Model-based Diagnostics for Propellant Loading Systems

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Abstract—The loading of spacecraft propellants is a complex, risky operation. Therefore, diagnostic solutions are necessary to quickly identify when a fault occurs, so that recovery actions can be taken or an abort procedure can be initiated. Model-based diagnosis solutions, established using an in-depth analysis and understanding of the underlying physical processes, offer the advanced capability to quickly detect and isolate faults, identify their severity, and predict their effects on system performance. We develop a physics-based model of a cryogenic propellant loading system, which describes the complex dynamics of liquid hydrogen filling from a storage tank to an external vehicle tank, as well as the influence of different faults on this process. The model takes into account the main physical processes such as highly non-equilibrium condensation and evaporation of the hydrogen vapor, pressurization, and also the dynamics of liquid hydrogen and vapor flows inside the system in the presence of helium gas. Since the model incorporates multiple faults in the system, it provides a suitable framework for model-based diagnostics and prognostics algorithms. Using this model, we analyze the effects of faults on the system, derive symbolic fault signatures for the purposes of fault isolation, and perform fault identification using a particle filter approach. We demonstrate the detection, isolation, and identification of a number of faults using simulation-based experiments.

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1. INTRODUCTION

The loading of cryogenic spacecraft propellants is an inherently risky and unsafe operation, especially in the case of hydrogen [1–4]. Therefore, diagnostic solutions are necessary to quickly identify when a fault occurs, so that recovery actions can be taken or an abort procedure can be initiated before system safety is compromised. Model-based diagnosis approaches enable quick and robust detection, isolation, and identification of faults, because they rely on a detailed model of system behavior under nominal and faulty conditions.

Applying a model-based approach requires an in-depth analysis and understanding of the underlying physical processes in order to produce an accurate and reliable system model. However, cryogenic propellant loading involves complex physical processes that are difficult to capture. In this paper, we develop a medium-fidelity, lumped-parameter dynamical model of propellant loading that takes into consideration a variety of complex multi-phase phenomena that govern the storage and transfer of cryogenic propellants, yet is simple enough to allow for physics analysis and numerical simulations of real loading systems [5]. We concentrate on a system of liquid hydrogen (LH2) filling that is functionally representative of the Space Shuttle refueling system. In this system, LH2 is stored on the ground in a spherical, insulated, double-walled storage tank (ST), and is transferred to the external vehicle tank (ET) through a network of pipes and valves. The model takes into account the main physical processes such as highly non-equilibrium condensation and evaporation of the hydrogen vapor, pressurization, and also the dynamics of liquid hydrogen and vapor flows inside the system in the presence of helium gas. Since the model incorporates faults in the system, it provides a suitable framework for model-based diagnostics and prognostics algorithms.

We apply a model-based diagnostic approach to the system using a combined qualitative-quantitative diagnosis methodology based on the approach of [6]. Deviations in measured values from model-predicted values are compared to qualitative predictions made using the system model for quick fault isolation. Fault identification is performed using particle filters for joint state-parameter estimation. Using simulation-based experiments, we demonstrate the detection, isolation, and identification of a number of faults.

The paper is organized as follows. Section 2 describes the propellant loading system and its physics model. Section 3 overviews the diagnosis approach. Section 4 describes the fault detection methodology. Section 5 discusses fault isolation, and Section 6 develops the fault identification approach. Section 7 verifies the approach with a number of simulation-based experiments. Section 8 concludes the paper.
2. SYSTEM MODELING

Model-based diagnosis algorithms utilize a model of the system for the fault detection, isolation, and identification tasks. We advocate a physics-based modeling approach, because an understanding of the physical processes in both the nominal and faulty cases is necessary for successful fault identification. Information on fault severity is necessary in order to make appropriate recovery actions in response to a fault, especially in the propellant loading domain. In this section, we develop the physics model of the LH2 propellant loading system. Since the focus of the paper is on diagnostics, we review the main features of the model, and refer to [4, 5] for additional details. We first summarize the filling protocol, followed by mathematical descriptions of the tank histories.

Filling Protocol

The purpose of the LH2 propellant loading system is to move LH2 from the ST to the ET. Fig. 1 shows a simplified, but functionally equivalent schematic of the system. Initially, the ullages of the tanks are at atmospheric pressure due to the presence of gaseous hydrogen (GH2). Before filling, the tanks are first pressurized. The ST is pressurized to 54.7 psia, then 80.7 psia through the use of the vaporizer, which boils off LH2 from the ST and returns the GH2 to the ullage of the ST. The ET is filled with gaseous helium (GHe) through the prepressurization valve, until it reaches 38.7 psia. The purpose of pressurization is two-fold. First, it limits potential boiling of the propellant by keeping a high vapor pressure in the ullage of the tanks. Second, the pressure difference between the ST and ET drives propellant from the ST to the ET in the absence of a pump.

