Simulations of cavitating turbopump inducers at their design flow rate are presented. Results over a broad range of $N_u$ numbers extending from single-phase flow conditions through the critical head break down point are discussed. The flow characteristics and performance of a subscale geometry designed for water testing are compared with the full-scale configuration that employs LOX. In particular, thermal depression effects arising from cavitation in cryogenic fluids are identified and their impact on the suction performance of the inducer quantified. The simulations have been performed using the CRUNCH CFD code that has a generalized multi-element unstructured framework suitable for turbomachinery applications. An advanced multi-phase formulation for cryogenic fluids that models temperature depression and real fluid property variations is employed. The formulation has been extensively validated for both liquid nitrogen and liquid hydrogen by simulating the experiments of Hord on hydrofoils; excellent estimates of the leading edge temperature and pressure depression were obtained while the comparisons in the cavity closure region were reasonable.

I. Introduction

Our focus is on the development of a computational framework to simulate cavitating liquid rocket turbomachinery that employ cryogenic working fluids. Liquid rocket systems are a subset of a broader class of pumps (e.g. refrigerant systems, boiler feed pumps, etc) where the operating temperature is elevated relative to the critical temperature of the fluid and thermodynamic effects of cavitation play an important role. At these operating temperatures, the ratio of liquid to vapor density is lower and consequently more liquid mass has to vaporize to sustain a cavity. Therefore evaporative cooling effects are more pronounced and result in the lowering of the mean fluid temperature in the cavitating region. Since the fluid thermodynamics properties (i.e. vapor pressure, density) are a strong function of temperature at these conditions, thermal effects suppress cavitation and lower the cavity pressure in a mean sense. Typically this results in improved mean performance of cryogenic pumps; liquid hydrogen systems being an extreme example where the pump may continue to generate head even when the fluid is boiling at the inlet.

The thermal effects of cavitation were studied extensively by numerous researchers through the 1970’s including: Stahl and Stepanoff, Ruggeri and Moore, Hord, and Brennen among others. Stahl and Stepanoff were the first to estimate head depression ($\Delta H_v$) values due to thermodynamic effects using the so-called ‘B-factor’ method based on a quasi-static theory where the temperature depression was estimated in terms of the ratio of the vapor volume to liquid volume. They provided a graph to evaluate NPSH corrections for hydrocarbons based on this methodology. More elaborate correlations, which included dynamic effects were given later by Ruggeri and Moore, Hord, and Holl. They collected extensive experimental data of cavity pressure and temperature depressions for a variety of model shapes and fluids and correlated the results using variants of the B-factor theory. The semi-empirical procedures outlined by Ruggeri and Moore continues to be used as an engineering tool for predicting the thermodynamic depression in pumps. A more rigorous numerical procedure was developed by Cooper where a barotropic equation of state was used to define the two-phase mixture and thermal effects.
effects were evaluated with a resulting non-dimensional vaporization parameter. Most of these techniques, however, require some degree of empiricism. Therefore, from a more fundamental modeling perspective, this discussion highlights the need for a generalized compressible formulation that takes the energy balance into account when simulating cavitation for cryogenic flows.

For turbomachinery simulations, CFD technology is currently limited to simulating mean cavitating performance at design conditions for idealized liquids (no thermodynamic effects). Typical simulations show comparisons with data at design conditions for the head coefficient and the critical \( N_a \) number at which performance breakdown occurs (Hosangadi\(^7\), Athavale and Singhal\(^6\), Dupont and Okamura\(^9\), Medvitz\(^10\)). However flows at off-design conditions, where large scale unsteadiness and high dynamic pressure loads are observed, cannot at this point be reliably predicted. Simulation of cavitation instabilities and rotational cavitation modes in pumps have not been simulated by any group to the best of our knowledge. Thus, to develop a CFD simulation framework that can eventually simulate unsteady, off-design performance of liquid rocket turbopumps that use cryogenic working fluids, significant development will be required from the current level of technology.

