Applying the General Path Model to Estimation of Remaining Useful Life

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\textbf{ABSTRACT}

The ultimate goal of most prognostic systems is accurate prediction of the remaining useful life of individual systems or components based on their use and performance. This class of prognostic algorithms is termed effects-based, or Type III, prognostics. A unit-specific prognostic model, called the General Path Model, involves identifying an appropriate degradation measure to characterize the system's progression to failure. A functional fit of this parameter is then extrapolated to a pre-defined failure threshold to estimate the remaining useful life of the system or component. This paper proposes a specific formulation of the General Path Model with dynamic Bayesian updating as one effects-based prognostic algorithm. The method is illustrated with an application to the prognostics challenge problem posed at PHM '08.

\textbf{1. INTRODUCTION}

Prognostics is a term given to equipment life prediction techniques and may be thought of as the "holy grail" of condition-based maintenance. Prognostics can play an important role in increasing safety, reducing downtime, and improving mission readiness and completion. Prognostics is one component in a full health management system (Figure 1). Health monitoring systems commonly employ several modules, including but not limited to: system monitoring, fault detection, fault diagnostics, prognostics, and management (Kothamasu et al., 2006 and Callan et al., 2006). System monitoring and fault detection modules are used to determine if a component or system is operating in a nominal and expected way. If a fault or anomaly is detected by the monitoring system, the diagnostic system determines the type, and in some cases, the severity of the fault. The prognostics module uses this information to estimate the Remaining Useful Life (RUL) of the system or component along with associated confidence bounds. With this information in hand, system operation may be adjusted to mitigate the effects of failure or to slow the progression of failure, thereby extending the RUL to some later point, such as a previously scheduled maintenance activity or the end of the planned mission.

Prognostic system development has been a daunting task for several reasons. One is that mission critical systems are rarely allowed to run to failure once degradation has been detected. This makes the existence of degradation data rare and the development of degradation-based models difficult. However, current individual-based, empirical prognostic techniques necessitate the availability of a population of exemplar degradation paths for each fault mode of interest. In some cases, physical models may be developed to generate simulated degradation data or may be used in a model-based prognostics framework to infer RUL (Pecht and...
Dasgupta, 1995, Valentin et al., 2003, and Oja et al. 2007). Second, if the components are subject to common fault modes which lead to failure, these fault modes are often designed out of the system through a proactive continuous improvement process. Third, very few legacy systems have the instrumentation required for accurate prognostics. In the absence of such instrumentation, accurate physics of failure models may be used to identify key measurements and systems may be re-instrumented.

This research focuses on RUL estimation for soft failures. These failures are considered to occur when the degradation level of a system reaches some predefined critical failure threshold, e.g. light output from fluorescent light bulbs decreases below a minimum acceptable level or car tire tread is thinner than some pre-specified depth. These failures generally do not concur with complete loss of functionality; instead, they correspond with the time when an operator is no longer confident that equipment will continue to work to its specifications.

Traditional reliability analysis, termed Type I prognostics, uses only failure time data to estimate a time to failure distribution (Hines et al., 2007). This class of algorithms characterizes the average lifetime of an average component operating in historically average conditions; it does not consider any unit-specific information beyond the current run time. As components become more reliable, few failure times may be available, even with accelerated life testing. Although failure time data become more sporadic as equipment reliability rises, often other measures are available which may contain some information about equipment degradation, such as crack length, tire pressure, or pipe wall thickness. Lu and Meeker (1993) developed the General Path Model (GPM) to model equipment reliability using these degradation measures, or appropriate functions thereof, moving reliability analysis from failure-time analysis to failure-process analysis. The GPM assumes that there is some underlying parametric model to describe component degradation. The model may be derived from physical models or from available historical degradation data. Typically, this model accounts for both population (fixed) effects and individual (random) effects.

Although GPM was originally conceived as a method for estimating population reliability characteristics, such as the failure time distribution, it has since been extended to individual prognostic applications (Upadhyaya et al., 1994). Most commonly, the fitted model is extrapolated to some known failure threshold to estimate the RUL of a particular component. This is an example of an Effects-based, or Type III, prognostic algorithm (Hines et al., 2007). This class of algorithms estimates the RUL of a specific component or system operating in its specific environment; it is the ultimate goal of prognostics for most mission critical components.

The following sections will present GPM theory including the original methodology for reliability applications and the extension to prognostics. In addition, a short discussion of dynamic Bayesian updating methods to incorporate prior information is given. Finally, an application of the proposed GPM methodology to the 2008 PHM Challenge problem is presented.