Filling progresses in stages with different filling rates, controlled by the position of the transfer line valve (in reality there are a number of valves between the tanks, but in this paper we consider a simplified representation consisting of a single valve). Slow fill begins first with a low flow rate and chilling of the ET. As the liquid drains out of the ST, its ullage pressure drops, so the vaporizer constantly maintains the ullage pressure to keep LH2 flowing to the ET. The flow through the vaporizer valve is modulated based on the error between the measured ST ullage pressure and the ST pressure set point. As the ET is filled, its ullage volume decreases, and, therefore, its ullage pressure increases. The ullage pressure in the ET is maintained using its vent valve, which opens and closes to maintain the pressure between 38.7 and 41.7 psia.

When the ET is 5% full, fast fill begins at a flow rate around 30 kg/s. When the ET is 72% full, the ullage pressure of the ST is reduced to 64.7 psia, reducing the flow rate. When the ET is 85% full, the fill rate is reduced further by partially closing the transfer line valve. When the ET is 98% full, topping begins at a lower flow rate. The ET vent valve is also opened, reducing the ET ullage pressure to 14.7 psia. Finally, at 100% full, topping ends and the tank is then continuously replenished until launch to replace the boil off. During replenish, the fill valve position is modulated to maintain the ET level at 100%.

Tank Modeling

For each tank, we consider three control volumes: the vapor, the liquid, and the vapor film, as shown in Fig. 2. By convention, positive mass/energy flows enter the CV, and negative flows exit the CV.

The vapor CV (v) consists of GH2 (subscript v) and GHe (subscript g), and is treated as a mixture of ideal gases with partial densities \( \rho_v \) and \( \rho_g \), as well as a common temperature \( T_v \), all related to each other by the following equations of state:

\[
\rho_v = \rho_v R_v T_v \quad (1)
\]

\[
\rho_g = \rho_g R_g T_v \quad (2)
\]

The liquid CV (l) is where, far from the surface, the temperature is equal to \( T_l \) and the liquid is treated as incompressible. \( T_l \) is treated as a constant at 20 K.

The vapor film CV (f) separates the liquid and gas phases. It is treated as saturated hydrogen vapor, whose temperature is
equal to that of the liquid/vapor interface, where \([1, 4, 7]\)
\[
T_f = T_C \left( \frac{p_v}{p_C} \right)^n, \tag{3}
\]
with \(p_C = 1.315 \text{ MPa}, T_C = 33.2 \text{ K}, \) and \(n = 5 \) \([7]\).

For the liquid CV, the net mass flow is defined by the external mass flow \(J_{le}\) (i.e., the flow across its boundaries other than through the interface), and the interphase (condensation-evaporation) flow \(J_{lv}\) (defined later):
\[
\dot{m}_l = J_{le} + J_{lv} \tag{4}
\]
Similarly, the GH2 and GHe mass conservation for CV \((v)\) is
\[
\dot{m}_v = J_{ve} - J_{lv} \tag{5}
\]
and
\[
\dot{m}_g = J_{ge}, \tag{6}
\]
where \(J_{ve}\) and \(J_{ge}\) are the external GH2 and GHe flows. For a given tank volume \(V\), the vapor volume is fully defined by the mass of liquid, since it has a constant density:
\[
V = V_v + V_l = V_v + m_l/p_l. \tag{7}
\]

The energy conservation law for the vapor CV yields:
\[
\frac{d(m_v u_v + m_g u_g)}{dt} = \dot{U}_v = \dot{Q}_{vf} - \dot{Q}_{fl} - W -
J_{lv} h_{vf} + J_{ve} (h_{ve} + v_{ve}^2/2) + J_{ge} (h_{ge} + v_{ge}^2/2), \tag{8}
\]
where \(u_{v(g)} = c_{v(g)} T_v\) are specific internal energies of the gases, \(\dot{U}_v\) is the net rate of change in internal energy of the mixed gas, \(\dot{Q}_{ve}\) is the net external heat flow into the CV \((v)\) through the tank walls, \(\dot{Q}_{vf}\) is the heat flow lost through the interface, \(W = -p_t \dot{V}_l/dt\) is related to the quasi-static power due to compression (expansion) of the CV \((v)\), \(h_{vf} J_{lv}\) is the interphase enthalpy flow (the specific enthalpy \(h_{vf}\) is equal to that of the saturated vapor), \(h_{ve}\) and \(h_{ge}\) are the specific enthalpies of the hydrogen and helium gas entering the CV, and \(v_{ve}\) and \(v_{ge}\) are the velocities of the incoming gases \([8]\). Here, the kinetic energies associated with both the GH2 and GHe mass flows entering the CV \((v)\) are taken into consideration, because the corresponding velocities \(v_{v(g)}\) are much greater than the one related to interphase flow. The specific enthalpies \(h_{v(g)} = u_{v(g)} + p_{v(g)}/\rho_{v(g)} = u_{v(g)} + R_{v(g)} T_v = c_{p_v(g)} T_v\). Here, \(c_{p_v(g)} = c_{V_v(g)} + R_{v(g)}\) and is the specific heat at constant pressure. The temperature of the mixed gas is then described by
\[
\dot{T}_v = \frac{1}{m_v c_{V_v}} (\dot{U}_v - \dot{m}_v c_{V_v} T_v - \dot{m}_g c_{V_g} T_v), \tag{9}
\]
where \(c_{V_v}\) and \(c_{V_g}\) are the specific heats at constant volume for the vapor and gas, and \(c_{V_v}\) is the specific heat at constant volume for the mixed gas.