The focus of this paper is on the simulation of cavitating, cryogenic inducers where real fluid property variations have a significant impact on the suction performance of turbopumps. To simulate this class of flows, a generalized multi-phase formulation has been developed that rigorously models thermal effects of phase change and the accompanying property variations. Thermal equilibrium is assumed and fluid thermodynamic properties are specified along the saturation line using the NIST-12 databank (Hosangadi and Ahuja\(^11\)). This framework will be the foundation for the eventual goal of our effort to model unsteady performance of liquid rocket pumps. Within this mind, the equations are cast in an acoustically accurate form and is an extension of earlier work by the authors for idealized liquids (Ahuja\(^12\)). The rigorous acoustic treatment of the gas/liquid interface captures the coupling of the phase change process with the dynamic pressure field which is an important attribute for unsteady cloud cavitation problems where high amplitude, localized pressure spikes can occur (Hosangadi\(^13\)).

The numerical code utilized in our simulations is the CRUNCH CFD\(^8\) code, which is a multi-element based unstructured code \(^14\)\(^15\). The underlying philosophy in the CRUNCH CFD\(^8\) code is to tailor the grid topology to resolving the dominant flow phenomena and the structural complexity of the problem. This is achieved by utilizing a combination of hexahedral, tetrahedral, prismatic and pyramidal elements in mesh construction. Such a framework is particularly attractive for complex turbomachine configurations, since high quality grids can be generated very efficiently with minimum skewness.

The computational framework described above has been applied to a full-scale main LOX pump inducer. Furthermore, to identify the impact of thermal effects on the inducer suction performance, a sub-scale configuration designed for water testing has also been simulated and compared with LOX configuration. The performance for both configurations is simulated at the design flow rate over a range of inlet pressures ranging from essentially single-phase flow all the way through head breakdown at the critical \( N_a \) number. Detailed analysis of the flowfield over this range of \( N_a \) numbers is undertaken; this includes ascertaining the location and shape of the vapor cloud, the detailed pressure distributions on the blade, and the velocity distributions both upstream and downstream of the inducer. In particular, the additional flow losses resulting under cavitating conditions and their effect on the velocity profile is identified. The effect of thermal depression on the breakdown \( N_a \) number is identified. We note that since all calculations presented here have been computed at the design flow rate large scale flow separation upstream of the inlet leading edge is not present and the total pressure rise is relatively steady.

### II. Multi-phase Equation System

The multiphase equation system is written in vector form as:

\[
\frac{\partial Q}{\partial t} + \frac{\partial E}{\partial x} + \frac{\partial F}{\partial y} + \frac{\partial G}{\partial z} = S + D,
\]

(1)

Here \( Q \) is the vector of dependent variables, \( E, F \) and \( G \) are the flux vectors, \( S \) the source terms and \( D \) represents the viscous fluxes. The viscous fluxes are given by the standard full compressible form of Navier Stokes equations. The vectors \( Q, E \) and \( S \) are given below with a detailed discussion on the details of the cavitation source terms to follow later.
Here, $\rho_m$ and $h_m$ are the mixture density and enthalpy respectively, and $\phi_g$ is the volume fraction or porosity of the vapor phase. The mixture energy equation has been formulated with the assumption that the contribution of the pressure work on the mixture energy is negligible which is a reasonable assumption for this flow regime. The source term for the vapor phase arises from rate of vapor mass generation due to cavitation $m_v$ and the corresponding source term for the energy equation is given as $m_v h_v$, where $h_v$ is the change in enthalpy resulting from the phase change and is a function of the local fluid temperature.

The mixture density and gas porosity are related by the following relations locally in a given cell volume:

$$\rho_m = \rho_g \phi_g + \rho_l \phi_l$$

$$1 = \phi_g + \phi_l$$

where $\rho_g, \rho_l$ are the physical material densities of the gas and liquid phase respectively and in general are functions of both the local temperature and pressure.

Thus far we have not made any statements defining the temperatures characterizing the liquid and vapor. In general, the liquid and vapor may not be in equilibrium locally and can have independent temperatures. Examination of temperature and pressure data for cavitation in Freon by Ruggeri 16 reveals that the saturation vapor pressure corresponding to the local fluid temperature in fact matches the local pressure measurement. This indicates local thermodynamic equilibrium that is exploited to significantly simplify Eqn. (1). The thermodynamic properties of the liquid and vapor in the cavity may now be defined by a single variable; the saturation temperature $T_s$. Hence, all thermodynamic properties (density, vapor pressure, viscosity, etc) of both the liquid and the vapor phase may be generated as a tabular function of the saturation temperature. In our study here, these properties were generated from the Standard thermodynamic database 12 available from NIST for pure fluids.

