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Wang, Yuechang; Dorgham, Abdel; Liu, Ying; Wang, Chun; Wilson, Mark C. T.; Neville, Anne; Azam, Abdullah

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An assessment of quantitative predictions of deterministic mixed lubrication solvers

Yuechang Wang, first author
Institute of Functional Surfaces, School of Mechanical Engineering, University of Leeds, Leeds LS2 9JT, UK
y.wang1@leeds.ac.uk

Abdel Dorgham, second author
Institute of Functional Surfaces, School of Mechanical Engineering, University of Leeds, Leeds LS2 9JT, UK
a.dorgham@leeds.ac.uk

Ying Liu, third author
State Key Laboratory of Tribology, Tsinghua University
State Key Laboratory of Tribology, Tsinghua University, Beijing 100084, China
liuying@mail.tsinghua.edu.cn

Chun Wang, forth author
Institute of Functional Surfaces, School of Mechanical Engineering, University of Leeds, Leeds LS2 9JT, UK
c.wang@leeds.ac.uk

Mark C.T. Wilson, fifth author
Institute of Functional Surfaces, School of Mechanical Engineering, University of Leeds, Leeds LS2 9JT, UK
m.wilson@leeds.ac.uk

Anne Neville, sixth author
Institute of Functional Surfaces, School of Mechanical Engineering, University of Leeds, Leeds LS2 9JT, UK
a.neville@leeds.ac.uk

Abdullah Azam, sixth author
Institute of Functional Surfaces, School of Mechanical Engineering, University of Leeds, Leeds LS2 9JT, UK
a.azam@leeds.ac.uk

ABSTRACT

1 Corresponding author. Tel: +86 10 6278 2336; Fax: +86 10 6278 2336.
The ability to simulate mixed lubrication problems has greatly improved, especially in concentrated lubricated contacts. A mixed lubrication simulation method was developed by utilizing the semi-system approach which has been proven to be highly useful for improving stability and robustness of mixed lubrication simulations. Then different variants of the model were developed by varying the discretization schemes used to treat the Couette flow terms in the Reynolds equation, varying the evaluation of density derivatives and varying the contribution of terms in the coefficient matrix. The resulting pressure distribution, film thickness distribution, lambda ratio, contact ratio, and the computation time were compared and found to be strongly influenced by the choice of solution scheme. This indicates that the output from mixed lubrication solvers can be readily used for qualitative and parametric studies, but care should be taken when making quantitative predictions.

Keywords: Mixed lubrication; Deterministic model; Numerical simulation

1. Introduction

Contacting surfaces in mechanical components are important for transmitting power and motion. To reduce the power loss and improve the efficiency and life of mechanical components, the contacting surfaces are usually lubricated [1]. In engineering practice, rough surfaces are universal and they affect the thickness of lubricant film within the contact. When the film thickness is not large enough to separate the two surfaces, solid to solid contact starts to occur and ultimately, the effect of roughness becomes dominant. Under such conditions, the lubrication state is referred to as the mixed lubrication regime where fluid lubrication and solid to solid contact occur simultaneously. Mixed lubrication is inevitable in engineering applications. Therefore, modelling and predicting the performance of the mixed lubricated contacts is one of the key challenges and popular topics in lubrication science, and great efforts have been made in this field.
Generally, there are two types of mixed lubrication models: stochastic (roughness defined by stochastic parameters) and deterministic (roughness defined at each grid node). One milestone in the stochastic mixed lubrication model development was made by Patir and Cheng [2]. Later, several improvements to the stochastic model were introduced [3-5]. Stochastic models use mean values, which results in loss of localized information, such as detailed pressure and film thickness variations which are important for studies on lubrication breakdown and failures. Therefore, stochastic models will not be discussed further in this study.

In the last two decades, more studies have focused on deterministic modelling approaches. The very first deterministic models considered simple geometry and roughness models [6-8] but later studies involved complex three-dimensional surface topographies [9-11]. The numerical solution methods for deterministic models have also improved overtime [12-15].

To solve the Reynolds equation, it is discretized to form a set of linear algebraic equations. The coefficient matrix of the discretized Reynolds equation is generally constructed from the Poiseuille flow terms alone. Deterministically solving a mixed lubrication model involves dealing with very thin lubricant films. Such thin films weaken the validity of Poiseuille flow and make the coefficient matrix lose its diagonal dominance, which results in loss of efficiency and accuracy of the linear algebraic solvers. A solution to this problem was proposed by Ai et al. [9] in the form of semi-system approach. The basic concept of the semi-system method is that both Poiseuille flow terms and Couette flow terms are used to construct the coefficient matrix, ensuring diagonal dominance even under severe contact conditions. Based on the semi-system method, Hu and Zhu [13]
proposed a mixed lubrication model for point contact elastohydrodynamic lubrication (EHL) problems which could handle the complete lubrication behavior from boundary to full film lubrication regimes. According to this model, when the film thickness was small enough, localized contact occurred. Parametric [16] and experimental [17] studies have proved the capability of this model.

The Reynolds equation is solved by first discretizing (several different discretization schemes available) and then solving the resulting linear algebra problem. Until now, it is presumed that all the different solution methods and discretization schemes produce identical results under similar working conditions. In the past, Liu et al. [14] studied the effects of differential scheme and mesh density on the point contact EHL film thickness predictions and recommended that for ultrathin films (below 10–20 nm), dense meshes should be used. They also suggested that the first-order backward schemes are better when dealing with these ultrathin films. Further, Zhu et al. [15] pointed out that the accurate calculation of roughness derivatives is critical for ensuring numerical accuracy. But no study addresses the issue of repeatability of predictions from a mixed lubrication (EHL) solver due to the different implementations of the mathematical model.

Therefore, the current study aims at developing this understanding by utilizing a series of key implementations. Various implementation cases are selected by considering, the information included into the coefficient matrix (main diagonal or tridiagonal), the different ways of dealing with the density derivative (numerical differentiation or differentiating its empirical equation) and finally linking these to the separate and combined implementations of the Couette terms. The results from these implementations are presented as pressure distribution, film thickness distribution, lambda ratio, and the
contact ratio. A comparison among the individual cases was made for the waviness surfaces as well as the numerically generated rough surfaces. To ensure the validity of the simulations, the results for the waviness surface are first compared against the work of Venner et al. [18]. The findings from the current study have direct implications on the use of semi-system method in lubrication simulations.

This paper is organized by first presenting the equations representing the mixed lubrication regime. Both dimensional and non-dimensional equation sets are given. Then the different discretization cases are outlined, and the derivation of the density term is discussed. Finally, six representative cases are chosen to analyze the consistency of the solver and the results are presented for all these cases. The predictions from the current study are compared against Venner’s work [18] to establish the validity of the current model. Finally, results are presented for the rough surface simulations and discussion is made on the use of the findings from the lubrication solvers for quantitative predictions.