If the film layer is considered negligibly thin so that one can ignore its mass \([1]\), then the energy balance equation for the CV \((f)\) can be written as
\[
\dot{Q}_{vf} - \dot{Q}_{fl} + J_{le} h_{lv} = 0, \tag{10}
\]
where \(\dot{Q}_{vf}\) is the heat flow from the vapor CV, \(\dot{Q}_{fl}\) is the heat flow to the liquid CV, and \(h_{lv}\) is the enthalpy (heat) of vaporization. Strictly speaking, \(h_{lv}\) depends on the saturated vapor temperature, such that it goes to zero when the surface temperature approaches the critical temperature \(T_C\) \([9]\). To take this effect into consideration, we use the following simple interpolation formula for \(T_f \leq T_C\):
\[
h_{lv}(T_f) = h_{lv}^0 \left( \frac{T_C - T_f}{T_C - T_l} \right)^{1/2}, \tag{11}
\]
where for liquid hydrogen, \(T_C = 33 \text{ K}\) and \(h_{lv}^0 \approx u_{lv}^0 = 4.5 \times 10^6 \text{ J/kg}\) at \(p = 1 \text{ atm}\) and \(T_l = 20 \text{ K}\) \([9]\).

The heat flows \(\dot{Q}_{fl}\) and \(\dot{Q}_{vf}\) are dominated either by conduction or convection, depending on the relative temperatures of the liquid, vapor, and film \([5]\). If \(T_f > T_l\) then conduction heat transfer defines \(\dot{Q}_{fl}\), else convection does. If \(T_v > T_f\), then conduction heat transfer defines \(\dot{Q}_{vf}\), else convection does. In the case of energy, we use the following approximations:
\[
\dot{Q}_{fl}^{\text{cond}} = A_f \alpha_{fl}^{\text{cond}} (T_f - T_l), \tag{12}
\]
\[
\dot{Q}_{vf}^{\text{cond}} = A_f \alpha_{vf}^{\text{cond}} (T_v - T_f), \tag{13}
\]
where \(A_f\) is the surface area of the interface, and the \(\alpha\) terms are heat transfer coefficients (see \([5]\)). Typically, conduction expressions contain complex integral relations, but here, we use this algebraic approximation that has proven adequate for our system \([5]\). For convection, we use
\[
\dot{Q}_{fl}^{\text{conv}} = A_f \alpha_{fl}^{\text{conv}} (T_f - T_l), \tag{14}
\]
\[
\dot{Q}_{vf}^{\text{conv}} = A_f \alpha_{vf}^{\text{conv}} (T_v - T_f). \tag{15}
\]

Both the liquid and vapor CVs absorb external \((e)\) heat from the tank walls. This heat is transferred by means of convection so that
\[
\dot{Q}_{w(t)e} = A_{w(t)} \alpha_{w(t)e} \left( T_w - T_{w(t)} \right), \tag{16}
\]
where \(A_{w(t)}\) are the internal tank surfaces in contact with vapor (liquid), and \(\alpha_{w(t)e}\) are the convection heat transfer coefficients \([5]\). The wall temperature \(T_w\) is governed by the heat flow passing through the walls from the environment \([1–3]\).

The temperature \(T_w\) of the tank wall, considered uniform, is defined by the heat exchange rate with the tank surroundings with the effective ambient temperature \(T_a\):  
\[
\dot{Q}_w = A_w \alpha_w (T_a - T_w), \tag{17}
\]
where \(A_w\) is the external tank surface area. The wall temperature is governed by the tank energy conservation:
\[
m_w c_w \dot{T}_w = \dot{Q}_w - \dot{Q}_{le} - \dot{Q}_{we}. \tag{18}
\]
Here, the heat transfer coefficients describe natural convection inside and outside the tank walls \([1, 10]\).
The Storage Tank

The above equations apply equally well to both the ST and the ET. We denote variables of the ST with a 1 subscript, and variables of the ET with a 2 subscript. For the ST, there is no GHe in the ullage volume, so \( p_{g1} = \rho_{g1} = 0 \). The external mass flow for the liquid consists of vaporizer flow \( J_{vap} \), transfer line flow \( J_{tr} \), and leak flow \( J_{leak,1} \):

\[
J_{le1} = -J_{vap} - J_{tr} - J_{leak,1}. \tag{19}
\]

\( J_{tr} \) is described by

\[
J_{tr} = \lambda_{tr}\sigma_{tr}\alpha_{tr}\sqrt{p_1 - p_2}, \tag{20}
\]

where \( \lambda_{tr} \) is the valve position (an input), \( \alpha_{tr} \) is the flow coefficient, \( p_1 = p_{v1} + \rho_{gh}h_{l1} \) is the total ST pressure where \( h_{l1} \) is the height of the liquid, \( p_2 = p_{v2} + \rho_{gh}h_{l2} \) is the total ET pressure, \( g \) is the acceleration due to gravity and \( \sigma_{tr} \) is a (dimensionless) multiplicative factor describing a transfer line blockage fault.