2. METHODOLOGY

As suggested by the “No Free Lunch” Theorem, no one prognostic algorithm is ideal for every situation (Koppen, 2004). A variety of models have been developed for application to specific situations or specific classes of systems. The efficacy of these algorithms for a new process depends on the type and quality of data available, the assumptions inherent in the algorithm, and the assumptions which can validly be made about the system. This research focuses on the general path model, an algorithm which attempts to characterize the lifetime of a specific component based on measures of degradation collected or inferred from the system.

2.1. The General Path Model

Lu and Meeker (1993) first proposed the General Path Model (GPM), an example of degradation modeling, to move reliability analysis methods from time-of-failure analysis to process-of-failure analysis. Traditional methods of reliability estimation use failure times recorded during normal use or accelerated testing to estimate a time of failure (TOF) distribution for a population of identical components. In contrast, GPM uses degradation measures to estimate the TOF distribution. The use of historical degradation measures allows for the direct inclusion of censored data, which gives additional information on unit-wise variations in a population.

GPM analysis begins with some assumption of an underlying functional form of the degradation path for a specific fault mode. The degradation of the $i^{th}$ unit at time $t_j$ is given by:

$$y_{ij} = \eta(t_j, \varphi, \theta_i) + \varepsilon_{ij}$$  \hspace{1cm} (1)

where $\varphi$ is a vector of fixed (population) effects, $\theta_i$ is a vector of random (individual) effects for the $i^{th}$ component, and $\varepsilon_{ij} \sim N(0,\sigma^2_e)$ is the standard measurement error term. Application of the GPM methodology involves several assumptions. First, the degradation data must be describable by a function, $\eta$; this function may be derived from physics-of-failure models or from the degradation data itself. In order to fit this model, the second
assumption is that historical degradation data from a population of identical components or systems are available or can be simulated. This data should be collected under similar use (or accelerated test) conditions and should reasonably span the range of individual variations between components. Because GPM uses degradation measures instead of failure times, it is also not necessary that all historical units are run to failure; censored data contain information useful to GPM forecasting. The final assumption of the GPM model is that there exists some defined critical level of degradation, $D$, which indicates component failure; this is the point beyond which the component will no longer perform its intended function with an acceptable level of reliability. Therefore, some components should be run to failure, or to a state considered failure, in order to quantify this degradation level. Alternatively, engineering judgment may be used if the nature of the degradation parameter is explicitly known.

Several methods are available to estimate the degradation model parameters, $\varphi$ and $\theta$. In some cases, the population parameters may be known in advance, such as the initial level of degradation. If the population parameters are unknown, estimation of the vector of population characteristics, $\varphi$, is trivial; by fitting the model to each exemplar degradation path, the fixed effects parameters can be taken as the mean of the fitted values for each unit. The variance of these estimates should be examined to ensure that the parameters can be considered to be fixed. If significant variability is present, the parameters should be considered random and moved to the $\theta$ vector. A two-stage method of parameter estimation was proposed by Lu and Meeker (1993) to estimate distribution parameters for the random effects.

In the first stage, the degradation model is fit to each degradation path to obtain an estimate of $\theta$ for that unit; these $\theta$s are referred to as stage-1 estimates. It is convenient to assume that the stage-1 estimates, or an appropriate transformation, $\Theta=H(\theta)$, is normally (or multivariate normally) distributed so that the random effects can be fully described using only a mean vector and variance-covariance matrix without significant loss of information. This assumption usually holds for large populations as a result of the central limit theorem; however, if it is not justifiable, the GPM methodology can be extended in a natural way to allow for other random effects distributions.

In the second stage, the stage-1 estimates (or an appropriate transformation thereof) are combined to estimate $\varphi$, $\mu_\theta$, and $\Sigma_\theta$. At this stage, if for any random parameter, $m$, the variance $\sigma_m^2$ is effectively zero, this parameter should be considered a fixed effects parameter and should be removed from the random parameter distribution.

In their seminal paper, Lu and Meeker (1993) describe Monte Carlo methods for using the GPM parameter estimates to estimate a time to failure distribution and corresponding confidence intervals. Because the focus of this paper is estimating time to failure of an individual component and not the failure time distribution of the population of components, these methods will not be described here.

