Planar jet stripping of liquid coatings: Numerical studies

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In this paper, we present a detailed example of numerical study of the film formation in the context of metal coating. Subsequently we simulate wiping of the film by a planar jet. The simulations have been performed using Basilisk, a grid-adapting, strongly optimized code. Mesh adaptation allows for arbitrary precision in relevant regions such as the contact line or the liquid–air impact zone, while coarse grid is applied elsewhere. This, as the results indicate, is the only realistic approach for a numerical method to cover the wide range of necessary scales from the predicted film thickness (hundreds of microns) to the domain size (meters). The results suggest assumptions of laminar flow inside the film are not justified for heavy coats (liquid zinc). As for the wiping, our simulations supply a great amount of instantaneous results concerning initial film atomization as well as film thickness.

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1. Introduction

1.1. Jet stripping of liquid coatings

We present here a numerical study of the liquid metal coating process. First, liquid film formation on a vertically climbing wall is simulated. Subsequently – in most cases in the same simulation – we simulate the wiping of the created film by a planar air jet. These processes are of major industrial significance e.g. in metallurgy (Takeishi et al., 1995), photography, painting and manufacturing of materials (Bajpai, 2018), where the need arises to control the thickness of the deposit. One of the means to establish this control is by the use of an airflow, for example with flat planar jets known as “air-knives”. These, located horizontally above the coat reservoir, will act by wiping the film in a controlled manner. However, the effect of the jets is not fully predictable when the airflow issuing from them becomes turbulent, especially around the product edges. The significant kinetic energy of the incoming turbulent airflow may cause unwanted coating atomization or defective coating around the product edges, forcing the operators to lower injected air velocity below certain thresholds – this are in practice found empirically. There is a sustained need for studies of such a configuration for the purposes such as process optimization.

Forming of the liquid film – the basis of the coat formation procedure – has been studied both experimentally and analytically by many authors, starting with the now classical results of (Landau and Levich, 1942). Analytic solutions were found e.g. by [Groenveld, 1970] who focused on withdrawal with “appreciable” inertial forces (relatively high Reynolds number (Re) flows) or (Spiers et al., 1973) who have modified the withdrawal theory of Landau and Levich, obtaining improved predictions for the film thickness that were also confirmed experimentally. Later, (Snoeijer et al., 2008) investigated extensively the film formation regimes in which bulges are formed, focusing on the transition between zero-flux and LL-type films.

As mentioned, in the process of coating, liquid is drawn from a reservoir onto a retracting sheet, forming a coat. The latter is characterized by phenomena such as longitudinal thickness variation (in 3D) or waves akin to that predicted by Kapitza & Kapitza (Cheng, 1994) (visible in two dimensions as well). While the industry standard configuration for Zinc coating is marked by coexistence of medium Capillary number (Ca=0.03) and film Reynolds number Re₁ > 2000, we present also parametric studies in order to establish if our numerical method influences the film regimes obtained in the target configuration. Note that metallurgical effects (solidification) are neglected, as they don’t play a role in the initial, rapid stages of film formation (Hocking et al., 2011).

As mentioned, significant Reynolds numbers in the air are expected in the wiping stage. Although the airflow effects on the coat

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can be studied using time averaging (Myrillas et al., 2013), certain instantaneous effects, such as the formation of bulges, edge effects or film defects will not be accounted for. Thus, numerical simulations are a promising tool to supplement experimental studies in this field. One of the first systematic accounts of the jet stripping of liquid coatings comes from (Ellen and Tu, 1984) who have shown analytically that not only is the pressure gradient acting on the film, but also the surface shear stress terms play an important role in the coating thickness modification. (Tuck, 1983) derived analytical expressions for a dependency between jet airflow velocity and resulting film thickness – assuming that only the pressure gradients play a role in the film deformation – and adopting the lubrication approximation for the film flow. The work (Takeishi et al., 1995) provided certain numerical solutions for velocity and shear profiles at the film-air interface during wiping (using a glycerine solution as the coating liquid).

The authors of (Hocking et al., 2011) analysed the problem numerically using a simplified model – including empirically determined shape functions – and a method of lines to study the modified equations of (Tuck, 1983). They concluded e.g. that disturbances of the coating (as bulges/dimples) above the impact zone will persist more likely for thinner coats, as thick ones `compensate’ for that with surface tension and solidification intensity.

In this work, we follow the DNS (Tryggvason et al., 2011) approach, i.e. we solve a complete set of Navier–Stokes equations describing the flow in both phases (in the one-fluid formulation (Delhaye, 1973)) with proper boundary conditions, if permitted by the computational code used. A similar approach has previously been adopted e.g. by (Lacanette et al., 2006), however their 2006 paper was limited to the two-dimensional Large Eddy Simulation (LES) approach. Still, they were able to recover the pressure profiles of an impinging jet, or predict splashing will take place below the impingement area. The authors of (Myrillas et al., 2013) performed a study very similar to (Lacanette et al., 2006) – but substituting dipropylene glycol for the coating liquid – yielding e.g. profiles of the film in the impingement zone. An even more basic 2D study using the VOF method was published in (Yu et al., 2014), yielding information e.g. about certain droplet trajectories after impact. In this paper, we continue such a numerical approach, this time applying a three-dimensional code with very high spatio-temporal resolutions and adaptive mesh refinement.

This paper is structured as follows. In the further parts of the Introduction, we outline the geometrical specification of the setup as well as its physical parameters. Section 2 deals with the mathematical description of the flow at hand. In Section 3, we briefly describe the computational methods chosen for the study. Subsequently, Section 4 presents all the results obtained from simulations, and the conclusions are presented in Section 5.

1.2 Problem specification

The investigated configuration is visible in Fig. 1. Dimensions visible in the leftmost illustration pertain to our target (or “industrial”) configuration. The coating liquid is drawn from the reservoir C at the bottom, and deposits on the vertical band A as the latter moves upwards. Subsequently, air injected from the nozzles B collides with the coated band A, interacts with the film deposit, and leaves the flow domain \( \Omega \) below and above the nozzle(s); outlets are drawn in Fig. 1 (left) with grayed lines.

As we can see in the side-view (Fig. 1 left), the nozzle-band distance \( d_{nf} \) is measured at \( d_{nf} = 10 \)mm in the industrial configuration. The nozzle diameter \( d \) is 1mm. The proportions in the two-dimensional schematic areforgone for presentation purposes, hence the vertically elongated domain shape is slightly more visible in the 3D rendering (Fig. 1 right). Gravity is taken into account, and the upward band velocity is in most cases taken at \( u_{wall} = 2 \)m/s. Except where noted, we have decided to choose liquid zinc as the coating liquid. Properties of \( 30 \text{Zn} \) are assumed, that is surface tension \( \sigma = 0.71[\text{N/m}] \), density \( \rho_l = 6500[\text{kg/m}^3] \) and viscosity \( \mu_l = 3.17 \cdot 10^{-3}[\text{Pa s}] \). Properties of the surrounding gas - which in all cases is air - are density \( \rho_g \approx 1.22[\text{kg/m}^3] \) and viscosity \( \mu_g = 2.1 \cdot 10^{-5}[\text{Pa s}] \).

As explained below, we introduce multiple sets of boundary conditions in three dimensions. To concisely refer to them, we introduce the following nomenclature to designate the investigated configurations. Two geometries considered will be termed \( G_i \) with \( i = 1, 2 \). If present, the second lower index may be used to designate the grid resolutions used. This index will equal the power of two corresponding to the maximum refinement used by the Basilisk code described further. And so, for example, \( G_{14} \) stands for the first configuration at \( 2^{14} \)-equivalent refinement level. Most of the distinguishing features of the two geometries have been delineated in Table 1. In case other quantities (such as injection velocity \( u_{inj} \)) are varied between configurations, it will be designated in parenthesis (e.g. \( G_{2+11} \left( u_{inj} = 42 \right) \) stands for the \( G_2 \) configuration on a \( 2^1 \)-equivalent grid with the air injection velocity equal to 42 m/s). Using the above terminology, we can now revisit Fig. 1: the configuration presented on the left-hand-side is recognized as \( G_2 \) in 2D, while the r-h-s of Fig. 1 depicts the three-dimensional \( G_1 \).