In the vaporizer, a certain amount of LH2 is evaporated and returned to the ST, thus controlling its ullage pressure. We assume that all liquid flow through the vaporizer is converted instantaneously to GH2. The flow is given by

\[
J_{vap} = \lambda_{vap}\sigma_{vap}\alpha_{vap}\sqrt{p_{v1} - p_{v2}} \tag{21}
\]

where \( \lambda_{vap} \) is the vaporizer valve position, \( \alpha_{vap} \) is the flow coefficient, \( p_{vap} \) is the pressure in the vaporizer (close to atmospheric pressure), and \( \sigma_{vap} \) is a multiplicative factor describing the vaporizer valve blockage fault. The vaporizer valve position is controlled by

\[
\lambda_{vap} = \min \left( 1, \max \left( 0, 10\frac{p_{v1}^* - p_{v1}}{p_{v1}^*} \right) \right). \tag{22}
\]

where \( p_{v1}^* \) is the desired ST ullage pressure.

The liquid leak flow is given by

\[
J_{leak,1} = A_{leak,1}\alpha_{leak,1}\sqrt{p_1 - p_{atrm}}. \tag{23}
\]

where \( A_{leak,1} \) is the the area of the leak hole, \( \alpha_{leak,1} \) is the leak hole flow coefficient, and \( p_{atrm} \) is atmospheric pressure.

In the vapor CV, \( J_{vc1} = -J_{v,leak,1} \), where

\[
J_{v,leak,1} = A_{v,leak,1}\alpha_{v,leak,1}\sqrt{p_1 - p_{atrm}}. \tag{24}
\]

Heat leaks constitute the external heat flow term, i.e., \( \dot{Q}_{ve1} = \dot{Q}_{leak,1} \).

The External Tank

For the ET, the external liquid flow is given by

\[
J_{te2} = J_{tr} - J_{leak,2} - J_{boil}, \tag{25}
\]

where \( J_{leak,2} \) is described similar to that for the ST, and \( J_{boil} = \dot{Q}_{te2}/h_{l1}(T_{w2} - T_{l2}) \) is responsible for intense LH2 evaporation as the ET walls are being initially chilled down during the beginning of the slow fill stage.

In the vapor CV, the external flows are given by

\[
\begin{align*}
J_{ve2} &= -J_{v,vent,2} - J_{v,leak,2} \tag{26} \\
J_{ge2} &= J_{g,pp} - J_{v,vent,2} - J_{g,leak,2} \tag{27}
\end{align*}
\]

where \( J_{g,pp} \) is the prepressurization flow of GHe, and

\[
J_{v(g),vent,2} = \lambda_{vent,2}\sigma_{vent,2}\frac{A_{vent,2}p_{atrm}\sqrt{\gamma(p_{v2} + p_{g2} - p_{atrm})}}{\gamma K_{vent,2}(p_{v2} + p_{g2})}. \tag{28}
\]

Here, the dimensionless flow coefficient \( K \) (the loss factor) can be found in Schmidt et al. [10] (see Tables 7-2 and 7-3 therein); a dimensionless relative valve position assumes values between \( \lambda_k = 1 \) (fully open) and \( \lambda_k = 0 \) (fully closed) [10]. During filling, the valve opens when the pressure exceeds 41.7 psia and closes when it falls below 38.7 psia. The vapor/gas and heat leaks may be defined as with the ST.

Nominal Dynamics

Fig. 3 summarizes the major results of the simulation of a nominal loading regime, which are based on the parameters, initial conditions, and filling protocol that are typical for LH2 loading systems. These values are provided in [5].

It can be seen that the LH2 level in the ST drops monotonically (Fig. 3a) as the level in the ET rises (Fig. 3d). The pressure \( p_{v1} \) in the ST (Fig. 3b) is determined by the loading dynamics and controlled by the vaporizer. Once achieved during slow fill, the pressure in the ST is maintained at approximately 80.7 psia up to the start of the reduced-pressure fast fill (see Figs. 3b and 3j), at which point it is maintained at 64.7 psia. Meanwhile, the ET ullage pressure (Fig. 3e) is oscillating due to the cycling of the vent valve that maintains the pressure between the lower and upper thresholds of 38.7 and 41.7 psia. The fluctuations in the ET ullage temperature (Fig. 3f) as well as in the mass flow rates (Figs. 3j and 3h) are driven by the ET pressure oscillations.

The LH2 partial pressure in the ET rises due to the continued hydrogen supply, while the GHe partial pressure drops because the helium is being permanently removed through the vent valve (Fig. 3i). In this case, due to the condensation blocking effect [4], the flow of the condensed vapor in the ET (Fig. 3h) is much smaller than that in the ST (Fig. 3g), because the vapor pressure is being maintained approximately equal to the equilibrium pressure of the condensed vapor at the temperature of LH2. The ullage temperature \( T_{v2} \) in the ET (Fig. 3f) increases initially due to the introduction of the GHe during the pressurization stage, then drops from the initial high value due to venting and near-wall boiling that generates relatively cold GH2 during filling. The liquid surface temperature \( T_{f1} \) in the ST increases (Fig. 3c) due to the vapor condensation at the interface. Simultaneously, the ST ullage temperature \( T_{e1} \) increases, mainly because the relatively hot
GH2 is supplied by the vaporizer as loading occurs. As a result, the ullage temperature approaches the temperature of LH2 saturated vapor at a pressure close to the final ST ullage pressure of approximately 5 atm (Fig. 3b).