2. Details of the mixed lubrication model

A typical set of basic equations that formulate a rough surface point contact EHL problem are given in this section [13]. The definitions of all the symbols used below are given in the Nomenclature.

The steady state Reynolds equation is used to focus our attention on the discretization schemes and not the transient effects:

\[
\frac{\partial}{\partial x} \left( \frac{\rho h^3}{12\eta} \frac{\partial p}{\partial x} \right) + \frac{\partial}{\partial y} \left( \frac{\rho h^3}{12\eta} \frac{\partial p}{\partial y} \right) = \frac{u_1 + u_2}{2} \frac{\partial (\rho h)}{\partial x}
\]

(1)

The x and y represent the coordinate axes, p is the pressure distribution, h is the film thickness distribution, \(\rho\) is the density, \(\eta\) is the viscosity, and \(u_1\) and \(u_2\) are the velocity of ball and disc surfaces. The terms on the L.H.S. of equation 1 represent the Poiseuille flow
terms while the terms on the R.H.S. represent the Couette flow terms.

**Film thickness equation:**

\[ h(x, y) = h_0 + \frac{x^2}{2R_x} + \frac{y^2}{2R_y} + \delta_1(x, y) + \delta_2(x, y) + v(x, y) \]  \hspace{1cm} (2)

In Eq. (2), the first term on the R.H.S. gives the minimum undeformed gap, \( h_0 \), the second and third terms represent the paraboloid Hertzian macro-geometry where \( R_x, R_y \) are radii of curvature in x and y direction, the fourth and fifth terms give the roughness (micro-geometry) of the contacting bodies (\( \delta_1, \delta_2 \)) while the last term is the deformation, \( v \).

**Elastic deformation equation** [19]:

\[ v(x, y) = \frac{2}{\pi E_e} \iint_{\Omega} \frac{p(\xi, \zeta)}{\sqrt{(x-\xi)^2 + (y-\zeta)^2}} d\xi d\zeta \]  \hspace{1cm} (3)

where \( E_e \) is the equivalent Young’s modulus.

**Load balance equation:**

\[ w = \iint_{\Omega} p(x, y) \, dx \, dy \]  \hspace{1cm} (4)

where \( w \) is the applied load.

These four equations (Eqs. (1-4)) together form the complete system of equations describing the point contact EHL problems. Due to the fact that since extremely high-pressure value is observed in the contact zone, the viscosity and density of the lubricant are also function of pressures. The Barus viscosity pressure equation [20] is used to represent the changes in viscosity and the Dowson-Higginson density pressure equation [21] represents changes in density in this study and are given below.

**Barus viscosity pressure equation:**

\[ \eta = \eta_0 \exp(\alpha p) \]  \hspace{1cm} (5)

where \( \alpha \) is the pressure-viscosity coefficient, and \( \eta_0 \) is the ambient viscosity.

**Density pressure equation:**
\[
\frac{\rho}{\rho_0} = 1 + \frac{0.6 \, p}{1 + 1.7 \, p}
\]  
(6)

where \( p \) is the pressure in GPa, and \( \rho_0 \) is the ambient density.

The detailed information about the non-dimensionalization for these equations can be found in Appendix A. The resulting non-dimensional Reynolds equation is,

\[
\frac{\partial}{\partial X} \left( \varepsilon_x \frac{\partial p}{\partial X} \right) + \frac{\partial}{\partial Y} \left( \varepsilon_y \frac{\partial p}{\partial Y} \right) = \frac{\partial \left( \tilde{\rho} H \right)}{\partial X} 
\]

(7)

It is important to mention that the Poiseuille flow terms are discretized by the second order central difference method and the discretization is given in Eq. (8), assuming that the discrete intervals are the same in \( x \) and \( y \) directions i.e. \((\Delta X = \Delta Y)\).

\[
\frac{\partial}{\partial X} \left( \varepsilon_x \frac{\partial p}{\partial X} \right) + \frac{\partial}{\partial Y} \left( \varepsilon_y \frac{\partial p}{\partial Y} \right) \approx \frac{1}{\Delta X^2} \left[ \varepsilon_{x_{i-1/2,j}} P_{i-1,j} - \left( \varepsilon_{x_{i-1/2,j}} + \varepsilon_{x_{i+1/2,j}} \right) P_{i,j} \right] + \frac{1}{\Delta X^2} \left[ \varepsilon_{y_{i-1/2,j}} P_{i,j-1} - \left( \varepsilon_{y_{i-1/2,j}} + \varepsilon_{y_{i+1/2,j}} \right) P_{i,j} \right] 
\]

(8)

For the Couette flow terms, the first order backward difference method is used, as suggested by Liu et al. [14]. But it should be pointed out that the Couette flow terms, \( \frac{\partial (\tilde{\rho} H)}{\partial X} \), can be expressed in two different ways: the combined form and the separate form. The combined form is given as,

\[
\frac{\partial (\tilde{\rho} H)}{\partial X} \approx \frac{(\tilde{\rho} H)_{i,j} - (\tilde{\rho} H)_{i-1,j}}{\Delta X} 
\]

(9)

This form is expected to give reasonable results for conditions where the effects of roughness is limited. When dealing with extreme cases or when the roughness dominates the contact behavior, the separate form generally gives better numerical accuracy and efficiency, and is given as,

\[
\frac{\partial (\tilde{\rho} H)}{\partial X} = \frac{\partial \tilde{\rho}}{\partial X} \tilde{H} + \frac{\partial H}{\partial X} \tilde{\rho} 
\]

(10)
Two further cases result due to the density term, $\frac{\partial \rho}{\partial X}$. Due to the fact that the density variation as a function of pressure is given by the formulation, Eq. (6) which is an empirical equation, the derivative of density can be estimated by either taking the derivative of this empirical equation (6) or using numerical differentiation. The latter method is given in Eq. (11).

$$\frac{\partial \rho}{\partial X} H + \frac{\partial H}{\partial X} \rho \approx \frac{\rho_{i,j} - \rho_{i-1,j}}{\Delta X} H_{i,j} + \frac{H_{i,j} - H_{i-1,j}}{\Delta X} \rho_{i,j} \tag{11}$$

Instead of calculating the derivatives of density by direct finite difference, Ai et al. [9] used the chain derivation rule, given in Eq. (12).

$$\frac{\partial \rho}{\partial X} H + \frac{\partial H}{\partial X} \rho - \frac{\partial (P)}{\partial P} \frac{\partial P}{\partial X} H + \frac{\partial H}{\partial X} \rho \approx \phi_{i,j} \frac{P_{i,j} - P_{i-1,j}}{\Delta X} + \rho_{i,j} \frac{H_{i,j} - H_{i-1,j}}{\Delta X} \tag{12}$$

where,

$$\phi_{i,j} = \frac{C_a H_{i,j}}{(1 + C_b P_{i,j})^2} \tag{12}$$

where $C_a$ and $C_b$ are the non-dimensional coefficient in Eq. (6) and their values are 0.6$p_h$ and 1.7$p_h$ respectively.