Our departure point is the full “industrial” configuration \( G_1 \), visible in Fig. 1 on the right. As sketched in Fig. 1, we orient the geometry so that \( y \) is the vertical direction, and air injection takes place along the \( x \) axis with nozzles extended in the \( z \) directions. As visible in Table 1 this configuration involves both “air-knife” nozzles; additionally there are outlet areas at the \( z+ \), \( z- \) and \( y+ \) domain walls. Split boundary conditions are used to ensure that fluid outflow takes place e.g. only above liquid bath level. As shown in Table 1, the thickness \( h_{noz} \) of the coated band A is kept at 1mm. The position of the coated wall along the \( x \) axis is given by \( x_{noz} \) in the Table: it is \( x \)-centered in the \( G_1 \) configuration, moved leftmost in \( G_3 \). In all cases, we impose the upward wall velocity \( u_{wall} = 2 \) (m/s). Due to the fact that the \( z \)-extent (depth) of the coated wall is smaller than the nozzle depth, the \( G_1 \) configuration allows the air issuing from both nozzles to collide. This ends the description of the \( G_1 \) configuration.

An additional configuration is rendered in Fig. 2. As with Fig. 1, note that the rendering is not fully up-to-scale: dimensions used in actual simulations are given in Table 1. This \( G_2 \) configuration has been created from \( G_1 \) by including only half of it (and a symmetry boundary condition at the \( x- \) direction. In other words, the \( G_2 \) configuration is a three-dimensional realization of the sketch presented on the left-hand-side of Fig. 1. Going into \( G_2 \), the depth \( z \)-extent of the coated wall has also been slightly decreased (from 15 to 5 cm) to limit computational cost of the simulation. Still, in the \( G_2 \) configuration the film is formed gravitationally and the airknife-liquid interaction is maintained. Since the coated wall is now centered at \( x = 0 \), only half of its thickness \( (x-span) \) is included in the \( G_2 \) configuration, which makes \( G_2 \) less suited for studies e.g. of the edge effects of the coated band. Instead, more computational resources can be directed at studying the air-liquid interactions. Of course, the \( G_2 \) includes only a single nozzle.

2. Description of the flow

2.1 Governing equations

In all the cases presented henceforth, the full Navier-Stokes equations:

\[
\frac{\partial \mathbf{u}}{\partial t} + (\mathbf{u} \otimes \mathbf{u}) = -\frac{1}{\rho} (\nabla \cdot (\mu \mathbf{D} - \mathbf{p})) + \mathbf{f},
\]

(1)
are solved, assuming the flow to be incompressible:
\[ \nabla \cdot \mathbf{u} = 0. \tag{2} \]

In (1), \( \mathbf{u} \) stands for the velocity vector and \( p \) signifies pressure. The liquid properties are designated by \( \mu \) and \( \rho \) for viscosity and density, respectively. Symbols \( \mathbf{I} \) and \( \mathbf{D} \) represent unitary and rate of strain tensors, respectively, with \( \mathbf{D} \) defined as
\[ \mathbf{D} = \nabla \mathbf{u} + \nabla^T \mathbf{u}. \]

Gravity is taken into account and represented by the body force \( \mathbf{f}_g \). Capillary forces are represented in (1) by \( \sigma \mathbf{n} \kappa \delta \), where \( \sigma \) is the surface tension coefficient, \( \kappa \) is the curvature of the interface \( S \), and \( \delta \) is a Dirac function defined only in the cells containing an interface. We adopt the one-fluid approach (Delhaye, 1973), in which density and viscosity can change at \( S \), and a pressure jump is possible there in case of non-zero surface tension. We will occasionally refer to the directions “up” and “down” which in both 2D and 3D simulations are to be associated with the \( y \)-axis. Moreover, we will occasionally denote the fluid properties with suffixes \( l \) and \( g \) (liquid/gas).

2.2. Film formation and the air-knife theory

In this subsection, we shall present a simplified theoretical description of the air-knife process, that allows us to estimate the final coating thickness. We employ the thin film approximation, which, followed by an order of magnitudes analysis and the associated simplifications, brings us to the following form of (1):
\[ \mu \frac{\partial^2 u_x}{\partial x^2} = \rho g + \frac{\partial p}{\partial y}. \tag{3} \]

Comparing magnitudes of pressure and shear stress imposed by the air flow with that of surface tension often results in dropping the latter from the model. The boundary conditions associated with (3) are the “no-slip” restriction at \( x = 0 \), where \( u_x = u_{\text{wall}} \), and an imposed shear stress on the surface \( (x = h) \) by the air flow:
\[ \tau_{xy} = \mu \frac{\partial u_y}{\partial x} \bigg|_{x=h}. \tag{4} \]
Upon integrating (3) and applying the boundary conditions, one obtains a parabolic profile inside the vertically moving film:

\[
 u_y(x) = -\frac{\rho g + \partial_y p}{2\partial_t} x(2h - x) + \frac{\tau_{xy}}{\mu_l} x + u_{wall}. 
\]

(5)

The volume flux through the film, by unit length is therefore:

\[
 q = \int_0^h u_y(x) \, dx = -\frac{\rho g + \partial_y p}{3\partial_t} h^3 + \frac{\tau_{xy}}{2\partial_t} h^2 + u_{wall} h. 
\]

(6)

There is a single length scale in the problem \( h_0 = [\mu_l u_{wall}/(\rho g)]^{1/2} \). Using this length, one can define a dimensionless flux,

\[
 Q := \frac{q}{u_{wall} h_0}, 
\]

(7)

a dimensionless film thickness,

\[
 T := \frac{h}{h_0}, 
\]

(8)

and a dimensionless shear stress,

\[
 S := \frac{\tau_{xy} h_0}{\mu_l u_{wall}}. 
\]

(9)

The dimensionless effective gravitational acceleration writes as:

\[
 G := 1 + \frac{\partial_y p}{\rho g}. 
\]

(10)

Subsequently, the dependency amongst these dimensionless groups arises from (6) and writes as:

\[
 Q = T \left( 1 - \frac{GT^2}{3} \right) + \frac{ST^2}{2}. 
\]

(11)

Note that (11) can also be used to study the gravitational film formation, that is the formation of a film by a withdrawal, before the action of the air-knives (\( S = 0 \) and \( G = 1 \)). We will employ this to estimate the Groenveld's thickness (denoted \( T_c \) here) of the film when studying its formation in Section 4.2 (see also (Groenveld, 1970)). In addition, one can readily derive the zero-flux thickness by additionally imposing \( Q = 0 \), thus obtaining \( T_0 = \sqrt{3} \).