![Figure 3. Nominal regime of LH2 propellant loading.](image)

3. Diagnosis Approach

We apply a model-based diagnosis approach using the physics model of the LH2 system. In this paper, we consider the problem of single fault diagnosis. The system may be described in the following general form:

\[
\begin{align*}
x(t) &= f(t, x(t), \theta(t), u(t), v(t)) \\
y(t) &= h(t, x(t), \theta(t), u(t), n(t)),
\end{align*}
\]

where \(x(t) \in \mathbb{R}^{n_x}\) is the state vector, \(\theta(t) \in \mathbb{R}^{n_\theta}\) is the parameter vector, \(u(t) \in \mathbb{R}^{n_u}\) is the input vector, \(v(t) \in \mathbb{R}^{n_v}\) is the process noise vector, \(f\) is the state equation, \(y(t) \in \mathbb{R}^{n_y}\) is the output vector, \(n(t) \in \mathbb{R}^{n_n}\) is the measurement noise vector, and \(h\) is the output equation.

Measurements are time-varying signals of \(y(t)\) obtained from the system sensors. In the LH2 system, we consider the following measurements for diagnosis: \(V_{l1}, p_1, T_{v1}, V_{l2}, p_2, T_{v2}, J_{tr}, \lambda_{cap},\) and \(\lambda_{vent,2}\).

We consider single, abrupt faults, modeled as unexpected step changes in system parameter values. We name faults by the
associated parameter and the direction of change, i.e., $\theta^+$ denotes a fault defined as an increase in the value of parameter $\theta$, and $\theta^-$ denotes a fault defined as a decrease in the parameter value. For the LH2 system, we consider liquid and gas leaks, heat leaks, and valve clogging. The liquid and vapor leaks are defined by the equivalent leak areas, nominally zero, so faults are defined as increases in these areas, denoted by $A^+_{leak,1}$, $A^+_{leak,2}$, and $A^+_{leak,2}$. The heat leaks are defined by the heat leak rate, nominally zero, so faults are increases in these values, denoted by $Q^+_{leak,1}$ and $Q^+_{leak,2}$. The valve clogging faults are described by the $\sigma$ parameters, which are nominally 1, so faults are decreases in these values (0 at a minimum), denoted by $\sigma^-_{tr}$, $\sigma^-_{vap}$, and $\sigma^-_{vent,2}$.

The diagnosis architecture is shown in Fig. 4. The system receives inputs $u(t)$ and produces outputs $y(t)$. The physics simulation runs simultaneously, producing predicted outputs $\hat{y}(t)$, given the inputs $u(t)$. Using statistical methods, the fault detection module decides when a measurement has deviated from its nominal value, triggering fault isolation. Measurement deviations are then used to quickly isolate faults $F$. Fault identification computes, for each fault $f \in F$, the value of the fault parameter that best fits the outputs of the system, and the candidate with the lowest output error is selected as the best candidate.

### 4. Fault Detection

In model-based fault detection, a model of the system provides reference outputs representing nominal system behavior. For each sensor output $y(t)$, we define the residual as $r(t) = y(t) - \hat{y}(t)$, where $\hat{y}(t)$ is the model-predicted output signal. Statistically significant deviations of the actual system outputs from the model-predicted outputs imply the presence of a fault. If the model is accurate, then fault detection thresholds can be small and faults detected quickly. The thresholds are dynamic in that they are defined with respect to nominal behaviors as a function of time. This is favorable to the current practice with such systems, where thresholds are static, preventing the detection of subtle deviations from nominal behavior that indicate faults.

We use the Z-test for robust fault detection using a set of sliding windows, as described in [11, 12]. The current mean of a residual signal, $\mu_r(t)$, is estimated over a small window $W_2$:

$$\mu_r(t) = \frac{1}{W_2} \sum_{i=t-W_2+1}^{t} r(i).$$

The variance of the nominal residual signal, $\sigma_r^2(t)$, is computed using a large window $W_1$ preceding $W_2$, by a buffer $W_{delay}$, ensuring that $W_1$ does not contain any samples after fault occurrence. The variance is computed using

$$\sigma_r^2(t) = \frac{1}{W_1} \sum_{i=t-W_2-W_{delay}-W_1+1}^{t-W_2-W_{delay}} (r(i) - \mu_r(t))^2,$$

where

$$\mu_r(t) = \frac{1}{W_1} \sum_{i=t-W_2-W_{delay}-W_1+1}^{t-W_2-W_{delay}} r(i).$$

A given confidence level determines the bounds $z^- < 0$ and $z^+ > 0$ for a two-sided Z-test. The fault detection thresholds, $\varepsilon_r^-(t)$ and $\varepsilon_r^+(t)$, are dynamically computed using

$$\varepsilon_r^-(t) = z^- \frac{\sigma_r(t)}{\sqrt{W_2}} - E,$$

$$\varepsilon_r^+(t) = z^+ \frac{\sigma_r(t)}{\sqrt{W_2}} + E,$$

where $E$ is a modeling error term. A fault is detected if $\mu_r(t)$ lies outside of the thresholds at time $t$. If $\mu_r(t) < \varepsilon_r^-(t)$, a $-$ symbol for the measurement is used by the fault isolation module, and if $\mu_r(t) > \varepsilon_r^+(t)$, a $+$ symbol is used. Generally, the parameters $W_1$, $W_2$, $W_{delay}$, the $z$ bounds, and $E$ must be tuned to optimize performance to minimize both false alarms and missed detections.

