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CFD WITH OPENSOURCE SOFTWARE

A course at Chalmers University of Technology Taught by Håkan Nilsson

Implementing a Zwart-Gerber-Belamri cavitation model

Developed for OpenFOAM-1806 and gnuplot 5.2

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December 20, 2018

Learning outcomes

The reader will learn:

How to use it:

• How to adapt and run the throttle tutorial.

The theory of it:

• The theory of Schnerr-Sauer (SS) and Zwart-Gerber-Belamri (ZGB) cavitation models.

How it is implemented:

• How to implement the ZGB cavitation model in the phaseChangeTwoPhaseMixtures class.

How to modify it:

• How to modify the mass transfer rates.

Prerequisites

The reader is expected to know the following in order to get maximum benefit out of this report:

- Fundamentals of fluid mechanics and bubble dynamics e.g. book by Brennen [1] or Franc and Michel [2].
- Some understanding of cavitation modelling, e.g. Schnerr and Sauer [3] and Zwart et. al. [4]
- Run standard document tutorials like throttle tutorial and/or cavitatingBullet tutorial.
- Basic post-processing in paraFoam have been used to visualize the results. However, paraFoam is not in the scope of this tutorial.

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Chapter 1

Introduction

This tutorial describes how to add the Zwart-Gerber-Belamri (ZGB) cavitation model to the phaseChangeTwoPhaseMixtures class used by interPhaseChangeFoam solver. Both the ZGB model and the default Schnerr-Sauer (SS) model are described in this chapter. The solver is described briefly in Chapter 2, followed by a simple test case with the default SS cavitation model. In Section 2.2, the ZGB cavitation model is implemented and the same case is solved with this additional model.

1.1 Bubble dynamics and cavitation models

The dynamics of a single bubble subjected to a pressure field can be described by

$$\frac{p_b - p_\infty}{\rho_l} = R \frac{d^2 R}{dt^2} + \frac{3}{2} \left(\frac{dR}{dt}\right)^2 + \frac{4\nu_l}{R} \frac{dR}{dt} + \frac{2S}{\rho_l R}$$
(1.1)

which is known as the Rayleigh-Plesset equation [1]. R = R(t) is the bubble radius, S is the surface tension coefficient, p_b is the bubble pressure and p_{∞} is the far-field pressure. ρ_l and ν_l is the liquid density and liquid kinematic viscosity respectively. The Rayleigh-Plesset equation assumes that bubbles are spherical and symmetric and that thermal effects are negligible. By further neglecting higher order terms and the effects of surface tension and viscosity, eq. (1.1) can be simplified to

$$\frac{dR}{dt} = \sqrt{\frac{2}{3}} \frac{p_b - p}{\rho_l} \tag{1.2}$$

Due to evaporation and condensation, mass transfer occur between the liquid- and vapor phase. This mass transfer is governed by a transport equation for the liquid fraction

$$\frac{\partial}{\partial t} \left(\alpha_l \rho_l \right) + \nabla \cdot \left(\alpha_l \rho_l \vec{v} \right) = \dot{m} \tag{1.3}$$

where ρ_l is the density of the liquid phase and \dot{m} is the mass transfer rate. In interPhaseChange-Foam, which later will be described, the mass transfer is decomposed to vaporization- and condensation terms

$$\dot{m} = \alpha_l \dot{m}_{\alpha}^- + (1 - \alpha_l) \, \dot{m}_{\alpha}^+ = \alpha_l \left(\dot{m}_{\alpha}^- - \dot{m}_{\alpha}^+ \right) + \dot{m}_{\alpha}^+ \tag{1.4}$$

where \dot{m}_{α}^{-} and \dot{m}_{α}^{+} are the destruction of liquid from evaporation and creation of liquid from condensation, respectively. α_{l} is the liquid volume fraction.