Our main interest however, is to find an estimate of the final coating thickness \( T_f \) due to the thinning effect of the air-knives. Examining the right-hand side of (11), we see that for a fixed value of the flux \( Q \), the cubic function for the thickness \( T \) admits two positive solutions \( T_e \) and \( T_s \). This is valid for every value of \( y \) since \( G \) and \( S \) are functions of that coordinate. There exists a transitional critical point at a certain \( y = y_c \), where the cubic function admits a double positive root \( T_c \). Thus, for \( y < y_c \), the physical thickness would be \( T_e \) whereas for \( y > y_c \), the selected thickness should be \( T_s \). In the spirit of what has been done in (Hocking et al., 2011), the condition for this critical transition is:

\[
 \frac{\partial Q}{\partial T} = 1 - G_e T_e^2 + S_c T_c = 0. 
\]

(12)

Knowing the pressure and the shear stress profiles, we expect this critical thickness to occur at \( y < y_c \), where \( S_c < 0 \) and \( G_e > 0 \). Therefore, a positive solution to (12) is:

\[
 T_c = \frac{S_c}{2G_e} \left( 1 - \sqrt{1 + \frac{4G_e S_c}{S_c^2}} \right). 
\]

(13)

We then approximate the pressure imposed by the air flow and the shear stress as

\[
 p \sim c_p \rho g u_{inj}^2, \quad \tau_{xy} \sim c_s \rho g u_{inj}^2, 
\]

(14)

where \( c_p \) and \( c_s \) are coefficients to be determined from numerical simulations. In addition, we define a length scale \( l_0 \) designating the area of the film over which the jet has the most effect. It is typically larger than the nozzle diameter due to the downstream spread of the jet, and can be readily determined from the numerical simulations. Hence, we approximate the gradient of pressure as \( \partial_y p \sim c_p \rho g u_{inj}^2/l_0 \). In an attempt to find the dominant term in the effective gravitational acceleration \( G \), we find that \( \partial_y p/(\rho g) - c_p \rho g / \mu_l \), where the \( Fr = u_{inj}^2/(g l_0) \) is the Froude number, typically very large for the standard operational parameters. Therefore, \( \partial_y p/(\rho g) \gg 1 \), and the effective gravitational acceleration can be written as \( G_e = \partial_y p/(\rho g) \). We now seek the simplification of solution (13). For that purpose, we approximate the term

\[
 G_e S_c^2 \sim (c_p/c_s^2)(u_{wall}/u_{inj})/(m Re) \]

(15)

where, \( m = \mu_g / \mu_l \) is the gas to liquid viscosity ratio, and \( Re = \rho u_{inj} l_0 / \mu_g \) is the Reynolds number based on the previously defined length scale \( l_0 \). At the standard operating conditions, \( Re \gg 1 \) and \( u_{inj} \gg u_{wall} \). Therefore, typically, \( G_e S_c^2 \ll 1 \), leading to a rough estimation of the critical thickness:

\[
 T_c \simeq -\frac{1}{S_c} = \frac{1}{|S_c|}. 
\]

(16)

Mass is conserved; hence, for a steady solution, the flux is constant throughout the film, and evaluated from (11) to be:

\[
 Q_e = \frac{G_e}{3S_c^2} - \frac{1}{2S_c} \simeq \frac{1}{2S_c}. 
\]

(17)

employing the previous order of magnitude analysis.

Downstream in the film, and far enough from the jet impingement area, the thickness, denoted as \( T_e \) has reduced due to the air-knife effect. The film is usually thin enough so that a constant velocity, that of the rising wall, may be assumed throughout. In (11), this yields \( G = S = 0 \). And the final coating thickness would therefore be:

\[
 T_e = Q_e = \frac{1}{2|S_c|}. 
\]

(18)

This falls in line with the findings of (Hocking et al., 2011), where the end result of a detailed calculation yields a ratio \( T_e/T_e \ll 2 \). Our estimates of the coating thickness will be given below (see Section 4.3), as well as summarized for the industrial parameters in Table 3.

3. Computational methods

In the research presented here we have applied the “Basilisk” computational code (Popinet, 2015), which is an in-house, GPL-licensed code whose main developer is one of the present authors (SP). It is a descendant of the “ Gerris” code (Popinet, 2009) and as the latter, it allows for the local adaptive mesh refinement (AMR) (Puclet and Saltzman, 1992) using the quad/oct-tree type mesh – regular, structured cubic meshes without refinement are also possible. The code is optimised for speed and capable of both OpenMP (single node) and MPI (multi-node) parallelism. Most recently, Basilisk has been applied e.g. to model compressible bubble dynamics (Fuster and Popinet, 2018), propose certain single-column models in meteorological simulations (van Hooft et al., 2018), or simulate turbidity currents (Yang et al., 2018). It is a multi-equation solver, making it de-facto a multi-physics code. Basilisk supplies a macro language (the Basilisk C) built atop the C99 standard of the C programming language. Upon compilation of the simulation case files, they are first parsed into clean C by the builtin parser/lexer (qcc). Then, the compilation is finalized with libc as the only dependence. This approach greatly enhances code portability.

The Navier-Stokes equations are solved using the projection method (Tryggvason et al., 2011) with a procedure similar to that
applied in Gerris (Popinet, 2003; 2009). Centered discretization is used for all scalar and vector fields, with additional face-centered values defined for \( \mathbf{u} \) which are used e.g. to ensure divergence-free condition during mesh refinement. For consistency reasons, advection term of (1) is defined and calculated on cell faces as is \( \nabla p \). Advection fluxes are obtained using the Bell-Collela-Glaz scheme (Bell et al., 1989). All discretizations are finite-differencing up to second order, unless noted otherwise. The Runge-Kutta scheme is used for time advancement. Optimisations of Poisson equation’s solution are achieved by implementing the Multigrid (MG) method (Brandt, 1984).

The Volume of Fluid (VOF) method (Tryggvason et al., 2011) is used to track the interface using geometric interface reconstruction (Anisiewski et al., 2014). In this method, fraction function \( C \) (equal to one or zero in either phases) is passively advected with the flow. Grid cells with fractional \( C \) values are those in which interface is geometrically reconstructed and represented by a line/planes (in two and three dimensions, respectively). Note that \( \mu \) and \( \rho \) are usually local functions of \( C \), obtained by averaging in the interface cells. The interface curvature \( \kappa \) is also computed from \( C \), using the Height-Functions (HF) method (Afkhami and Bussmann, 2008; Popinet, 2009) with a proper treatment close to solution boundaries. Depending on the the local resolution, \( \kappa \) may be obtained by the HF method using full stencils (optimal) or one of two fallback options: parabola fitting of height functions and parabola fitting of the VOF centroids.

Basilisk’s procedure for local mesh adaptation employs the wavelet transform of a given scalar field to assess the latter’s discretization error. If the error is above the user-specified threshold, the grid is locally refined by subdividing it onto four (quad-tree) and eight (octree) sub-cells and performing a prolongation of the coarser-mesh scalar onto children cells to obtain their initial values (the inverse process is termed restriction). For the simulations presented herein, we use \( u \) and \( C \) fields’ error as the refinement criteria with \( 1 \cdot 10^{-3} \) and \( 1 \cdot 10^{-2} \) error thresholds, respectively.

### 3.1. Ensuring momentum conservation in two-Phase flow

The momentum-conserving methods (Vaudor et al., 2017) derive from a variant of the VOF (Hirth and Nichols, 1979) method originally proposed in (Rudman, 1998) to treat two-phase flows with considerable density ratios. The idea is that instead of the simple incompressibility assumption

\[
\nabla \cdot \mathbf{u} = 0.
\]

one writes the mass transport equation in full, as is done in the compressible formulation (Pilliod, 1996):

\[
\frac{\partial \rho}{\partial t} + \nabla \cdot (\rho \mathbf{u}) = 0,
\]

using also the conservative form of the Navier-Stokes equations (not shown) (Vaudor et al., 2017), which contain the momentum term

\[
\nabla \cdot (\rho \mathbf{u} \otimes \mathbf{u}).
\]

Subsequently, in implementation, we calculate density from the fraction function definition:

\[
\rho = \rho_0 C + (1 - C) \rho_0.
\]

which is a reversal of the traditional approach (Hirth and Nichols, 1979). Expression (22) is nontrivial only in the interface cells. The way in which the momentum-conserving methods differ from the traditional two-phase Navier-Stokes equation models using VOF is that subsequently, the \( \rho(C)U \) products found in both (20) and (21) are calculated consistently in the same control volumes. This can be non-trivial if staggered grid discretizations are used, and can be solved either by grid-cell subdivisions (Rudman, 1998) or using sub-fluxes of the fraction function (Vaudor et al., 2017). Thus consistency between transports of mass and momentum are ensured numerically, resulting in a far more robust computation.