### 5. Fault Isolation

We utilize a qualitative diagnosis methodology that isolates faults based on the transients they cause in system behavior, manifesting as deviations in observed measurement values from nominal measurement values [6]. The transients are abstracted using qualitative + (increase), − (decrease), and 0 (no change) values to form fault signatures. Fault signatures represent these measurement deviations from nominal behavior as the immediate (discontinuous) change in magnitude, and the first nonzero derivative change.

![Figure 4. Diagnosis architecture.](image-url)
The fault signatures can be derived automatically from a graph-based representation of the system model, known as a temporal causal graph (TCG) [6]. A TCG captures dynamic system variables as nodes in a graph, and the qualitative relations between them as edges. These edges are labeled with \( dt \), representing integration, +1, denoting a proportionality, −1, denoting an inverse proportionality, and system parameters. Faults that are modeled as parameter changes appear on edges, allowing the qualitative effects of parameter changes to be propagated over the system variables. Propagation to measured variables reveals the qualitative effects of faults on measurements.

A partial TCG for the LH2 system, showing some key variables of the ST, is shown in Fig. 5. Variables within dashed boxes are measured. Here, we show only the \( \sigma_{\text{vap}} \) fault. A forward propagation algorithm may be used to derive the fault signatures. Details may be found in [6], so here, we use the vaporizer valve clogging fault as an example to illustrate the general procedure. According to the TCG, a decrease in \( \sigma_{\text{vap}} \) will lead to a decrease in \( J_{\text{v}} \), and subsequently, a decrease in \( J_{\text{t}} \), and, due to the −1 label, an increase in \( J_{\text{r}} \). The decrease in \( J_{\text{v}} \) will lead to a first-order decrease (due to the \( dt \) label) in \( m_{\text{v}} \) and, subsequently, \( p_{\text{v}} \), which is a measured variable. This change then propagates as a decrease in \( J_{\text{t}} \), and then towards ET variables. The increase in \( J_{\text{r}} \) will lead to a first-order increase in \( m_{\text{l}} \) and \( V_{\text{l}} \), which is measured, leading to an increase in \( J_{\text{t}} \). This conflicts with the previously predicted decrease from the other path with the same derivative order, resulting in an ambiguity denoted with a *.

We can use the simulation and knowledge of the system dynamics to resolve these ambiguities. In this case, we know that the hydrostatic pressure due to the height of the liquid is negligible compared to \( p_{\text{v}} \) due to the very low density of LH2. Therefore, we know that the decrease effect will dominate. The propagation continues to all system variables.

Fault signatures for the LH2 system are shown in Table 1. Due to the dynamics of the system, all faults appear as smooth changes in the measured values, so only the first change presents useful diagnostic information, hence, we show only the first symbol derived from forward propagation. Recall that the derived symbols represent deviations from nominal behavior, so, for example, the + symbol for \( V_{\text{l}} \) caused by \( A_{v,\text{leak},1}^+ \) does not necessarily mean that \( V_{\text{l}} \) increases, rather, it increases with respect to its nominal value, i.e., it decreases at a slower rate. The remaining * symbols are for those in which the qualitative effect may be different depending on the state of the system. Note also that this is a hybrid system, i.e., it has continuous dynamics mixed with discrete dynamics due to the valves, and we show only the signatures for the mode where the vent valve is closed. When the vent valve opens, some of the signatures will flip. A systematic framework for dealing with this issue is discussed in detail in [13].

Using the signatures, we can perform diagnosability analysis to determine the effectiveness of the isolation step. We can see that in most cases, the set of signatures produced distinguishes most faults. One exception is the pair \( A_{v,\text{leak},1}^+ \) and \( \sigma_{\text{vap}} \), which have all the same signatures, therefore, fault identification will have to distinguish between them, since quantitatively, they should have different effects on the measured variables. If we measure \( J_{\text{vap}} \), then the faults can be distinguished, because \( \sigma_{\text{vap}} \) would produce a decrease in \( J_{\text{vap}} \), whereas \( A_{v,\text{leak},1}^+ \) would produce an increase. The faults \( \dot{Q}_{\text{leak},2}^+ \) and \( \sigma_{\text{vent},2} \) are similarly indistinguishable.