Finally, eq. (1.3) is implemented in terms of volume fraction and decomposed volume transfer rates, \dot{V}^- and \dot{V}^+ ,

$$\frac{\partial \alpha_l}{\partial t} + \nabla \cdot (\alpha_l \vec{v}) = \left(\nabla \cdot \vec{v} + \dot{V}^- - \dot{V}^+ \right) \alpha_l + \dot{V}^+ \tag{1.5}$$

where the volume transfer rate, \dot{V} , and velocity divergence, $\nabla \cdot \vec{v}$,

$$\dot{V} = \left(\frac{1}{\rho_l} - \alpha_l \left(\frac{1}{\rho_l} - \frac{1}{\rho_v}\right)\right) \dot{m}$$
(1.6)

$$\nabla \cdot \vec{v} = \left(\frac{1}{\rho_l} - \frac{1}{\rho_v}\right) \dot{m} \tag{1.7}$$

have been used. The divergence of the velocity field, eq. (1.7), also appears on the RHS of the pressure correction equation. In this case, the mass transfer can be decomposed to

$$\dot{m}_p = \frac{\dot{m}}{(p - p_v)} = \dot{m}_p^- - m_p^+ \tag{1.8}$$

where p_v is a threshold pressure (the saturation pressure for the liquid). For a full description of the discretized equations and the decomposition of mass transfer rates, refer to Asnaghi [5].

1.1.1 Schnerr-Sauer cavitation model

Schnerr and Sauer [3] derived an expression for the net mass change rate from the generalized vapor transport equation. This model is implemented in OpenFoam through the mass source terms \dot{m}_{α}^{-} and \dot{m}_{α}^{+}

$$\dot{m}_{\alpha}^{-} = C_v \left(1 + \alpha_{nuc} - \alpha_l\right) \frac{3\rho_v \rho_l}{\rho_m R_B} \sqrt{\frac{2}{3\rho_l} \frac{1}{|p - p_v|}} \min\left(p - p_v, 0\right)$$
(1.9)

$$\dot{m}_{\alpha}^{+} = C_{c} \alpha_{l} \frac{3\rho_{v} \rho_{l}}{\rho_{m} R_{B}} \sqrt{\frac{2}{3\rho_{l}} \frac{1}{|p - p_{v}|}} max \left(p - p_{v}, 0\right)$$
(1.10)

which follow the decomposition in eq. (1.4). ρ_v is the vapor density. ρ_m is the mixture density, a volume weighted average of the liquid and vapor density. α_{nuc} is a numerical nucleation term that is introduced to initialize cavitation. Without the presence of this term, eq. (1.9) equals zero for all pressures in a pure liquid and cavitation would never occur. The relation between nucleation site volume fraction, α_{nuc} , and bubble radius, R_B , is described according to

$$\alpha_{nuc} = \frac{\frac{1}{6}n_b \pi d_{nuc}^3}{1 + \frac{1}{6}n_b \pi d_{nuc}^3} \tag{1.11}$$

$$R_B = \left(\frac{3}{4\pi n_b} \frac{1 + \alpha_{nuc} - \alpha_l}{\alpha_l}\right)^{\frac{1}{3}} \tag{1.12}$$

where n_b is the bubble number density and d_{nuc} is the nucleation site diameter. It can be seen that the mass transfer rates for evaporation and condensation are differentiated by the coefficients C_v and C_c , respectively. The default value is however 1 for both coefficients.

The phase change terms in the pressure correction equation are defined similarly, but according to the decomposition in eq. (1.8),

$$\dot{m}_{p}^{-} = -C_{v}\alpha_{l}\left(1 + \alpha_{nuc} - \alpha_{l}\right)\frac{3\rho_{v}\rho_{l}}{\rho_{m}R_{B}}\sqrt{\frac{2}{3\rho_{l}}\frac{1}{|p - p_{v}|}neg\left(p - p_{v}\right)}$$
(1.13)

$$\dot{m}_{p}^{+} = C_{c}\alpha_{l}\left(1 - \alpha_{l}\right)\frac{3\rho_{v}\rho_{l}}{\rho_{m}R_{B}}\sqrt{\frac{2}{3\rho_{l}}\frac{1}{|p - p_{v}|}pos\left(p - p_{v}\right)}$$
(1.14)

where the neg function returns -1 when the input is negative and zero otherwise, and the pos function returns 1 when the input is positive and zero otherwise.