Once the Poiseuille and the Couette flow terms have been discretized, a linear equation set with unknown pressures, $P$ is obtained. The discrete Reynolds equation can be represented by only taking $P_{i,j}$, $P_{i+1,j}$, and $P_{i-1,j}$ as the unknowns and all terms related to $P_{i,j-1}$ and $P_{i,j+1}$ are moved to the R.H.S, which makes the coefficient of the linear equation set a tridiagonal matrix and can easily be solved. The form of the linear equation set is shown below.

$$\alpha_{i,j}^{(s)} P_{i-1,j}^{(s+1)} + \beta_{i,j}^{(s)} P_{i,j}^{(s+1)} + \gamma_{i,j}^{(s)} P_{i+1,j}^{(s+1)} = b_{i,j}^{(s)} \tag{13}$$

Where $\alpha_{i,j}^{(s)}$, $\beta_{i,j}^{(s)}$, $\gamma_{i,j}^{(s)}$, and $b_{i,j}^{(s)}$ are the coefficients, (s) denotes the iteration index.

It should be noted that the terms involving $P_{i,j-1}$ and $P_{i,j+1}$ are moved to the R.H.S with the
term $b_{i,j}$. The expressions of the coefficients are listed in Appendix B. An iterative procedure is used to solve Eq. (13). Typically, the coefficient matrix in Eq. (13) is only constructed from the Poiseuille flow terms alone and this significantly reduces the numerical stability, especially in the case of ultrathin lubricant films. The main diagonal terms in the coefficient matrix become very small, and it losses its diagonal dominance and convergence becomes difficult. To overcome this reduced stability, the semi-system method constructs the coefficient matrix in Eq. (13) from both the Poiseuille and Couette flow terms. To consider the Couette flow terms, the calculation procedures of the elastic deformation $v$, are taken into account by representing it as a function of the unknown pressures.

The non-dimensional elastic deformation $V$ is determined by the pressure distribution. The discrete equation used to compute the elastic deformation is

$$V_{i,j} = \sum_{k=1}^{M-1} \sum_{l=1}^{N-1} D_{k,i}^{i,j} P_{k,l}$$  \hspace{1cm} (14)

In this equation, $D$ is called the influence matrix and is calculated based on Eq. (3). $M$ and $N$ are the number of discretization points in the x and y directions, respectively. The detailed procedure for obtaining the influence coefficients, $D$ can be found in work [22]. Eq. (14) can be expressed as a function of unknown pressures, $P_{ij}$, $P_{i+1,j}$, and $P_{i-1,j}$, and the deformation is also expressed in the form of multiple diagonal terms which can then be added to Eq. (13) to implement the semi-system method. Whether the influence coefficient has been written in terms of the pressure, $P_{ij}$ alone or in terms of the three pressures, $P_{ij}$, $P_{i+1,j}$, and $P_{i-1,j}$, two different formulations result: one with main diagonal terms only and the other with main as well as secondary diagonals, respectively. Huang et al. [23] only extracted the main diagonal terms, but Ai et al. [9] also extracted the secondary diagonal
terms.

Based on the above discussion, different possibilities for formulating the system are possible by combining the single and multiple diagonal formulations, the separate and combined approaches and the different density derivatives. A total of six choices result for implementing the semi-system method and are summarized in Table 1. The numbers ‘0’ and ‘1’ in the table represent the different combinations. ‘1’ means that the method described in the header of the particular column was used and ‘0’ means that the method mentioned in the header of that particular column was not used. All the detailed expressions and formulae for these different implementations are provided in Appendix B.

3. Contact configuration and numerical simulation details

Two sets of simulation cases were considered. The first set of simulations were performed for the waviness surface and the second set of simulation cases used computer generated rough surfaces.

The input parameters of the EHL problem and waviness surface were extracted from the work of Venner et al. [18]. The equation for generating waviness surface is,

\[ R_w(X, Y) = A_x \cos \left(2\pi \frac{X - X_s}{\Omega} \right) \]  

Where \( R_w \) is the waviness surface data which is used as roughness data in Eq. (2), \( A_x \) is the non-dimensional waviness amplitude, \( X_s \) is the location of waviness start in x direction, \( \Omega \) is the non-dimensional wavelength. The waviness amplitude is 0.08 \( \mu \)m and wavelength is 59 \( \mu \)m [18]. In this paper, the starting location of waviness is at the start location of the calculation domain. The calculation domain is \( X_s \leq X \leq X_e \) and \( Y_s \leq Y \leq Y_e \), where \( X_s = -2.5 \), \( X_e = 1.5 \), \( Y_s = -2 \), \( Y_e = 2 \). The discrete grid density is \( 1024 \times 1024 \), unless otherwise stated. This grid density has been proved to be dense enough in literature [14,15].
The parameters of the EHL problem are listed in Table 2. The values of dimensionless speed parameter $U$ selected in this paper varies from $4 \times 10^{-16}$ to $4 \times 10^{-11}$ where the results for $U = 4 \times 10^{-11}$ correspond to the simulations results of Venner et al. [18] and are used to validate the implementation methods and the computer codes used in this paper.

Besides the waviness surface, numerically generated Gaussian isotropic rough surfaces were used in this work. The rough surfaces were generated based upon the method from Wu [24], and the generated rough surface data, as used in this study, is also provided as supplementary data with this paper. The attached data is a random number matrix $(1024 \times 1024)$ following Gaussian distribution with a unit root-mean-square variance (Rq). Rough surfaces with any other Rq values can be obtained by multiplying the data with that specific Rq value. In the current study, the Rq used is the same as the amplitude of the waviness surface.

