Bayesian Modelling and Analysis of Utility-based Maintenance for Repairable Systems

A thesis submitted to the University of Dublin, Trinity College in partial fulfillment of the requirements for the degree of Doctor of Philosophy

Department of Statistics, Trinity College Dublin

September 2016

Shuaiwei Zhou
I dedicate this thesis to my wonderful and supportive parents
Declaration

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Shuaiwei Zhou

Dated: September 12, 2016
Abstract

This thesis focuses on modelling and inference for maintenance systems for the purpose of utility optimisation. Providing standardised notation throughout, we first demonstrate the motivation for investigating the problem of modelling and inference for maintenance systems and briefly state the problems which are to be explored. The definitions and terminology, which are also used within the general domains of science and engineering, have been presented in terms of statistical representation.

We propose a Bayesian method to optimise the utility of a two phase maintenance system sequentially by dynamic programming method. In particular, the parameters of the failure distribution for the system of interest are analysed within the Bayesian framework. Utility-based maintenance is modelled in several modified models, including imperfect preventive maintenance, time value of money effect in maintenance, maintenance for systems with discrete failure time distributions, maintenance for parallel redundant systems, of which all follow numerical examples. A hybrid approach combining myopic and dynamic programming method is proposed to solve multi-phase maintenance systems.

The Bayesian dynamic programming is carried out through the gridding approach to solve the issue arising from nested series of maximisations and integrations over a highly non-linear space. The core of gridding method, the increment is studied extensively. We also utilise and modify the approach proposed by Baker (2006) to analyse the effect of risk aversion on the variability of system in cash flows.

The potential generalisation of the current models has been discussed and the future work concerning complicated models and efficient computation methods have also been indicated.
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I would like to express my utmost and sincere gratitude to my two wonderful supervisors, Professor Simon P. Wilson and Professor Brett Houlding, for their invaluable guidance and constant devotion to me during my PhD research. No matter how stupid the questions I asked or how naïve I was, they were always extremely patient to explain and explore potential ideas with me.

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Shuaiwei Zhou

Trinity College, Dublin

September 2016
## Abbreviations

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<tr>
<td>ARA</td>
<td>Absolute Risk Aversion</td>
</tr>
<tr>
<td>CARA</td>
<td>Constant Absolute Risk Aversion</td>
</tr>
<tr>
<td>CBM</td>
<td>Condition-based Maintenance</td>
</tr>
<tr>
<td>CDF</td>
<td>Cumulative Distribution Function</td>
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<tr>
<td>CE</td>
<td>Certainty Equivalent</td>
</tr>
<tr>
<td>CHF</td>
<td>Cumulative Hazard Function</td>
</tr>
<tr>
<td>CM</td>
<td>Corrective Maintenance</td>
</tr>
<tr>
<td>CR</td>
<td>Cost Rate</td>
</tr>
<tr>
<td>CRRA</td>
<td>Constant Relative Risk Aversion</td>
</tr>
<tr>
<td>DARA</td>
<td>Decreasing Absolute Risk Aversion</td>
</tr>
<tr>
<td>DP</td>
<td>Dynamic Programming</td>
</tr>
<tr>
<td>DRRA</td>
<td>Decreasing Relative Risk Aversion</td>
</tr>
<tr>
<td>GA</td>
<td>Genetic Algorithm</td>
</tr>
<tr>
<td>H-M-DP</td>
<td>Hybrid Myopic Dynamic Programming</td>
</tr>
<tr>
<td>IARA</td>
<td>Increasing Absolute Risk Aversion</td>
</tr>
<tr>
<td>IID</td>
<td>Independent and Identically Distributed</td>
</tr>
<tr>
<td>IPM</td>
<td>Imperfect Preventive Maintenance</td>
</tr>
<tr>
<td>IRRA</td>
<td>Increasing Relative Risk Aversion</td>
</tr>
<tr>
<td>PDF</td>
<td>Probability Density Function</td>
</tr>
<tr>
<td>PM</td>
<td>Preventive Maintenance</td>
</tr>
<tr>
<td>PMF</td>
<td>Probability Mass Function</td>
</tr>
<tr>
<td>PPM</td>
<td>Perfect Preventive Maintenance</td>
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<tr>
<td>RRA</td>
<td>Relative Risk Aversion</td>
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Chapter 1

Introduction

1.1 Background and Motivation

In the current climate of globalisation, competition as well as varying demands from stakeholders, increasing pressure on manufacturing performance has been one of the main driving forces in the growth of manufacturing industries (Tsang 2002). From this point of view maintenance is a significant activity in industrial practice, resulting in the importance of maintenance optimisation. Maintenance aims to combat the inevitable degradation of systems over their operational lifetime and keep them in working order. Therefore, maintenance plays an important role in sustaining and improving systems availability, which in turn affects the productivity of the system of interest.

Recently, more attention has been directed towards improving and optimising maintenance in manufacturing systems as inappropriate maintenance could result in huge cost and risk (Holmberg et al. 2010). Maintenance costs can reach anywhere between 15% and 70% of production costs (Wang et al. 2008), which also indicates that there is still a large potential for increasing the productivity in current maintenance practices. In some industries, a slight improvement in throughput could result in significant economic impact.

In modern times, the complexity of maintenance systems has increased drastically, see Duffuaa et al. (2001). This is partly due to modern manufacturing systems which involve numerous interactions and dependencies between components. It is evident that analytical and mathematical approaches are limited in solving such complex maintenance problems. When it comes to maintenance optimisation methods, sequential
analysis is applied to use accumulating evidence to make advantageous early decisions. In the context of system engineering, this could help save cost and even improve system performance. The Bayesian method of sequential analysis is to make decisions that minimise the expected value of some loss function which can be viewed as a function of corresponding inputs and outputs, see DeGroot (1970) and Brockwell and Kadane (2003). In this thesis, we focus the study on Bayesian sequential analysis applied to maintenance optimisation of repairable systems.

The study of system maintenance has attracted increasing attention in recent years because of a need from industry for increasing the reliability and availability of systems whilst decreasing the associated costs. Percy and Kobbacy (1996) pioneered work in preventive maintenance modelling from a Bayesian perspective. Damien et al. (2007) analysed a single item maintenance in a Bayesian semi-parametric setting, which solves the drawbacks of other models failing to capture the true underlying relationships in the data. However, their analysis is based on a pre-defined finite time horizon, for example, see Baker (2010); in other words, the maintenance time phases are pre-defined which is not practical in reality; in our work, on the contrary, the maintenance time phases are also pre-defined depending on a particular system but random and flexible, which meets the maintenance scheduling programme. Nonparametric methods have also been investigated in system maintenance. Gilardoni et al. (2013) use a power-law-process parametric method by incorporating the nonparametric maximum likelihood estimate of an intensity function to estimate the optimal preventive maintenance policy. However, all these approaches fail to consider sequential maintenance which requires more complicated modelling and longer computation time.

Maintenance based on prognostics is a prior event analysis and action. By means of incorporating prognostics into the maintenance decision making process, one could carry out a maintenance forecast based on known characteristics as well as the evaluation of the significant parameters of the item. With regard to maintenance objectives, as in most of the literature cost-based optimisation framework is taken (Van Horenbeek et al. 2010). However, focus should not only be on costs as risk preference is simply ignored if only cost-oriented objective is taken into account. Utility functions used to measure risk preferences are ubiquitous in economic research. The little published work that is the exception occurs in warranty and inventory, see Padmanabhan and
Rao (1993); Keren and Pliskin (2006). In fact, the field of maintenance and reliability is a suitable area to apply risk-averse policies because there are numerous cash flows occurring stochastically. A drawback of taking cost per unit time as a criterion of optimality is that two policies might then be equally attractive, even if for one of them the annual maintenance spend were much more variable than that for the other. What might be seen by some as over-maintenance, in the sense that mean cost per unit time is not minimised, could be optimal as a risk-averse policy, in which the large unscheduled losses from failure have such a dis-utility that very frequent maintenance is carried out. Clearly, a policy that minimised cost per unit time would be unsatisfactory for a maintenance engineer who could not convince management that periods of high loss were an unavoidable part of an optimal long-term policy or for an enterprise that could not survive because of short-term cash flow problems. Thus, extra maintenance activity is an insurance policy against large losses occurring over a period.

Models and methodologies proposed in this thesis are primarily suited for large industrial purposes, for example, an automatic manufacturing system, a robotic process, or a computer server for the non-life essential services, in which cases failure is neither rare or frequent, maintenance itself is not cheap or trivial, but failure is a considerable expense, though not exorbitantly so. Hence, this approach is not suitable to apply to maintenance of systems with very high risk aversion properties, e.g., a nuclear power facility, an off-shore oil field, or a life support system. It is also not worthwhile applying to trivial systems where the computational cost of performing this analysis outweighs any savings.

1.2 Structure and Main Contributions

The overall research goal of this thesis is to develop a utility-based prognostic maintenance optimisation methodology within a Bayesian statistical framework, which uses historical information and predictions in remaining lifetime of repairable systems.

The following is an overview of the structure of this thesis along with the main research contributions.

- Chapter 1 introduces the research background and motivation, and briefly outlines the structure and main contributions of the thesis.
- Although research background and motivation is given in Chapter 1, Chapter 2 gives a detailed review on maintenance modelling and analyses as well as the fundamental concepts in systems maintenance, from reliability measures to classical failure time distributions. We highlight the essential publications which are highly related to the research questions in this thesis.

- For those beyond the statistical research community, Chapter 3 briefly presents the Bayesian perspective on modelling, and continues to introduce the foundational concepts of the dynamic programming method as well as utility theory, which are the maintenance optimisation methodologies in this research.

- Chapter 4 solves the sequential maintenance problem under the policy of perfect preventive maintenance by a dynamic programming method utilising the idea proposed by Brockwell and Kadane (2003), whereby a grid is constructed in the maintenance and failure time space, over which the utility functions of expected cost per unit time are evaluated. This method has a computation time which is linear in the number of phases in the sequential problem.

- Chapter 5 extends the previous sequential preventive models to imperfect preventive maintenance, taking account of the time value of money, modelling preventive maintenance in discrete time setting as well as maintenance for parallel systems, analysing the effect of failure time distribution assumptions on preventive maintenance time and proposing an adaptive approach to solving multi-phase systems’ maintenance. Sensitivity analysis via the parameters of sequential preventive maintenance models will also be carried out in the chapter.

- Chapter 6 utilises and modifies the approach proposed by Baker (2006) to investigate and analyse the effect of risk aversion on the variability of system in cash flows from a certainty-equivalent point of view.

- Chapter 7 states the major conclusions and contributions of this research and suggests future work directions.
Chapter 2

Repairable Systems and Maintenance

In this chapter we introduce the difference between non-repairable systems and repairable systems, and classify the maintenance policies and review related modelling methods.

2.1 Repairable Systems

2.1.1 Basic Terminology and Examples

A repairable system is a system that can be restored to an operating condition by some repair process instead of replacement of the entire system. For example, an automobile is a repairable system because most failures, such as the inability to start because of a bad starter, can be fixed without replacing the entire automobile. Repair does not have to involve replacement of any parts. For instance, the automobile may fail to start because of a bad connection with the battery. In this case cleaning the cables and their connectors with the battery may solve the problem. On the other hand, a light socket is not considered as a repairable system. The only way to repair a burned-out light is to replace the bulb; in other words, replace the entire system.

A non-repairable system is one which is discarded after failure. A light bulb is a non-repairable system for example. Today, with automated production processes being implemented in industry, many products that used to be repaired after failure are now
discarded when they fail. Consider for example a small desk-top fan which can be purchased for less than 10 euro at a discount shop. When such a unit fails, we would probably discard it and buy another, because the cost of fixing it is greater than that of purchasing a new one. Many electrical systems are now non-repairable, or they are more expensive to repair than to replace.

A few definitions used in repairable systems are given below.

**Definition 2.1 Global time**  
Failure of a repairable system is measured in global time if the failure times are recorded as time since the initial start-up of the system. Failures in global time will be denoted by $X_1 < X_2 < \cdots$.

**Definition 2.2 Local time**  
Failure times of a repairable system are measured in local time if the failure times are recorded as time since the previous failure. Failures in local time will be denoted by $T_1, T_2, \ldots$.

Local time is mainly used in the following work, unless explicitly stated.

**Definition 2.3 Deterioration and Improvement**  
We say that a repairable system is deteriorating if the times between failure tend to get shorter with ageing. If the times between failure tend to increase, then we will say that the system is improving.

### 2.1.2 Reliability Measures

There are similarities and differences between repairable and non-repairable systems. A few issues are clarified to understand repairable system behaviour as follows.

For a non-repairable system the lifetime of the system is a random variable. As there is no repair, the system would be discarded after its one and only failure, and if it does not have an impact on the performance of a similar system located elsewhere, then the assumption that different systems have lifetimes that are independent is reasonable. Also, if many copies of the system were produced by the same manufacturing process, then it is also reasonable to assume that the system lifetimes have the same distribution. These two assumptions can be combined into one statement that says the lifetimes are independent and identically distributed (IID) from some distribution having cumulative distribution function (CDF) $F(x)$. 
Definition 2.4 Cumulative Distribution Function The cumulative distribution function (CDF) of a random variable \( X \) is defined to be the function

\[
F(x) = P(X \leq x).
\]

Since the lifetime must be nonnegative the probability distribution must have positive probability on the positive axis only. In other words, \( F(x) = 0 \) for \( x < 0 \).

Definition 2.5 Survival Function The survival function \( S(x) \), also called the reliability function, is the probability that a system will carry out its mission through time \( x \).

The survival function evaluated at \( x \) is just the probability that the failure time is beyond time \( x \). Thus the survival function is related to the CDF in the following way:

\[
S(x) = P(X > x) = 1 - P(X \leq x) = 1 - F(x). \tag{2.1}
\]

Definition 2.6 Probability Density Function The probability density function (PDF) is defined to be the derivative of the CDF, provided that the derivative exists. That is,

\[
f(x) = \frac{d}{dx} F(x) = -\frac{d}{dx} S(x).
\]

Another way to express the PDF is through the limit

\[
f(x) = \lim_{\Delta x \to 0} \frac{F(x + \Delta x) - F(x)}{\Delta x} = \lim_{\Delta x \to 0} \frac{P(x < X \leq x + \Delta x)}{\Delta x}. \tag{2.2}
\]

Another important function related to, but distinct from the PDF, is the hazard function.

Definition 2.7 Hazard Function The hazard function is

\[
h(x) = \lim_{\Delta x \to 0} \frac{P(x < X \leq x + \Delta x | X > x)}{\Delta x}. \tag{2.3}
\]

This is the limit of the probability per unit time that a unit fails (for the first and only time) in a small interval given that it has survived to the beginning of the interval. Compare the definition of the hazard function \( h(x) \) in (2.3) with the result for the pdf
given in (2.2). These are nearly the same, except one is a conditional probability and
the other is not.

One property of a PDF is that it must integrate to 1; that is, since we are dealing
with random variables that have all the probability on the nonnegative axis,

$$\int_0^\infty f(x)dx = 1.$$ 

The hazard is defined as the limit of a conditional probability, but it is not a conditional
probability density function. The hazard function does not need to integrate to 1, and
in fact, for most distributions we study, the hazard will not integrate to 1 but infinity
(see Cumulative Hazard Function). For a system whose hazard function is increasing,
this means that (in the limit) the probability of failure in a small interval divided by
the length of the interval is increasing with time. Thus if we take a small fixed length
of time, such as one hour, an increasing hazard would mean that the probability of
failing in this one hour, given that the system survived past the start of that hour,
increases with the age of the system. In this case we say that the system is wearing
out. Compare this definition with that of deterioration for a repairable system. We
say that a repairable system deteriorates when the times between failures tend to get
smaller, and we say that a non-repairable system is wearing out if the hazard function
is increasing. A non-repairable system with a decreasing hazard function is said to
experience burn-in. The term “deteriorate” will be reserved for repairable systems
and the term “wear out” will be reserved for non-repairable systems. Similarly, the
terms “improvement” and “burn-in” will be reserved for repairable and non-repairable
systems, respectively. Also note, for a continuous random variable the hazard function
can be defined as

$$h(x) = \frac{f(x)}{S(x)}.$$ 

Knowing any one of the pdf $f(x)$, the cdf $F(x)$, the survival function $S(x)$, or the
hazard function $h(x)$ is enough to find all of the others.

**Definition 2.8 Cumulative Hazard Function** The quantity

$$H(x) = \int_0^x h(x)dx$$

is called the cumulative hazard function (CHF).
As $t$ tends to infinity, i.e., $S(x)$ tends to 0, the cumulative hazard function increases without bound, which implies that $h(x)$ must not decrease too quickly, otherwise, $H(x)$ will converge.

### 2.1.3 Classical Failure Distributions

The next section covers some of the commonly used distributions for lifetime, including the exponential, the Weibull, and the gamma.

#### Exponential Distribution

The simplest model for lifetimes is the exponential distribution.

**Definition 2.9 Exponential distribution**  *The exponential distribution is a continuous distribution having pdf*  

$$f(x) = \lambda \exp(-\lambda x), \quad x > 0$$

*and cdf*  

$$F(x) = P(X \leq x) = \int_0^x \lambda \exp(-\lambda t) dt = 1 - \exp(-\lambda x), \quad x > 0. \quad (2.4)$$

We write $X \sim \text{EXP} (\lambda)$, where $\lambda$ is often referred to as a rate parameter, to indicate that the random variable $X$ has an exponential distribution with a CDF given by (2.4).

The mean and variance of the exponential distribution are $\frac{1}{\lambda}$ and $\frac{1}{\lambda^2}$, respectively. The most distinctive feature of the exponential distribution is that it is the only continuous distribution with the memoryless property.

**Definition 2.10 Memoryless property**  *A distribution has the memoryless property if*  

$$P(X > t + x \mid X > t) = P(X > x).$$

In other words, if the distribution has the memoryless property, then for instance, the probability that an old unit survives one more day will equal the probability that a brand new unit will survive one day. The memoryless property imposes some strong assumptions about the way units age.

Another unique feature of the exponential distribution is that it is the only continuous distribution with a constant hazard function.
Weibull Distribution

We discuss here the Weibull distribution for several reasons. First, it is probably the most widely used distribution for lifetimes. Second, if repairs bring a system back to a good-as-new state and the times between failures $X_1, X_2, \ldots$ are independent, then the assumption that the times between failures are iid Weibull random variables may be reasonable because Weibull is a versatile distribution that can take on the characteristics of other types of distributions.

**Definition 2.11** The Weibull distribution has survival function

$$S(x) = \exp \left\{ -\left( \frac{x}{\alpha} \right)^\eta \right\}, \quad x > 0.$$  \hfill (2.5)

If $X$ is a random variable with this survival function, then we will write $X \sim \text{WEI}(\eta, \alpha)$, where $\eta$ and $\alpha$ are the shape and scale parameters, respectively.

The cdf, pdf and hazard functions are therefore given as follows:

$$F(x) = 1 - S(x) = 1 - \exp \left\{ -\left( \frac{x}{\alpha} \right)^\eta \right\}, \quad x > 0$$  \hfill (2.6)

$$f(x) = F'(x) = \frac{\eta}{\alpha} \left( \frac{x}{\alpha} \right)^{\eta-1} \exp \left\{ -\left( \frac{x}{\alpha} \right)^\eta \right\}, \quad x > 0$$  \hfill (2.7)

$$h(x) = \frac{f(x)}{S(x)} = \frac{\frac{\eta}{\alpha} \left( \frac{x}{\alpha} \right)^{\eta-1} \exp \left\{ -\left( \frac{x}{\alpha} \right)^\eta \right\}}{\exp \left\{ -\left( \frac{x}{\alpha} \right)^\eta \right\}} = \frac{\eta}{\alpha} \left( \frac{x}{\alpha} \right)^{\eta-1}, \quad x > 0.$$  \hfill (2.8)

The hazard function $h$ is increasing when $\eta > 1$ and decreasing when $\eta < 1$. When $\eta = 1$, the hazard function is the constant function $h(x) = 1/\alpha$. Thus, the exponential distribution is a special case of the Weibull distribution that occurs when $\eta = 1$. 

Figure 2.1: Comparison of Weibull Distributions with Different Shape Parameters $\eta$: Probability Density Functions (top-left), Cumulative Distribution Functions (top-right), Survival Functions (bottom-left), Hazard Functions (bottom-right).

When the scale parameter $\alpha = 1$, Figure 2.1 shows a number of Weibull probability density functions, cumulative distribution functions and corresponding hazard functions, respectively.

The mean and variance of the Weibull can be expressed in terms of the gamma function which is defined below.

Definition 2.12 Gamma Function For $a > 0$ the gamma function is defined to be

$$\Gamma(a) = \int_0^\infty x^{a-1}e^{-x}dx.$$  

The next theorem gives the mean and variance of the Weibull distribution in terms of the gamma function.

Proposition 2.1 If $X \sim WEI(\eta, \alpha)$, then

$$E(X) = \alpha \Gamma \left( 1 + \frac{1}{\eta} \right)$$  \hspace{1cm} (2.9)
See Rigdon and Basu (2000) for proofs.

**Gamma Distribution**

The gamma distribution is another useful model for the lifetime of systems.

**Definition 2.13** The pdf for the gamma distribution can be written as

\[
f(x) = \frac{x^{\eta-1}}{\theta^\eta \Gamma(\eta)} \exp(-x/\theta), \quad x > 0.
\]

We will write \( X \sim \text{GAM}(\eta, \theta) \) if the random variable \( X \) has this pdf, where \( \eta \) and \( \theta \) are the shape and scale parameters, respectively. Another useful form for the gamma pdf is obtained by substituting \( 1/\lambda \) for \( \theta \); this gives

\[
f(x) = \frac{\lambda^\eta x^{\eta-1}}{\Gamma(\eta)} \exp(-\lambda x), \quad x > 0.
\] (2.11)

The cdf and the survival function, and hence also the hazard function, cannot be written in closed form. We can write the cdf as

\[
F(x) = \int_0^x \frac{\lambda^\eta \omega^{\eta-1}}{\Gamma(\eta)} \exp(-\lambda \omega) d\omega.
\]

If we make the transformation \( y = \lambda \omega \), then this becomes

\[
F(x) = \int_0^{\lambda x} \frac{\lambda^\eta (y/\lambda)^{\eta-1}}{\Gamma(\eta)} e^{-y/\lambda} \frac{1}{\lambda} dy
\]

\[
= \frac{\lambda^\eta}{\lambda^{\eta-1}\Gamma(\eta)} \frac{1}{\lambda} \int_0^{\lambda x} y^{\eta-1} e^{-y} dy
\]

\[
= \frac{1}{\Gamma(\eta)} \int_0^{\lambda x} y^{\eta-1} e^{-y} dy.
\]

The hazard function is therefore

\[
h(x) = \frac{f(x)}{1 - F(x)}
\]

\[
= \frac{\frac{\lambda^\eta x^{\eta-1}}{\Gamma(\eta)} \exp(-\lambda x)}{1 - \frac{1}{\Gamma(\eta)} \int_0^{\lambda x} y^{\eta-1} e^{-y} dy}.
\]

This hazard function is increasing when \( \eta > 1 \), decreasing when \( \eta < 1 \), and constant when \( \eta = 1 \), when the corresponding pdf is that of the exponential distribution.
When the scale parameter $\theta = 1$, Figure 2.2 shows a number of gamma probability density functions, cumulative distribution functions and corresponding hazard functions, respectively.

### 2.2 Maintenance Modelling

Traditional repairable systems assume a whole range of performance levels, varying from perfect functioning to complete failure, and assuming the repair is perfect. However, many manufacturing systems suffer increasing wear with usage, age or deterioration, that is, perfect functioning is not always satisfied. Therefore, maintenance management, as an important policy for a repairable system, is widely used to keep systems in good condition, to decrease failures, and increase system availability. Based on the European standard (EN 13306:2010), a definition of maintenance management is given as follows:
Definition 2.14 **Maintenance Management** depicts all activities of the management that determine the maintenance objectives, strategies and responsibilities, and implementation of them by such means as maintenance planning, maintenance control, and the improvement of maintenance activities and economics.

According to the definition above, the major steps to maintenance modelling can be summarised as:

1. Determine the maintenance objective(s).

2. Define or select maintenance policies according to measures of system performance.

3. Plan, control and improve maintenance.

### 2.2.1 Maintenance Policies

The availability and usability play a crucial part in a system’s performance because any breakdowns and holdups can seriously impede its performance. At the same time, idle systems negatively affect the ratio between fixed cost to output. The reduced output induced by system breakdowns would result in less production as well as less profitability which can be regarded as an inefficiency for the system. Moreover, complex systems usually require a significant startup time after an interruption occurs. Possibly during this period of time, goods that do not meet acceptable levels, *e.g.*, scrap or goods of minor quality are produced, as a result, one cannot obtain her or his expected profit since these products cannot be sold or have to be sold at reduced prices. Thus, efficient operation of a system requires well-scheduled maintenance to avoid interruptions as much as possible and to recover from breakdowns quickly.

For a manufacturing system, wear-out, ageing and deteriorating will have a negative impact on the function of the system, which results in the consequence that the system cannot fulfil its capability. Maintenance is introduced to counteract those negative effects from an economic point of view. Therefore, maintenance actions plays an essential role in sustaining and possibly improving a system’s availability, which in return will improve the productivity of the system considered. In general, maintenance policies and strategies are commonly categorised into three domains: Corrective Maintenance (CM), Preventive Maintenance (PM) and Condition based Maintenance (CBM).
Corrective maintenance is initiated when the system sees a breakdown which results in a stop for a system working and induces considerable cost. Corrective maintenance is usually named repair, restoration or replacement of failed components. This maintenance policy is often applied to systems of which failure is not costly and do not result in disastrous situations, for components with constant failure rate, *e.g.*, if a failure time of components is assumed to follow an exponential distribution.

Preventive maintenance is implemented for the purpose of minimising the negative impact of an unexpected breakdown. Generally speaking, preventive maintenance usually involves less resource consumption compared to that of corrective maintenance and it can be designed in the production plans of the system of interest. PM includes all partial or complete overhauls, such as filter cleaning, oil charging, *etc.* in order to prevent a critical failure that is costly before it actually occurs. It can be seen that preventive maintenance makes sense in the situation when the failure rate of a unit or component is increasing in time. Unlike CM that is unexpected, preventive maintenance can usually be properly planned and prepared. Although preventive maintenance is incorporated to prevent critical failures in system designing, sometimes failure may still be seen. As a result, it is usually suggested to combine both corrective maintenance and preventive maintenance tasks.

However, when the operation schedules and environmental variables change in practice, exhaustive or unnecessary use of preventive maintenance can occur. To make sure that preventive maintenance is taken only when it is required, condition based maintenance was introduced by incorporating inspections of the system of interest in pre-determined intervals to determine the system’s operation condition. Depending on the outcome of an inspection, relevant maintenance tasks can be implemented. It is worthwhile noting that CBM is sometimes analysed in the field of PM ([Manzini et al., 2010](#)).

The preventive maintenance policies include time based PM ([Roux et al., 2008](#)) in which PM is conducted every \( t \) units of time and age based PM ([Chen et al., 2006](#)) where PM is carried out every \( t \) units of operating time. There are other alternatives of preventive maintenance, *e.g.*, for non-repairable systems, group block replacements where units or components would be replaced if it failed whereas the other working components would be replaced at pre-determined schedule ([Roux et al., 2008](#)).
Condition based maintenance has received less attention probably because it is relatively new compared to CM and PM. However, thanks to the fact that the inspection is less costly, one is encouraged to implement CBM (Xiang et al., 2012). If a system is designed to serve for a long period, one inspection monitor can be installed if it is relatively cheaper. Van Horenbeek and Pintelon (2013) proposed a prognostic maintenance by combining CBM with the prediction about the states of components to see if a threshold is expected to be reached before the following scheduled inspection. If it does, the component is replaced immediately. Although it can be seen that there is an increasing application of CBM in practice (Wang et al., 2008), it is less studied in the literature.

In reviewing the literature, we find limited effort was taken to compare different maintenance policies. Xiang et al. (2012) investigated a repairable system under preventive maintenance and condition based maintenance policies and found that condition-based maintenance is superior to scheduled maintenance paradigm via simulation. Van Horenbeek and Pintelon (2013) studied five different maintenance policies (i.e., CM, block PM, age based PM, CBM with inspection and CBM under continuous monitoring) on one machine and their noted effects.

2.2.2 Maintenance Effectiveness

Preventive maintenance comprises all maintenance activities which are not triggered by a system failure. Not only the mode of maintenance task (preventive maintenance or corrective maintenance) and its associated maintenance interval impact the failure rate, but also its level of quality (effectiveness of maintenance task). The state after a maintenance action is performed on a component is assumed to be: perfect, imperfect, minimal, worse or worst (Pham and Wang, 1996). See Table 2.1.
<table>
<thead>
<tr>
<th>Maintenance Policy</th>
<th>System State</th>
<th>Failure Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preventive maintenance</td>
<td>The system state is restored to be “as good as new”</td>
<td>Decreasing of the failure rate</td>
</tr>
<tr>
<td>Imperfect maintenance</td>
<td>A maintenance action that restores the system to a state somewhere between “as good as new” and “as bad as old”</td>
<td>Decreasing of the failure rate</td>
</tr>
<tr>
<td>Minimal Maintenance</td>
<td>The system state is “as bad as old”</td>
<td>No effect on the failure rate</td>
</tr>
<tr>
<td>Worse Maintenance</td>
<td>System is in operating state worse than just prior to the maintenance action</td>
<td>Increasing of the failure rate</td>
</tr>
<tr>
<td>Worst maintenance</td>
<td>System breaks down right after maintenance action</td>
<td>Increasing of the failure rate</td>
</tr>
</tbody>
</table>

Table 2.1: State assumptions after maintenance

Imperfect, worse or worst maintenance can be caused by repairing parts of a system mistakenly. In addition to one of the best known models proposed by Brown and Proschan (1983), Pham and Wang (2006) reviewed other approaches to model imperfect maintenance, such as \((p, q)\) rule, \((p(t), q(t))\) rule, improvement factor, and virtual age approach, etc. Briefly, with probability \(p\), the item is restored to the as good as new state or otherwise to the as bad as old state with probability \(q = 1 - p\). A novel approach to modelling imperfect preventive maintenance will be proposed in Chapter 5.

Quality levels of worse and worst maintenance are related to maintenance and repair induced failures. Preventive maintenance actions, as cleaning or greasing, mitigate the deterioration effect of some failure mechanics and restore components to a “as good as new” condition with respect to some failure mechanisms only. All other failure
mechanisms will remain unaffected. Hence, Lin et al. (2001) introduced the concept of two categories of failure mechanisms, maintainable failure mechanisms and non-maintainable failure mechanisms. Preventive maintenance will affect maintainable failure mechanisms exclusively, whereas non-maintainable mechanisms remain unaltered. Zequeira and Berenguer (2005) stated that the maintainable and non-maintainable failure rates are dependent and restored the system of interest to a condition between as good as new and as bad as old via their proposed preventive maintenance actions.

In the literature, three approaches for modelling the impact of preventive maintenance on the failure rate have been studied extensively; a failure rate model by Lie and Chun (1986) and Nakagawa (1986), Nakagawa (1988), an age reduction model by Canfield (1986) and Malik (1979) and a hybrid model by Lin et al. (2001).

In maintenance modelling, most researchers model a system as a whole unit, without considering the effect of deterioration and failure on the subsystems. On the other hand, machines were modelled as subsystems by some researchers. Van Horenbeek and Pintelon (2013) modelled a simplified version only considering one subsystem in a few machines and analysed the structural and stochastic dependencies. Roux et al. (2008) evaluated the impact of three maintenance policies under an assumption that a system has only two independent components. One of the assumptions in maintenance modelling is assuming all the units or components of a system are identical and independent. There are other less rigorous assumptions and we will include some of them in our research:

- Perfect maintenance: preventive maintenance is assumed to be done perfectly and it is often referred to as ‘systems are are good as new’ after preventive maintenance is conducted. In this thesis, perfect preventive maintenance (PPM) is equivalent to corrective maintenance (CM) in the sense that they have the same consequence with regard to the state of the system after maintenance.

- Time of implementing maintenance is assumed to be constant and often is even reduced to be negligible implying that maintenance is carried out instantaneously.

- Costs of all relevant maintenance tasks are assumed to be known as constant and the cost of CM is always more than that of PM.

- Relevant resources such as spare components are assumed to be sufficient to
• System failures are assumed to be detected immediately.

2.2.3 Sequential Maintenance

Under sequential maintenance, systems may be maintained at unequal time intervals, in contrast to periodic maintenance in which the maintenance interval is fixed and unique.

Barlow and Proshan (1965) design a sequential maintenance policy for a finite time span. Under this maintenance policy, the time for which maintenance is scheduled depends on the time remaining rather than the pre-determined and identical maintenance time. In addition, the next maintenance interval is determined by minimising the average cost per unit time during the remaining time. Therefore, this policy does not determine systems’ future maintenance intervals at the initial time of system performance, which adds flexibility and reduces systems costs.

Nguyen and Murthy (1981) model preventive maintenance under a sequential policy in which a failure needed to be corrected has not occurred by a reference time $t_i$, where $t_i$ is the maximum time that a system should be fully replaced without extra maintenance after the $(i - 1)^{th}$ repair; in other words, the system is replaced after $(k - 1)$ repairs. The system is repaired (or replaced at the $k^{th}$ repair) at the time of failure or at time $t_i$, whichever occurs first.

Nakagawa (1986) propose and compare periodic and sequential preventive maintenance policies for the system with minimal repair at failure and compute the maintenance intervals in a Weibull distribution case, indicating that a sequential preventive maintenance policy is superior to a periodic one. Schutz et al. (2011) extend previous research by considering a system performing a wide range of missions over a finite planning horizon and a dynamic system failure law is taken into account to modelling the different missions with various characteristics depending on the operational conditions.

Lin et al. (2000) propose general sequential imperfect preventive maintenance models, in which the effective remaining time of the system would be reduced whilst the hazard rate would be adjusted after each preventive maintenance. We refer it to as a hybrid sequential preventive maintenance model because it considers both time reduction and
hazard adjustment on systems.

There are also studies by Dieulle et al. (2003) calculating the long-time expected cost per unit of time via considering if system’s state is above or below a threshold and assuming deterioration as a gamma process; and Schutz et al. (2011) investigating the periodic and sequential preventive maintenance policies over a finite planning horizon. However, due to the highly mathematical formalisation of their modelling, it is not straightforward to be applied in practice.

2.3 Maintenance Optimisation

2.3.1 Optimisation Approaches

As the complexity of maintenance systems increases, maintenance optimisation methods have also developed from classical methods to modern or non-traditional methods.