Several limitations and areas of future work of the GPM are identified by Meeker et al. (1998). Some of these areas have been addressed in work by other authors. First, the authors cite the need for more accurate physics of failure models. While such models are helpful for understanding degradation mechanisms, they may not be strictly necessary for RUL estimation. In fact, if exemplar data sets cover the range of likely degradation paths, it may be adequate to fit a function which does not explain failure modes but accurately models the underlying relationships. With this idea, neural networks have been applied to GPM reliability analysis (Chinnam, 1999 and Girish et al., 2003).

In addition, the GPM was originally developed for reliability analysis of only one fault mode. In practical applications, the system of interest may consist of several components each with different fault modes, or of one component with several possible, even simultaneous fault modes. These multiple degradation paths may be uncorrelated, in which case extension of the GPM is trivial: reliability of a component for all degradation modes is simply the product of the individual reliabilities, and RUL can be considered some function of the RULs for each fault mode, such as the minimum. If, however, the degradation measures are correlated, extension of the GPM is more complicated. For example, in the case of tire monitoring, several degradation measures may contain information about tire reliability, including tread thickness, tire pressure, tire temperature and wall material characteristics. However, it is easy to see that these measures may be correlated; a higher temperature would cause a higher pressure etc. The case of multiple, competing degradation modes is beyond the scope of the current work. A discussion of the problem can be found in Wang and Coit (2004).

### 2.2. GPM for Prognostics

The GPM reliability methodology has a natural extension to estimation of remaining useful life of an individual component or system; the degradation path model, $y$, can be extrapolated to the failure threshold, $D$, to estimate the component’s time of
failure. This type of degradation extrapolation was proposed early on by Upadhyaya et al. (1994). In that work, the authors used both neural networks and nonlinear regression models to predict the RUL of an induction motor. The prognostic methodology used for the current research is described below.

First, exemplar degradation paths are used to fit the assumed model. The stage-1 parameter estimates are used to evaluate the random-effects distributions, to determine the mean population random effects, the mean time to failure (MTTF) and their associated standard deviations, and to estimate the noise variance in the degradation paths. The MTTF distribution can be used to estimate the time of failure for any component which has not yet been degraded.

As data are collected during use, the degradation model can be fit for the individual component. This specific model can be used to project a time of failure for the component. Because of noise in the degradation signal, the projected time of failure is not perfect. A prediction interval (PI) about the estimated parameters can be evaluated as:

$$\theta \in \left[ \hat{\theta} - t_{n-1,\alpha/2} \frac{s}{\sqrt{1 + \frac{1}{n} \hat{\theta}}}, \hat{\theta} + t_{n-1,\alpha/2} \frac{s}{\sqrt{1 + \frac{1}{n} \hat{\theta}}} \right]$$

where $t_{n-1,\alpha/2}$ is the Student’s t-distribution, $n$ is the number of observations used to fit the model, and $s$ is the standard deviation of the degradation model parameters for normally distributed, uncorrelated parameters; if this assumption is not met, the method can be extended to estimate PIs for other distributions. The standard deviation of the parameters can be estimated through traditional linear regression techniques. The range of model parameters can be used to project an PI about the estimated time of failure.

The methodology described considers only the data collected on the current unit to fit the degradation model. However, prior information available from historic degradation paths can be used for initial model fitting, including the mean degradation path and associated distributions. This data can provide valuable knowledge for fitting the degradation model of an individual component, particularly when only a few data points have been collected or the collected data suffer from excessive noise. The following section outlines a dynamic Bayesian updating method for including prior information in degradation model fitting.

### 2.3. Incorporating Prior Information

The current research investigates using Bayesian methods to include prior information for linear regression problems. However, as discussed above, the GPM methodology can be applied to nonlinear regression problems as well as other parametric modeling techniques such as neural networks. Other Bayesian methods must be applied to these types of models, but such application is beyond the scope of the current research. For a complete discussion of Bayesian statistics including other Bayesian update methods, the interested reader is referred to Carlin and Louis (2000) and Gelman et al. (2004). In addition, work by Robinson and Crowder (2000) focuses on Bayesian methods for nonlinear regression reliability models.

A brief review of Bayesian update methods for linear regression is given here; a more complete discussion can be found in Lindley and Smith (1972) as well as the texts cited above. Bayesian updating is a method for combining prior information about the set of model parameters with new data observations to give a posterior distribution of the model parameters (Figure 2). This allows both current observation and past knowledge to be considered in model fitting.