The equation system as formulated in Eqn. (1) is very stiff since the variations in density are much smaller than the corresponding changes in pressure. Therefore to devise an efficient numerical procedure we wish to transform Eqn. (1) to a pressure based form where pressure rather than density is the variable solved for. An acoustically accurate two-phase form of Eqn. (1) is first derived, followed by a second step of time-scaling or preconditioning to obtain a well-conditioned system. We begin by defining the acoustic form of density differential for the individual gas and liquid phase as follows:

$$d\rho_g = \frac{1}{c_g^2} dP, \quad d\rho_l = \frac{1}{c_l^2} dP$$

Here $c_g$ is the isothermal speed of sound in the pure gas phase, and $c_l$ is the corresponding isothermal speed of sound in the liquid phase, which is a finite-value. We note that in Eqn. (5) the variation of the density with temperature has been neglected in the differential form. This assumption was motivated by the fact that the temperature changes are primarily due to the source term and not by the pressure work on the fluid i.e. the energy equation is a scalar equation. This simplifies the matrix algebra for the upwind flux formulation significantly, at the potential expense of numerical stability in a time-marching procedure. However, more importantly, there is no
impact on the accuracy since the fluid properties themselves are taken directly from the thermodynamic data bank for each fluid.

Following the discussion above, the differential form of the mixture density $\rho_m$ using Eqn. (5) is written as,

$$d\rho_m = (\rho_t - \rho_L) d\phi_t + \frac{1}{c_p^2} dP$$

$$+ \left( \frac{1}{c_p^2} = \frac{\phi_t}{c_p^2} + \frac{\phi_L}{c_p^2} \right)$$

(6)

Here, $c_p$ is a variable defined for convenience and is not the acoustic speed, $c_m$, in the mixture, which will be defined later. Using Eqn. (6), Eqn. (1) may be rewritten as:

$$\Gamma \frac{\partial \rho}{\partial t} + \frac{\partial \rho u}{\partial x} + \frac{\partial \rho w}{\partial y} + \frac{\partial \rho h}{\partial z} = S + D_v$$

(7)

and,

$$Q_v = \left[ \rho, u, v, w, \phi_t, \phi_L, k, e \right]^T$$

(8)

The numerical characteristics of the Eqn. (7) are studied by obtaining the eigenvalues of the matrix, $\Gamma^{-1} \left( \frac{\partial \rho}{\partial t} \right)$. The eigenvalues of the system are derived to be:

$$\lambda = (u + c_m, u - c_m, u, u, u, u, u, u)$$

(9)

where $c_m$ turns out to be the well-known, harmonic expression for the speed of sound in a two-phase mixture and is given as:

$$\frac{1}{c_m^2} = \rho_m \left[ \frac{\phi_t}{c_t^2} + \frac{\phi_L}{c_L^2} \right]$$

(10)

The behavior of the two-phase speed of sound indicates that at either limit the pure single-phase acoustic speed is recovered. However, away from the single-phase limits, the acoustic speed rapidly drops below either limit value and remains at the low-level in most of the mixture regime. As a consequence, the local Mach number in the interface region can be large even in low speed flows.

To obtain an efficient time-marching numerical scheme, preconditioning is now applied to the system in Eqn. (7), in order to rescale the eigenvalues of the system so that the acoustic speeds are of the same order of magnitude as the local convective velocities.

III. Cavitation Source Terms

In the present effort, the cavitation source term is defined via a simplified non-equilibrium, finite rate form as follows:

$$m_i = K_f \rho_t \phi_t + K_s \rho_s \phi_s$$

(11)

where the constant $K_f$ is the rate constant for vapor being generated from liquid in a region where the local pressure is less than the vapor pressure. Conversely, $K_s$ is the rate constant for reconversion of vapor back to liquid in
regions where the pressure exceeds the vapor pressure. Here, the rate constants are specified using the form given by Merkle\textsuperscript{17}.