Classical optimisation methods (Rao, 2009) are analytical and use differential calculus to find the optimal value. For example, scatter search (Chen et al. 2006), Nelder-Mead method (Roux et al. 2008, 2013), cyclic coordinate method (Xiang et al. 2012), the modified Powell method (Marquez et al. 2003), Fibonacci algorithms (Asadzadeh and Azadeh 2014) and local search (Gupta and Lawsirirat 2006; Triki et al. 2013), have been applied to simple manufacturing systems. However, it has been criticised that optimising maintenance through classical methods lacks the analysis of objective functions and solution space. Thus it poses difficulty in justification of the optimisation methods.

In order to deal with the complexity increase of maintenance systems, modern or non-traditional methods have been utilised (Deb 2005, Rao 2009). Two modern optimisation methods are mainly applied, which are Genetic Algorithms (GA) and Simulated Annealing (SA). In fact the use of SA is only reported in a few articles. In this sense, exploration of other modern optimisation methods to systems maintenance problems could be another research branch.

The most reported modern optimisation method in maintenance, GA, is based upon the process of natural selection in biology and has been applied to various optimisation problems (Rao 2009). SA originates from the idea of the annealing process in metallurgy to harden metals. In other words, metals are melted in a high temperature at the
beginning and then cooled gradually under a fully controlled environment to obtain desirable shapes or properties. This method has been used to solve a wide variety of problems, such as the ones with continuous, discrete and mixed-integer variables (Rao, 2009).

Dynamic programming (DP), see Bellman (1954), is a method of solving a complex problem by breaking it down into series of simpler subproblems (different parts of the original problem) and then combining the solutions to the subproblems to obtain an overall solution to the original problem. There are two advantages using this formulation. First, dynamic programming enables computing the optimal solution in some cases which usually only applies to smaller problems. Due to the curse of dimensionality, computing the optimal solution to larger problems cannot be done in a reasonable and feasible amount of time (Powell, 2007). Second, dynamic programming can produce optimal theoretical results which could indicate the behaviour of the optimal policy in the proposed models, for example, see Ding et al. (2002).

In a general assumption about state, action and parameter spaces, Rieder (1975) consider a non-stationary Bayesian dynamic decision model which can be reduced to a non-Markovian decision model with known transition probabilities. As a pioneering work in Bayesian dynamic programming, his work provides criteria of optimality and the existence of Bayes policies. Nicolato and Runggaldier (1999) combine burn-in, which is used to cope with the problem of “infant mortality” in system running periods, with identical multi-component systems, and propose a Bayesian dynamic programming method to make decisions on the optimal maintenance interval and best burn-in time.

In other words, dynamic programming is an optimisation approach that transforms a complex problem into a sequence of simpler problems; its essential characteristic is the multistage nature of the optimisation procedure, which provides a general framework employed to solve particular aspects of a more general formulation. In the decision tree problem, we often call it “Roll-Back”, see Ross (1995). As in the problem we will discuss in the next chapter, we divide the system’s running procedure into a few stages and based on the information we learn from running the system, we model the functional form of the system utility and then attempt to maximise it.

The problem addressed in the next chapter is associated with a two-phase sequential
problem. We consider the system running from a global perspective, which means we assume the whole running procedure of the system, and make a decision based on the optimal utility of the system. On the contrary, myopic decision making means that we can do local utility optimisation based on the information we previously obtained. Although we would not obtain the optimal utility for the system, optimisation problems can be simplified and are also easier to implement in practice, and we can simply regard it as a “roll-forward” method.

2.3.2 Optimisation Objectives

Cost minimisation is one of the most reported optimisation objectives in maintenance studies, for example, minimising total cost or expected cost per unit time. Increasing preventive maintenance cost could result in over-maintenance for systems while increasing hazard rate and its consequences would end up with under-maintenance. Note that usually corrective maintenance cost is fixed and higher than preventive maintenance cost (Gupta and Lawsirirat, 2006; Roux et al., 2008; Xiang et al., 2012). However, it is not sufficient to simply only consider maintenance cost as maintenance is part of systems and various costs are associated with systems processing. Thus some researchers have incorporated other costs, in addition to maintenance cost, into maintenance objectives. For instance, a penalty per unit time for systems unavailability (Alrabghi and Tiwari, 2013), or the cost resulting from unsatisfactory products (Oyarbide-Zubillaga et al., 2008).

On the contrast to maintenance cost, Roux et al. (2013) propose maximising systems availability as the optimisation objective. They argue that it is more justified as production costs are more dominant in the total cost of systems running. However, such interpretation would ignore the fact that maintenance costs can be higher than production costs (Wang et al., 2008).

However, it is suggested to design the objective as maximising production throughput because maximum system availability does not necessarily guarantee maximum production throughput (Lei et al., 2010) because a system would not be in a working state due to various reasons such as lack of raw materials or supporting tools.

On the requirements of different engineering problems, optimising several objectives simultaneously for repairable systems has drawn more attention, e.g., minimising
average cost per unit time and maximising systems availability. There are mainly three ways to realise this aim: firstly, put several objectives into one objective function, however, this method requires transformation in a universal unit among different objectives; secondly, assign weights to different objectives based on decision maker’s preference. Although transformation is not required in this method, the decision maker has to trade-off among objectives; thirdly, simply solve several objectives simultaneously using multi-objective optimisation algorithms, e.g., Oyarbide-Zubillaga et al. (2008) implement a Non-dominated Sorting Genetic Algorithm to minimise costs as well as maximise system production profits.

The objective proposed in this thesis is utility-based maintenance. From the maintenance engineer’s point of view, by incorporating risk preferences into maintenance modelling, management of systems could avoid short-term problems such as short-term cash flow problems by considering optimal long-term policy such as systems availability. For example, there are two policies that are equally preferred based on minimising average cost per unit time, however, it is likely that one of them has more variable cash flows than the other, which would result in instability.
Chapter 3

Statistical Methodology and Utility

In operational research and management science we face uncertainty frequently, whether in the form of uncertain demand for production, uncertain performing time in systems, or uncertainty about parameters calibrating a simulation model, etc. Despite the fact that we are facing uncertainty, we usually are able to collect some related information which could reduce the effects of such uncertainty. This would have two impacts: on one hand, gathering information helps us make better decisions which could produce better outcomes; on the other hand, the costs induced by collecting information, such as time and money spent, or even opportunity cost due to collecting one piece of information whereas we could have had another, could increase. How to find the optimal balance between these benefits and costs is what we are attempting to explore. Two methods for formulating such problems are Bayesian methods and Dynamic Programming, respectively.

3.1 Bayesian Modelling

In the twentieth century, the so called ‘frequentism’ has dominated the statistical philosophy. Under the framework of the frequentist approach, the hypothetical long term proportion of the time that an event of interest occurs is expressed via probability and the parameters of probability models are regarded to be unknown but fixed numerical quantities. Pearson studied the problems of the frequentist style extensively in the early twentieth century. One can find most of the foundations for frequentist modelling and inference in the classical work by Fisher (1922).
Bayesian modelling arises earlier than the frequentist methodology actually. It is being dated back to a thought experiment conducted by Bayes (1763) who threw balls onto a square table, and to the ‘inverse probability’ found by Laplace (1774) and later replaced as ‘Bayesian’ in the 1950s by Fisher (Fienberg, 2006). Both Bayes and Fisher took the uniform prior distribution: it was a outcome of the ball throwing experiment in Bayes’ case, while it was considered as an intuitive axiom which is a ‘principle of insufficient reason’ by Fisher. Yet, Bayesian modelling and inference has not drawn much attention until it reappeared and was recognised in the modern form thanks to Jeffreys (1939) and Savage (1954), among other researchers.

Briefly the Bayesian concept combines both objective probability that has a similar interpretation compared to the probabilities in frequentism and subjective probability that is taken to express a degree of belief from a personalistic point of view. The treatment of regarding the parameters of probability models as realisations of a random variable or not differentiate Bayesian modelling from frequentist methodology, and enable one to put direct statements of probability for these parameters.

Let us consider a sequence of observations \( x = \{x_1, \ldots, x_n\} \) and a corresponding probability model that is believed to be the generating model for the data, with a probability density function \( f_X(x \mid \psi) \) in which the parameter(s) \( \psi \) is made explicitly to model the dependence. It is worthwhile noting that \( \psi \) may be a vector of parameter in general. \( \psi \) is considered as the unknown realisation of a random variable in the Bayesian modelling, as a result we can consider the joint density function of the random variables \( X \) and \( \Psi \):

\[
f_{X,\Psi}(x, \psi) = f_{X \mid \Psi}(x \mid \psi) f_{\Psi}(\psi)
\]

(3.1)

where \( f_{X \mid \Psi}(x \mid \psi) := f_X(x \mid \psi) \) for simplicity.

As a result, given data \( x \), one can apply Bayes’ Theorem to directly derive the uncertainty expression for \( \psi \) as:

\[
f_{\Psi \mid X}(\psi \mid x) = \frac{f_X(x; \psi) f_{\Psi}(\psi)}{\int_{\Omega} f_X(x; \psi) f_{\Psi} d\psi} \propto f_X(x; \psi) f_{\Psi}(\psi)
\]

(3.2)

where \( \Omega \) is the sample space of \( \Psi \). Notice that the denominator is independent of \( \psi \) thanks to the integral, so the proportionality \( \propto \) follows. In the Bayesian modelling,
is referred to as the posterior distribution of parameter(s) $\psi$ and encapsulates all knowledge and information concerning the unknown parameters.

However, in practice the integral in the denominator of (3.2) which is the normalising constant, does not often have an analytically tractable form. Due to this fact, it hindered the learning and application of Bayesian modelling for quite a long period of time. With the rapid development of computing techniques such as Markov chain Monte Carlo in modern time, Bayesian modelling has become easier to implement since the 1990s.

### 3.1.1 Likelihood function

In (3.2), the first term in the numerator is usually written as $L(\psi; x) := f_X(x \mid \psi)$ to indicate the fact that the observed data $x$ are fixed and $\psi$ is in fact stochastic. Fisher (1922) terms this the likelihood, arguing that it was all necessary to do inference through simply maximising $L(\psi; x)$ with regard to $\psi$ and one can obtain the maximum likelihood estimate (MLE), $\hat{\psi}$. However, due to the fact that $\int_{\Omega} L(\psi; x) d\psi \neq 1$, we can not take the likelihood as a probability distribution function for $\psi$, which makes expressing the uncertainty about the point estimate $\hat{\psi}$ somewhat more ambiguous than the expression in the form of posterior distribution in Bayesian modelling.

In favour of the ‘fiducial probability’ of $\psi$, Fisher (1930) defined $\frac{\partial}{\partial \psi} F_X(x; \psi)$ as a way to connect $\psi$ to a direct probability uncertainty. However, after extensive criticism, e.g. Lindley (1958), almost nobody refers to this idea in the modern literature. Therefore, based on asymptotic analysis, the expression of uncertainty for $\hat{\psi}$ is limited to be required to state the upper/lower confidence bound for non-standard models, which is argued to contain the ‘true’ $\psi$ in terms of a specified hypothetical long run proportion.

Still only in the way of the likelihood can the data enter the posterior distribution to be utilised. And as the amount of data collected increases, the likelihood part dominates the posterior distribution more, resulting in approximate agreement between maximum likelihood estimate for frequentists and highest a posteriori density values for Bayesians, which is the mode of the posterior distribution. In Bayesian modelling, any two models with the same likelihood $L(\psi; x)$ are believed to result in the same inference because the likelihood expresses all the information from the data, which is
called the ‘likelihood principle’.

3.1.2 Prior distribution

The prior distribution is the term $f_\Psi(\psi)$ in numerator from (3.2). Because the parameters are treated as random variables in Bayesian modelling, a Bayesian is required to specify corresponding distributions for the random variables which are independent of the data $x$ in (3.1). The prior distribution is taken to represent the information that a decision maker has known prior to observing any possible data and is usually regarded as the most controversial part in Bayesian methodology.

The controversy arises owing to the fact that the choice of a prior distribution is always a subjective decision. However, it is argued that the implementation of a prior distribution adds flexibility of modelling. For example, we can choose a ‘sceptical prior’ which is suggested to assign a lower probability to some favourable outcomes. Thus, the probability weighs in favour of that outcome becomes even more compelling. However, we can think that the chosen model that is believed to be where the data are generated, $f_X(\cdot; \psi)$ is also a somewhat subjective choice.

**Common prior choices**

In Bayesian modelling and inference, the prior is often chosen from some parametric family of distributions and the parameters of the prior distribution are referred to as hyper-parameters. There are commonly a few ways to choose a prior distribution for the model specification. In general, these method include the following: subjective priors, objective priors, empirical priors, priors from experts and conjugate priors.

In subjective probability, a particular individual chooses or specifies a prior to capture her or his belief under examination as good as possible. It is worthwhile mentioning that even a very vague prior can be useful, because the results during the inference are mathematically and rationally updated for the belief concerning $\psi$ with observations coming in increasingly.

On the contrary, objective priors are chosen as a convenience to capture ‘ignorance’ about $\psi$ a priori. These priors with hyper-parameters are often set to have very high variance. and have good frequentist properties. Alternatively, as it can be seen in (3.2), the prior exists in both numerator and denominator, then any multiplication of
the priors do not make any difference, as a result, some practitioners choose some priors which do not integrate to 1 in a controversial fashion. It has to be noticed that one can no longer guarantee the posterior to be a legitimate probability distribution because of that. For example, if one choose a flat prior $f_{\psi}(\psi) = c \quad \forall \psi \ (c > 0)$, then it will lead to solutions corresponding to Fisher’s fiducial probabilities, which is improper in our Bayesian setting. Problems of improper priors are that there is the danger of over-interpreting them since they are not probability densities, and also do not necessarily ensure a proper posterior. To address the issue that Bayesian inference is sensitive to the parameterisation for modelling, Jeffreys (1946) developed Jeffreys prior, which is specified as $f_{P_{\psi}}(\psi) \propto J(\psi)^{1/2}$, where $J(\psi)$ is the Fisher information with respect to $\psi$. Jeffreys prior enables the posterior to be invariant to re-parameterisation for the model, yet this can still have problems of improper priors, which are that there is the danger of over-interpreting them since they are not probability densities and also do not necessarily ensure a proper posterior.

Empirical priors lie in the field of so called ‘Empirical Bayes’. Its basic idea is to learn some of the parameters of the prior from the data. Let us consider a hierarchical Bayesian model (to be introduced as follows) with parameter $\Psi$ and hyper-parameter $\lambda$:

$$f_X(x | \lambda) = \int f_X(x | \psi)f_{\psi}(\psi | \lambda) \, d\psi,$$

and estimate the hyper-parameter via maximum likelihood estimate as:

$$\hat{\lambda} = \arg \max_{\lambda} f_X(x | \lambda).$$

This method has its advantage of being robust because it overcomes some limitations of mis-specification of the prior. However, it double counts the data, which results in a likelihood principle violation of the relationship between data and hypothesis.

Specifying priors from experts is actually a research domain itself that is called ‘prior elicitation’. For simplicity, this might be conducted by pooling the experts’ opinion about the parameters of the model onto a credible range and applying a normal prior with fixed $\gamma\%$ points for upper and lower bounds. This method requires the parameters of the model to be well interpreted to the experts, otherwise, it would add more questions if the experts have no understanding of probability theory. Garthwaite et al. (2005) and O’Hagan et al. (2006) are relatively recent reviews of techniques to address this issue.
If a prior distribution is multiplied by the likelihood, resulting in an expression that is algebraically from the same family as the prior distribution, up to a normalising constant, we call this conjugacy and the prior is called a conjugate prior distribution. Its advantage arises since the normalising constant can be written down via inspection without conducting integral in the denominator in (3.2). Particularly, if the likelihood is from the exponential family and written as:

\[ f_X(x \mid \psi) := f(x)g(\psi) \exp\{\Phi(\psi)^T s(x)\} \]

where \( \Phi \) is a vector of natural parameter, \( s(x) \) is a sufficient statistic, and \( f, g \) are positive functions of \( x \) and \( \psi \), respectively. If the conjugate prior is taken from exponential family as:

\[ f_\psi(\psi) = h(\eta, \nu)g(\psi)^\eta \exp\{\Phi(\psi)^T \nu\} \]

where \( \eta \) and \( \nu \) are hyper-parameters and \( h \) is the normalising function, then the posterior distribution for \( n \) independent exponentially distributed data points is also conjugate, with hyper-parameters \( \eta + n \) and \( \nu + \sum_i s(x_i) \) and has the computationally convenient form as follows.

\[ f_\psi(\psi \mid x_1, \ldots, x_n) = \left( \eta + n, \nu + \sum_i s(x_i) \right) g(\psi)^{\eta+n} \exp\left\{ \Phi(\psi)^T \left( \nu + \sum_i s(x_i) \right) \right\} \]

### 3.1.3 Posterior analysis

In the Bayesian modelling and inference setting, one can fully conduct an analysis based on the posterior distribution to deal with all the questions of interest. Usually, this can include probability density function plots, summary statistics such as expectation and variance, the modal value of the posterior, or the intervals of the highest posterior density. When the interest lies in some functional form of the parameters of models, it often can also be dealt with comparatively straightforwardly.

It is worthwhile noting that models can become increasingly complex in a frequentist framework, such as when there exist nuisance parameters in some elements of \( \psi \). This issue can be beautifully dealt with via standard probability theory. For instance, if \( \psi = (\delta, \lambda) \) in which \( \delta \) is the parameter of interest whilst \( \lambda \) is a nuisance parameter required to construct the complete model. In a Bayesian framework, we can obtain the
posterior distribution of $\delta$ by simply integrating out the nuisance parameter:

$$f_{\Delta}(\delta \mid x) = \int_{\Omega} f_{\Psi \mid X}(\delta, \lambda \mid x) d\lambda$$

Once the posterior distribution for a parameter of a model is obtained, it can be used further to explore other probabilities. For example, a posterior predictive distribution can be derived to provide the predictive probability distribution by taking into account the uncertainty in $\psi$ for a future data point, $x^*$, which is from the same data generating process incorporating all the information:

$$f_{X^* \mid X}(x^* \mid x) = \int_{\Omega} f_X(x^* \mid \psi) f_{\Psi \mid X}(\psi \mid x) d\psi$$  \hspace{1cm} (3.5)

In a frequentist framework, it is usual to replace the parameter $\psi$ with a point estimate $\hat{\psi}$ in order to derive the predictive distribution:

$$f_{X^* \mid X}(x^* \mid x) := f_X(x^* \mid \hat{\psi}).$$  \hspace{1cm} (3.6)

However, (3.6) fails to interpret the uncertainty in $\hat{\psi}$.

### 3.1.4 Hierarchical Bayesian models

When it comes to modeling more complex real problems in practice, the simple specification of the likelihood and prior functions indicates its limitation. Hierarchical modelling arises to address this issue in a natural expression under a Bayesian framework.

In the previous illustration, we have stated that the observations are believed to be from the data generating process $f_X(\cdot \mid \psi)$ with parameter $\psi$ where we can treat $\psi$ as a realisation of a random variable $\Psi$. And the core of Bayesian modelling, the posterior presents the probability distribution for a particular realisation $\psi$ expressing the characteristic of the process. Let us consider multiple realisations of the parameters, denoted as $\psi_i$, which will result in the data in different groups, e.g., $x_{i1}, \cdots, x_{im_i}$ are the data in group $i$ from the parameter $\psi_i$. We can place a hyper-prior for the prior distribution $f_{\Psi}(\cdot)$ under the Bayesian idea. Therefore, we take the estimation of $\psi_i$ as part of our Bayesian modelling process via hierarchically conditional probabilities as follows in an example.
Assume we have the data $x = \{x_{11}, \ldots, x_{mn}\}$ where the first index indicates which group the data are from and the second one tells us the number of observations in that group. We define the following to specify the hierarchical model:

$$f_{X|\Psi}(\cdot \mid \psi_i) \quad (3.7)$$
$$f_{\Psi|\Lambda}(\cdot \mid \lambda) \quad (3.8)$$
$$f_{\Lambda}(\cdot) \quad (3.9)$$

where (3.7) is the probability model of the data generating process for group $i$. Note that for different hierarchies, the parametric probability functions are from the same family, and only differ in the terms of different parameters for each group (as mentioned previously, each $\psi_i$ can be in a vector form). (3.8) is assigned as the prior distribution for the parameters $\Psi$, within which the hyper-parameter $\lambda$ is made explicitly. Note that the hyper-parameter $\lambda$ is not specified either, but given a hyper-prior probability distribution (3.9). This complete model includes parameters $\xi = (\psi_1, \ldots, \psi_n, \lambda)$, so that the posterior is expressed as follows:

$$f_{\Xi|Y}(\xi \mid x) \propto f_{Y|\Xi}(x \mid \xi) \cdot f_{\xi}(\xi)$$

$$= \underbrace{f_{Y}(x \mid \psi)}_{\text{first term}} \cdot \underbrace{f_{\Psi}(\psi \mid \lambda) \cdot f_{\Lambda}(\lambda)}_{\text{second term}}$$

The simplification of the first term to the usual likelihood is thanks to the conditional independence of $x$ and $\lambda$ given $\psi$, and the joint prior in the second term decomposes naturally because of the model formulation.

Then the interest will usually be in the posterior predictive probability distribution of $\psi$, since this incorporates all the information learned about that parameter from observing the data points:

$$f_{\psi^*|X}(\psi^* \mid x) = \int f_{\Psi}(\psi^* \mid \lambda) \cdot f_{\Lambda|X}(\lambda \mid x) \, d\lambda \quad (3.10)$$

The above is simply one of the possible hierarchical models, but it can be much more complicated when dealing with other sophisticated models in practice, then for modelling the dependencies one can simply adopt directed acyclic graphs (DAG) which makes identification of conditional dependencies and independencies more clear.
3.1.5 Bayesian method in maintenance

Under the non-Bayesian framework, when there is little learning from the system operation or rare evidence to judge the characteristics of a system or its components, it would be questionable to carry out maintenance policies, such as corrective maintenance (CM), and preventive maintenance, because it is difficult to judge if the system or its components are critical or not on a priori grounds. At the same time, applying maintenance policies may cause a significant drift of the system reliability, and it can fail to capture the dynamics of systems when a new maintenance policy is carried out without considering Bayesian learning.

Due to the uncertainty characteristics of most systems, such as unknown lifetime distribution or known distribution but with uncertain parameters, it is necessary to model these uncertainties to conduct reasonable maintenance policies. There have been increasing applications of Bayesian methods in maintenance modelling, which may be categorised as follow:

- Bayesian inference

Researchers consider Bayesian theoretical approaches to determine the preventive maintenance policies through inference on the uncertain parameters and establishing formal expression and updating the uncertain parameters. For example, Percy and Kobbacy (1996) consider the analysis of a delayed renewal process and a delayed alternating renewal process incorporating exponential failure time into their models. By adopting Bayesian methods, they propose the predictive distribution for down time, the number of corrective maintenances, and cost per unit time, instead of calculating the estimated renewal function (expected value). However, there are some restrictions in their models, for example, only the exponential failure time distribution was considered and the downtimes were assumed to be constant. Our models in the next chapter will be dealing with globally optimal maintenance times rather than the preventive maintenance interval considered by Percy and Kobbacy (1996). By incorporating minimal repair, maintenance and replacement, Sheu et al. (2001) formulated the expected cost per unit time by analysing the uncertain parameters of a Weibull distribution, which was named “adaptive preventive maintenance”. Kim et al. (2007) adopt
the Bayesian method to the unknown parameters for a Weibull failure time distribution of a sequential preventive maintenance model, which is defined in the context of a cycle. Within a cycle, minimal repair is conducted after a failure and the effective performing time of the system and its hazard rate are both adjusted to modelling the deterioration of the system. At the end of a cycle, a full replacement of the system is carried out. However, their methods did not consider the statistical learning connection between cycles and thus ignored the future possibility of system performance.

• Bayesian computation and decision making

Bayesian inference and computation for stochastic processes in maintenance problems have been rising in recent years. The underlying methodologies in this field are conducting Bayesian inference and computation for systems modelled by varying stochastic process, such as geometric process, gamma process, etc. Chen et al. (2010) carried out for the geometric process with the exponential distribution and lognormal distribution and applied a Gibbs sampler and Metropolis algorithm to calculate the Bayesian estimators of the parameters in the geometric process. Optimal maintenance decisions under imperfect inspection were proposed by Kallen and van Noortwijik (2005), in which a gamma process is used to model the corrosion damage mechanism and Bayes’ theorem is applied to update prior knowledge concerning the corrosion rate. However, the decision model was based on periodic inspection and replacement policy, which is not regarded as a globally optimal maintenance.

• Bayesian network

Bayesian network has been applied in studies concerning maintenance modelling. In order to estimate the future state of a system after a maintenance action based on conditional probabilities, Kang and Golay (1999) proposed a model with influence diagrams. Moreover, Celeux et al. (2006) developed a questioning procedure to elicit expert opinions by collecting information and building up the network structure. By considering the factors that could induce uncertainty during maintenance actions, De Melo and Sanchez (2008) applied Bayesian networks to the prediction of delays for software maintenance projects.
• Applications

Bayesian methods have also been applied in other domains. For example, Durango-Cohen and Madanat (2008) used a quasi-Bayes approach to optimise the inspection and make maintenance decisions for infrastructure facilities under performance model uncertainty by taking a mixture of known models, of which the mixture proportions are assumed to be random variables with probability densities updated over time. When there are limited data and information, Zhang and Wang (2014) used Bayesian linear methods to combine the subject expert knowledge with the available limited data to estimate the unknown parameters of models and applied it in infrastructure assets. Their optimisation objective is still to minimise the cost per unit time rather than the utility that is the optimisation objective of this thesis.

In the Bayesian modelling as follows, prior information or expert judgement will be combined with the available updated system operating characteristics (Bayesian approach) and the risk aversion (utility) and economic consequences (relevant costs) taken into account (statistical decision theory) to optimise the maintenance strategies.

3.2 Dynamic Programming

Dynamic programming is an optimisation method that solves a complex problem by transforming it into a sequence of simpler subproblems. In general, dynamic programming provides a framework for analysing and solving many types of problems, although it requires one to identify whether the particular problem can be analysed via dynamic programming.

3.2.1 An Elementary Example

We first introduce the philosophy and concept of dynamic programming by an elementary example used by Chinneck (2015).

Assume that the number of students that fail a course depends on the number of demonstrators allocated to each session of the course, see Table 3.1. If there are 6 demonstrators available and there are 4 sessions of this course. The problem is:
How many demonstrators should be allocated to each session of this course to have the fewest student fail?

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Table 3.1: Number of failed students for a course with 4 sessions and 6 available demonstrators.

First we formulate the problem in terms of characteristics in dynamic programming:

- **stages**: 1st solve section 4, 2nd solve section 3 and 4, 3rd solve section 2, 3 and 4, 4th solve section 1, 2, 3 and 4.
- **state at a stage**: number of demonstrators available to be allocated.
- **decision**: how many demonstrators to be allocated to section $i$, $i = 1, 2, 3, 4$.
- **decision update to state**: number of available demonstrators to allocate is reduced corresponding to the decision.
- **recursive value relationship**.

Related notations for this problem are presented below.

$x_i$ number of demonstrators allocated to section $i$

$F_i(x_i)$ number of students who fail the course in section $i$ given $x_i$ demonstrators

$d_i$ number of demonstrators available at the beginning of stage $i$

$f_i(d_i)$ best possible solution from stage $i$ to the end, i.e., $\min_{x_i} \{F_i(x_i) + f_{i+1}(d_i - x_i)\}$

Since there is a total of 6 demonstrators we have a possibility of having 0 to 6 demonstrators available to be allocated. We may assign 0 to 6 demonstrators to section 4; of course, we cannot allocate more demonstrators to section 4 than there are available. Let’s fill the rest of the Table 3.2 with the values from Table 3.1. 15 students
will fail if no demonstrator is allocated to section 4; 8 students will fail if 1 demonstrator is allocated to section 4; 6 students will fail if 2 demonstrators are allocated to section 4; 3 students will fail if 3 demonstrators are allocated to section 4; 2 students will fail if 4 demonstrators are allocated to section 4; 1 student will fail if 5 demonstrators are allocated to section 4; no student will fail if 6 demonstrators are allocated to section 4.

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Table 3.2: Dynamic Programming Example: Stage 4.

The recursive relationship for this stage is $f_4(d_4) = \min_{x_4} \{F_4(x_4)\}$. Therefore, the $f_4(d_4)$ is the smallest value in each row.

We will now move to Stage 3: section 3 and 4. Again, we can have 0 to 6 demonstrators available to allocate and we allocate 0 to 6 demonstrators to this section. The recursive relationship at this stage is $f_3(d_3) = \min_{x_3} \{F_3(x_3) + f_4(d_3 - x_3)\}$. Let’s look at the case where we have 2 available demonstrators to allocate ($d_3 = 2$) and we choose to allocate 1 demonstrator to section 3 ($x_3 = 1$). This gives us a value of $f_3(2) = F_3(1) + f_4(2 - 1) = F_3(1) + f_4(1)$. From Table 3.1 we get $F_3(1) = 16$ since 16 students will fail if 1 demonstrator is allocated to section 3, so $f_3(2) = 16 + f_4(1)$. From Table 3.2 $f_4(1) = 8$, i.e., when $d_4 = 1$, the fewest number of students that will fail is 8, so $f_3(2) = 16 + 8 = 24$, and we can write this into Table 3.3 where $d_3 = 2$ and $x_3 = 1$. Let’s fill in the rest of Table 3.3. Now let’s find $f_3(d_3)$ for each state by selecting the minimum value for each row.
We will now move on to stage 2, section 2, 3 and 4 of this course. Since stage 3 includes section 3 and 4, we will not need Table 3.2 until we retrieve the solution. Let’s look at the stage 2 shown in Table 3.4. The recursive relationship at this stage is $f_2(d_2) = \min_{x_2} \{F_2(x_2) + f_3(d_2 - x_2)\}$. Let’s look at the case where we have 4 demonstrators available to allocate ($d_2 = 4$) and we choose to allocate only 1 to section 2 ($x_2 = 1$). The recursive relationship becomes $f_2(4) = F_2(1) + f_3(4 - 1) = F_2(1) + f_3(3)$. We get $F_2(1)$ from Table 3.1, 20 students will fail if 1 demonstrator is allocated to section 2. We get $f_3(3)$ from the Table 3.3, i.e., if we have 3 demonstrators to allocate to section 3 and 4, what is the smallest number of students that will fail? So $f_2(4) = 20 + 21 = 41$. We enter this value into Table 3.4 where $d_2 = 4$ and $x_2 = 1$. Now we will add the rest of the values to Table 3.4.

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**Table 3.3:** Dynamic Programming Example: Stage 3.

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<td>-</td>
<td>-</td>
<td>54</td>
</tr>
<tr>
<td>2</td>
<td>49</td>
<td>49</td>
<td>51</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>49</td>
</tr>
<tr>
<td>3</td>
<td>46</td>
<td>44</td>
<td>44</td>
<td>47</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>44</td>
</tr>
<tr>
<td>4</td>
<td>40</td>
<td>41</td>
<td>39</td>
<td>40</td>
<td>43</td>
<td>-</td>
<td>-</td>
<td>39</td>
</tr>
<tr>
<td>5</td>
<td>37</td>
<td>35</td>
<td>36</td>
<td>35</td>
<td>36</td>
<td>40</td>
<td>-</td>
<td>35</td>
</tr>
<tr>
<td>6</td>
<td>35</td>
<td>32</td>
<td>30</td>
<td>32</td>
<td>31</td>
<td>33</td>
<td>38</td>
<td>30</td>
</tr>
</tbody>
</table>

**Table 3.4:** Dynamic Programming Example: Stage 2.

Now we will look at stage 1, section 1, 2, 3 and 4. We will not need the Table 3.3.
again, until we retrieve the solution. In stage 1 (Table 3.5), the number of demonstrators available is 6, no demonstrators have been allocated before this stage, so we know that there are exactly 6 demonstrators available. The recursive relationship for this stage is 

\[ f_1(d_1) = \min_{x_1} \{ F_1(x_1) + f_2(d_1 - x_1) \}. \]

Let’s look at the case where we choose to allocate 3 demonstrators to section 1, then \( f_1(6) = F_1(3) + f_2(6 - 3) = F_1(3) + f_2(3). \)

From Table 3.1, \( F_1(3) = 5 \), i.e., 5 students will fail if we allocated 3 demonstrators to section 1. From Table 3.4, we get \( f_2(3) = 44 \), which means if we have 3 demonstrators to allocate to section 2, 3 and 4, the smallest number of students that will fail is 44, so \( f_1(6) = 5 + 44 = 49 \) and we put it in Table 3.5. Now let’s fill in the rest of the table and \( f_1(d_1) \) is the smallest value in this row.

<table>
<thead>
<tr>
<th>( d_1 )</th>
<th>( x_1 = 0 )</th>
<th>( x_1 = 1 )</th>
<th>( x_1 = 2 )</th>
<th>( x_1 = 3 )</th>
<th>( x_1 = 4 )</th>
<th>( x_1 = 5 )</th>
<th>( x_1 = 6 )</th>
<th>( f_1(d_1) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>47</td>
<td>46</td>
<td>48</td>
<td>49</td>
<td>51</td>
<td>55</td>
<td>61</td>
<td>46</td>
</tr>
</tbody>
</table>

**Table 3.5:** Dynamic Programming Example: Stage 1.

Now let’s retrieve the solution. Looking at stage 1 (Table 3.5), we know the least number of students that will fail, i.e., allocating 6 demonstrators to this course, there will be 46 students who will fail.

We will trace back through the solution to obtain the allocation of demonstrators. We start by looking at stage 1 (Table 3.5). How many demonstrators should we allocate to section 1? The smallest number of students that will fail is 46, which occurs when \( x_1 = 1 \), i.e., we are supposed to allocate 1 demonstrator to section 1. Now we will look at stage 2 (Table 3.4), \( d_2 \) at stage 2 is 5 since we start with 6 demonstrators and have allocated 1 to section 1. The smallest number of students that will fail if we have 5 demonstrators to allocate to section 2, 3 and 4 is 35, which occurs when either 1 or 3 demonstrators are allocated to section 2, in other words, there are two possible allocations that will give us the best solution. Let’s look at stage 3 (Table 3.3): given the first partial solution, 2 demonstrators have been allocated so we have 4 demonstrators left for section 3 and 4, so the smallest number of students that will fail is 15 when \( x_3 = 3 \); given the second partial solution, 4 demonstrators have been allocated so 2 are available, thus the smallest number of students that will fail is 24, which occurs when \( x_3 = 1 \). Now let’s go forward to stage 4 (Table 3.2). Both partial solutions have allocated 5 demonstrators to section 1, 2 and 3, leaving only 1 demonstrator for section 3.
4, hence the smallest number of students that will fail is 8, which occurs when $x_4 = 1$. As a result, there are two schemes of allocation of demonstrators, $A$ and $B$ that see the least number of failed students, at 46,

<table>
<thead>
<tr>
<th>Sessions</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allocation $A$</td>
<td>1</td>
<td>1</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Allocation $B$</td>
<td>1</td>
<td>3</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

Now we have solved this problem successfully by dynamic programming and the characteristics of dynamic programming will be illustrated via this example.