### 3.2. Implementation of embedded solids

The problem geometry illustrated in Fig. 1 is nontrivial, due to the fact that flow is expected to take place around walls of the coated band, as well as the edges and corners defining the flat nozzle, i.e. space containing embedded (or immersed) solids, and the computational code used must allow for this. We use a rudimentary technique of locally modifying the velocity field for this purpose. Local modification of scalar fields is a relatively simple technique used when simulating large-scale systems involving solids (Lin-Lin et al., 2016). It is a strongly simplified variant of the Immersed Boundary Method (IBM) of Peskin (Peskin, 2002), which does not modify the grid data structure and is thus compatible with MPI protocol. If we denote the interior of the solid contained by boundary \( \Gamma \) by \( \mathbb{S} \) we can note:

\[
\forall x \in \mathbb{S} \cup \Gamma : \mathbf{u}(x) = 0,
\]

that is, all velocity components are set to zero within the solid. As long as no conditions are needed for \( x \in \Gamma \), the crude approximation provided by (23) yields satisfactory results (Lin-Lin et al., 2016). A moving wall can be prescribed by using a non-zero \( \mathbf{u}_{\text{wall}} \) right-hand side in (23). Note however, that pressure \( p \) is not modified in any way inside the solid \( \mathbb{S} \) which, in principle, may result in its incorrect values especially at the boundary \( \Gamma \). This could be addressed by locally modifying pressure gradients, which in a physical sense is equivalent to defining a certain force which would only be nonzero at the boundary \( \Gamma \). This however complicates the technique to a degree comparable with the implementation of domain reshaping, as optimally, it should be followed by removal of the interior points from the grid.

Instead, we note that for the geometries presented – even the most complicated \( \mathbb{S} \) setup – the domain interior is merely a sum of cuboids: it contains no inclined nor curved surfaces. The no-slip condition at the surface of the substrate wall moving with velocity \( \mathbf{u}_{\text{wall}} \) can be reasonably approximated using (23).

### 3.3. Spatially restricted refinement

To limit the associated CPU cost of the grid refinement, we have employed an additional technique of spatially restricted refinement (for short, we will use the abbreviation ‘SRR’ below). Using SRR is straightforward. The quad/oct-tree data structure in Basilisk results in subdivisions of cells into four/eight sub-cells as the grid is refined in two or three dimensions respectively. The entire domain is a 0-level (parent/root) cell with four/eight 1-level sub-cells and so on. If the refinement criterion is locally fulfilled, Basilisk will keep refining the grid until it reaches the maximum allowed level. The SRR technique locally limits this maximum grid level using a spatial criterion. This means larger discretization errors are intentionally allowed far from regions of interest. The latter regions have to be predefined before the simulation. Then, dynamic grid refinement will act as usual, the only difference being that refinement to the maximum level will take place only in chosen domain sub-areas while outside of them a lower maximum level is forced. This tactic of refinement situates the presented simulation between the block-based (Lakehal, 2010) and point-based (Popinet, 2009) mesh refinement. Its drawback is the increase in the globally calculated numerical dissipation.
4. Results

4.1. Simulation planning

The full, three-dimensional airknife configuration poses numerous challenges for reasons of code stability, CPU cost or the wide range of simulated physical scales. Due to this challenging character, we have tackled the case progressively, including the following steps.

We have commenced with the film formation studies, both in two and three dimensions, the results of which will be presented in Section 4.2. This lets us compare the obtained thickness with the analytical prediction, as well as shed some light on the turbulent, or transitory, character of the film formation itself.

Subsequently, we include a study of the gas dynamics, namely of the impinging air jet without including the liquid. Here, the goal is to validate the single-phase Basilisk result against the known approximation of the u profile. Results can be found in Section 4.3.

Once the two above phenomena are investigated, we present, in Section 4.4 the full, three-dimensional configuration results. This Section contains additionally a relaxed/industrial parameters comparison for $G_1$. Subsequently, the $G_2$ results are presented in Section 4.5, featuring the discussion on varying the injection velocity $u_{inj}$ in this configuration.

4.2. Film formation studies

As mentioned in Section 2.2, the dimensionless zero-flux film thickness can be readily obtained by $\sqrt{3h_0}$ leading to

$$h_0 = \sqrt{\frac{3\mu \bar{u}_{wall}}{\rho f}}.\tag{24}$$

Using (24) we arrive at $h_0 = 5.46 \cdot 10^{-4}\mu m$ for the industrial parameters. With this value, we can make a rough attempt at assessing the Reynolds number characterizing the liquid film formation in the industrial process, that is

$$Re_f(h_0) = \frac{\rho h_0 \bar{u}_{wall}}{\mu_f} = 2240.\tag{25}$$

This value is close to a critical $Re_f$ delineating the laminar and turbulent film formation regimes (see also Table 2). Another estimate is possible if the $h_c$ thickness is used. Note that its dimensionless counterpart $T_c$ was mentioned above in the context of (11). For the high-Re regime, (Groenheld, 1970) proposes $T = 0.52$ and $Q = 0.47$ which results in

$$h_c = 1.63 \cdot 10^{-4}\mu m.\tag{26}$$

To get (26), one uses (7) and (8) and returns to dimensional quantities through a multiplication by $h_0$. Once $h_c$ is used to calculate the Reynolds number, a value of $Re_f = 672$ is found, so three times smaller than $Re_f(h_0)$. To complete the picture we denote that the definition of $Re_f$ introduced by (Colina-Marquez et al., 2016) results in $Re_f = 2600$. To summarize, these estimates of the Reynolds number suggest a possibility of a turbulent character of the film formation process, as the numbers are between second and third order. One could conclude that the coating process used in the industry is a man-made system at the edge of criticality. This is confirmed in the results presented below.

We start our investigation of the film formation regime with the two-dimensional case. Fig. 3 presents the interface geometry at six instances of time where $t \in [0, 0.14]$. For this 2D simulation, the $u_{wall}$ velocity is implemented as a boundary condition; the simulation used the $2^{14}$-equivalent grid. Fig. 3b presents the same interfaces, but using log $x$ for the horizontal axis. First off, it is visible from Fig. 3 that the uptake of the film by the moving wall is somewhat faster than expected from the value of $u_{wall}$. We attribute this to the imposed contact angle of the VOF interface (Afkhami et al., 2018). The wavy character of the film becomes visible towards the $t = 0.14s$ mark, especially visible in Fig. 3b. Groenheld’s thickness prediction $h_c$ is marked with a dashed line in both subfigures, a good overall fit is observed at least in the upper parts of the film. Apparent bulges on the film for $t > 7.8 \cdot 10^{-2}s$ are consistent with a transitory or turbulent regime, we will examine this effect more closely in three dimensions below.

At this grid resolution, the Basilisk grid cells are of the size $\Delta x = L_c/2^{14} \approx 39\mu m$ for the 2D simulation depicted in Fig. 3. This translates to four grid points inside the film with thickness $h_c$ and allows us to get a very rough estimation of the velocity profile inside the forming film, as discussed in the context of the next figure.
Fig. 4 presents the creation of a boundary layer in time (in the \( t > 0.1457s \) range) of the same flow. The profiles have been sampled at \( h = 0.14m \) or \( 0.04m \) above the reservoir. The velocity profile remains parabolic, however it clearly becomes steeper for \( t > 0.1s \) with an apparent plateau extending for \( x > 5 \times 10^{-4} \) suggesting a detachment of the layer adjacent to the plate (Snoeijer et al., 2008). In Fig. 4 we additionally compare the profile for \( t = 0.1457s \) with the analytical expression (5) (dots). Consistency is visible especially very close to the wall, suggesting that the final profiles lend themselves well to those assumed in (Groenewold, 1970), as hinted previously by Fig. 3. Additionally, it is observable in Fig. 4 (inset) that the transition through the profile described by (5) occurs for \( t \in [7.85 \times 10^{-2}s, 1.45 \times 10^{-1}s] \) which coincides with the moment the boundary layer starts forming. This serves as an argument that the film evolution is reasonably well described by the high-Re theory. Moreover, in Fig. 4 profiles are sampled only for \( C = 0 \) (i.e. inside the liquid film). Thus, for each of the lines, the abscissa of its right-hand end-point corresponds to the film thickness \( h(y) \) at \( y = 0.14m \). As one can observe e.g. for \( t \approx 7.8 \times 10^{-2}s \) we have \( h(0.14, t) = 6 \times 10^{-3}m \) whereas for \( t = 1.457 \times 10^{-1}s \) the thickness drops, suggesting a bulge has passed over the point and retracted.