A linear regression model is given by:

$$Y = bX$$

The model parameters are estimated using the pseudo-inverse formula as:

$$b = (X^T \Sigma_y^{-1} X)^{-1} X^T \Sigma_y^{-1} Y$$

$$\Sigma_y = \begin{bmatrix} \sigma_{y1} & 0 & \cdots & 0 \\ 0 & \sigma_{y2} & \cdots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \cdots & \sigma_{ym} \end{bmatrix}$$

where $\Sigma_y$ is the variance-covariance noise matrix, which gives an indication of the accuracy of each entry in the $Y$-vector. It is important to note that the linear regression model is not necessarily a linear model, but is linear-in-parameters. The data matrix...
can be populated with any function of degradation measures, including higher order terms, interaction terms, and functions such as \( \sin(x) \) or \( e^x \). If prior information is available for a specific model parameter, i.e. \( \beta_j \sim N(\beta_{jm}, \sigma_{\beta_j}^2) \), then the matrix \( X \) should be appended with an additional row with value one at the \( j^{th} \) position and zero elsewhere, and the \( Y \) matrix should be appended with the \textit{a priori} value of the \( j^{th} \) parameter.

\[
Y^* = \begin{bmatrix}
  y_1 \\
  y_2 \\
  \vdots \\
  y_n \\
  \beta_{jm}
  \end{bmatrix} \quad X^* = \begin{bmatrix}
  x_{11} & x_{12} & \cdots & x_{1n} \\
  x_{21} & x_{22} & \cdots & x_{2n} \\
  \vdots & \vdots & \ddots & \vdots \\
  x_{n1} & x_{n2} & \cdots & x_{nn} \\
  0 & \cdots & 1 & 0
  \end{bmatrix}
\]

Finally, the variance-covariance matrix is augmented with a final row and column of zeros, with the variance of the \textit{a priori} information in the diagonal element.

\[
\Sigma^*_j = \begin{bmatrix}
  \sigma_{y_1}^2 & 0 & \cdots & 0 & 0 \\
  0 & \sigma_{y_2}^2 & \cdots & 0 & \vdots \\
  \vdots & \vdots & \ddots & \vdots & \vdots \\
  0 & \cdots & 0 & \sigma_{y_n}^2 & 0 \\
  0 & 0 & \cdots & 0 & \sigma_{\beta_{jm}}^2
  \end{bmatrix}
\]

If knowledge is available about multiple regression parameters, the matrices should be appended multiple times with one row for each parameter.

It is convenient to assume that the noise in the degradation measurements is constant and uncorrelated. Some \textit{a priori} knowledge of the noise variance is available from the exemplar degradation paths. If this assumption is not valid for a particular system, then other methods of estimating the noise variance may be used; however, it has been seen that violating this assumption does not have a significant impact on RUL estimation. In addition, it is also convenient to assume that the noise measurements are uncorrelated across observations of \( y \); this allows the variance-covariance matrix to be a diagonal matrix consisting of noise variance estimates and \textit{a priori} knowledge variance estimates. If this assumption is not valid, including covariance terms is trivial; again, these terms can be estimated from historical degradation paths.

After \textit{a priori} knowledge is used in conjunction with \( n \) current data observations to obtain a posterior estimate of degradation parameters, this estimate becomes the new prior distribution for the next estimation of regression parameters. The variance of this new knowledge is estimated as:

\[
\sigma_{\text{post}}^2 = \frac{1}{2} \sigma_{\text{prior}}^2 + \frac{n}{2} \sigma_{\text{data}}^2
\]

The Bayesian information may be used to dynamically update the model fit as new data become available for each desired RUL estimate.

### 2.4 Combined Monitoring and Prognostic Systems

Figure 3 shows a combined monitoring, fault detection, and prognostics system similar to the one used in this research. The monitoring system employs an Auto-Associative Kernel Regression (AAKR) model for monitoring and the Sequential Probability Ratio Test (SPRT) for fault detection. Both of these methods are described in broad detail below. The interested reader is referred to (Hines et al., 2008 and Garvey et al., 2007) for a more complete discussion of AAKR and (Wald, 1943) for SPRT.

Auto-Associative models can generally be considered an error correction technique. These models compare a new observation to those seen in the past to estimate how the system “should” be running. These corrected predictions can be compared to the measured data to identify faulted operation. Several auto-associative architectures are available, including auto-associative neural networks, auto-associative kernel regression, and multivariate state estimation technique (Hines et al., 2008). This research employs the AAKR algorithm for system monitoring.