1.1.2 Zwart-Gerber-Belamri cavitation model

In the ZGB cavitation model, the mass transfer rates are given as

$$\dot{m}_{\alpha}^{-} = -C_{v}r_{nuc}\frac{3\rho_{v}}{R_{B}}\sqrt{\frac{2}{3\rho_{l}}\frac{1}{|p-p_{v}|}}min\left(p-p_{v},0\right)$$
(1.15)

$$\dot{m}_{\alpha}^{+} = C_{c} \frac{3\rho_{v}}{R_{B}} \sqrt{\frac{2}{3\rho_{l}} \frac{1}{|p - p_{v}|}} max \left(p - p_{v}, 0\right)$$
(1.16)

where r_{nuc} is the nucleation site volume fraction. Like the SS model, it is based on the simplified Rayleigh-Plesset equation but assumes a constant nucleation site volume. Thus, both the nucleation site volume fraction, r_{nuc} , and nucleation site radius, R_B , are model constants. According to Zwart et. al. [4], these constants are $r_{nuc} = 5.0 \times 10^{-4}$ and $R_B = 1.0 \times 10^{-6}$ m by default. Following the same decomposition as in the SS model, the phase change terms in the pressure correction equation are

$$\dot{m}_{p}^{-} = -C_{v}r_{nuc}\alpha_{l}\frac{3\rho_{v}}{R_{B}}\sqrt{\frac{2}{3}\frac{|p-p_{v}|}{\rho_{l}}}neg\left(p-p_{v}\right)$$
(1.17)

$$\dot{m}_{p}^{+} = C_{c} \left(1 - \alpha_{l}\right) \frac{3\rho_{v}}{R_{B}} \sqrt{\frac{2}{3} \frac{|p - p_{v}|}{\rho_{l}}} pos\left(p - p_{v}\right)$$
(1.18)

Chapter 2

interPhaseChangeFoam solver

The interPhaseChangeFoam solver is a fully transient solver. Turbulence is generic, i.e. it can be solved using either laminar, RAS or LES. It is based on the PIMPLE pressure correction. It solves for two isothermal, incompressible fluids using a Volume of Fluids (VoF) approach where momentum equations are solved for the mixture. interPhaseChangeFoam supports mass transfer through cavitation and the available models are Merkle, Kunz, and Schnerr-Sauer. The Schnerr-Sauer model is implemented as described in Section 1.1.1. Note that interPhaseChangeFoam solves for the liquid fraction, which will affect how the mass transfer rate is implemented.

interPhaseChangeFoam

_Allwclean _Allwmake _alphaControls.H _alphaEqn.H _alphaEqnSubCycle.H createFields.H _interPhaseChangeDyMFoam/ _interPhaseChangeFoam.C _Make/ _pEqn.H phaseChangeTwoPhaseMixtures/ Kunz/ lnInclude/ Make/ _Merkle/ _phaseChangeTwoPhaseMixture/ SchnerrSauer/ UEqn.H

The interPhaseChangeFoam directory contains both Allwmake and Allwclean. You can also find a solver for dynamic meshing, interPhaseChangeDyMFoam, which will not be considered in this tutorial. There are equations for p, U and alpha, which will be described later. One interesting thing is that there is a class named phaseChangeTwoPhaseMixtures with its own Make directory. This class is located in the solver, which is not the more common approach in OpenFOAM. We will come back to both this class and the equations, but first take a look in interPhaseChangeFoam.C.

```
interPhaseChangeFoam.C
                                                                       mixture->correct();
Info<< "\nStarting time loop\n" << endl;</pre>
                                                                        #include "alphaEqnSubCycle.H"
while (runTime.run())
                                                                        interface.correct();
{
    #include "readTimeControls.H"
                                                                        #include "UEqn.H"
    #include "CourantNo.H"
    #include "setDeltaT.H"
                                                                        // --- Pressure corrector loop
                                                                        while (pimple.correct())
    runTime++:
                                                                        ſ
                                                                            #include "pEqn.H"
    Info<< "Time = " << runTime.timeName() << nl << endl;</pre>
                                                                        }
    // --- Pressure-velocity PIMPLE corrector loop
                                                                        if (pimple.turbCorr())
    while (pimple.loop())
                                                                        {
                                                                            turbulence->correct();
        #include "alphaControls.H"
                                                                        }
                                                                   }
        surfaceScalarField rhoPhi
        (
                                                                   runTime.write();
            IOobject
                                                                   runTime.printExecutionTime(Info);
                                                               }
                 "rhoPhi",
                 runTime.timeName(),
                                                               Info<< "End\n" << endl;</pre>
                mesh
            ).
            mesh.
                                                               return 0:
            dimensionedScalar(dimMass/dimTime, Zero)
        );
                                                           }
```

