STABILITY OF COUPLING METHODS FOR CONJUGATE HEAT TRANSFER

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We suggest a novel approach for coupled computations of conjugate heat transfer, considering the exchange of the boundary conditions between fluid and solid Within the multi-physics environment COOLFluiD 3, developed at the von Kármán Institute for Fluid Dynamics, we included four different coupling strategies. In all methods, boundary conditions are exchanged until equal temperatures and heat fluxes at the interface from the solid to the fluid domain. The first method sets a temperature distribution to the fluid solver that predicts a heat flux distribution imposed to the solid solver. The second method sets a heat flux distribution to the fluid solver computing a temperature field for the solid. A third method imposes a temperature field to the fluid returning a Robin boundary condition to the solid using the wall heat transfer equation. Based on a stability analysis for the existing coupling procedures, we postulate a new method, imposing a heat flux distribution to the fluid solver that returns a Robin boundary condition to the solid solver. The stability of all methods only depend on the dimensionless Biot number, the ratio of conductive to convective thermal resistance. For flat plate computations, the result of each method is in good agreement with an analytical solution. We compare the novel coupling strategy with the established methods. Considering the stability, the new approach is advantageous, especially for high Biot numbers. Further, it converges faster concluding that it can improve efficiency and accuracy of conjugate heat transfer computations.

1 INTRODUCTION

Many engineering design processes require to predict temperature distributions, e.g. the life of a turbine blade reduces by half with an increased metal temperature of 30 Kelvin [1]. In case of a complex flow field, the temperature prediction is improved if the fluid and solid temperature computations are coupled. Besides the need for two different solvers, the challenge arises through the different time scales in the solid and the fluid that can vary by orders of magnitude and increase the computational cost.

2 NUMERICAL METHOD AND COUPLING PROCEDURES

The flow equations are solved by the second order accurate Pressure Stabilized Petrov Galerkin Finite Element method with a Streamline Upwind Petrov Galerkin stabilization. The time is integrated with the second order accurate Crank-Nicholson scheme. The convective terms are linearized in time as described in [?]. The solid conduction is solved with a steady state Finite Element Method. Many problems need to compute fluid and solid part simultaneously taking the influence of one domain on the other into account, which we call a conjugate problem. There are two main approaches to solve a conjugate problem. A monolithic approach with one solver for both domains and a coupled approach with different solvers for each domains. Besides the monolithic approach there are different partitioned approaches as we detailed in [2] and summarize in the next sections.

2.1 THE FLUX FORWARD TEMPERATURE BACK METHOD

The flux forward temperature back method (often referred to as the FFTB method, named from the perspective of the fluid domain) imposes a wall temperature distribution to the fluid solver. After the fluid calculation the resulting heat flux distribution is imposed to the solid conduction solver, which returns an updated temperature in turn imposed to the fluid domain. The exchange of boundary conditions is repeated until continuous temperatures and heat fluxes between both domains.

$$\underbrace{fluid}_{\longleftarrow} \underbrace{\stackrel{q_{wall}}{\longleftarrow}}_{T} \underbrace{solid}_{}$$

Figure 1: FFTB coupling.

2.2 THE TEMPERATURE FORWARD FLUX BACK METHOD

The temperature forward flux back method (often referred to as the TFFB method, named from the perspective of the fluid domain) imposes a heat flux distribution as boundary condition to the fluid solver. The resulting temperature is imposed to the solid solver, which returns a new heat flux distribution to the fluid domain.

$$\underbrace{fluid}_{\longleftarrow} \underbrace{T}_{q_{wall}} \underbrace{solid}$$

Figure 2: TFFB coupling.

2.3 THE HEAT TRANSFER COEFFICIENT FORWARD TEMPERATURE BACK METHOD

The heat transfer coefficient forward flux back method (referred to as the hFTB method, named from the perspective of the fluid domain) starts with an initial temperature distribution as boundary condition for the fluid computation. It uses Newton's law of convective heat transfer to return a Robin (mixed type) boundary condition to the solid domain:

$$\underbrace{fluid}_{T} \xrightarrow{h, T_{fluid}} \underbrace{solid}_{T}$$

Figure 3: hFTB coupling.

$$q_{wall}^{i} = h \cdot (T_{wall} - T_{fluid}). \tag{1}$$

Again, boundary conditions are exchanged until convergence.

2.4 THE HEAT TRANSFER COEFFICIENT FORWARD FLUX BACK METHOD

Based on the stability analysis of chapter 3 we propose a fourth method (referred to as the hFFB method, named from the perspective of the fluid domain), which imposes a heat flux distribution to the fluid domain returning mixed type boundary conditions to the solid solver.

$$\underbrace{fluid} \xrightarrow{q_{wall}} \underbrace{fluid}_{q_{wall}} \underbrace{solid}$$

Figure 4: hFFB coupling.

3 STABILITY ANALYSIS

All four methods have different stability and convergence properties. We derived a stability criterion for coupled simulations of conjugate heat transfer, based on the relations between temperature and heat flux at the interface of solid and fluid domain [2].

