and \( D(6m) = 0.1 \)). A 5 % level of noise is added, and the identified damage indices are shown in Figure 3.2-(d), in which only the damage at location 2.5 m \( (D(2.5m) = 0.3) \) is identifiable; damage at the 6 m location is indistinguishable from the noise.
Figure 3.2: Identified damage indices of a simply-supported beam using the equilibrium gap method. (a): no noise, (b): 2.5% noise, (c): 5% noise, (d): 5% noise.

3.5.1.2 Validation of the Data Discrepancy Functional Method

The previously described problem is solved again using the data discrepancy functional method and the results are shown in Figure-3.3. Both the equilibrium gap and data discrepancy functional methods are in good agreement with the baseline solution.
In a second example, a 10 m long, simply-supported beam is simulated to validate the data discrepancy functional method. The beam is assumed to have a constant stiffness; a concentrated load of 5 kN is applied at the mid-span. The beam is discretized using 100 beam elements; hence, the continuous stiffness distribution function is represented by 100 discrete unknown stiffness parameters. A FE simulation is used to calculate the deflection profile. A Gaussian noise level of 5% is added to the synthetic displacement field that is to be used as input for the identification.

For the first validation test, we try to recover the stiffness of a beam with constant stiffness. The initial estimate is based on a uniform stiffness factor of 0.5 (i.e. half of the true stiffness value). The results from the application of the adjoint optimization method are illustrated in Figure 3.4. In the second test case, the beam is assumed to have a continuous parabolic stiffness distribution. Again, a constant stiffness factor of 0.5 is
used to initiate the iteration process. The results of the data discrepancy-based method in comparison to the reference stiffness distribution are shown in Figure 3.5.

The results shown in Figures 3.4 and 3.5 illustrate the performance of the data discrepancy formulation, and show that, at a distance from the supports, the identification is in good agreement with the expected stiffness values. The displacements of a simply supported beam close to the support are very small (zero at the support), and the lack of information in this region is not capable to improve the initial guess locally. This explains the drift between the expected results and the results from the simulation near the ends of the beam.

Figure 3.4: Identified stiffness factor distribution for a simply-supported beam with constant stiffness using the data discrepancy functional method
3.5.2 Experimental examples

To demonstrate the performance of the two proposed methods, we propose a series of experimental tests on a cantilever beam. An aluminum cantilever beam with four pre-defined damage locations is tested and the associated inverse problems are solved for validation. The beam’s cross-section is a HSS of $1 \times 1$ inch, and the thickness of the wall section plate is 1.58 mm. The material modulus of elasticity is measured from the static deflection of undamaged beam and it is found to be close to 72GPa.

In all four cases, the deflection profile is obtained using a close range photogrammetric method presented earlier in the present paper. The reference frame used to measure the deformation was the image of the beam under its self-weight. Although in the deformation under the self weight is negligible in the present case, this choice allows us to exclude the deformation of the beam under its own dead load. The stiffness distribution of the beam from the fixed base to the loading point is to be

![Figure 3.5: Identified stiffness factor distribution for a simply-supported beam with parabolic stiffness variation.](image-url)
identified, while the stiffness distribution in the region of the free tip beyond the loading is not taken into consideration, since no information is available in that region.

3.5.2.1 Test # 1: Step-wise damage detection

The first test setup is shown in Figure 3.6. The total length of the beam is \( L = 130 \text{ cm} \). A concentrated load of 21 \( N \) is applied at a location 90 cm away from the fixed end. To induce a predefined damage, the bottom face of the HSS section is cut out between 50 cm and 70 cm from the fixed end as illustrated. The stiffness in the damaged region is reduced to 0.83\% of the original cross-section \( (D = 0.17) \).

![Figure 3.6. The cantilever beam with a distributed damage.](image)

The identified damage variable profile using the equilibrium gap method is shown in Figure 3.7, it is clear that the results reproduce the expected damage location and level with very good accuracy. The recovered stiffness factor of each element using the data discrepancy method and reference values are plotted in Figure 3.8. The step damage is clearly visible despite the oscillatory nature of the identified stiffness distribution; the amplitudes of oscillations are generally less than 5\%, with exception of regions near the beam’s ends. The larger errors in the two ends can be attributed to the measurement
errors of the displacement close to the loading point and the lack of information close to the fixed end (theoretical displacements close to zero).

Figure 3.7. Damage detection of the cantilever beam with distributed damage using the equilibrium gap method.

Figure 3.8. Identified stiffness factor distribution of the cantilever beam with distributed damage using the data discrepancy formulation.
Figure 3.9 illustrates the performance of the adjoint based optimization algorithm for solving the inverse problem formulated using the data discrepancy functional. Figure 3.9(a) shows the convergence of the cost function minimization, whereas Figure 3.9(b) illustrates that at convergence, the solution is independent of the initial guess of the beam’s stiffness. The intermediate results during the iteration processes are shown in Figure 3.9(c) and 3.9(d) as they illustrate the convergence process.
3.5.2.2 Test # 2: Single saw-cut damaged beam

In this example, a single saw-cut was created on the tested beam to simulate damage in the form of a crack, as shown in Figure 3.10. The length of the beam is $L = 95.5\, \text{cm}$, and a concentrated load of $17\, \text{N}$ is applied at a distance of $89\, \text{cm}$ away from the support. The location of the saw-cut is $50\, \text{cm}$ from the base; the cut is $1.5\, \text{mm}$ wide and $15\, \text{mm}$ in depth, corresponding to a $86\%$ reduction in stiffness ($D = 0.86$).
Using the equilibrium gap method, the damage distribution along the beam is shown in Figure 3.11. The sharp peak is captured by the solution, indicating a clearly concentrated damage. The recovered stiffness factor profile using the data discrepancy method is plotted in Figure 3.12. The identification of the single concentrated damage is also clear, however small amplitudes oscillations similar to test 1 are observed. The larger errors close to the free-end can be attributed to the errors in the displacement measurement close to the loading point.
3.5.2.3 Test # 3: Discrete three saw-cuts damaged beam

The objective of this test is to check the capability of the two algorithms to detect multiple cracks. Three discrete saw-cuts were created; the beam dimension is shown in Figure 3.13. The experimental data are as follows: the length of the beam is $L = 95.5 \, \text{cm}$, and the concentrated load of $17 \, \text{N}$ is applied at $89 \, \text{cm}$ from the fixed end. The first cut is
30.6 cm away from the base, with a depth of 1.5 mm, corresponding to 17% stiffness reduction; the second cut is located at 50 cm away from the base, having a depth of 15 mm, corresponding to a 86% reduction in stiffness; the third cut is at 70 cm from the base, and is of 2 mm in depth, corresponding to 20% stiffness reduction. All introduced cuts have a width of 1.5 mm.

![Figure 3.13. The cantilever beam with three saw-cuts damage.](image)

The results of the equilibrium gap and the data discrepancy methods are illustrated in Figures 3.14 and 3.15, respectively. The sharp peaks are clear in the identified damage variables obtained by the equilibrium gap method (Figure 3.14), the identification of the location and magnitude of the damage is in good agreement with the expected results. From the stiffness factors obtained using the data discrepancy method, shown in Figure 3.15, it can be seen that although the location and magnitude of the damage have been captured, oscillations of the solution are still visible.
3.5.2.4 Test #4: Combination of discrete and step cuts damaged beam

In this example, a step-cut damage was added to the previously discussed beam; a step-cut is added between the second and the third saw-cuts (Figure 3.16). The depth of
the step cut is 2 \text{ mm}, equal to the depth of the original third saw-cut at the location of 70 cm from the base; thus, the third saw-cut becomes part of the step-cut, but the second cut still displays as a discrete cut attached to the continuous step material removal.

![Figure 3.16. Cantilever beam with double saw-cuts and a distributed damage.](image)

The identified damage indices and stiffness factors, using the two proposed methods, are shown in Figure 3.17 and Figure 3.18. The results of the equilibrium gap method show two sharp peaks and the flat stiffness region of the step damage are clearly identified. The overall predictions are consistent with expected damage location and levels. We observe an error close to the location of the loading larger than in the rest of the beam, this error can be attributed to the measurement variability in this region and the influence of the concentrated load. On the other hand, the results of the data discrepancy method display a similar trend of oscillations as was observed in the previous example; however, the damage locations and magnitude are easily identifiable.
Figure 3.17. Damage detection of the beam with saw-cuts and step damage using the equilibrium gap method.

Figure 3.18. Identified stiffness factor distribution of the beam with saw-cuts and step damage using the data discrepancy formulation.

The identified damage indices and stiffness factors of several critical sections in the four test cases are transformed to stiffness ($EI$) and are displayed in Tables 3.3 to 3.6. Note that the identified values of saw-cuts are taken as the peak value around the
location, but the identified step-cut or original beam section stiffness values are taken as average of five values around the approximate location since there can be some oscillations in the identified stiffness of a constant section, especially for the data-discrepancy formulation.

Table 3.3: Identified stiffness of test #1

<table>
<thead>
<tr>
<th>Stiffness (unit: $10^8$ MPa · mm$^4$)</th>
<th>$E1_1$</th>
<th>$E1_2$</th>
<th>$E1_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expected values</td>
<td>5.5259</td>
<td>4.5865</td>
<td>5.5259</td>
</tr>
<tr>
<td>Equilibrium gap</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Identified values</td>
<td>5.5143</td>
<td>4.4533</td>
<td>5.4651</td>
</tr>
<tr>
<td>Errors</td>
<td>-0.2%</td>
<td>-2.9%</td>
<td>-1.1%</td>
</tr>
<tr>
<td>Data discrepancy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Identified values</td>
<td>5.4485</td>
<td>4.3655</td>
<td>5.4430</td>
</tr>
<tr>
<td>Errors</td>
<td>-1.4%</td>
<td>-4.8%</td>
<td>-1.5%</td>
</tr>
</tbody>
</table>
Table 3.4: Identified stiffness of test #2

<table>
<thead>
<tr>
<th>Stiffness (unit: $10^8 \text{MPa} \cdot \text{mm}^4$)</th>
<th>$EI_1$</th>
<th>$EI_2$</th>
<th>$EI_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expected values</td>
<td>5.5259</td>
<td>0.7737</td>
<td>5.5259</td>
</tr>
<tr>
<td>Equilibrium gap</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Identified values</td>
<td>5.5093</td>
<td>0.7644</td>
<td>5.4430</td>
</tr>
<tr>
<td>Errors</td>
<td>-0.003</td>
<td>-1.2%</td>
<td>-1.5%</td>
</tr>
<tr>
<td>Data discrepancy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Identified values</td>
<td>5.6088</td>
<td>2.4739</td>
<td>5.3049</td>
</tr>
<tr>
<td>Errors</td>
<td>1.5%</td>
<td>219%</td>
<td>-4.0%</td>
</tr>
</tbody>
</table>


Table 3.5: Identified stiffness of test #3

<table>
<thead>
<tr>
<th>Stiffness (unit: $10^8 \text{ MPa} \cdot \text{mm}^4$)</th>
<th>$E_1$</th>
<th>$E_{I2}$</th>
<th>$E_{I3}$</th>
<th>$E_{I4}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expected values</td>
<td>4.5865</td>
<td>0.7737</td>
<td>5.5259</td>
<td>4.4207</td>
</tr>
<tr>
<td>Equilibrium gap</td>
<td>Identified values</td>
<td>4.3577</td>
<td>0.7710</td>
<td>5.4706</td>
</tr>
<tr>
<td>Errors</td>
<td>-5.0%</td>
<td>-0.35%</td>
<td>-1.0%</td>
<td>0.4%</td>
</tr>
<tr>
<td>Data discrepancy</td>
<td>Identified values</td>
<td>4.4041</td>
<td>1.6528</td>
<td>5.1943</td>
</tr>
<tr>
<td>Errors</td>
<td>-4.0%</td>
<td>113.6%</td>
<td>-6%</td>
<td>-6.9%</td>
</tr>
</tbody>
</table>

Table 3.6: Identified stiffness of test #4

<table>
<thead>
<tr>
<th>Stiffness (unit: $10^8 \text{ MPa} \cdot \text{mm}^4$)</th>
<th>$E_1$</th>
<th>$E_{I2}$</th>
<th>$E_{I3}$</th>
<th>$E_{I4}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expected values</td>
<td>4.5865</td>
<td>0.7737</td>
<td>4.4207</td>
<td>5.5259</td>
</tr>
<tr>
<td>Equilibrium gap</td>
<td>Identified values</td>
<td>4.3649</td>
<td>0.7869</td>
<td>4.6141</td>
</tr>
<tr>
<td>Errors</td>
<td>-4.8%</td>
<td>1.7%</td>
<td>4.4%</td>
<td>-2.0%</td>
</tr>
<tr>
<td>Data discrepancy</td>
<td>Identified values</td>
<td>4.3384</td>
<td>1.3964</td>
<td>4.5865</td>
</tr>
<tr>
<td>Errors</td>
<td>-5.4%</td>
<td>80.5%</td>
<td>3.8%</td>
<td>-5.0%</td>
</tr>
</tbody>
</table>
3.6 Conclusions

Two inverse problem based formulations are proposed to identify damage in Euler-Bernouilli beams using the equilibrium gap concept and a data discrepancy functional. The static deflection profile of the beam is obtained from close range photogrammetric technique. A quasi-continuous measurement of the deflection became possible with the use of digital image processing based technology. We describe a technique to measure the displacement of beams based on an edge detection algorithm. The first inverse problem formulation uses the equilibrium gap principle along with a finite element forward problem solver. An over-determinate algebraic system is obtained and solved in the least squares sense with a TTV regularization scheme. The second formulation is based on a data discrepancy expression of the measured and model based deflection. The minimization of the functional is obtained through an adjoint method and a TTV regularization.

The proposed methodology is validated by a series of synthetic data generated from simulations of a damaged beam. The identification of the location and level of damage is verified and the effect of noise is reported. The reconstruction of a continuously varying stiffness beam is also validated. Four tests were conducted on beams with different damage scenarios. The two methodologies were validated and showed overall good performance within a laboratory conditions. In each test, the location and level of damage were identified with a good level of accuracy. However, the formulation based on the equilibrium gap functional performed better and showed closer solutions to the real damage state. The data discrepancy based formulation depicts a slight oscillatory phenomenon in the solution, but the results are acceptable from a
practical point of view. A validation on more complex systems, such as rehabilitated reinforced concrete beams is underway.

As for any inverse problem formulation, the quality of the data measurements is essential to the performance of the algorithm. Two aspects need to be addressed in future work: (i) develop a reliable methodology for deflection measurements using digital image correlation, and (ii) include error uncertainty in the formulation of the inverse problems. These two enhancements will allow practical use of the proposed methodologies for the health monitoring of structures.

The proposed methodology deals with two broad activities: (i) periodic non-destructive damage localization and severity estimation; and (ii) the assessment of structural safety based on the results of the non-destructive damage detection.

Once the identification of the structure has been accomplished, subsequent analyses, e.g. the determination of load capacity, rating, reliability and useful life determination of the structure can then be conducted.

3.7 Extension and envision:

Field testing and identification is different from lab setting. In the published results, some are validated using pure simulated data to small-scale structures (Banan, Banan et al., 1994b) (Sanayei, Imbaro et al., 1997) , some using test data in lab experiments on small scale structures; nonetheless, in-situ testing is challenging.

Experimental design, testing, acquisition of data, data processing and data quality assessment, all involve some difficulties. Acquiring a topologically correct a priori FE
model can also be a problem, boundary conditions are needed to be identified first before damage identification.

There is a long history of research in full-scale testing for assessment of highway bridge (Phares, Rolander et al., 2001) (Bakht and Jaeger, 1990). While for large and exotic bridges the SHM systems are more an academic performance, for smaller bridges the global response is more sensitive to defects, visual inspection is less frequent and SHM systems can make a real contribution (Heywood, Roberts et al., 2000).
CHAPTER FOUR: MATERIAL PARAMETER IDENTIFICATION USING FULL-FIELD MEASUREMENTS

Give me matter, and I will construct a world out of it. (Immanuel Kant)

4.1 Introduction and literature review of related research

4.1.1 General review

Modern design and performance evaluation requires realistic simulation of structural and material behaviours; the corresponding theories are classified as models of elasticity, plasticity, hyperelasticity, rate-dependent viscoelasticity/viscoplasticity, as well as additional models of continuum damage and fracture mechanics at material and structural levels. For background material about the different classes of inelastic constitutive models for isotropic materials and numerical solution schemes, we cite, for example: (Kojic and Bathe, 2005; Neto, Peric et al., 2008).

In today’s practice of inelastic FE analysis using commercial programs, extensive libraries of material models are available. Constitutive parameters associated with inelastic models are not necessarily available in standard material databases, but are essential input data for any finite element simulation. Therefore, unknown material parameters must be identified experimentally. Traditionally, the identification of material parameters is performed using standard testing procedures on regularly shaped cut-outs or machined specimens. In these tests, it is generally assumed that the mechanical fields are homogeneous, and that the experimental specimen gauge response can be associated
to the simulated material response through simple fitting of the analytical relationships (Bell, 1984).