### 3.2.2 Characteristics

There are three most important characteristics in dynamic programming which are *stages*, *states* and *recursion*.

**Stages**

The essential part of dynamic programming method is to recognise and restructure the optimisation problems into a multiple of *stages* and only solve one stage subproblem at a time sequentially. The solution of each one-stage subproblem assist to define the characteristics of the next one-stage problem in the sequence, though each one-stage subproblem is solved via a normal optimisation problem.

The stages usually present the different time periods in a problem’s analysing procedure, which means the stage duration is constrained by the length of the problem to be analysed. The problem of determining the optimum preventive maintenance time in this thesis is to be stated as a dynamic programming problem. The decision variable is the scheduled preventive maintenance time at the beginning of each phase. As the system considered will be performing for a few planned phases; the objective is to maximise the total expected utility of system performance; however, there are no resources (cost budgets) constraints in our assumption. If we can only determine the optimum preventive maintenance times for each phase of the system at the beginning of system performing, we could restructure the problem into a few stages based on the number of planned phases, of which each represents the decision regarding the optimum maintenance time at the beginning of each phase.
However, the stages of a dynamic programming do not have time implications necessarily. In the example illustrated in §3.2.1 the problem of allocating 6 available demonstrators to 4 sessions of the course is restructured into 4 stages, which are 1st solve section 4, 2nd solve section 3 and 4, 3rd solve section 2, 3 and 4, 4th solve section 1, 2, 3 and 4. The decision variable $x_i$ is the number of demonstrators to be allocated to each session of the course to make the least failed number of student in this course. It is worthwhile noting that problems without time implications are comparatively difficult in practice to be restructured in stages via dynamic programming.

**States**

The *states* correspond to each stage of the optimisation problem in dynamic programming. The states indicate the information needed to fully analyse the outcomes that the current decision has upon the future situations. In the demonstrator allocation problem in §3.2.1 each stage has only one variable representing the state: $d_i$ the number of demonstrators available to be allocated at each stage. In our problem of determining the optimum preventive maintenance time, the situation that failure time $T_f$ which can be observed or not before scheduled maintenance time $T_m$ for each phase $i$ that is the state variable.

The elicitation of the state variable in a dynamic programming problem is very critical. However, there do not exist standard rules to specify it in particular and it usually requires one to study the problem through dynamic programming in a somewhat creative and subtle way. Based on practical implementation, it is suggested to select the states of a dynamic programming problem based on the following criteria.

- The states should reflect sufficient information for one to make decisions in the future regardless of how the problem has arrived at the current state.

  In the demonstrator allocation problem in §3.2.1 the state variable, the number of available demonstrators $x_i$ for each session $i$, does meet this criterion because it does not consider how one has allocated the demonstrators prior to the current section $i$.

- As the computation expenditure related to dynamic programming is higher with the number of state variables in a dynamic programming problem, it is suggested
to specify the number of states as small as possible. As a result, the limited property of dynamic programming restricts the application of dynamic programming.

**Recursive Optimisation**

To solve a dynamic programming problem with \( N \) stages (or \( N \) subproblems) through sequentially solving a one stage problem each time until the overall or global optimisation is done, it requires the recursion process which is the last important characteristic of the dynamic programming method. This process is usually based on a *backward induction* technique, in which one is supposed to analyse the first stage of a dynamic programming problem and then move back one stage each time until all the stages have been analysed. In the demonstrator allocation problem in §3.2.1, this process is carried out via the recursion relationship \( f_i(d_i), \text{ i.e., } \min_{x_1} \{F_i(x_i) + f_{i+1}(d_i - x_1)\} \), from the stage 4 to stage 1. Alternatively, the recursion process can also be based on *forward induction* technique, in other words, one solves the initial stage and moves forward to the next one each time until the overall problem is solved. However, in some dynamic programming problems, only one of these two induction techniques can be used, *e.g.*, when dealing with problems involving uncertainty, only backward induction is applied.

Dynamic programming must observe the *principle of optimality*, that no matter what the current state and decision are, the following decision must be optimal with regard to the state following the currently made decision.

### 3.2.3 Formalisation under Uncertainty

We formalise dynamic programming under uncertainty as follows. Given a state and corresponding stage, under uncertainty in dynamic programming, the current decision does not completely determine the next state of the problem for the next stage, which is different from a dynamic programming problem under certainty in which the next state of the problem entirely depends on the current state and decision. Given the current state of the process \( s_n \) with \( n \) following stages and the current decision \( d_n \), with an uncertain event expressed by a random variable \( Z_n \) of which \( z_n \) is a realisation which is not determined by the decision maker. The return function for the current stage can depend on this random variable \( Z_n \), *i.e.,*

\[
 f_n(d_n, s_n, Z_n), \tag{3.11}
\]
where \( f_n(\cdot) \) is the return of the process when there are \( n \) extra stages to go.

The next state of the process \( s_{n-1} \) has \((n-1)\) stages to go and we define the transition function \( t_n(\cdot) \) as

\[
s_{n-1} = t_n(d_n, s_n, Z_n).
\] (3.12)

The consequence of the random variable \( Z_n \) is controlled by a probability distribution \( P_n(z_n | d_n, s_n) \), which can be conditional on the state, the current stage, and the current decision. One can only select the optimal decision after knowing the consequence of the uncertain event prior to the current decision making.

In dynamic programming under uncertainty, because the return and following stage is potentially unknown at each stage, we have to optimise the expected return over all the stages of the problem by considering a sequence in which possible decisions can be made and the consequences of uncertainties become available. Backward induction is applied to find the optimal policy in this situation, whereas forward induction cannot since it is impossible to give specified values for the states at the following stages which are dependent on the uncertainty from the current stage. However, via backward induction, there is no such issue because the states without following stages to go are analysed first, and then the states having one stage to go are evaluated through the optimum expected value of decision.

Backward induction can be illustrated as follows. First compute the optimum value of return at stage 0, which means this is the stage with no following stages to go. Second compute the expected value of return of stage 1 for each uncertain event via their corresponding and following state by their probabilities. We continue this process in a similar way until arriving at the final stage.

The optimum expected value for dynamic programming under uncertainty can be written in recursion form as follows:

\[
r_n(s_n) = \max \mathbb{E} \left\{ f_n(d_n, s_n, Z_n) + r_{n-1}(s_{n-1}) \right\},
\] (3.13)

where \( s_{n-1} = t_n(d_n, s_n, Z_n) \).

### 3.3 Utility Theory

In this section, we introduce basic properties of utility functions and the notation of expected utility in a short summary that is sufficient for our purposes in this thesis. We

Utility theory is concerned with a decision maker’s preferences or values which can be represented in numerically useful ways under assumptions about a decision maker’s preferences [Fishburn 1968]. A utility theory is usually based on a decision maker’s preference-indifference relation $\preceq$ (read “is not preferred to”), and a set $X$ of elements $x, y, z, \cdots$ (interpreted as decision alternatives). If $x, y$ and $z$ are in $X$, then they are assumed to have the following properties:

1. Complete. For all $x$ and $y$ in $X$, either $x \preceq y$ ($x$ is not preferred to $y$) or $y \preceq x$ or both.

2. Reflexive. For all $x$ in $X$, $x \preceq x$, i.e., $x$ should be preferred at least as much as itself.

3. Transitive. For all $x, y$ and $z$ in $X$, if $z \preceq y$ and $y \preceq x$, then $z \preceq x$.

Note that $x \prec y$ ($y$ is preferred to $x$) is strict preference and $x \sim y$ ($x$ is indifferent to $y$) is indifference, which are other relations of $x$ and $y$.

### 3.3.1 Utility Functions and Probabilities

A utility function $U(x)$ is a twice differentiable function of resource $x$, where $x > 0$. If its first derivative $U'(x) > 0$ (i.e., an increasing function) and second derivative $U''(x) < 0$ (i.e., a concave function), we say it is a legitimate utility function.

Under different circumstances, if the decision maker has reasonable preferences about the decision, then a utility function can be used to represent these preferences. i.e., a function $U : X \rightarrow R$ such that $x \preceq y$ if and only if $U(x) \leq U(y)$. However, the decision problem has a special structure when uncertainty is incorporated into decision making. In general, how a decision maker values decisions in one state as compared to another will depend on the probability that the state in question will actually occur.

For this reason, we will write the utility function as depending on the probabilities as well as on monetary return sets. Suppose that we are considering two mutually exclusive states such as buying item 1 or item 2. Let $c_1$ and $c_2$ represent the costs induced by decision 1 (buying item 1) and decision 2 (buying item 2), then let $\pi_1$ and
\( \pi_2 \) be the probabilities that induced by decision 1 and decision 2 actually is made and their corresponding costs occur, the utility function is written as

\[
U(c_1, c_2, \pi_1, \pi_2) = \pi_1 c_1 + \pi_2 c_2
\]

If the two decisions are mutually exclusive, so that only one of them can happen, then \( \pi_2 = 1 - \pi_1 \). But we will still write out these probabilities in order to keep symmetry.

Given this notation, we can write the utility function for decision as \( U(c_1, c_2, \pi_1, \pi_2) \). This is the function that represents the decision maker’s preference over each decision.

There are several classes of utility functions suitable for describing various types of decision makers’ economic behaviour. We examine some examples of well known classes: the quadratic, logarithmic, iso-elastic and negative exponential utility functions.

**Quadratic Utility Function**

**Definition 3.1** A quadratic utility function is of the form

\[
U(x) = ax - bx^2.
\]

Since its first derivative \( U'(x) = a - 2b > 0 \), when \( x < a/2b \) and second derivative \( U''(x) = -2b < 0 \), this is a legitimate utility function.

A quadratic utility function is mainly used in the context of permanent income and life cycle hypotheses [Bergman 2005].

**Logarithmic Utility Function**

**Definition 3.2** A logarithmic utility function is of the form

\[
U(x) = \log(x).
\]

This is a legitimate utility function as its first derivative \( U'(x) = x^{-1} > 0 \) and second derivative \( U''(x) = -x^{-2} < 0 \).
**Iso-Elastic Utility Functions**

**Definition 3.3** A class called iso-elastic utility functions have the following form

\[
U(x) = \begin{cases} 
\frac{x^{1-a} - 1}{1-a} & \text{for } a > 0, a \neq 1; \\
\log(x) & \text{the limiting case for } a = 1.
\end{cases}
\]

These functions have the property of *iso-elasticity*, which means that we get the same utility function (up to a positive affine transformation) if the cost is scaled by some constant \(k\). Formally,

For all \(k > 0\),

\[
U(kx) = f(k)U(x) + g(k),
\]

for some function \(f(k) > 0\) which is independent of \(x\) and some function \(g(k)\) which is independent of \(x\) as well, see Appendix B for proof.

This iso-elasticity property implies that if a given percentage cost budget is optimal for the current level of budgets, then the same percentage cost budget allocation is optimal for all the other levels of budgets as well.

**Negative Exponential Utility Function**

**Definition 3.4** A negative exponential utility function is of the form

\[
U(x) = -\exp\{-ax\}.
\]

Since the first derivative \(U'(x) = a \exp\{-ax\} > 0\) and the second derivative \(U''(x) = -a^2 \exp\{-ax\} < 0\), this one is also a legitimate utility function.

The class of negative exponential utility functions has an interesting property that it is invariant under any additive cost transformation, *i.e.*, for any constant \(k\),

\[
U(k + x) = f(k)U(x) + g(k),
\]

for some function \(f(k) > 0\) which is independent of \(x\) and some function \(g(k)\) which is independent of \(x\) as well, see Appendix B for proof.

**3.3.2 Expected Utility**

It is natural to weight each cost induced by a decision with the corresponding probability that it will be made. This gives us a utility function of the following form

\[
U(c_1, c_2, \pi_1, \pi_2) = \pi_1 c_1 + \pi_2 c_2.
\]
This expression is actually known as the *expected value*, which is simply the average level of cost that would happen.

One form that the utility function might take is the following:

\[
U(c_1, c_2, \pi_1, \pi_2) = \pi_1 V(c_1) + \pi_2 V(c_2).
\] (3.14)

This means that utility can be written as a weighted sum of some function of each cost, \(V(c_1)\) and \(V(c_2)\), where the weights are in fact the probabilities \(\pi_1\) and \(\pi_2\). Thus equation (3.14) represents the expected utility, of the pattern of cost \((c_1, c_2)\) induced by the relevant decisions \(d_1, d_2\).

We refer to a utility with the form described above as an expected utility function or a utility function that has an expected utility property. When we say that a decision maker’s preferences can be represented by an expected utility function, or that the decision maker’s preferences have the expected utility property, we mean that we are able to choose a utility function that has the additive form described in equation (3.14). And this form also turns out to be especially convenient. It has been proved that an expected utility function has the property of uniqueness, *i.e.*, it is unique up to an affine transformation, which simply means that we can apply an affine transformation to it and obtain another expected utility function that describes the same preferences (Varian, 1992).

The expected utility function can also be subjected to some kinds of monotonic transformation and still have the expected utility property. A function \(V(U)\) is a positive affine transformation if it can be written in the form: \(V(U) = \alpha U + \beta\) where \(\alpha > 0\), which indicates that it not only represents the same preferences but it also still has the expected utility property. It is straightforward to extend a utility function to the case of a finite number of costs induced by decisions. If cost \(c_i\) is associated with probability \(p_i\), for \(i = 1, 2, \ldots, n\), then the expected utility is

\[
\mathbb{E}U(C) = \sum_{i=1}^{n} p_i U(c_i)
\]

And it also holds for continuous probability distribution. If \(f(c)\) is defined as a probability density function on cost \(c\), then the expected utility can be written as

\[
\mathbb{E}U(C) = \int U(c) f(c) \, dc
\]
3.3.3 Risk Aversion

Based on the attitude to risk, we distinguish risk averse, risk neutral, and risk seeking decision makers. Their utility functions are concave, affine, and convex, correspondingly. Most decision makers are assumed to be risk averse and it is often convenient to have a measure of risk aversion.

The coefficient of risk aversion is a special measure reflecting the character and degree of a decision maker’s risk aversion. Intuitively, the more concave the expected utility function is, the more risk averse the decision maker tends to be. We could measure risk aversion by the second derivative of the utility function. However, this definition is sensitive to changes in the utility function: if we consider any positive multiple of the utility function, the second derivative changes but the decision maker’s behaviour does not. If we normalise the second derivative by dividing by the first, we get a reasonable measure known as Arrow-Pratt absolute risk aversion coefficient (Arrow, 1965; Pratt, 1964). The most common measures are the coefficients of absolute risk aversion (ARA) and relative risk aversion (RRA).

Coefficients of Risk Aversion

Definition 3.5 The coefficient of absolute risk aversion at a point $x$ pertaining to a utility function $U$ is defined as

$$\lambda_A(x) = -\frac{U''(x)}{U'(x)}.$$  \hfill (3.15)

Utility functions with a constant absolute risk aversion coefficient are called constant absolute risk aversion (CARA) utility functions.

Definition 3.6 The coefficient of relative risk aversion at a point $x$ pertaining to a utility function $U$ is defined as

$$\lambda_R(x) = -x\frac{U''(x)}{U'(x)} = -x\lambda_A(x).$$  \hfill (3.16)

Utility functions with a constant relative risk aversion coefficient are called constant relative risk aversion (CRRA) utility functions.
Decreasing & Increasing Risk Aversion

**Definition 3.7** If the absolute risk aversion \( \lambda_A(x) \) is decreasing, then we say the decreasing absolute risk aversion (DARA) is present, i.e., the following inequality holds,

\[
\frac{\partial \lambda_A(x)}{\partial x} = -\frac{U'(x)U''(x) - [U''(x)]^2}{[U'(x)]^2} < 0. \tag{3.17}
\]

Also the increasing absolute risk aversion (IARA) is present if \( \frac{\partial \lambda_A(x)}{\partial x} > 0 \).

**Definition 3.8** If the relative risk aversion \( \lambda_R(x) \) is decreasing, then we say the decreasing relative risk aversion (DRRA) is present, i.e., the following inequality holds,

\[
\frac{\partial \lambda_R(x)}{\partial x} < 0. \tag{3.18}
\]

Also the increasing relative risk aversion (IRRA) is present if \( \frac{\partial \lambda_R(x)}{\partial x} > 0 \).

Thus, among the utility functions introduced in §3.3.1, negative exponential utility exhibits constant absolute risk aversion (CARA) and increasing relative risk aversion (IRRA); both the absolute and relative risk aversions of quadratic utility function are increasing; for logarithmic and iso-elastic utility functions, they both exhibit decreasing absolute risk aversion(DARA) and constant relative risk aversion (IRRA). Corresponding results are present in Table 3.6.

<table>
<thead>
<tr>
<th>Utility Function</th>
<th>Coefficient of ARA</th>
<th>Coefficient of RRA</th>
<th>DARA/IARA</th>
<th>DRRA/IRRA</th>
</tr>
</thead>
<tbody>
<tr>
<td>( ax - bx^2 )</td>
<td>( \frac{1}{x+a/2b} )</td>
<td>( \frac{x}{x+a/2b} )</td>
<td>( \frac{1}{(ab/2-x)^2} &gt; 0 )</td>
<td>( \frac{2ab}{(a-2bx)^2} &gt; 0 )</td>
</tr>
<tr>
<td>( \log(x) )</td>
<td>( \frac{1}{x} )</td>
<td>1</td>
<td>( -\frac{1}{x^2} &lt; 0 )</td>
<td>0</td>
</tr>
<tr>
<td>( \frac{x^{1-a}-1}{1-a} )</td>
<td>( \frac{a}{x} )</td>
<td>( a )</td>
<td>( -\frac{a}{x^2} &lt; 0 )</td>
<td>0</td>
</tr>
<tr>
<td>( -\exp{-ax} )</td>
<td>a</td>
<td>( ax )</td>
<td>0</td>
<td>( a &gt; 0 )</td>
</tr>
</tbody>
</table>

**Table 3.6**: Properties of Utility Functions

It is worth noting that a utility function \( U \) exhibits constant absolute risk aversion (CARA) if the absolute risk aversion coefficient does not depend on the resource or \( \lambda'_A(x) = 0 \), and decreasing absolute risk aversion (DARA) is present if decision makers with more resource are less absolutely risk averse than those with less resource or \( \lambda'_A(x) < 0 \). We notice that there is a natural assumption that most decision makers have decreasing absolute risk aversion, e.g., quadratic utility functions, which present
increasing absolute risk aversion, are avoided by economists because quadratic utility functions imply unrealistic behaviour in practice in the sense of absolute risk aversion.

### 3.3.4 Utility in Maintenance

In the context of the problem of maintenance optimisation, let us denote $c$ as the random variable representing systems’ average operating cost. If the random cost $c$, depends among other stochastic or deterministic factors, upon a decision variable $j$ and $J$ is the set of all feasible decisions $j$, the optimal maintenance time solves the problem

$$\max_{j \in J} E(U(c(j))).$$  \hspace{1cm} (3.19)

The crucial thing here is the right choice of the utility function and its parameters, reflecting in particular decision makers’ attitude to risk. Usually, the parameters entering the utility functions are estimated using some statistical methods or psychological experiments.

Kapliński (2013) briefly considers the economic and psychological aspects of decision making in maintenance and repair and discusses risk assessment criteria such as expected value and maximisation of expected utility, of which research results suggest different attitudes towards risk would influence the choice of decisions. For example, a maintenance engineer and production manager would have very different risk preferences: the former would prefer to maintain systems frequently whereas the latter would prefer to keep systems performing consecutively.

Baker (2010) proposes a new concept of minimising the disutility of cost per unit time instead of cost per unit time in maintenance modelling, which provides a maintenance policy that is optimal under risk aversion. But this paper only advocates use of the exponential utility function, thus it would be interesting to explore the use of a different utility function than the exponential.

Houlding and Coolen (2011) address some of the foundational issues of adaptive utility when utility is uncertain, seen from the perspective of a Bayesian statistician, which generalise the traditional utility concepts of value of information and risk aversion. In Houlding and Coolen (2012) they extend their work by combining the decision making with uncertain utility and nonparametric predictive inference, by means of which they present the Nonparametric Predictive Utility Inference (NPUI) suggestion
as a possible strategy for the problem of utility induction in cases of extremely vague information. Meanwhile, Houlding and Coolen (2007) examine how the possibility to learn preferences can be of interest for decisions in the area of reliability, which offers a generalisation of the classical Bayesian approach by adaptive utility for sequential decision making.

Flood et al. (2010) use a Bayesian Network to model the downtime of a system and employ the posterior distribution within a decision analysis. They give an example by computing the expected utilities for a warranty policy and an adaptive form of the former through simulating samples of system downtime from the posterior distribution. Taking maximising the expected utility as the objective, the optimally acceptable downtime range of a system is found simply by using the optim function in R under a continuous decision space.

In software reliability, as software is more frequently used, the reliability of the software increases, which is different from the systems commonly modelled with increasing failure rates (IFR). For example, McDaid and Wilson (2001) propose a decision theoretic solution to the problem of deciding the optimal length of the software testing period by using an error detection model and a sensible utility.

Overall, utility application in maintenance is a relatively new area and few works have been done under maintenance optimisation for repairable systems based on utility theory, thus it is very valuable to explore utility-based maintenance modelling.
Chapter 4

Sequential Preventive Maintenance

A system maintenance policy specifies how the maintenance activities should be scheduled and executed. Each maintenance action is taken to keep the repairable system at the required operation level and it can be minimal repair, perfect maintenance or replacement, etc.

Many models concerning maintenance describe a periodic maintenance policy, where the maintenance frequency and times are pre-determined [Barlow and Proschan, 1965], or fixed prior to modelling set up [Schutz et al., 2011]. This policy has its advantages, for example, it is less complicated to implement in practice if the system maintenance is based on calendar time and as a result, it is popular for practitioners. However, this policy also has its disadvantages due to its inflexibility. The main issue is that this policy is not a globally optimal maintenance policy, which could result in exceptionally expensive costs if the system fails to perform on a desired level due to inadequate and not-in-time maintenance. A Bayesian situation with fixed maintenance times started with the work by [Percy and Kobbacy, 1996]. Damien et al. (2007) analysed a single item maintenance in a Bayesian semi-parametric setting, which solves the drawbacks of other models failing to capture the true underlying relationships in the data, but still with a pre-defined time horizon. Another example is [Baker (2010)] who considers failures of a system under some maintenance policy but where the system may potentially reach a regeneration point $T$. However, again the maintenance time phases are pre-defined, which is not practical in reality.

Myopic maintenance modelling methodology focuses on the next maintenance phase based on the previous and current system status, which fails to consider the possible
maintenance series in the future. As a result, this methodology is not a globally optimal method either. In the study of this chapter, maintenance time phases are initially pre-defined, depending on the particular system, but are flexible and updated with data.

The objective of Chapter 4 is to determine the optimal maintenance schedule times by proposing sequential maintenance models through adopting the Bayesian approach on certain random or unknown parameters of failure distributions. The Bayesian approach could be quite flexible when the failure distributions of the system is either unknown or contains uncertain parameters, which is common in most of the practical situations.

This chapter starts with the problem setting for two-phase maintenance systems, discusses the choice of utility functions, models two-phase systems maintenance by stochastic dynamic programming, and utilises a gridding method to solve problems arising from classical optimisation methods. A few numerical examples follow. Note that the “time” in this thesis is regarded as “local time” (Definition 2.2).

4.1 The Problem Setting

For simplicity and clarification, let us consider a general two-phase repairable system subject to deterioration while running, and which ultimately leads to failure. Under an assumption that, following any failure, a repair is completed instantaneously though it is not necessary, i.e., the duration time of repairs is assumed to be negligible, a natural idea, and one which has already been much studied in the literature (Kobbacy and Murthy 2008; Li and Pham 2006; Wang and Pham 2006), is then to try to prevent some failures of the system by maintaining it.

Here we assume that the initial (or unmaintained) system evolves according to failure times following an arbitrary probability distribution and seek to determine a maintenance policy designed for the system. At the beginning of the process we are required to make a decision about the maintenance time $T_{m1}$. Then we could face the scenario that the failure of the system is observed or not (i.e., failure occurs at $T_{f1}$ which is before or after the maintenance time $T_{m1}$). Based on different scenarios we should be able to set the updated maintenance time for the system accordingly, depending

\[\text{The term “phase” is equivalent to “period” in this thesis.}\]
on if failures are observed or not. In general, our model falls into the category of condition-based maintenance, but with scheduled maintenance times being subject to random failure that induces change in maintenance.

Costs of the system, such as failure cost, repair cost and maintenance cost, are also assumed to depict the properties of the system evolution. These costs are assumed to be constant here for the purpose of simplicity though this is also not a necessary assumption. Our aim is to find the optimal maintenance times according to the running of the system based on the criterion of maximising the expected utility per unit time (i.e., here the negative expected cost per unit time because the payoff is negative cost in this problem). The utility, of course can be altered and incorporated in a more general or specified horizon according to various contexts (either theoretical, practical, or both).

The system considered here has two processing time periods described through a decision tree, see Figure 4.1. At the decision nodes (represented by squares) $T_{m1}$, $T_{m2}$, one has to decide the optimal maintenance time for the associated phase. A phase is defined as the period between the successive occurrence of a failure or a maintenance. Chance nodes (represented by circles) are used to describe the possibility that systems go to various situations, in particular $C_1$ is the chance node for phase 1 and $C_{2i}$ ($i = 1, 2$) are the chance nodes for phase 2. $T_{fj}$ ($j = 1, 2$) are the potential failure times for each phase, while $T_{mi}$ ($i = 1, 2$) is the optimal maintenance times we are attempting to find. Prior to the system running, we need to determine the maintenance time $T_{m1}$ for the first time period, afterwards, the system starts running, where it may face two circumstances: failures can be before the maintenance time $T_{m1}$; or after the maintenance time $T_{m1}$ (right-censored). Then after the maintenance or repair, the system goes to the second running period, again with the same potential consequences as above. All the maintenance time decisions are made according to the average cost of the system processing (average cost per unit time, i.e., cost rate, is expressed at the end of each branch), of which are $C_r$, $C_m$ and $C_f$ representing costs of repair, maintenance and failure of the system, respectively. The utility is a function of cost rates.
Interest centres on the value of the unknown maintenance times $T_{m_i}$ ($i = 1, 2$) for each phase, given observing failure or not. In a non-sequential setting, one could simply choose how many phases the system has and then find a single maintenance time so as to optimise the expected cost or utility per unit time of the system. This method can be repeated for all potential phases. The sequential setting, however, allows the choice of maintenance times to depend on the data observed at successive points, and also on what may be learned in the future.

The cost rate $CR(\cdot)$ is a function with respect to the operation time $T_o$ of the system and the corresponding cost $C$ induced during $T_o$, defined as

$$CR_{T_o} = CR_{T_o}(T_o, C) = \frac{C}{T_o} \quad (4.1)$$

A utility function is defined as $U(CR_{T_m})$ over cost rate $CR_{T_m}$ induced by corresponding maintenance time $T_m$. As a result, the expected utility of cost rate can be written as follows,

$$\mathbb{E}(U(CR_{T_m})) = \int U(CR_{T_m})f(CR_{T_m}) dCR_{T_m} \quad (4.2)$$
where \( f(CR_{Tm}) \) is a probability density function defined on cost rate \( CR_{Tm} \).

In this problem, the expected utility of cost rate at chance node \( CN_1 \) for a one-phase system is

\[
\mathbb{E}_{CN_1}(U(CR_{Tm_1})) = \int U(CR_{Tm_1}(T_1, C_1)) \times f(CR_{Tm_1}(T_1, C_1)) \, dCR_{Tm_1}
\]

\[
= \int_0^{T_{m_1}} U_{T_{m_1}} \left( \frac{C_f + C_r}{t_{f_1}} \right) \times f_{T_{f_1}}(t_{f_1}) \, dt_{f_1} + U_{T_{m_1}} \left( \frac{C_m}{T_{m_1}} \right) \times f_{T_{f_1}}(t_{f_1} > T_{m_1})
\]

where \( U_{T_{m_1}} \) is the utility function with regard to \( T_{m_1} \). \( T_1 \) and \( C_1 \) are the operating time of the one-phase system and corresponding cost, \( i.e. \), \( (C_f + C_r) \) would be induced if \( T_1 = T_{f_1} \) while \( C_m \) would be present if \( T_1 = T_{m_1} \). In other words, the expected utility at chance node \( CN_1 \) is the sum of utility when the system fails before or after maintenance weighted by corresponding probabilities.

The main difference between non-sequential and sequential problem solving is that whilst both use conditional probability based on the evolution of the system by the point of maintenance scheduling, the latter also considers all the possible future states instead of only the next state given any previous history. In other words, the sequential scheme puts all future maintenance schedules into the modelling.

### 4.2 Utility Functions

A few utility functions have been introduced in \[3.3.1\] and their aversion properties in \[3.3.3\] because of the risk aversion degrees of different utility functions, only for very small levels of risk aversion will the same maintenance policy be given via different utility functions, which approximates to minimising the cost rate. The utility functions introduced in \[3.3.1\] are mathematically tractable, however, one may ask if there are other utility functions presenting more realistic and practical maintenance policies. With the rapid development of computation techniques and software, one tends to prefer approximate solutions to exactly mathematical ones in statistics and operational research.

The average cost per unit time, or cost rate, is used herein, which is functionally written as \( C/T \), where \( C \) and \( T \) are associated costs and corresponding times.
utility function captures both the core issues of cost and time. Consider two maintenance policies 1 and 2 which have cost rates $CR_1$ and $CR_2$, with current cost budget $B$ which is planned for maintenance in particular. Then one tends to prefer maintenance policy 1 if $E[U(B - CR_1)] > E[U(B - CR_2)]$ for every $B$. Pranzagl (1959) and Bell (1988) have proved that the negative exponential and linear utility function families are the only ones for which preference holds regardless of budget $B$.

Bell and Fishburn (2001) suggest that the following utility

$$U(x) = x - \alpha \exp\{-\beta x\}, \quad \alpha > 0, \beta > 0 \quad (4.5)$$

should be used because its form of sum of linear and exponential has a few properties:

- It is a legitimate utility function because $U'$ is positive and $U''$ is negative.
- It meets a contextual uncertainty condition, that is larger uncertainties in $B$ should be preferred to be solved to small ones before decision making (Bell, 1995).
- It satisfies the so-called “one-switch” rule, which means preference for one of two maintenance policies is allowed to change only once as the cost budget $B$ increases.
- Its absolute risk aversion is decreasing as

$$\frac{\partial^2 U(x)}{\partial x^2} = -\frac{\alpha \beta^3 \exp(\beta x)}{(\alpha \beta + \exp(\beta x))^2} < 0.$$

However, it can be problematic to apply this utility function in practice: it adds difficulty in eliciting the two risk aversion parameters $\alpha$ and $\beta$; it is required to know a decision maker’s cost budget $B$; the optimal maintenance policy would change to a riskier one as the cost budget $B$ increases. As a result, the negative exponential utility function family will be used in this thesis to optimise the maintenance time of systems, though it is still worthwhile studying these drawbacks. Through an affine transformation of the original negative utility function, the following utility function will be used:

$$U(x) = \frac{1 - \exp(-\eta x)}{\eta} \quad (4.6)$$

where $\eta$ is the risk aversion parameter.

As it can be seen in Figure [1.2], this utility function becomes more risk averse when the risk aversion parameter $\eta$ increases. Note that $x$ will be treated in terms of cost rate function $CR$ in our maintenance modelling.
4.3 Modelling Maintenance

Let $T$ follow an arbitrary or unknown distribution, of which there exists its probability density function $f(t)$ with parameter(s) set $\Theta$, written as $f(t | \Theta)$.

Its hazard function is $h(t | \Theta)$, and depending on the functional property of parameters $\Theta$, the hazard function can be decreasing, constant or increasing. Systems considered here have been assumed to have increasing hazard rate when they are working, which meets the characteristics of most industrial systems in practice though this is not necessary for general applications.

To solve the sequential maintenance optimisation problem, a generalised form of the stochastic dynamic programming algorithm for this specific problem is given as follows.

Based on a Bayesian framework, we give hyper-parameter(s) $\Delta$ with a specific distribution $g(\Theta | \Delta)$ for the parameters $\Theta$, and through observing data, we update our belief with respect to parameters $\Theta$.

4.3.1 Myopic Modelling

Myopic is the most elementary modelling method for sequential maintenance. Myopic modelling optimises the cost per unit time, but does not explicitly use forecasted
information or any direct representation of decisions about maintenance times in the future; in other words, it makes no explicit attempt to capture the impact of a current maintenance time on the future. In its most basic form, a myopic policy can be given by

$$T_m^{\text{myopic}} = \arg \max U(CR_{T_m})$$  \hspace{1cm} (4.7)$$

where $T_m^{\text{myopic}}$ is the optimum maintenance time and $U(\cdot)$ is a utility function of cost rate.

In the two-phase sequential maintenance problem, the optimum maintenance time for phase one under myopic modelling is

$$T_{m_{CN1}}^{\text{myopic}} = \arg \max \left\{ \int_0^{t_{f_1}} U_{T_{m_1}} \left( \frac{C_f + C_r}{T_{f_1}} \right) \times f_{T_{f_1}}(t_{f_1}) dt_{f_1} + U_{T_{m_1}} \left( \frac{C_m}{T_{m_1}} \right) \times f_{T_{f_1}}(t_{f_1} > T_{m_1}) \right\}$$  \hspace{1cm} (4.8)$$

Note that the function here is the expected utility of cost rate.