Considering a one dimensional conjugate heat transfer problem, we specify a solid temperature at one boundary of the solid domain and the temperature of the fluid. The problem is to find the temperature T_{wall} and the heat flux q_{wall} at the interface. With the heat transfer coefficient h, the thermal conductivity λ_s and the solid domain width L it is defined by:

$$q_{wall} = \frac{\lambda_s}{L} (T_s - T_{wall}) \qquad \text{on } \Omega_s,$$

$$q_{wall} = h(T_{wall} - T_{fluid}) \qquad \text{on } \Omega_f,$$
(2)

representing heat fluxes resulting from the solid domain (Ω_s) and the fluid domain (Ω_f) computations. Heat fluxes equal at the interface and we have:

$$\frac{\lambda_s}{L}(T_s - T_{wall}) = h(T_{wall} - T_{fluid}). \tag{3}$$

With the Biot number $Bi = \frac{hL}{\lambda_s}$ the previous equation becomes:

$$T_s - T_{wall} = Bi(T_{wall} - T_{fluid}). (4)$$

Thus, the wall temperature is:

$$T_{wall} = \frac{Ts + Bi \cdot T_{fluid}}{1 + Bi}.$$
 (5)

In the next sections, we continue with the stability analysis for each of the four methods.

3.1 STABILITY OF THE FFTB METHOD

In the FFTB method, a wall temperature T_{wall}^0 is imposed to the fluid domain that varys by a value α_0 from the correct wall temperature:

$$T_{wall}^0 = T_{wall} + \alpha_0. (6)$$

The heat flux solid is then:

$$q_{wall}^{0} = h(T_{wall}^{0} - T_{fluid})$$

$$= h(T_{wall} - T_{fluid}) + h \cdot \alpha_{0}$$

$$= q_{wall} + h\alpha_{0}.$$
(7)

Using the new heat flux imposed to the solid results in a new wall temperature:

$$T_{wall}^{1} = T_{s} - \frac{L}{\lambda_{s}} \cdot q_{wall}^{0}$$

$$= T_{s} - \frac{L}{\lambda_{s}} \cdot q_{wall} - \alpha^{0} \frac{hL}{\lambda_{s}}$$

$$= T_{wall} - \alpha_{0} \cdot Bi.$$
(8)

At the i-th iteration, the temperature is given by:

$$T_{wall}^{i} = T_{wall} + \alpha_0 \cdot (-Bi)^{i}. \tag{10}$$

The heat flux results in:

$$q_{wall}^{i} = q_{wall} + \alpha_0 \cdot (-Bi)^{i} \cdot h. \tag{11}$$

As we can see, convergence is only achieved, if |Bi| < 1.

3.2 STABILITY OF THE TFFB METHOD

In the TFFB method, a wall temperature T_{wall}^0 is imposed to the solid domain that varys by a value α_0 from the correct wall temperature:

$$T_{wall}^0 = T_{wall} + \alpha_0. (12)$$

The heat flux is then:

$$q_{wall}^{0} = \frac{\lambda_{s}}{L} (T_{s} - T_{wall}^{0})$$

$$= \frac{\lambda_{s}}{L} (T_{s} - T_{wall}) + \frac{\lambda_{s}}{L} \cdot \alpha_{0}$$

$$= q_{wall} + \frac{\lambda_{s}}{L} \alpha_{0}.$$
(13)

Using the new heat flux imposed to the fluid results in a new wall temperature:

$$T_{wall}^{1} = T_{fluid} + \frac{q_{wall}^{0}}{h}$$

$$= T_{fluid} + \frac{q_{wall}}{h} - \alpha^{0} \frac{\lambda_{s}}{hL}$$

$$= T_{wall} - \frac{\alpha_{0}}{Bi}.$$
(14)

At the i-th iteration, the temperature is given by:

$$T_{wall}^{i} = T_{wall} + \alpha_0 \cdot \left(-\frac{1}{Bi}\right)^{i}.$$
 (16)

The heat flux results in:

$$q_{wall}^{i} = q_{wall} + \alpha_0 \cdot \left(-\frac{1}{Bi}\right)^{i} \cdot \frac{\lambda_s}{L}.$$
 (17)

As we can see, convergence is only achieved, if |Bi| > 1.

3.3 STABILITY OF THE hFTB METHOD

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4 FLAT PLATE TEST CASE

Schematic diagram for conjugate heat transfer test case:

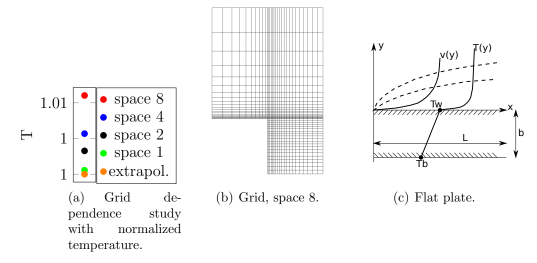
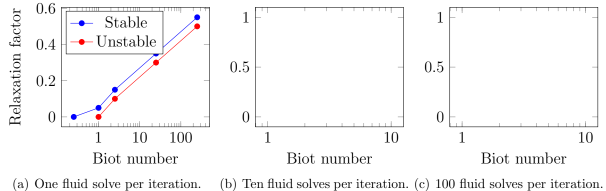


Figure 5: Grid for computational setup.

5 RESULTS AND DISCUSSION



(a) One find solve per iteration. (b) Ten find solves per iteration. (c) 100 find solves per iteration

Figure 6: Relaxation factor needed for stable computations of the FFTB method as a function of the Biot number for different fluid solves per iteration.

Relative convergence 10^{0} $\begin{array}{c} & & & \\ & 10^{-3} \\ & &$

Figure 7: Relaxation factor needed for stable computations of the FFTB method as a function of the Biot number for different fluid solves per iteration.

6 CONCLUSIONS

- found a stability criterion for cht computations
- confirm the stability criterion, dependence on Biot number by validation flat plate test case
- influence of Navier-Stokes computations per iteration
- one coupling procedure (hFTB) does not converge as the others
- flux back methods converge faster for many Navier-Stokes solves per iteration
- suggest a novel coupling procedure

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