However, even in the case of simple tests, obtaining a perfectly homogeneous deformation field is not certain. For example, the accurate estimation of plastic material properties from a simple tensile test can be difficult due to the non-uniform stress/strain distribution in the necking zone. Generally, it is not possible to determine the hardening parameters from direct measurement of specimen elongations. In order to calculate the stress accurately, the Bridgman correction needs to be applied, and this requires additional measurements of the contractions and curvature of the necking zone (Bridgman, 1952). To overcome these limitations, Rodic et al. used an inverse approach, referred to as the “error minimization concept”, to estimate the hardening parameters in a Nadai-type constitutive law (Rodic, Gresovnik et al., 1995). In this approach, the parameters are estimated by minimizing an error function between the experimental and predicted load-displacement responses. Consequently, the necking distortion present in the specimens is implicitly included in the analysis; this approach does not require additional measurements to be obtained from the necking zone as is required for application of the Bridgman correction. Rodic and Gresovnik developed an identification system based on the finite element code, “Elfen”, and an inverse program, “Inverse”, to solve a minimization problem (Rodic and Gresovnik, 1998). A similar method was adopted by Mahken and Stein, but they included the observed contours of the necking zone during the loading process as additional experimental information to calculate the elasto-plastic material parameters of a mild steel (Mahnken and Stein, 1997).
A technique called the “mixed numerical experimental” method parallels the error minimization concept, but the deformation mode does not need to be pre-determined, so general specimen geometries and loading conditions can be chosen (Hendricks, 1991; Meuwissen, 1998; Meuwissen, Oomens et al., 1998). For example, Meuwissen (1998) applied the mixed technique to identify the mechanical parameters of aluminum and steel alloy materials, where the specimens were plate with holes so the geometries were not standard and deformation was inhomogeneous. Measurements included displacement fields on the surface of specimens determined by the grid or grating method; the combination of Von Mises yield criterion with nonlinear hardening laws was validated. This family of identification procedures derives from formulation as an optimization problem to minimize the difference between the computed and the measured responses. Typically, these procedures employ a pseudo-Newton method to search for the minimum, and dedicated gradient computation schemes, such as direct differentiation or finite difference, are required (Meuwissen, 1998; Meuwissen, Oomens et al., 1998; Mahnken, 2004) (Springmann and Kuna, 2003; Springmann and Kuna, 2005) (Zentar, Hicher et al., 2001) (Mahnken, 2000) (Mahnken, 2002).

Subsequently, due to the rapid development of finite element analysis of nonlinear materials and optimization techniques that accompanied the extraordinary development of computer hardware and software in the last twenty years, this method has been studied and applied extensively to determine different material parameters in a variety of tests. The mixed numerical experimental technique has been examined under names such as the “inverse identification approach”, or the “simultaneous determination of material

In addition to gradient-based local deterministic methods, the application of evolutionary algorithms to solve optimization problems, and identify inelastic material parameters with finite element model updating based on uni-axial load-displacement data has been investigated (Hwang, Wu et al., 2010) (Müller and Hartmann, 1989; Furukawa and Yagawa, 1997) (Munoz-Rojas, Cardoso et al., 2010). For example, Dusunceli et al. proposed a formulation based on error minimization of the stress-strain curve from homogeneous uni-axial tension tests along with a genetic algorithm-based solver (Dusunceli, Colak et al., 2010).

Alternatively, artificial neural networks (ANN) have been used to determine material properties from load-displacement data (Aguir, BelHadjSalah et al., 2011) (Mahnken, 2004) (Lefik and Schrefler, 2002). The ANNs can be trained using FEA simulations to build a mapping from measurement data to material parameters. ANNs are tolerant of measurement errors. The ANN model is used as an alternative to the finite element calculations to evaluate the objective functions.

One may classify the different methods of correlating test data and model parameters as:

1) methods based on theoretical/analytical relationships (analytical fitting);
2) methods based on numerical simulations of the test (mixed-numerical experimental methods); and
3) methods based on empirical relationships, including empirical relationships using artificial neural networks.
The mixed-numerical experimental method provides not only more flexibility in test design, but also more accurate and complete assessment of constitutive models. For example, this method allows the design of experiments in which conditions are much closer to those of practical situations; this is especially important in mechanical component analysis since it minimizes the extrapolation of results obtained under testing conditions to practical conditions. Thus, the mixed numerical-experimental method is a perfect candidate for designing experiments that approximate real-world conditions. This is especially important if in-situ testing is required, in which the specimen is tested within the assembly.

Generally, forward simulation is performed with finite element analysis, so this inverse identification is actually a problem of finite element model updating (see Chapter 3 for discussion of FE model updating in damage identification). Since the 1980s, finite element model updating has emerged as an important aspect of the design of mechanical systems and civil structures, especially in the development of automotive and aerospace systems; it quickly became the most popular branch of model-updating (Keane and Nair, 2005; Arora, 2007).

4.1.2 Outline of the proposed methodology

The research presented in this chapter considers the problem of identification of material plasticity parameters. We developed an identification method based on the minimization of a least-squares error function between the inhomogeneous displacement fields measured by digital image correlation (DIC) and a finite element simulation under a given load. The material parameters are identified simultaneously by means of a direct
derivative-free optimization method that uses the finite element code as a black-box procedure.

The three components of this inverse identification procedure are: (i) nonlinear finite element analysis, (ii) direct optimization algorithm, and (iii) digital image correlation (DIC) test results. One of the fundamental aims of this work is to reformulate the algorithm to minimize the cost function, and thus remove the need to evaluate the gradient of the cost function. Derivative-free optimization methods allow us to avoid the tedious coding of numerical gradient computation associated with a material constitutive model. The second objective of this work is to exploit the use of a full-field measurement technique based on DIC as input. DIC is capable of capturing heterogeneous deformation fields and provides an excellent source of data for both deterministic and statistical analysis. Furthermore, both finite element analysis and DIC are now standard tools widely available in industry. Therefore, the proposed methodology can be easily adopted for practical use. Finally, this work employs statistical studies of nonlinear regression analysis in this inverse identification procedure to verify and validate the identified material parameters. DIC provides a large amount of data and thus allows the use of statistical inference; whereas homogeneous tests seldom provide enough data for statistical inference unless they are associated with random simulations (Seibert, Lehn et al., 2000a) (Harth, 2003) (Harth, Schwan et al., 2004) (Harth and Lehn, 2007).

Taken together, finite element analysis, optical full-field measurement techniques (such as DIC or advanced grid methods), and direct optimization techniques comprise a standard tool for the determination of material properties. Moreover, since these
components of the proposed inverse identification procedure (not only finite element analysis, but also the DIC technique and finite element model updating through direct optimization), have become commercialized, they are currently available for general industrial use.

4.1.3 The advantages of using DIC

Different optical measurement techniques have been used in material characterization studies. For example, an optical method called “silhouette analysis” was used to obtain the section profile of a cylindrical specimen during necking; these data, along with load-displacement curves, were used to identify parameters in a damage model (Broggiato, Campana et al., 2007). Likewise, the moiré method was used to obtain planar displacements (Kreißig, Benedix et al., 2007), while a grid method was used by Meuwissen and Mahken (Meuwissen, 1998) (Mahken, 1998). However, the commercially most successful image processing-based measurement technique is the digital image correlation (DIC) method (Louban, 2009) (see Chapter 2 for a list of current commercial DIC systems and software). Sutton et al. presented details of the DIC technique and a review of its applications in a recent book (Sutton, Orteu et al., 2009).

Many researchers have noted the advantages of using inhomogeneous displacement fields in material parameter identification (Kleuter, Menzel et al., 2007) (Cooreman, Lecompte et al., 2008). Recently, investigators in France developed a methodology based on the principle of virtual work, the “virtual fields method (VFM)”, that allows the development of simple or more complex analytical models for the constitutive equations of the material under investigation. The VFM assumes that the input will consist of full-field measurement data obtained using DIC or the grid method.
In addition, inhomogeneous displacement fields must be provided for the VFM; otherwise, the resulting linear system of equations becomes ill-conditioned as the data may not contain sufficient information for more than two parameters. Intuitively, although homogeneous displacement fields are adequate for identification of the elastic modulus, the deformation must be heterogeneous for the determination of elasto-plastic parameters. Section 4.8 presents further discussion of the issue of identifiability.

There is an additional advantage arises from the large amount of data generated by DIC: the valid use of statistical inference. For example, the estimate of the covariance is valid when the amount of data is large. On the other hand, unless used in conjunction with random simulation, homogeneous tests seldom provide enough data to support statistical inference (Seibert, Lehn et al., 2000a) (Harth, 2003) (Harth, Schwan et al., 2004) (Harth and Lehn, 2007).

Another significant advantage to using DIC is that the identification of parameters in material models can be performed with non-standard specimens and even with original components, so in-situ testing becomes possible. The use of non-standard specimens under non-standard loading conditions is not only an intellectual curiosity, but is particularly useful in some practical applications. For example, Partheepan et al used miniature specimens, thus avoiding the removal of large material samples, for the evaluation of current material properties of an in-service component (Partheepan, Sehgal et al., 2008).
4.1.4 Organization of this chapter

Chapter 4 is organized as follows: Section 4.2 outlines the formulation of continuous functions in solid mechanics and gives an overview of popular material constitutive models that are widely used in engineering finite element analysis. Section 4.3 outlines the approaches used in the current work, and describes the identification method in detail; including the least squares solution and the validation procedures based on theories in nonlinear regression analysis. Section 4.4 summarizes the relevant theories in nonlinear regression related to this work. Section 4.5 presents the inverse identification problem using nonlinear least squares estimation. Section 4.6 is an overview of state-of-the-art numerical optimization methods, with emphasis on direct derivative-free techniques developed in recent years. Section 4.7 describes the statistical inference procedures for the DIC test data; these procedures employ linearized covariance analysis (LCA) of uncertainty for the identified parameters and sampling-based statistical inferences of the response fit.

Identifiability issues are discussed in Section 4.8. Numerical and experimental examples are presented in Section 4.9; the first validation is based on synthetic data, and the experimental example deals with a cast iron bearing cap component, where the displacement field was obtained using the ARAMIS DIC system (ARAMIS). Four direct optimization methods are compared using these examples, including (i) the derivative-free CONDOR algorithm which is based upon the trust-region algorithm and response surface exploration techniques, (ii) the Nelder-Mead simplex method, (iii) a response surface method in FE software, and (iv) a pseudo-Newton method using finite difference approximation to gradients. The fourth method is usually considered as first-order,
gradient-based method with an approximate gradient evaluation; this type of methods have been used in material parameter identification. However, since in this type of methods, the FE simulation process can actually be treated as black-box, we consider it also as a derivative-free method; i.e. in this thesis the term “derivative-free” is associated with methods in which the objective-function evaluation process can be treated as an external black box. Section 4.10 summarizes the findings of this work.

4.2 Overview of elasto-plasticity

In this section, we summarize the general theory of elasto-plasticity. By definition, the theory of elasto-plasticity is concerned with solids that, after being subjected to a loading program, may sustain permanent (or plastic) deformations after complete unloading. Elasto-plasticity theory has been successfully applied to metals, and it is nowadays theoretically consolidated and well-studied phenomenological constitutive model.

The basic components of the general elasto-plastic constitutive model include:

- The strain decomposition principle where the total strain $\epsilon$ is assumed to be the sum of an elastic, $\epsilon_e$, and a plastic components, $\epsilon_p$: $\epsilon = \epsilon_e + \epsilon_p$.

- A yield criterion usually expressed in the stress space, $\Phi(\sigma)$, that defines the onset of plasticity. Classical yield criteria widely used in engineering practice include Tresca, von Mises, Mohr-Coulomb and Drucker-Prager criteria.

- A plastic flow rule defining the evolution of the plastic strain, the flow rule is usually defined as a flow potential. In the case of associative flow rule, the flow rule is described from the gradient of the yield function
which leads to the fact that the plastic strain rate is a tensor normal to the yield surface in the stress space.

- A hardening law, characterizing the evolution of the yield surface (i.e. deformation after yielding occurred). The hardening rule describes the evolution of the yield function over time, usually identified from a uniaxial specimen.

- To express the irreversibility of the plastic deformation, the Kuhn-Tucker conditions must be applied.

The identification of an associative elasto-plastic constitutive model consists of finding the parameters that define the yield function and the hardening surface. In FE analysis, the hardening law is generally expressed as a one-dimensional function describing the full stress-strain curve during uniaxial loading (Ramberg and Osgood, 1943) (ASTM, 2011).

For example, in the case of the linear hardening law: $\sigma = \sigma_0 + H \epsilon_p$, where $\sigma$ is the current yield stress, $\sigma_0$ is the initial yield stress, $\epsilon_p = \epsilon - \epsilon_e$ is the equivalent plastic strain, $\epsilon$ is the total strain and $\epsilon_e$ is the elastic strain, $H$ is the hardening modulus. In this simple model, the parameters that need determined are $\sigma_0$ and $H$. In most FEA programs, the data required for an elasto-plastic behaviour is a multi-linear function that is the best fitting of the stress-strain curve. There exist a variety of nonlinear hardening laws for metals including the Nadai hardening law: $\sigma = \sigma_0 + c \epsilon_p^n$, defined by the three parameters $\sigma_0$, $c$ and $n$.; the rigid-plastic power hardening law: $\sigma = c \epsilon_p^n$; and the Ramberg-Osgood
hardening law: \( \epsilon = \frac{\sigma}{E} + K \left( \frac{\sigma}{E} \right)^n \), where \( E \) is the Young’s modulus, and \( K \) and \( n \) are the two material parameters.

### 4.3 General methodology

Material parameter identification is a particular type of parameter identification problem. Parameter identification problems arise in all fields of engineering science. Consequently, parameter identification has attracted considerable research interest since the mid-seventies; a number of books describe standard treatment of parameter identification using deterministic and statistical methods (Bard, 1974) (Beck and Arnold, 1977) (Bates and Watts, 1988) (Tarantola, 2004) (Seber and Wild, 2005). In the current research, we focus on a formulation for the identification of constitutive parameters based on the least-squares cost function measuring the gap between a FE model prediction and the results of experimental measurements. The least-squares estimation is a fundamental approach to system identification problems, and has already become a classic method frequently practiced by scientists and engineers in a wide variety of applications.

The principle components of the inverse identification technique are presented schematically in Figure 4.1:
In this procedure, the post-processing phase performs a coordinate transformation and interpolation to correlate the coordinates used in the DIC experiment with those of the FE simulation results. When commercial FEA software is involved in the optimization process, the data flow between the optimization algorithm and the FEA code is maintained through file exchanges. This process requires the optimization algorithm to modify the input file iteratively and submit the modified data to the FEA software. Modern commercial design software packages developed for coupling with FE analysis (e.g. iSIGHT or HYPERSTUDY) have interfaces that simplify the data flow process.
The research presented in this chapter uses ANSYS software for the FE analyses. MATLAB programs implementing CONDOR and Nelder-Mead simplex algorithms are used as well, and the interaction with the ANSYS FE software is performed through modification of the ANSYS APDL modeling file. For purposes of comparison, a response surface methodology included in the ANSYS DesignSpace software package is also used; in this case, the interaction between the optimization algorithm and the FE code is internal to the software.

4.4 Nonlinear regression theory: parameter estimation using statistical inference

Due to unavoidable systematic inaccuracies inherent in loading, boundary conditions, and measurement devices, the recorded experimental data includes errors. Therefore, the sensitivity and inferences on the calculated parameters represents important information to the user regarding the precision of the combined testing/modeling procedures.

The parametric nonlinear regression model is written as: \( Y = f(x, \theta) + \varepsilon \); where the response vector \( Y \) is observed for each value of the independent variable \( x \), it is assumed that the true regression relationship between \( Y \) and \( x \) is a sum of a systematic (physical) part and a random part; generally, the true system function is approximated by a parametric function \( f \), called the regression function, that depends on unknown parameters \( \theta \). The function \( f \) does not need to be known explicitly, in many cases it is a function of the solution of differential equations. \( \varepsilon \) is a random error equal, by construction, to the discrepancy between observation \( Y \) and \( f \) .
The problem is to estimate the unknown parameter vector $\theta$. A natural and popular choice is the least squares estimator, and the LS estimator is also the maximum likelihood estimator (MLE) in the case of Gaussian observations.

The inverse identification process described in Section 4.3 can be interpreted as a statistical parameter estimator using nonlinear regression. Statistical inference can be employed to study the reliability of the parameter identification, and to provide an assessment of the quality of the results (Huet, Bouvier et al., 2004). In using statistical inference, a set of parameters ($\theta$) are considered as random variables; thus, their accuracy is related to the shape of the probability distribution function. Accuracy is often expressed in terms of a confidence region for the parameter vector, or as the confidence interval of its components. A $100(1-\alpha)\%$ confidence region is a region of parameter space that contains the true parameters with a probability of $1 - \alpha$; therefore, $1 - \alpha$ is the confidence level for this region (Huet, Bouvier et al., 2004).

There are three methods for statistical inference:

I. Likelihood approach,
II. Sampling theory approach,
III. Bayesian approach.

All three methods produce the same point estimates for $\theta$. They also produce similar confidence regions of reasonable parameter values.

In nonlinear regression inference, the success of parameter identification can be checked by: 1) expectation function fit and 2) parameter fit. If a model gives a smaller residual response and/or smaller confidence regions in the expectation function and
response function, the model is said to be phenomenologically better, or the model parameter set yields a better description of the model’s behaviour.

The most accurate way to perform a sensitivity analysis is through Monte Carlo methods in which properties of the distributions of random variables are investigated by use of simulated random numbers (Gentle, 2003). In the random simulation, scattered data from simulated identification experiments is collected, and analyzed statistically (Mosegaard and Sambridge, 2002). For example, researchers used stochastic search and Monte Carlo analyses to study viscoplasticity models; their test data were homogeneous responses and their models were ordinary differential equations (ODE) solved numerically using the Runge-Kutta method. This type of forward analysis is easy to solve, and computationally inexpensive, therefore stochastic techniques can be applied without difficulty (Seibert, Lehn et al., 2000a) (Harth, Schwan et al., 2004) (Harth and Lehn, 2007).

Nevertheless, due to the complex geometry and nonlinear FEA, we cannot recommend this approach for industrial applications. With current computing power, if optimization and validation require between 20 to 50 FE analyses, then this approach is affordable; however, if these processes require 5000 FE analyses, the computational cost is too high. For the same reason, global stochastic optimization methods are not applicable to this problem.