AAKR is a non-parametric, empirical technique. Exemplar historical observations of system operation are stored in a data matrix. As a new observation is collected, it is compared to each of the following:
the exemplar observations to determine how similar the new observation is to each of the exemplars. This similarity is quantified by evaluating the distance between the new observation and the exemplar. Most commonly, the Euclidean distance is used:

\[ d_i = \sqrt{\sum_{j=1}^{m} (X_j - x_{i,j})^2} \]  

(8)

where \( d_i \) is the distance between the new observation, \( X \), and the \( i^{th} \) exemplar, \( x_i \). The distance is converted to a similarity measure through the use of a kernel. Many kernels are available; this research employs the Gaussian kernel:

\[ s_i = \exp \left( \frac{d_i^2}{h^2} \right) \]  

(9)

where \( s_i \) is the similarity of the new observation to the \( i^{th} \) exemplar and \( h \) is the kernel bandwidth, which controls how close vectors must be to be considered similar. Finally, the “corrected” observation value is calculated as a weighted average of the exemplar observations:

\[ \hat{X}_i = \frac{s_i X_i}{s_i} \]  

(10)

Monitoring system residuals are then generated as the difference between the actual observation and the error-corrected prediction. These residuals are used with a SPRT to determine if the system is operating in a faulted or nominal condition. As the name suggests, the SPRT looks at a sequence of residuals to determine if the time series of data is more likely from a nominal distribution or a pre-specified faulted distribution. As new observations are made, the SPRT compares the cumulative sum of the log-likelihood ratio:

\[ s_i = s_{i-1} + \log \frac{g_i}{\hat{X}_i} \]  

(11)

to two thresholds, which depend on the acceptable false positive and false negative fault rates:

\[ a \log \frac{1}{\alpha} \]

\[ b \log \frac{1}{\beta} \]  

(12)

where \( \alpha \) is the acceptable false alarm (false positive) rate and \( \beta \) is the acceptable missed alarm (false negative) rate. For this research, false alarm and missed alarm rates of 1% and 10% respectively are used. If \( s_i < a \), then the null hypothesis cannot be rejected; that is, the system is assumed to be operating in a nominal condition. If \( s_i > b \), then the null hypothesis is rejected; that is, the system is assumed to be operating in a faulted condition. When a determination is made, the sum, \( s_i \), is reset to zero and the test is restarted.

After a fault is detected in the system, the prognostic system can be engaged to determine the RUL for the system. As discussed above, the GPM methodology uses a measure of system degradation, called a prognostic parameter, to make prognostic estimates. An ideal prognostic parameter has three key qualities: monotonicity, prognosability, and trendability.

Monotonicity characterizes the underlying positive or negative trend of the parameter. This is an important feature of a prognostic parameter because it is generally assumed that systems do not undergo self-healing, which would be indicated by a non-monotonic parameter. This assumption is not valid for some components such as batteries, which may experience some degree of self repair during short periods of nonuse, but it tends to hold for mechanical systems or for complex systems as a whole.

Prognosability gives a measure of the variance in the critical failure value of a population of systems. A wide spread in critical failure values can make it difficult to accurately define a critical failure threshold and to extrapolate a prognostic parameter to failure. Prognosability may be very susceptible to noise in the prognostic parameter, but this effect may be reduced by traditional variance reduction methods such as parameter bagging and data denoising.

Finally, trendability indicates the degree to which the parameters of a population of systems have the same underlying shape and can be described by the same functional form.

The population of noise-free prognostic parameters shown in Figure 4 exhibits the three desired features. The parameters are monotonic: they all generally trend upward through time. They are prognosable: the parameter value at failure for each unit is at approximately the same value, as indicated by the red markers. Finally, they are trendable: each parameter appears to follow the same upward exponential or quadratic trend.

Monitoring system residuals, or combinations of residuals, are natural candidates for prognostic parameters because they inherently measure the deviation of a system from normal operation. The following section investigates the application of this monitoring/prognostic method to the 2008 PHM challenge problem.
3. PHM '08 Challenge Application