\begin{align}
K_b &= \begin{cases} 
0 & \text{if } p < p_i \\
\frac{1}{t_b \Delta L_b} \left[ \frac{p - p_i}{2 \rho_i Q_\infty^2} \right] & \text{if } p > p_i
\end{cases} \\
K_f &= \begin{cases} 
0 & \text{if } p < p_i \\
\frac{1}{t_f \Delta L_f} \left[ \frac{p - p_i}{2 \rho_i Q_\infty^2} \right] & \text{if } p > p_i
\end{cases}
\end{align}

\[ p_i = p_\infty - \frac{1}{2} \rho_\infty Q_\infty^2 \text{ Cav. No.} \]

\[ t_f = \text{Time constant for vapor formation} \]

\[ t_b = \text{Time constant for liquid reconversion} \]

\[ \text{Cav. No.} = \frac{p_\infty - p_i}{\frac{1}{2} \rho_\infty Q_\infty^2} \]

We note that for steady attached cavitation this simplified form may be adequate since the cavitation time scales do not interact with the fluid time scales if the cavitation rate constants are fast enough. For unsteady cavitation modeling, however, it becomes essential to integrate bubble dynamics within a dense cloud framework wherein both the number density and mean local radius of the bubbles in an evolving cloud are tracked. The development of a more rigorous non-equilibrium source term model is a topic of ongoing research.

IV. Validation Study For Cavitation In Cryogenic Fluids

The multi-phase formulation described above has been validated extensively by simulating experiments by Hord\textsuperscript{1} on a cavitating hydrofoil. Hord performed sub-scale tests using both liquid nitrogen and hydrogen in a blow-down tunnel. The details of the tunnel and the hydrofoil geometry are given in Fig. 1. The tunnel width is 1 inch while the hydrofoil width is 0.312 inches. Hence considerable blockage effects from the tunnel wall are present and this necessitated modeling the tunnel geometry in the simulations. To ensure that the tunnel blockage effects were being correctly modeled we performed single-phase, non-cavitating simulations and have compared it to Hord's non-cavitating data in Fig. 2. Excellent agreement is obtained giving confidence that the tunnel interaction is being captured. We note that the single-phase solution is insensitive to the Reynolds number since both liquid hydrogen and nitrogen show identical pressure profiles.

Cavitating results in liquid nitrogen for one flow condition are discussed here. We refer the reader to Hosangadi and Ahuja\textsuperscript{11} for additional validation studies in both liquid nitrogen and hydrogen. The operating range of liquid nitrogen varies from roughly 70K - 100K. The variation of properties along the saturation line for liquid and vapor densities as well as the vapor pressure are shown in Fig. 3. At 89 K, the slope of vapor pressure curve indicates a 16 KPa increase for a 1 K change in temperature.

The general characteristics of the cavitating flowfield are shown in Figure 4(a-d) for tunnel conditions in Run 290C (Table I).
Figure 4a shows the temperature profile in the cavity. The strong temperature depression at the leading edge of the cavity is evident with the gradual temperature recovery due to condensation in the rear of the cavity. The vapor volume fraction is qualitatively compared with a typical flow visualization of the flow (note that the flow conditions at which this visualization was done are not reported in the Hord’s report). The overall shape and features of the cavity appear to be similar. The computed pressure field indicates strong interaction between the cavity and the tunnel wall which is expected due to the relative scales of the geometry.

Table 1. Run Conditions For Liquid Nitrogen Cases.

<table>
<thead>
<tr>
<th>RUN NUMBER</th>
<th>FREESTREAM TEMP. (K)</th>
<th>FREESTREAM VEL. (M/S)</th>
<th>CAVITATION NUMBER</th>
<th>CAVITY LENGTH (CM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>289C</td>
<td>88.64</td>
<td>23.5</td>
<td>1.55</td>
<td>2.29</td>
</tr>
<tr>
<td>290C</td>
<td>83.06</td>
<td>23.9</td>
<td>1.70</td>
<td>1.9</td>
</tr>
<tr>
<td>293A</td>
<td>77.64</td>
<td>24.0</td>
<td>1.75</td>
<td>1.52</td>
</tr>
<tr>
<td>294F</td>
<td>77.94</td>
<td>9.8</td>
<td>1.78</td>
<td>1.52</td>
</tr>
</tbody>
</table>