4.5 Least-squares formulation and solution of the identification problem

In the mixed numerical-experimental method using a least squares estimator, the set of parameters $\theta$ of the material model are determined by minimizing the difference
between a test data set \( d \) and numerically simulated results denoted \( M(x; \theta) \) in the sense of an error norm. Usually, a classical least squares objective function is used:

\[
\min f(\theta) = \frac{1}{2} \| M(x; \theta) - d \|_2^2 \quad M(x; \theta) \in \mathbb{R}^N \quad \theta \in \mathbb{R}^K \quad d \in \mathbb{R}^N \quad (4.1)
\]

Obviously, the objective function \( f(\theta) \) is a scalar function of the unknown material parameters \( \theta \); the numerically simulated response, referred to as model, \( M(x; \theta) \) is a function of the material parameters \( \theta \). Here, \( N \) is the number of measured response data and \( K \) is the number of parameters. For most practical purposes; and to keep generality, the response \( M(x; \theta) \) is assumed to be generated by finite element simulations. The variables defined in Equation (4.1) include: \( x \), the discrete space locations where the measurements are carried out and, \( \theta \) which denotes the space of material parameters \( \theta \) associated with a specific constitutive model; this space defines the practical constraints for each parameter in the set \( \theta \). In formulation (4.1) the \( L_2 \) norm is used; however, different norms can also be considered as it was shown previously that the identification results are barely affected by the choice of the norm (Seibert, Lehn et al., 2000b).

The cost function \( f(\theta) \) in (4.1) can be rewritten as:

\[
f(\theta) = \frac{1}{2} R^T(\theta) R(\theta) = \frac{1}{2} \sum_{i=1}^{N} R_i^2(\theta) \quad (4.2)
\]

where \( R(\theta) = \{R_i(\theta)\}^T, i = 1, ..., N, \) denotes a column vector of residual components between the calculated and measured values; and \( R_i(\theta) = M_i(\theta) - d_i \), where index \( i \) refers to a particular space location \( x_i \).

The necessary optimality condition, \( \nabla f(\theta^*) = 0 \), leads to the normal equation:

\[
f^T J \theta^* = J^T R \quad (4.3)
\]
where $J$ denotes the Jacobian matrix of the residuals with respect to the parameters; that is:

$$J_{ij} = \frac{\partial R_i}{\partial \theta_j} \quad (4.4)$$

If the model mapping $M(x; \theta)$ is linear, this normal equation leads to a direct solution of the unknown parameters $\theta$. In general, the mapping is implicit and nonlinear; therefore, an iterative procedure needs to be used.

Without loss of generality, the problem expressed by Equation (4.1) can be considered as finite element model updating through minimization of the cost function defined in (4.2). This general formulation is flexible as different types of experimental data can be combined. For example, DIC data and load-displacement data can be combined and input together, or DIC data at different loading stages and/or under different loading configurations can be defined as an input ensemble. In the present work, we used DIC measurements as input, but the solution methodology can be generalized to data acquired with different measurement techniques.

In practice, the data $d$ is unavoidably affected by measurement errors, and a weight can be applied to be cost function. Theoretically, the ideal weight used should be the inverse of the covariance matrix of the observed data. In the current study, we assumed that the noise from different sources of error is independent and identically distributed; the cost function with these ideal weights can be expressed as:

$$f(\theta) = \frac{1}{2} \sum_{i=1}^{N} \frac{R_i^2(\theta)}{\sigma_i^2} \quad (4.5)$$

where $\sigma_i^2$ is the expected error variance of measurement data $d_i$. Thus, the least squares estimate is identical to the maximum likelihood one.
From examination of currently available optimization algorithms, one may distinguish between schemes that make use of the gradient of the objective function, and those that do not rely on gradient estimations. Most of the existing mixed numerical-experimental approaches use gradient-based optimization methods, and the gradient is generally derived analytically using either direct differentiation or adjoint formulation techniques, or calculated numerically using finite difference. An iterative method is adopted along with the gradient information to solve the nonlinear least-squares problem (Kreißig, Benedix et al., 2007) (Mahnken, 2004) (Springmann and Kühhorn, 2009).

In this study, direct optimization methods were preferred to solve the least squares identification problem expressed by Equation (4.2). Note that the terms direct search, derivative-free search, and zero-th order search are sometimes used interchangeably in the literature. According to Conn et al, for state-of-the-art research and development, derivative-free optimization methods are best for dealing with noisy data; the performance of these methods is notably better than gradient-based methods using finite difference approximations, especially in cases of noisy data (Conn, Scheinberg et al., 2009).

Powell (1964) originally proposed a derivative-free version of the nonlinear conjugate gradient method consisting of the construction of a sequence of at most $K + 1$ one-dimensional searches ($K$ being the number of unknown parameters); each one-dimensional search is conducted by finding the exact minimum of a quadratic interpolant (Powell, 1964). The interpolant used in Powell’s method is similar to the response surface methodology (RSM) in stochastic optimization.
In Monte Carlo analyses, model parameters simulate real world conditions by varying randomly, whereas response surface methodology combines global and local analyses to create a simple mathematical equation of the complex and implicit relationship between different factors.

In the last twenty years, with the exception of a few heuristic-based searches and cluster-based random searches, direct optimization algorithms developed extensively primarily due to the efficient use of response surface exploration in combination with different efficient optimization, line-search and trust-region algorithms. Recently, considerable research progress in code development of derivative-free Newton-based methods has taken place, including DFO (Conn, Scheinberg et al., 1997), Powell’s UOBYQA (Powell, 2002), CONDOR (Berghen, 2004) (Berghen and Bersini, 2005), and NEWUOA (Powell, 2006).

Berghen developed CONDOR (COnstrained, Non-linear, Direct, parallel Optimization using trust Region method for high-computing load function) for expensive optimizations, such as objective functions evaluated from the output of nonlinear FEA or Computational Fluid Dynamics (CFD) simulations. CONDOR was designed mainly for unconstrained optimization problems and for easy (box) constraints. The algorithms in CONDOR are based on a derivative-free trust region method using a surrogate model that approximates the objective function by a quadratic polynomial, which is then minimized by sequential quadratic programming (SQP). Descent conditions from trust region methods enforce convergence properties. CONDOR has been compared to DFO (Ugur, Karasozan et al., 2008) and genetic algorithms (Harth, Sun et al., 2007) in designs involving flow problems, and its performance was found to be better.
In the present study, the CONDOR method was used to minimize the least squares cost function and to obtain the list of unknown parameters. For comparison, we also used the original Nelder-Mead simplex method, a pseudo-Newton method based on finite difference derivatives, and a response surface method to illustrate the superiority of a derivative-free algorithm in the context of parameter identification.

The use of derivative-free optimization methods provides the level of generality and modularity necessary for the optimization tool if the objective is to use the finite element code as a black box. Some of the modern derivative-free methods are considered globally convergent, since they include techniques such as line-search, reinforcing good geometry and trust region, thus ensuring a global optimum. Note, however, that because of the lack of rigorous convergence properties of derivative-free methods, the solutions must be considered local. One simple way to address this issue is to vary the starting point and the trust region radius to validate the final optimal solution.

In summary, for the solution of the optimization problem, the following recommendations are made:

- Use justification to give box-constraints to define the acceptable search region.
- If possible, have more than one method to solve the problem.
- Start from several different starting points.

### 4.6 Overview and selection of optimization techniques

The material parameter identification requires the solution of a nonlinear optimization problem:
minimize $f(\theta); \quad \theta \in R^K$

$s.t. \quad \theta_i \in [\theta_i^l, \theta_i^u], i = 1, ..., K$

where $\theta \in R^K$ is the vector of unknown parameters, and $\theta_i^l$ and $\theta_i^u$ are vectors of lower and upper bounds, respectively. Comparing to the general nonlinear optimization problem, there is no present of equality and inequality constraints.

The major concern is that the evaluation of $f(\theta)$ is computationally expensive. This fact rules out the application of stochastic optimization methods, such as evolutionary algorithms and other heuristic-based global optimization methods. Intuitively, the most straightforward way to tackle the computational cost issue is to use a strategy that employs computationally cheap surrogate models for the objective to solve an approximation to $f(\theta)$. Surrogate-assisted search algorithms are the choice.

They can be classified into zero-order methods, first order-methods, and second-order methods. The zero-order methods use only the objective function values to determine the optimum values. The first-order methods use the objective function and the gradient of the objective function to construct search directions during iterations. The second-order methods use second-derivatives (Hessian) to construct the search directions.

Nonlinear least squares problems can be solved in two ways: 1) as nonlinear least squares optimization problems or 2) as PDE-constrained optimization problems. In this study, only the first category of these solutions is considered. Various numerical solution methods for optimization problems are available. For example, one may distinguish schemes by whether or not the solution uses gradients of the objective function. In general, published accounts describing mixed-numerical-experimental approaches used
gradient-based optimization methods. Usually formulas to evaluate a gradient were
derived analytically, and an iterative method was adopted along with the gradient
information to solve the nonlinear least-squares problem (Kreißig, Benedix et al., 2007)
(Mahnken, 2004) (Springmann and Kühhorn, 2009). The most frequently used iterative
approaches for solving nonlinear least-square problems include the pseudo-Newton
methods, such as the Gauss-Newton method or the BFGS method, and trust region
methods, such as the Levenberg-Marquardt method. In these approaches, the Hessian
$\nabla^2 f(\theta)$ is not required, which is a great advantage over complete Newton methods since
derivation of analytical Hessian is more complicated than analytical gradients. A
summary of the fundamental aspects of gradient-based methods for solving the
optimization problem follows.

The iterative schemes can be expressed as:

$$
\theta^{k+1} = \theta^k - \alpha^j H^j \nabla f(\theta^j)
$$

(4.6)

where $\alpha^j$ is the step size at the $k^{th}$ iteration and $H^k$ is the pseudo-Newton iteration
matrix at the $k^{th}$ iteration.

The pseudo-Newtonian iteration matrices defined for each method are:

<table>
<thead>
<tr>
<th>Method</th>
<th>$H$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gauss-Newton</td>
<td>$(J^T J)^{-1}$</td>
</tr>
<tr>
<td>Levenberg-Marquardt</td>
<td>$(J^T J + \gamma I)^{-1}$</td>
</tr>
<tr>
<td>BFGS</td>
<td>$H(H^{-1}, \nabla f, \theta)$</td>
</tr>
</tbody>
</table>
Gradient methods are efficient and often require fewer forward simulations. In the case of practical applications where the data are associated with noise, the convergence rate of gradient-based methods depends closely on the initial estimate of the parameters. Apart from the problem of local minimum, the major difficulty in using gradient-based methods lies in the implementation of gradient-evaluation procedures. These are tedious and require a great deal of additional effort in addition to the FE analysis. Furthermore, users generally do not have access to commercial codes; this prevents users of complex commercial numerical analysis programs from obtaining reliable derivative information for the requested numerical analysis. Another hindrance is that the evaluation of gradients can vary for different material models, element types, deformation levels, integration schemes, or even the type of response data.

In the FE model updating and material identification literature, gradient evaluations are often referred to as sensitivity analyses. A sensitivity analysis using DIC-measured surface strains was reported by (Cooreman, Lecompte et al., 2007). This particular analysis applies only to simple tensile tests, however, and since the return-mapping scheme was considered in the derivation, it cannot be directly applied to FE programs using other schemes, such as the effective stress integration scheme in ADINA (ADINA). Some researchers have used finite difference approximation of the gradients as an alternative (Meuwissen, 1998) (Meuwissen, Oomens et al., 1998) (Haber, Tortorelli et al., 1993) (Springmann and Kuna, 2003) (Ghouati and Gelin, 2001), but this approach is considered unreliable, especially when the noise in the experimental data is not negligible (Springmann and Kuna, 2003) (Ghouati and Gelin, 2001) (Frąś, Nowak et al.,
Of course, the poor reputation of this method may stem from the fact that its simplicity is not persuasive for application to scholarly research.

Gradient methods are more efficient, and require fewer forward simulations. On the other hand, they are local methods; if the objective function is multi-modal (the multi-modality can be essential or caused by noise in the data), convergence is highly dependent on the initial estimates of the parameters. A local minimum solution is easily found; a global minimum may be found using multiple initial guesses.

There are also a variety of direct search methods based on heuristics, which are often without theoretical justification; these include random search, evolutionary algorithms, and taboo search. The most significant advantage of these methods is that they are generally global; however, they are also often slow. In other words, these methods are ‘slow, but sure’. Simulated annealing and genetic algorithms are among the leading candidates for global optimization applications. Nevertheless, these methods are also problem-specific, and require user-intervention and learning. Consequently, standard implementation of these methods has yet to coalesce. The cluster-oriented controlled random search method proposed by Price (Price, 1983) was used by (Harth and Lehn, 2007) to identify material parameters in Chaboche’s viscoplastic model for solution of an error minimization problem with data from homogeneous tension/compression tests and creep tests. Since the strain was homogeneous, the model was an ODE model solved using Runge-Kutta methods. The random search direct method requires extensive forward simulations, but is affordable for ODE models. However, if an FE model is used, and the model is geometrically complicated and nonlinear, the random search will become unaffordable with current computational resources.
In this paper, direct optimization methods are used to solve the least square identification problem. The phrase “direct search” was coined by Hooke and Jeeves in (Hooke and Jeeves, 1961); they proposed a pattern search method based on heuristics. The terms direct search, derivative-free search, and zero-th order search are sometimes used interchangeably, especially in finite element design software. There is no unanimity among researchers in their use of these terms. This is of no great consequence in practice, and to some extent simply reflects historical developments.

Conn et al. defined derivative-free methods as “methods without explicit approximation to derivatives of objective or constraints” (Conn, Scheinberg et al., 2009). However, optimization with finite difference derivatives is excluded from Conn et al.’s definition, and in our study, “derivative-free” means any method that treats the FEA forward solution as a black box. In this way, finite difference gradient-based pseudo-Newtonian methods can also be considered as derivative-free. Under this condition, the requirements from a user are minimal, since the simulation code is used in the manner of a black-box. According to (Conn, Scheinberg et al., 2009), with current state-of-the-art derivative-free optimization methods one can expect to successfully address problems: 1) which do not possess more than 100 variables, 2) problems which are reasonably smooth, and 3) problems in which the evaluation of the function is expensive and computed with noise.

There are four main classes of methods for designing a derivative-free method:

I. Coordinate search: these methods are slow, robust and capable of handling noise. The HYPERSTUDY/HYPERWORK program implemented a
simple version of this type of method, which is called “alternative-direction method” in the menu.

II. Nelder-Mean simplex method: the most popular method, but is not robust or reliable in some cases. The MATLAB program implemented an original version of this method in the Optimization toolbox (Nelder and Mead, 1965).

III. Implicit filtering algorithm: a line-search algorithm that imposes sufficient decrease along a quasi-Newton direction. The main difference from derivative-based methods is that the true gradient is replaced by the simplex gradient; so, this method resembles to some extent a pseudo-Newton method using finite difference approximation to gradients. This implementation of pseudo-Newton methods is deemed by Kelley to be particularly well-equipped to handle noisy functions since there is implicit filtering of noise due to the use of the simplex gradient corresponding to the gradient of a regression model and to an inaccurate line-search (Kelley, 1999). To our knowledge, this class of methods has not been implemented in any FE/design software.

IV. Interpolation-based trust region approach: trust region-based algorithms in which interpolation models are built from polynomial interpolation or regression. The interpolation is often called response surface and the term “response surface method” is frequently used in FEA and engineering design software (e.g. ANSYS includes several response surface methods). However, in the derivative-free optimization literature, the term “response
surface” is rarely used, and is generally reserved for experimental design and stochastic optimization.

Most recent progress in derivative-free optimization has focused on the fourth of these method categories. Powell’s celebrated method (1964) is a derivative-free version of the nonlinear conjugate gradient method. It consists in each stage of a sequence of \( K + 1 \) one-dimensional searches, and each one-dimensional search is conducted by finding the exact minimization of a quadratic interpolant. This method is thus a first-order method with a Q-property. The interpolant used in Powell’s method is actually the response surface methodology from stochastic optimization. Response surface is the idea that if the function evaluation is exact, one can use finite difference to approximate the derivative. If the function evaluation is uncertain, however, one can design an appropriate experiment and perform a regression analysis to obtain a surrogate of the original function, and thus estimate the derivative and update the search directions.

Powell’s work has spawned considerable research and code development of derivative-free Newtonian-based methods, including DFO (Conn, Scheinberg et al., 1997), Powell’s UOBYQA (Powell, 2002), and CONDOR (Berghen, 2004) (Berghen and Bersini, 2005), and recently, NEWUOA (Powell, 2006). Conn et al. (1997) constructed a multivariate DFO algorithm that uses a surrogate model for the objective function within a trust region method. In that work, points were sampled to obtain a well-defined quadratic interpolation model, and descent conditions from trust region methods enforced convergence properties. The trust region method can be used to globalize Newton-based methods and to avoid most of the local instability within Newton-based methods. The CONDOR method for high-computing load function developed by Berghen, is based on
the UOBYQA of Powell, and was developed mainly for unconstrained optimization problems and for easy (box) constraints. The algorithms are based on a derivative-free trust region method approximating the objective function by a quadratic polynomial, which is then minimized by a sequential quadratic programming (SQP) method. This method was developed for expensive optimizations, such as objective functions evaluated from the output of nonlinear FE or CFD simulations, and is based on the assumption that the function evaluation is time-consuming. Details of this method can be found in (Berghen, 2004) (Berghen and Bersini, 2005). DFO was applied as a black-box optimization routine in optimizing energy systems (Lee, Terlaky et al., 2001) and for helicopter rotor blade design (Scheinberg, 2000). Numerical tests in both of these reports show that DFO is faster and more accurate than derivative-based methods such as the quasi-Newton methods. However, CONDOR was also compared to DFO (Ugur, Karasozzen et al., 2008) and genetic algorithms (Harth, Sun et al., 2007) in designs involving flow problems, and its performance was found to be better. Several derivative-free methods are available in software or freeware; (Conn, Scheinberg et al., 2009) list a collection of freely-available derivative-free programs.