As mentioned, interPhaseChangeFoam is a PIMPLE solver. In the outer loop, before the pressure correction, the interfaces are corrected by calling alphaEqnSubCycle.H. and solving the (liquid) phase transport equation. The phase equation is found in alphaEqn.H. On the LHS we have, in order, the transient term $\frac{\partial \alpha}{\partial t}$ solved with first order Euler Scheme, the convective term $\nabla \cdot (\alpha \vec{v})$ solved with upwind flux, and the convective source term $\nabla \cdot (\alpha_l \vec{v})$ (on the RHS in eq. (1.5)). The other source terms can be seen on the right according to eq. (1.5). As indicated by the names of the functions, vDotvAlphal and vDotcAlphal are volume source terms.

```
alphaEqn.H
Pair<tmp<volScalarField>> vDotAlphal =
    mixture->vDotAlphal();
const volScalarField& vDotcAlphal = vDotAlphal[0]();
const volScalarField& vDotvAlphal = vDotAlphal[1]();
const volScalarField vDotvmcAlphal(vDotvAlphal - vDotcAlphal);
fvScalarMatrix alpha1Eqn
    fv::EulerDdtScheme<scalar>(mesh).fvmDdt(alpha1)
  + fv::gaussConvectionScheme<scalar>
    (
         mesh,
         phi,
         upwind<scalar>(mesh, phi)
    ).fvmDiv(phi, alpha1)
   - fvm::Sp(divU, alpha1)
    fvm::Sp(vDotvmcAlphal, alpha1)
    vDotcAlphal
):
```

The volume source terms vDotvAlphal (vaporization) and vDotcAlphal (condensation) are components of the pair called vDotAlphal. This is found in the phaseChangeTwoPhaseMixtures class, which is a part of the interPhaseChageFoam solver.

```
phaseChangeTwoPhaseMixture.C
Foam::Pair<Foam::tmp<Foam::volScalarField>>
Foam::phaseChangeTwoPhaseMixture::vDotAlphal() const
{
    volScalarField alphalCoeff(1.0/rho1() - alpha1_*(1.0/rho1() - 1.0/rho2()));
   Pair<tmp<volScalarField>> mDotAlphal = this->mDotAlphal();
    return Pair<tmp<volScalarField>>
    (
        alphalCoeff*mDotAlphal[0],
        alphalCoeff*mDotAlphal[1]
   );
}
Foam::Pair<Foam::tmp<Foam::volScalarField>>
Foam::phaseChangeTwoPhaseMixture::vDotP() const
{
    dimensionedScalar pCoeff(1.0/rho1() - 1.0/rho2());
    Pair<tmp<volScalarField>> mDotP = this->mDotP();
    return Pair<tmp<volScalarField>>(pCoeff*mDotP[0], pCoeff*mDotP[1]);
}
```