The iterative pressure calculation procedure stops when the convergence criteria have been met. The convergence criteria for pressure distribution and load balance calculation are defined based on Eq. (16) and Eq. (17), respectively.

$$\sum \sum |P^{(s+1)} - P^{(s)}| \leq \epsilon_p$$  \hspace{1cm} (16)

$$\sum \sum |W^{(s+1)} - W| \leq \epsilon_w$$  \hspace{1cm} (17)

$W$ in Eq. (17) is the non-dimensional applied load and $W^{(s+1)}$ is the calculated load capacity at the newest iteration step. The terms $\epsilon_p$ and $\epsilon_w$ are the threshold values which for both pressure distribution and load balance calculation were set as $1 \times 10^{-6}$. Moreover, to improve the numerical calculation efficiency, as suggested by Liu et al. [14], when the
dimensionless film thickness values were less than $1 \times 10^{-6}$, solid contact was assumed to occur. It should be noted that the criterion used here for establishing a solid contact still lacks firm theoretical basis. Zhu [1] discussed this issue and tried to connect this threshold value to the molecular length of lubricant. To ensure that the selection of this criteria has no effect on the outcomes presented in the current study, the value of minimum film thickness for establishing solid contact was kept the same for the different implementations of numerical models presented in this study. Another important aspect about the asperity contact is that the contact behavior is resolution dependent [25]. Thus, the input roughness data and the size of solution domain are kept fixed for comparing corresponding simulations.

With ultrathin lubricant films, it is hard to plot fine film thickness distribution in the Hertzian contact zone by common plotting methods as the different regions with different film thickness values cannot be identified due to the very small overall film thickness values involved. Venner et al. [18] used pseudo interference graphs to tackle this problem. The dimensionless film thickness data was represented by the corresponding intensity values based on Eq. (18).

$$I_H(X, Y) = 0.5 + 0.5 \cos \left( \frac{2\pi H(X, Y)}{\Lambda} \right)$$

(18)

where $\Lambda$ is non-dimensional wavelength for which the default value used in this study is 0.05.

For the simulation results with rough surface, the lambda ratio ($\lambda$) and contact ratio ($A_c$) are used to identify different contact configurations. In this work, the lambda ratio is not defined as the minimum film thickness value divided by the composite surface roughness as the minimum film thickness loses generality when rough surface is used.
Instead, the average film thickness value in the Hertzian contact zone ($h_{ac}$) is used. The corresponding equation is

$$ \lambda = \frac{H_{ac}}{A} $$

(19)

The contact ratio is defined as the number of contact grid points divided by the total number of grid points in the Hertzian contact zone as shown in Eq. (20).

$$ A = \frac{N_c}{N_{Hertz}} $$

(20)

where $N_c$ is the number of contact points in the Hertzian contact zone, and $N_{Hertz}$ is the total number of grid points of the Hertzian contact zone.

Relative computation time is calculated for different implementing methods with the same EHL simulation parameters. The relative computation time is defined as the ratio of computation time of different methods and the computation time when (0,0,0) method is used. In addition, the actual computation time (in seconds) is also provided for the (0,0,0) implementation. It should be noted that both the actual and relative computation time are not comparable with different speed parameter values, because some iteration coefficients were modified to get converged results within acceptable time when the speed parameter was changed. All the simulations were conducted with MATLAB 9.7 on a laptop with a 2.5-GHz Intel Core I7 processor.

4. Simulation results and discussion

4.1 Waviness surface (validation and discussion of results)

In Fig. 1, the pressure and film thickness profiles are presented at $Y = 0$, from the present work for all six implementation cases. All these schemes produce similar results with slight differences in the pressure peak height towards the exit of the contact region. In Fig. 2, the (0,0,0) implementation case is considered and the pressure and film thickness
profiles are plotted. Three different grid sizes i.e. 256×256, 512×512, and 1024×1024 were used to ensure that the differences were not related to the grid. The pressure and film thickness results are also extracted from reference [18] and plotted in Fig. 2 for comparison. It can be readily seen that the qualitative and quantitative match is very good but, once again, slight differences exist in the prediction of pressure peak height towards the exit of the contacting region.

To further investigate the effect of the numerical implementations, the contours of corresponding pressure and film thickness distributions are shown in Fig. 3 and Fig. 4, respectively. It is evident from these contour plots that the converged results from all the six implementations are the same. It is important to note that in Fig. 4, the method proposed by Venner (given in Eq. (16) [18]) is used to plot film thickness contours.

Next, the speed parameter in the simulations was reduced from $U = 4 \times 10^{-11}$ to lower values of $U = 4 \times 10^{-12}$, $4 \times 10^{-13}$, and $4 \times 10^{-14}$ successively. Fig. 5 shows the corresponding pressure and film thickness profiles at $Y = 0$ for these different speeds. It shows that as the speed decreases, the discrepancies between the different implementing methods increases. In order to show this effect more comprehensively, pressure and film thickness distributions in the Hertzian contact zone with $U = 4 \times 10^{-14}$ are plotted in Fig. 6 and Fig. 7 respectively. These plots correspond to the profiles in Fig. 5. No big differences in the pressure distribution results with different implementing methods can be observed. However, the implementation method used has dramatic influence on the film thickness distribution.

In the following, the results from these different implementations are analyzed to access the key reasons for the discrepancies among the output. To facilitate discussion, the
different implementations have been labeled as (a) to (f) in Fig. 7. A close look at the film thickness distribution in (a) and (b) indicates that the different derivation methods adopted for evaluation of the density derivative does not affect the output of the solver. Also, looking at (a) and (e), (c) and (f), respectively, it can be clearly seen that whether the separate or combined form of derivation is used, the output of the solvers is unaffected. But when comparing (a), (b), (c) and (e), it can be readily seen that the output is very different for the case of (c) than for (a), (b) and (e). A look at the implementation schemes suggests that the key difference is the inclusion or exclusion of the effect of Couette flow in the secondary diagonal terms. The results in (c) are significantly different from the other implementations because the effect of Couette terms is only included in the main diagonal. Therefore, for the waviness surfaces, the output of the solver is not reliable when the effect of Couette terms is only included in the main diagonal terms.

An interesting phenomenon can be observed by looking at the output (c) and (d). These two methods only differ in the definition of the density derivative (differentiating the empirical equation or using finite differences) but the output from both the cases is different. Also, it seems that the output from (c) is not correct. Therefore, it can be inferred that the solution method can be stabilized if the density derivative is defined through the chain rule (see Eq. (12)). This stability can be linked to the additional pressure derivative that appears in the derivation of density.

The next task is to compare the time involved in each of these implementation methods. Table 3 lists the relative simulation time and the actual computation time for the (0,0,0) method. With $U = 4 \times 10^{-11}$, using (0,0,0) method results in the fastest convergence, and the relative computation time values obtained by implementing other five methods are
almost the same. Similarly, when \( U = 4 \times 10^{-12} \) was used, all the six implementation methods result in nearly the same computation time. With lower speed parameter values (\( U = 4 \times 10^{-12} \) and \( U = 4 \times 10^{-13} \)), the computation time for using (0,0,0) method is the shortest. One interesting phenomenon here which does not occur with higher speed is that all the implementation methods that only consider the effect of Couette flow on the primary diagonal, indexed as (0,1,0), (0,1,1), and (1,1,0) result in near thirty times higher computation effort. The other two factors; combined or separate form of Couette flow term and the different density derivative implementation methods, have far little effect on the computation time. This indicates that for the waviness surfaces with lower speed parameter, even including the effect of Couette flow in the primary diagonal alone could dramatically reduce the computation efficiency. These results also indicate that overall using (0,0,0) method results in the highest computational efficiency (shortest computation time).