The present study used the CONDOR method to minimize the least squares cost function and to obtain the requested list of parameters. As mentioned above, this method is a trust region method combined with response surface technique that finds the minimum $\mathbf{x}^* \in \mathbb{R}^n$ of an objective function $F(\mathbf{x}) \in \mathbb{R}$. For the purpose of comparison, the original Nelder-Mead simplex method, a pseudo-Newtonian method based on the finite difference derivative, and a response surface method in ANSYS were also used in this study. The use of derivative-free optimization methods provides generality and
modularity to the approach considered here. One advantageous by-product of this approach is that direct methods are usually more robust and more global than their derivative-counterparts.

4.7 Statistical inference with DIC data

Estimating the level of uncertainty associated with the identification of the parameters is a new trend in material testing. It is good practice in any measurement to evaluate and report the uncertainty associated with the test results. A customer who wishes to know the limits within which the reported result may be assumed to lie may require a statement of uncertainty, or the test laboratory itself may wish to develop a better understanding of which particular aspects of the test procedure have the greatest effect on results so that this may be monitored more closely. A Code of Practice (CoP) was developed by UNCERT (a project funded by the European Commission’s Standards, Measurement and Testing programme under reference SMT4-CT97-2165) to simplify the way in which uncertainties are evaluated. This CoP is one of seventeen produced by the UNCERT consortium for the estimation of uncertainties associated with mechanical tests on metallic materials.

The inverse identification process outlined in Section 4.4 can be interpreted as a statistical parameter estimator using nonlinear regression. Statistical inference can then be employed to study the reliability of the identification and to provide a measure of quality. In using statistical inference, the material parameters set \( \theta \) are considered random variables, so their accuracy is related to the shape of the probability distribution function. Generally, accuracy is defined in terms of a confidence region for the
parameter vector, or as the confidence interval of its components. A $100(1-\alpha)\%$ confidence region is a region of parameter space that contains the real parameters with a probability of $1-\alpha$, where $1-\alpha$ is the confidence level of this particular region.

In deterministic inverse problem formulation, $\mathbf{M}(\boldsymbol{\theta})$ is called the model; in statistical inverse problem formulation of nonlinear regression, it is called the expectation function. In nonlinear regression inference, the success of the identification can be checked by two measures: (i) expectation function fit and (ii) parameter fit. If a model gives a smaller residual in the response function, and/or a smaller confidence region in the expectation function and response function, the model is said to be phenomenologically better (i.e., the model parameter set yields a better description of the model’s behaviour).

The most accurate way to perform the statistical analysis is through Monte Carlo (MC) methods. The Monte Carlo method has been used to validate the linearized covariance analysis (LCA) for nonlinear parameter estimation problems (Grimstad, Kolltveit et al., 2001). However, this method requires the solution of a large number of related parameter identification problems. Considering the complex geometry and nonlinearity involved in the identification of plasticity models, the Monte Carlo approach is not feasible for the present study.

4.7.1 Nonlinear regression inference using linear approximation

From a statistical perspective, the identification of constitutive laws using full-field measurements can be considered as a problem where the sample is essentially the complete population. The LCA method is the most computationally efficient method and
the most used one. It was shown previously that, for normally distributed errors, LCA is equivalent to the likelihood method (Uusipaikka, 2009) (Seber and Wild, 2005).

The covariance matrix $P$ of the model parameters can be approximated from the linearization of the cost function $f(\theta)$ around the solution $\tilde{\theta}$:

$$f(\theta) \approx f(\tilde{\theta}) + J(\tilde{\theta})(\theta - \tilde{\theta})$$

(4.6)

where $J(\tilde{\theta})$ is the Jacobian of the cost function $f(\theta)$ evaluated at the solution $\tilde{\theta}$. The matrix $P$ is then given by:

$$P = \text{cov}(\theta) \approx s^2 \left(J^T(\tilde{\theta})J(\tilde{\theta})\right)^{-1}$$

(4.7)

Recalling expansion (2), the residual mean square estimate, $s^2$, is given as follows:

$$\sigma^2 \approx s^2 = \frac{R^T(\tilde{\theta})R(\tilde{\theta})}{N-K}$$

(4.8)

where $N$ is the sample size and $K$ is the number of parameters.

The associated correlation matrix is defined as:

$$C_{ij} = \frac{p_{ij}}{\sqrt{p_{ii}p_{jj}}}$$

(4.9)

The off-diagonal elements of the $C_{ij}$ matrix represent the correlation between two parameters $\theta_i$ and $\theta_j$. A strong correlation often indicates over-parameterization due to either a fault or true redundancy within the model. The diagonal elements of the covariance matrix $P$ represent the estimated values of the variances of the parameters $\theta_j$.

$$\sigma^2_{\theta_j} \approx p_{jj}$$

(4.10)

which is a measure of confidence for the identified parameters $\theta_j$.

The $100(1 - \alpha)\%$ joint confidence region is an ellipsoid and can be defined as:
\( (\theta - \hat{\theta})^T P (\theta - \hat{\theta}) \leq KF(K, N - K; \alpha) \)  \hspace{1cm} (4.11)

where \( F(K, N - K; \alpha) \) is the upper \( \alpha \) quantile for Fisher’s F-distribution with \( K \) and \( N - K \) degrees of freedom.

An approximate \( 100(1 - \alpha)\% \) marginal confidence interval for a given parameter \( \theta_i \) is expressed by:

\[
\hat{\theta}_i \pm s(\hat{\theta}_i) t \left( N - K; \frac{\alpha}{2} \right)
\]  \hspace{1cm} (4.12)

where \( t \left( N - K; \frac{\alpha}{2} \right) \) is the upper \( \alpha/2 \) quantile for Student’s t-distribution with \( N - K \) degrees of freedom and \( s(\hat{\theta}_i) \) is the approximate standard deviation associated with \( \hat{\theta}_i \):

\[
s(\hat{\theta}_i) = \sqrt{P_{ii}}
\]  \hspace{1cm} (4.13)

Expression (4.12) indicates that the marginal posterior density for a single parameter \( \theta_i \) is a univariate Student’s t-distribution with location parameter \( \hat{\theta}_i \), scale parameter \( P_{ii} \), and degrees of freedom \( N - K \). Similarly, the marginal posterior density of the model response is also a Student’s t-distribution.

The \( 100(1 - \alpha)\% \) confidence interval is approximated as:

\[
(\theta - \hat{\theta})^T P^{-1}(\theta - \hat{\theta}) \leq \chi^2_{K, \alpha}
\]  \hspace{1cm} (4.14)

where \( \chi^2_{K, \alpha} \) is the upper \( \alpha \) quantile for the \( \chi^2 \)-distribution with \( K \) degrees of freedom.

A popular formulation of the confidence interval is given as:

\[
|\theta_i - \hat{\theta}_i| \leq z_{\alpha/2} \sqrt{P_{ii}}
\]  \hspace{1cm} (4.15)

where \( z_{\alpha/2} \) is the upper \( \alpha/2 \) quantile for the Gaussian distribution.
If the model-data combination is not highly nonlinear, the minimum of the paraboloid approximation is close to the minimum of the true cost function, and the inference can be considered reliable.

The LCA-based inference can also be used in other ways, including model selection and design of experiments (DOE); this is best achieved through Bayesian inference. The Bayesian marginal posterior density for material parameters $\theta$ (Seber and Wild, 2005) is formulated as:

$$p(\theta|d) \propto \left(1 + \frac{(\theta - \hat{\theta})^T P^{-1}(\theta - \hat{\theta})}{N-K}\right)^{-N/2}$$ (4.16)

This probability distribution is in the form of a $K$-variate Student’s t-density with location parameter $\hat{\theta}$, scaling matrix $P$, and $N - K$ degrees of freedom. If a non-informative prior density for $\theta$ is assumed, the Bayesian identification and Bayesian inference will be identical to the results based on sampling theory given above.

Bayesian inference possesses a great advantage because prior information can be naturally incorporated into the identification process, and model selection is also convenient (Koch, 2007). However, this aspect is for the focus of future work, and will not be addressed in this thesis.

4.7.2 Calculation of Sensitivity

The sensitivity of the response function is required in the LCA inference, the expectation fit check, and the calculation of curvature measures; the gradient is obtained by finite differentiation. While the cost function involves non-negligible errors associated with the measurement data, the response function is evaluated using the
numerical model representing the true physics, the modeling error induced by the finite element approximation can be ignored and the numerical results can thus be assumed to be noise-free.

The gradient is approximated by means of finite differences and requires an extra FE simulation for each of the unknown parameters of the constitutive law with a perturbed value for the parameter under consideration. The sensitivity can be determined as follows:

\[
\frac{dj}{d\theta_i} \approx \frac{\Delta j}{\Delta \theta_i} = \frac{J(\theta_i + \Delta \theta_i)}{\Delta \theta_i}
\]

(4.17)

A perturbation size of \(\Delta \theta_i = 0.001 \theta_i\) is used for each parameter with the exception of the Poisson’s ratio, where a step size of \(\Delta \nu_i = 0.1 \nu_i\) is used.

4.7.3 Checking response fit

The residual \(R(\hat{\theta})\) has been used directly as the criterion for checking the convergence, and for comparing the model’s fit. However, the residual values depend on the units of the model and response functions, loading level and magnitude of response, and also the number of data points, \(N\); thus, it is not a convenient measure of model fit and convergence.

A relative offset measure for checking convergence and model adequacy in nonlinear regression is given by (Bates and Watts, 1988):

\[
I_M = \frac{\left(\|Q_1^T(d-M(\theta))\|/\sqrt{K}\right)/\sqrt{N-K}}{\left(\|Q_2^T(d-M(\theta))\|/\sqrt{N-K}\right)}
\]

(4.18)

where \(Q_1\) and \(Q_2\) are respectively, the first \(K\) and the last \(N-K\) columns of the \(Q\) matrix in the QR-decomposition of Jacobian \(J\), denoted \(Q\). This index is related to the cotangent
of the angle that the residual vector makes with the tangent plane, so that a small relative offset corresponds to an angle near $90^\circ$, and validates the model’s accuracy. Bates and Watts suggested the use of the criterion $I_M \leq 0.001$, reasoning that any inference will not be affected because the current parameter vector is less than 0.1% of the radius of the confidence region disk from the least squares point (Bates and Watts, 1988). Considering that the Jacobian is evaluated using finite difference in the present work, the criterion $I_M < 0.01$ is used to check response fit.

4.8 Identifiability issues

4.8.1 General

One of the fundamental problems in any inverse analysis is the justification and assessment of the credibility of inversely identified results. The difficulty inherent in the substantiation of a specific material identification problem is that without reference values for comparison, how can we trust the identified material model parameters, especially for in-situ determination of material properties where no “reference” for parameter values exists?

The principle that the parameters in a model can be consistently estimated is often referred to as the identifiability of a model (Hsia, 1977). Identifiability is necessary to draw consistent statistical inferences. The problem of identifiability and the stability of numerical results are examined in this section; scepticism is the attitude we take in addressing this issue.
4.8.2 Ill-posed or well-posed

The definition of well-posed problems given by Hadamard refers to the existence, uniqueness and stability of solutions to differential equations (Tikhonov and Arsenin, 1977). Thus, in the case of inverse problems, the concept of well-posedness is related to the existence and uniqueness of the solution to an inverse problem; the data must be sufficient (i.e. contain enough information for simultaneous determination of all the unknown parameters), and the global optimal must be unique and stable when there is noise in the data.

The problem of material parameter identification has been described as ill-posed (Mahnken, 2004); in practice, however, the material parameter identification problem must be well-posed, with a few specific exceptions, such as the non-uniqueness of parameters in the creep test for viscoplasticity (Seibert, Lehn et al., 2000a) (This particular model is phenomenological, and is a summation of an arbitrary number of Dirichlet-Prony exponentials chosen by the user). One indicator of ill-posedness is the need for regularization. There should not be any regularization terms in the least square objective function (4.1) for an unbiased solution of material parameters. In general, the number \(N\) of samples of experimental data has to be much larger than the number \(K\) of parameters; when these conditions are met, the resulting material parameter identification problem is usually a well-posed problem.

Essentially, whether the problem is ill-posed or well-posed lies in whether the problem is a function parameter identification problem or a functional identification problem (i.e. problem of identification of a parameterized continuous field). The problem outlined in Chapter 3 is indeed ill-posed, since it is a parameterized functional
identification problem. Functional identification problems are affluent in geophysical and medical tomography (Snieder and Trampert, 1999). In functional identification problems, whether the forward problem is represented as a linear system $Am = d$ or a nonlinear system $A(m) = d$, where $A$ defines a nonlinear operator, it is ill-posed if the information contained in data $d$ alone cannot guarantee a unique and stable solution of model parameters $m$.

4.8.3 Model and parameter identifiability

Kataygiotis and Beck (1998) made a distinction between “model identifiability” and “parameter identifiability” (Kataygiotis and Beck, 1998). The former has also been called “structural identifiability” (Cobelli and DiStefano, 1980; Hof, 1998). The model identification problem is to find all of the structural models within a specified model class that are able to produce the same output within a set of observed degrees of freedom when the models are all subjected to the same input. However, in our material parameter identification problem, the model is selected via the judgement of the engineer, not through the identification process; thus, we need only focus on the parameter identifiability issue.

4.8.4 Verification and validation (V&V)

The assessment of the credibility of a numerical procedure is typically called verification and validation (V&V), and is closely connected to the problem of identifiability. The verification process can be defined as verification of the model response obtained from data, and the validation process as the comparison of the model
response with data that was not used in the identification. In the professional literature, verification is usually defined as the process of assessing software correctness and the numerical accuracy of the solution to a given model; validation is the process of assessing the physical accuracy of a model (ASME, 2006).

A more straightforward definition of V&V is that verification is “solving the equations right”, and validation is “solving the right equations” (Oberkampf and Roy, 2010). In material identification, verification can be defined as the correct solution of the inverse problem (i.e. the global optimal of the multi-modal LS objective function is determined), and validation is that the problem is defined correctly (i.e. the data contain enough information, the global optimum is unique, and the FE model (mesh, element type, boundary conditions, etc.) represents the reality well, so that the nonlinear regression model (data-model set) is correctly defined).

Thus, V&V can be described as the process of providing evidence of the correctness and accuracy of the parameter identification results; V&V procedures are founded on the concept of quantitative accuracy assessment. While V&V do not entirely answer the question of identification credibility, they are the keys to establishing credibility. Traditionally, one needs to have accurate benchmarks or reference values for comparison. Reference values are not generally available for complex material model parameters, however, especially for individual components. For this reason, numerical studies and statistical tools are used to assess uncertainty during V&V of the inverse identification process.

In material parameter identification, V&V is concerned with two issues: (i) to verify that the solution is a global optimum; and (ii) to validate the global optimum by
ensuring that it corresponds to a set of parameters that are similar enough to the correct values. When using local optimization methods, techniques based on the re-identification of the parameters using different initial solutions is widely used to verify that the optimum is a global solution. Although derivative-free methods, such as CONDOR, are considered to be robust and globally convergent, the use of multiple-initial estimates is still recommended for V&V. Alternatively, re-identification of the parameters using synthetic data can also serve as a verification tool for assessing the identifiability of a given experimental/estimation solution.

4.8.5 Quality and nature of information from data

Besides the problem of local minimum, the quality and nature of the available data also needs to be considered carefully. There are two causes for unsuccessful identifications: (i) the minimization process leads to a local minimum; and/or (ii) the data are insufficient. For example, the parameters of the Rousselier damage mechanics model were identified by updating a FE model to match a simulated 2D displacement field on a tensile specimen using synthetic data at every FE node (Springmann and Kuna, 2003). If only one material parameter was under consideration, the identification was successful; however, simultaneous updating of two or more parameters was less successful because, in some cases, only a local optimum could be found. One possible reason for unsuccessful identification may be due to the fact that the information contained in the collected data is insufficient for simultaneous updating of all of the parameters.

The nature of the data can lead to over-parameterization; i.e. when one or more of the mechanical properties associated with the parameters are not activated during the
experiment. For example, if, when attempting the identification of a hardening law in an elasto-plastic constitutive model, the material remains largely in the elastic range under a given load. In the case of DIC, the large amount of data with an elastic nature may overwhelm the small fraction of data in the plastic range. This situation is called fault redundancy induced within the experiment.

A quantitative approach to determining identifiability is based on Fisher information; a parameter vector is identifiable from a particular set of observed responses if the associated Fisher information matrix is nonsingular (Bos, 2007). The Fisher information is related to the sensitivity of the Jacobian with respect to parameters.

\[
F_\theta = E(S_\theta S_\theta^T) = E \left( \frac{\partial q(w; \theta)}{\partial \theta} \frac{\partial q(w; \theta)}{\partial \theta^T} \right)
\]

is the Fisher information matrix of the observed data \(d\), where \(q(w; \theta)\) is the log-probability density function of the response, and \(S_\theta\) is the Fisher score vector. The Fisher score and Fisher information are connected to the sensitivities of the response to the unknown parameters.

However, local sensitivity at a single point may not be sufficient; the singularity of Fisher information not only depends on the local sensitivity at a point in parameter space, but also depends on the global sensitivity in a finite region. Global sensitivity analysis techniques deal with the entire range of variation of the input parameters. Sensitivity analysis allows one to study the relationships of response variances and parameter variances, and to identify adequacies in the numerical-experimental setting. In particular, global sensitivity analysis allows analysts to perform model calibration, model validation, and decision making; i.e. global sensitivity analysis enables any process where
it is useful to know which of the variables contribute most to the variability of the output. One should note that while sensitivity analysis quantifies the influence of the input parameters on the model’s response, uncertainty analysis is used to evaluate statistical parameters, confidence intervals and probability laws for the model’s responses.

Thus, inadequacy and faulty parameter redundancy due to the quality and nature of the data can be detected through global sensitivity analysis and/or statistical inference (Patelli, Pradlwarter et al., to appear) (Saltelli, Ratto et al., 2008).

4.8.6 Curvature measure of nonlinearity: checking the adequacy of linearized covariance analysis (LCA)

Uncertainty quantification is an important aspect of V&V. Linearized covariance analysis (LCA) is used to check the fit of parameters, and provides a quantitative measure of uncertainty. LCA is an approximation of a nonlinear problem. Whether this approximation is valid, and produces results that are adequate is addressed in this section by adopting an attitude of scepticism.