This section presents an application of the proposed GPM prognostic method to the PHM Challenge data set. The efficacy of the method is analyzed based on the given cost function for the 218 test cases. RUL estimates far from the actual value are penalized exponentially. The cost function is asymmetric; RUL predictions greater than the actual value are penalized more heavily than those which predict failure before it happens. The cost for each case is given by the following formula:

\[ \text{score}(d < 0) = \exp\left(\frac{-d}{13}\right) - 1 \]

\[ \text{score}(d > 0) = \exp\left(\frac{d}{10}\right) - 1 \]

where \( d \) is the difference between the estimated and the actual RUL. If \( d \) is negative, then the algorithm underestimates the RUL leading one to end operation before failure occurs; if \( d \) is positive, then the algorithm overestimates the RUL and results in a greater penalty because one may attempt to operate the component longer than possible and thereby experience a failure. The following sections give a brief description of the simulated data set used for the challenge problem, then outline the data analysis and identification of an appropriate prognostic parameter for GPM trending. Finally, the application of the GPM method and Bayesian updating are presented with final results given for the described method. The performance of the GPM algorithm with and without Bayesian updating is compared.

3.1. PHM Challenge Data Set Description

The PHM Challenge data set consists of 218 cases of multivariate data that track from nominal operation through fault onset to system failure. Data were provided which modeled the damage propagation of aircraft gas turbine engines using the Commercial Modular Aeropropulsion System Simulation (C-MAPSS). This engine simulator allows faults to be injected in any of the five rotating components and gives output responses for 58 sensed engine variables. The PHM Challenge data set included 21 of these 58 output variables as well as three operating condition indicators. Each simulated engine was given some initial level of wear which would be considered within normal limits, and faults were initiated at some random time during the simulation. Fault propagation was assumed to evolve in an exponential way based on common fault propagation models and the results seen in practice. Engine health was determined as the minimum health margin of the rotating equipment, where the health margin was a function of efficiency and flow for that particular component; when this health indicator reached zero, the simulated engine was considered failed. The interested reader is referred to Saxena et al. (2008) for a more complete description of the data simulation.

The data have three operational variables – altitude, Mach number, and TRA – and 21 sensor measurements. Initial data analysis resulted in the identification of six distinct operational settings; based on this result, the operating condition indicators were collapsed into one indicator which fully defined the operating condition of the engine for a single observation (flight). In addition, ten sensed variables were identified whose statistical properties changed through time and were well correlated (linear correlation coefficient of at least 0.7, shown in Figure 5) to each other. In this way, the 24 sensor data set was reduced to 11 variables, with original variable numbers: 1 (the operating condition indicator), 5, 6, 7, 12, 14, 17, 18, 20, 23, and 24.

The GPM method uses degradation information, either directly measured or inferred, to estimate the system RUL. Initial analysis of the raw data does...
not reveal any trendable degradation parameter. That is, no sensed measurement has an identifiable trend toward failure. Figure 6 is a plot of the eleven variables that were determined to statistically change with time. These variables were used to develop a monitoring and prognostics system. Visual inspection of the data does not indicate any obvious trends toward failure. The monitoring system provides much greater sensitivity to subtle changes that may be indicative of failure.

3.2. Monitoring and Prognostics Results

An AAKR model is used to determine the expected values of the eleven variables of interest. The baseline model is developed using the first 15% of each run as training data; this assumes that faults occur after at least 15% of each run is completed. This assumption is not universally valid, but seems to be reasonable for this data set. Based on the AAKR predictions, a residual is calculated between the nominal prediction and the actual value. These residuals are potential candidates for inclusion into the degradation parameter.

Figure 7 is a plot of sensed variable 17 and the corresponding residual for five of the training cases. The final value for each of the five cases is indicated in the lower plot by red asterisks. In this case, the residual does not provide a useful prognostic parameter. The residual is not trendable; that is, the five cases show several distinct residual shapes. In addition, the residual's prognosability is not high. The residuals end at very different values for each case. This could be indicative of different failure modes, but is not directly useful as a degradation parameter. For the purpose of this analysis, it is assumed that all fault modes may be lumped together into one prognostic model. Therefore, a single degradation parameter is desired to prognose all systems.

Several of the residuals grow in a similar manner with time for all the units and have failure values without much variation. These residuals can be used as a degradation parameter by trending them through regression and extrapolating the functional fit to some degradation threshold to give an estimate of RUL. The top plot in Figure 8 shows one such sensed variable while the lower plot is the residual between the predicted and measured value. It shows the unfiltered residual for 5 different training cases. This variable is a good prognostic parameter because the corresponding monitoring system residuals of each of the training systems have the same basic shape and failures occur at approximately the same negative value. The task is to model the degradation parameter and predict the failure point when only a subset of the case is given. Five residuals were found to have a similar shape with well clustered values at failure.
By combining the five degradation parameters with similar shapes, an average parameter was developed. The five residuals are combined in a weighted average, where each residual weight is inversely proportional to its variance. Figure 9 is a plot of one of the candidate residuals, and Figure 10 is the averaged degradation parameter resulting from a fusion of the five residuals for all 218 cases. As the plots show, the residual parameters have very similar shapes for each training case. However, the single residual is contaminated with greater noise and has a relatively larger spread in the final parameter value. By combining several similar residuals, the spread in the failure value relative to the range of the parameter is significantly reduced, as shown in the second figure. This is sometimes referred to as parameter bagging and is a common variance reduction technique.