The volume source terms vDotAlphal are related to the mass source terms mDotAlphal with the coefficient alphaCoeff, corresponding to eq. (1.6). The mass source terms are also part of the phaseChangeTwoPhaseMixtures class, but specific for each individual cavitation model e.g. Schnerr-Sauer which is found in SchnerrSauer.C.

```
SchnerrSauerr.C
Foam::tmp<Foam::volScalarField>
Foam::phaseChangeTwoPhaseMixtures::SchnerrSauer::pCoeff
 const volScalarField& p
) const
volScalarField limitedAlpha1(min(max(alpha1_, scalar(0)), scalar(1)));
volScalarField rho
(
    limitedAlpha1*rho1() + (scalar(1) - limitedAlpha1)*rho2()
):
return
    (3*rho1()*rho2())*sqrt(2/(3*rho1()))
   *rRb(limitedAlpha1)/(rho*sqrt(mag(p - pSat()) + 0.01*pSat()));
}
Foam::Pair<Foam::tmp<Foam::volScalarField>>
Foam::phaseChangeTwoPhaseMixtures::SchnerrSauer::mDotAlphal() const
ſ
    const volScalarField& p = alpha1_.db().lookupObject<volScalarField>("p");
    volScalarField pCoeff(this->pCoeff(p));
    volScalarField limitedAlpha1(min(max(alpha1_, scalar(0)), scalar(1)));
    return Pair<tmp<volScalarField>>
    (
        Cc_*limitedAlpha1*pCoeff*max(p - pSat(), p0_),
        Cv_*(1.0 + alphaNuc() - limitedAlpha1)*pCoeff*min(p - pSat(), p0_)
   ):
}
```

SchnerrSauer.C returns a pair mDotAplhal with the mass transfer rates. The member functions pCoeff and mDotAlphal together make the mass source terms seen in eq. (1.9)-(1.10). limitedAlpha1 is just a bounded liquid fraction, since this value should be between 0 and 1. 0.01*pSat() is a small addition that prevents zero in the denominator. rRb is the reversed bubble radius $\frac{1}{R_B}$ according to eq. (1.12). In mDotAlphal we can see alphaNuc, mentioned earlier in eq. (1.11), which prevents zero value in the denominator. The volume fraction of nucleation sites, alphaNuc, is a function of the bubble number density, n, and the initial bubble diameter, dNuc, which are both model constants. rRb and alphaNuc are also specified in SchnerrSauer.C, but are excluded here for brevity.

2.1 throttle tutorial

This section describe how to pre-process, run and post-process a case involving incompressible flow through a cavitating jet. The throttle geometry consists of two volumes connected through a narrow section, shown in Figure 2.1. At the left side is a pressure inlet, and the right side a pressure outlet. If the pressure difference between inlet and outlet is large enough, the change in dynamic pressure in the narrow section will induce cavitation.



Figure 2.1: Geometry of the throttle tutorial case.

The following steps describe how to copy the throttle tutorial to the run directory and how to modify it to solve with interPhaseChangeFoam. Start by copying and renaming the tutorial files. Clean up some files that are not used.

```
cp -r $FOAM_TUTORIALS/multiphase/cavitatingFoam/RAS/throttle \
$FOAM_RUN/throttleSchnerr
cd $FOAM_RUN/throttleSchnerr
rm All*
rm system/topo*
rm system/refineMeshDict
```

To run the tutorial with interPhaseChangeFoam, some changes are needed in initial- and boundary conditions and in the system directory.

2.1.1 Boundary and initial conditions

Boundary and initial conditions are found in the 0 directory where the following variables are found

alpha.vapour	р
k	rho
nut	U
omega	

The interPhaseChangeFoam solver uses the dynamic pressure and the liquid fraction. It can be seen from the boundary conditions that cavitatingFoam, which is the default solver for this tutorial, solves for the vapor volume fraction instead. Rename the initial conditions for the pressure and replace the initial conditions for the phase fraction. Reduce the outlet pressure to make sure that cavitation occurs.

```
mv 0/p 0/p_rgh
sed -i s/"uniform 100e5;"/"uniform 50e5;"/g 0/p_rgh
rm 0/alpha.vapour
cp -r $FOAM_TUTORIALS/multiphase/interPhaseChangeFoam/cavitatingBullet/0.orig/alpha.water \
0/alpha.water
sed -i s/"inletOutlet"/"fixedValue"/g 0/alpha.water
sed -i s/"inletValue"/"value"/g 0/alpha.water
sed -i s/"bullet"/"frontBack"/g 0/alpha.water
```

sed -i s/"zeroGradient"/"empty"/g 0/alpha.water sed -i s/"symmetry"/"zeroGradient"/g 0/alpha.water