The most reliable check of the adequacy of LCA is through the use of Monte Carlo analyses. However, as noted previously, the research presented here is based on the assumption that Monte Carlo analyses are not affordable, so curvature measures of nonlinearity are substituted as an alternative to Monte Carlo techniques in this study. Curvature measures are able to detect only incorrect results that are false positives (type-II errors). In other words, it is possible for nonlinearity measures to give a false alarm indicating a problem with the LCA when the LCA is actually adequate; however, nonlinearity measures will correctly detect all instances when the LCA is inadequate. In
fact, type-I error rarely happens in material testing and parameter identification if sound judgement is employed; so, focusing on avoidance of type-II error is sufficient for our purposes.

When deciding whether to reject the null hypothesis that $H_0: \theta = \hat{\theta}$, there are two fundamental errors to be identified by inference:

- Type-I error: the hypothesis is actually true, but we think it is wrong, i.e. the LCA is adequate, but we think that there is a problem with the solution.
- Type-II error: the hypothesis is actually wrong, but we think it is true, i.e. the LCA is inadequate, but we think that the solution is correct.

In the context of statistical regression analysis, the cost function $f(\theta)$, is considered as an expectation function. The noise associated with measurement and modeling errors is assumed to have spherical normal distribution. The expectation function can be considered as an $N$-dimensional response surface. The vector $M(x; \theta)$ defines a $K$-dimensional surface, and is called the expectation surface in the response space. The Least Square estimates thus correspond to a point on the expectation surface $\hat{M} = M(\hat{\theta})$, which is the closest point to data vector $d$. For nonlinear models, the expectation surface is curved and bounded (Green, 1988) (Seber and Wild, 2005).

The LCA involves two distinct approximations (Bates and Watts, 1988):

I. The planar assumption, where the expectation surface $M(\theta)$ near $\hat{M}(\hat{\theta})$ is approximated by its tangent plane at $\hat{M}(\hat{\theta})$.

II. The uniform coordinate assumption, in which a linear coordinate system $J(\theta - \hat{\theta})$, is imposed to the approximation of tangent plane defined in 1).
To validate these two assumptions, one can use the curvature of the expectation surface at the solution $\hat{\theta}$. The curvature of the expectation surface depends on the testing and simulation procedures, and will also depend on the type and amount of data used in the identification procedure. For example, an identification based on a load-displacement curve is expected to produce a higher curvature than a technique using full-field data from DIC measurements.

In summary, the LCA uses local information to generate a tangent plane with a linear coordinate system defined by the derivative vectors, projects the residual vector onto that tangent plane, and then maps the tangent plane coordinates onto the parameter plane using linear mapping. If we assume that the tangent plane forms a good approximation to the expectation surface near $\hat{\theta}$, then the likelihood region for $\hat{\theta}$ corresponds to a disk on the tangent plane with a radius proportional to $\sqrt{S(\theta)}$ with $S(\theta) = \| M(\theta) - d \|^2$.

While LCA is used to characterize the quality of the estimates using inference intervals, the adequacy of the LCA can also be checked. The relative curvature measure enables us to assess the intrinsic and parameter-effects (PE) nonlinearity, and the validity of LCA inference. The LCA method is based on the linearization of the model function at a given point; it is of fundamental importance therefore, to verify the validity of the LCA. An accurate method of assessing the validity of LCA, and the global sensitivity analysis is with Monte Carlo analysis, but as noted previously, this method is computationally expensive due to the complex geometry and nonlinearity of the model. The curvature measures of nonlinearity approach are a potential alternative to the Monte Carlo method.
Carlo approach. Curvature measures of nonlinearity are approximations, but are considerably less expensive to compute. In this study, the curvature measures are used to check the validity of LCA inference; as noted previously, this approach can prevent type-II errors, but not type-I errors. If the LCA is deemed to be adequate, then the validity is assured; on the other hand, if the curvature measures of nonlinearity indicate that the LCA is inadequate, it does not necessarily follow that the LCA inference is wrong.

Bates and Watts (1980) employed differential geometry to construct a relative intrinsic and a parameter-effects curvature measure, which provide global measures of nonlinearity of the model (Bates and Watts, 1980). Threshold values for curvature measures of nonlinearity have been published for which application of the LCA may be justified as inadequate, but not necessarily wrong, based on the shape of linearized confidence regions (Bates and Watts, 1988) (Seber and Wild, 2005) (Grimstad, Kolltveit et al., 2001) (Haines, O'Brien et al., 2004). Donaldson and Schnabel (Donaldson and Schnabel, 1987) performed extensive evaluation of LCA predictions for a series of different nonlinear models using Monte Carlo analyses; the recommended thresholds for the curvature measures of nonlinearity constructed by Bates and Watts were found to give good indications as to when the LCA may be insufficient (Donaldson and Schnabel, 1987).

The use of nonlinearity measures to validate LCA, and thus avoid type-II errors, is proposed in this chapter, but is not implemented in the examples. It is left for future investigation in which a Monte Carlo analysis can be performed to determine an accurate threshold for this specific problem, for comparison of the results of LCA, curvature measures of nonlinearity and Monte Carlo approaches.
4.8.7 Summary of procedures addressing the identifiability issue

In summary, the fundamental elements that build credibility into computational results are: (a) the quality of the physics modeling; (b) V&V activities; (c) uncertainty quantification; and (d) sensitivity analyses. All four of these elements are necessary to establish credibility, and none is sufficient in itself. The first element is a pre-requrement for the estimator. Approaches addressing the latter three can be summarized as follows:

1) One may try different optimization algorithms (if required) with different initial values to ensure that the minimization achieves a global minimum.

2) Re-identify the model: perform a numerical study before testing to re-identify parameters of a given material model. This is the verification procedure in V&V. This numerical test needs to be based on both synthetic noise-free data and noise-polluted data to ensure the well-posedness of the inverse problem defined by the data-optimizer combination. In other words, not only the existence and uniqueness of the solution, but also the stability of the solution need to be assured. From the results of numerical tests, one can determine whether or not all of the unknown model parameters could be identified from the response of the selected system. Unlike functional identification problems, the material parameter identification problem must be a well-posed problem.

3) Validation using different data sets: depending on the type of test, the test data can be divided as \( D = D_1 + D_2 \). The material parameters are
determined based on data set $D_1$ (parameter identification), and investigation of the model quality can be performed using another set of data $D_2$ (validation) for extrapolation. Extrapolation (validation) is an assessment performed by using the identified parameters to predict the response of the material in experiments that were not used to supply data for the identification. If the difference is significant in all or part of the responses, then this is an indication of potential problems in the choice of model, the experimental setup, or the design of the identification process. Furthermore, if the errors between predicted and measured responses show a deterministic pattern that cannot be explained by stochastic distribution of errors, this also indicates potential problems.

4) Inference using LCA: to consider the parameters as random variables. The transfer process of uncertainty from data to the identified parameters is studied, and a quantitative evaluation of uncertainty is determined.

5) In the optimization process, checking expectation (response) fit using the offset index $I_M$ in addition to the convergence check inherent in the adopted optimization algorithm. This gives a non-dimensional indication of the level of response fit. If the final residual $R(\hat{\theta})$ is too high as demonstrated by a high value of non-dimensional index $I_M$, it is an indication to increase the complexity of the model (i.e., increase the number of parameters $K$), to reconsider the underlying physics for selection of a different model, or to re-examine the modeling process and test setup. For example, the Bauschinger effect can be included, or
exponential hardening changed to piecewise linear hardening laws, bilinear hardening changed to multi-linear hardening, or a power type Norton law could be used. More complicated models may be involved, such as anisotropic models (Hill models).

6) Curvature measure of nonlinearity: for checking whether the LCA inference is adequate or not. This procedure completely removes type-II errors from material parameter identification, which is of greater concern than type-I errors.

7) If necessary, perform a global sensitivity analysis.

The above procedures can be implemented with software for general purpose material parameter identification in industrial applications.

4.9 Examples and results

4.9.1 Numerical example

The first example uses artificial data generated from numerical simulation; the intent is to test the optimization algorithm and demonstrate the proposed methodology. We consider a plate with a hole subjected to a tension load above the elasticity limit where the deformation is heterogeneous (Figure 4.2). The boundary value problem is solved using a standard finite element method with reference parameters. The goal is to re-identify the parameters using synthetic displacement data collected at 25 scattered points on the surface of the specimen from evenly distributed grids that simulate DIC measurements, since DIC measurement is based on a regular grid pattern defined by a fixed number of pixels.
The material is assumed to be isotropic; the elastic behavior is characterized by Young’s modulus, and the hardening curve is assumed to be linear isotropic characterized by two parameters. Young’s modulus, Poisson’s ratio, the initial flow stress and the tangent modulus after yielding were assumed to be $E = 70000$ MPa, $\nu = 0.2$, yield stress $\sigma_f = 243$ MPa, and tangent modulus $E_t = 2127$ MPa. The objective of this identification process is to identify the four material parameters, $\theta = (E, \nu, \sigma_f, E_t)$, that collectively characterize the isotropic hardening behavior. Equation (2) is used as the cost function, and in this case, we have $K = 4$, and $N = 25$. The optimization was performed as one single process to simultaneously identify all four of the parameters.

A plane stress condition is assumed; the right edge was subjected to stress increasing from zero to a maximum value of 133.65 MPa, and then the plate was unloaded. The peak value was selected so that the mean stress over the section with the hole is 10% above the yield stress. The plasticity range during the final loading step is
shown in Figure 4.3. Second-order triangular elements with four Gauss points were used in the simulation.

Several inverse analyses were performed with different initial estimates, and the results show that the solution produces a global minimum; this is the first indication that the problem is not ill-posed and that the solution is stable. To evaluate the sensitivity of these methods to uncertainty in the measurement, a 5% white noise measure was added to the synthetic data. The noise was defined as:

$$\text{noise}(x) = NRAND \cdot \alpha \cdot u(x)$$

(4.18)

where $NRAND$ is a Gaussian random distribution with zero-mean and a unit standard deviation; $\alpha = 5\%$ is the applied noise level; and $u(x)$ is the displacement response at location $x$. The corresponding identification solution is given in the fifth column of
Table 4.1; while the initial estimates used to start the optimization process is given in the fourth column.

<table>
<thead>
<tr>
<th>noise</th>
<th>parameter</th>
<th>Target values</th>
<th>Initial guesses</th>
<th>Obtained values (CONDOR)</th>
<th>Variance (LCA) $\sigma_{\theta_j}^2$</th>
<th>relative offset measure $I_M$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0%</td>
<td>$E_1$</td>
<td>70000</td>
<td>50000</td>
<td>70001</td>
<td>2.86</td>
<td>0.114</td>
</tr>
<tr>
<td></td>
<td>$\nu$</td>
<td>0.2</td>
<td>0.3</td>
<td>0.215</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>$\sigma_y$</td>
<td>243</td>
<td>500</td>
<td>243</td>
<td></td>
<td>2.5</td>
</tr>
<tr>
<td></td>
<td>$E_2$</td>
<td>2127</td>
<td>10000</td>
<td>2126.8</td>
<td>11.9</td>
<td></td>
</tr>
<tr>
<td>5%</td>
<td>$E_1$</td>
<td>70000</td>
<td>50000</td>
<td>70006.1</td>
<td>2.86</td>
<td>0.116</td>
</tr>
<tr>
<td></td>
<td>$\nu$</td>
<td>0.2</td>
<td>0.3</td>
<td>0.216</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>$\sigma_y$</td>
<td>243</td>
<td>500</td>
<td>240.4</td>
<td></td>
<td>12.7</td>
</tr>
<tr>
<td></td>
<td>$E_2$</td>
<td>2127</td>
<td>10000</td>
<td>2141.2</td>
<td>31.4</td>
<td></td>
</tr>
</tbody>
</table>

For comparison, two additional direct search methods were also used; namely, (i) the Nelder-Mead simplex method and (ii) the genetic algorithm in the MATLAB toolboxes. Table 4.1 summarizes the results. Both methods produced similar solutions to the method proposed here, but with different levels of efficiency. The proposed solution required nine iterative steps of optimization when using CONDOR, 29 iterative steps of optimization using the Nelder-Mead simplex method, and 3500 function evaluations using the MATLAB genetic algorithm.
Table 4.2: Comparison of identification solution with and without synthetic noise to Nelder-Mead and Genetic Algorithm (GA) methods (load factor = 1.1)

<table>
<thead>
<tr>
<th>Noise</th>
<th>Parameter</th>
<th>Target values</th>
<th>Initial estimates</th>
<th>Obtained values (Nelder-Mead)</th>
<th>Obtained values (GA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0%</td>
<td>$E_1$</td>
<td>70000</td>
<td>50000</td>
<td>70001.5</td>
<td>70002</td>
</tr>
<tr>
<td></td>
<td>$\nu$</td>
<td>0.2</td>
<td>0.3</td>
<td>0.217</td>
<td>0.216</td>
</tr>
<tr>
<td></td>
<td>$\sigma_y$</td>
<td>243</td>
<td>500</td>
<td>243</td>
<td>242</td>
</tr>
<tr>
<td></td>
<td>$E_2$</td>
<td>2127</td>
<td>10000</td>
<td>2126.5</td>
<td>2129</td>
</tr>
<tr>
<td>5%</td>
<td>$E_1$</td>
<td>70000</td>
<td>50000</td>
<td>70008</td>
<td>70005</td>
</tr>
<tr>
<td></td>
<td>$\nu$</td>
<td>0.2</td>
<td>0.3</td>
<td>0.217</td>
<td>0.214</td>
</tr>
<tr>
<td></td>
<td>$\sigma_y$</td>
<td>243</td>
<td>500</td>
<td>238</td>
<td>240</td>
</tr>
<tr>
<td></td>
<td>$E_2$</td>
<td>2127</td>
<td>10000</td>
<td>2143</td>
<td>2141</td>
</tr>
</tbody>
</table>

It is clear that the identification results depend highly on the quality of the input data. The plasticity mechanism must be fully activated in order to identify the model parameters. To study this aspect of the proposed method, the identification was repeated with simulated inhomogeneous displacement field data at the loading stage where the mean stress is 5% above the yield stress. The identified model parameters calculated without synthetic noise are shown in Table 4.3. The deterioration of the identified plasticity parameters is evident. It is interesting that the evaluations for the initial tangent
modulus $E_1$ and Poisson’s ratio $\nu$ did not degrade, however plasticity-related parameters were affected; both the standard error of the mean and the variance increased for these parameters.

Table 4.3: Identification solution without synthetic noise (load factor = 1.05) $I_M=0.0001$

<table>
<thead>
<tr>
<th>parameter</th>
<th>Target values</th>
<th>Initial guesses</th>
<th>Optimized values (method 1)</th>
<th>Variance</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E_1$</td>
<td>70000</td>
<td>50000</td>
<td>70001</td>
<td>2.3</td>
</tr>
<tr>
<td>$\nu$</td>
<td>0.2</td>
<td>0.3</td>
<td>0.212</td>
<td>0.115</td>
</tr>
<tr>
<td>$\sigma_y$</td>
<td>243</td>
<td>500</td>
<td>254</td>
<td>46</td>
</tr>
<tr>
<td>$E_2$</td>
<td>2127</td>
<td>10000</td>
<td>3028</td>
<td>342</td>
</tr>
</tbody>
</table>

In the original identification we assumed the associated von Mises flow rule. To further study influences on the proposed plasticity model, the identification was repeated when the underlying model was changed to use a Tresca yield criterion. The parameters identified using noiseless data are shown in Table 4.4. Since the model selected differs from the “true” mechanism, there was deterioration in the identification of the plasticity parameters. This is the problem with “model selection”.
Table 4.4: identification solution without synthetic noise (load factor = 1.1)(Tresca yield criterion) $I_M = 0.0002$

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Target values</th>
<th>Initial estimates</th>
<th>Optimized values</th>
<th>Variance</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E_1$</td>
<td>70000</td>
<td>50000</td>
<td>70001</td>
<td>2.78</td>
</tr>
<tr>
<td>$\nu$</td>
<td>0.2</td>
<td>0.3</td>
<td>0.25</td>
<td>0.13</td>
</tr>
<tr>
<td>$\sigma_y$</td>
<td>243</td>
<td>500</td>
<td>244</td>
<td>11.5</td>
</tr>
<tr>
<td>$E_2$</td>
<td>2127</td>
<td>10000</td>
<td>2124</td>
<td>14.7</td>
</tr>
</tbody>
</table>

4.9.2 Example 2: identification of model parameters of a cast iron component

In this example, a grey cast iron engine bearing cap subjected to vertical compressive loading is analyzed (Figure 4.4). The load was applied on a Tinius–Olsen hydraulic universal testing machine with a load capacity of 300 kN. A maximum load of 150 KN was applied to the top of the bearing cap. The ARAMIS-DIC system was used to measure the deformation and the strain on the surface of the specimen. An automatic data acquisition system coupled with the testing machine was also used to collect the data for the load—displacement curve.

A random pattern was applied to the specimen’s surface; a flat white paint was applied to the surface in order to reduce the glare in the camera image. A black speckle pattern was then applied using spray paint in order to provide contrast with the white surface coating, and to provide surface target points from which strain and displacement measurements were evaluated using digital image correlation photogrammetry. Speckle
points were approximately 3 to 5 pixels in diameter, and the correlated subset size chosen was 15 pixels. According to (Tong, 2005) and (Robert, Nazaret et al., 2007), 5% of RMS error is a conservative estimate of the error in measurement for a speckle pattern with speckle sizes smaller than the subset size.

The test data was comprised of the force-load curves and the three-dimensional displacement/strain fields on the surface of the component, measured by using the ARAMIS commercial DIC system. Two CCD cameras with a 510 mm object-camera distance were used to acquire synchronized stereo images of the component under compression at different load levels during the test (Figure 4.5).