Similarly for chance nodes $CN_{21}$ and $CN_{22}$, the optimum maintenance times under myopic modelling can be expressed as follows, respectively:

$$T_{m_{2CN21}}^{\text{myopic}} = \arg \max \left\{ \int_0^{t_{f_2}} U_{T_{m_2}} \left( \frac{C_f + C_r}{T_{f_2}} \right) \times f_{T_{f_2}}(t_{f_2} | t_{f_1}) dt_{f_2} + U_{T_{m_2}} \left( \frac{C_m}{T_{m_2}} \right) \times f_{T_{f_2}}(t_{f_2} > T_{m_2} | t_{f_1}) \right\}$$  \hspace{1cm} (4.9)$$

$$T_{m_{2CN22}}^{\text{myopic}} = \arg \max \left\{ \int_0^{t_{f_2}} U_{T_{m_2}} \left( \frac{C_f + C_r}{T_{f_2}} \right) \times f_{T_{f_2}}(t_{f_2} | t_{f_1} > T_{m_{1CN1}}^{\text{myopic}}) dt_{f_2} + U_{T_{m_2}} \left( \frac{C_m}{T_{m_2}} \right) \times f_{T_{f_2}}(t_{f_2} > t_{m_2} | t_{f_1} > T_{m_{1CN1}}^{\text{myopic}}) \right\}$$  \hspace{1cm} (4.10)$$

Briefly, the procedure to find the optimum maintenance times for two-phase systems under myopic modelling are as below:

1. Maximise the expected utility function of cost rate for phase one to find the optimum maintenance time, $T_{m_1}^*$;

2. Similarly, maximise the expected utility function of cost rate for phase two and find the optimum maintenance times, $T_{CN_{21},m_2}$ and $T_{CN_{22},m_2}$, given additional data with regard to the outcome of the first phase.

From the equations above, we can see that the maintenance times obtained are not globally optimal as the myopic method only utilises previous information to make
decisions based on the current state of knowledge instead of taking future states into account. In order to find the globally optimum maintenance time, it is required to propose other methods.

4.3.2 Stochastic Dynamic Programming

In contrast to myopic modelling, dynamic programming considers the two-phase sequential maintenance problem from the beginning of modelling; in other words, the optimum maintenance time for chance node $CN_1$ is solved by considering all the possible states in the second phase, *i.e.*, the system fails before planned perfect preventive maintenance ($T_{f_2} \leq T_{m_2}$) or afterwards ($T_{f_2} > T_{m_2}$).

Based on the framework of dynamic programming under uncertainty in §3.2.3, the return functions for the maintenance time optimisation problem for stage 1 $s_1$ and stage 2 $s_2$ can be written in the form of (3.11) as

$$
\gamma_2 \left( d_2 = T_{m_2}, s_1, Z_2 = \begin{cases} t_{f_2} < T_{m_2} | t_{f_1} < T_{m_1} \\
T_{f_2} > T_{m_2} | t_{f_1} < T_{m_1} \\
T_{f_2} < T_{m_2} | t_{f_1} > T_{m_1} \\
T_{f_2} > T_{m_2} | t_{f_1} > T_{m_1} \end{cases} \right), \quad (4.11)
$$

$$
\gamma_1 \left( d_1 = T_{m_1}, s_2, Z_1 = \begin{cases} t_{f_1} < T_{m_1} \\
T_{f_1} > T_{m_1} \end{cases} \right), \quad (4.12)
$$

where $\gamma_i$, $d_i$ and $Z_i$, $i = 1, 2$, are best possible solutions from stage $i$ to the end, decision variables and state variables, respectively.

As a result, we suggest use “roll-back” method, where we solve the optimal maintenance times for phase two first, *i.e.*, $T_{m_2}$ for chance nodes $CN_{21}$ and $CN_{22}$; and then input the maximum expected utility of cost rate for phase two back into the expected utility function of cost rate for phase one, and hence solve to obtain the globally optimum maintenance time $T_{m_1}$ for phase one.

$$
T_{m_2}^{\text{DP, CN}_{21}} = \arg \max \left\{ \int_0^{T_{m_2}} U_{T_{m_2}} \left( \frac{2(C_f + C_r)}{T_{f_1} + T_{f_2}} \right) \times f_{T_{f_2}}(t_{f_2} | t_{f_1}) dt_{f_2} + U_{T_{m_2}} \left( \frac{C_f + C_r + C_m}{T_{f_1} + T_{m_2}} \right) \times f_{T_{f_2}}(t_{f_2} > T_{m_2} | t_{f_1}) \right\} \quad (4.13)
$$

$$
T_{m_2}^{\text{DP, CN}_{22}} = \arg \max \left\{ \int_0^{T_{m_2}} U_{T_{m_2}} \left( \frac{C_m + C_f + C_r}{T_{m_1} + T_{m_2}} \right) \times f_{T_{f_2}}(t_{f_2} | t_{f_1} > T_{m_1}) dt_{f_2} + U_{T_{m_2}} \left( \frac{2C_m}{T_{m_1} + T_{m_2}} \right) \times f_{T_{f_2}}(t_{f_2} > T_{m_2} | t_{f_1} > T_{m_1}) \right\} \quad (4.14)
$$
Accordingly, the maximum expected utility of cost rate for chance nodes $CN_{21}$ and $CN_{22}$ are as follows, respectively:

$$E(CN_{21}) = \int_{0}^{T_{m2_{CN_{21}}}} U_{T_{m2}} \left( \frac{2(C_f + C_r)}{T_f + T_{f2}} \right) \times f_{T_{f2}}(t_{f2} \mid t_{f1}) \, dt_{f2}$$

$$+ U_{T_{m2}} \left( \frac{C_f + C_r + C_m}{T_f + T_{m2_{CN_{21}}}} \right) \times f_{T_{f2}}(t_{f2} > T_{m2_{CN_{21}}} \mid t_{f1})$$

(4.15)

$$E(CN_{22}) = \int_{0}^{T_{m2_{CN_{22}}}} U_{T_{m2}} \left( \frac{C_m + C_f + C_r}{T_{m1} + T_{f2}} \right) \times f_{T_{f2}}(t_{f2} \mid t_{f1} > T_{m1}) \, dt_{f2}$$

$$+ U_{T_{m2}} \left( \frac{2C_m}{T_{m1} + T_{m2_{CN_{22}}}} \right) \times f_{T_{f2}}(t_{f2} > T_{m2_{CN_{22}}} \mid t_{f1} > T_{m1})$$

(4.16)

For phase one, replace the expected utility of cost rate $E(CN_{21})$ and $E(CN_{22})$ back into the expected utility function of cost rate for phase one, i.e.,

$$T_{m1_{CN_{1}}}^{DP} = \arg \max \left\{ \int_{0}^{T_{m1}} U_{T_{m1}} \left( \frac{E(CN_{21})}{T_{f1}} \right) \times f_{T_{f1}}(t_{f1}) \, dt_{f1} \\
+ U_{T_{m1}} \left( \frac{E(CN_{22})}{T_{m1}} \right) \times f_{T_{f1}}(t_{f1} > t_{m1}) \right\}$$

(4.17)

Then the globally optimum maintenance time for phase one by dynamic programming method is $T_{m1_{CN_{1}}}^{DP}$.

Briefly, the procedure to find the optimum maintenance times for two-phase systems under dynamic programming modelling are as below:

1. Maximise the expected utility function of cost rate for phase two to find the optimum maintenance times, $T_{m2}^{*}$ and corresponding maximum expected utility of cost rate $E(T_{m2}^{*})$;

2. Then plug the obtained maximum expected utility of cost rate $E(T_{m2}^{*})$ in the expected utility function of cost rate for phase one to find the globally optimum maintenance time $T_{m1}^{DP}$.

### 4.3.3 Bayesian Weibull Modelling

For example, let failure time $T_f$ follow a Weibull distribution with scale parameter $\kappa$ and shape parameter $\theta$. The probability density function (pdf) of $T_f$ is given by

$$f_{T_f}(t_f \mid \kappa, \theta) = \begin{cases} \frac{\theta \kappa}{\kappa} (\frac{t_f}{\kappa})^{\theta-1} \exp \left\{ - \left( \frac{t_f}{\kappa} \right)^{\theta} \right\} & t_f > 0, \kappa > 0, \theta > 0; \\
0 & \text{otherwise.} \end{cases}$$
Its hazard function is \( h_{T_f}(t_f \mid \kappa, \theta) = \frac{\theta}{\kappa} \left( \frac{t_f}{\kappa} \right)^{\theta-1} \), so depending on the shape parameter \( \theta \), the hazard function can be decreasing, constant or increasing. The Weibull distribution has been widely used in practice for modelling the failure time of systems, see Houlding and Wilson (2011); Singpurwalla and Wilson (1999).

The hierarchical Bayesian method is applied to modelling systems considered in this thesis. The modelling centres on the uncertainty of shape parameter \( \theta \) in the Weibull probability distribution. \( \theta \) is assumed to follow a truncated normal distribution in order to make sure that \( \theta \) can well express the property of hazard function, i.e., increasing, decreasing or constant. The hyper-parameter \( \mu \) in the truncated normal distribution is assumed to be a uniform prior because one knows it is between a range from knowledge by experts, but no other information is available about its location. In specifying the model, define the following:

\[
\begin{align*}
T_f & \sim \text{Weibull} \ (\theta, \kappa) \quad (4.18) \\
\theta & \sim \text{Truncated Normal} \ (\mu, \sigma \mid a, b) \quad (4.19) \\
\mu & \sim \text{Uniform} \ (a_0, b_0) \quad (4.20)
\end{align*}
\]

(4.18) is the Weibull probability model with shape parameter \( \theta \) and scale parameter \( \kappa \).

(4.19) is the prior distribution of the shape parameter \( \theta \), which is a normal distribution truncated at \( a \) and \( b \), where \(-\infty < a < b < \infty\), with mean \( \mu \) and standard deviation \( \sigma \), with the hyper-parameters made explicitly as \( \mu \). Its probability density function (pdf) \( f \), for \( a \leq \theta \leq b \), is given by

\[
f(\theta \mid \mu, \sigma, a, b) = \frac{1}{\sigma \phi \left( \frac{x-\mu}{\sigma} \right)} \Phi \left( \frac{b-\mu}{\sigma} \right) - \Phi \left( \frac{a-\mu}{\sigma} \right) \quad (4.21)
\]

where \( \phi(\zeta) = \frac{1}{\sqrt{2\pi}} \exp \left( -\frac{1}{2} \zeta^2 \right) \) is the probability density function of the standard normal distribution \( N(0,1) \) and \( \Phi(\cdot) \) is its cumulative distribution function. See Figure 4.3 for the probability density function of \( \theta \). The left graph shows normal distributions truncated at 1 with different sets of parameters, with mean \( \mu = 1, 2, 3, 4 \) and standard deviation \( \sigma = 1 \); it can be seen that \( \theta \) has higher probability near the mean and lower probability near the tail, with smaller mean \( \mu \). The right graph shows normal distributions truncated at various points at 0.5, 1, 1.5, 2 with same mean \( \mu = 2 \) and standard deviation \( \sigma = 1 \); we may see that \( \theta \) has higher probability near the mean when the truncation point is closer to the mean \( \mu \).
Figure 4.3: Comparison of truncated normals with various mean (left) and truncating points (right).

Crucially, \( \mu \) is not specified directly, but has hyper-prior distribution (4.20) which is a uniform distribution with parameters \( a_0 \) and \( b_0 \). Its probability density function (pdf) is given by

\[
f(\mu | a_0, b_0) = \begin{cases} 
\frac{1}{b_0 - a_0} & \theta \in [a_0, b_0]; \\
0 & \text{otherwise.}
\end{cases}
\]

This full model involves parameters \( \xi = (\theta, \mu) \). The mean and variance of these distributions can be expressed as

\[
\mathbb{E} (T_{f_i} | \theta, \kappa) = \kappa \Gamma \left( 1 + \frac{1}{\theta} \right) \tag{4.22}
\]

\[
\text{Var} (T_{f_i} | \theta, \kappa) = \kappa^2 \left[ \Gamma \left( 1 + \frac{2}{\theta} \right) - \left( \Gamma \left( 1 + \frac{1}{\theta} \right) \right)^2 \right] \tag{4.23}
\]

\[
\mathbb{E} (\theta | a, b) = \mu + \frac{\phi \left( \frac{a - \mu}{\sigma} \right) - \phi \left( \frac{b - \mu}{\sigma} \right)}{\Phi \left( \frac{b - \mu}{\sigma} \right) - \Phi \left( \frac{a - \mu}{\sigma} \right)} \tag{4.24}
\]

\[
\text{Var} (\theta | a, b) = \sigma^2 \left[ 1 + \frac{a - \mu}{\sigma} \phi \left( \frac{a - \mu}{\sigma} \right) - \frac{b - \mu}{\sigma} \phi \left( \frac{b - \mu}{\sigma} \right) - \frac{\phi \left( \frac{a - \mu}{\sigma} \right) - \phi \left( \frac{b - \mu}{\sigma} \right)}{\Phi \left( \frac{b - \mu}{\sigma} \right) - \Phi \left( \frac{a - \mu}{\sigma} \right)}^2 \right] \tag{4.25}
\]
\[ \mathbb{E}(\mu) = \frac{1}{2}(a_0 + b_0) \]  
\[ \text{Var}(\mu) = \frac{1}{12}(b_0 - a_0)^2 \]  
\[ 4.26 \]  
\[ 4.27 \]

Systems we discuss here are assumed to have increasing hazard when they are running, which meets the characteristics of most industrial systems in practice though this is not necessary for general applications.

To solve the sequential maintenance optimisation problem, a generalised form of the stochastic dynamic programming algorithm for this specific problem is given below.

Let the failure times of the system follow a Weibull distribution with shape parameter \( \theta \) and scale parameter 1 (for simplicity as our concern is about shape parameter \( \theta \)). We give a prior for \( \theta \) as a truncated normal distribution \( N(\mu, 1) \) truncated at 1 as the hazard function would decrease sharply at the initial time if the shape parameter \( \theta \) is less than 1. The mean of truncated normal distribution of \( \theta \) is given by \( \mathbb{E}(\mu) = 2 \) when the hyperparameter distribution of \( \mu \) is a uniform distribution with \( a_0 = 1 \) and \( b_0 = 3 \). Thus, related functional forms can be written as

\[
f(\theta \mid \mu = 2, \sigma = 1, a = 1, b = \infty) = \frac{\phi(\theta - 2)}{\Phi(1)} = \frac{1}{\sqrt{2\pi}} \exp \left\{ -\frac{1}{2}(\theta - 2)^2 \right\} \Phi(1) \]

\[
f_{T_{f_i}}(t_{f_i} \mid \theta) = \theta(t_{f_i})^{\theta-1} \exp \left\{ -(t_{f_i})^\theta \right\}, \quad i = 1, 2 \]

\[
f_{T_{f_1}}(t_{f_1} > T_{m_1} \mid \theta) = \exp \left\{ -(T_{m_1})^\theta \right\} \]

\[ 4.28 \]
\[ 4.29 \]
\[ 4.30 \]

where \( \phi(\zeta) = \frac{1}{\sqrt{2\pi}} \exp(-\frac{1}{2}\zeta^2) \) is the probability density function of the standard normal distribution and \( \Phi(\cdot) \) is its cumulative distribution function. The expectation and variance of failure \( T_{f_i}, i = 1, 2 \) with respect to \( \theta \) are

\[
\mathbb{E}(T_{f_i} \mid \theta, \kappa = 1) = \Gamma \left( 1 + \frac{1}{\theta} \right) \]

\[
\text{Var}(T_{f_i} \mid \theta, \kappa = 1) = \Gamma \left( 1 + \frac{2}{\theta} \right) - \left( \Gamma \left( 1 + \frac{1}{\theta} \right) \right)^2 \]

\[ 4.31 \]
\[ 4.32 \]

As we can see in Figure 4.4, with the increasing of \( \theta \), the expectation of \( T_f \) increases and variance of \( T_f \) decreases, which means the system is less likely to fail with the increase of \( \theta \).
The prior distribution of $\theta$ is assumed to be a truncated normal distribution truncated at 1 with mean 2 and standard deviation 1, see Figure 4.5. The expectation and variance of $\theta$ are as

\[
\begin{align*}
\mathbb{E} (\theta \mid \mu = 2, \sigma = 1, a = 1) &= \mu + \sigma \frac{\phi \left( \frac{a - \mu}{\sigma} \right)}{1 - \Phi \left( \frac{a - \mu}{\sigma} \right)} \\
&= 2 + \frac{\phi(1)}{\Phi(1)} \quad (4.33) \\
\text{Var} (\theta \mid \mu = 2, \sigma = 1, a = 1) &= \sigma^2 \left[ 1 - \frac{\phi \left( \frac{a - \mu}{\sigma} \right)}{1 - \Phi \left( \frac{a - \mu}{\sigma} \right)} \left( \frac{\phi \left( \frac{a - \mu}{\sigma} \right)}{1 - \Phi \left( \frac{a - \mu}{\sigma} \right)} - \frac{a - \mu}{\sigma} \right) \right] \\
&= 1 - \frac{\phi(1)}{\Phi(1)} - \left( \frac{\phi(1)}{\Phi(1)} \right)^2 \quad (4.34)
\end{align*}
\]

Figure 4.5: Prior distribution of $\theta$: truncated normal distribution $N(2, 1)$ truncated at 1.
It can be seen from Figure 4.6 that the expectation and variance of parameter $\theta$ increases and decreases respectively, when the truncating point $a$ is moving from 1 to 2. Then the prior distribution of $\theta$ approaches to a left skewed and leptokurtic truncated normal distribution.

![Figure 4.6: Expectation of $\theta$ (left) and variance of $\theta$ (right) given $a$ and $b = \infty$.](image)

Failure time $T_{f_1}$ follows a Weibull distribution with shape parameter $\theta$ and scale parameter 1. By integrating over $\theta$, we may obtain the marginal distribution of $T_{f_1}$.

$$f_{T_{f_1}}(t_{f_1}) = \int_{\theta} f_{T_{f_1}}(t_{f_1} \mid \theta) f(\theta) \, d\theta$$

$$= \int_1^\infty \frac{\theta \exp\{-\frac{1}{2}(\theta - 2)^2 - (t_{f_1})^\theta\} (t_{f_1})^{\theta-1}}{\sqrt{2\pi}} \left(1 - \frac{1}{2} \text{Erfc} \left( -\frac{1}{2\sqrt{2}} \right) \right) \, d\theta$$

where Erfc($\cdot$) is the complementary error function.$^2$

Figure 4.7 shows the marginal density function of failure time $T_{f_1}$.

$^2$Note that the error function is $\text{Er}(\rho) = \frac{2}{\sqrt{\pi}} \int_0^\rho \exp\{-u^2\} \, du$ and $\text{Erfc}(\rho) = 1 - \text{Er}(\rho) = \frac{2}{\sqrt{\pi}} \int_\rho^\infty \{ -u^2 \} \, du$.  

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4.3.3.1 One-Phase System

For a one-phase system, the expected utility of cost rate at decision node $CN_1$ is

$$
E_{CN_1}(U(CR_{T_{m_1}})) = \int_0^{T_{m_1}} U_{T_{m_1}} \left( \frac{C_f + C_r}{t_{f_1}} \right) \times f_{T_{f_1}}(t_{f_1}) \, dt_{f_1} + U_{T_{m_1}} \left( \frac{C_m}{T_{m_1}} \right) \times f_{T_{f_1}}(t_{f_1} > T_{m_1}) \tag{4.36}
$$

$$
= \int_0^{T_{m_1}} \int_1^{\infty} U_{T_{m_1}} \left( \frac{C_f + C_r}{t_{f_1}} \right) \times \frac{\theta \exp\{-\frac{1}{2}(\theta - 2)^2 - (t_{f_1})^\theta\}(t_{f_1})^{\theta-1}}{\sqrt{2\pi} \left( 1 - \frac{1}{2}\text{Erfc} \left( -\frac{1}{\sqrt{2}} \right) \right)} \, dt_{f_1} \, d\theta + U_{T_{m_1}} \left( \frac{C_m}{T_{m_1}} \right) \times \exp \left\{ -(T_{m_1})^\theta \right\} \tag{4.37}
$$

where $U(\cdot)$ is the exponential utility function defined in (4.6), which represents the utility induced by mean cost per unit time here. This expected utility function has one random variable $T_{f_1}$.

4.3.3.2 Two-Phase System

For a two-phase system, the conditional probabilities have the following forms:
\[
f_{T_{f_2}}(t_{f_2} \mid t_{f_1}) = \frac{f(t_{f_1}, t_{f_2})}{f(t_{f_1})} = \frac{\int \exp \left\{ -\frac{1}{2} (\theta - 2)^2 - (t_{f_2})^\theta - (t_{f_1})^\theta \right\} \, d\theta}{\int \exp \left\{ -\frac{1}{2} (\theta - 2)^2 - (t_{f_2})^\theta - (t_{f_1})^\theta \right\} \, d\theta}
\]

\[
f_{T_{f_2}}(t_{f_2} \mid t_{f_1} > T_{m_1}) = \frac{f(t_{f_2}, t_{f_1} > T_{m_1})}{f(t_{f_1} > T_{m_1})} = \frac{\int \exp \left\{ -\frac{1}{2} (\theta - 2)^2 - (t_{f_2})^\theta - (T_{m_1})^\theta \right\} \, d\theta}{\int \exp \left\{ -\frac{1}{2} (\theta - 2)^2 - (T_{m_1})^\theta \right\} \, d\theta}
\]

\[
f_{T_{f_2}}(t_{f_2} > T_{m_2} \mid t_{f_1}) = \frac{f(t_{f_2} > T_{m_2}, t_{f_1})}{f(t_{f_1})} = \frac{\int \exp \left\{ -\frac{1}{2} (\theta - 2)^2 - (t_{f_2})^\theta - (T_{m_2})^\theta \right\} \, d\theta}{\int \exp \left\{ -\frac{1}{2} (\theta - 2)^2 - (T_{m_2})^\theta \right\} \, d\theta}
\]

\[
f_{T_{f_2}}(t_{f_2} > T_{m_2} \mid t_{f_1} > T_{m_1}) = \frac{f(t_{f_2} > T_{m_2}, t_{f_1} > T_{m_1})}{f(t_{f_1} > T_{m_1})} = \frac{\int \exp \left\{ -\frac{1}{2} (\theta - 2)^2 - (t_{f_2})^\theta - (T_{m_1})^\theta \right\} \, d\theta}{\int \exp \left\{ -\frac{1}{2} (\theta - 2)^2 - (T_{m_1})^\theta \right\} \, d\theta}
\]
Correspondingly, the expected utility of cost per unit time for decision nodes $CN_{21}$, $CN_{22}$ and $CN_1$ can be written as

$$
E_{CN_{21}}(U(CR_{T_{m_2}}))
= \int_0^{T_{m_2}} U_{T_{m_2}} \left( \frac{2(C_f + C_r)}{t_{f_1} + t_{f_2}} \right) \times f_{T_{f_2}}(t_{f_2} \mid t_{f_1}) \, dt_{f_2} \\
+ U_{T_{m_2}} \left( \frac{C_f + C_r + C_m}{t_{f_1} + T_{m_2}} \right) \times f_{T_{f_2}}(t_{f_2} > T_{m_2} \mid t_{f_1})
$$

(4.42)

$$
E_{CN_{22}}(U(CR_{T_{m_2}}))
= \int_0^{T_{m_2}} U_{T_{m_2}} \left( \frac{C_m + C_f + C_r}{T_{m_1} + t_{f_2}} \right) \times f_{T_{f_2}}(t_{f_2} \mid t_{f_1} > T_{m_1}) \, dt_{f_2} \\
+ U_{T_{m_2}} \left( \frac{2C_m}{T_{m_1} + T_{m_2}} \right) \times f_{T_{f_2}}(t_{f_2} > T_{m_2} \mid t_{f_1} > T_{m_1})
$$

(4.43)

$$
E_{CN_1}(U(CR_{T_{m_1}}))
= \int_0^{T_{m_1}} E_{CN_{21}}(U(CR_{T_{m_2}}))f_{T_{f_1}}(t_{f_1}) \, dt_{f_1} \\
+ E_{CN_{22}}(U(CR_{T_{m_2}}))f_{T_{f_1}}(t_{f_1} > T_{m_1})
$$

(4.44)

We can see that these conditional probabilities are complicated and do not have analytically mathematical form.

Correspondingly, the expected utility of cost per unit time for decision nodes $CN_{21}$, $CN_{22}$ and $CN_1$ can be written as
Similarly, due to the complicated mathematical forms of conditional probabilities, the expected utility of cost per unit time for decision nodes $CN_{21}$ and $CN_{22}$ do not have analytically mathematical forms either.

By integrating with respect to $\theta$ in $f(t_{f_1}, t_{f_2}), f(t_{f_1}), f(t_{f_1} > T_{m_1}, t_{f_2})$ and $f(t_{f_1} > T_{m_1})$, we can obtain $f(t_{f_2} | t_{f_1})$ and $f(t_{f_2} | t_{f_1} > T_{m_1})$ and plug them into the expected utility functions of average cost per unit time. Applying numerical methods, one could maximise functions to find the optimal maintenance time for $T_{m_2}$ given observed $t_{f_1}$ or determined $T_{m_1}$.

Then the problem of finding the optimal maintenance $T_{m_2}$ for the second phase can be graphed as follows, see Figure 4.8. If we observe a failure at $t_{f_1}$, in order to find the optimal $T_{m_2}$ given observed $t_{f_1}$, we can simply refer to the coloured part of Figure 4.8 in which $t_{f_1} < T_{m_1}$, then find the optimal $T_{m_2}$ on the $T_{m_2}$ axis. Thus, a two dimensional problem with respect to $t_{f_1}, T_{m_1}$ is transformed into a one dimensional problem with regard only to the relation of $t_{f_1}$ and $T_{m_1}$. 

$$
\times \left( \int_{0}^{T_{m_2}} U_{T_{m_2}} \left( \frac{2(C_r + C_f)}{t_{f_1} + t_{f_2}} \right) \frac{\int_{1}^{\infty} \exp \{ -\frac{1}{2}(\theta - 2)^2 - (t_{f_1})^\theta - (t_{f_2})^\theta \} \ d\theta}{\int_{1}^{\infty} \exp \{ -\frac{1}{2}(\theta - 2)^2 - (t_{f_1})^\theta \} \ d\theta} \ dt_{f_2} \\
+ U_{T_{m_2}} \left( \frac{C_r + C_f + C_m}{t_{f_1} + T_{m_2}} \right) \frac{\int_{1}^{\infty} \exp \{ -\frac{1}{2}(\theta - 2)^2 - (t_{f_1})^\theta - (T_{m_2})^\theta \} \ d\theta}{\int_{1}^{\infty} \exp \{ -\frac{1}{2}(\theta - 2)^2 - (t_{f_1})^\theta \} \ d\theta} \ dt_{f_1} \\
+ \exp (- (T_{m_1})^\theta) \right) \\
\times \left( \int_{0}^{T_{m_2}} U_{T_{m_2}} \left( \frac{C_m + C_r + C_f}{t_{f_2} + T_{m_1}} \right) \frac{\int_{1}^{\infty} \exp \{ -\frac{1}{2}(\theta - 2)^2 - (t_{f_2})^\theta - (T_{m_1})^\theta \} \ d\theta}{\int_{1}^{\infty} \exp \{ -\frac{1}{2}(\theta - 2)^2 - (T_{m_1})^\theta \} \ d\theta} \ dt_{f_2} \\
+ U_{T_{m_2}} \left( \frac{2C_m}{T_{m_1} + T_{m_2}} \right) \frac{\int_{1}^{\infty} \exp \{ -\frac{1}{2}(\theta - 2)^2 - (T_{m_1})^\theta - (T_{m_2})^\theta \} \ d\theta}{\int_{1}^{\infty} \exp \{ -\frac{1}{2}(\theta - 2)^2 - (T_{m_1})^\theta \} \ d\theta} \right)
$$
4.3.4 Gridding Approach

As we can see in Figure 4.1, the branches of the decision tree increase as $2^n$, where $n$ is the number of phases or time periods, which results in issues of computation time; in addition, even for a simple two-phase maintenance optimisation, based on dynamic programming, the optimum maintenance time for phase one is determined by the subsequent optimum times of phase two requiring solutions of nested series of maximisations and integrations over a highly non-linear space, which have no analytical forms.

It is worthwhile to mention that Houlding et al. (2015) proposed a conjugate class of utility functions for sequential decision problems. However, due to the fact that different utility functions are integrated by different intervals instead of the whole real line as it was applied, we cannot simply apply the method to our modelling. Hence, a gridding method for sequential decision problems such as in Brockwell and Kadane (2003) is considered. One can construct an approximation to the expected cost per unit time by evaluating it at the points of a grid and storing the results for the current phase; then one can go back to the previous phase and compute the expected cost per unit time for the previous phase by also evaluating at grid points. With this step finished, it is not necessary to keep the value of the current phase, and the related storage space
can be released. However, in our decision-making process, it is necessary to keep the value of maintenance time $T_m$ for each phase. This process can be repeated until one has found the optimal decision (initial maintenance time here) for the beginning time point.

To be more precise, we introduce the notation as follows: select the lower and upper bounds $b_i^l$ and $b_i^u$, with $b_i^u > b_i^l$, as well as a number of subdivisions $n_i$, for $i = 1, \ldots, K$, which is the number of phases. Define grid points

$$b_i = (b_i^l, b_i^l + \delta, b_i^l + 2\delta, \ldots, b_i^l + n_i\delta = b_i^u)$$

where the grid intervals $\delta_i$ are given by

$$\delta_i = (b_i^u - b_i^l)/n_i$$

For the gridding method, the formula in Equation 4.35 can be written in discrete form as:

$$p_{T_{f1}}(t_{f1}) = \sum_\theta p_{T_{f1}}(t_{f1} | \theta)p(\theta)$$

$$\mathbb{E}_{CN21}(U(CR_{T_{m2}})) = \sum_{T_{f2}=0}^{T_{m2}} U_{T_{m2}} \left( \frac{2(C_f + C_r)}{t_{f1} + t_{f2}} \right) \times p_{T_{f2}}(t_{f2} | t_{f1})$$

$$+ U_{T_{m2}} \left( \frac{C_f + C_r + C_m}{t_{f1} + T_{m2}} \right) \times p_{T_{f2}}(t_{f2} > T_{m2} | t_{f1})$$

$$\mathbb{E}_{CN22}(U(CR_{T_{m3}})) = \sum_{T_{f2}=0}^{T_{m2}} U_{T_{m2}} \left( \frac{C_m + C_f + C_r}{T_{m1} + t_{f2}} \right) \times p_{T_{f2}}(t_{f2} | t_{f1} > T_{m1})$$

$$+ U_{T_{m2}} \left( \frac{2C_m}{T_{m1} + T_{m2}} \right) \times p_{T_{f2}}(t_{f2} > T_{m2} | t_{f1} > T_{m1})$$

where

$$p_{T_{f2}}(t_{f2} | t_{f1}) = \frac{p(t_{f1}, t_{f2})}{p_{T_{f1}}(t_{f1})} = \frac{\sum_\theta p_{T_{f2}}(t_{f2} | \theta)p_{T_{f1}}(t_{f1} | \theta)p(\theta)}{\sum_\theta p_{T_{f1}}(t_{f1} | \theta)p(\theta)}$$

$$p_{T_{f2}}(t_{f2} | t_{f1} > T_{m1}) = \frac{p(t_{f1} > T_{m1}, t_{f2})}{p_{T_{f1}}(t_{f1} > T_{m1})}$$
\[
\sum_{\theta} p_{T_{f_2}}(t_{f_2} \mid \theta)p_{T_{f_1}}(t_{f_1} > T_{m_1} \mid \theta)p(\theta)
\]
\[
\sum_{\theta} p_{T_{f_1}}(t_{f_1} > T_{m_1} \mid \theta)p(\theta)
\]

By this gridding method, and for our example, we set \(\delta_\theta\) and \(\theta\) is from 1 to 10; \(\delta_{T_f}\) as 0.1 and \(t_{f_1}, t_{f_2}\) is from 1 to 6.0.

**Figure 4.9:** Prior distribution of \(\theta\) (left) and marginal distribution of \(T_{f_1}\) (right) by the gridding method.

**Figure 4.10** and **Figure 4.11** show probabilities of \(T_{f_2}\) conditional on \(T_{f_1}\): depending on various \(t_{f_1}\) from 0.1 to 6.0, it can be seen that the modes of \(p_{T_{f_2}}(t_{f_2} \mid t_{f_1})\) move from left to right and reversely after reaching a certain threshold node, which demonstrates the dynamics of conditional probabilities.
Figure 4.10: Probability of $T_{f2}$ conditioning on $t_{f1}$, i.e., $p_{T_{f2}}(t_{f2} | t_{f1} = i)$, where $i = 0.1, \ldots, 6$, by the gridding method.
Figure 4.11: Comparison of $p_{Tf_1}(t_{f_1})$ (red) and $p_{Tf_2}(t_{f_2} \mid t_{f_1} = i)$ (green), where $i = 0.1, 0.5, 1.0, 1.1, 1.5, 2.0, 2.1, 2.5, 3.0, 3.1, 3.5, 4.0, 4.1, 4.5, 5.0, 5.1, 5.5, 5.9$.

To find the threshold node, a three-dimensional graph is plotted as Figure 4.12.
Figure 4.13 is the projection, in which numbers alongside the dots are $T_{f_2}$ which produce the maximum probability conditioning on corresponding $T_{f_1}$. From Figure 4.12 and Figure 4.13, we can see that the node is at when $p_{T_{f_2}}(t_{f_2} = 0.9 | t_{f_1} = 1)$.

**Figure 4.12**: 3D scatter plot for maximal probability of $T_{f_2}$ conditioning on $t_{f_1}$, i.e., $\max p_{T_{f_2}}(t_{f_2} | t_{f_1})$, with vertical lines for each point.

**Figure 4.13**: Maximal probability of $T_{f_2}$ conditioning on $t_{f_1}$, i.e., $\max p_{T_{f_2}}(t_{f_2} | t_{f_1})$. 
Figure 4.14 presents $T_f^2$ that has maximal probability conditioning on $t_f^1$, i.e.,
\[
\arg \max_{t_f^2} p_{T_f^2}(t_f^2 \mid t_f^1);
\]
the corresponding probabilities (rounded to 2 decimals) are shown alongside dots that represent $T_f^2$ which maximise the probabilities given a specific $t_f^1$. It can be seen that a few $t_f^2$ have the same maximal probability conditional on various $t_f^1$, which is because our gridding methods have limited precision. While Figure 4.15 shows the $t_f^2$ that has maximal probability and its corresponding probability value, for example, for varying $T_f^1$, the maximum conditional probability is about 0.08 when $T_f^2 = 4$.

**Figure 4.14:** $T_f^2$ that has maximal probability conditioning on $t_f^1$, i.e.,
\[
\arg \max_{t_f^2} p_{T_f^2}(t_f^2 \mid t_f^1);
\]
the corresponding probabilities (rounded to 2 decimals) shown alongside dots that represent $T_f^2$ which maximise the probabilities given a specific $t_f^1$. 
Figure 4.15: Maximal probability of $T_{f2}$ and the corresponding $T_f$; corresponding $t_{f1}$ shown alongside dots.