With the relative computation time required and the accuracy for each of these cases in mind, it is believed that, for the waviness surfaces, the (0,0,0) implementation that considers separate form, effect of Couette flow included in the secondary diagonals as well and the density derivative implemented through finite differencing, is the most suitable for computations and is recommended.

4.2 Numerically generated isotropic Gaussian surface

Next, a critical assessment is made on the effect of these implementations on the output of mixed lubrication solvers when numerically generated rough surfaces are used. The 3-dimensional visualization and roughness profiles along \( X = 0 \) and \( Y = 0 \) of the generated surface is shown in Fig. 8. In the first set of simulations, the speed parameter is fixed as \( U \)
= 4×10^{-13}. This speed value is selected to ensure that it is a full film case for all the implementations. The simulations with lower speed where asperity contact is impossible to avoid, will be presented later in this section.

The pressure and film thickness profiles at Y = 0 are presented in Fig. 9 for all the different implementation methods. Contrary to the waviness surface case, there are obvious differences among the different profiles corresponding to the each of the methods used. The contours of pressure and film thickness within the Hertzian contact zone are further shown in Fig. 10 and Fig. 11. All these distributions were normalized to the interval between 0 and 1 to generate these graphs with the same color scale. The corresponding lambda ratio and relative computation time are also compared and are listed in Table 4. The actual computation time of using (0,0,0) method is also provided.

The film thickness distribution plots in Fig. 11 are much clearer than the pressure distribution plots in Fig. 10. As the differences can be seen more clearly in Fig. 11, only the differences in the film thickness predictions are discussed. Once again, to facilitate discussion, the different implementations in Fig. 11 have been labelled as (a) to (f). First, it is important to note that the output of the solvers for the rough surface simulations is quite different from the output by the respective solvers for the case of waviness surfaces. Comparing in pairs, (a) and (e), like (c) and (f), suggest that for rough surface simulations, the use of combined or separate form of the Couette terms does affect the output of the solver. Moreover, comparing the pair, (a) and (c), the pair (b) and (d) and the pair (e) and (f), it can be seen that, unlike the waviness surface results, whether the effect of Couette terms is included in the main diagonal alone or the secondary diagonal as well, the output from the solvers is not affected. To access the effect of density derivative, a comparison of
the pair (a) and (b) and the pair (c) and (d), suggests that the implementation of the density derivative also affects the output of rough surface simulations. These results provide further evidence that the quantitative predictions from the mixed lubrication solvers should be treated cautiously.

To further provide evidence to support our argument, the lambda ratio values are listed in Table 4. These values also show that the output is solver or implementation dependent. Corresponding to the film thickness distribution plots in Fig. 11, there are totally three groups of lambda ratio values, (a):0.77 and (c):0.77, (b):1.76 and (d):1.76, (e):0.26 and (f):0.27, in which the group (a) and (c) has moderate lambda ratio and is used as the benchmark for comparison. Comparing group (e), (f) and group (a), (c), it can be seen that the use of the combined form of Couette terms dramatically reduces the film thickness which results in a much smaller lambda ratio. Moreover, comparing group (b), (d) and group (a), (c), it can be concluded that if the chain derivation rule for calculating the derivatives of density is used, the film thickness is significantly enhanced. As the differences among these three groups of lambda ratios are rather big, it is difficult to recommend which kind of implementation method is best.

A comparison of the computation time taken to reach convergence by each of these implementations suggests that the method (0,0,0) is still the most efficient. Furthermore, (0,1,0), (0,1,1), and (1,1,0) cases still require longer computation time than other cases. The trend is similar to that observed in the case of waviness surfaces at low speeds. This indicates that for the generated rough surfaces without asperity contact, only including the effect of Couette flow in the primary diagonal could reduce the computation efficiency.
In order to test the different implementing methods with asperity contact, an ultralow speed parameter value of \( U = 4 \times 10^{-16} \) is imposed, and the simulations are repeated. Fig. 12 shows the pressure and film thickness profiles at \( Y = 0 \) for all the different implementations. The contours of contact area are plotted in Fig. 13 instead of contours for pressure and film thickness. This is to ensure that the effect of asperity contact is actually compared. Table 5 lists the lambda ratio, contact ratio, and relative computation time values with the actual computation time of using the \((0,0,0)\) implementation. From Fig. 12, the discrepancies in pressure and film thickness profiles is less compared to the case of no asperity contact for the different implementations. The different implementations have been labelled again to facilitate discussion. Based on the contact area graphs (Fig. 13), the pair (a) and (e) and the pair (c) and (f), show, similar to the case of no asperity contact that the use of combined or separate form of Couette terms in the solvers does change the output. Next, looking at the pair, (a) and (c), the pair, (b) and (d) and the pair (e) and (f) suggests that the inclusion of the effect of Couette flow in the main diagonal alone or the main as well as secondary diagonals, does not affect the output of the solver. Finally, the pair (a) and (b) and the pair (c) and (d) suggest that the different formulations of the density derivative give different results, but the effect is less pronounced compared to the case of no asperity contact.

According to Table 5, the lambda ratio and contact ratio values could be divided into three groups as well, which is similar to the lambda ratio values in Table 4. Interestingly, as the effect of different formulations of the density derivative terms is less obvious compared to when no asperity contact occurs. This can be clearly seen when comparing case (a), (b), (c), and (d) as these result in nearly the same lambda ratio and contact ratio values. Moreover, it could be inferred that these outputs have convergent trend as the
severity of contact increases. Then, the question comes to how to choose between the combined form and the separated form of the Couette term. Although some researchers preferred the separated form [9,10,13,15], there is still no solid evident that which one is the most accurate and should be recommended.

Next, the computation time in Table 5 is analyzed. Unlike the two cases presented above (waviness surface and the rough surface without asperity contact), the (0,0,1) implementation is the most efficient (shortest computation time). Interestingly, the computation time for the (0,0,0) implementation (shortest for the waviness and no asperity contact case) is even longer than the (0,1,0) and the (0,1,1) cases. Another important thing to notice is that the differences in relative time among all the different implementations is not as great as the case of waviness surface or the case of rough surface with no asperity contact.