The quantitative comparisons of pressure and temperature depression in the cavity are compared with experimental data in Fig. 5. Note that the pressure values plotted are \((P - P_{\infty})\). For a non-cryogenic case this value would be zero in the cavity, while values below zero in the cryogenic case indicate pressure depression due to thermal effects. In general excellent comparison is obtained for the leading edge temperature depression of approximately 2.5 K. The temperature recovery within the cavity compares well both in their slope as well as the
length of the cavity. The temperature rise in the cavity closure region shows some differences; the computed solution recovers to the freestream value more quickly than does the data, which doesn’t quite fully recover to the freestream value. As per the discussion by Hord\(^1\), this probably was due to the unsteady effects in the cavity closure region whereby the thermocouples were not always enclosed in vapor giving erroneous readings. The instrumentation error given for the chromel-gold thermocouples is 0.20 K and the computed results are within the uncertainty of the experiments.

The comparison of the pressure depression in Fig. 5a also indicates excellent overall comparison with data and within the instrumentation error bar of 0.69 \(N/cm^2\). The leading edge pressure depression is 23.5 percent relative to the freestream vapor pressure. It further illustrates why temperature effects have a substantial impact on the performance of cryogenic pumps. The experimental data plotted includes both the actual pressure measured (symbol: circle) as well as the saturation pressure values (symbol: square) corresponding to the temperature measurements. The close match between the actual pressure and saturation pressure values indicates that the thermodynamic equilibrium assumption is valid for liquid nitrogen flows.

![Pressure depression and Temperature depression](image)

**Fig. 5.** Comparison of Cavitation Hydrofoil Simulations with Data for Liquid Nitrogen.

### V. CRYOGENIC INDUCER SIMULATIONS

Simulations of a full-scale LOX inducer are presented here and compared with a sub-scale configuration designed for water testing. The sub-scale water configuration has a mass flow rate of 250.3 lbm/s, a rotational speed of 4800 rpm, and an inlet water temperature of 85°F. The sub-scale inducer has four blades with an inlet hub diameter of 1.654 inches and tip diameter of 6.016 inches. The tip gap clearance is 0.022 inches. The simulations were carried out by generating a multi-element unstructured grid for the complete 4-bladed configuration. The bulk of the grid cells between the blades is composed of hexahedral cell elements while tetrahedral and prismatic cells are used in the tip gap region as well as the region between hexahedral blocks to reduce grid skewness. The total grid size was approximately 4.5 million cells and the solution was obtained by decomposing the grid on 32 processors on a parallel machine.

The full-scale LOX inducer was simulated over a range of \(N_u\) numbers ranging from 10000 to 30000 to model inducer performance from mildly cavitating conditions through head breakdown. Figure 6 shows the vapor volume fraction iso-surface over four different \(N_u\) numbers. We note the formation of the primary cavitation front at the leading edge which grows preferentially near the shroud as the \(N_u\) number increases. However, interestingly we also evidence the formation of a second cavitation front at a bend in the blade where the blade goes to its full thickness. The corresponding pressure distribution on the blade suction side surface is shown in Fig. 7. The loss of pressure generation corresponds to the advancement of the vapor formation as expected. The temperature depression levels due to thermal effects of cavitation are shown in Fig. 8. Lower temperatures occur in regions where vaporization of liquid takes place. This figure indicates that the temperature depression is localized around the leading edge of the blade; the primary location of vapor formation, therefore, is at the leading edge while in the remainder of the vapor cloud condensation back to liquid is taking place. This is consistent with the attention given to the leading edge shape in design tools for inducers.

The pressure distribution on the both the pressure and suction side of the blade surfaces are shown at different \(N_u\) numbers in Fig. 9 for both the water and the LOX case at a location close to the blade tip. We observe that the

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loading profiles for the water and LOX configuration are similar at lower N,, numbers but that the water case breaks down earlier. As cavitation sets in, the pressure on the suction side drops to the nominal vapor pressure value and this front advances along the suction side consistent with the vapor cloud location. An interesting point to note is that while the pressure levels on the pressure side drop under cavitating conditions, the leading edge continues to take on load up until an N,, number of 22000. At 25000, the water case has broken down but the LOX configuration is continuing to generate head indicating that thermal effects have improved the mean performance of the pump.