We can also use the signatures to perform measurement selection. The measurement of \( \lambda_{\text{vap}} \) always has the same signature as \( p_{\text{v}} \), therefore one of these can be eliminated without loss of diagnosability. However, both quantities are relatively easy to measure, so there is value in keeping them both. \( \lambda_{\text{vap}} \) is typically more sensitive to faults than \( p_{\text{v}} \), since \( p_{\text{v}} \) is controlled by the vaporizer, which may mask faults. Simi-
larly, both $p_2$ and $\lambda_{\text{vent},2}$ are not needed. Changes in $p_2$ are reflected in changes in the switching frequency of the vent valve, and therefore provide the same information, although changes in the vent valve position can be detected much more easily, again because the control of $p_2$ leads to fault masking in that measurement. It is also beneficial to keep as many measurements as possible, because this gives more information for fault identification. Further, the qualitative fault signatures say nothing about fault magnitude. Some faults may not produce large enough changes in some variables for the changes to be detected, therefore, having more measurements helps to alleviate this problem.

6. Fault Identification

Fault identification is initiated immediately after the initial set of fault candidates is produced after fault detection. Each candidate has its own identification module that updates its estimate at every time step. Identification is performed using the particle filtering algorithm for joint state-parameter estimation [14, 15], which has seen previous application in model-based diagnosis and prognosis algorithms [16–19]. The magnitude of the fault parameter $\theta$ is estimated along with the system state.

The identification module must compute $p(x_k, \theta_k | y_{0:k})$. A general solution to this problem is the particle filter, which may be directly applied to nonlinear systems with non-Gaussian noise terms. Particle filters offer approximate (sub-optimal) solutions to the state estimation problem for systems where optimal solutions are unavailable or intractable [14, 15]. In particle filters, the state distribution is approximated by a set of discrete weighted samples, called particles. As the number of particles is increased, performance increases and the optimal solution is approached. Due to the highly nonlinear dynamics of the LH2 loading system, particle filters are favored over other estimation approaches such as the extended Kalman filter.

The particle approximation to the state distribution is given by

$$\{(x_i^k, \theta_i^k), w_i^k\}_{i=1}^N,$$

where $N$ denotes the number of particles, and for particle $i$, $x_i^k$ denotes the state vector estimate, $\theta_i^k$ denotes the parameter vector estimate, and $w_i^k$ denotes the weight. The posterior density is approximated by

$$p(x_k, \theta_k | y_{0:k}) \approx \sum_{i=1}^N w_i^k \delta(x_k^i, \theta_k^i) (dx_k d\theta_k),$$

where $\delta(x_k^i, \theta_k^i) (dx_k d\theta_k)$ denotes the Dirac delta function located at $(x_k^i, \theta_k^i)$.

We use the sampling importance resampling (SIR) particle filter with systematic resampling [14, 20]. The pseudocode for a step of the filter is shown as Algorithm 1. Each particle is propagated forward to time $k$ by sampling new parameter and state values. The particle weight is assigned using $y_k$. The weights are then normalized, and then the particles are resampled (see [14]).

Note that the parameters $\theta_k$ evolve by some unknown process that is independent of the state $x_k$. The particle filter algorithm requires some type of evolution to the parameters. We use a random walk, i.e., for parameter $\theta$, $\theta_k = \theta_{k-1} + \xi_{k-1}$, where $\xi_{k-1}$ is Gaussian noise. The particles generated with parameter values closest to the true values should match the outputs better, and, therefore, be assigned higher weight, thus allowing the particle filter to converge to the true values. The selected variance of the random walk noise affects both the rate of this convergence and the estimation performance after convergence. Using the simulation, we can determine appropriate values for the random walk variances.

Under the single fault assumption, we can run a set of parallel particle filters, one for each consistent fault candidate. This reduces the dimensionality of the estimation task over combined estimation of all fault parameters, and allows the particle filters to be much more efficient. The particle filters are initiated at the point of fault detection, using the model-estimated state at that time point for initialization of the states. The fault parameter estimate starts at its value during nominal operation (e.g., $0$ for leak areas). When the fault isolation module reduces the set of candidates $F$, the identification task continues only for those faults remaining in $F$.

Because diagnosability may be limited, fault identification must also be used to help refine fault candidates. The particle filter for the true fault candidate should estimate the correct fault parameter and track the faulty outputs with low error, whereas the particle filters for the incorrect faults will not track and result in large error. We compute the mean squared output error for each candidate from the point of fault detection to the present time. The candidate with the lowest output error $\epsilon$ is considered to be the true candidate.
7. RESULTS