In summary, the implementation method for the semi-system method has dramatic influence on the simulation output. A key point to stress is that the effect varies from case to case and is highly dependent upon the problem being simulated. For the waviness surface, like the rough surface, as the lubricant film gets thinner, the discrepancies become greater. However, when using numerically generated isotropic Gaussian rough surfaces, unlike the waviness surfaces, modifying the diagonal or secondary diagonal terms does not affect the output. In this case, other parameters like, the combined or separated form of the Couette terms and the chain derivation rule for the derivatives of density are the key factors affecting the output. In case of severe asperity contact, when the speed reduces, only the combined or separated derivatives of the Couette term matters and the effect of the different formulations for the density derivative is negligible.
A comparison of computational times further complicates the process of selection of the adequate implementation method as, once again, it is dependent upon the problem at hand. For the simulation cases presented above, as long as the waviness surfaces or rough surfaces (without asperity contact) are considered, the (0,0,0) implementation is computationally the most efficient but when rough surfaces with asperity contact are considered, the (0,0,1) implementation gives the least computation time.

Finally, combining the effect of different implementation methods on the accuracy and efficiency of simulation output, the (0,0,0) implementation method is recommended for waviness surfaces. However, it is hard to say which implementation method should be recommended when generated rough surfaces are used.

5. Conclusions

In this paper, mixed lubrication solvers were developed by considering the combined or separate form of the Couette flow terms, including the effect of the Couette flow terms in the main diagonal alone or the secondary diagonal as well and formulating the density derivative by either using finite differencing or using chain rule. These different implementations were tested for waviness surfaces and numerically generated rough surfaces. The output from the solver was found to be highly influenced by the choice of the implementation. The main conclusions can be summarized by the following points:

1) For the waviness surfaces, the inclusion of the effect of Couette terms in the diagonal terms alone gives different results compared to when their effect is included in the secondary diagonals as well. And combining the accuracy and efficiency of the solver, the (0,0,0) implementation method is recommended.

2) For the rough surfaces, if the asperity contact is not considered (full film cases)
the formulation of the density derivative is the key factor affecting the output while the inclusion of Couette terms in the main diagonal alone or secondary diagonals could be neglected.

3) For the rough surfaces, if the asperity contact is considered (mixed contact) the solver output was found to be only slightly affected by the choice of density derivative formulation but the combined or separate formulations of the Couette terms influences the solver output most.

4) Irrespective of the accuracy of the output, for the rough surface simulations (full film) the computational efficiency was found to be the best for the implementation, (0,0,0) which involves separate form of Couette flow terms, includes the effect of Couette flow terms in the main as well as secondary diagonals and not using the chain rule for density formulation. For the rough surface (mixed contact) the computationally best implementation, (0,0,1), had density derivative formulated using chain rule.

Therefore, when using EHL and mixed lubrication solvers care should be taken when using their output for quantitative predictions. And more attention should be paid to how to get enough accurate output through deterministic mixed lubrication solvers.

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### NOMENCLATURE

| Symbol | Description |
|--------|-------------|
| $A_x$  | non-dimensional waviness amplitude |
| $b$    | radius of Hertzian contact |
| $C_a$, $C_b$ | dimensionless coefficients for pressure-density equation |
| $D_{i,j}^{k,l}$ | influence coefficient used in elastic deformation computation |
| $E_e$  | equivalent Young’s modulus |
| $h$, $H$ | dimensional and non-dimensional film thickness |
| $h_0$  | reference value in film thickness equation |
| $N_c$, $N_{Hertz}$ | number of contact points and total number of points of the Hertzian contact zone |
| $p$, $P$ | dimensional and non-dimensional pressure |
| $p_h$  | maximum Hertzian contact pressure |
| $R_x$, $R_y$ | radii of curvature in x and y directions |
| $R_w$  | waviness surface |
| $R_q$  | root-mean-square variance |
| $u_1$, $u_2$ | velocity of ball and disc surfaces |
| $u_s$  | entrainment speed |
| $U$    | dimensionless speed parameter $U = \eta_0 u_s / (2E_s R_s)$ |
| $v$, $V$ | dimensional and non-dimensional elastic deformation |
w \quad \text{dimensional load}

x, X \quad \text{dimensional and non-dimensional coordinates in rolling direction}

X_s \quad \text{start location of the computation domain and waviness in x direction}

X_e \quad \text{end location of the computation domain in x direction}

y, Y \quad \text{dimensional and non-dimensional coordinates in the perpendicular direction to the rolling direction}

Y_s, Y_e \quad \text{start and end location of the computation domain in y direction}

\alpha \quad \text{pressure-viscosity coefficient}

\alpha_{i,j}^{(s)} \beta_{i,j}^{(s)} \gamma_{i,j}^{(s)} b_{i,j}^{(s)} \quad \text{coefficients and right side in the discrete Reynolds equation}

\delta_1, \delta_2 \quad \text{rough surfaces}

\Delta X, \Delta Y \quad \text{dimensionless mesh spacing in x and y direction}

\varepsilon_x, \varepsilon_y \quad \text{dimensionless coefficients for Poiseuille flow}

\eta, \eta_0 \quad \text{viscosity and ambient viscosity}

\lambda \quad \text{ratio of average film thickness in the Hertzian contact zone and the amplitude or Rq of surfaces}

\rho, \rho_0 \quad \text{density and ambient density}

\bar{\rho} \quad \text{dimensionless density}

\phi_{i,j} \quad \text{dimensionless product of derivation of density and film thickness}

\Omega \quad \text{non-dimensional wavelength}
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**Figure Captions List**

Fig. 1  Pressure and film thickness profiles at Y = 0 with six implementation methods

Fig. 2  Comparison of the pressure and film thickness profiles at Y= 0 of Venner’s results [18] and present simulated results

Fig. 3  Pressure distributions with six implementation methods (Waviness surface, U = 4×10^{-11})

Fig. 4  Film thickness distributions with six implementation methods (Waviness surface, U = 4×10^{-11})

Fig. 5  Pressure and film thickness profiles at Y = 0 with six implementation methods (Waviness surface, (a) U = 4×10^{-12}, (b) 4×10^{-13}, and (c) 4×10^{-14})

Fig. 6  Pressure distributions in the Hertzian contact zone with six implementation methods (Waviness surface, U = 4×10^{-14})

Fig. 7  Film thickness distributions in the Hertzian contact zone with six implementation methods (Waviness surface, U = 4×10^{-14}, λ=5×10^{-5})

Fig. 8  Visualization of generated rough surface (a) 3D view (b) Profile along X= 0 (c) Profile along Y=0

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Fig. 10 Pressure distributions in the Hertzian contact zone with six different implementation methods (Rough surface, U = 4×10^{-13})
Fig. 11  Film thickness distributions in the Hertzian contact zone with six implementation methods (Rough surface, \( U = 4 \times 10^{-13} \))

Fig. 12  Pressure and film thickness profiles at \( Y = 0 \) with six different implementation methods (Rough surface, \( U = 4 \times 10^{-16} \))