2.1.2 Transport properties

Replace /constant/transportProperties and remove /constant/thermodynamicProperties (since interPhaseChangeFoam solves for isothermal fluids). Add the gravity g which is used to subtract the hydrostatic pressure. Leave the default $k-\omega$ SST turbulence model.

```
rm constant/thermodynamicProperties
rm constant/transportProperties
cp -r $FOAM_TUTORIALS/multiphase/interPhaseChangeFoam/cavitatingBullet\
/constant/transportProperties constant/.
cp -r $FOAM_TUTORIALS/multiphase/interPhaseChangeFoam/cavitatingBullet/constant/g \
constant/.
```

2.1.3 Discretization schemes and solution control

As mentioned, interPhaseChangeFoam solves for the liquid fraction and not the vapor fraction. Solver settings for the liquid fraction, and dynamic pressure, are found in system/fvSolution and system/fvSchemes. Change solvers in system/fvSolution to

ol	vers					precondition	ner	GAMG;
-	"alj {	pha.water.*"				tolerance relTol		1e-6; 0:
		cAlpha	0;					
		nAlphaCorr	2;			nVcycles		2;
		nAlphaSubCycles	1;			·		
		1 5				smoother		DICGaussSeidel;
		MULESCorr	yes;					
		nLimiterIter	5;		};			
					to	lerance	1e-7	7;
		solver	<pre>smoothSolver;</pre>		re	lTol	0;	
		smoother	symGaussSeidel;		ma	xIter	50;	
		tolerance	1e-8;	}				
		relTol	0;					
		maxIter	10;	"1	corr	.*"		
	};			{				
					\$p.	_rgh;		
	"U.	*"			re	lTol	0;	
	{			}				
		solver	<pre>smoothSolver;</pre>					
		smoother	symGaussSeidel;	Pł	ni			
		tolerance	1e-6;	{				
		relTol	0;		\$p.	_rgh;		
	};				re	lTol	0;	
				};				
	p_r	gh						
	{			"	Uk	omega)"		
		solver	GAMG;	{				
		tolerance	1e-8;		so	lver	smoo	othSolver;
		relTol	0.1;		sm	oother	sym(GaussSeidel;
					to	lerance	1e-(08;
		smoother	DICGaussSeidel;		re	lTol	0.1	;
				}				
		_		",	U k	omega)Final"		
		maxIter	50;	ł		_		
	};				SO.	lver	smoo	othSolver;
					sm	oother	sym(GaussSeidel;
	p_r	ghFinal			to	lerance	1e-(08;
	ł		200		re	1101	0;	
		solver	PCG;	۲ ۲				
		preconditioner		1				
		1						

and add after solvers

In /system/fvSchemes, add the following divSchemes.

div(phi,alpha)	Gauss	vanLeer;
div(phirb,alpha)	Gauss	linear;

2.1.4 Run and post-process

Create a blockMesh and run interPhaseChangeFoam.

blockMesh >& log_blockMesh&
interPhaseChangeFoam >& log_run&

At the final time step, the velocity shown in Figure 2.2a has got stabilized and the jet takes a symmetric shape. From the water fraction shown in Figure 2.2b, it can be seen that the throttle cavitates.

Add a singleGraph file to the system directory to extract pressure values along a center line through the domain.

```
touch system/singleGraph
```

Change the content of singleGraph to:

```
singleGraph
ſ
    start
             (0.0 \ 0.0 \ 0.0);
    end
             (0.017 \ 0.0 \ 0.0);
    fields
            (p_rgh U);
    #includeEtc "caseDicts/postProcessing/graphs/sampleDict.cfg"
    setConfig
    {
        axis
                 distance;
    }
    // Must be last entry
    #includeEtc "caseDicts/postProcessing/graphs/graph.cfg"
}
```

Run the post processing with the singleGraph function.

```
postProcess -func singleGraph >& log_postProcess&
```

Plot the pressure using gnuplot. It can be seen in Figure 2.3 that the pressure in the throttle reaches the vapor pressure, which explains the cavitation inception.

```
gnuplot
set style data linespoints
plot "postProcessing/singleGraph/0.0019/line_p_rgh.xy"
```



Figure 2.2: Last time step for throttle tutorial solved with interPhaseChangeFoam and Schnerr-Sauer cavitation model.