A second order polynomial model can be used to model the degradation parameter. While an exponential model may be more physically appropriate, the quadratic model is more robust to noise and better describes the data fit for the chosen prognostic parameter. For the methodology proposed, the model must be linear in parameters; however, simple exponential models, such as $y = \exp(ax + b)$ parameterized as $\ln(y) = ax + b$, cannot be used with negative $y$-values, because the natural logarithm of a negative number is undefined in the real number system. This adds unnecessary complexity to the modeling method. Quadratic equations, on the other hand, are naturally linear in parameters and can be used without significant concern for the effects of noise on the model fit. Shifting the prognostic parameter to the positive quadrant by adding 25.0 to every value eliminates the problem of taking the logarithm of negative values; however, the quadratic fit results in a lower fitting error than the exponential fit, with mean squared errors of 1.53 and 2.33 respectively. Because of its robustness to noise and reduced modeling error, the quadratic fit is chosen for this research.

Figure 11 gives an example of a polynomial fit of the prognostic parameter with the time the model crosses the critical failure threshold indicated. The threshold of -13.9 was chosen as the upper 95% level of the distribution of failure values for the known failed cases. This gives an estimated system reliability of 95%, which is a conservative estimate of failure time and reduces the possibility of overestimating RUL leading to in-service failure. The time between the last sample and the estimated time of failure is the estimate of RUL, as indicated by the blue area. For this case, the estimated RUL is exactly correct, with an estimated remaining life of 36 cycles.

The GPM methodology presented works well if many observations are available to fit a model to the degradation parameter as in case 106 shown above. However, when only a few observations have been collected, the model fit is highly susceptible to noise. To counteract this, the Bayesian updating
The method described previously is used to include prior information about the degradation parameter fit.

For the current problem, quadratic models (eqn. 14) were fit to the full degradation parameter for each of the 218 training cases.

\[ d = p_1 t^2 + p_2 t + p_3 \]  

(14)

The means and standard deviations for the three parameters \((p_i)\) are given in Table 1. The parameters should be considered random effects because their standard deviations represent a significant proportion of the mean parameter value. The large variance seen in \(p_3\) is assumed to correspond to the random level of initial degradation. The variance of the degradation parameter can be estimated from the training examples by smoothing each example path and subtracting the smoothed path from the actual path. This gives an estimate of the noise; the noise variance can be estimated directly as the variance of this data set. For this data, the noise variance in the degradation parameter is estimated to be 0.0588 units.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Mean</th>
<th>Std Dev</th>
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</thead>
<tbody>
<tr>
<td>(p_1)</td>
<td>-0.0001</td>
<td>4.30E-05</td>
</tr>
<tr>
<td>(p_2)</td>
<td>0.0075</td>
<td>0.0028</td>
</tr>
<tr>
<td>(p_3)</td>
<td>-0.2057</td>
<td>0.37</td>
</tr>
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</table>

Table 1: Prior Distribution for Quadratic Parameters

Figure 12 gives an example of a degradation case which is not well fit by the non-Bayesian approach. Few observations (~30% of the total lifetime) are available, and those available have noise levels, which preclude appropriate model fitting. The same data set, fit with the Bayesian approach described and the prior distribution estimates given above is shown in Figure 13. As can be seen, the Bayesian fit reflects the shape seen in the historical degradation paths. The RUL estimate obtained with the Bayesian approach is 135 cycles, versus an undeterminable estimate obtained from the non-Bayesian approach. The actual RUL after the first 84 observations is 170 cycles, resulting in an RUL error of approximately 20%. While this error is still high, it is within a reasonable accuracy considering the amount of data available and will improve as more data are collected.