By means of photogrammetrical procedures and image processing, the data acquisition system evaluated the three dimensional displacement field associated with each respective loading step. Although the ARAMIS system provides three-dimensional displacements and two-dimensional strains on the surface, only the horizontal strains ($\mu_x$) and vertical strains ($\mu_y$) were used in the identification of model parameters. In DIC measurement, the primary results consist of displacements, and the deformation data are obtained by post-acquisition processing techniques. Equation 4.2 was used as the cost function, where the data vector $d$ consists of 120 measurement points inside the rectangle (shown in Figure 4.10), and the corresponding numerical responses $M_i(\theta)$ are the simulated results from the same points. It was found that strains in this region are sensitive to the material parameters, and that the DIC-measured data are reliable in this region (located in the center of the images). In total, 53 images were taken during the loading process at one second time intervals. The load was measured with a Tinius-Olsen
load cell and recorded by the ARAMIS data acquisition system using a BNC connector cable. Each strain stage represents a complete data set for load and deformation.

The bearing surface under the test specimen was greased to reduce friction; therefore, in the FE model, the bottom faces of the two feet of the bearing cap were constrained to prevent vertical movement, while horizontal movement was kept free; symmetry along the direction of thickness was adopted to reduce the computational cost. ANSYS (Version-9) software was used to generate the FE model for a nonlinear cast iron material (Figure 4.6). A solution of horizontal strain of this model is presented in Figure 4.7, which is comparable to the measured quantities in Figure 4.10.

Figure 4.4: the bearing cap
Figure 4.5: photo of the bearing cap on the testing machine and two cameras taking images

Figure 4.6: FE model of the bearing cap
Cast iron materials are known to exhibit different behaviours under tension than they do under compression; the yielding point under tension is significantly lower than the yielding point under compression. The tension and compression behaviour past yielding also differ. Usually, the past yielding tangent modulus for tension is larger than it is for compression. The curve shown in Figure 4.8 is a typical stress-strain curve for cast iron materials. In practice, piecewise-linear models are used to approximate the continuous stress-strain curves after yielding. The most commonly used is two different bilinear curves for tension and compression, respectively, as shown in Figure 4.9. In this case, the material parameters to be identified can be assembled in the set $\theta = (E, E_t, E_c, \epsilon_t, \epsilon_c)^T$, where $E$ is the initial Young’s modulus before yielding, $E_t$ is the tangent modulus after tensile yielding, $E_c$ is the tension modulus after yielding under compression, $\epsilon_t$ is the yielding strain under tension, and $\epsilon_c$ is the yielding strain under compression.
Figure 4.8: typical stress-strain curves for cast iron materials under tension and compression

Figure 4.9: a bilinear approximation model for cast iron material
Figure 4.10: typical images from a DIC test result in ARAMIS; data measured inside the red rectangle are used in the identification process.

Figure 4.10 presents typical DIC test results using the ARAMIS system; the upper image is the strain in the horizontal direction, the lower left image is the picture of the specimen at the corresponding loading stage, and the lower-right image presents the
strain history of three points at different locations. This figure illustrates clearly the heterogeneous nature of the strain field.

After the compression test, a permanent displacement between two end points (A and B in Figure 4.10 was measured as 3.37 mm using vernier calipers; this indicates the presence of plastic deformation. The load and strain data at Stage 50 were used to update the FE model; the compressive load recorded at this stage was 143 kN, and the measured strains were well beyond the elastic limit of 0.2%. Four parameters (i.e., the tension yield strain, the compression yield strain, the tangent modulus after yielding under tension, and the tangent modulus after yielding under compression) were selected as the material parameters represented in two different bilinear models (for tension and compression, respectively). To simplify the identification process, the initial tangent modulus $E$, identical for both tension and compression, was identified first, using the data measured in Stage 10. The load at Stage 10 was 46 kN; the behavior of the whole specimen remained elastic, which is evident in the full-field strain data.

As a first attempt at validating the outlined approach, we re-identified the model parameters using synthetic data generated by FE analysis. The numerical data were collected at the same locations as the test data. Table 4.5 summarizes the results; the target values are the inputs for the constitutive law used by the ANSYS program to generate the displacement and strain fields. It is clear that the results obtained with noiseless data, as well as those contaminated by 5% synthetic noise, were conclusive with respect to identifiability and also demonstrated the usefulness of LCA inference. Although outcomes were similar, the variance of the parameter values obtained with noisy data increased slightly.
Table 4.5: Re-identified parameters with and without synthetic noise

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Target values</th>
<th>Initial estimates</th>
<th>without synthetic noise $I_M = 0.0019$</th>
<th>with 5% synthetic noise $I_M = 0.0044$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Estimated values</td>
<td>Variance</td>
</tr>
<tr>
<td>$E_1$</td>
<td>70000</td>
<td>50000</td>
<td>70002</td>
<td>5.6</td>
</tr>
<tr>
<td>$\epsilon_t$</td>
<td>0.3</td>
<td>0.2</td>
<td>0.305</td>
<td>0.09</td>
</tr>
<tr>
<td>$\epsilon_c$</td>
<td>0.4</td>
<td>0.2</td>
<td>0.396</td>
<td>0.08</td>
</tr>
<tr>
<td>$E_t$</td>
<td>56000</td>
<td>50000</td>
<td>56050</td>
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</tr>
<tr>
<td>$E_c$</td>
<td>55000</td>
<td>50000</td>
<td>55046</td>
<td>74</td>
</tr>
</tbody>
</table>

**Correlating DIC to FEA**

The calibration process in ARAMIS system provide physical coordinates for each pixel point in the image; this data must be mapped to the FE model coordinates and the corresponding FE solutions are extracted at that coordinates. In this work, the selected reference points to build the mapping were the four physical points shown in Figure 4.10: two corner points (A and B), the top of the semi-circle (C), and the top of the cap (D). These points can be identified in the image taken before loading; furthermore, their pixel coordinates within the image can be easily identified, and the mapping between the image and the finite element coordinates can be established for transformation. In some cases, when no physical target points were clearly visible in the image taken by camera (such as points A and B in Figure 4.10), pre-printed target points applied to the specimen surface were used to define the transformation.
Results and analysis

The elastic modulus was identified using data from an elastic stage (Stage 10), and the results are shown in Table 4.6. The other parameters were identified using data from Stage 50; the identified yielding stress points for tension and compression, and tangent moduli after yielding are tabulated in Table 4.7 (The variance is given in brackets under each identified value). Three different optimization methods were used to obtain the parameters: 1) CONDOR; 2) the response surface method using quadratic interpolation, denoted as response surface in ANSYS DesignSpace; and 3) the BFGS pseudo-Newtonian method with finite difference approximated numerical gradient, denoted as numerical gradient.

The best parameter identification obtained for Stage 50 (in comparison to the reference values from Stage 10 shown in Table 4.6) was for the elasticity modulus $E$ apart from the “1-parameter” case shown in the first row of Table 4.7. In the “1-parameter” case, the response surface procedure assumes that the only design variable to be updated is Young’s modulus $E$, meaning that the whole material is elastic in this loading stage. Obviously, the identification of $E$ differs greatly from the reference values (Table 4.6; 70037 [Stage 10] compared to 51251 [Stage 50]), and the inadequacy of the response surface model can also be seen by comparing the $I_M$ values [0.0041 [Stage 10] versus 0.1125 [Stage 50]). The outcomes for the response surface model (second row), the numerical gradient model (third row), and CONDOR model (fourth row) are compared in Table 4.7 using the “2-parameter” procedure, in which two parameters, strain threshold and hardening modulus, are considered simultaneously. In other words, the models are assumed to be formulations of a bilinear hardening law that does not
distinguish the difference between compression and tension. Although the two-parameter procedure is still an inadequate approach to modelling the cast iron material considered here, the differences between the “2-parameter” and the “4-parameter” cases are less evident than those observed with the elasticity model in the “1-parameter” case, suggesting that “2-parameter” models are closer to the “true” one. In the “4-parameter” cases (compared in the bottom three rows of Table 4.7), all four parameters \((E_t, E_c, \epsilon_t, \epsilon_c)\) were included in the identification of material parameters using the three derivative-free methods; the table shows the minimum variances and the minimum objective function values, along with minimum \(I_M\) measures.

The differences between the outcomes of the three derivative-free methods were found to be small, but consistent (Table 4.7). The “numerical gradient” method, had the worst performance indicators, as can be seen from comparison of the objective function values and \(I_M\) measures for the three methods (numerical gradient>response surface>CONDOR). Although the algorithmic details of the “response surface” method are not clear, its performance was found to be better than that of the “numerical gradient” method. Nevertheless, the CONDOR method was shown to be the best method for identifying model parameters in a cast iron component. The identification of model parameters was consistent with reference values for an elastic stage, and CONDOR produced better performance indicators (objective function values and \(I_M\) measures) than the other methods.
Table-4.6: Parameter identification with Stage 10 measurement data (elastic response)

<table>
<thead>
<tr>
<th>Method</th>
<th>$E$ (MPa)</th>
<th>Objective</th>
<th>Variance</th>
<th>$I_M$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-parameter (response surface)</td>
<td>70037</td>
<td>1.6428</td>
<td>25.6</td>
<td>0.0041</td>
</tr>
<tr>
<td>1-parameter** (CONDOR)</td>
<td>70037</td>
<td>1.6427</td>
<td>25.5</td>
<td>0.0040</td>
</tr>
<tr>
<td>1-parameter (numerical gradient)</td>
<td>70101</td>
<td>1.6744</td>
<td>54.2</td>
<td>0.0058</td>
</tr>
</tbody>
</table>

**: the best model
A sensitivity analysis was conducted to study the effects of possible measurement error on parameter identification. The sensitivity of results containing error from the mapping transformation between the FE and the DIC coordinates were considered. The
errors in the x- and y- coordinate transformation (expressed as a percentage) are denoted as $dx$ and $dy$, respectively. The variation of the identified initial tangent modulus $E$ and the objective function value are plotted in Figure 4.11. Figure 4.12 shows the variation of the initial tangent modulus, the tangent modulus for tension and compression as well as the initial yield strain for tension and compression.

Figure 4.11: the sensitivity of the identification of one parameter $E$. (Left: the sensitivity surface of identified $E$ with respect to the errors in the x- and y-translation; Right: sensitivity of the surface of the objective function with respect to the errors in the x- and y- translation)
Figure 4.12: the sensitivity of the identification of four parameters \((E_t, E_c, \epsilon_t, \epsilon_c)\) and the objective function.

It can be seen in these figures that transformation errors introduce uncertainty into the identification results. Nonetheless, the sensitivity surfaces are relatively flat,
especially for transformation errors near the center (zero coordinate). This indicates that the identification function is well-posed with respect to coordinate transformation errors.

### 4.10 Summary of this study

The problem of parameter identification in plasticity constitutive models is formulated as an inverse problem (i.e., mixed-numerical experimental inverse identification of material model parameters) in this study. The data were obtained from full-field measurement of the displacement/strain on the surface of structures using the digital image correlation (DIC) technique. Our objective was to develop a procedure that uses derivative-free optimization algorithms available in commercial FE software (e.g. ABAQUS, ANSYS, COMSOL, MARK, etc.); the derivative-free optimization algorithms were used to determine the parameters by minimizing the gap between measured data and simulated responses.

The proposed methodology is general and appropriate for any material model implemented in commercial software or codes. The method consists of three building blocks: (i) commercial finite element analysis, (ii) direct optimization algorithms, and (iii) digital image correlation test results. The latter two are currently highly commercialized and available for general industrial use.

The focus of this study was on checking the quality and reliability of the solutions. Methodology for validating experiments was presented and demonstrated. A systematic verification and validation process, using statistical and sensitivity analyses, was proposed for use in cases where no reference values of the true parameters are available a priori. If we state that the null hypothesis \( H_0 \) is that the identified parameters
are similar to the “true” parameters, then the proposed validation process can prevent Type-2 error (i.e., acceptance of incorrectly identified parameters), which is the most important requirement for industrial applications.

In conclusion, the direct optimization technique, finite element simulation, and digital image correlation can be combined into a useful, efficient and industrially-applicable technique for the accurate selection and validation of material models and model parameters. Furthermore, all of the techniques involved in the proposed methodology are readily available on the commercial market.
CHAPTER FIVE: HYPERELASTICITY MODEL IDENTIFICATION FOR RUBBER AND RUBBER-LIKE SOLIDS

“*The life of rubber is like the life of human.*” 
(Anonymous)

5.1 Introduction and literature review of related research

Components comprised of rubbery materials play very important roles in engineering processes and products. The use of sophisticated mathematical constitutive models capable of accurate representation of stress/deformation responses (e.g. finite element analysis) is a key ingredient in the design of engineering components and structures under general loading conditions. Usually, the mechanical behavior of rubber-like materials is characterized by hyperelastic constitutive models in which the existence of a strain energy function is postulated; elastic materials that possess a strain energy function are called ‘Green-elastic’ or ‘hyperelastic’ (Ciarlet, 1988; Doghri, 2000).

Strain energy functions are usually defined in terms of the strain invariants (e.g. the polynomial forms), or in terms of the principal stretches (e.g. the Ogden forms). Various types of strain energy functions have been proposed; the most popular of these include the Mooney-Rivlin model and the Yeoh model, based on the phenomenological framework of finite elasticity, and the Arruda-Boyce model, which is founded on the statistical mechanics-based kinetic theory of polymer chain deformations (Boyce and Arruda, 2000; Saccomandi and Ogden, 2004).

There is not a single constitutive model currently available, which can reproduce all aspects of the behavior of real rubber. The selection of the type of strain energy
function and the essential parameters of the model must be determined through appropriate laboratory and/or field tests. In contrast to most material constitutive models, the parameters in rubber constitutive models often have no physical counterparts/meanings, and therefore cannot be measured directly; in general, they must be estimated through an inverse approach (i.e., by fitting the model to experimental data, or by empirical correlations, such as the correlation between rubber stiffness and hardness) (Gent, 2001).

Traditional laboratory techniques for determining the appropriate form of strain energy function and corresponding parameters require homogeneous deformation tests performed on cutout standard samples; these include tests of uniaxial tension and compression, planar shear, and equibiaxial tension and compression. The task is to find the strain energy model that fits the observational data exactly, and will behave reasonably and predictably in other deformation modes.

Initially, elementary methods were utilized to determine the material constants of an isotropic material from simple observational data (Ogden, 1972), such as Treloar’s (Treloar, 1944) data for samples cut from a single sheet of vulcanized natural rubber. Several researchers have used Treloar’s experimental data in simple tension, pure shear, and equibiaxial tension to develop their models for rubber-like materials. For example, Ogden (Ogden, 1972) used a simplified stress-deformation function to model Treloar’s (1944) data; \( P = \sum_{i=1}^{N} \mu_i (\lambda^{-1+\alpha_i} - \lambda^{-1+c\alpha_i}) \), where \( P \) represents the force per unit undeformed area, \( \lambda \) is the principal stretch, \( \mu_i \) and \( \alpha_i \) are the material parameters, and \( c \) is related to deformation type which is equal to -0.5, -2, -1 for simple tension, equibiaxial tension, and pure shear, respectively. Thus, Ogden (1972) developed an ad hoc method
to obtain parameters based on elementary test data, employing the fact that at small strain values, only one term is dominant. The obvious limitation to this approach is that the model provides good fit for only a limited range of strain with such sets of material parameters.

Recently, researchers improved the estimation by using optimization algorithms, such as the Levenberg-Marquardt least-square optimization algorithm, to determine the material constants for each set of the homogeneous deformation data. Many studies that focus on the identification of Rivlin- or Ogden-type polynomial functions use the load-displacement curves or stress-strain relations of a specially shaped sample to formulate a least-square problem. Analytical sensitivity relations can be derived for Rivlin- and Ogden-type strain energy functions; Rivlin models are polynomial functions based on the invariants of the Right Cauchy-Green tensor, and Ogden models are power functions of the eigenvalues of the right stretch tensor with real exponents (Benjeddou, Jankovich et al., 1993) (Gendy and Saleeb, 2000). The major drawback is that the optimization procedure is based on the analytical solution for the deformation observed in simple experiments with a specific form of hyperelastic strain energy; it does not generalize to deformations that differ from those used by the researcher to develop the model or to a different stress-deformation function (Gendy and Saleeb, 2000).

Physical parameters for an in vivo hyperelastic model in living soft tissue were identified in a similar fashion by using the analogy between experimental and predicted numerical results (Tsuta, Yamazaki et al., 1996). The tests were performed on a regular circular area of soft tissue on the human forehead; force-displacement data were collected, and the analytical gradient of displacement for a specific model-type was determined and
parameters were computed for the two-dimensional axi-symmetric model using a steepest
descent method. Currently, the most typical approach used is the general FE-based
procedure for determination of the constitutive law of rubber-like hyperelasticity (Wang
and Lu, 2003). First, uniaxial tension and compression tests of rubber specimens are
conducted; then, tension and compression test simulations with a one-element model and
with FE models of the experimental specimen are performed, and suitable rubber-like
hyperelastic constitutive laws are obtained. Finally, candidate constitutive laws for
hyperelastic and rubber-like materials are selected through comprehensive comparison of
the simulation results (from the FE analysis for real working conditions) with the
experimental observations.