We now shift our attention to three-dimensional cases. Using the \((2^{12})^3\)-equivalent grid, we have performed a three-dimensional simulation \( G_{212} \) to study film formation. Its results are presented in Fig. 5, which could be seen as a 3D analog of the interface geometry presented above in Fig. 3a. Similar time instance, \( t = 1.48 \times 10^{-1}s \) is chosen in Fig. 5. A heavily “rugged” film surface is easily recognizable in Fig. 5a, in which it has been colored by the vertical velocity component \( u_y \). We can observe – from the \( u_y \) values the film is colored with – distinct liquid boundary layer develops directly adjacent to the wall, traveling with velocity \( u_{wall} \). This is fully consistent with liquid velocity profiles presented in Fig. 4 for \( t = 1.45 \times 10^{-1}s \). As we get further away from the boundary layer, the velocity at which the film is climbing drops sharply; Fig. 5a indicates also that surface material crumbles back into the bath (blue areas close to the reservoir height). We have included, as an inset (Fig. 5b) an isosurface for the zero vertical velocity \( (u_y = 0) \), rendered in turquoise against the gray interfacial surface. (Note that \( u_y = 0 \) occurs as well in the gas far from the coated wall. For this reason, parts of the isosurface were removed from Fig. 5b artificially to not obscure the view of the coated wall area.)

In this way, we are able to approximate the stagnation height for \( t = 1.48 \times 10^{-1}s \) as 0.13m e.g. 0.03m above the bath level. Above this height, all flow is upwards. The interface deformations visible throughout the height of the film surface seem sufficiently resolved and not numerically induced. For example, halfway through the film height in Fig. 5 film thickness is of order 0.01m (or eighty times the grid size at 12 levels of grid refinement).

Another simulation is presented, for the \( G_{11} \) configuration, in Fig. 6. Even using a slightly less refined grid (11 levels of refinement, or \( 2048^3\)-equivalent), we still observe a wrinkling of the interface as well, mostly on the coated band edge. In this stage of the flow, the band is fully coated, while some “dimples” appear close to the reservoir surface once zinc is drained. Only a very thin layer of zinc is deposited close to the band edges, as can be seen by the surface color which corresponds to \( u_{wall} = 2 \). The surface of the film undergoes progressive distortion starting from the side of coated band. This applies especially to the coated \( x^+ \) and \( x^- \) walls, in which wrinkling appears progressively further from the
band edges. The turbulent nature of the film is suggested by surface disturbances, along with the fully three-dimensional character of the wrinkles/waves. To our knowledge, this is the first published result of a 3D coating simulation including the edge, and $Re_f$ is far higher than the previously published 2D results (Myrillas et al., 2013; Lacanette et al., 2006).

We continue our examination of the physics of three-dimensional film formation with Fig. 7, which contains velocity profiles for the vertical component ($u_v$) and the transverse component ($u_t$) along the wall height – only the height range of $y \in [0, 0.2]$ is included, as all $t$ values included are smaller than $t = 0.15$s. At that time, the liquid reaches roughly to $y = 0.3$m, consistent with Fig. 3. Four instantaneous profiles are presented with separate point types. Each of the profiles in Fig. 7 has been $z$-averaged so they represent information from the entire width of the coated plate. Also, profiles include data only for $x < 0.001$m (across the film), in other words the measurement window includes only the direct proximity of the coated wall. In Fig. 7a, we observe a transition from a rather smooth $u_v$ profile at $t = 1.5 \cdot 10^{-2}$s to a much more varied, at final pictured stages. Notably, we observe a stagnation region forming close to the bath level (itself drawn with a dashed line) which is consistent with interface geometry observed in Fig. 6. It is expected that $u_t < 0$ velocities are present in this region further from the wall – this however has not been captured with the profile measurement window. Average $(u_y)_{\bar{z}}$ values are consistent with Fig. 6 as well (note that gas velocity is also taken into account in Fig. 7). We now focus our attention on the curve for $(u_y)_{\bar{z}}$ at $t = 0.15$s (red color in Fig. 7a). This curve, although calculated using a three-dimensional simulation, is comparable with Fig. 4 (curve for $t = 0.1457$s). If, using the latter of the mentioned curves, one calculates a mean value (for $x \in [0, 1]$) of $u_y$, it is equal to 1.22 m/s. This value should be at least comparable with Fig. 7a taken for $t = 0.15$s and $y = 0.14$m; in fact, we find $(u_y)_{\bar{z}}(0.14) \approx 1.1$ which is within ten percent of the two-dimensional simulation. The slight discrepancy might be attributed to the $z$-averaging in three dimensions; e.g. presence of the coated band edge, as well as wrinkles pictured in Fig. 6.

Further evidence of the strictly three-dimensional character of the film is found in Fig. 7b, showcasing this time the profiles of the transverse velocity component $u_t$. While close to the beginning of the flow at $t = 0.015$s (blue squares) this component is nearly zero (approximately two-dimensional flow), $u_t$ oscillates with increasing amplitude in the entrainment zone as time progresses, and remains negative everywhere below the bath level. That is to say the net flow of the liquid layers contacting the coated wall is from the coated edge towards the symmetry plane ($z = 0$). As the film forms and its top edge moves further away from the bath, transverse net flow is positive, i.e. towards the coated edge, which is consistent with Fig. 6 and explains the rugged surface of the film in the edge neighborhood. Summarizing, it is possible that at this $Re_f$ values, three-dimensional effects are strongly present and decisive in determining the liquid flow character.

Finally, note also that Fig. 7 features the most resolved of the 3D simulations presented in the paper, at $2^{13}$ which is (locally) equivalent to a grid with $8192^3$ points.

4.3. Single-phase impinging jet study

This subsection presents results of a 2D numerical study of the $G_2$ configuration (Fig. 2). However, for the test presented here, the liquid phase (along with the reservoir) is removed, and we focus on the gas phase. Calculations presented here have been motivated by the need to find the coefficients $c_i$ and $c_p$ in (14), which such a simplified (and cheaper) configuration allows. Additionally, it provides validation for the Basilisk code as shown below.

This simulation used the “industrial” parameters and a $2^{11}$- equivalent grid. Results are presented in Fig. 8. As we can see from Fig. 8b, the airflow develops symmetrically with the nozzle in its axis. The turbulent character of the flow is not immediately visible in the figure due to the application of temporal averaging - the technique was used only in this simulation, as discussed below.

Fig. 8a presents the profile of the velocity component $u_\tau$ normalized by the mean value $u_{\bar{m}}$, taken at $3 \cdot d_{nf}/4$ distance from the stagnation point. To plot this profile, we have applied a combination of time- and ensemble-averaging in order to ensure smoothness (15 simulations were ensemble-averaged). Time-spans used

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1 Due to CPU time and memory restrictions we have not continued this simulation into the air injection stages.
point directly opposing the nozzle outlet. Zooming into Fig. 8b reveals that pressure changes sign very close to the wall, which is consistent with velocity curves predicted by (27) and the existence of boundary layer. According to (Tuu and Wood, 1996), peak pressure evolves with the distance from the nozzle as

$$p_i = \frac{7d}{\mu_f^2}.$$

Values presented in Fig. 8b are about half of predicted by (28), meaning that the potential core is not resolved well enough for $G_{2,11}$. This warrants an increase in refinement, and is one of the reasons for using at least 12 levels of refinement in majority of the simulations.