The advantage of including prior information via dynamic Bayesian updating is to improve RUL estimates when very few observations are available, the data are very noisy, or both. A comparison of the performance through time of the GPM algorithm and the GPM with Bayesian updating, hereafter referred to as GPM/Bayes, is given in Figure 14. In this analysis, the two methodologies were applied to each of the training cases using only a fraction of the full lifetime. The models were applied to subsets of each lifetime in 5% increments, i.e. the models were run using 5% of the full lifetime, 10%, 15%, etc. The RUL error at each percentage was calculated across the 218 full training cases to determine how the error decreases as more data become available. As was seen in the example case above, the non-Bayesian method may result in an undeterminable RUL. In fact, for the data used here, nearly half the runs resulted in an indeterminate RUL estimate using the GPM methodology without Bayesian updating for runs using less than half the total lifetime. For these cases, the RUL is estimated using a Type I, or traditional reliability-based, method in order to give an estimate of RUL prediction error. The mean residual life is found at each time using a Weibull fit of the failure times and the current lifetime (Figure 15).

\[
MRL(t) = \frac{1}{R(t)} \int_0^t R(s)ds
\]

(15)

where \(R(t)\) is the reliability function at time \(t\). In practice, the prognostic method would likely fall back to a more rudimentary method such as this if the Type III model did not produce a reasonable answer.
The GPM/Type I model which does not include prior information gives an average error of approximately 55% when only 5% of the full lifetime is available and relies on the Type I method for approximately half of the cases. Conversely, the GPM/Bayes method gives approximately 25% error and is able to predict an RUL for every case. As Figure 14 shows, the average error of both methods decreases as more data becomes available and eventually converges to approximately equal error values when the available data overpowers the prior information in the GPM/Bayes model.

4. CONCLUSION

This paper presented a method for performing prognostics on individual components or systems. The General Path Model (GPM) method is used to extrapolate a prognostic parameter curve to a predefined critical failure threshold to obtain an estimate of the Remaining Useful Life (RUL). In cases where only a few data points are available or the data are contaminated by significant noise, a Bayesian method was introduced. The Bayesian method includes prior information about the prognostic parameter distribution to "force" the functional fit to follow the trend seen in historic systems. The method was applied to the 2008 PHM conference challenge problem to illustrate its efficacy.

The given application utilized the results of a condition monitoring and fault detection system to characterize the degradation in a specific system. A prognostic parameter was generated from a subset of the monitoring system residuals; monitoring system residuals are well-suited components of a prognostic parameter because they naturally characterize the deviation of a system from nominal condition. A parametric, linear-in-parameters regression fit of the time series prognostic parameter was extrapolated to the critical failure threshold to give an estimate of the system RUL. A Bayesian updating method was applied to allow for the inclusions of prior information, which improves model performance particularly when faced with small amounts of data or extremely noisy data. The results show that the GPM/Bayes method greatly improved RUL predictive performance over a conventional regression solution.

The need for a diversity of algorithms suggests that development of a large variety of prognostic methods can only strengthen the field. While the algorithm described here may not be the best performing method for this dataset, it has several key advantages that the winning PHM Challenge algorithms lack, which may make it better suited for other applications. The proposed GPM/Bayes algorithm is qualitatively compared to the three best performing algorithms at the 2008 PHM Challenge in the following discussion.

The similarity-based approach described in (Wang et al., 2008) shares several assumptions with the GPM method, namely (1) run-to-failure data from multiple units are available and (2) the history of each training unit ends at a soft failure, but may begin at some random level of initial degradation. However, this similarity-based method suffers the same deficiency that all similarity-based models suffer; it is only applicable within the range of data used for training. The proposed GPM/Bayes method will trend toward the training data when the Bayesian information is dominant, early in equipment degradation or when data are very noisy, but as more data become available, the method will accommodate degradation paths outside those seen in training. Additionally, the proposed similarity method requires storage of a large bank of historical data. This may not be a problem for large computer systems, but it can become cumbersome for onboard prognostic algorithms and systems with many fault
modes requiring many historical paths. Conversely, the GPM/Bayes method requires storage for only the regression model to be fit and the Bayesian prior information.

The second and third place submissions both focused on recurrent neural networks for prognostic estimation (Heimes, 2008; Peel, 2008). Neural networks require a certain level of expertise and finesse to develop. While they are very powerful modeling tools, neural networks lack the accessibility of the GPM/Bayes method or other regression models. A well-developed neural network may outperform many other prognostic algorithms, but development is not a trivial task. Neural network approaches should not be discounted by any means, but the advantage of the GPM/Bayes method is its relative simplicity.

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REFERENCES


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