We illustrate the diagnosis process with the $A_{\text{v, leak, 1}}^+$ fault injected at 2600 s with magnitude $3 \times 10^{-3}$ m$^2$. Note that from diagnosability analysis, the observed signatures will be consistent with both the ST vapor leak fault $A_{\text{v, leak, 1}}^+$ and the vaporizer valve clogging fault $\sigma_{\text{vap}}$, so the fault identification stage will have to resolve the ambiguity. Relevant measurements are shown in Fig. 6. The diagnoser, running at a sample time of once per second, detects the fault at 2605 s due to an observed decrease in $p_1$. The initial list of consistent faults is \{ $A_{\text{v, leak, 1}}^+, A_{\text{v, leak, 2}}^+, A_{\text{l, leak, 1}}^+, A_{\text{l, leak, 2}}^+$, $\sigma_{\text{vap}}$ \}. At 2608 s, a decrease in $J_{tr}$ is detected, followed by an increase in $\lambda_{\text{vap}}$ at 2612 s and a decrease in $T_{v1}$ at 2690 s. The candidate set remains the same. At 2703 s, a decrease in $V_{l2}$ is detected, which rules out the ET vapor/gas leak fault $A_{\text{v, leak, 2}}^+$. At this point, the output error of the $A_{\text{v, leak, 1}}^+$ fault is an order of magnitude less than for all other remaining candidates. At 2806 s, a detected increase in $V_{l1}$ rules out $A_{\text{l, leak, 1}}^+$ and $A_{\text{l, leak, 2}}^+$, resulting in the candidate list \{ $A_{\text{v, leak, 1}}^+, \sigma_{\text{vap}}$ \}. By this time, the particle filter for $A_{\text{v, leak, 1}}^+$ has also converged on the correct value of the fault magnitude, as shown in Fig. 7. The fault was detected within 5 seconds of its occurrence, and identification confirmed the true fault with correct magnitude within 200 s. It is important to quickly discriminate between a vapor leak in the ST and a clogging of the vaporizer, because the former requires an abort, whereas the system can continue fueling in a safe manner with the latter. As discussed in Section 5, measuring $J_{\text{vap}}$ would allow a faster discrimination of these two faults.

Diagnosis results over the complete set of faults are shown in Table 2. Here, $\Delta t_d$ denotes the time to detect the fault, $\Delta t_i$ denotes the time to isolate, taken as the last time at which the candidate set is reduced, $F_{id}$ denotes the output of the fault identification module at the end of the scenario, and $f^*$ denotes the final output of the diagnoser, i.e., the identified fault with the lowest error. For the fault identification stage, 50 particles were used per particle filter, which seemed to offer a reasonable trade-off between computation time and identification accuracy.

Overall, the results are fairly good. In all cases, the correct fault was identified with sufficient accuracy. In some cases, the effects of the faults are very subtle at first, and it takes some time before the changes they produce can be distinguished from the sensor noise. For the vent valve clogging fault, the effects are visible only when the valve is open, so if the fault first appears when the valve is closed, it will take time for the fault to be detected. The $\sigma_{\text{tr}}$ fault was detected immediately because the change in $J_{tr}$ was significant. Without this measurement, it would take some time for the fault to be visible in the liquid volumes of the tanks.

For some faults, the operation of the vaporizer masks the change in ST pressure, therefore, it is important to monitor the vaporizer valve position in order to obtain a more sensitive

![Figure 6](image1.png)

Figure 6. Predicted and observed outputs for the $A_{\text{v, leak, 1}}^+$ fault injected at 2600 s with magnitude $3 \times 10^{-3}$ m$^2$.

![Figure 7](image2.png)

Figure 7. Estimated fault magnitudes and output error for the $A_{\text{v, leak, 1}}^+$ fault injected at 2600 s with magnitude $3 \times 10^{-3}$ m$^2$. The $A_{\text{v, leak, 1}}^+$ estimator converges to the true value by 2800 s. The $\sigma_{\text{vap}}$ estimator converges by 2800 s also, but it cannot track the outputs, as shown by the corresponding error $e_{\sigma_{\text{vap}}}$, which is greater than $e_{A_{\text{v, leak, 1}}}$ at 2800 s.
detection. Changes in the vent valve switching frequency are also easy to observe, but one must wait for the upper pressure threshold to be reached before it can be determined whether it closes early or late. We also observed that faults in the ET detection. Changes in the vent valve switching frequency are historical data of faulty situations. Such data can be difficult to acquire. We considered only single faults here, and in the future, we will extend the framework to multiple faults. Sensor faults were not considered here, but can be easily incorporated into a model-based framework by including models of the sensors [12]. Using the physics model for prognostics and loading optimization is also of interest.

We would like to validate the approach by using available historical data of faulty situations. Such data can be difficult to find, and, further, there is only a limited capability to inject faults into the system for the purpose of diagnosis algorithm validation, due to the high cost of a fueling operation and the explicit danger that fault injection presents in such a system. Therefore, an accurate simulation with which to validate diagnosis algorithms, such as that used here, is very valuable.

8. CONCLUSIONS

In this paper, by applying our previously developed physics-based model of a propellant loading system [5], we analyzed the effects of faults, and applied a model-based diagnosis approach to fault detection, isolation, and identification. The detection stage compares observed and model-predicted outputs to detect faults. The isolation stage compares model-predicted fault transients to observed measurement deviations to quickly isolate faults and reduce the complexity of the identification stage. The fault identification stage uses particle filters to estimate the values of fault parameters and resolve isolation ambiguities.

The simulation results showed that with such a model-based diagnosis approach, faults can be quickly detected, isolated, and identified, and this knowledge can be used to determine if the system can continue loading safely or if an abort is required. We considered only single faults here, and in the future we will extend the framework to multiple faults. Sensor faults were not considered here, but can be easily incorporated into a model-based framework by including models of the sensors [12]. Using the physics model for diagnostics and loading optimization is also of interest.

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<table>
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<tr>
<th>Fault</th>
<th>Magnitude</th>
<th>$\Delta t_d$</th>
<th>$\Delta t_i$</th>
<th>$F_{id}$</th>
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<td>$\sigma_{\text{vent}} = 0.50$, $e = 9.55$</td>
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**REFERENCES**