By fitting the simplified $p_{0t}$ and $\tau_{0t}$ curves, constructed using (14) to the results obtained in this subsection, we were able to establish the values of the $c_t$ and $c_p$ coefficients mentioned in Section 2.2. For the industrial parameters, they amount to $c_p = .90$, $c_t = 0.00325$. These values are plugged into (18) through (15), and the resulting $T_{01}$ value is then made dimensional as $h_1 = T_{01}h_0$. We thus arrive at the dimensional coating thickness $h_1$ value roughly between 20 and 50 microns.

4.4. Three-dimensional wiping simulations. configuration $G_{1}$.

In this section, results for the full $G_1$ configuration are discussed. In most cases, the simulations ran at the $2^{12}$-equivalent resolution, i.e. $\Delta x \approx 100 \mu m$. Thus the grid size is below both $h_{00}$ and $h_c$, but above the expected $h_n$ values. However, due to specifics of the geometric VOF method (Anisiewski et al., 2014), it is possible to represent a film with thickness below $\Delta x$, including advection of such a film with the uniform velocity defined in that cell.

The $G_1$ results within this Section will include variations in boundary conditions, which we refer to as industrial and relaxed parameters. For reference, Table 2 contains parameters for both industrial and relaxed variants of the considered problem. The most important differences between them include an order of magnitude lower liquid density, and a higher $u_{wall}$ in the relaxed set: both of these contribute to sway the balance between gravity and liquid uptake towards the latter. This subsequently leads to a thicker film formed, thus decreasing associated CPU cost needed to perform the simulations. For the same reasons, in the gas phase, velocity $u_{gas}$ is decreased twofold in relaxed parametrisation. This results for example in the zero-flux $h_{00}$ thickness of the film in relaxed parameters being fourteen times that of its value for the industrial parameters. Additional difference between the relaxed and industrial configurations is the coated plate thickness: it is assumed 5mm for the relaxed variant and 1mm in industrial. Nozzle wall thickness is configured analogically. Both changes facilitate the implementation of the simulation geometry in the relaxed case, meaning that coarser grids suffice to implement formulation (23) as more grid-points end up contained in the $\tilde{\Omega}$ region.

We begin with a discussion of the $G_{1,12}$ simulation, which is first presented in Fig. 9 displaying the interface geometry at $t = 0.162s$. This is the stage of the flow at which the film has already been formed on the coated band, and right after the airflow, issuing from the planar nozzles, impacts it. For comparative purposes, the flow for the relaxed configuration is shown in Fig. 9b. For this image, the value of $t$ is obviously different, and the images have been matched via the width of the impact zone. In Fig. 9, nozzle locations are drawn using shading and black outlines in this view – this is done in post-processing and only for illustrative purposes.

\footnote{\textsuperscript{2} Naturally, higher resolutions are required to represent velocity variation within the film.}
Additionally, a cut-plane is positioned in the back-drop (parallel to \( z = 0 \) coordinate) colored by vorticity. For the industrial configuration, at \( t \approx 0.16 \), a relatively wide impact zone is already visible, with individual droplets ejected from the film, as well as rich wrinkling. The structure of the air trace is three-dimensional: even if its character is homogeneous above the nozzle, below it we see two zones with larger traces. Additionally, the edge area is visibly atomized.

Fig. 9b emphasizes the consequences of certain geometrical differences between the industrial and relaxed parameter sets. A thicker coated plate is visible. The coat on the plate edge is seemingly not disturbed except in the impact area where it interacts directly with the turbulent structures resulting from the collision of air emanating from the opposing nozzles. Moreover, some liquid deposits on the nozzle walls (the nozzles are not rendered in Fig. 9b), partly obscuring the view. Large amounts of the coating material crumble down below the impact zone, resembling the “peeling” effect observed in (Myrillas et al., 2013).

The obvious difference between the relaxed and industrial parametrisations for the wiping process is the degree of visible atomization. Comparing the adimensional numbers as seen in Table 2, the airflow \( \text{Re} \) is three times higher in the industrial case (based on \( u_{\text{inj}} \) and \( d \)). The effects are demonstrated in Fig. 10, which displays the relaxed case in the earlier (than in Fig. 9) stages of the macroscopic flow evolution. Significant atomization level is visible compared to Fig. 9a.

Continuing the flow analysis for the industrial case, we turn our attention to Fig. 11 which displays the film geometry at approximately 0.17s. By this time, the lower bulge (A) starts forming below the nozzle level leading to the onset of back-flow into the reservoir. The distance between the points A and B in Fig. 11 can be referred to as the impact zone, at \( t \approx 0.017s \) we estimate its width at 0.04m. The wavy edge film structure visible in Fig. 11 is an early effect of film formation. However, inspecting Fig. 11 we may conclude that the edge film is nearly entirely atomized in the impact zone. In Fig. 11 the same film is seen, looking parallel to the \( z \)-axis, centered at the impact zone. A certain perspective shortcut effect takes place in Fig. 11, as droplets close to the viewpoint, i.e. on the plate edge, seem bigger than those far from it. Besides, the view contains an apparent accumulation of droplets from all the plate’s depth.

The total CPU cost of the 3D, 2\(^{12}\) simulation of the \( G_1 \) configuration is approximated at 122000 CPUh, only twice the cost of 2D simulations presented in previous subsections. However, the maximum refinement level is four times lower here, and the simulated physical time is only about one tenth that of the 2D simulation. The \( G_1 \) configuration includes both the airknives and the edge of the coated band – which, potentially, can produce an abundance of information about the physics of the flow. However, its CPU cost is nearly prohibitive if longer physical times were to be simulated.

\(^3\) Note that in Fig. 10, the shape of the nozzles is distinguishable by looking at the cut-plane in the back.
Thus, we include a more detailed analysis using the $G_2$ configuration, results of which will be presented below.

4.5. The $G_2$ configuration results

In this section, we present results pertaining to the $G_2$ configuration. It enables us to focus on the air-liquid impact study in more detail since, as long as the mean flow is considered, the $G_1$ configuration has an inherent symmetry. In $G_2$, placing the coated wall in the corner of a cubic domain, we use the SRR technique to coarsen the grid proportionally to the distance from the coated walls. This, as the simulations below confirm, has proven sufficient to dampen the turbulent flow far from the zone of interest and prevent backflows. Thus, we simulate only a single nozzle and one side of the coated band, decreasing the CPU cost. All other assumptions are carried over from the $G_1$ configuration.

In Fig. 12, we present a visualisation of the macroscopic shape of the interface for the $G_2$ simulation performed using a $2^{12}$-equivalent grid. This simulation corresponds to industrial parameters (and is the $G_2$-analog of the $G_1$ results mentioned above e.g. in Fig. 11). In three sub-figures, instantaneous shapes are visible for (a) $t = 1.656 \cdot 10^{-3}$ s (b) $t = 1.677 \cdot 10^{-3}$ s and (c) $t = 1.747 \cdot 10^{-3}$ s. Each of the pictures presents two separate view: an isometric one on the left-hand-side, and a side-view (looking along the $z$-axis) on the right-hand-side. The cut-plane positioned at the $z$-domain wall is colored with $\omega$. Varying the cell size in the vorticity cut-plane is, of course, a consequence of employing the SRR technique to limit adaptivity in regions above and below the nozzle. Full resolution is maintained everywhere inside the planar nozzle and within $1 \text{mm}$ of the coated wall. As visible in the r.h.s. images of Fig. 12b and Fig. 12c, the grid coarsening affects interfacial formations as well: the ejected droplets and ligaments are represented with a coarser grid further away from the coated wall.

Directly after the air contacts the liquid in Fig. 12a, we note a distinct imprint of the nozzle shape on it. Three transversal bulges are formed: one below the nozzle, one directly opposing the air outlet and lastly, a small bulge is formed above the nozzle. A mere two milliseconds later, as shown in Fig. 12b, the central bulge - whose liquid is “trapped” by the airflow, has been completely atomized, turning it into a cloud of droplets and ligaments. (This is shown particularly well in the side-view.) This last result is consistent with that of (Yu et al., 2014), who have investigated (in 2D) a flow characterized by a higher Weber Number (We) of 13.5 with a lower density ratio. Their results show a (C) distribution consistent with a cloud of droplets – with temporal averaging, it is displayed as a bulge.