To understand the cavitation loss mechanisms better, we plot the velocity profiles downstream of the blade exit for the non-cavitating case as well as the cavitating case at N,, of 25000 in Fig. 10. The axial velocity plots for both the non-cavitating and the cavitating case show a slowdown near the shroud due to losses. However, the slow down in the axial velocity is more pronounced in the cavitating case indicating larger losses. The differences in the axial velocity also lead to differences in the flow angle at the exit. The loss mechanism in the cavitating case was further investigated by plotting the streamlines near the shroud (Fig. 11). The streamlines in the cavitating case are affected strongly by the blockage due to the vapor cloud. The flow passage gets constricted at the inlet causing the flow to accelerate and squeeze through the narrower effective flow passage. Conversely at the cavity closure region, the fluid experiences a sudden expansion in flow area as the blockage ends. This causes larger flow turning losses that are reflected in the exit velocity profiles.

**Fig. 6.** Vapor Isosurface ($\Phi = 0.4$) Contours at Various N,, Numbers in LOX Inducer.

**Fig. 7.** Blade Suction Side Pressure at Various N,, Numbers in LOX Inducer.

**Fig. 8.** Blade Suction Side Temperature at Various N,, Numbers in LOX Inducer.
The comparison of the head coefficient generated by the inducer as a function of the $N_a$ number are plotted in Fig. 12 for both the full-scale LOX case as well as the sub-scale water test. We note that although we did not discuss the details of the sub-scale water simulations, the qualitative flow features for the water configuration were very similar to the LOX case. In particular, the location and shape of the vapor cloud had similar features although the vapor cloud in the LOX case evolved at a slower pace due to thermal depression effects. The effect of the thermal depression is apparent on the critical $N_a$ number at which breakdown occurs. In the water case breakdown occurs in the range of 23000-24000 while in the LOX case breakdown occurs around 30000. Furthermore, in the LOX case, the slope of head drop-off is more gradual as the breakdown point is approached probably due to the moderating influence of the thermal effects. The single-phase result for the LOX case shows a slightly lower head rise than the single-phase result for water despite the higher Reynolds number for the LOX case. This may due to the much higher tip-gap losses in the LOX case; we note that the LOX case was run with the tip gap from the sub-scale geometry being scaled up by the same scale factor.
VI. SUMMARY

A generalized multi-phase formulation has been developed for cavitation in "real" fluid flows (e.g., cryogenic fluids) that operate at temperatures close to their critical temperatures. The formulation is based on a thermal equilibrium assumption between the vapor and liquid in the cavitating region. Fluid thermodynamic properties are specified along the saturation line as a function of temperature using the NIST-12 data bank. The energy equation for the mixture is solved in conjunction with the mass and momentum conservation and the evaporative cooling effects of cavitation are accounted for rigorously.

Detailed validation has been presented to quantify temperature and pressure depression in liquid nitrogen and hydrogen by simulating the cavitating hydrofoil experiments of Hord'. In general excellent comparison of the
leading edge pressure and temperature depression was observed for all cases. The temperature profiles in the cavity closure show more variation and this is in part due to the unsteadiness in the data, which was not characterized. The multi-phase formulation was used to simulate a full-scale LOX inducer as well as the corresponding subscale configuration designed for water testing. The head breakdown appears to be in the range of 23000-24000 Ns for the water model while the inducer breaks down at 30000 Ns for the LOX. The increase in suction performance is attributed to effect of thermal depression in LOX. Furthermore the head breakdown in the LOX case was found to be more gradual than in the water case particularly near the breakdown point. The overall cavitation characteristics, however, were otherwise similar between the LOX and the water case and in particular a second cavitation front was observed at a bend in the blade where it goes to its full thickness. The temperature depression was observed primarily at the leading edge with the largest temperature drop being at the leading edge tip. The velocity profiles downstream of the blade exit were analyzed. Both the single-phase and cavitating cases showed a slow down in the axial velocity near the shroud. However the slow down was much more in the cavitating case due to the additional losses in the cavitating case. These additional losses were attributed to the increased flow turning losses in the cavitating case due to blockage from the vapor cavity.

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VIII. REFERENCES