Figure 4.16 and Figure 4.17 illustrate the dynamics of $p_{T_{f2}} (t_{f2} | t_{f1} > T_{m1})$, which has similar property compared with that of $p_{T_{f2}} (t_{f2} | t_{f1})$. 
Figure 4.16: Probability of $T_{f_2}$ conditioning on $t_{f_1} > T_{m_1}$, i.e., $p_{T_{f_2}}(t_{f_2} | t_{f_1} > T_{m_1})$, where $T_{m_1} = 0.1, \ldots, 6$, by the gridding method.
Figure 4.17: Comparison of \( p_{T^{f_1}}(t_{f_1}) \) (red) and \( p_{T^{f_2}}(t_{f_2} \mid t_{f_1} > T_{m_1}) \) (green), where \( T_{m_1} = 0.1, 0.5, 1.0, 1.1, 1.5, 2.0, 2.1, 2.5, 3.0, 3.1, 3.5, 4.0, 4.1, 4.5, 5.0, 5.1, 5.5, 5.9 \).
In summary, these graphs demonstrate the dynamic property of conditional probabilities used to calculate corresponding cost rate and utility in Figure 4.1. However, due to the discrete property of the gridding method, only approximate presentation can be shown.

4.3.5 Pseudocode

Here a general pseudocode for the gridding method is presented as follows, regardless of failure time distribution assumptions.

1. Pre-defined variables

   (a) Give costs of failure $C_f$, repair $C_r$, preventive maintenance $C_m$ and increment $\delta$.

   (b) Generate a sequence of possible values for parameter $\theta$ of a distribution $D_\theta$ from $\theta_{\text{min}}$ to $\theta_{\text{max}}$ by increment $\delta$.

   (c) Generate discrete probability mass of $\theta$ from an arbitrary distribution $D_\theta$.

   (d) Generate a sequence of possible failure time values of $T_{f1}$, $T_{f2}$ and maintenance time values $T_{m1}$, $T_{m2}$ and define the lengths of them as $l_{T_{f1}}$, $l_{T_{f2}}$, $l_{T_{m1}}$, $l_{T_{m2}}$.

2. Compute $p(t_{f2} | t_{f1})$

   (a) Define a matrix $M^{l_{T_{f1}} \times l_{T_{f2}}}$ for the joint mass distribution of $T_{f1}$ and $T_{f2}$, where rows are possible $T_{f1}$ and columns are possible $T_{f2}$.

   (b) Calculate the joint probability of $T_{f1}$ and $T_{f2}$, i.e., $p(t_{f1}, t_{f2})$, by generating failure probability from an arbitrary distribution $D_f$ and integrating over parameter $\theta$.

   (c) Sum the rows the matrix $p(t_{f1}, t_{f2})$ which is the marginal probability of $T_{f1}$, i.e., $p(t_{f1})$.

   (d) Obtain the conditional probability matrix $p(t_{f2} | t_{f1})$ by $\frac{p(t_{f1}, t_{f2})}{p(t_{f1})}$.

3. Compute $p(t_{f2} > T_{m2} | t_{f1})$

   (a) Define a matrix $p(t_{f2} > T_{m2} | t_{f1})$ with 0 entries having $l_{T_{f1}}$ rows ($i$) and $l_{T_{m2}}$ columns ($j$).
(b) For each row, update the entries by sum of the columns of \( p(t_{f_2} \mid t_{f_1}) \) from \( j + 1 \) to \( l_{T_{m_2}} \) and obtain the joint probability of \( t_{f_2} > T_{m_2} \) and \( t_{f_1} \), i.e., \( p(t_{f_2} > T_{m_2}, t_{f_1}) \).

(c) Obtain the conditional probability matrix \( p(t_{f_2} > T_{m_2} \mid t_{f_1}) \) by \( \frac{p(t_{f_2} > T_{m_2}, t_{f_1})}{p(t_{f_1})} \).

4. Compute \( p(t_{f_2} \mid t_{f_1} > T_{m_1}) \)

(a) Define a vector representing \( p(t_{f_1} > T_{m_1}) \), of which the length is \( l_{T_{m_1}} \).

(b) Define a matrix for the joint probability of \( t_{f_1} T_{m_1} \) and \( t_{f_2} \), i.e., \( p(t_{f_1} > T_{m_1}, t_{f_2}) \), of which rows are possible \( T_{m_1} \) and columns are possible \( T_{f_2} \).

(c) For a given \( T_{m_1} \), sum \( p(t_{f_1}) \) of which \( t_{f_1} > T_{m_1} \), which gives \( p(t_{f_1} > T_{m_1}) \).

(d) For a given \( T_{m_1} \), sum \( p(t_{f_1}, t_{f_2}) \) of which \( t_{f_1} > T_{m_1} \) and corresponding \( t_{f_2} \), which gives \( p(t_{f_2}, t_{f_1} > T_{m_1}) \).

(e) Obtain the conditional probability matrix \( p(t_{f_2} \mid t_{f_1} > T_{m_1}) \) by \( \frac{p(t_{f_2}, t_{f_1} > T_{m_1})}{p(t_{f_1} > T_{m_1})} \).

5. Compute \( p(t_{f_2} > T_{m_2} \mid t_{f_1} > T_{m_1}) \)

(a) Define a matrix \( p(t_{f_2} > T_{m_2} \mid t_{f_1}) \) with 0 entries having \( (l_{T_{m_1}} - 1) \) rows \( (i) \) and \( l_{T_{m_2}} \) columns \( (j) \).

(b) For each row, update the entries by sum of the columns of \( p(t_{f_2} \mid t_{f_1} > T_{m_1}) \) from \( j + 1 \) to \( l_{T_{m_2}} \) and obtain \( p(t_{f_2} > T_{m_2}, t_{f_1} > T_{m_1}) \).

(c) Obtain the conditional matrix \( p(t_{f_2} > T_{m_2} \mid t_{f_1} > T_{m_1}) \) by \( \frac{p(t_{f_2} > T_{m_2}, t_{f_1} > T_{m_1})}{p(t_{f_1} > T_{m_1})} \).

6. Calculate the expected utility of cost rate (the expected cost per unit time) at chance node \( CN_{21} \) and \( CN_{22} \), i.e., \( E_{CN_{21}}(U(CR_{T_{m_2}})) \) and \( E_{CN_{22}}(U(CR_{T_{m_2}})) \)

(a) \( E_{CN_{21}}^{Upper} \) is a utility matrix related to \( t_{f_1} \) and \( t_{f_2} \leq T_{m_2} \), with \( l_{T_{f_1}} \) rows and \( l_{T_{m_2}} \) columns.

(b) \( E_{CN_{21}}^{Lower} \) is a utility matrix related to \( t_{f_1} \) and \( t_{f_2} > T_{m_2} \), with \( l_{T_{f_1}} \) rows and \( l_{T_{m_2}} \) columns.

(c) For each row of matrix \( E_{CN_{21}}^{Upper} \) (i.e., given a possible \( T_{f_1} \)), each entry is the utility given a possible \( T_{f_2} \), which gives \( \frac{C_r + C_u}{t_{f_1} + t_{f_2}} \times p(t_{f_2} \mid t_{f_1}) \).

(d) For each row of matrix \( E_{CN_{21}}^{Lower} \) (i.e., given a possible \( T_{f_1} \)), each entry is the utility given a possible \( T_{m_2} \), which gives \( \frac{C_r + C_u + C_m}{t_{f_1} + T_{m_2}} \times p(t_{f_2} > T_{m_2} \mid t_{f_1}) \).
(e) Sum the two matrices above, i.e., $E_{CN_{21}}^{Upper}$ and $E_{CN_{21}}^{Lower}$, which gives the expected utility of expected cost per unit time at chance node $CN_{21}$, i.e., $E_{CN_{21}}(U(CR_{Tm_2}))$.

(f) Apply the same process to $E_{CN_{22}}^{Upper}$ and $E_{CN_{22}}^{Lower}$ with different cost structures and conditional probability matrices to obtain $E_{CN_{22}}(U(CR_{Tm_2}))$.

7. Calculate the expected cost per unit time at chance node $CN_1$, i.e., $E_{CN_1}^{T_{f_1}}$.

(a) For each row of matrix $E_{CN_{21}}(U(CR_{Tm_2}))$, find the maximum utility and corresponding $T_{m_2}$, which gives the optimal $T_{m_2}$ that produces maximum utility, given a $T_{f_1}$.

(b) For each row of matrix $E_{CN_{22}}(U(CR_{Tm_2}))$, find the maximum utility and corresponding $T_{m_2}$, which gives the optimal $T_{m_2}$ that produces maximum utility, given a $T_{m_1}$.

(c) Sum the maximum utilities above pairwisely with their corresponding probabilities, i.e., $p(t_{f_1})$ and $p(t_{f_1} > T_{m_1})$, which gives the $E_{CN_1}(U(CR_{Tm_1}))$.

(d) Find the optimal maintenance time $T_{m_1}$ that produces maximum utility from $E_{CN_1}(U(CR_{Tm_1}))$.

4.4 Numerical Examples

The results and related simulations are shown in this section according to previous models proposed. Maintenance time discussed herein is “local time” (Definition 2.2).

4.4.1 PPM for Two-Phase System

For a two-phase system, we assume the failure cost as $C_f = 2$, repair cost as $C_r = 1$ and maintenance cost as $C_m = 0.5$, which will also apply to other examples.

To compare the results obtained through the dynamic programming framework, the results under a myopic framework are presented as well. Within the myopic framework, the decision maker optimises the maintenance time at the current state, but does not explicitly use forecasting information or any direct representation of decisions in the future. As a result the calculation is expected to be less time consuming than that.
within the dynamic programming framework. However, its solution would not be globally optimal since it ignores possible learning.

From Figure 4.18 we are able to see that the expected utility at chance node $CN_{21}$ is decreasing due to the branch assumption of $t_{f_1} \leq T_{m_1}$ indicating more cost induced, while the expected utility at chance node $CN_{22}$ is increasing thanks to the branch assumption of $t_{f_1} > T_{m_1}$. By looking for the maximum value of the expected utility at chance node $CN_1$, we are able to find the corresponding maintenance time, which is the optimal maintenance time for $CN_1$, $T_{m_1}$.

![Figure 4.18](image)

**Figure 4.18**: Expected utilities for two-phase systems at chance nodes $CN_{21}$, $CN_{22}$ and $CN_1$ under the dynamic programming method.

From Table 4.1 we can see that the optimal maintenance time $T_{m_1}$ at chance node $CN_1$, by the dynamic programming method is 0.5, which is larger than that by the myopic method that is 0.3, whereas the expected utility is higher, which can be explained as by considering all possible future decisions and previous information, and as a result, one can reduce the cost spent on system maintenance, which in other words improves its utility. When it comes to the second phase, if failure is not observed before maintenance at $T_{m_1}$ at local time 0.5 for the first phase, then maintenance time for the second phase is supposed to be at local time 0.3, which can be explained as: early maintenance decisions can be riskier so as to gain information whilst later decisions (here the last decisions) have no or little use to gain extra information because the
system considered here has two phases only.

<table>
<thead>
<tr>
<th>Maintenance Time</th>
<th>Dynamic Programming Expect Utility</th>
<th>Myopic Maintenance Time</th>
<th>Expected Utility</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_{m_1}$</td>
<td>$ECR_{CN_{11}}$</td>
<td>$T_{m_1}$</td>
<td>$ECR_{CN_{11}}$</td>
</tr>
<tr>
<td>0.5</td>
<td>-2.56</td>
<td>0.3</td>
<td>-3.23</td>
</tr>
</tbody>
</table>

$$T_{m_2}(T_{f_1} \leq T_{m_1}) \quad ECR_{CN_{21}} \quad T_{m_2}(T_{f_1} \leq T_{m_1}) \quad ECR_{CN_{21}}$$

<table>
<thead>
<tr>
<th></th>
<th>$ECR_{CN_{21}}$</th>
<th>$ECR_{CN_{21}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9(0.1)</td>
<td>-8.75</td>
<td>0.9(0.1)</td>
</tr>
<tr>
<td>0.8(0.2)</td>
<td>-6.86</td>
<td>0.8(0.2)</td>
</tr>
<tr>
<td>0.7(0.3)</td>
<td>-5.73</td>
<td>0.7(0.3)</td>
</tr>
<tr>
<td>0.7(0.4)</td>
<td>-4.95</td>
<td>-</td>
</tr>
<tr>
<td>0.6(0.5)</td>
<td>-4.37</td>
<td>-</td>
</tr>
</tbody>
</table>

$$T_{m_2}(T_{f_1} > T_{m_1}) \quad ECR_{CN_{22}} \quad T_{m_2}(T_{f_1} > T_{m_1}) \quad ECR_{CN_{22}}$$

<table>
<thead>
<tr>
<th></th>
<th>$ECR_{CN_{22}}$</th>
<th>$ECR_{CN_{22}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.3</td>
<td>-1.61</td>
<td>0.4</td>
</tr>
</tbody>
</table>

**Table 4.1:** Optimal corrective maintenance (CM) time and corresponding expected cost for each chance node based on dynamic programming and myopic methods. Bracketed figures are failure time $T_{f_1}$ with respect to $T_{m_1}^*$ and $T_{m_2}^*$, numbers in brackets representing corresponding failure times.

As we can see in Figure 4.19 and Table 4.2, for very low risk aversion, in other words, when the risk aversion parameter $\eta \to 0$, exponential utility function will give the same maintenance policy, i.e., same maintenance time by maximising expected utility of cost rate, which tends towards minimising the expected cost per unit time or cost rate.
Figure 4.19: Expected costs for two-phase systems at chance nodes $CN_{21}$, $CN_{22}$ and $CN_1$ under the dynamic programming method.

<table>
<thead>
<tr>
<th>Maintenance Time</th>
<th>Expected Cost Rate</th>
<th>Expected Utility $\eta = 0.001$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T^*_m$</td>
<td>$ECR_{CN_1}$</td>
<td>$T^<em><em>m$, $E</em>{T^</em><em>m} U(CR</em>{CN_1})$</td>
</tr>
<tr>
<td>0.5</td>
<td>2.55</td>
<td>0.5, -2.56</td>
</tr>
<tr>
<td>$T_m (T_f \leq T_m)$</td>
<td>$ECR_{CN_{21}}$</td>
<td>$T_m (T_f \leq T_m)$, $E_{T_m} U(CR_{CN_{21}})$</td>
</tr>
<tr>
<td>0.9(0.1)</td>
<td>8.69</td>
<td>0.9(0.1), -8.75</td>
</tr>
<tr>
<td>0.8(0.2)</td>
<td>6.83</td>
<td>0.8(0.2), -6.86</td>
</tr>
<tr>
<td>0.7(0.3)</td>
<td>5.71</td>
<td>0.7(0.3), -5.73</td>
</tr>
<tr>
<td>0.7(0.4)</td>
<td>4.93</td>
<td>0.7(0.4), -4.95</td>
</tr>
<tr>
<td>0.6(0.5)</td>
<td>4.36</td>
<td>0.6(0.5), -4.37</td>
</tr>
<tr>
<td>$T_m (T_f &gt; T_m)$</td>
<td>$ECR_{CN_{22}}$</td>
<td>$T^<em><em>m (T_f &gt; T_m)$, $E</em>{T^</em><em>m} U(CR</em>{CN_{22}})$</td>
</tr>
<tr>
<td>0.3</td>
<td>1.61</td>
<td>0.3, -1.61</td>
</tr>
</tbody>
</table>

Table 4.2: Optimum Perfect Preventive Maintenance (PPM) time through different optimisation objectives, i.e., expected cost and expected utility for each chance node based on dynamic programming method. Bracketed figures are failure time $T_f$ with respect to $T_m$ and $T_{m2}$, numbers in brackets representing corresponding failure times.

To investigate the impact of risk aversion parameter $\eta$ on the decision making about
the optimal maintenance time, $\eta$ is increased gradually to compare the corresponding optimal maintenance time and expected utility for chance node $CN_1$ of the two-phase repairable system. As it is shown in Table 4.3, with the decision maker becoming more risk averse (i.e., larger $\eta$), one is supposed to preventively maintain the system earlier and the expected utility decreases accordingly. Figure 4.20 shows the characteristic of decreasing, in which the expected utility is transformed via an exponential function $\exp(-\cdot)$ to be presented in the right graph.

<table>
<thead>
<tr>
<th>Risk Aversion Parameter $\eta$</th>
<th>Maintenance Time</th>
<th>Expected Utility</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.001</td>
<td>0.5</td>
<td>-2.56</td>
</tr>
<tr>
<td>0.100</td>
<td>0.4</td>
<td>-3.61</td>
</tr>
<tr>
<td>0.200</td>
<td>0.4</td>
<td>-7.71</td>
</tr>
<tr>
<td>0.300</td>
<td>0.4</td>
<td>-43.49</td>
</tr>
<tr>
<td>0.400</td>
<td>0.4</td>
<td>-522.29</td>
</tr>
<tr>
<td>0.500</td>
<td>0.3</td>
<td>-7977.54</td>
</tr>
<tr>
<td>0.600</td>
<td>0.3</td>
<td>-131705.40</td>
</tr>
<tr>
<td>0.700</td>
<td>0.2</td>
<td>-2257595.00</td>
</tr>
<tr>
<td>0.800</td>
<td>0.2</td>
<td>-39617496.00</td>
</tr>
<tr>
<td>0.900</td>
<td>0.2</td>
<td>-706949065.00</td>
</tr>
</tbody>
</table>

**Table 4.3**: Optimal perfect preventive maintenance (PPM) time and corresponding expected utility for chance node $CN_1$ by dynamic programming conditioning on various risk aversion parameter $\eta$ of an exponential utility function.

**Figure 4.20**: Risk aversion parameter $\eta$ on utility.

According to the results obtained via the dynamic programming method, one can
determine the optimal maintenance times for each phase of the two phase system by referring to Figure 4.21. For example, one determines the maintenance time for phase 1 as 0.5; if a failure occurs at time 0.2, the maintenance time for phase 2 is chosen as 0.8 at local time, \(i.e.,\ 1.0(0.2 + 0.8)\) at global time; if no failure happens before 0.5, the maintenance for phase 2 is decided as 0.3 at local time, \(i.e.,\ 0.8(0.5 + 0.3)\) at global time.

Figure 4.21: Optimal Maintenance Decision Tree for Two-Phase Systems.

Figure 4.22 shows the prior and posterior probability distributions of parameter \(\theta\) depending on the observed failure \(T_{f1}\) and conducted preventive maintenance at \(T_{m1} = 0.5\) and indicates that the posterior probability distribution of \(\theta\) is more right-skewed and has higher mode if failure occurs earlier in the first phase.
Figure 4.22: Posterior probability density of $\theta$ conditioning on $T_{f1}(\leq 0.5)$ and $T_{f1} > 0.5$ compared with prior probability density of $\theta$.

4.4.2 Simulation

A simulation is an experiment performed on a model. In this context, the experiment is understood as varying parameters or changing the setup of the model. While the formal modelling of systems using mathematics tries to find analytical solutions, computer-based simulation can provide solutions to problems which are beyond the limits of analytical approaches. Computer-based simulation uses approximation techniques (e.g. numerical integration) to find results to problems which have no analytical solution, can combine discrete and continuous system characteristics, or interact with the simulator. The simulation results in this thesis coordinate with the theoretical ones obtained by the gridding method.

By the law of large numbers, with the number $N$ increasing in simulations, the average of the simulation results from a large number of simulations should be close to the expected cost rate for chance node $CN_1$, $E CR_{CN_1}$, obtained by dynamic programming.
$$ECR_{CN_1}: \mu_{CR}$$

<table>
<thead>
<tr>
<th>$N$</th>
<th>Simulated Expected Cost Rate $\bar{SE}$</th>
<th>Variance $\sigma^2 = E[(\bar{SE} - \mu_{CR})^2]$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1000</td>
<td>2.430923</td>
<td>1.53014e-05</td>
</tr>
<tr>
<td>5000</td>
<td>2.596228</td>
<td>3.262069e-07</td>
</tr>
<tr>
<td>20000</td>
<td>2.517805</td>
<td>6.777306e-08</td>
</tr>
<tr>
<td>50000</td>
<td>2.540612</td>
<td>3.925825e-09</td>
</tr>
<tr>
<td>100000</td>
<td>2.548292</td>
<td>4.006983e-10</td>
</tr>
</tbody>
</table>

Table 4.4: Two-Phase Maintenance System Simulation Results.

As can be seen in Table 4.4, the variance of the simulated expected cost rate becomes smaller with the increasing number of simulations, $N$. In other words, the expected cost rate can be approximated via a large number of simulations.

Related R code is also attached as follows.
n <- 1000  # number of theta
sim.theta <- rtruncnorm(n, a=1, b=Inf, mean=2, sd=1)

# generate tf1 and tf2 which follow Weibull distribution given shape parameter theta
tf1 <- rep(0, rep=n)
tf2 <- rep(0, rep=n)

k <- 1
while (k<=n) {
  x <- rweibull(1, shape=sim.theta[k], scale=1)
y <- rweibull(1, shape=sim.theta[k], scale=1)
  # x <- rgamma(1, shape=sim.theta[k], scale=1)
  # y <- rgamma(1, shape=sim.theta[k], scale=1)
  if ((round(x, 1) != 0) && (round(y, 1) != 0)) {
    tf1[k] <- x
    tf2[k] <- y
    k=k+1
  }
}

# compute simulated cost for chance node CN1
cost.cn <- rep(0, n)
for (i in 1:n) {
  if (round(tf1[i], 1) <= cn1.tm1.opt) {
    if (round(tf2[i], 1) <= cn21.tm2.opt[round(tf1[i], 1)*10]) {
      cost.cn[i] <- (1+alpha)*(Cf+Cr)/(round(tf1[i], 1)+round(tf2[i], 1))
    } else if (round(tf2[i], 1) > cn21.tm2.opt[round(tf1[i], 1)*10]) {
      cost.cn[i] <- (Cf+Cr+alpha*Cm)/(round(tf1[i], 1)+cn21.tm2.opt[round(tf1[i], 1)*10])
    }
  } else if (round(tf1[i], 1) > cn1.tm1.opt) {
    if (round(tf2[i], 1) <= cn22.tm2.opt) {
      cost.cn[i] <- (Cm+alpha*(Cf+Cr))/(cn1.tm1.opt+round(tf2[i], 1))
    } else if (round(tf2[i], 1) > cn22.tm2.opt) {
      cost.cn[i] <- (1+alpha)*Cm/(cn1.tm1.opt+cn22.tm2.opt)
    }
  }
}

# compute simulated expected cost for chance node CN1
mean(cost.cn)
Chapter 5

Sequential Maintenance Extension

Possible extensions to the sequential preventive maintenance models in Chapter 4 are discussed in this chapter as follows: imperfect preventive maintenance, preventive maintenance modelling with the time value of money, modelling preventive maintenance in a discrete time setting as well as maintenance for parallel maintenance, analysing the effect of failure time distribution assumptions on preventive maintenance time and proposing an adaptive approach to solving multi-phase systems’ maintenance. Sensitivity analysis via the parameters of sequential preventive maintenance models will also be carried out in this chapter.

5.1 Imperfect Maintenance

As stated previously in Chapter 2, both perfect preventive maintenance and imperfect preventive maintenance have been implemented and discussed. By definitions and assumptions, after perfect preventive maintenance, the system comes to a brand new state, of which the failure time distribution would be the same as it was initially run. When it comes to imperfect preventive maintenance, the system would be maintained to some level which is not as good as a brand new system, but still is able to keep the system running on an acceptable range and level. So it is reasonable to assume that the rate of failure time distribution would increase after imperfect preventive maintenance, in other words, systems would deteriorate if imperfect preventive maintenance were

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1Extensions to the sequential preventive maintenance models in this chapter are based on two-phase systems and results are presented with regard to maintenance time $T_{m_1}$, unless explicitly stated otherwise. $T_{m_2}$ can be obtained by referring to Figure 4.21.
conducted, i.e., $p_{T_{f_2}}(t_{f_2} | t_{f_1} > T_{m_1}) > p_{T_{f_2}}(t_{f_2} | t_{f_1})$.

According to a review by Nakagawa (2012), existing imperfect preventive maintenance models can be categorised as: age reduction models, hazard rate reduction models, and hybrid models of both. Via age reduction models, the virtual age of a system would reduce to $t - \delta$ from $t$ after imperfect preventive maintenance, i.e., the hazard function would change from $h(t)$ to $h(t - \delta)$, whereas the hazard function would change from $h(t)$ to $\tau h(t)$ ($\tau > 1$) after imperfect preventive maintenance by hazard rate reduction modelling. And hybrid method combines them to model the hazard rate from $h(t)$ to $\tau h(t - \delta)$. A novel method by manipulating the probability matrix $p_{T_{f_2}}(t_{f_2} | t_{f_1} > T_{m_1})$ is proposed in this section.

Based on the original conditional failure probability matrix $p_{T_{f_2}}(t_{f_2} | t_{f_1})$ by the gridding method, the conditional failure time matrix $p_{T_{f_2}}(t_{f_2} | t_{f_1} > T_{m_1})$ for perfect preventive maintenance $M$ is obtained by (§4.3.5 page 82). Then imperfect preventive maintenance probability manipulation is conducted based on this matrix: for each row, take the last arbitrary $\beta$ percent values and reassign them proportionally to each element left, a new matrix $M^{\text{new}}$ is created, which can be regarded as a way of expressing the assumption that the failure rate increases after maintenance. For simplicity, the matrix $M$ with entry $M_{ij} = p_{T_{f_2}}(T_{f_2} = t'_{f_2} | t_{f_1} > t_{m_1})$ is expressed as an example as follows

$$M = \begin{bmatrix} 0.943 & 0.048 & 0.006 & 0.003 \\ 0.800 & 0.151 & 0.038 & 0.011 \\ 0.653 & 0.232 & 0.084 & 0.031 \\ 0.644 & 0.237 & 0.087 & 0.032 \end{bmatrix}$$

where $t'_{f_2}$ is the $j^{th}$ possible value of $T_{f_2}$ and $t_{m_1}^i$ is the $i^{th}$ possible value of $T_{m_1}$; and the sum of each row is 1.

Now, for each row, the last entries 0.003, 0.011, 0.031, 0.032 are replaced with 0 and added proportionally to the other entries for each row. For example, for the first row, $0.943 + 0.003 \times \frac{0.943}{0.943+0.048+0.006}$, $0.048 + 0.003 \times \frac{0.048}{0.943+0.048+0.006}$ and $0.173 + 0.003 \times$
which becomes the first row of matrix $M^{\text{new}}$,

$$
M^{\text{new}} = \begin{bmatrix}
0.9458 & 0.0481 & 0.0060 & 0.0000 \\
0.8089 & 0.1527 & 0.0384 & 0.0000 \\
0.6739 & 0.2394 & 0.0867 & 0.0000 \\
0.6653 & 0.2448 & 0.0899 & 0.0000
\end{bmatrix}
$$

and the sum of each row for matrix $M^{\text{new}}$ is still 1.

Generally, by manipulating the conditional probability matrix $M \, p_{T_{f_2} \mid t_{f_1} > T_{m_1}}$, one can take the last $\beta$ amount of probability values for each row of $M$ and reallocate them to other entries of $M$ for each row proportionally according to their original values in $M$. This can be regarded as an alternative method to deal with conditional probability in imperfect preventive maintenance. We can see from Figure 5.1 that the system is more likely to fail in the second phase if imperfect preventive maintenance is carried out than that under perfect preventive maintenance policy because the system is not as good as new that can only be achieved by perfect preventive maintenance.

![Figure 5.1](image-url)

**Figure 5.1**: Comparison of probabilities of $T_{f_2}$ conditioning on optimal $T_{m_1}$ obtained through PPM and IPM.
As we can see in Figure 5.2 through the manipulation of the conditional probabilities, the system is more likely to fail early at the second phase if it is imperfectly preventively maintained at $T_{m1}$ as 0.5.

![Figure 5.2](image)

**Figure 5.2**: Comparison of probabilities of $T_{f2}$ conditioning on observed $T_{f1}$ and implemented imperfect preventive maintenance at $T_{m1}$.

Table 5.1 presents the comparison of optimal maintenance time by dynamic programming and myopic methods through imperfect preventive maintenance modelling via various preventive maintenance power $\beta$. Imperfect preventive maintenance at power $\beta = 0$ is equivalent to perfect preventive maintenance. From Table 5.1, we can see that expected cost rates increase with respect to the increasing PM power $\beta$ because the system is more likely to fail under imperfect preventive maintenance compared with perfect preventive maintenance. It may be generally concluded that under the imperfect preventive maintenance modelling policy systems tend to be maintained later ($T_{m1} = 0.6$ when $\beta = 55$ vs $T_{m1} = 0.5$ when $\beta = 0$) while expected cost per unit time is increasing due to the system having the high possibility to fail early and being less reliable. More interesting is that one is supposed to maintain systems even
later because once maintained the system is less reliable under the imperfect preventive maintenance policy. For example, the expected cost rate increases 11.65% under the imperfect preventive maintenance policy with PM power $\beta = 55$ compared with its counterpart under the perfect preventive maintenance policy which is when $\beta = 0$.

<table>
<thead>
<tr>
<th>PM Power $\beta$</th>
<th>Chance Node</th>
<th>Dynamic Programming</th>
<th>Myopic</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Maintenance Time</td>
<td>Expected Cost Rate</td>
</tr>
<tr>
<td>0</td>
<td>$CN_1$</td>
<td>0.5</td>
<td>2.55</td>
</tr>
<tr>
<td></td>
<td>$CN_{22}$</td>
<td>0.3</td>
<td>1.61</td>
</tr>
<tr>
<td>10</td>
<td>$CN_1$</td>
<td>0.5</td>
<td>2.57</td>
</tr>
<tr>
<td></td>
<td>$CN_{22}$</td>
<td>0.3</td>
<td>1.64</td>
</tr>
<tr>
<td>20</td>
<td>$CN_1$</td>
<td>0.5</td>
<td>2.57</td>
</tr>
<tr>
<td></td>
<td>$CN_{22}$</td>
<td>0.3</td>
<td>1.64</td>
</tr>
<tr>
<td>30</td>
<td>$CN_1$</td>
<td>0.5</td>
<td>2.57</td>
</tr>
<tr>
<td></td>
<td>$CN_{22}$</td>
<td>0.3</td>
<td>1.64</td>
</tr>
<tr>
<td>40</td>
<td>$CN_1$</td>
<td>0.5</td>
<td>2.58</td>
</tr>
<tr>
<td></td>
<td>$CN_{22}$</td>
<td>0.3</td>
<td>1.64</td>
</tr>
<tr>
<td>50</td>
<td>$CN_1$</td>
<td>0.5</td>
<td>2.67</td>
</tr>
<tr>
<td></td>
<td>$CN_{22}$</td>
<td>0.2</td>
<td>1.77</td>
</tr>
<tr>
<td>55</td>
<td>$CN_1$</td>
<td>0.6</td>
<td>2.85</td>
</tr>
<tr>
<td></td>
<td>$CN_{22}$</td>
<td>0.3</td>
<td>1.78</td>
</tr>
</tbody>
</table>

Table 5.1: Optimal Imperfect Preventive Maintenance (IPM) time and corresponding expected cost for chance nodes $CN_1$ and $CN_{22}$ conditioning on various PM power parameter $\beta$ based on dynamic programming and myopic methods.

Because the gridding method used in this study is an approximate optimisation approach and it is more computationally expensive to apply this method when having smaller gridding intervals, the optimal maintenance times presented in Table 5.1 are almost identical. However, one can expect different optimal maintenance times when the precision of grids increases.
5.2 Time Value of Money

In finance, the net present value (NPV) is defined as the sum of the present values of incoming and outgoing cash flows over a period of time. According to this idea, we can consider the payoff and cost as incoming and outgoing cash flows of the maintenance system, respectively; in other words, a payoff or cost now is more valuable than an identical payoff or cost in the future, which is because of a discount factor.

Discounting of future costs has been widely used in finance, where net present value (NPV) is used as criterion for assessing alternative policies. In our maintenance problem, if a cost induced by a maintenance policy can be delayed for a time $\delta t$, this part of money from the cost budget can be invested for that period of time and earn corresponding interest with rate $r$ or one does need not to borrow that amount of money for a period $\delta t$ with rate $r$. This time value of money effect can be modelled through discounting costs induced by a particular maintenance at time $\delta t$ ahead by a discounting factor $\exp(-r\delta t)$ which is the continuous compounding of interest. So we can re-define the cost rate function as

$$CR(t) = \frac{f(\text{cost to time } t)}{t} = \sum_i \exp(-rt_i)\text{cost}_i$$

(5.1)

where $t_i$ is the time at which cost $i$ occurs, $r$ is the inflation rate or factor for costs and $t$ is the system performing time. Here it is assumed that it becomes less expensive to be maintained or fail in the future and the utility function meets the requirement of risk aversion in this case. In other words, one can replace the previous cost $C_i$ ($i$ is the phase number) in Chapter 4 by $\exp(-rt_i)C_i$, where $t_i$ is the local time that cost $C_i$ happens at.

In the maintenance decision making problem, by implementing a discount factor in utility modelling, the maintenance would be able to depict the reality more accurately in terms of taking time effect into the maintenance modelling procedure.

Given the time effect parameter $r$ varying between 0 and 1, the time factor’s effect on systems’ perfect preventive maintenance optimisation is explored.

$^2r$ is equivalent to the rate of return in finance.
Table 5.2: Optimal Perfect Preventive Maintenance (PPM) time and corresponding expected cost rate for chance node CN₁ conditioning on various time effect parameter $r$ based on dynamic programming. ✓ indicates the cost rate comparison to its previous one.