The atomization process results in most of the liquid droplets being ejected out of the field of view. Some examples of fast-moving “glider” droplets are visible as traces just below the nozzle in Fig. 12b and c. In the meantime, the lower and upper bulges move away from the nozzle. In Fig. 12c, we note that the upper bulge has, by $t = 1.747 \cdot 10^{-3}$ advanced approximately $10 \text{mm}$ upwards, and has been considerably smoothed. Compared to the lower bulge, there is almost no atomized material near the upper one. Meanwhile, as suggested by the right-hand-side view in Fig. 12c, the material below the nozzle is partly stripped from the wall and immediately atomized. Fig. 12b suggests that most of the droplets in the impact zone originate from the atomized middle bulge material. Subsequently, the number of droplets below the nozzle in Fig. 12c is far smaller than that visible in Fig. 12b, suggesting that atomization visible in Fig. 12b is a transient phenomenon. Above the level of the nozzle and between the bulges, a thin film is formed, covered by a three-dimensional wave structure as visible in Fig. 12c.

Atomization of the film occurring at the first instance of the air-liquid contact might be investigated by looking at the dimensionless numbers characterizing this interaction. While the film Reynolds number $Re_f$ characterizes mostly the film formation, we formulate the Weber number $We$ involving gas velocity, as follows:

$$We = \frac{\rho_g u_0^2 h}{\sigma}.$$  \hspace{1cm} (29)  

with $h$ standing for film thickness. Definition (29) is first applied to industrial parameters characterized by $u_{inj} = 200$.

Feeding the zero-flux thickness (24) into (29) one obtains $We(h_{00}) = 38.1$. If, instead, we settle upon using Groenveld’s thickness $h_G$, (29) yields $We = 11.4$. Both these values seem consistent with a regime in which atomization might be expected.

Values of the film thickness calculated using various definitions are given in Table 3, which contains also resulting values of the film Reynolds number as well as the Weber number.

As the atomization effect has not been reported previously (Myrillas et al., 2013; Ellen and Yu, 1984) we have decided to study it further. This is motivated by the fact that similar liquid

\footnote{For the “relaxed” configuration presented previously, using $h_{00}$ is more justified as the film Reynolds number is three times lower. Doing so, we obtain $We(h_{00}) = 7.57.$}
breakup could be induced numerically, e.g. by inconsistent momentum transfers (Vaudor et al., 2017), curvature calculation errors, or by not accounting for interactions between liquid-gas interface and vortical structures in the latter phase (Aniszewski, 2016). Thus, we have included a simulation configured as $G_{12}(u_{inj} = 42)$ by decreasing air injection velocity. This configuration is characterized by $We (h_{inf}) = 1.68$ and $We (h_c) = 0.5$ which, again, places the system just below the “edge of criticality” as in a context of $Re_f$ in our film formation study. We thus follow up with an examination of the flow at th decreased injection velocity.

Fig. 13 presents the $G_{12}(u_{inj} = 42)$ simulation roughly 2 milliseconds after air-liquid impact. In the Figure, white shaded iso-surface represents the liquid interface, while several navy blue areas depict the (un-coated) moving wall. The far view presented in Fig. 13a confirms again the turbulent film character below the impact zone (denoted “1” in the Figure). The interface geometry in the entrainment region 1 is comparable to Fig. 6, and should not be associated with the air-liquid interaction. The impact zone is visible above as an area with horizontal wrinkles (marked “2” Fig. 13a). Looking closer at the impact zone we note a small number of gaps (denoted “3”) in the film, mainly close to the edges of the coated band. Defects may results from the expected film thickness being not fully resolved. There are however visible edge coating defects (denoted “4” in Fig. 13b) not likely associated with airflow. This is consistent with Fig. 6 and seems to suggest that not only increased resolutions are required in the neighbourhood of the coated edge, but possibly specific formulation of the boundary conditions at the sharp solid edge (singularity).

Regarding the atomization phenomenon at the instance of air-liquid contact, comparing Fig. 13 with the r.h.s. images in Fig. 12 we note the nearly complete lack of atomized structures for $u_{inj} = 42 m/s$ and $We = 1$.

Another look at the low Weber number flow is provided by Fig. 14. Three sub-figures present the same flow as pictured in Fig. 13 at instances of time $t \in \{0.1736, 0.178\}$. Fig. 14a presents the interface geometry in the impact z1 almost directly following the first air-liquid contact. Effects of the decrease in the Weber number are instantly recognizable: no distinct horizontal liquid bulges are formed along the z direction; instead, smaller-wavelength disturbances are showing within a gradually broadening region, as seen in Fig. 14c. The side-views included in the Figure show a significant decrease in the number of droplets, which we quantify below in Fig. 16. Overall image of the flow is different than that for $We > 10$. Juxtaposing Fig. 12b with Fig. 14c we note the complete absence of the droplet cloud below the impact area. We suspect that in the low-We regime the film is merely disturbed by the airflow, while areas above the nozzle are continuously fed liquid; hence no permanent film thinning should be expected in such flows.

Indeed, looking at the film thickness profiles presented in Fig. 15, we note that the $z$-averaged film thickness in the vicinity of the impact zone (the nozzle level is marked with an arrow around $y = 0.3 m$) has been altered but not significantly diminished, except the area some 15mm above the nozzle. In the Figure,
h(y) profiles are shown for two instances: \( t = 1.687 \times 10^{-1} \) (continuous line) and \( t = 1.783 \times 10^{-1} \) (black dots). Groenveld’s thickness \( h_C \) is drawn in dashed line for comparison. The profiles visible in Fig. 15 show clearly a bulge for \( y \in [0.265, 0.295] \). This formation cannot be simply associated with the jet influence, as it is present as well in the profile prior to impact; it is more likely that it results from uneven coating. At this height, the film is characterized by dimples (Snoeijer et al., 2008) – visible in Fig. 14a, and in 2D in Fig. 3 above \( y = 0.3 \) – and the coat is of three-dimensional character, being on average thicker closer to the \( z \)− edge. This is reflected in the profiles below the impact zone. As for the consequences of the impact itself, \( h(y) \) oscillates in the vicinity of \( y = 0.3 \) which is fully consistent with the instantaneous images in Fig. 14a–c and indicates alternating areas of thinning and thickening of the film. (Note that the zero-level shift in Fig. 15 is a correction for the wall thickness of 5 · 10^{-4}m.)

By representing \( p_{en}(y) \) and \( \tau_{yy} \) using gas velocity as in (14), we note that their magnitudes, for \( u_{ij} = 42 \), are one order below those for \( u_{ij} = 200 \), which in the context of (14) amounts to a higher \( h_u \). While at first sight it would seem consistent with results presented in Fig. 14 and Fig. 15, a far longer simulation would be required to establish the actual post-impact film thickness \( h_u \) (by widening the area in which a thinner film would be established) which is not the objective of this parameter study.

As mentioned, the thickness is diminished for \( y \in [0.31, 0.32] \), with the thinner area coated by two slight bulges. These formations are visible in Fig. 14 as horizontal sets of bulges above the impact zone. In our opinion, both the film thinning for \( y \approx 0.314 \) as well as the bulges are artifacts of the collision of a large horizontal vortical structure with the film. A similar effect should be observed in a longer timescale. Namely, individual, spatially distinct "craters" are probably created on the film surface by individual vortical structures - separated by distances resulting from the jet flapping frequency.