Although parameters can be estimated from the results of a single test, tests
conducted at different deformation modes are usually required to obtain a strain energy
function that is adequate for predicting material behaviour under a variety of deformation
modes. As an alternative to homogeneous experiments, Hartmann, Mars and Fatemi
proposed specially-designed experiments that would be conducted using a novel
specimen, a short, thin-walled cylindrical specimen subjected to combined axial and twist
displacements, in order to study the mechanical response of rubber-like materials under
Hartmann identified parameters in Rivlin’s hyperelasticity model using tension, torsion,
and combined tension-torsion tests with cylindrical rubber specimens where the
analytical solutions to the extension and torsion of a cylindrical body subjected to
incompressibility are known (Hartmann, 2001). The solution to the resulting boundary
value problem is known, and is used to formulate the least-square optimization algorithm.
Sometimes it is difficult to machine an appropriate homogeneous specimen, however, due to the availability of backup components or the size of the component. Furthermore, it is always more desirable to test the behaviour of rubber materials in their installed size and shape within the component, rather than using cut-out standard samples. Nevertheless, determination of the elastic properties of rubber-like materials can be considered as an inverse problem of identification in each of these cases. Moreover, since the essential requirement for laboratory testing is to simulate field conditions as closely as possible, the principal advantage of in-situ tests is that they assess rubber behaviour under natural conditions, thus avoiding sample disturbance. In-situ tests also tend to be more economical than laboratory tests, and in some cases, can provide results that are more representative due to the higher density of collected data.

The research presented in this chapter explores a method that combines Finite Element (FE) model updating with full-field measurement using digital image correlation (DIC) to determine the in-service material properties of mechanical and structural components. From a mathematical point of view, FE model updating involves the minimization of the response gap between the analytical and experimental response data. From a statistical point of view, FE model updating is essentially a nonlinear regression problem fitting the data using a nonlinear function. For example, FE model updating using least-square optimization algorithms was used to estimate Mooney-Rivlin parameters for soft tissue from force-displacement data at the indenter obtained during an in vivo animal experiment; a 3D FE model simulated the force at the indenter and an optimization program updated the parameters and ran the simulation iteratively (Seshaiyer and Humphrey, 2003).
Full-field optical techniques for displacement or strain measurements are now widely used in experimental mechanics. The main techniques reported are photoelasticity, geometric moiré, moiré interferometry, holographic interferometry, speckle interferometry (ESPI), grid method and DIC (Rastogi, 2000). Due to its simplicity and versatility, the DIC method is one of the most commonly used techniques. Many applications for DIC can be found in the professional literature, including research related to the heterogeneous deformation of foams (Wang and Cuitino, 2002), large deformation of polymers (Chevalier, Calloch et al., 2001) (Parsons, Boyce et al., 2004), and measurement of small strains in fiber-reinforced refractory castables (Robert, Nazaret et al., 2007). The DIC method can be used with a single camera (standard DIC) to measure in-plane displacement/strain fields on planar objects, or with two cameras (3D DIC) to measure 3D displacement/strain fields on any 3-D object (Rastogi, 2000).

The DIC technique provides a number of benefits; DIC measures whole field displacement/strain on the surface of a component, generates massive amounts of experimental data in a single test (replacing traditional techniques involving multiple tests with multidimensional loading paths), and thus provides enough information to formulate an inverse problem. The full-field measurement in DIC allows better characterization of the behavior of materials and the response of structural components to external loadings. The problem of ill-conditioning, which is often encountered in FE-model updating, can be avoided to a large extent. Furthermore, although the deformation includes multiple modes, the FE-model formulated using DIC includes coefficients that reproduce the complex deformation modes exactly; thus, the estimate function will be better at predicting the behavior of the material. The large amount of full-field data from
DIC is conducive to statistical inference; for example, the estimate of covariance is valid when the amount of data is large. On the other hand, unless used in conjunction with random simulation, homogeneous tests seldom provide enough data for statistical inference (Seibert, Lehn et al., 2000a) (Harth, 2003) (Harth, Schwan et al., 2004) (Harth and Lehn, 2007). The scientific literature dealing with different aspects of the identification of material parameters for constitutive equations and structural parameters for analysis models is quite extensive, and was recently reviewed (Mahnken, 2004). Mahnken and Stein suggested that full-field optical tests such as grid methods or DIC can overcome the problem of ill-posedness, which is often a major difficulty in inverse identification problems, and to account for non-uniform stress and strain distribution during experiments (Mahnken and Stein, 1996); they presented a unified strategy for material parameter identification in the context of the FE-method that accounts for non-uniform stress and strain distribution in the test specimen. Their work was limited to plane stress samples, however, and required complete measurements in a continuous region.

In this chapter, a commercial FE program (ARAMIS) and a derivative-free optimization tool (CONDOR) are integrated to solve the inverse identification problem. The use of the derivative-free optimization program is not due solely to intellectual curiosity, but is particularly useful in industrial applications. One of the major difficulties in using optimization-based inverse identification besides the problem of local minimum–lies in the implementation of gradient-evaluation procedures. It is usually difficult or impossible to get reliable derivative information for the requested numerical analysis when using complex commercial numerical analysis programs because users
generally do not have access to the codes. Thus, the gradient-evaluation procedures are tedious and require a great deal of additional effort beyond the numerical analysis provided by commercial software. Furthermore, the evaluation of the gradients involved in the analysis can differ for various material models, element types, integration schemes, and degree of deformation or type of response data. For example, a sensitivity analysis was performed using DIC-measured surface strains (Cooreman, Lecompte et al., 2007); this analysis is limited to simple tensile tests, and as the return-mapping scheme is considered in the derivation, cannot be applied directly to FE programs using other integration schemes (e.g. ADINA, which uses an effective stress integration scheme).

The CONDOR method, a trust-region based derivative-free optimization method using multivariate polynomial interpolation, is examined in the analysis presented in this chapter. This algorithm is designed to minimize simple constrained (box constraints) functions whose evaluations are considered to be expensive and whose derivatives are not available for computation. The strain/displacement data measured on part of the surface of a three-dimensional component was used to update the FE model in order to obtain the strain energy function parameters. This approach does not require a complete set of response measurements, but there is no strict mathematical theory that guarantees the validity of the solution. The problem of identifiability is discussed, and nonlinear regression inferences are used to support the validity of the identifications. To demonstrate this inverse approach, several hyperelastic models were identified for the rubber blocks in an engine mount. Specifically, the research presented in this chapter studies the identification of constitutive models for rubberlike materials using an inverse
approach with data from DIC measurements. The analysis is illustrated with a simulated rubber block and experimental observations of an engine block.

This chapter is structured in the following way: Section 5.1: The introduction and review. Section 5.2: Basic description of hyperelastic constitutive models for rubberlike materials. Section 5.3: The approaches used in this work. Section 5.4: Examples and results: including a simulated rubber block and experimental results of an engine block. Section 5.5: Summary and conclusions.

5.2 Hyperelastic models for rubber-like materials

This section presents a short review of current hyperelasticity models for the representation of elasticity behaviour in rubber and rubber-like materials. The underlying hypothesis of hyperelasticity is that there is a scalar-valued potential function, the strain energy density function $W$, which is a function of the strain state, and whose derivative with respect to a particular strain component gives the corresponding stress component by setting:

$$ F(x) = \left( \frac{\partial x^0}{\partial x} \right)(x) = Id + \nabla u(x) = \text{deformation gradient} \quad (5.1) $$

$$ T(x) = T^\phi(x^\phi(x))F(x)^{-T} \det(F(x)) = \text{first Piola-Kirchhoff stress tensor} \quad (5.2) $$

where $x \in \Omega$ ($\Omega$ is the computation domain of the problem) is generic point of the reference configuration, with coordinates $x_i$; and $x^\phi = \phi(x)$ is the generic point of the deformed configuration, with coordinates $x_i^\phi$. The strain energy is a function of the
deformation gradient $F$ (relative to some fixed reference configuration), and is written $W(F)$ per unit volume (Ciarlet, 1988).

The tensor $C = F^T F$ (in $M^3 = E \otimes E$) is called the right Cauchy-Green tensor; it measures the length of an elementary vector after deformation in terms of its definition in the reference configuration. The tensor is symmetric and positive-definite by construction.

Corresponding to the right Cauchy-Green strain tensor $C$, there is also the left Cauchy-Green tensor

$$B = FF^T$$

(5.3)

By construction, $C$ is a symmetric positive definite tensor in $E \otimes E$, which implies that $C$ has three strictly positive eigenvalues $(\lambda_i^2(C))_{i=1}^3$. Three invariants can be defined for $C$

$$I_1(C) = \text{tr}(C) = \lambda_1^2 + \lambda_2^2 + \lambda_3^2$$

(5.4)

$$I_2(C) = \frac{1}{2} \{ (\text{tr} C)^2 - \text{tr}(C^2) \} = \text{tr}(\text{cof}(C)) = \lambda_1^2 \lambda_2^2 + \lambda_2^2 \lambda_3^2 + \lambda_3^2 \lambda_1^2$$

(5.5)

$$I_3(C) = \det(C) = \lambda_1^2 \lambda_2^2 \lambda_3^2$$

(5.6)

A deformation is isochoric if and only if, it preserves volumes ($dx^o = dx$). This deformation is characterized by the deformation gradient $F$ which satisfies the incompressibility constraint

$$\det F(x) = 1 \quad \forall x \in \Omega$$

(5.7)
Thus, the hyperelastic theory states that the first Piola-Kirchhoff stress tensor $T$ is given as:

$$T = \frac{\partial W}{\partial F} = 2F \frac{\partial W}{\partial C}$$  \hspace{1cm} (5.8)

The Cauchy stress tensor $T^\varphi$ is automatically symmetric, and is now given by:

$$T^\varphi = (\det F)^{-1} TF^T = 2(\det F)^{-1} F \frac{\partial W}{\partial C} F^T$$  \hspace{1cm} (5.9)

As for compressible materials, the axiom of indifference implies that $W$ is a function of the right Cauchy-Green strain tensor only (Tallec, 1994). The constitutive law therefore becomes

$$T = 2F \frac{\partial \tilde{W}}{\partial C} - pF^{-T}$$  \hspace{1cm} (5.10)

$$T^\varphi = 2F \frac{\partial \tilde{W}}{\partial C} F^T - pI$$  \hspace{1cm} (5.11)

Similarly, for isotropic materials, $W$ is again a function of the invariants of $C$ only. Thus, in this case we have simply

$$\frac{\partial W}{\partial C} = \left( \frac{\partial W}{\partial I_1} + I_1 \frac{\partial W}{\partial I_2} \right) I - \frac{\partial W}{\partial I_2} C$$  \hspace{1cm} (5.12)

$$T = 2 \left( \frac{\partial W}{\partial I_1} + I_1 \frac{\partial W}{\partial I_2} \right) F - 2 \frac{\partial W}{\partial I_2} FC - pF^{-T}$$  \hspace{1cm} (5.13)

In all of these constitutive laws, the hydrostatic pressure $p$ is an additional unknown element, determined by the Lagrange multiplier associated with the additional nonlinear kinematic constraint

$$J = \det F = 1$$  \hspace{1cm} (5.14)
Various forms of the strain energy function $W$ have been proposed for rubber-like materials. The choice of $W$ usually reflects the experience and preferences of the individual researchers rather than objective criteria. In general, the proposed models can be categorized into two classes: (1) phenomenological descriptions; and (2) descriptions based on statistical mechanics (Horgan and Saccomandi, 2006).

Phenomenological descriptions (based on continuum mechanics analysis) assume that rubber-like materials in the undeformed state are isotropic; i.e., the long molecular chains are distributed randomly, which allows formulation of a strain energy density description in volume element units. The strain energy function is either: 1) a function of the principal invariants of the right Cauchy-Green strain tensor; or 2) a symmetric function of the principal stretches. The former includes the popular Mooney-Rivlin model and the Yeoh model, which are appropriate for strains of medium magnitude; while the latter includes the Ogden model, which is considered to be an effective model for very large strains (Horgan and Saccomandi, 2006).

The strain energy functions for rubber-like materials based on statistical mechanics assume that the elastic restoring force is related to the decrease of entropy, and that elongation of the material’s fibers reduces disorder within the material. Essentially, these models make assumptions about the length and direction of molecular chains; the constitutive models are obtained via statistical analysis (Treloar, 1975). The Arruda-Boyce and Van der Waals are important forms of the models in this class (Boyce and Arruda, 2000) (Horgan and Saccomandi, 2004).
The Arruda-Boyce model of strain energy is the most popular of the \( W \) functions based on statistical molecular theory. The strain energy function is expressed in terms of \( I_1 \) and needs only two parameters, namely \( \mu \) and \( \lambda_m \), for incompressible rubber material, and one additional parameter, \( D \), for a compressible case (Arruda and Boyce, 1993). The strain energy function is:

\[
W = \mu \sum_{i=1}^{5} \frac{C_i}{\lambda_m^{2i-3}} \left( \bar{I}_i - 3 \right) + \frac{1}{D} \left( \frac{J^2 - 1}{2} - \ln J \right)
\]  

(5.15)

where: \( C_1 = \frac{1}{2}, C_2 = \frac{1}{20}, C_3 = \frac{11}{1050}, C_4 = \frac{19}{7050}, C_5 = \frac{519}{673750} \). The Arruda-Boyce strain energy model is also called the eight-chain model. The values for \( C_i \) (\( i = 1, \ldots, 5 \)) are obtained from statistical thermodynamics, and therefore they are physically meaningful. The parameter \( \mu \) represents the initial shear modulus of the material. The parameter \( \lambda_m \) is the locking stretch, which location is around the sharpest point on the stress-strain curve. This type of strain energy function has been shown to be accurate for engineering most elastic materials.

The polynomial form of the strain energy function is written as:

\[
W = \sum_{i+j=1}^{N} C_{ij} (\bar{I}_1 - 3)^i (\bar{I}_2 - 3)^j + \sum_{i=1}^{N} \frac{1}{D_i} (J - 1)^{2i}
\]  

(5.16)

This function denote \( \bar{I}_1 \) and \( \bar{I}_2 \) as the first and second invariants of the deviatoric part of right-Cauchy-Green deformation tensor, respectively; \( J = \text{det}(F) \) is the ratio of deformed volume over the undeformed volume of the material, and \( F \) is the deformation gradient. The parameter \( N \) is the order of the polynomial chosen for a specific material. The parameters \( D_i \) determines the compressibility of the material; the material is
incompressible if all values of $D_i$ are equal to zero. For polynomial models, the initial shear modulus $\mu_0$ and volume modulus $k_0$ are determined by the first-order polynomial ($N = 1$) no matter how many orders are selected for the polynomial model.

\[ \mu_0 = 2(C_{10} + C_{01}), \quad \text{and} \quad k_0 = \frac{2}{D_i} \]  \hspace{1cm} (5.17)

The famous Mooney-Rivlin strain energy form can be obtained by retaining only the linear part of strain energy, that is, $N = 1$.

\[ W = C_{10}(\overline{T}_1 - 3) + C_{01}(\overline{T}_2 - 3) + \frac{1}{D_i}(J - 1)^2 \]  \hspace{1cm} (5.18)

The Mooney-Rivlin form is a very popular one; although it cannot model the sharp upturn of the stress-strain curve under large strains, it is effective for small and medium strain ranges. The first term of this form dictates a linear shear modulus model - the neo-Hookean model:

\[ U = C_{10}(\overline{T}_1 - 3) \]  \hspace{1cm} (5.19)

Treloar constructed the so-called neo-Hookean form of the strain energy on the basis of Gaussian statistics and molecular network theory (Treloar, 1975) as:

\[ W = \frac{1}{2}\mu(I_1 - 3) \]  \hspace{1cm} (5.20)

where $\mu$ is the shear modulus in the ground state. It is easy to see that the Neo-Hookean model can be obtained by retaining only the first term in Equation (5.18). Actually the neo-Hookean form is the first $W$ function proposed by Treloar in 1943.

The reduced polynomial forms are obtained from the polynomial models by setting all of the $C_{ij} = 0$ for $j \neq 0$.
\[ U = \sum_{i=1}^{N} C_{10} (\bar{I}_1 - 3)^i + \sum_{i=0}^{N} \frac{1}{D_i} (J - 1)^{2i} \] (5.21)

A special form of the reduced polynomial is the Yeoh model which is the reduced polynomial function at \( N=3 \):

\[ U = \sum_{i=1}^{3} C_{10} (\bar{I}_1 - 3)^i + \sum_{i=0}^{3} \frac{1}{D_i} (J - 1)^{2i} \] (5.22)

The Yeoh model, is another popular model, and is effective for modeling the large deformation of rubber-like materials.

To obtain accurate estimates of material parameters, the data space of \( \lambda_1 - \lambda_2 \) must be sampled thoroughly. While this is not easily done using traditional tests on homogeneous stress-strain fields, it is easy to do with tests of the original components, so that the test configuration generates a complicated strain/displacement pattern. The mechanical response of a rubber-like material is then defined by choosing a constitutive model with a strain energy function that fits the experimental behavior of the tested components. Several of the most widely applied hyperelastic constitutive models are summarized in Table-5.1.

Table 5.1: Popular constitutive models for compressible rubberlike materials in FE analysis

<table>
<thead>
<tr>
<th>Constitutive models based on statistical mechanics</th>
<th>Number of material parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Arruda-Boyce model</td>
<td>3</td>
</tr>
<tr>
<td>2. Van der Waals model</td>
<td>4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Phenomenological models</th>
<th>Number of material parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Polynomial models (N-th order)</td>
<td>2N</td>
</tr>
<tr>
<td>2. Reduced polynomial models (Yeoh)</td>
<td>N</td>
</tr>
</tbody>
</table>
The number of items included in the polynomial and reduced-polynomial models can be infinitely high, making it possible to represent stress-strain behaviors of any shape. High order strain energy functions are of little practical value, however, because rubber materials are not sufficiently reproducible to allow one to evaluate a large number of coefficients with any accuracy. Generally, the extra terms usually only do a good job in fitting experimental errors. Therefore, the Mooney-Rivlin model remains the most widely used strain energy function in FEA, and should be the first choice due to its simplicity and robustness (Saccomandi and Ogden, 2004). The Mooney-Rivlin model is used in the example presented in Section 5.4.1.