Figure 2.3: Pressure along a centre line.

2.2 ZGB cavitation model

Since the ZGB cavitation model is not available by default, it has to be added to the phaseChangeTwoPhaseMixtures class.

2.2.1 Implementation and compilation

Start by copying the existing cavitation models in phaseChangeTwoPhaseMixtures to your user directory. Copy the SS cavitation model (which we will use as a starting point) and rename the files.

```
cp -r $FOAM_APP/solvers/multiphase/interPhaseChangeFoam/phaseChangeTwoPhaseMixtures \
$WM_PROJECT_USER_DIR/src/phaseChangeTwoPhaseMixtures
cd $WM_PROJECT_USER_DIR/src/phaseChangeTwoPhaseMixtures
cp -r SchnerrSauer Zwart
cd Zwart
mv SchnerrSauer.C Zwart.C
mv SchnerrSauer.H Zwart.H
```

Replace each 'SchnerrSauer' with 'Zwart' in Zwart.C and Zwart.H.

sed -i s/"SchnerrSauer"/"Zwart"/g Zwart.*

Make sure that Make/files ends with the following two lines:

... Zwart/Zwart.C

```
LIB = $(FOAM_USER_LIBBIN)/libphaseChangeTwoPhaseMixtures
```

This ensures that the Zwart model is compiled and the binaries are put in the user directory, to not interfere with the original installation. Try using wmake to see that the model compiles. Note

that it is still identical to the SS cavitation model. When the compilation is successful, the user library phaseChangeTwoPhaseMixtures will be used as default instead of the library in the original installation. You can check which library will be used by interPhaseChangeFoam.

ldd `which interPhaseChangeFoam` | grep phaseChange

It can be seen that the phaseChangeTwoPhaseMixtures library is read from the user directory.

libphaseChangeTwoPhaseMixtures.so =>/home/marcjans/OpenFOAM/marcjans-v1806/ \
platforms/linux64GccDPInt32Opt/lib/libphaseChangeTwoPhaseMixtures.so \
(0x00007fca9277d000)

When comparing eq. (1.9)-(1.10) to eq. (1.15)-(1.16), it can be seen that the differences are minor. The ZGB model only considers the vapor density while the SS model also includes the fraction between liquid- and mixture density. For small vapor fractions, however, this fraction will be close to unity. Furthermore, there is a difference in the contribution of the phase volume fractions. Since the SS model in OpenFOAM uses coefficients for evaporation and condensation, this is similar to the ZGB model. The default values are different though, where ZGB uses $C_v = 50$ and $C_c = 0.01$.

As mentioned, the ZGB model uses constant values for nucleation site volume fraction, r_{nuc} , and nucleation site radius, R_B . Start by removing the private member data for n_, dNuc_, rRb_ and alphaNu_c in Zwart.H. Add the following new member data between Cv_ and p0_.

```
//- Nucleation site volume
dimensionedScalar rNuc_;
//- Nucleation site radius
dimensionedScalar Rb_;
```

Continue with Zwart.C. Remove the constructors for n_ and dNuc_. Add in the constructor, between Cv_ and p0_:

```
rNuc_("rNuc", dimless, phaseChangeTwoPhaseMixtureCoeffs_),
Rb_("Rb", dimLength, phaseChangeTwoPhaseMixtureCoeffs_),
```