<table>
<thead>
<tr>
<th>Time Effect Parameter $r$</th>
<th>Maintenance Time</th>
<th>Expected Cost Rate</th>
<th>Cost ✓</th>
</tr>
</thead>
<tbody>
<tr>
<td>$r = 1.0$</td>
<td>0.5</td>
<td>2.55</td>
<td>-</td>
</tr>
<tr>
<td>$r = 0.9$</td>
<td>0.5</td>
<td>2.43</td>
<td>4.96%</td>
</tr>
<tr>
<td>$r = 0.8$</td>
<td>0.4</td>
<td>2.29</td>
<td>5.55%</td>
</tr>
<tr>
<td>$r = 0.7$</td>
<td>0.4</td>
<td>2.15</td>
<td>6.36%</td>
</tr>
<tr>
<td>$r = 0.6$</td>
<td>0.4</td>
<td>2.00</td>
<td>6.89%</td>
</tr>
<tr>
<td>$r = 0.5$</td>
<td>0.4</td>
<td>1.85</td>
<td>7.50%</td>
</tr>
<tr>
<td>$r = 0.4$</td>
<td>0.3</td>
<td>1.69</td>
<td>8.53%</td>
</tr>
<tr>
<td>$r = 0.3$</td>
<td>0.3</td>
<td>1.51</td>
<td>10.71%</td>
</tr>
<tr>
<td>$r = 0.2$</td>
<td>0.3</td>
<td>1.31</td>
<td>13.01%</td>
</tr>
<tr>
<td>$r = 0.1$</td>
<td>0.2</td>
<td>1.07</td>
<td>18.25%</td>
</tr>
</tbody>
</table>

From Table 5.2 we are able to conclude generally that with the time effect more dominating ($r \to 0$), systems tend to be maintained earlier because it would be much more expensive to conduct maintenance in the future, which meets our expectation; we may also notice that the expected cost rate decreases significantly with maintenance actions being conducted earlier. For example, when $r$ changes from 0.9 to 0.8, maintenance time would be scheduled 0.1 time units in advance and the expected cost rate would be decreased by 5.55%; and if $r$ changes from 0.2 to 0.1, although the maintenance time would be also scheduled 0.1 time units earlier, the expected cost rate would be decreased more significantly by 18.25%. These results imply the crucial role that the time value of money effect plays in the maintenance modelling.

Models and results presented in this section suggest that the consideration of NPV is crucial for preventive maintenance optimisation problem in practice, which is because money is usually borrowed from banks to carry out maintenance in practical circumstances. Therefore, a NPV could be achieved by deferring the maintenance cost to the future, which should be taken into account in the maintenance optimisation process.
5.3 Maintenance in Discrete Time

In the survival analysis of repairable systems’ maintenance, the time to failure is not always observed in a continuous time setting. For instance, in practice tyres of fighter aircraft are preventively replaced after about 4 ∼ 14 flights (Nakagawa, 2012). In some situations, the lifetimes of a system are recorded depending on the number of cycles that it is working, so there is not a calendar or clock involved, e.g., the failure time data of a toy manufacturing system may be collected each manufacturing cycle. In other cases, its lifetimes are not defined at the exact clock time but are statistically observed monthly, seasonly, or yearly, for example. Thus it is interesting and worthwhile considering system maintenance in a discrete time setting.

Consider the time over an indefinitely long cycle $n (n = 1, 2, \ldots)$ that a single unit should be operating. A unit is replaced at cycle $N (N = 1, 2, \ldots)$ after its installation or at failure, whichever occurs first. Let $\{P_n\}_{n=1}^{\infty}$ denote the discrete failure distribution that a unit fails at cycle $n$. Cost $(C_f + C_r)$ is incurred for the system that is replaced and cost $C_m (< C_f + C_r)$ is incurred for the non-failed system that is preventively maintained. Then, the expected cost rate for a one phase system is given by

$$C(T_N) = \frac{(C_f + C_r) \sum_{j=1}^{T_N} P_j + C_m \sum_{j=T_N+1}^{\infty} P_j}{\sum_{j=1}^{T_N} \sum_{i=j}^{\infty} P_i} \quad (T_N = 1, 2, \ldots) \quad (5.2)$$

Let $h_n \equiv \frac{P_n}{\sum_{j=n}^{\infty} P_j}$ $(n = 1, 2, \ldots)$ be the hazard rate of the discrete Weibull distribution and $\mu$ be the mean discrete failure time, i.e., $\mu = \sum_{n=1}^{\infty} n P_n < \infty$. Then, the Bayesian dynamic programming method in the continuous time process in chapter 4 can be applied to modelling the optimal preventive maintenance if the failure time is assumed to be discrete.

Assume the failure time of a system follows a discrete Weibull distribution (Nakagawa and Osaki 1975). Khalique (1989) discussed the statistical properties of a few discrete failure time distributions. Formally, $T_f \sim \text{Discrete Weibull}(p, \theta)$ indicates that the random failure time $T_f$ has a discrete Weibull distribution with real parameter $p$ ($0 < p < 1$), and shape parameter $\theta$. Its probability mass function is

$$P_{T_f}(t_f) = (1 - p)^{(t_f)^p} - (1 - p)^{(t_f+1)^p}, \quad t_f = 0, 1, 2, \ldots \quad (5.3)$$

The probability mass function for three different real parameter settings ($p = 0.01, 0.05, 0.1, 0.5, \ \theta = 2$) is illustrated below.
Figure 5.3: Discrete Weibull Probability Mass Functions (top-left: $p = 0.01, \theta = 2$; top-right: $p = 0.1, \theta = 2$; bottom-left: $p = 0.5, \theta = 2$; bottom-right: $p = 0.9, \theta = 2$).

From Figure 5.3, we can see that discrete Weibull failures are more likely to happen for the system if $p$ is smaller.

The cumulative distribution on the support of $T_f$ is
\[
\mathbb{P}(T_f) = \mathbb{P}(T_f \leq t_f) = 1 - (1 - p)^{(t_f+1)^\theta}, \quad t_f = 0, 1, 2, \ldots \quad (5.4)
\]
then the survival function is
\[
S(t_f) = \mathbb{P}(T_f > t_f) = (1 - p)^{(t_f)^\theta}, \quad t_f = 0, 1, 2, \ldots. \quad (5.5)
\]

The hazard function for discrete Weibull failure time is
\[
h(t_f) = \frac{\mathbb{P}(t_f)}{S(t_f)} = 1 - (1 - p)^{(t_f+1)^\theta - (t_f)^\theta}, \quad (5.6)
\]
the first derivative of $h(t_f)$ with respect to $t_f$ is
\[
\frac{\partial h(t_f)}{\partial t_f} = -(1 - p)^{-(t_f)^\theta + (1+t_f)^\theta} \left(-\theta(t_f)^{-1+\theta} + \theta(1 + t_f)^{-1+\theta}\right) \log(1 - p). \quad (5.7)
\]

It is obvious to see that (5.7) $> 0$ if and only if when $\theta > 1$, which represents an increasing hazard function.
We can apply the dynamic programming method to a two phase system following a discrete Weibull failure time distribution with the same assumptions concerning the other parameters in Chapter 4. Corresponding results are shown in Table 5.3.

<table>
<thead>
<tr>
<th>Parameter $p$</th>
<th>Maintenance Time</th>
<th>Expected Cost Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.01</td>
<td>1</td>
<td>0.52</td>
</tr>
<tr>
<td>0.05</td>
<td>1</td>
<td>0.63</td>
</tr>
<tr>
<td>0.09</td>
<td>1</td>
<td>0.73</td>
</tr>
<tr>
<td>0.13</td>
<td>1</td>
<td>0.83</td>
</tr>
<tr>
<td>0.17</td>
<td>1</td>
<td>0.93</td>
</tr>
<tr>
<td>0.21</td>
<td>1</td>
<td>1.03</td>
</tr>
<tr>
<td>0.25</td>
<td>1</td>
<td>1.13</td>
</tr>
<tr>
<td>0.29</td>
<td>1</td>
<td>1.23</td>
</tr>
<tr>
<td>0.33</td>
<td>1</td>
<td>1.33</td>
</tr>
<tr>
<td>0.37</td>
<td>1</td>
<td>1.43</td>
</tr>
<tr>
<td>0.41</td>
<td>1</td>
<td>1.53</td>
</tr>
<tr>
<td>0.45</td>
<td>1</td>
<td>1.63</td>
</tr>
<tr>
<td>0.49</td>
<td>1</td>
<td>1.73</td>
</tr>
</tbody>
</table>

**Table 5.3**: Optimal perfect preventive maintenance (PPM) time and corresponding expected cost for chance node $CN_1$ by dynamic programming conditioning on various parameter $p$ of a discrete Weibull failure distribution.

As we can see in Table 5.3, with the parameter $p$ increasing, which means the system is less likely to fail following a discrete Weibull distribution, the expected cost rate is also increasing despite having the same maintenance time of 1 for all cases. This means that under the same maintenance time schedule, one can reduce the expected cost rate significantly if the system is more likely to have a discrete Weibull failure time. Maintenance time for the second phase can be achieved via the same procedure proposed in Chapter 4.

This section is an elementary exploration to discrete failure probability distribution. Hence, one can consider other discrete distributions in practice.
5.4 Maintenance for Parallel Systems

Systems studied so far have only one unit. Now let us consider a parallel redundant system that consists of \( N (N \geq 2) \) identical units and the system fails when all its units fail. Assume each unit has a failure distribution \( F(t) \) with finite mean \( \mu \).

Suppose that a one phase system is replaced at system failure or at planned time \( T(0 < T < \infty) \), whichever occurs first. Then, we have the expected cost per unit time for a one phase system as

\[
C(T_m; N) = \int_0^{T_m} \frac{C_f + Cr}{t} dF^N(t) + \int_{T_m}^\infty \frac{C_m}{t} dF^N(t)
\]

where \( C_f + Cr \) is the cost of replacement (failure cost and repair cost) at system failure, \( C_m \) is the cost of perfect preventive maintenance at scheduled maintenance time \( T_m \) with \( C_m < C_f + Cr \).

For example, one is supposed to determine the optimal preventive maintenance time for a two unit parallel redundant system with two phases, of which two units are identical and follow a Weibull probability distribution with the shape parameter \( \theta \) from a normal distribution truncated at 1 with mean 2 and standard deviation 1 and scale parameter 1, which are same assumptions presented for two-phase maintenance models in Chapter 4. The decision tree is the same as in Figure 4.1 (page 56).

Based on the gridding method, the expected cost rates for chance nodes \( CN_{21} \) and \( CN_{22} \) are expressed as

\[
\mathbb{E}_{CN_{21}}(CR_{T_{m_2}}; N) = \sum_{T_{f_1}=0}^{T_{m_2}} \left( \frac{2(C_f + Cr)}{t_f_1 + t_f_2} \right) \times \left( p_{T_{f_2}}(t_{f_2} > t_{f_1}) \right)^N
\]

\[
\mathbb{E}_{CN_{22}}(CR_{T_{m_2}}; N) = \sum_{T_{f_1}=0}^{T_{m_2}} \left( \frac{C_m + C_f + Cr}{T_{m_1} + t_f_2} \right) \times \left( p_{T_{f_2}}(t_{f_2} > T_{m_2} > t_{f_1}) \right)^N
\]

where \( N \) is the number of parallel units of systems.
As it can be seen \((5.10)\) and \((5.11)\), the conditional probability matrices for the two unit system are transformed via component wise matrix multiplication. Thus, one can apply the dynamic programming method to solving the optimal preventive maintenance problem for two unit (and even multi-unit simply via matrix multiplication) redundant parallel systems.

<table>
<thead>
<tr>
<th>Maintenance Time</th>
<th>Expected Cost Rate</th>
<th>One-unit System (N = 1)</th>
<th>Maintenance Time</th>
<th>Expected Cost Rate</th>
<th>Parallel System (N = 2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(T_{m_1})</td>
<td>(ECR_{CN_1})</td>
<td>(T_{m_1})</td>
<td>(ECR_{CN_2})</td>
<td>(T_{m_2})</td>
<td></td>
</tr>
<tr>
<td>0.5</td>
<td>2.55</td>
<td>0.5</td>
<td>0.57</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(T_{m_2}(T_{f_1} \leq T_{m_1}))</td>
<td>(ECR_{CN_2})</td>
<td>(T_{m_2}(T_{f_1} \leq T_{m_1}))</td>
<td>(ECR_{CN_2})</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.9(0.1)</td>
<td>8.69</td>
<td>0.8(0.1)</td>
<td>17.64</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.8(0.2)</td>
<td>6.83</td>
<td>0.8(0.2)</td>
<td>10.32</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.7(0.3)</td>
<td>5.71</td>
<td>0.7(0.3)</td>
<td>6.93</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.7(0.4)</td>
<td>4.93</td>
<td>0.7(0.4)</td>
<td>5.04</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.6(0.5)</td>
<td>4.36</td>
<td>0.6(0.5)</td>
<td>3.89</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(T_{m_2}(T_{f_1} &gt; T_{m_1}))</td>
<td>(ECR_{CN_2})</td>
<td>(T_{m_2}(T_{f_1} &gt; T_{m_1}))</td>
<td>(ECR_{CN_2})</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3</td>
<td>1.61</td>
<td>0.5</td>
<td>1.55</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 5.4: Optimal perfect preventive maintenance (PPM) time and corresponding expected cost rate for each chance node by dynamic programming for one-unit systems and two-unit redundant parallel systems. Bracketed figures are failure time \(T_{f_1}\) with respect to \(T_{m_1}\) and \(T_{m_2}\), numbers in brackets representing corresponding failure times.

Table 5.4 shows that the expected cost per unit time for the two unit redundant parallel system is 0.57, which is 77.85% cheaper than that for the one unit system (2.55). Hence, one could consider a two unit redundant system if its implementing cost rate is less than 1.96. In other words, one can take the expected cost of 1.96 as a cut-off to determine if a parallel redundant system should be implemented in this case.
5.5 PPM under failure time distribution assumptions

Here we simply compare the failure distribution assumption’s effect on system perfect preventive maintenance optimisation.

For example, we assume the system failure time follows Weibull and gamma distribution, respectively. These two distributions have the same shape parameter $\theta$ (denoted as $\theta_W$ and $\theta_G$, respectively) that has a normal distribution prior with mean 2 and standard deviation 1 truncated at 1. For a legitimate comparison, it is assumed that the failure times $T_f^W$ and $T_f^G$ from two distributions have the same expectation. Formally,

$$
\begin{align*}
E(T_f^W) &= E(T_f^G), \\
\kappa_W \Gamma \left(1 + \frac{1}{\theta_W} \right) &= \kappa_G \theta_G, \\
E \left( \kappa_W \Gamma \left(1 + \frac{1}{\theta_W} \right) \right) &= E \left( \kappa_G \theta_G \right), \\
\kappa_W E \left( \frac{\Gamma \left(1 + \frac{1}{\theta_W} \right)}{\Gamma \left(1 + \frac{1}{\theta_W} \right)} \right) &= \kappa_G E \left( \frac{\theta_G}{\Gamma \left(1 + \frac{1}{\theta_W} \right)} \right),
\end{align*}
$$

where $\kappa_W$ and $\kappa_G$ are the scale parameters for Weibull and gamma distributions, respectively.

$\kappa_W$ is obtained through the following steps:

1. $E(\theta_G) = 2 + \frac{\varphi(1)}{\Phi(1)} = 2.2876 \, (4.33, \text{page } 66)$;

2. Generate a large number of $N$ random number for $\theta_W$, denoted as $(\theta_W^i)$, $i = 1, \cdots, N$, from a normal distribution truncated at 1 with mean 2 and standard deviation 1 ($\kappa_G = 1$), which is the same distribution of $\theta_G$;

3. Apply gamma function $\Gamma(\cdot)$ to each $\left(1 + \frac{1}{(\theta_W^i)}\right)$, $i = 1, \cdots, N$;

4. Compute the mean of $\Gamma \left(1 + \frac{1}{(\theta_W^i)}\right)$, $i = 1, \cdots, N$, which is 0.8993309 and replaces $E \left( \Gamma \left(1 + \frac{1}{\theta_W} \right) \right)$;

5. $\kappa_W$ is obtained as 2.543669.
Then we are able to apply our gridding method to obtain marginal probability of failure time $T_f$ from Weibull distribution with shape parameter $\theta_W$ and scale parameter $\kappa_W = 2.543669$ and gamma distribution with shape parameter $\theta_G$ and scale parameter $\kappa_G = 1$. As they are shown in Figure 5.4, systems under gamma distribution assumption are more likely to fail earlier than that under Weibull distribution assumption, on the condition of same expected failure time for the two distributions.

![Graph showing Weibull and gamma distributions](image)

**Figure 5.4:** Marginal Weibull and gamma probabilities of $T_f$, $p_W(t_f)$ and $p_G(t_f)$ with same expectation, i.e., $E(T_f^W) = E(T_f^G)$.

We can see the difference in effect by looking at the probability density functions of the two distributions:

$$f^W_{T_f}(t_f \mid \kappa_W, \theta_W) = \frac{\theta_W}{\kappa_W} \left( \frac{t_f}{\kappa_W} \right)^{\theta_W-1} e^{-\left( \frac{t_f}{\kappa_W} \right)^{\theta_W}} \left\{ - \left( \frac{t_f}{\kappa_W} \right)^{\theta_W} \right\}$$

$$\propto (t_f)^{\theta_W-1} \exp \left\{ - \left( \frac{t_f}{\kappa_W} \right)^{\theta_W} \right\}$$

$$f^G_{T_f}(t_f \mid \kappa_G, \theta_G) = \frac{(t_f)^{\theta_G-1}}{(\theta_G)^{\kappa_G} \Gamma(\theta_G)} e^{-\frac{t_f}{\theta_G}} \left\{ - \frac{t_f}{\theta_G} \right\}$$

$$\propto (t_f)^{\theta_G-1} \exp \left\{ - \frac{t_f}{\theta_G} \right\}$$
By ignoring all the normalising constants, we can see that the probability density function of the Weibull distribution drops off much more quickly for shape parameter $\theta_W > 1$ than the gamma distribution. In the case where $\theta_W = \theta_G = 1$, they both reduce to the exponential distribution. More importantly, the hazard increases for the Weibull distribution but tends to be a constant for the gamma distribution.

A comparison of maintenance time and expected cost rate obtained according to different failure distribution assumptions are presented in Table 5.5.

<table>
<thead>
<tr>
<th>Weibull failure time</th>
<th>gamma failure time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maintenance Time</td>
<td>Expected Cost Rate</td>
</tr>
<tr>
<td>$T_{m_1}$</td>
<td>$ECR_{CN_1}$</td>
</tr>
<tr>
<td>1.2</td>
<td>1.00</td>
</tr>
<tr>
<td>$T_{m_2}(T_{f_1} \leq T_{m_1})$</td>
<td>$ECR_{CN_21}$</td>
</tr>
<tr>
<td>2.3(0.1)</td>
<td>4.86</td>
</tr>
<tr>
<td>2.2(0.2)</td>
<td>3.97</td>
</tr>
<tr>
<td>2.1(0.3)</td>
<td>3.44</td>
</tr>
<tr>
<td>2.0(0.4)</td>
<td>3.07</td>
</tr>
<tr>
<td>2.0(0.5)</td>
<td>2.78</td>
</tr>
<tr>
<td>1.9(0.6)</td>
<td>2.56</td>
</tr>
<tr>
<td>1.8(0.7)</td>
<td>2.37</td>
</tr>
<tr>
<td>1.8(0.8)</td>
<td>2.22</td>
</tr>
<tr>
<td>1.8(0.9)</td>
<td>2.08</td>
</tr>
<tr>
<td>1.7(1.0)</td>
<td>1.97</td>
</tr>
<tr>
<td>1.7(1.1)</td>
<td>1.86</td>
</tr>
<tr>
<td>1.6(1.2)</td>
<td>1.77</td>
</tr>
<tr>
<td>$T_{m_2}(T_{f_1} &gt; T_{m_1})$</td>
<td>$ECR_{CN_22}$</td>
</tr>
<tr>
<td>0.8</td>
<td>0.65</td>
</tr>
</tbody>
</table>

Table 5.5: Optimal Perfect Preventive Maintenance (PPM) time and corresponding expected cost rate for each chance node by dynamic programming based on Weibull and gamma failure time assumptions. Bracketed figures are failure time $T_{f_1}$ with respect to $T_{m_1}$ and $T_{m_2}$, numbers in brackets representing corresponding failure times.

From Table 5.5 we can conclude that systems with gamma failure time distribution
tend to be preventively maintained earlier (the maintenance time $T_{m_1} = 1$ under gamma failure time assumption and $T_{m_1} = 1.2$ under Weibull failure time assumption) and about 66.24% more expensive (systems under gamma failure time assumption would induce higher expected cost rate (1.66) than that under Weibull failure time assumption (1.00)). In this sense, it is essential to have appropriate failure time assumptions about the systems studied in practice because failure time assumptions could result in different maintenance policies.

5.6 Hybrid Myopic-Dynamic Programming

Maintenance modelling and optimisation for two-phase systems have been studied so far. In practice, however, one may be faced with more complex systems with multi-phases. Due to the problem of the “curse of dimensionality”, the model becomes much more complicated and the computation time increases dramatically when modelling systems with even more phases and the number of conditional probabilities is $2^n$, where $n$ is the number of phases for a system. To partially solve this problem, a hybrid myopic-dynamic programming method is proposed for multi-phase systems.

Consider a three-phase system with the same setting and assumptions as the two-phase system in §4.1. A decision tree concerning the optimal maintenance times for this system is shown in Figure 5.5. A decision maker has to determine the optimal maintenance times $T_{m_1}$, $T_{m_2}$ and $T_{m_3}$ for each phase before the system starts working.

The hybrid myopic-dynamic programming (H-M-DP) can be briefly explained as follows. First, we apply the results in the form of the optimal maintenance times from the two-phase systems to the first two phases of the three-phase systems; in other words, we solve part DP-1 in Figure 5.5 using the dynamic programming method. Second, conditioning on the optimal $T_{m_1}$ we obtained previously, we solve part DP-2 in Figure 5.5 also by treating the posterior of related parameters (in this case, the posterior of $\theta$) as a new prior and implementing dynamic programming as a function of $T_{f_1}$. 
Figure 5.5: Decision tree for three-phase system with sequential problem with shading indicating a range of possible outcomes for the preceding chance node; Box DP-1 and DP-2 show the break into two period problems.

In order to apply the dynamic programming method to finding the optimal maintenance times for this three-phase system, one is required to have the conditional probabilities rooting from chance nodes $CN_{31}$, $CN_{32}$, $CN_{33}$ and $CN_{34}$, which are eight $(2^3)$ conditional probabilities as below.

1. \[
    f_{T_f}(t_{f_3} | t_{f_2}, t_{f_1}) = \frac{f(t_{f_3}, t_{f_2}, t_{f_1})}{f(t_{f_2}, t_{f_1})} = \frac{\int \theta f(t_{f_3} | t_{f_2}, t_{f_1}, \theta) f(t_{f_2} | t_{f_1}, \theta) f(t_{f_1} | \theta) f(\theta) d\theta}{\int \theta f(t_{f_2} | t_{f_1}, \theta) f(t_{f_1} | \theta) f(\theta) d\theta} = \frac{\int \theta f(t_{f_3} | \theta) f(t_{f_2} | \theta) f(t_{f_1} | \theta) f(\theta) d\theta}{\int \theta f(t_{f_2} | \theta) f(t_{f_1} | \theta) f(\theta) d\theta}
\]

2. \[
    f_{T_f}(t_{f_3} > T_{m_3} | t_{f_2}, t_{f_1}) = \frac{f(t_{f_3} > T_{m_3}, t_{f_2}, t_{f_1})}{f(t_{f_2}, t_{f_1})} = \frac{\int \theta f(t_{f_3} > T_{m_3}, t_{f_2}, t_{f_1}, \theta) f(t_{f_2} | t_{f_1}, \theta) f(t_{f_1} | \theta) f(\theta) d\theta}{\int \theta f(t_{f_2} | t_{f_1}, \theta) f(t_{f_1} | \theta) f(\theta) d\theta}
\]
\[
\begin{align*}
\int_0 f(t_{f_3} > T_{m_3} \mid t_{f_2}, t_{f_1}, \theta) f(t_{f_2} \mid t_{f_1}, \theta) f(t_{f_1} \mid \theta) f(\theta) d\theta &= \\
\int_0 f(t_{f_3} > T_{m_3} \mid \theta) f(t_{f_2} \mid \theta) f(t_{f_1} \mid \theta) f(\theta) d\theta &= \\
\int_0 f(t_{f_3} > T_{m_3} \mid \theta) f(t_{f_2} \mid \theta) f(t_{f_1} \mid \theta) f(\theta) d\theta &= 1
\end{align*}
\]

3. \(f_{T_{f_3}}(t_{f_3} \mid t_{f_2} > T_{m_2}, t_{f_1})\)
\[
\int_0 f(t_{f_3} \mid t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_3} \mid t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_1} \mid \theta) f(\theta) d\theta = \\
\int_0 f(t_{f_3} \mid t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_3} \mid t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_1} \mid \theta) f(\theta) d\theta = 1
\]

4. \(f_{T_{f_3}}(t_{f_3} > T_{m_3} \mid t_{f_2} > T_{m_2}, t_{f_1})\)
\[
\int_0 f(t_{f_3} > T_{m_3} \mid t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_1} \mid \theta) f(\theta) d\theta = \\
\int_0 f(t_{f_3} > T_{m_3} \mid t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_1} \mid \theta) f(\theta) d\theta = 1
\]

5. \(f_{T_{f_3}}(t_{f_3} > T_{m_3} \mid t_{f_2}, t_{f_1} > T_{m_1})\)
\[
\int_0 f(t_{f_3} > T_{m_3} \mid t_{f_2}, t_{f_1} > T_{m_1}) f(t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_1} \mid \theta) f(\theta) d\theta = \\
\int_0 f(t_{f_3} > T_{m_3} \mid t_{f_2}, t_{f_1} > T_{m_1}) f(t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_1} \mid \theta) f(\theta) d\theta = 1
\]

6. \(f_{T_{f_3}}(t_{f_3} > T_{m_3} \mid t_{f_2}, t_{f_1} > T_{m_1})\)
\[
\int_0 f(t_{f_3} > T_{m_3} \mid t_{f_2}, t_{f_1} > T_{m_1}) f(t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_1} \mid \theta) f(\theta) d\theta = \\
\int_0 f(t_{f_3} > T_{m_3} \mid t_{f_2}, t_{f_1} > T_{m_1}) f(t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_1} \mid \theta) f(\theta) d\theta = 1
\]

7. \(f_{T_{f_3}}(t_{f_3} > T_{m_3} \mid t_{f_2} > T_{m_2}, t_{f_1} > T_{m_1})\)
\[
\int_0 f(t_{f_3} > T_{m_3} \mid t_{f_2} > T_{m_2}, t_{f_1} > T_{m_1}) f(t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_1} \mid \theta) f(\theta) d\theta = \\
\int_0 f(t_{f_3} > T_{m_3} \mid t_{f_2} > T_{m_2}, t_{f_1} > T_{m_1}) f(t_{f_2} > T_{m_2}, t_{f_1}) f(t_{f_1} \mid \theta) f(\theta) d\theta = 1
\]

8. \(f_{T_{f_3}}(t_{f_3} > T_{m_3} \mid t_{f_2} > T_{m_2}, t_{f_1} > T_{m_1})\)
\[
= \frac{f(t_f > T_{m_3}, t_f > T_{m_2}, t_f > T_{m_1})}{f(t_f > T_{m_2}, t_f > T_{m_1})} = \int_{\theta} f(t_f > T_{m_3} \mid t_f > T_{m_2}, t_f > T_{m_1}, \theta) f(t_f > T_{m_2} \mid t_f > T_{m_1}, \theta) f(t_f > T_{m_1} \mid \theta) f(\theta) \, d\theta
\]

The conditional probabilities above are with regard to \(T_{f_3}\), which are conditioning on two joint probabilities, e.g., the calculation for \(p_{T_{f_3}}(t_f, t_f, t_f)\). This complex form adds a challenge to computing. As a result, the gridding approach to solving the dynamic programming problem requires the multi-dimensional matrix formalisation.

This combination of the myopic method and dynamic programming can be applied to those expanded problems with multi-phase systems’ optimal maintenance determination.

In the proposed hybrid myopic-dynamic programming method, first we solve part DP-1 by dynamic programming and get \(T_{m_1}\) as 0.5.

Figure 5.6: Comparison of probabilities of \(T_{f_2}\) conditioning on varying \(T_{f_1}\) and optimal \(T_{m_1}\).
Making use of this result, in Figure 5.6 we can see that \( p(t_{f_2} \mid t_{f_1} = i), i = 0.1, \ldots, 0.5 \), is mostly on the right and below \( p(t_{f_1}) \), which tells us that the system is more likely to fail earlier if a system failure is observed.

Let us compare the prior and posterior distribution of parameter \( \theta \). By Bayes’ theorem, the posterior of \( \theta \) can be shown as \( p(\theta \mid t_{f_1} \leq T_{m_1}) \) and \( p(\theta \mid t_{f_1} > T_{m_1}) \):

\[
p(\theta) = \frac{\phi(\theta - 2)}{\Phi(1)} \tag{5.12}
\]

\[
p(\theta \mid t_{f_1} \leq 0.5) = \frac{p(t_{f_1} \leq 0.5 \mid \theta)p(\theta)}{p(t_{f_1} \leq 0.5)} \tag{5.13}
\]

\[
p(\theta \mid t_{f_1} > 0.5) = \frac{p(t_{f_1} > 0.5 \mid \theta)p(\theta)}{p(t_{f_1} > 0.5)} \tag{5.14}
\]

Figure 5.7 shows the prior and posterior distributions of parameter \( \theta \) given the failure time is left censored and right censored, respectively. If a failure occurs before the scheduled maintenance though it is unknown by how much, the posterior of \( \theta \) is updated in the red line indicating a right-skewed property and a higher mode compared to the prior.
Figure 5.7: Posterior probability density of $\theta$ conditioning on $T_{f1} \leq T_{m1}(0.5)$ and $T_{f1} > T_{m1}(0.5)$ compared with prior probability density of $\theta$.

Under the different conditions of failures happening before scheduled maintenance $T_{m1}$ and maintenance is carried out at $T_{m1}$, the distribution of the shape parameter of the Weibull distribution is updated and treated as the prior distribution for the next phase of the system.

Depending on the observed failure times $T_{f1}$ or schedule maintenance at $T_{m1}$ at the first phase, we apply the dynamic programming method to find the optimal maintenance times for chance node $CN_{21}$ and $CN_{22}$ by considering all the possible situations in the third phase.
Figure 5.8: Expected cost rates at chance node $CN_{21}$ conditioning on $t_{f1} \leq T_{m1}$ (top-left: $t_{f1} = 0.1$, top-right: $t_{f1} = 0.2$, middle-left: $t_{f1} = 0.3$, middle-right: $t_{f1} = 0.4$ and bottom-left: $t_{f1} = 0.5$; and expected cost rates at chance node $CN_{22}$ conditioning on $t_{f1} > T_{m1}$ (bottom-right) under the H-DP-M method.

Figure 5.8 shows expected cost rates at chance node $CN_{21}$ conditioning on $t_{f1} \leq T_{m1}$ (top-left: $t_{f1} = 0.1$, top-right: $t_{f1} = 0.2$, middle-left: $t_{f1} = 0.3$, middle-right: $t_{f1} = 0.4$ and bottom-left: $t_{f1} = 0.5$; and expected cost rates at chance node $CN_{22}$ conditioning on $t_{f1} > T_{m1}$ (bottom-right) under the H-DP-M method.

The optimal maintenance times of the three phase system for chance nodes $CN_{21}$ and $CN_{22}$ and corresponding expected cost rates depending what have been observed in the first phase are shown in Table 5.6.
<table>
<thead>
<tr>
<th>Chance Node</th>
<th>History</th>
<th>H-M-DP (Three-Phase)</th>
<th>DP (Two-Phase)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Maintenance Time</td>
<td>Expected Cost Rate</td>
</tr>
<tr>
<td>$CN_{21}(T_{j_1} \leq T_{m_1})$</td>
<td>$T_{j_1} = 0.1$</td>
<td>0.7</td>
<td>6.19</td>
</tr>
<tr>
<td></td>
<td>$T_{j_1} = 0.2$</td>
<td>0.7</td>
<td>5.64</td>
</tr>
<tr>
<td></td>
<td>$T_{j_1} = 0.3$</td>
<td>0.6</td>
<td>5.18</td>
</tr>
<tr>
<td></td>
<td>$T_{j_1} = 0.4$</td>
<td>0.6</td>
<td>4.79</td>
</tr>
<tr>
<td></td>
<td>$T_{j_1} = 0.5$</td>
<td>0.6</td>
<td>4.46</td>
</tr>
<tr>
<td>$CN_{22}(T_{j_1} &gt; T_{m_1})$</td>
<td>$T_{m_1} = 0.5$</td>
<td>0.4</td>
<td>2.37</td>
</tr>
</tbody>
</table>

Table 5.6: Optimal Perfect Preventive Maintenance (PPM) time and corresponding expected cost rate of chance nodes $CN_{21}$ and $CN_{22}$ for three-phase maintenance systems based on Hybrid Myopic-Dynamic Programming and myopic methods.

From Table 5.6, we see that through hybrid myopic-dynamic programming the future information about the system performance is taken into our maintenance modelling, then the maintenance times at $CN_{21}$ are earlier than in the same chance node for the two phase system, which results in lower expected cost rate compared with dynamic programming for the two-phase system.

According to the results obtained via the hybrid myopic-dynamic programming method, one can determine the optimal maintenance times for each phase of the three phase system by referring to Figure 5.9. For example, one determines the maintenance time for phase 1 as 0.5; if a failure occurs at time 0.2, the maintenance time for phase 2 is chosen as 0.7; if no failure happens before 0.7, the maintenance for phase 3 is decided as 0.8.
Figure 5.9: Optimal Maintenance Decision Tree for Three-Phase Systems
5.7 Sensitivity Analysis

The impact of parameter variation on maintenance policies, \textit{i.e.}, sensitivity analysis, is considered and presented in this section.

5.7.1 Gridding Increments

In Chapter 4, the gridding method was implemented to solve the optimisation problem in sequential preventive maintenance models, and the increment parameter was set up as 0.1 as default. This section is intended to analyse and understand the core of the gridding method, \textit{i.e.}, the gridding increments’ impact on decisions concerning the optimal preventive maintenance times and the relationships between them.

First, the optimal perfect preventive maintenance times based on two different gridding increments, $\delta = 0.05, 0.1$, are compared as an example.

As we can see in Table 5.7: under $\delta = 0.05$, we preventively maintain systems earlier (0.45 time units) compared with (0.5 time units) under $\delta = 0.1$, thanks to the increment intervals; the former also produces lower expected cost rate (2.52) than that (2.55) of the latter, which is a 1.18\% reduction. Depending on characteristics of systems and available budget, this could be a significant cost reduction for a particular repairable system.