5.3 Methodological and analytical approaches used for this project
The research presented in this chapter is based on the same methodology as the research presented in Chapter 4. Briefly, the ARAMIS commercial DIC system was used to measure displacements and strains on the surface of the test component. A finite element model of the component was developed and correlated with the experimental measurements. Direct optimization algorithms and regression analysis techniques were employed to evaluate and validate this nonlinear optimization problem. Procedural details are provided in Chapter 4, sections 4.2 to 4.7. The results of applying this approach to hyperelasticity in rubberlike materials (a rubber block and an engine mount with a rubber component) are described in Section 5.4.
5.4 Results and illustrative examples

5.4.1 Rubber block

Firstly, a simulated rubber block was used to assess the robustness of the optimization procedure. This example was solved as a plane strain problem using COMSOL software (COMSOL); Figure 5.1 shows the simulated strain contours. The block was compressed between a stationary plane surface and a rigid indenting cylinder; the rigid cylinder started with a gap of 1 mm between the cylinder and the object, and was lowered 8 mm. The lower straight part of the block was glued to the underlying body, so all displacements were constrained there.

Figure 5.1: The simulated strain contours of a rubber block using COMSOL
The rubber material used in this experiment is hyperelastic and was approximated as a Mooney-Rivlin material with $C_{10} = 0.37MPa$ and $C_{01} = 0.11MPa$. The material is almost incompressible, so the bulk modulus was set to 104 MPa, and the mixed formulation option was used. The simulated surface normal strains along a vertical direction were used for updating the model to define the parameters (Table 5.2).

Table 5.2: numerical test of the inverse problem ($I_M = 0.001$)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Reference value</th>
<th>Re-identified value</th>
<th>Relative error</th>
<th>Variances</th>
</tr>
</thead>
<tbody>
<tr>
<td>$C_{10}$</td>
<td>0.37</td>
<td>0.3704</td>
<td>0.11%</td>
<td>0.02</td>
</tr>
<tr>
<td>$C_{01}$</td>
<td>0.11</td>
<td>0.1106</td>
<td>0.55%</td>
<td>0.02</td>
</tr>
<tr>
<td>$1/D_1$</td>
<td>104</td>
<td>103.6</td>
<td>-0.38%</td>
<td>4.8</td>
</tr>
</tbody>
</table>

Studies examining the performance of DIC suggest that a 5% RMS error in strain measurement is a safe estimation (Tong, 2005) (Robert, Nazaret et al., 2007). Therefore, 5% synthetic noise was added to the simulated strains. The results show that this procedure produces good solutions even with potential noise in the DIC measurements (Table 5.3).
Table 5.3: numerical test of the inverse problem with 5% synthetic noise ($I_M = 0.0004$)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Reference value</th>
<th>Re-identified value</th>
<th>Relative error</th>
<th>Variances</th>
</tr>
</thead>
<tbody>
<tr>
<td>$C_{10}$</td>
<td>0.37</td>
<td>0.3647</td>
<td>-1.43%</td>
<td>0.09</td>
</tr>
<tr>
<td>$C_{01}$</td>
<td>0.11</td>
<td>0.1189</td>
<td>8.09%</td>
<td>0.05</td>
</tr>
<tr>
<td>$1/D_1$</td>
<td>104</td>
<td>106.3</td>
<td>2.21%</td>
<td>17.4</td>
</tr>
</tbody>
</table>

5.4.2 Engine mount

An engine mount with a rubber component was tested in the second numerical example. The FE simulation is performed using commercial FE software ANSYS (ANSYS). In the FE model, totally 9454 solid185 hyperelastic elements are used to mesh one rubber block in the engine mount; symmetry condition is used to reduce the computation cost in the FE simulation.

Figure-5.2 shows a typical experimental setup using ARAMIS DIC system. A random speckle pattern using black and white spray paints was coated on the surface of the specimen. The two cameras are shown mounted on a stable base with a special support tripod and support bars that allowed flexibility in positioning the cameras. External light was used to provide optimal exposure. The lighting is especially important as the contrast between white and black speckles can determine the success of a measurement. Likewise, stabilizing the camera, in order to minimize movement and vibration while capturing images, is particularly important. Camera movement causes errors in the camera calibration and deformation calculations. Preliminary trials were conducted to optimize the camera placement and the random speckle pattern.
A small region on the randomly speckled surface of the object was measured using high zoom ratios along with the 1-inch² (25.4 mm) pre-manufactured calibration block that comes with the ARAMIS DIC system. Random speckle patterns were created using 15-pixel subsets that were three times larger than the mean speckle size (5-6 pixels). (DIC measurement depends on the surface speckle pattern of the component to be measured. Um and Kim (2007) studied the effects of subset size, subset shape, and step size on correlation error. Robert et al. (2007) examined the effects of factors such as speckle size and geometry and image noise in speckle patterns. In general, the findings indicate that if the size of a large speckle is larger than the subset, poor correlation scores will result. Thus, a practical rule of thumb is that the subset should be three times larger than the mean speckle size).

The DIC measurement depends on the surface speckle pattern of the component to be measured. Um and Kim conducted experimental research of a paper tensile specimen to study the effect of subset size, subset shape, and step size, on the correlation error (Um and Kim, 2007). Robert et al studied the effect of speckle patterns, i.e., factors such as speckle size and geometry, image noise, etc. (Robert, Nazaret et al., 2007). It was found that if the size of a large speckle is larger than the subset, bad correlation scores will results. A practical rule of thumb is that the subset 3 times larger than the mean speckle size. A subset size of 15 pixels was used in our DIC test; an optimal speckle size should be 5 to 6 pixels.

The frontal surface of the mount was first painted white, and then a black speckle pattern was randomly sprayed onto the white painted surface. Efforts were made to obtain an evenly distributed homogeneous pattern of black paint drops on the undeformed
surface. Inhomogeneous strain/displacement fields on the rubber surface under loading were obtained using the ARAMIS DIC system (ARAMIS). The inhomogeneous strain histories under incremental loading at multiple points were used as input to select the best model and to identify the corresponding material parameters. In this test, the loads were applied to the top of the engine mount, incrementally from 0.0 kN to 13.8 kN; the DIC measurements were conducted at six discrete load-cases, (i.e. 3.6, 6.0, 7.8, 10.8, 12.2, and 13.8 kN load steps). As a good model must be able to predict the behaviour of rubber under a large range of strains, DIC measurements from all six of these load cases were used in the least-squares objective function to update the FE model.

Figure 5.2: The engine mount and the DIC test using the ARAMIS DIC system
Therefore, the least-squares objective problem is written as:

$$\text{Find } \theta \in \Theta : \min f(\theta) = \sum_{p \in P} \sum_{i=1}^{N} \frac{1}{2} \left\| M(\theta) - d_{exp} \right\|^2$$

(29)

where \( P \) is the set of load cases used in the identification; and \( N \) is the number of effective measurement points of the DIC tests. In these DIC measurements, each image contains 1024-by-1280 pixels; grids with subsets of 15-by-15 pixels were used in the analysis. As noted previously, the optimal speckle size is 5 to 6 pixels for the subset size of 15 pixels used in our DIC tests. For speckles larger than this size, the RMS error of strain/displacement measurements increases with the increase in speckle size.

Since the test is performed on a real component rather than a symmetrical cutout, patterns are not well-controlled. The paint was applied to the engine mount manually, and there were also unique scratches, and marks or labels on the surface of the mount. Therefore, regions on the surface with speckles larger than the subset size were deleted from the measured results. Thus, a typical DIC measurement (Figure 5.3) may contain blank areas (regions where the speckle pattern is not adequate for accurate DIC measurement). Therefore, the effective measurement points used in Equation (29), must be chosen from DIC test results that exclude these blank regions.

Figure 5.3: The geometry model and finite element model of the engine mount
The material parameters for several hyperelastic models were identified using the FE model updating procedure described in Section 5.3. Before conducting the identification analysis, a numerical study was performed to validate this test: a three-parameter Mooney-Rivlin hyperelastic model was incorporated into the finite element model, and the parameters were re-identified using the simulated surface displacements as the input measurements. The re-identified parameters are tabulated in Table 5.4.

Table 5.4: numerical test of the inverse problem using simulated surface displacements 
\( (I_M = 0.0006) \)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Reference value</th>
<th>Re-identified value</th>
<th>Relative error</th>
<th>Variances</th>
</tr>
</thead>
<tbody>
<tr>
<td>( C_{10} )</td>
<td>0.66666</td>
<td>0.672623</td>
<td>0.89%</td>
<td>0.072</td>
</tr>
<tr>
<td>( C_{01} )</td>
<td>0.33333</td>
<td>0.310546</td>
<td>-6.84%</td>
<td>0.079</td>
</tr>
<tr>
<td>( D_1 )</td>
<td>0.111111E-4</td>
<td>0.122127E-4</td>
<td>9.91%</td>
<td>0.042E-4</td>
</tr>
</tbody>
</table>

To study the effect of possible errors in DIC measurement, an error with 5% standard deviation was given to the simulated displacements; parameters are then re-identified using the simulated noisy surface displacements as the measurements. According to (Tong, 2005) (Robert, Nazaret et al., 2007), 5% of RMS error is a safe estimate of the error in measurement for a speckle pattern with speckle sizes less than the subset size.
Table 5.5: numerical test of the inverse problem using simulated surface displacements with synthetic noise ($I_M = 0.0013$)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Reference value</th>
<th>Re-identified value</th>
<th>Relative error</th>
<th>Variances</th>
</tr>
</thead>
<tbody>
<tr>
<td>$C_{10}$</td>
<td>0.66666</td>
<td>0.67812</td>
<td>1.72%</td>
<td>0.091</td>
</tr>
<tr>
<td>$C_{01}$</td>
<td>0.33333</td>
<td>0.31735</td>
<td>-4.79%</td>
<td>0.099</td>
</tr>
<tr>
<td>$D_1$</td>
<td>0.111111E-4</td>
<td>0.12472E-4</td>
<td>12.25%</td>
<td>0.048E-4</td>
</tr>
</tbody>
</table>

Tables 5.4 and 5.5 show that, for the identification of parameters in a Mooney-Rivlin hyperelastic model, the accuracy of the identified $C_{10}$ parameter is very good, while the potential errors in identification of the $C_{01}$ parameter and the $D_1$ parameter are relatively high. The identified strain energy function is sufficient to describe the behavior of this rubber, however, because the contribution of the $C_{10}$ parameter is dominant in the Mooney-Rivlin strain energy function. Furthermore, the effects of parameter $C_{01}$ and the compressibility parameter $D_1$ are relatively insignificant for a strain range up to 150%, so this level of error can be regarded as acceptable in FE analysis.
a) Photo of the left rubber block surface used in DIC

b) Horizontal displacement obtained via DIC
c) Vertical displacement obtained via DIC

Figure 5.4: The DIC measurements of displacement in the 7.8 kN load case

Figure 5.5: The FE simulation of displacement at the 7.8 kN load case using identified Mooney-Rivlin model

The parameters identified for four different strain energy functions using the experimental data are tabulated in Table 5.6. The estimated variance for each parameter is given in brackets. Note that, as the measure derives from different models in this case,
the variance is not a meaningful measure for model selection. The measure $I_M$ can be used to select a model that best fits the data in a statistical sense.

Table 5.6: The identified model parameters using experimental data

<table>
<thead>
<tr>
<th>Neo-Hookean</th>
<th>Mooney-Rivlin</th>
<th>Arruda-Boyce</th>
<th>Yeoh</th>
</tr>
</thead>
<tbody>
<tr>
<td>$I_M = 0.0205$</td>
<td>$I_M = 0.0072$</td>
<td>$I_M = 0.0070$</td>
<td>$I_M = 0.0094$</td>
</tr>
<tr>
<td>1.892152 (0.412)</td>
<td>0.578129 (0.124)</td>
<td>1.422176 (0.215)</td>
<td>0.62174 (0.102)</td>
</tr>
<tr>
<td>0.165373 (0.107)</td>
<td>0.452273 (0.147)</td>
<td>4.244912 (0.771)</td>
<td>-0.06631 (0.023)</td>
</tr>
<tr>
<td>0.123581E-03 (0.076E-3)</td>
<td>0.144236E-02 (0.068E-2)</td>
<td>0.14527 (0.062)</td>
<td>0.17412E-03 (0.026E-3)</td>
</tr>
</tbody>
</table>

Figure 5.6: The experimental and simulated load-displacement curves

The load-displacement curve was chosen to verify the identification. The simulated load-displacement curves using FE-identified model parameters are compared with the load-displacement curve derived from experimentally measured parameters in
Figure 5.6. The Neo-Hookean model does not perform well for high load ranges because it is a linear shear modulus model. The simulated load-displacement curves generated by the three other FE-identified models compare well with the experimental load-displacement curve. In particular, the Mooney-Rivlin and Arruda-Boyce models produced the best outcomes in terms of accurately simulating the experimental load-displacement curve. Thus, designers can use FE-identified models, such as the Mooney-Rivlin and Arruda-Boyce models in this case, to assess a static stiffness measure for the design of a component under different load configurations and strain ranges. Moreover, the examination of relative error when artificial noise was added to the test data indicates that the level of error in FE-models is comparable to that seen in experimental tests.

The Mooney-Rivlin strain energy function parameters, which were identified after introducing noise (5% error to the DIC experimental data), are given in Table 5.7. The relative error shown in this table was calculated with respect to the model parameters identified without artificial noise in the test data. Therefore, according to this numerical analysis, the identification of model parameters is not overly sensitive to measurement error; since error of ±5% is well within the range of potential error in a well-controlled DIC test, this level of error is acceptable for numerical analysis. Likewise, the speckle points applied to the surface were approximately 3 to 5 pixels (less than 0.05mm) in diameter; the corresponding subset size chosen was 15 pixels. According to Tong and Robert, RMS error of 5% is a safe estimate of the measurement error for a speckle pattern with speckle sizes smaller than the subset size (Tong, 2005) (Robert, 2007)).
Table 5.7: Sensitivity of the identified model parameters with introduced noise in the test data

<table>
<thead>
<tr>
<th>Identified Mooney-Rivlin model without artificial noise in test data</th>
<th>Identified Mooney-Rivlin model with artificial noise in test data</th>
<th>Relative error</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.578129</td>
<td>0.581362</td>
<td>0.56%</td>
</tr>
<tr>
<td>0.452273</td>
<td>0.474483</td>
<td>4.91%</td>
</tr>
<tr>
<td>0.123581E-03</td>
<td>0.136529E-03</td>
<td>10.48%</td>
</tr>
</tbody>
</table>

5.5 Summary

This chapter provides a methodology for rapid model selection and identification of model parameters for rubber-like materials in structural components. This method is based on an inverse problem approach which uses displacement/strain data measured by digital image correlation (DIC) as input. A least-squares objective function is minimized by use of a trust region-based derivative-free optimization method. An engine mount with rubber components was tested to demonstrate and validate this approach.

The concept proposed in this chapter is a flexible methodology for the identification of effective material models of rubber-like materials. The identifiability of model parameters and numerical stability issues arising from this approach are discussed; several numerical analyses are recommended to validate model identification.

In particular, this chapter considers the identification of constitutive parameters of rubberlike materials under monotonic loading. Future research would include the identification of constitutive parameters of rubber-like materials under cyclic loading, enabling study of parameters defining hysteresis and the Mullins effect. The Mullins effect is closely related to the fatigue of elastomeric parts used in engineering
applications. Quantitative study of the Mullins effect is thus a necessary step towards the scientific evaluation of the fatigue life of rubber and rubber-like materials.
CHAPTER SIX: CONCLUSION AND FUTURE WORK

Contribution is sometimes a reward in itself. Recognition of the value of an idea is a further reward. Edward de Bono

This thesis solves two inverse problems: (i) reconstruction of stiffness distribution of beams which can serve as damage identification and structural monitoring technique, and (ii) the identification of nonlinear material model parameters of components/specimens which is a topic in more and more demanding as engineers today have the ambition to describe comprehensively the nonlinear deformation processes of materials and structures. The two inverse problems are different, one is ill-posed functional identification and one is well-posed parameter identification, although the former must also be solved in parametric ways.

In this work, optically measured beam deflection profiles and full-field displacements/strains on the specimen surface using digital cameras are used as inputs to inverse identification processes. The author believes that optical measuring techniques are very promising, and whence combined with modern parameter identification techniques, offers great potentials to engineering problems including damage and material identification.

A novel methodology of damage identification is developed using static deflection measurements. Two FE model updating methods for beams were proposed in this work; they can be used as damage identification methods. The methods presented in this work depart from traditional damage identification techniques by trying to
reconstruct the complete stiffness distribution of a beam. Although a baseline model is required, only boundary conditions need to be accurately modeled.

Throughout this work, a number of “future work” have appeared in text. This is not an indication of a lack of effort; rather, it reflects the fact of the research,

1) Measurements: photogrammetry, camera calibration, metric cameras, other measurement techs, fibre optic, inclinometers, etc.

2) Additional field testing: the primary focus of future efforts should be on continued testing with field testing on real structures, to see how well the methods will perform in practice. Loading is a problem, especially bridges, issues like lighting, requirement of stable mounting base, calibration of cameras may arise and need to be studied in practice. Another interesting issue can be the permanent mounting of inspecting cameras and data transition through cables, in a time that cameras are installed everywhere for security of humans, it would a more though to use cameras for safety of structures.

3) Combination of different damage identification techniques: local+global, dynamic + static, irregularity detection from static deflection measurements.
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Li Li was born in 1973 in Hefei, China. He completed a engineer degree in construction management at the Anhui University, Hefei, China, in 1992 before he worked as a highway and bridge construction superintendent and project manager in Anhui Provincial Road and Bridge Construction Company, Anhui Provincial Ministry of Highway Transportation from 1992 to 1995. He went on to the Xi’an Highway University, Xi’an, China, where he obtained a Master of Engineering degree in Bridge and Tunnel Engineering in 1998. Afterwards he had worked as bridge engineer at a consulting company in Beijing, China.

Li Li came to Canada in 2003 and obtained a Master of Applied Science degree in Civil Engineering in 2005 at the University of Windsor, Windsor, Ontario. Presently Li Li is a candidate for Doctor of Philosophy degree with specialization in Structural Engineering at the University of Windsor, Windsor, Ontario; he finished his final defense on September 11 and expects to graduate in October 2012.