Fig. 13  Contact area in the Hertzian contact zone with six implementation methods (Rough surface, \( U = 4 \times 10^{-16} \))
Table Caption List

Table 1  Six implementation methods of the semi-system method

Table 2  Parameters of the EHL problem

Table 3  Relative simulation time (Waviness surface)

Table 4  Lambda ratio and relative simulation time (Rough surface, $U = 4 \times 10^{-13}$)

Table 5  Lambda ratio, contact ratio and relative simulation time (Rough surface, $U = 4 \times 10^{-16}$)
Appendix A

The steady state Reynolds equation:

$$\frac{\partial}{\partial x}\left(\rho h^3 \frac{\partial p}{\partial x}\right) + \frac{\partial}{\partial y}\left(\rho h^3 \frac{\partial p}{\partial y}\right) = \frac{u_1 + u_2}{2} \frac{\partial (\rho h)}{\partial x}$$  \hspace{1cm} A(1)

is non-dimensionalized by introducing the following dimensionless variables:

$$X = \frac{x}{a}, Y = \frac{y}{b}, H = \frac{h R_x}{a^2}, P = \frac{P}{p_0}, \bar{\eta} = \frac{\eta}{\eta_0}, \bar{\rho} = \frac{\rho}{\rho_0}$$

where a and b are the radius of Hertzian contact in two principal directions. In point contact, a equals b. $p_h$ is the maximum Hertzian contact pressure value. $R_x$ is the radius of curvature in the x direction, and $\eta_0$ and $\rho_0$ are respectively the ambient viscosity and density of the lubricant.

The non-dimensional form of A(1) is then

$$\frac{\partial}{\partial X}\left(\varepsilon_x \frac{\partial P}{\partial X}\right) + \frac{\partial}{\partial Y}\left(\varepsilon_y \frac{\partial P}{\partial Y}\right) = \frac{\partial (\bar{\rho} H)}{\partial X}$$  \hspace{1cm} A(2)

where

$$\varepsilon_x = \varepsilon_y = \frac{a^3 p_h}{12 \eta_0 u_x R_x^2} \frac{\bar{\rho} H^3}{\bar{\eta}}$$

$$u_x = \frac{u_1 + u_2}{2}$$

Film thickness equation:

$$h(x, y) = h_0 + \frac{x^2}{2R_x} + \frac{y^2}{2R_y} + \delta_1(x, y) + \delta_2(x, y) + \gamma(x, y)$$  \hspace{1cm} A(3)

In non-dimensional form:
\[ H(x, y) = \frac{h(x, y)R}{a^2} = H_{\infty} + \frac{X^2}{2} + \frac{Y^2}{2} + (\delta_1(x, y) + \delta_2(x, y) + \nu(x, y)) \frac{R}{a^2} \] \quad A(4)

Elastic deformation equation:

\[ v(x, y) = \frac{2}{\pi E} \int_{\Omega} \frac{p(\xi, \zeta)}{\sqrt{(x - \xi)^2 + (y - \zeta)^2}} d\xi d\zeta \] \quad A(5)

In non-dimensional form:

\[ V = \frac{2}{\pi} \int_{\Omega} \frac{P(\xi, \zeta)}{\sqrt{(X - \xi)^2 + (Y - \zeta)^2}} d\xi d\zeta \] \quad A(6)

The Hertzian contact formulas are used as follows:

\[ a = b = \left( \frac{3wR}{2E} \right)^{\frac{1}{3}}, \quad p_h = \frac{3w}{2\pi a^2} \] \quad A(7)

Load balance equation:

\[ w(t) = \int_{\Omega} p(x, y) dx dy \] \quad A(8)

In non-dimensional form:

\[ W = \frac{2\pi}{3} \] \quad A(9)

Barus viscosity pressure equation:

\[ \eta = \eta_0 \exp(\alpha p) \] \quad A(10)

In non-dimensional form:

\[ \bar{\eta} = \exp(\alpha p_h P) \] \quad A(11)

Density pressure equation:
\[
\frac{\rho}{\rho_0} = 1 + \frac{0.6p}{1 + 1.7p} \quad \text{A(12)}
\]

In non-dimensional form:

\[
\bar{\rho} = 1 + \frac{\bar{C}_a \bar{P}}{1 + \bar{C}_b \bar{P}} \quad \text{A(13)}
\]

where

\[
C_a = 0.6p_h, \quad C_b = 1.7p_h \quad \text{A(14)}
\]
Appendix B

Expressions of the discrete Reynolds equation according to six discrete methods.

The discrete equation set has the form shown below.

\[ \alpha_{i,j}^{(s)} p_{i-1,j}^{(s+1)} + \beta_{i,j}^{(s)} p_{i,j}^{(s+1)} + \gamma_{i,j}^{(s)} p_{i+1,j}^{(s+1)} = b_{i,j}^{(s)} \]

Index No. (1 1 0)

\[ \alpha_{i,j}^{(s)} = e_{i-1/2,j}^{(s)} \]

\[ \beta_{i,j}^{(s)} = \left[ -\left( e_{i-1/2,j}^{(s)} + e_{i+1/2,j}^{(s)} + e_{i,j-1/2}^{(s)} + e_{i,j+1/2}^{(s)} \right) \right] - \left( \bar{\rho}_{i,j}^{(s)} D_{i,j}^{i,j} - \bar{\rho}_{i-1,j}^{(s)} D_{i-1,j}^{i-1,j} \right) \Delta X \]

\[ \gamma_{i,j}^{(s)} = e_{i+1/2,j}^{(s)} \]

\[ b_{i,j}^{(s)} = -\left( e_{i,j-1/2}^{(s)} p_{i,j-1}^{(s)} + e_{i,j+1/2}^{(s)} p_{i,j+1}^{(s)} \right) + \left[ (\rho H)_{i,j} - (\rho H)_{i-1,j} - \left( \bar{\rho}_{i,j}^{(s)} D_{i,j}^{i,j} - \bar{\rho}_{i-1,j}^{(s)} D_{i-1,j}^{i-1,j} \right) \right] \Delta X \]