Second, the precision of the increment parameter $\delta$ is further investigated to understand the underlying impact of gridding increments. In this case, the increment parameter $\delta$ decreases, \textit{i.e.}, the gridding interval becomes larger. As we can see in Table 5.8 with the increment parameter increasing from 0.005 to 0.100, there is little change with respect to the optimal maintenance times ranging from 0.4 to 0.5, \textit{i.e.}, it does not change our optimal decisions significantly. However, we note that the expected cost rate also increases with the increasing of increment parameter $\delta$. It is worthwhile noting that the expected cost rate is not strictly increasing with the increase of $\delta$, \textit{e.g.}, it decreases from 2.5479 to 2.5475 when $\delta$ increases from 0.085 to 0.090. That is because it is sampled at different points with the change of the accuracy of discrete approximation to a continuous distribution.
<table>
<thead>
<tr>
<th>$\delta = 0.05$</th>
<th>$\delta = 0.1$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Maintenance Time</strong></td>
<td><strong>Expected Cost Rate</strong></td>
</tr>
<tr>
<td>$T_{m_1}$</td>
<td>$EC_{CN_1}$</td>
</tr>
<tr>
<td>0.45</td>
<td>2.52</td>
</tr>
<tr>
<td>$T_{m_2}(T_{f_1} \leq T_{m_1})$</td>
<td>$EC_{CN_{21}}$</td>
</tr>
<tr>
<td>1.00(0.05)</td>
<td>10.97</td>
</tr>
<tr>
<td>0.90(0.10)</td>
<td>8.96</td>
</tr>
<tr>
<td>0.85(0.15)</td>
<td>7.75</td>
</tr>
<tr>
<td>0.80(0.20)</td>
<td>6.89</td>
</tr>
<tr>
<td>0.75(0.25)</td>
<td>6.24</td>
</tr>
<tr>
<td>0.75(0.30)</td>
<td>5.72</td>
</tr>
<tr>
<td>0.70(0.35)</td>
<td>5.29</td>
</tr>
<tr>
<td>0.70(0.40)</td>
<td>4.92</td>
</tr>
<tr>
<td>0.65(0.45)</td>
<td>4.60</td>
</tr>
<tr>
<td>$T_{m_2}(T_{f_1} &gt; T_{m_1})$</td>
<td>$EC_{CN_{22}}$</td>
</tr>
<tr>
<td>0.3</td>
<td>1.70</td>
</tr>
</tbody>
</table>

**Table 5.7:** Comparison of optimal perfect preventive maintenance (PPM) time and corresponding expected cost for each chance node based on different increment parameter $\delta$. Bracketed figures are failure time $T_{f_1}$ with respect to $T_{m_1}$ and $T_{m_2}$, numbers in brackets representing corresponding failure times.
<table>
<thead>
<tr>
<th>Increment Parameter $\delta$</th>
<th>Maintenance Time</th>
<th>Expected Cost Rate</th>
<th>Computation Time (seconds)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.005</td>
<td>0.460</td>
<td>2.49</td>
<td>577.03</td>
</tr>
<tr>
<td>0.010</td>
<td>0.460</td>
<td>2.49</td>
<td>99.63</td>
</tr>
<tr>
<td>0.015</td>
<td>0.465</td>
<td>2.49</td>
<td>52.95</td>
</tr>
<tr>
<td>0.020</td>
<td>0.460</td>
<td>2.50</td>
<td>11.71</td>
</tr>
<tr>
<td>0.025</td>
<td>0.475</td>
<td>2.50</td>
<td>6.29</td>
</tr>
<tr>
<td>0.030</td>
<td>0.450</td>
<td>2.51</td>
<td>3.78</td>
</tr>
<tr>
<td>0.035</td>
<td>0.455</td>
<td>2.51</td>
<td>2.60</td>
</tr>
<tr>
<td>0.040</td>
<td>0.480</td>
<td>2.51</td>
<td>1.84</td>
</tr>
<tr>
<td>0.045</td>
<td>0.450</td>
<td>2.52</td>
<td>1.36</td>
</tr>
<tr>
<td>0.050</td>
<td>0.450</td>
<td>2.52</td>
<td>1.02</td>
</tr>
<tr>
<td>0.055</td>
<td>0.440</td>
<td>2.53</td>
<td>0.79</td>
</tr>
<tr>
<td>0.060</td>
<td>0.480</td>
<td>2.53</td>
<td>0.64</td>
</tr>
<tr>
<td>0.065</td>
<td>0.455</td>
<td>2.53</td>
<td>0.53</td>
</tr>
<tr>
<td>0.070</td>
<td>0.490</td>
<td>2.54</td>
<td>0.43</td>
</tr>
<tr>
<td>0.075</td>
<td>0.450</td>
<td>2.54</td>
<td>0.39</td>
</tr>
<tr>
<td>0.080</td>
<td>0.480</td>
<td>2.54</td>
<td>0.34</td>
</tr>
<tr>
<td>0.085</td>
<td>0.425</td>
<td>2.55</td>
<td>0.27</td>
</tr>
<tr>
<td>0.090</td>
<td>0.450</td>
<td>2.55</td>
<td>0.23</td>
</tr>
<tr>
<td>0.095</td>
<td>0.475</td>
<td>2.55</td>
<td>0.21</td>
</tr>
<tr>
<td>0.100</td>
<td>0.500</td>
<td>2.55</td>
<td>0.19</td>
</tr>
</tbody>
</table>

| Table 5.8: Optimal perfect preventive maintenance (PPM) time and corresponding expected cost for chance node $CN_1$ conditioning on various increment parameter $\delta$ based on dynamic programming. |

Because of the impact on expected cost rates by the increment parameter $\delta$, as we are able to see from Table 5.8, the compiling time decreases dramatically from 577.03 to 99.63 seconds when the increment parameter $\delta$ changes from 0.005 to 0.010, and it does not change much after $\delta = 0.020$.

Other indices concerning optimal decisions are also considered. From the top-right graph in Figure 5.10 we can see that the optimal maintenance time fluctuates with
respect to the increment parameter $\delta$ but not very significantly. And from the bottom-left graph in Figure 5.10 the absolute expected cost rate increases with respect to the increment parameter $\delta$, while the relative expected cost, i.e., the expected cost rate per compiling time unit, increases seemingly exponentially. As a result, decision makers in practice choose an appropriate increment interval to trade off between accuracy and the cost budget that they have.

![Graphs showing sensitivity analysis](image)

**Figure 5.10:** Sensitivity analysis concerning gridding intervals: Compiling time (top-left); Optimal Maintenance Time (top-right); Expected Cost Rate (bottom-left); Expected Cost Rate per Compiling Time Unit (bottom-right).

Gridding increments and choice of interval depend on the practical problems and it is often a trade-off between accuracy and efficiency. For example, smaller intervals would increase the accuracy to find the closest maintenance time to theoretical results, however, it would add computing time. Therefore, it is a decision made via cooperation among different stakeholders to seek the optimum design of system maintenance.
5.7.2 Cost Structure

The related costs induced by system performing is one of the core issues considered in our maintenance modelling. So the impact of cost structure on corresponding systems must also be considered.

In previous studies, the perfect preventive maintenance cost $C_m$ is assumed to be less than that of a failure and a repair, $C_f + C_m$. From Table 5.9, we increase the perfect preventive maintenance cost $C_m$ from 0.5 to 3 with fixed failure and repair cost, in other words, the cost difference between failure and maintenance is shrinking. As we can see, the maintenance time increases from 0.5 to 6 which is the maximum range of the time assumption and the expected cost rate increases as well, which means there is less advantage to maintain the system preventively if one has to spend a higher cost to carry it out.

<table>
<thead>
<tr>
<th>Cost Difference between Failure &amp; Maintenance $\Delta$</th>
<th>Maintenance Cost $C_m$</th>
<th>Failure and Repair Cost $C_f + C_r$</th>
<th>Maintenance Time</th>
<th>Expected Cost Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta = 2.5$</td>
<td>0.5</td>
<td>3.0</td>
<td>0.5</td>
<td>2.55</td>
</tr>
<tr>
<td>$\Delta = 2.0$</td>
<td>1.0</td>
<td>3.0</td>
<td>0.7</td>
<td>3.33</td>
</tr>
<tr>
<td>$\Delta = 1.5$</td>
<td>1.5</td>
<td>3.0</td>
<td>0.9</td>
<td>3.76</td>
</tr>
<tr>
<td>$\Delta = 1.0$</td>
<td>2.0</td>
<td>3.0</td>
<td>1.4</td>
<td>3.95</td>
</tr>
<tr>
<td>$\Delta = 0.5$</td>
<td>2.5</td>
<td>3.0</td>
<td>5.2</td>
<td>3.96</td>
</tr>
<tr>
<td>$\Delta = 0.0$</td>
<td>3.0</td>
<td>3.0</td>
<td>6.0</td>
<td>3.96</td>
</tr>
</tbody>
</table>

Table 5.9: Optimal perfect preventive maintenance (PPM) time and corresponding expected cost for chance node $CN_i$ conditioning on cost difference between failure and maintenance based on dynamic programming.

The cost structure is much more complicated in practice. However, it is usually one of the most appealing areas in management. Within business operations, decision makers should have the ability to budget and control cost at a desired level. In particular, maintenance may account for a large proportion in the total budget. In practice, the decision makers will work with the cost accountants and use statistical modelling to build up a comprehensive costing database to budget and control the maintenance cost depending on the complexity of systems.
5.7.3 Prior Sensitivity

We conduct our parameter analysis under the Bayesian framework, so in this section, we analyse the impact of prior assumptions on the system maintenance time decision making.

The expected value for the shape parameter of the Weibull distribution is set from 1 to 10. As we can see in Table 5.10, with the increasing of the prior expectation for the shape parameter $\theta$, the expected cost rate decreases significantly from 3.05 to 0.91, but it has little impact on the decision about the optimal maintenance time, which simply ranges from 0.5 to 0.6. That is because a larger expectation of $\theta$ indicates a higher hazard (failure) rate: one can reduce the expected cost rate by applying a same maintenance policy (here same maintenance time) to a system with a higher possibility to fail.

<table>
<thead>
<tr>
<th>$E(\theta_{\text{prior}})$</th>
<th>Maintenance Time</th>
<th>Expected Cost Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.5</td>
<td>3.05</td>
</tr>
<tr>
<td>2</td>
<td>0.5</td>
<td>2.55</td>
</tr>
<tr>
<td>3</td>
<td>0.5</td>
<td>2.00</td>
</tr>
<tr>
<td>4</td>
<td>0.5</td>
<td>1.57</td>
</tr>
<tr>
<td>5</td>
<td>0.5</td>
<td>1.30</td>
</tr>
<tr>
<td>6</td>
<td>0.6</td>
<td>1.16</td>
</tr>
<tr>
<td>7</td>
<td>0.6</td>
<td>1.05</td>
</tr>
<tr>
<td>8</td>
<td>0.6</td>
<td>0.98</td>
</tr>
<tr>
<td>9</td>
<td>0.6</td>
<td>0.93</td>
</tr>
<tr>
<td>10</td>
<td>0.6</td>
<td>0.91</td>
</tr>
</tbody>
</table>

Table 5.10: Optimal perfect preventive maintenance (PPM) time and corresponding expected cost for chance node $CN_1$ conditioning on various prior mean of parameter $\theta$ based on dynamic programming.

In Table 5.11, the standard deviation for the shape parameter $\theta$ does not have a significant effect on making the optimal decision about the maintenance time, which ranges from 0.4 to 0.6, or the expected cost rate, which ranges from 2.60 to 1.84.
<table>
<thead>
<tr>
<th>$\text{SD}(\theta_{\text{prior}})$</th>
<th>Maintenance Time</th>
<th>Expected Cost Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\sigma=0.1$</td>
<td>0.4</td>
<td>2.60</td>
</tr>
<tr>
<td>$\sigma=0.3$</td>
<td>0.4</td>
<td>2.64</td>
</tr>
<tr>
<td>$\sigma=0.5$</td>
<td>0.4</td>
<td>2.68</td>
</tr>
<tr>
<td>$\sigma=0.7$</td>
<td>0.5</td>
<td>2.65</td>
</tr>
<tr>
<td>$\sigma=0.9$</td>
<td>0.5</td>
<td>2.59</td>
</tr>
<tr>
<td>$\sigma=1.0$</td>
<td>0.5</td>
<td>2.55</td>
</tr>
<tr>
<td>$\sigma=2.0$</td>
<td>0.5</td>
<td>2.22</td>
</tr>
<tr>
<td>$\sigma=3.0$</td>
<td>0.5</td>
<td>1.98</td>
</tr>
<tr>
<td>$\sigma=4.0$</td>
<td>0.5</td>
<td>1.84</td>
</tr>
</tbody>
</table>

Table 5.11: Optimal perfect preventive maintenance (PPM) time and corresponding expected cost for chance node $CN_1$ conditioning on various prior standard deviation of parameter $\theta$ based on dynamic programming.

The findings above helps practitioners to implement our method in practice because of the simpler interpretation to expectation than that to standard deviation.

5.7.4 Utility Function Forms

The utility functional forms also need to be considered as it would produce different risk aversion.

Let us examine the maintenance time decision under uncertainty following the Cobb-Douglas utility function \cite{Cobb and Douglas 1928} written for one phase system for simplicity:

$$U \left( \frac{C_f + C_r}{T_f}, \frac{C_m}{T_m}, f_{T_f}(t_f \leq T_m), f_{T_f}(t_f > T_m) \right)$$

$$= \left( \frac{C_f + C_r}{T_f} \right)^{f_{T_f}(t_f \leq T_m)} \left( \frac{C_m}{T_m} \right)^{f_{T_f}(t_f > T_m)}.$$ (5.15)

Apply logarithm to each side of (5.15),

$$\log U = f_{T_f}(t_f \leq T_m) \log \left( \frac{C_f + C_r}{T_f} \right) + f_{T_f}(t_f > T_m) \log \left( \frac{C_m}{T_m} \right).$$ (5.16)

Then we can use this monotonically transformed utility function to find the optimal maintenance time and compare it with the expected cost rate objective function.
As we can see in Table 5.12, for chance node $CN_1$, one is suggested to maintain the system at local time 0.5 under log utility function and expected cost rate objective (non-log utility function).

<table>
<thead>
<tr>
<th>Non-Log Utility Function</th>
<th>Log Utility Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maintenance Time</td>
<td>Expected Cost</td>
</tr>
<tr>
<td>$T_{m_1}$</td>
<td>$EC_{CN_1}$</td>
</tr>
<tr>
<td>0.5</td>
<td>2.55</td>
</tr>
</tbody>
</table>

$T_{m_2}(T_f \leq T_{m_1})$ $EC_{CN_21}$ $T_{m_2}(T_f \leq T_{m_1})$ $EC_{CN_21}$
| 0.9(0.1) | 8.69 | 1.1(0.1) | 1.91 |
| 0.8(0.2) | 6.83 | 0.9(0.2) | 1.75 |
| 0.7(0.3) | 5.71 | 0.9(0.3) | 1.61 |
| 0.7(0.4) | 4.93 | 0.8(0.4) | 1.49 |
| 0.6(0.5) | 4.36 | 0.7(0.5) | 1.39 |

$T_{m_2}(T_f > T_{m_1})$ $EC_{CN_22}$ $T_{m_2}(T_f > T_{m_1})$ $EC_{CN_22}$
| 0.3 | 1.61 | 0.5 | 0.32 |

Table 5.12: Optimal perfect preventive maintenance (PPM) time and corresponding expected cost for each chance node by dynamic programming based on non-Log and Log utility functions. Bracketed figures are failure time $T_f$ with respect to $T_{m_1}$ and $T_{m_2}$, numbers in brackets representing corresponding failure times.

However, the expected cost rate (0.62) for log utility function is much less than that (2.55) for non-log utility function. That is because the logarithm transformation of the expected cost rate function conveys risk aversion to the decision maker.

That is because the logarithm transformation of the expected cost rate function presents risk aversion on the decision making. Under the non-log utility function, because the decision maker is risk neutral, she or he is supposed to maintain the system later compared to her or his counterpart with risk aversion, however, the maintenance time is also 0.5, which means the system is maintained earlier than it is supposed to be, as a result, the expected cost rate is higher than that under log transformation.
5.8 Parallel Computing

We have seen that the increment parameter $\delta$ is an essential element in the gridding method to optimising the maintenance time via dynamic programming in §5.7.1. When $\delta$ is smaller, on the one hand, it will increase the accuracy of the gridding method, which leads one to approach the theoretically optimal maintenance time. On the other hand, it results in a problem of being computationally expensive.

Parallel computing is a method of carrying out a large number of calculations simultaneously based on the principle that a complicated large problem can be independently transformed into smaller and simpler ones. A few applications of parallel computing in maintenance have been investigated. For example, Yang et al. (2012) build up a parallel computing platform for multiobjective simulation optimisation of bridge maintenance planning that can span tens of years, and also investigate the proposed framework through a practical case, which shows the superiority to GA method.

Here the possible solutions to implement parallel computing in further research are discussed. For example, we need to compute joint probability mass of $T_{f1}$ and $T_{f2}$, see Figure 5.11:

\[
\begin{align*}
\text{Matrix } & \quad \begin{pmatrix}
0 & 1 \\
1 & 0 \\
1 & 0 \\
0 & 1 \\
0 & 1 \\
1 & 0 \\
0 & 1 \\
1 & 0 \\
\end{pmatrix} \\
\text{Figure 5.11: Parallel Computing for } & \quad p(t_{f2}, t_{f1})
\end{align*}
\]

If there are only two possible values of for $T_{f1}$ and $T_{f2}$, we can calculate them separately in four different processes independently and then combine them to form...
the matrix for the joint probability mass.

With the development of cloud computing, decision makers could share computer processing resources and data to make better and coherent decisions in a comprehensive and efficient manner. Although discussion of cloud computing is beyond the scope of this thesis, it is interesting to mention that there have been some investigation on parallel computing on maintenance. For instance, Vert et al. (2015) review models and algorithms of a pipeline parallel computing process in intelligent scheduling. This parallel computing process models the computing nodes’ change of resource consumption, which can help decision makers to see which nodes have the highest usage in computing. Through being implemented in a virtual cloud environment, it provides better security and higher computing.
Chapter 6

Risk Aversion in Maintenance

The concept of risk aversion plays an important part in economics and finance. In maintenance modelling, risk aversion would induce a more costly maintenance policy but could potentially avoid huge expenditures. In this context, if one is more risk averse, she or he would tend to maintain the system more frequently or earlier. In this chapter, the approach proposed by Baker (2006) is used and modified to analyse the effect of risk aversion on the variability of systems in cash flows.

6.1 Utility Functions

Baker (2006) introduces the concept of risk aversion in maintenance policies. Instead of dealing with the expected utility per unit time, we model the expected cost per unit time and modify the approaches to modelling in (Baker 2006) incorporating risk aversion.

Risk aversion is usually modelled by a concave utility function, which can be expressed as $U(x)$, which is the utility of a sum of resources $x$, where $U' > 0$ and $U'' < 0$, which are the first and second derivatives of the utility function $U(x)$.

In this chapter, the exponential utility function is used, and defined as

$$U = \frac{1 - \exp(-\eta x)}{\eta},$$

where $\eta > 0$ is the parameter measuring the risk aversion. Note that

$$\lim_{\eta \to 0} \frac{1 - \exp(-\eta x)}{\eta} = x$$

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and its Taylor series expansion at \( x = 0 \) is
\[
x - \frac{\eta}{2} x^2 + \frac{\eta^2}{6} x^3 - \frac{\eta^3}{24} x^4 + \frac{\eta^4}{120} x^5 - \frac{\eta^5}{720} x^6 + O(\eta^6)
\]
In other words, as \( \eta \to 0 \), \( U \to x \), and to first order in \( \eta \),
\[
U \simeq x - \frac{\eta}{2} x^2.
\]
(6.2)

There are a few advantages of using this utility function:

1. The exponential distribution has the memoryless property. The utility function in (6.1) is defined with constant risk aversion, in other words, the amount of resource that one prepares to risk is not a function of the initial resource \( x_0 \); this utility function does not depend on the resources spent on other fields either. As a result, using of this utility function with this feature would avoid considering one’s resources and other activities.

2. To first order in the risk aversion parameter \( \eta \), all utility functions on the whole real line will be in the form given in (6.2). Therefore, using exponential utility function will produce more general results under lower risk aversion.

3. Most utility functions generally have more than one parameters as shown in (§3.3.1, page 44), which would add much complexity on modelling.

As this thesis deals with costs induced by system performance, we define cost \( C = -x \) having disutility
\[
-\frac{U}{\eta} = \frac{\exp(\eta C) - 1}{\eta}.
\]
(6.3)

The certainty equivalent is defined as a guaranteed return that one would accept, instead of taking a chance on a higher but uncertain and risky return. In our modelling, we interpret it as the sum of resources that one definitely gains or loses which would have the same expected utility as the variable cash flows via the system performance. Therefore, if the system is performing for time \( T \), the exponential utility function can be expressed as
\[
\frac{\exp(\eta \cdot CE \cdot T) - 1}{\eta} = \frac{\mathbb{E}\exp(\eta C) - 1}{\eta},
\]
where \( CE \) is the certainty equivalent per unit time, or
\[
CE = \frac{\log \{ \mathbb{E}\exp(\eta C) \}}{\eta T}.
\]
(6.4)
6.2 Risk-averse Maintenance Modelling

We consider the preventive maintenance policy for the system modelled in chapter 4, which would be maintained perfectly at an optimum maintenance time $T_m$ or at failure $T_f$, whichever occurs first, i.e., the length of system’s first phase is $T_m$ or $T_f$, whichever is smaller, denoted as $T_i$. Let the cost over the $i$th phase be $F_i$, where $F_i$ is a random cash flow during the phase $i$; in other words, $F_i$ induced by preventive maintenance or failure and repair, of which the cost is $C_m$ and $C_f + C_r$, respectively. The certainty equivalent per unit time for phase $i$ is

$$CE_i = \frac{\log \mathbb{E}\{\exp(\eta F_i)\}}{\eta T_i}$$

For simplicity, the certainty equivalent per unit time for each phase is

$$CE = \frac{\log \mathbb{E}\{\exp(\eta F)\}}{\eta T}$$  \hspace{1cm} (6.5)

by dropping the phase subscript $i$.

Note that the expression $\mathbb{E}\exp(\eta F)$ in equation (6.5) is the moment generating function (m.g.f) of the random variable $F$ with risk aversion parameter $\eta$.

When the value of the risk aversion parameter $\eta$ is small, from formula (6.2) $CE$ reduces to

$$CE \simeq \frac{\mathbb{E}(F) + \frac{\eta}{2}\text{Var}(F)}{T},$$

which shows that the cost per unit time is modified by the risk aversion parameter $\eta$ and increases with the variance of the cost induced in each phase.

Also $\log \mathbb{E}\{\exp(\eta F)\}$ is the cumulant generating function, hence

$$CE = \frac{\sum_{j=1}^{\infty} \eta^{j-1} k_j / j!}{T}$$  \hspace{1cm} (6.6)

where $k_j$ is $j^{th}$ cumulant of the cost for each phase. As we can see, with the risk aversion parameter $\eta$ increasing, certainty equivalent $CE$ also increasingly weights towards higher cumulants, such as skewness and kurtosis.

As we model sequential preventive maintenance with unfixed maintenance time, the length of each phase $T_i$ is not equal. As a result, equation (6.5) becomes

$$\exp(\eta \cdot CE \cdot T) = \sum_{i=1}^{N} \{\log \mathbb{E}(\exp(\eta F))\}^i P_i = G(\log \mathbb{E}(\exp(\eta F)))$$
for a system with $N$ phases, where $P_i$ is the probability that $i$ phases have occurred by time $T$, therefore,

$$CE = \frac{\log \{G(\log \mathbb{E} \exp(\eta F))\}}{\eta T},$$

(6.7)

where $\log G(\cdot)$ is the cumulant generating function for the number of phases. Cox (1962) gives the first few cumulants when expanding equation 6.7 as $T \rightarrow \infty$,

$$CE = \left(1 \frac{1}{k_1}\right) \frac{\log \mathbb{E} \exp(\eta F)}{\eta} + \frac{1}{2!} \left(\frac{k_2}{k_1^2}\right) \frac{(\log \mathbb{E} \exp(\eta F))^2}{\eta}$$

$$+ \frac{1}{3!} \left(\frac{3k_3^2}{k_1^3} - \frac{k_3}{k_1}\right) \frac{(\log \mathbb{E} \exp(\eta F))^3}{\eta}$$

$$+ \frac{1}{4!} \left(\frac{k_4}{k_1^3} + \frac{15k_3^2}{k_1^4} - \frac{10k_2k_3}{k_1^3}\right) \frac{(\log \mathbb{E} \exp(\eta F))^4}{\eta} + \cdots. \quad (6.8)$$

For the system considered here which is perfectly maintained at time $T_m$ or on failure $T_f$. The cost per unit time is

$$C(T_m) = \frac{C_f + (C_m - C_f - C_r)S(T_m)}{T} \quad (6.9)$$

where $S(\cdot)$ is the survival function and $T_m$ is optimum maintenance time obtained in chapter 4 by dynamic programming approach. We consider the first term in the expansion 6.8 to approximate $CE$.

Define $I$ as an indicator variable, where $I = 1$ denotes system failure in $(0, T_m]$ and $I = 0$ denotes system has survived to time $T_m$ with failure occurring. Therefore,

$$CE(T_m) = \frac{\log \{\mathbb{E} \exp(\eta((C_f + C_r)I + C_m(1 - I)))\}}{\eta T} + \cdots, \quad (6.10)$$

where $T$ is the length of the phase.

Baker (2006) gives the expansion of the exponential as

$$\mathbb{E} \exp((C_f + C_r - C_m)\eta I) = S(T_m) + \exp(\eta(C_f + C_r - C_m))(1 - S(T_m)),$$

and rearrange equation 6.10

$$CE(T_m) = \frac{C_f + C_r}{T} + \frac{\log \{1 + S(T_m)(\exp(\eta(C_m - C_f - C_r)) - 1)\}}{\eta T} + \cdots. \quad (6.11)$$

Then can use the equation 6.11 to analyse the relationship between risk aversion parameter $\eta$ and the corresponding certainty equivalent.
6.3 Numerical Examples

We consider the sequential preventive maintenance model in chapter 4 where the system is perfectly maintained preventively at $T_m$ or at failure time $T_f$ following a Weibull distribution with survival function $S(t) = \theta t^{\theta - 1}$, whichever occurs first, whilst the corresponding cost is $C_m$ and $C_f + C_r$, respectively. From Table 4.1 (page 86), the optimum maintenance time $T_m$ for phase one is 0.5 and corresponding possible failure times $T_f$ are 0.1, 0.2, 0.3, 0.4, 0.5.

Replace $C_f$ as 2, $C_r$ as 1 and $C_m$ as 0.5 in equation 6.11,

$$ CE(T_m = 0.5 \mid T_f, \theta, \eta) = \frac{3}{T_f} + \log \left\{ \frac{1 + S(0.5)(\exp(-2.5\eta) - 1)}{\eta T_f} \right\} + \cdots $$

$$ = \frac{3}{T_f} + \log \left\{ \frac{1 + \theta 0.5^{\theta - 1}(\exp(-2.5\eta) - 1)}{\eta T_f} \right\} + \cdots (6.12) $$

As we can see in equation 6.12, the certainty equivalent is associated with failure time $T_f$, shape parameter $\theta$ for Weibull distribution with scale parameter 1 and risk aversion parameter $\eta$.

The shape parameter $\theta$ of the Weibull distribution is set as 2 and analysed the relationship between the risk aversion parameter $\eta$ and the certainty equivalent $CE$. As we can see in Figure 6.1 first, the certainty equivalent $CE$ increases with increasing risk aversion; second, the certainty equivalent $CE$ decreases with failure time increasing, i.e., when the system is more reliable. In other words, one intends to spend less resource if the system is more reliable.

![Figure 6.1: Certainty-Equivalent with the risk-aversion parameter $\eta$ under different operating times.](image-url)
6.4 Discussions

This chapter investigates maintenance optimisation via risk aversion from a certainty-equivalent point of view. The aim in this section is to discuss relevant areas of interest for further research.

- As can be seen in previous studies, utility function with very low risk aversion provides the same maintenance policy, which tends towards minimising the average cost per unit time. Although one would be interested to know if there are other utility functions can be applied in similar maintenance optimisation modelling, it has been proved that only the linear and exponential utility functions have properties that are reasonable to apply in practice. However, it could be an interesting area to investigate depending on practical scenarios, *e.g.*, nonparametric utility function modelling.

- The findings that minimising the average cost per unit time is almost equivalent to maximising the expected utility of cost rate and the certainty-equivalent cost increases as risk aversion increases bring up the question of determining the risk aversion parameter of the utility function. Baker (2010) suggests that one may refer to a plot showing the relationship of certainty-equivalent cost per unit time that is estimated from cost data and varying values of risk aversion parameter. The certainty-equivalent cost per unit time that is not very sensitive to risk aversion should be preferred.
Chapter 7

Conclusions and Future Work

7.1 Conclusions

From the utility point of view, the maintenance problems for repairable systems, including corrective and preventive maintenance were mainly discussed. Under the framework of Bayesian methodology, the parameters of models are random rather than fixed as for the non-Bayesian method, while the former often suits in most of the practical situations, the failure distribution is either unknown or contains several unknown parameters. In such case, the Bayesian approach can be quite effective to estimate these unknown parameters by assigning prior distributions to them.

Starting from a two-phase system maintenance model, we explored the system’s perfect preventive maintenance by dynamic programming, compared it with a myopic method, and found the dynamic programming superior in terms of optimising maintenance time. When the risk aversion parameter in the utility function is very small, our utility-based maintenance optimisation would reduce to minimising the expected cost per unit time.

Under different failure time distributions, we explored the effect of failure time assumptions on the optimal maintenance time, and that a gamma failure time assumption would result in later maintenance time.

Considering time effect on maintenance optimisation, we found maintenance tends to be done earlier with the future cost becoming more expensive. Several modified maintenance optimisation models are proposed such as modelling systems with discrete failure distribution, parallel redundant systems.
By manipulating a conditional probability matrix, we explore and compare the difference between perfect preventive maintenance and imperfect preventive maintenance and we may generally conclude that under an imperfect preventive maintenance modelling framework systems tend to be maintained later because the system would enter into a state with higher probability to fail if imperfect preventive maintenance were conducted.

To avoid the problem of dimension increasing and to speed up computation time, we proposed a hybrid myopic-dynamic programming method. By using the posterior distribution for the parameter of interest, we see that the hybrid myopic-dynamic programming maintenance time at \( CN_{21} \) are earlier, which results in lower expected cost rate compared with dynamic programming for a two-phase system.

### 7.2 Applications

The utility-based maintenance strategies and policies proposed in this thesis are intensively mathematically and theoretically modelled with a number of assumptions. Although they are not straightforward to apply in practice, they provide new perspectives on modelling preventive maintenance with modification. The prognostic maintenance policies derived from historical information and system evolution require decision makers to understand thoroughly the characteristics of the system of interest and to anticipate all possible consequences as much as possible.

As a result, models and methodologies proposed in this thesis are more suitable for large industrial and business purposes. The corresponding systems are supposed to be well documented for decision makers to extract critical information and parameters for further maintenance modelling. For example, decision makers should be able to have access to advice from an expert if required, because expert information is essential to justify related prior parameters to implement Bayesian dynamic programming. At the same time, the potential future states of a system should be finite and of a relatively small number. For instance, a personal computer’s possible states could be ‘working’, ‘sleeping’, or ‘down’, which are accountable states.

Because the methodologies in this thesis are sensitive to the number of system states, it would not be reasonable to apply them to highly dynamic systems, such as
machines dealing with signal processing. Other systems that could potentially be of interest to apply modelling methods in this thesis are an automatic manufacturing system, a robotic process, or a computer server for the non-life essential services, in which cases failure is neither rare or frequent, maintenance itself is not cheap or trivial, but failure is a considerable expense, though not exorbitantly so. Hence, this approach is not suitable to apply to maintenance of systems with very high risk aversion properties, e.g., a nuclear power facility, an off-shore oil field, or a life support system. It is also not worthwhile applying to trivial systems where the computational cost of performing this analysis outweighs any savings.

In practice, there are few systems that would meet the requirements of the theoretical modelling, though it is the decision makers’ prerogative to adjust relevant parameters, define overall and critical objectives, and utilise available information to seek the practical maintenance policies appropriate to the properties of an individual system.

### 7.3 Outlook

Maintenance optimisation for repairable systems is part of maintenance theory in engineering, and it also requires more attention and effort to deal with increasing issues concerning it.

1. This research is based on utility to optimise maintenance systems. However, it can be required to optimise several objectives simultaneously, which impose difficulty and challenges.

2. The utility function is arbitrarily assigned in this research from a maintenance engineering perspective. In situations where the utility function is unknown, other theory in utility, for example, adaptive utility, can be explored.

3. Failure time distributions are in parametric form in this thesis, so it could be interesting to explore semi-parametric and non-parametric assumptions in future research.

4. Systems considered in this research are relatively simple, however, as the complexity of systems in modern society increases, high demand to deal with the
maintenance on complex systems requires us to explore this field, despite the huge mathematical and computational challenges.

5. As economies grow in modern society, there is a vast increasing demand for resources, including human and material. Hence it is worthwhile considering these constraints in future maintenance optimisation modelling.
Appendix A

Glossary

\[ \preceq, \prec, \sim \] Binary preference relations
\( \theta \) Shape parameter of Weibull lifetime distribution
\( \kappa \) Scale parameter of Weibull lifetime distribution
\( \eta \) Risk aversion parameter
\( B \) Budget
\( C_f \) Cost induced by failure
\( C_m \) Cost induced by maintenance
\( C_r \) Cost induced by repair
\( CN_i \) Chance node \( i \)
\( D_f \) Arbitrary probability distribution
\( EU(\cdot) \) Expected utility function
\( T_i \) Local time \( i \)
\( T_{fi} \) Failure time for phase \( i \)
\( T_{mi} \) Maintenance time for phase \( i \)
\( U(\cdot) \) Utility function
\( X_i \) Global time \( i \)
Appendix B

Mathematical Proofs

B.1 Iso-elastic Utility Functions

A class called iso-elastic utility functions have the following form

\[ U(x) = \begin{cases} 
\frac{x^{1-a}-1}{1-a} & \text{for } a > 0, \ a \neq 1; \\
\log(x) & \text{the limiting case for } a = 1.
\end{cases} \]

These functions have the property of iso-elasticity, which means that we get the same utility function (a positive affine transformation) if the cost is scaled by some constant \( k \). Formally,

For all \( k > 0 \),

\[ U(kx) = f(k)U(x) + g(k), \]

for some function \( f(k) > 0 \) which is independent of \( x \) and some function \( g(k) \) which is independent of \( x \) as well.