Remove the entire member functions rRb and alphaNuc. Replace the member functions pCoeff and mDotAlpha so the mass transfer rates correspond to eqs. (1.15)-(1.16). Replace mDotP according to eqs. (1.17)-(1.18).

```
Foam::tmp<Foam::volScalarField>
Foam::phaseChangeTwoPhaseMixtures::Zwart::pCoeff
(
        const volScalarField& p
) const
{
        return
        (3*rho2())*sqrt(2/(3*rho1()))
            /(Rb_*sqrt(mag(p - pSat()) + 0.01*pSat()));
}
```

```
Foam::Pair<Foam::tmp<Foam::volScalarField> >
Foam::phaseChangeTwoPhaseMixtures::Zwart::mDotAlphal() const
ſ
    const volScalarField& p = alpha1_.db().lookupObject<volScalarField>("p");
    volScalarField limitedAlpha1(min(max(alpha1_, scalar(0)), scalar(1)));
    volScalarField pCoeff(this->pCoeff(p));
    return Pair<tmp<volScalarField> >
    (
        Cc_*pCoeff*max(p - pSat(), p0_),
        Cv_*rNuc_*pCoeff*min(p - pSat(), p0_)
   );
}
Foam::Pair<Foam::tmp<Foam::volScalarField> >
Foam::phaseChangeTwoPhaseMixtures::Zwart::mDotP() const
ſ
    const volScalarField& p = alpha1_.db().lookupObject<volScalarField>("p");
    volScalarField pCoeff(this->pCoeff(p));
    volScalarField limitedAlpha1(min(max(alpha1_, scalar(0)), scalar(1)));
    return Pair<tmp<volScalarField> >
    (
        Cc_*(1.0 - limitedAlpha1)*pos(p - pSat())*pCoeff,
        (-Cv_)*rNuc_*limitedAlpha1*neg(p - pSat())*pCoeff
   );
}
```

Finally, remove n_ and dNuc_ from the read function and add rNuc_ and Rb_. Compile the class again using wmake.

```
bool Foam::phaseChangeTwoPhaseMixtures::Zwart::read()
{
    if (phaseChangeTwoPhaseMixture::read())
    {
        phaseChangeTwoPhaseMixtureCoeffs_ = optionalSubDict(type() + "Coeffs");
    phaseChangeTwoPhaseMixtureCoeffs_.lookup("Cc") >> Cc_;
        phaseChangeTwoPhaseMixtureCoeffs_.lookup("Cv") >> Cv_;
        phaseChangeTwoPhaseMixtureCoeffs_.lookup("rNuc") >> rNuc_;
        phaseChangeTwoPhaseMixtureCoeffs_.lookup("Rb") >> Rb_;
        return true;
   }
    else
    {
        return false;
    }
}
```

2.2.2 Pre- and post processing

Go to the run directory and copy the throttle tutorial. Remove the old time directories.

```
run
cp -r throttleSchnerr throttleZwart
cd throttleZwart
rm -r 0.*
rm -r postProcessing
```

In the constant/transportProperties, change phaseChangeTwoPhaseMixture from SchnerrSauer to Zwart to use the new cavitation model. Add the new model coefficients after the section with SchnerrSauerCoeffs.

```
ZwartCoeffs {

        Cc 0.01;

        Cv 50;

        rNuc 5.0e-04;

        Rb 1.0e-06;

}
```

Note the additional coefficients rNuc and Rb and that the values for Cc and Cv are changed. Solve the case, now with the new ZGB cavitation model.

interPhaseChangeFoam >& log_run&

Run the post processing.

postProcess -func singleGraph >& log_postProcess&

Plot the pressure using gnuplot. Both the pressure distribution and the vapor fraction shown in Figure 2.4 are similar to the previous results. It should be noted that the models are sensitive to their various parameters and have to be tuned for each specific case. Validation of the models and their results are not in the scope of this tutorial.

```
gnuplot
set style data linespoints
plot "postProcessing/singleGraph/0.0019/line_p_rgh.xy"
```



Figure 2.4: Last time step for throttle tutorial solved with interPhaseChangeFoam and Zwart et. al. cavitation model.

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- [4] P. J. Zwart, A. G. Gerber, and T Belamri. A two-phase flow model for predicting cavitation dynamics. In *Proceedings of the International Conference on Multiphase Flow (ICMF 04)*, Yokohama, Japan, 2004.
- [5] Abolfazl Asnaghi. Developing computational methods for detailed assessment of cavitation on marine propellers. PhD thesis, Department of Shipping and Marine Technology, Chalmers University of Technology, 2015.

Study questions

- 1. How do you check which libraries are used by a specific solver, e.g. interPhaseChangeFoam?
- 2. What are the main differences between the Schnerr-Sauer model and the Zwart-Gerber-Belamri model?
- 3. What is the role of α_{nuc} ?
- 4. Why do you need to change the phase fraction boundary conditions for the throttle, cf. cavitatingBullet? (disregard the different patch names)