Index No. (1 0 0)

\[ \alpha_{i,j}^{(s)} = e_{i-1/2,j}^{(s)} - \left( \bar{\rho}_{i,j}^{(s)} D_{i-1,j}^{i-1,j} - \bar{\rho}_{i+1,j}^{(s)} D_{i+1,j}^{i-1,j} \right) \Delta X \]

\[ \beta_{i,j}^{(s)} = \left[ -\left( e_{i-1/2,j}^{(s)} + e_{i+1/2,j}^{(s)} + e_{i,j-1/2}^{(s)} + e_{i,j+1/2}^{(s)} \right) \right] - \left( \bar{\rho}_{i,j}^{(s)} D_{i,j}^{i,j} - \bar{\rho}_{i-1,j}^{(s)} D_{i-1,j}^{i-1,j} \right) \Delta X \]

\[ \gamma_{i,j}^{(s)} = e_{i+1/2,j}^{(s)} - \left( \bar{\rho}_{i,j}^{(s)} D_{i,j}^{i-1,j} - \bar{\rho}_{i+1,j}^{(s)} D_{i+1,j}^{i-1,j} \right) \Delta X \]

\[ b_{i,j}^{(s)} = -\left( e_{i,j-1/2}^{(s)} p_{i,j-1}^{(s)} + e_{i,j+1/2}^{(s)} p_{i,j+1}^{(s)} \right) - \left( \bar{\rho}_{i,j}^{(s)} D_{i,j}^{i-1,j} - \bar{\rho}_{i-1,j}^{(s)} D_{i-1,j}^{i-1,j} \right) \Delta X \]

Index No. (0 1 0)

\[ \alpha_{i,j}^{(s)} = e_{i-1/2,j}^{(s)} \]
\[ \beta^{(s)}_{i,j} = \left[ -\left( e^{(s)}_{i-1/2,j} + e^{(s)}_{i+1/2,j} + e^{(s)}_{i,j-1/2} + e^{(s)}_{i,j+1/2} \right) - \left( \tilde{p}^{(s)}_{i,j} D_{i,j} - \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} + \tilde{p}^{(s)}_{i,j-1} D_{i,j-1} - \tilde{p}^{(s)}_{i-1,j-1} D_{i-1,j-1} \right) \Delta X \right] \]

\[ \gamma^{(s)}_{i,j} = e^{(s)}_{i-1/2,j} \]

\[ b^{(s)}_{i,j} = -\left( e^{(s)}_{i,j-1/2} P_{i,j-1} + e^{(s)}_{i,j+1/2} P_{i,j+1} \right) + \left[ \rho \left( H_{i,j} - H_{i-1,j} \right) - \left( \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} - \tilde{p}^{(s)}_{i-1,j-1} \right) P_{i,j} \right] \Delta X \]

Index No. (0 1 1)

\[ \alpha^{(s)}_{i,j} = e^{(s)}_{i-1/2,j} + \phi^{(s)}_{i,j} \Delta X \]

\[ \beta^{(s)}_{i,j} = \left[ -\left( e^{(s)}_{i-1/2,j} + e^{(s)}_{i+1/2,j} + e^{(s)}_{i,j-1/2} + e^{(s)}_{i,j+1/2} \right) - \left( \tilde{p}^{(s)}_{i,j} D_{i,j} - \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} + \phi^{(s)}_{i,j} \right) \Delta X \right] \gamma^{(s)}_{i,j} = e^{(s)}_{i-1/2,j} \]

\[ b^{(s)}_{i,j} = -\left( e^{(s)}_{i,j-1/2} P_{i,j-1} + e^{(s)}_{i,j+1/2} P_{i,j+1} \right) + \left[ \rho \left( H_{i,j} - H_{i-1,j} \right) - \left( \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} - \tilde{p}^{(s)}_{i-1,j-1} \right) P_{i,j} \right] \Delta X \]

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\[ \alpha^{(s)}_{i,j} = e^{(s)}_{i-1/2,j} - \left( \tilde{p}^{(s)}_{i,j} D_{i,j} - \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} \right) \Delta X - \left( \tilde{p}^{(s)}_{i,j} D_{i,j} - \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} \right) \Delta X \]

\[ \beta^{(s)}_{i,j} = \left[ -\left( e^{(s)}_{i-1/2,j} + e^{(s)}_{i+1/2,j} + e^{(s)}_{i,j-1/2} + e^{(s)}_{i,j+1/2} \right) - \left( \tilde{p}^{(s)}_{i,j} D_{i,j} - \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} + \phi^{(s)}_{i,j} \right) \Delta X \right] \gamma^{(s)}_{i,j} = e^{(s)}_{i-1/2,j} - \left( \tilde{p}^{(s)}_{i,j} D_{i,j} - \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} \right) \Delta X \]

\[ b^{(s)}_{i,j} = -\left( e^{(s)}_{i,j-1/2} P_{i,j-1} + e^{(s)}_{i,j+1/2} P_{i,j+1} \right) + \left[ \rho \left( H_{i,j} - H_{i-1,j} \right) - \left( \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} - \tilde{p}^{(s)}_{i-1,j-1} \right) P_{i,j} \right] \Delta X \]

\[ b^{(s)}_{i,j} = -\left( e^{(s)}_{i,j-1/2} P_{i,j-1} + e^{(s)}_{i,j+1/2} P_{i,j+1} \right) + \left[ \rho \left( H_{i,j} - H_{i-1,j} \right) - \left( \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} - \tilde{p}^{(s)}_{i-1,j-1} \right) P_{i,j} \right] \Delta X \]

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\[ \alpha^{(s)}_{i,j} = e^{(s)}_{i-1/2,j} + \phi^{(s)}_{i,j} \Delta X - \left( \tilde{p}^{(s)}_{i,j} D_{i,j} - \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} \right) \Delta X \]

\[ \beta^{(s)}_{i,j} = \left[ -\left( e^{(s)}_{i-1/2,j} + e^{(s)}_{i+1/2,j} + e^{(s)}_{i,j-1/2} + e^{(s)}_{i,j+1/2} \right) - \left( \tilde{p}^{(s)}_{i,j} D_{i,j} - \tilde{p}^{(s)}_{i-1,j} D_{i-1,j} + \phi^{(s)}_{i,j} \right) \Delta X \right] \]
\[
\gamma_{i,j}^{(s)} = \varepsilon_{i+1/2,j}^{(s)} - \left(\bar{\rho}^{(s)}_{j} D_{i+1,j}^{(s)} - \bar{\rho}^{(s)}_{j} D_{i+1,j}^{(s-1)}\right) \Delta X
\]

\[
b_{i,j}^{(s)} = -\left(\varepsilon_{i,j-1/2}^{(s)} P_{i,j-1}^{(s)} + \varepsilon_{i+1/2,j}^{(s)} P_{i+1,j}^{(s)}\right) + \left[\frac{\rho_{j} \left( H_{j} - H_{i-1,j} \right) - \left(\bar{\rho}^{(s)}_{i,j} D_{i,j}^{(s)} - \bar{\rho}^{(s)}_{i,j} D_{i+1,j}^{(s-1)}\right) \rho_{j} P_{i,j}}{-\left(\bar{\rho}^{(s)}_{i,j} D_{i+1,j}^{(s)} - \bar{\rho}^{(s)}_{i,j} D_{i+1,j}^{(s-1)}\right) P_{i+1,j}}\right] \Delta X
\]