Fig. 16 presents the droplet volume distribution for \( G_{2,12}(u_{ij} = 200) \) (pink bars) and \( G_{2,12}(u_{ij} = 42) \) (yellow bars). In the Figure, minimum cell volume \((\Delta x)^3\) at the finest grid level is denoted with a black vertical line. Clearly, both simulations involve a significant number of "sub-grid" VOF "debris" – grid-cells containing non-zero fraction function values that cannot be geometrically reconstructed. This is due to the fact that, firstly, turbulent airflow contributes to droplet breakup which continues until grid resolution becomes insufficient. Secondly, the SRR technique makes this mechanism act much more often which can be seen e.g. in the r.h.s. image of Fig. 12c. Focusing our attention on the resolved droplets, we note in Fig. 16 that at low injection velocity there is about 15 resolved droplets in total (yellow bars) which is qualitatively different than at higher air velocity (red bars).

An attempt to characterize the influence of the impinging gas flow onto the internal velocities of the liquid film is presented in Fig. 17. In the Figure, we are looking at the approximated vertical component of the liquid velocity obtained using the VOF fraction function \( C \), which is equal to 1 inside the liquid. In other words the product \( C(x) u_y \) disappears in the gas phase, and Fig. 17 shows its profile in the direct neighbourhood of the impact zone \( y \in [0.28, 0.32] \) i.e. two centimeters below and above the impact zone. Two simulations are included with \( u_{ij} = 42 \) and 200 m/s. Since both flows have slightly different characteristic time scales due to higher \( We \) in the latter, we have compared instantaneous profiles at the instance corresponding to the impact zone width approx-
imately equal 0.01m (as e.g. in Fig. 14c). The curves have been averaged over a sampling window 1 mm thick. For both flows, pre-impact velocity distribution in the analysing window is similar with \( u_y \approx 1 \), which is caused by a film thinning in the analysing window slightly below the impact zone (i.e. the film is not perfectly flat even before the impact). After the impact, in case of the high-We simulation (inverted triangles in Fig. 17) one immediately observes the downward flow caused by the gas in the impact zone. Strong upward movement is visible above it. In the case of lower \( u_{inj} \), the average \( u_y \) values remain positive, suggesting the air knife wiping is far weaker for chosen injection parameters.

This concludes our investigation of the influence of the Weber number on the atomization process - we conclude that the atomization effect visible at higher We is a correct result. The simulated air-liquid system responds as expected to the decrease in dimensionless numbers, while other simulation parameters (e.g. grid resolution) are kept constant. We finish our analysis of the \( G_2 \) simulations with a brief remark on the computational efficiency. Thus, the approximated computational cost for the \( G_{2,12} \) Basilisk simulations presented e.g. in Fig. 16 was \( 4.67 \cdot 10^5 \) CPU-hours (for the simulation with \( u_{inj} = 42m/s \)) and \( 4.8 \cdot 10^5 \) CPU-hours (for the \( u_{inj} = 200m/s \) simulation).

5. Conclusions

In this paper, we have presented a novel set of simulations of a very demanding, two-phase fluid flow whose characteristics closely correspond to that of air and liquid Zinc. The boundary conditions correspond to the air knife jet-wiping process in hot-dip coating. In many aspects this is a pioneering work: to our knowledge, the only similar calculations published have described a two-dimensional case with RANS/LES performed for the airflow (Myrillas et al. 2013) or investigated the film formation only (Snoeijer et al. 2008) or, possibly, included a predefined film (i.e. not formed gravitationally). Similarly, for reasons of numerical stability (Vaudor et al., 2017), virtually all preceding attempts included a much decreased density ratio between the phases. Obviously, a multitude of practical applications of similar results exist e.g. in metallurgical and automotive industries, however they are strictly proprietary and can not be consulted by the general public.

None of these simplifications apply here: calculation accuracy for the methodology presented here is limited only by available computational resources dictating the grid resolution. Full resolution of the turbulent flow (i.e. below the Kolmogorov scale) is still too expensive. However, thanks to grid adaptivity, we are able to achieve DNS in limited areas: this claim can be further substantiated e.g. by considering the Hinze scale \( l_H \), defined as the ratio of turbulent kinetic energy and surface tension, and estimated for the industrial parameters at \( l_H \approx 1.76 \cdot 10^{-3}m \). At the grid resolution used in most cases presented here (12 levels of refinement) we obtain \( l_H/\Delta x \approx 14 \), proving that \( l_H \) is resolved. We can thus claim energy transfer from gas to liquid is at largely resolved, although of course we do not hold such claims for the turbulent flow in the gas itself.

Our results show that - as expected in metal foundry practice - the airflow inflicts a pressure gradient at the liquid layer, and “punctures” it to a degree controlled by the Re in the air, and a properly defined We. This gradient restricts the liquid feeding from reservoir, thinning the deposit. Our calculations possibly fall short of resolving the upper film thickness \( h_f \) in the full \( G_1 \) and \( G_2 \) geometries due to the lack of grid resolution. However, the \( G_2 \) case clearly displays the thinning effect, it is also observable in the \( G_1 \) case performed with decreased \( \rho_l \).

We have observed levels of atomization of the liquid metal that were not previously reported in the literature. This phenomenon, to our knowledge, has also not been observed experimentally, which leads us to believe it is a purely transient effect, taking place only as a consequence of the initial gas-liquid impact event. It is predicted for \( We \approx 38 \), while for We values closer to one, liquid wrinkling is observed. The appearance of atomization seems thus predicted correctly, instead of being induced numerically. An additional observation is that the liquid material is “milled” (atomized) by the airknife before falling back to the reservoir, and that liquid-liquid collisions are aplenty. This is already visible at the \( 2^{12} \) level (Figs. 9b and 11), and suggests that the coat-thinning mechanism is far more turbulent in its nature than known previously.

The two geometries introduced in the paper focus on the two- and three-dimensional coat formation and nozzle interactions (\( G_1 \)), coat thinning and edge effects (\( G_2 \)) and the character of gas-liquid impact. For future research we would envision working preferably with the \( G_2 \) configuration in two and three dimensions, with increasing resolutions (and simulated time-spans), preferably until full resolution of the \( h_f \) thickness is feasible.

Disclaimer

On the 9th of August 2019, that is while this paper was already in its review stages, a potentially serious bug was reported in the Basilisk solver by the user Petr Karnakov\(^7\). The bug caused a redundant multiplication of the surface tension force by the \( \sigma \) coefficient (i.e. \( \sigma^2 \) instead of \( \sigma \)), which was activated only in the areas of strongly under-resolved interface. The well-resolved areas, using the HF method with full sized stencils, were not impacted, nor the secondary backup solution using parabola fitting of height functions. Only the tertiary backup method, i.e. parabola fitting of VOF centroids introduced the additional multiplication. The bug was present in the Basilisk code repository between December 13, 2017, and August 19, 2019.

In this paper, the simulations of 2D and 3D film formation presented in Section 4.2 are not affected by the bug, as they were performed before Dec 13, 2017. However, the buggy surface tension calculation might have influenced results of the Section 4.4 in which we study the air wiping process. To verify that, we have substantially elongated the redaction process by rerunning part of the simulations, and assessing the result obtained with the bug-free code. In particular, the \( G_{2,12}(u_{inj} = 200) \) simulation presented in Fig. 12 and others, has been repeated using a bug-free version of Basilisk. The results of the repeated simulations have proven to be statistically and qualitatively identical to those already presented. We thus stand by the validity of the results presented in the paper, including the wiping process simulations discussed above.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Credit authorship contribution statement

Wojciech Aniszewski: Conceptualization, Methodology, Software, Formal analysis, Investigation, Data curation, Writing - original draft, Writing - review & editing, Visualization. Youssef Saade: Formal analysis, Investigation, Writing - review & editing, Software. Stéphane Zaleski: Supervision, Project administration, Funding acquisition, Conceptualization, Investigation, Methodology. Stéphane Popinet: Software, Supervision, Conceptualization, Methodology.

\(^7\) https://groups.google.com/d/msg/basilisk-fr/8Ub0LKQWyDc/NtBsk7p2EwAJ
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Supplementary material

Supplementary material associated with this article can be found, in the online version, at 10.1016/j.jjmultiphaseflow.2020.103399

References


8 See www.basilisk.fr/srj/view.html