First consider the case when \( a \neq 1 \),

\[ U(kx) = \frac{(kx)^{1-a}-1}{1-a} \\
= k^{1-a} \left( \frac{x^{1-a}-1}{1-a} \right) + \frac{k^{1-a}-1}{1-a} \\
= k^{1-a}U(x) + \frac{k^{1-a}-1}{1-a} \]

The log function can be written as

\[ U(kx) = \log(kx) \\
= \log(k) + \log(x) \\
= U(x) + \log(k) \]
B.2 Negative Exponential Utility Functions

A negative exponential utility function is of the form

\[ U(x) = -\exp\{-ax\}. \]

Since the first derivative \( U'(x) = a \exp\{-ax\} > 0 \) and the second derivative \( U''(x) = -a^2 \exp\{-ax\} < 0 \), this one is also a legitimate utility function.

The class of negative exponential utility functions has an interesting property that it is invariant under any cost transformation, i.e., for any constant \( k \),

\[ U(k + x) = f(k)U(x) + g(k), \]

for some function \( f(k) > 0 \) which is independent of \( x \) and some function \( g(k) \) which is independent of \( x \) as well, it can be verified as below:

\[
U(k + w) = -\exp\{-a(k + x)\} \\
= -\exp\{-ka\} \exp\{-ax\} \\
= \exp\{-ka\} U(x)
\]

B.3 Certainty Equivalent

In general, the certainty equivalent \( CE \) for a maintenance policy whose consequence is given by a random variable \( X \) is:

\[ CE = U^{-1}(E(U(X))) \]

\[ U(CE) = E(U(X)) \]

If a decision maker with utility function \( U \) has the current cost budget less than \( CE \), she will regard the system less reliable and tend to maintain the system earlier; if her current cost budget is more than \( CE \), she will think the system is more reliable and tend to maintain the system later; and if her current cost budget is exactly \( CE \), she will be thinking there is no difference between maintain the system earlier or not.

Decision makers use utility functions to compare different decisions to each other. In this sense, we can rescale a utility function via multiplying it by a positive constant and/or adding any other positive or negative constant, which is called a positive affine
Two utility functions produce the same results if they are connected via a positive affine transformation.

Suppose we have constants $\alpha > 0$ and $\beta$ and a utility function $U$; another utility function $V$ is defined via $U, \alpha$ and $\beta$ as:

$$V(x) = \alpha U(x) + b$$

Since

$$V'(x) = \alpha U'(x) > 0 \quad \text{because} \quad \alpha > 0 \quad \text{and} \quad U'(x) > 0$$
$$V''(x) = \alpha U''(x) < 0 \quad \text{because} \quad \alpha > 0 \quad \text{and} \quad U''(x) < 0,$$

$V$ is a legitimate utility function.

If one can make a decision with consequence that is given by a random variable $X$, let $CE$ be the certainty equivalent with utility function $U$, then

$$CE = U^{-1}(E(U(X)))$$
$$U(CE) = E(U(X))$$
$$V(CE) = \alpha U(CE) + \beta$$
$$= \alpha E(U(X)) + \beta$$
$$= E(V(X))$$
$$CE = V^{-1}(E(V(X)))$$

So we regard the two utility functions $U$ and $V$ the same.

### B.4 Risk Aversion

We say that two utility functions are the same if they are connected via a positive affine transformation

$$V(X) = \alpha U(X) + \beta$$

The first and second derivatives of both sides are

$$V'(X) = \alpha U'(X)$$
$$U''(X) = \alpha U''(X)$$
By division, we obtain
\[ \frac{V''(X)}{V'(X)} = \frac{U''(X)}{U'(X)} \]

Assume that there exists such a pair of functions \( U \) and \( V \) for which \( \frac{V''(X)}{V'(X)} = \frac{U''(X)}{U'(X)} \) holds and let

\[ g(X) = \frac{V'(X)}{U'(X)} \]

Its first derivative
\[
\begin{align*}
g' &= (V'(U')^{-1})' \\
    &= V''(U')^{-1} + V'(-1)(U')^{-2}U'' \\
    &= \frac{V''}{U'} - \frac{V'U''}{(U')^2} \\
    &= \frac{V''U' - V'U''}{(U')^2} \\
    &= \frac{0}{(U')^2} \\
    &= 0
\end{align*}
\]

Since \( g' = 0 \), \( g(X) = \alpha \) for some constant \( \alpha \), then

\[ V'(X) = \alpha U'(X) \]

Integrals of both sides are
\[
\int V''(X) \, dx = \int \alpha U'(X) \, dx \\
V(X) = \alpha U(X) + \beta \quad \text{for some constant } \beta
\]

Now we show that if and only if their second derivatives and first derivatives are the same, the two utility functions can be regarded as same.
Appendix C

Computational Notes

The statistical computation in this thesis are written in R code [R Core Team 2015], running on a MacBook Pro (mid 2012) with an Intel Core i7 2.9 GHz CPU.

```
# myrcode1
# two-time period system

# predefined variables
Cf <- 2  # failure cost
Cr <- 1  # repair cost
Cm <- 0.5 # maintenance cost
alpha <- 1 # time effect parameter, discount rate: <=1

library(truncnorm)
delta <- 0.1 # increment
theta.poss <- seq(from=1, to=10, by=delta) # possible values of theta
theta.mass <- dtruncnorm(theta.poss, a=1, b=Inf, mean=2, sd=1)
theta.prob <- theta.mass/sum(theta.mass) # probability of theta

# possible values and lengths of tf1, tf2, tm1 and tm2
tf1.poss <- seq(from=delta, to=6, by=delta)
tf2.poss <- seq(from=delta, to=6, by=delta)
tm1.poss <- seq(from=delta, to=6, by=delta)
tm2.poss <- seq(from=delta, to=6, by=delta)

tf1.n <- length(tf1.poss)

# compute tf2gtf1.prob: p(tf2|tf1)
# define a matrix for the joint mass distribution of tf1 & tf2
# rows are possible tf1 and columns are possible tf2
tf2gtf1.upper <- matrix(rep(0, tf1.n*tf2.n), nrow=tf1.n, ncol=tf2.n)
for (i in 1:tf1.n) {
  ...
}
```
for (j in 1:t2.n) {
  t2gtf1.upper[i, j] <- sum(dweibull(t2.poss[i], shape=theta.poss, scale=1) * dweibull(t2.poss[j], shape=theta.poss, scale=1) * theta.prob)
  # t2gtf1.upper[i, j] <- sum(dgamma(t2.poss[i], shape=theta.poss, scale=1) * dgamma(t2.poss[j], shape=theta.poss, scale=1) * theta.prob)
}

joint prob of tf1 and tf2: p(tf1, tf2)
t2gtf1.upper.prob <- t2gtf1.upper/sum(t2gtf1.upper)
# marginal prob of tf1: p(tf1)
t2gtf1.lower.prob <- rowSums(t2gtf1.upper.prob)
# conditional prob of tf2 given tf1: p(tf2|tf1)
t2gtf1.prob <- exp(log(t2gtf1.upper.prob)-log(t2gtf1.lower.prob))

# plot t2gtf1.prob: p(tf2|tf1)
# convert matrix t2gtf1.prob to a data frame with possible tf2
t2gtf1.prob.frame <- data.frame(t( t2gtf1.prob ), Tf2=t2.poss)
# convert the data frame from "wide" format to "long" format
library(reshape2)
t2gtf1.prob.long <- melt(t2gtf1.prob.frame,
  id.vars="Tf2",
  variable.name="Tf1",
  value.name="Density")
levels(t2gtf1.prob.long$Tf1) <- t2.poss # set the variables as possible tf1
library(grDevices)
library(ggplot2)
ggplot(data=t2gtf1.prob.long, # tf1 from delta to 6
  aes(x=Tf2, y=Density, colour=Tf1)) +
  geom_line()
# ggplot(data=t2gtf1.prob.long[1:1800, ], # tf1 from delta to 3
#  aes(x=Tf2, y=Density, colour=Tf1)) +
#  geom_line()
# ggplot(data=t2gtf1.prob.long[1801:3600, ], # tf1 from 3+delta to 6
#  aes(x=Tf2, y=Density, colour=Tf1)) +
#  geom_line()
max.prob.t2 <- apply(t2gtf1.prob, 1, max)
max.t2 <- delta*apply(t2gtf1.prob, 1, which.max)
# create a data frame
# highest probability, max tf2 given tf1
hpt2gtf1 <- data.frame(t1.poss, max.t2, max.prob.t2)

# for each t1.poss, the highest prob of tf2
ggplot(hpt2gtf1, aes(x=t1.poss, y=max.prob.t2)) +
  geom_point() +
  geom_line()
geom_text(aes(x= tf1.poss +0.08, label = max.tf2), size=2, hjust=0) +
xlab("Tf1") + ylab("Highest probability value w.r.t Tf2")
# for each tf1.poss, the tf2 that has the highest prob
ggplot(hpmtf2gtf1, aes(x= tf1.poss, y=max.tf2)) +
geom_point() +
geom_text(aes(y=max.tf2+0.01, label = round(max.prob.tf2, 2)),
  size=2, vjust=0) +
xlab("Tf1") + ylab("Tf that has the highest probability")
# for each tf1.poss, the tf2 that has the highest prob and its prob
ggplot(hpmtf2gtf1, aes(x=max.tf2, y=max.prob.tf2)) +
geom_point() +
geom_text(aes(x=max.tf2+0.015, label = tf1.poss),
  size=2, hjust=0) +
xlab("Tf2 that has the highest probability given Tf1") +
ylab("Highest probability value w.r.t Tf2")
# # scatter 3d plot for tf1.poss, max.tf2 and max.prob.tf2
library(scatterplot3d)
# scatterplot3d(tf1.poss, max.tf2, max.prob.tf2,
107  # xlab="Tf1", ylab="Tf2 that has highest density given Tf1",
108  # zlab="Density")
109
110 # compute centf2gtf1.prob, i.e., p(tf2>tm2|tf1)
111 centf2gtf1.prob <- matrix(rep(0, tf1.n*tm2.n), nrow= tf1.n, ncol=tm2.n)
112 for (i in 1:tf1.n) {
113  for (j in 1:(tm2.n-1)) {
114    centf2gtf1.prob[i, j] <- sum(tf2gtf1.prob[i, (j+1):tm2.n])
115  }
116 }
117
118 # compute tf2gtm1.prob, i.e., p(tf2|tf1>tm1)
119 # define a vector representing p(tf1>tm1), of which the length is tm1.n
120 tf2gtm1.lower.prob <- rep(0, tm1.n)
121 # define a matrix for the joint prob of tf1>tm1 and tf2, i.e., p(tf1>tm1, tf2)
122 # rows are possible tm1 and columns are possible tf2
123 tf2gtm1.upper.prob <- matrix(rep(0, tm1.n*tf2.n), nrow=tm1.n, ncol=tf2.n)
124 for (i in 1:tm1.n) {
125  for (j in 1:tf2.n) {
126    tf2gtm1.lower.prob[i] <- sum(tf2gtf1.lower.prob[tf1.poss>tm1.poss[i]])
127    # tf2gtm1.lower.prob is p(tf1>tm1)
128    # for a given tm1, sum p(tf1) of which tf1>tm1
129    for (j in 1:tf2.n) {
130      tf2gtm1.upper.prob[i, j] <- sum(tf2gtf1.upper.prob[tf1.poss>tm1.poss[i],j])
131      # tf2gtm1.upper.prob is the joint p(tf1>tm1, tf2)
132      # for a given tm1, sum p(tf1, tf2) of which tf1>tm1 and corresponding tf2
133    }
134  }
135}
136 tf2gtm1.prob <- exp(log(tf2gtm1.upper.prob[-tm1.n,])-log(tf2gtm1.lower.prob[-tm1.n]))
137 # tf2gtm1.prob is P(tf2|tf1>tm1)
138 # use [-tm1.n] to disgard the last one as it is NaN
\# compute centerf2gtm1, i.e., p(tf2|tm2|tf1>tm1)
centf2gtm1.prob <- matrix(rep(0, (tm1.n-1)*tm2.n), nrow=tm1.n-1, ncol=tm2.n)
for (i in 1:(tm1.n-1)) {
  for (j in 1:(tm2.n-1)) {
    centf2gtm1.prob[i, j] <- sum(centf2gtm1.prob[i, (j+1):tm2.n])
  }
}

\# for preventive maintenance (PM)
\# we assume that the failure rates increase after PM
\# i.e., p(tf2|tf1>tm1) > p(tf2|tf1)
\# for each row, the number of entries to be re-allocated
\# prop is an integer number from 1 to ncol-1,
\# which is the number of prob(s) taken out from each row
prop <- 55
prop.tf2gtm1.prob <- matrix(rep(0, (tm1.n-1)*tf2.n), nrow=tm1.n-1, ncol=tf2.n)
\# for each row of tf2gtm1.prob, i.e., P(tf2|tf1>tm1),
\# take the sum of entries from prop+1 to ncol,
\# allocate it to other entries from 1 to prop
\# depending their proportions in their own sum;
\# replace entries from prop+1 to ncol with 0.
for (i in 1:(tm1.n-1)) {
  prop.tf2gtm1.prob[i, ] <- c(prop.tf2gtm1.prob[1, 1:(tf2.n-prop)] +
    sum(prop.tf2gtm1.prob[1, (tf2.n-prop+1):tf2.n])*tf2gtm1.prob[1, 1:(tf2.n-prop)]/
    (sum(prop.tf2gtm1.prob[1, 1:(tf2.n-prop)])),
    rep(0, prop))
}
\# compute new p(tf2>tm2|tf1>tm1)
acenprop.tf2gtm1.prob <- matrix(rep(0, (tm1.n-1)*tm2.n),
  nrow=tm1.n-1, ncol=tm2.n)
for (i in 1:(tm1.n-1)) {
  for (j in 1:(tm2.n-1)) {
    acenprop.tf2gtm1.prob[i, j] <- sum(acenprop.tf2gtm1.prob[i, (j+1):tm2.n])
  }
}
par(mfrow=c(3,2))
plot(tf1.poss, tf2gtf1.lower.prob, type="l", col="black", lty=1,
xlab="Tf2", ylab="p(tf2|tf1>1)"
lines(tf2.poss, tf2gtm1.prob[10, ], col="red", lty=2)
lines(tf2.poss, prop.tf2gtm1.prob[10, ], col="blue", lty=3)
abline(v=tf2.poss[which.max(tf2gtf1.lower.prob)], col="black", lty=1)
abline(v=tf2.poss[which.max(tf2gtm1.prob[10, ])], col="red", lty=2)
abline(v=tf2.poss[which.max(prop.tf2gtm1.prob[10, ])], col="blue", lty=3)
plot(tf1.poss, tf2gtf1.lower.prob, type="l", col="black", lty=1,
plot(tf1.poss, tf2gtf1.lower.prob, type="l", col="black", lty=1, 
    xlab="Tf2", ylab="p(tf2|tf1>2)"
lines(tf2.poss, tf2gtm1.prob[20, ], col="red", lty=2)
lines(tf2.poss, prop.tf2gtm1.prob[20, ], col="blue", lty=3)
abline(v=tf2.poss[which.max(tf2gtf1.lower.prob)], col="black", lty=1)
abline(v=tf2.poss[which.max(tf2gtm1.prob[20, ])], col="red", lty=2)
abline(v=tf2.poss[which.max(prop.tf2gtm1.prob[20, ])], col="blue", lty=3)
plot(tf1.poss, tf2gtf1.lower.prob, type="l", col="black", lty=1, 
    xlab="Tf2", ylab="p(tf2|tf1>3)"
lines(tf2.poss, tf2gtm1.prob[30, ], col="red", lty=2)
lines(tf2.poss, prop.tf2gtm1.prob[30, ], col="blue", lty=3)
abline(v=tf2.poss[which.max(tf2gtf1.lower.prob)], col="black", lty=1)
abline(v=tf2.poss[which.max(tf2gtm1.prob[30, ])], col="red", lty=2)
abline(v=tf2.poss[which.max(prop.tf2gtm1.prob[30, ])], col="blue", lty=3)
plot(tf1.poss, tf2gtf1.lower.prob, type="l", col="black", lty=1, 
    xlab="Tf2", ylab="p(tf2|tf1>4)"
lines(tf2.poss, tf2gtm1.prob[40, ], col="red", lty=2)
lines(tf2.poss, prop.tf2gtm1.prob[40, ], col="blue", lty=3)
abline(v=tf2.poss[which.max(tf2gtf1.lower.prob)], col="black", lty=1)
abline(v=tf2.poss[which.max(tf2gtm1.prob[40, ])], col="red", lty=2)
abline(v=tf2.poss[which.max(prop.tf2gtm1.prob[40, ])], col="blue", lty=3)
plot(tf1.poss, tf2gtf1.lower.prob, type="l", col="black", lty=1, 
    xlab="Tf2", ylab="p(tf2|tf1>5)"
lines(tf2.poss, tf2gtm1.prob[50, ], col="red", lty=2)
lines(tf2.poss, prop.tf2gtm1.prob[50, ], col="blue", lty=3)
abline(v=tf2.poss[which.max(tf2gtf1.lower.prob)], col="black", lty=1)
abline(v=tf2.poss[which.max(tf2gtm1.prob[50, ])], col="red", lty=2)
abline(v=tf2.poss[which.max(prop.tf2gtm1.prob[50, ])], col="blue", lty=3)
plot(tf1.poss, tf2gtf1.lower.prob, type="l", col="black", lty=1, 
    xlab="Tf2", ylab="p(tf2|tf1>5.9)"
lines(tf2.poss, tf2gtm1.prob[59, ], col="red", lty=2)
lines(tf2.poss, prop.tf2gtm1.prob[59, ], col="blue", lty=3)
abline(v=tf2.poss[which.max(tf2gtf1.lower.prob)], col="black", lty=1)
abline(v=tf2.poss[which.max(tf2gtm1.prob[59, ])], col="red", lty=2)
abline(v=tf2.poss[which.max(prop.tf2gtm1.prob[59, ])], col="blue", lty=3)
# p(tf1) vs p(tf2|tf1>tm1*=0.5)(CM) vs p(tf2|tf1>tm1*=0.6)(PM)
par(mfrow=c(1,1))
plot(tf2.poss, prop.tf2gtm1.prob[6, ], type="l", col="blue", lty=2, lwd=2.5, 
    xlab="Tf2", ylab="p(tf2|tf1>tm1*)"
lines(tf2.poss, tf2gtm1.prob[5, ], col="red", lty=2, lwd=2.5)
lines(tf1.poss, tf2gtf1.lower.prob, col="black", lty=1, lwd=2.5)
legend(3.5, 0.30, 
    c("p(tf1)", "p(tf2|tf1>tm1*=0.5) (CM)", "p(tf2|tf1>tm1*=0.6) (PM)"), 
    lty=c(1, 2, 2),
# dynamic programming

# calculate expected cost per unit time at chance node CN21 and CN22, i.e., euc21 and euc22.

# euc21.upper is a cost matrix related to tf1 and tf2<=tm2, 
# for each row (given a possible tf1), 
# each entry is the cost given a possible tm2; 
# e.g., the euc21.upper[1, 2] is the average cost per unit time 
# when tf1=tf1.poss[1], and tf2=tf2.poss[1], tf2.poss[2], 
# where tm2=tm2.poss[2], 
# i.e., euc21.upper[1, 2]=utility(tf1=tf1.poss[1], tf2=tf2.poss[1])*utility(tf1=tf1.poss[1], tf2=tf2.poss[2]).

# euc21.upper <- matrix(rep(0, tf1.n*tm2.n), nrow=tf1.n, ncol=tm2.n)
for (i in 1:tf1.n) {
  euc21.upper[i, 1] <- (1+alpha)*(Cf+Cr)/(tf1.poss[i]+tf2.poss[1])
  for (j in 2:tm2.n) {
  }
}

# euc21.lower is a cost matrix related to tf1 and tf2>tm2, 
# for each row (given a possible tf1), 
# each entry is the cost given a possible tm2; 
# e.g., the euc21.lower[1, 2] is the average cost per unit time 
# when tf1=tf1.poss[1] and tm2=tm2.poss[2], 
# i.e., euc21.lower[1, 2]=utility(tf1=tf1.poss[1], tm2=tm2.poss[2]).

euc21.lower <- matrix(rep(0, tf1.n*tm2.n), nrow=tf1.n, ncol=tm2.n)
for (i in 1:tf1.n) {
  for (j in 1:(tm2.n-1)) {
    euc21.lower[i, j] <- (Cf+Cr+alpha*Cm)/(tf1.poss[i]+tm2.poss[j])
    for (centf2gtf1.prob[i, j]
  }
}

# add the two matrices above, i.e., euc21.upper+euc21.lower, 
# give the matrix of expected cost per unit time at chance node CN21, 
# i.e., euc21, 
# which is associated with given tf1 and tm2, 
# e.g., euc21[1, 2] is the expected cost per unit time
# when $\text{tf}_1 = \text{tf}_1.\text{poss}[1]$ and $\text{tm}_2 = \text{tm}_2.\text{poss}[2]$.

euc21 <- euc21.upper + euc21.lower

# $\text{cn}_21.\text{tm}_2$ is the $\text{tm}_2$ that minimise the cost
# $\text{cn}_21.\text{min}$ is the minimal cost

cn21.tm2 <- rep(0, tf1.n)
cn21.min <- rep(0, tf1.n)
for (i in 1:tf1.n) {
  # for a given $\text{tf}_1$,
  # the $\text{tm}_2$ that minimise the cost and associated cost.
  cn21.tm2[i] <- tm2.poss[which.min(euc21[i,])]
  cn21.min[i] <- min(euc21[i,])
}

# $\text{euc}_22.\text{upper}$ is a cost matrix related to $\text{tm}_1$ and $\text{tf}_2 < \text{tm}_2$,
# for each row (given a possible $\text{tm}_1$),
# each entry is the cost given a possible $\text{tm}_2$;
# e.g., the $\text{euc}_22.\text{upper}[1,2]$ is the average cost per unit time
# when $\text{tm}_1 = \text{tm}_1.\text{poss}[1]$, and $\text{tf}_2 = \text{tf}_2.\text{poss}[1]$, $\text{tf}_2.\text{poss}[2]$, $\text{tm}_2 = \text{tm}_2.\text{poss}[2]$,
# i.e., $\text{euc}_22.\text{upper}[1,2] = \text{utility}(\text{tm}_1 = \text{tm}_1.\text{poss}[1], \text{tf}_2 = \text{tf}_2.\text{poss}[1]) +
# \text{utility}(\text{tm}_1 = \text{tm}_1.\text{poss}[1], \text{tf}_2 = \text{tf}_2.\text{poss}[2])$.

euc22.upper <- matrix(rep(0, (tm1.n-1)*tm2.n), nrow = tm1.n-1, ncol = tm2.n)
for (i in 1:(tm1.n-1)) {
  euc22.upper[i, 1] <- (Cm + alpha*(Cf + Cr))/
  (tm1.poss[i]+tf2.poss[1])*prop.tf2gtm1.prob[i, 1]
  for (j in 2:tm2.n) {
    euc22.upper[i, j] <- euc22.upper[i, j-1] +
    (Cm + alpha*(Cf + Cr))/(tm1.poss[i]+tf2.poss[j])*
    prop.tf2gtm1.prob[i, j]
  }
}

# $\text{euc}_22.\text{lower}$ is a cost matrix related to $\text{tm}_1$ and $\text{tf}_2 > \text{tm}_2$,
# for each row (given a possible $\text{tm}_1$),
# each entry is the cost given a possible $\text{tm}_2$;
# e.g., the $\text{euc}_22.\text{lower}[1,2]$ is the average cost per unit time
# when $\text{tm}_1 = \text{tm}_1.\text{poss}[1]$ and $\text{tm}_2 = \text{tm}_2.\text{poss}[2]$,
# i.e., $\text{euc}_22.\text{lower}[1,2] = \text{utility}(\text{tm}_1 = \text{tm}_1.\text{poss}[1], \text{tm}_2 = \text{tm}_2.\text{poss}[2])$.

euc22.lower <- matrix(rep(0, (tm1.n-1)*tm2.n), nrow = tf1.n-1, ncol = tm2.n)
for (i in 1:(tm1.n-1)) {
  for (j in 1:(tm2.n-1))
    euc22.lower[i, j] <- (1+alpha)*Cm/
    (tm1.poss[i]+tm2.poss[j])*cenprop.tf2gtm1.prob[i, j]
}

# add the two matrices above, i.e., $\text{euc}_22.\text{upper} + \text{euc}_22.\text{lower}$,
# give the matrix of expected cost per unit time at chance node CN22,
# i.e., euc22,
# which is associated with given tm1 and tm2,
# e.g., euc22[1, 2] is the expected cost per unit time
# when tm1=tm1.poss[1] and tm2=tm2.poss[2].
euc22 <- euc22.upper+euc22.lower

cn22.tm2 <- rep(0, tm1.n-1)
cn22.min <- rep(0, tm1.n)
for (i in 1:(tm1.n-1)) {
  # for a given tf1,
  # the tm2 that minimise the cost and associated cost.
  cn22.tm2[i] <- tm2.poss[which.min(euc22[i, ])]
  cn22.min[i] <- min(euc22[i, ])
}

# calculate expected cost per unit time at chance node CN1, i.e., cn1.
# compute p(tf1>tm1), i.e., centf1.prob
# p(tf1) is tf2gtf1.lower.prob
centf1.prob <- rep(0, tm1.n)
for (i in 1:(tm1.n-1)) {
  centf1.prob[i] <- sum(tf2gtf1.lower.prob[(i+1):tm1.n])
}

# upper21 is a cost vector related to tf1<=tm1,
# each element is the cost given a possible tm1;
# e.g., the upper21[2] is the average cost per unit time
# when tf1=tf1.poss[1], tf1.poss[2], where tm1=tm1.poss[2],
# i.e., upper21[2]=utility(tf1=tf1.poss[1])+utility(tf1=tf1.poss[2]).
upper21 <- rep(0, tm1.n)
upper21[1] <- (cn21.min*tf2gtf1.lower.prob)[1]
for (i in 2:tm1.n) {
  upper21[i] <- upper21[i-1]*(cn21.min*tf2gtf1.lower.prob)[i]
}

# lower22 is a cost vector related to tm1,
# each element is the cost given a possible tm1;
# e.g., the lower22[2] is the average cost per unit time
# when tm1=tm1.poss[2],
# i.e., lower22[2]=utility(tm1.poss[2]).
lower22 <- cn22.min*centf1.prob

# add two vectors, i.e., upper21+lower22
# give the vector of expected cost per unit time at chance node CN1,
# i.e., en1,
# which is associated with given tm1,
# e.g., cn1[2] is the expected cost per unit time
# when tm1=tm1.poss[2].
cn1 <- upper21+lower22

ucn1.frame <- data.frame(Tm1=tm1.poss, CN21=upper21, CN22=lower22, CN1=cn1)
# convert the data frame from "wide" format to "long" format
library(reshape2)
ucn1.long <- melt(ucn1.frame,
                   id.vars="Tm1",
                   variable.name="ChanceNodes",
                   value.name="ExpectedCost")
levels(ucn1.long$Tm1) <- tm1.poss

ggplot(data=ucn1.long,
       aes(x=Tm1, y=ExpectedCost, colour=ChanceNodes)) +
  geom_line() +
  xlab("Tm1") + ylab("Expected Cost") +
  labs(fill="Chance Nodes")

# cn1.tm1 is the tm1 that minimise the cost at chance node CN1
# cn1.min is the minimal cost at chance node CN1
cn1.tm1 <- tm1.poss[which.min(cn1)]
cn1.min <- min(cn1)
cn1.tm1.opt <- cn1.tm1 # the optimal maintenance time for phase 1

cn1.opt <- cn1.min # the minimal expected cost for the entire system

# go forward from phase 1 to phase 2
# given optimal maintenance time for phase 1, i.e., cn1.tm1.opt

# cn21.tm2.opt and cn21.opt are
# the optimal maintenance time and minimal expected cost for phase 2
# when tf1<=cn1.tm1.opt.
cn21.tm2.opt <- cn21.tm2[tf1.poss<=cn1.tm1.opt]
cn21.opt <- cn21.min[tf1.poss<=cn1.tm1.opt]

# cn22.tm2.opt and cn22.opt are
# the optimal maintenance time and minimal expected cost for phase 2
# when tf1>cn1.tm1.opt.
cn22.tm2.opt <- cn22.tm2[tm1.poss==cn1.tm1.opt]
cn22.opt <- cn22.min[tm1.poss==cn1.tm1.opt]

# print results
print(cn1.tm1.opt)
print(cn1.opt)
print(tf1.poss[tf1.poss<=cn1.tm1.opt]) # tf1 that is smaller than tm1
print(cn21.tm2.opt)
print(cn21.opt)
print(cn22.tm2.opt)
print(cn22.opt)

# myopic

# for chance node CN1

# mc1.upper is a cost vector related to tf1<=tm1,
# each element is the cost given a possible tm1;
# e.g., the mc1.upper[2] is the average cost per unit time
# when tf1=tf1.poss[1], tf1.poss[2], where tm1=tm1.poss[2],
# i.e., mc1.upper[2]=utility(tf1=tf1.poss[1])*utility(tf1=tf1.poss[2]).
mc1.upper <- rep(0, tm1.n)
mc1.upper[1] <- (((Cf+Cr)/tf1.poss)*tf2gtf1.lower.prob)[1]
for (i in 2:tm1.n) {
  mc1.upper[i] <- mc1.upper[i-1]+(((Cf+Cr)/tf1.poss)*tf2gtf1.lower.prob)[i]
}

# mc1.lower is a cost vector related to tm1,
# each element is the cost given a possible tm1;
# e.g., the mc1.lower[2] is the average cost per unit time
# when tm1=tm1.poss[2],
# i.e., mc1.lower[2]=utility(tm1.poss[2]).
mc1.lower <- (Cm/tm1.poss)*centf1.prob
mc1 <- mc1.upper + mc1.lower
mc1.tm1 <- tm1.poss[which.min(mc1)] # the optimal tm1
mc1.min <- min(mc1) # the minimal cost for CN1

# for chance node CN21

# tf1=optimal tm1
mtf1.n <- length(subset(tf1.poss, tf1.poss<=mc1.tm1))
mc21.upper <- matrix(rep(0, mtf1.n*tm2.n), nrow=mtf1.n, ncol=tm2.n)
for (i in 1:mtf1.n) {
  mc21.upper[i, 1] <- ((1+alpha)*(Cf+Cr)/(tf1.poss[i]+tf2.poss[1]))*
    tf2gtf1.prob[i, 1]
  for (j in 2:tm2.n) {
    mc21.upper[i, j] <- mc21.upper[i, j-1]*
      ((1+alpha)*(Cf+Cr)/(tf1.poss[i]+tf2.poss[j]))*
      tf2gtf1.prob[i, j]
  }
}
mc21.lower <- matrix(rep(0, mtf1.n*tm2.n), nrow=mtf1.n, ncol=tm2.n)
for (i in 1:mtf1.n) {
  for (j in 1:(tm2.n-1))
    mc21.lower[i, j] <- (Cf+Cr+alpha*Cm)/(tf1.poss[i]+tm2.poss[j])*
      centf2gtf1.prob[i, j]
}
mc21 <- mc21.upper + mc21.lower
mc21.tm2 <- rep(0, mtf1.n)
mc21.min <- rep(0, mtf1.n)
for (i in 1:mtf1.n) {
  mcn21.tm2[i] <- tm2.poss[which.min(mcn21[i, ])]
  mcn21.min[i] <- min(mcn21[i, ])
}

# for chance node CN22
# tf1 > optimal tm1
mcn22.upper <- rep(0, tm2.n)
mcn22.upper[1] <- (Cm+alpha*(Cf+Cr))/(mcn1.tm1+tf2.poss[1])*prop.tf2gtm1.prob[which.min(mcn1), 1]
for (i in 2:tm2.n) {
  mcn22.upper[i] <- mcn22.upper[i-1]*
  (Cm+alpha*(Cf+Cr))/(mcn1.tm1+tf2.poss[i])*prop.tf2gtm1.prob[which.min(mcn1), i]
}
mcn22.lower <- (1+alpha)*Cm/(mcn1.tm1+tm2.poss)*cenprop.tf2gtm1.prob[which.min(mcn1), ]
mcn22 <- mcn22.upper + mcn22.lower
mcn22.tm2 <- tm2.poss[which.min(mcn22)]
mcn22.min <- min(mcn22)

print(mcn1.tm1)
print(mcn1.min)
print(mcn21.tm2)
print(mcn21.min)
print(mcn22.tm2)
print(mcn22.min)

####
# simulation
set.seed(2014)

n <- 1000 # number of theta
sim.theta <- rtruncnorm(n, a=1, b=Inf, mean=2, sd=1)

# generate tf1 and tf2 which follow Weibull distribution
# given shape parameter theta
# tf1 <- rep(0, rep=n)
tf1 <- rep(0, rep=n)
tf2 <- rep(0, rep=n)

k <- 1
while (k<=n) {
  x <- rweibull(1, shape=sim.theta[k], scale=1)
  y <- rweibull(1, shape=sim.theta[k], scale=1)
  # x <- rgamma(1, shape=sim.theta[k], scale=1)
  # y <- rgamma(1, shape=sim.theta[k], scale=1)
  if ((round(x, 1) != 0) && (round(y, 1) != 0)) {
    tf1[k] <- x
    tf2[k] <- y
  }
  k <- k + 1
}
# compute simulated cost for chance node CN1

cost.cn <- rep(0, n)
for (i in 1:n) {
  if (round(tf1[i], 1) <= cn1.tm1.opt) {
    if (round(tf2[i], 1) <= cn21.tm2.opt[round(tf1[i], 1)*10]) {
      cost.cn[i] <- (1+alpha)*(Cf+Cr)/(round(tf1[i], 1)+round(tf2[i], 1))
    }
  } else if (round(tf2[i], 1) > cn21.tm2.opt[round(tf1[i], 1)*10]) {
    cost.cn[i] <- (Cf+Cr+alpha*Cm)/(round(tf1[i], 1)+cn21.tm2.opt[round(tf1[i], 1)*10])
  }
  else if (round(tf1[i], 1) > cn1.tm1.opt) {
    if (round(tf2[i], 1) <= cn22.tm2.opt) {
      cost.cn[i] <- (Cm+alpha*(Cf+Cr))/(cn1.tm1.opt+round(tf2[i], 1))
    } else if (round(tf2[i], 1) > cn22.tm2.opt) {
      cost.cn[i] <- (1+alpha)*Cm/(cn1.tm1.opt+cn22.tm2.opt)
    }
  }
}

# compute simulated expected cost for chance node CN1

mean(cost.cn)

pm_2tp.R
References


