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Numerical simulation of droplet impact on wettability-patterned surfaces / Russo, A; Icardi, M; Elsharkawy, M; Ceglia, D; Asinari, P; Megaridis, Cm. - In: PHYSICAL REVIEW FLUIDS. - ISSN 2469-990X. - ELETTRONICO. - 5:7(2020), pp. 1-22. [10.1103/PhysRevFluids.5.074002]

Availability: This version is available at: 11583/2842759 since: 2020-08-19T17:16:25Z

Publisher: AMER PHYSICAL SOC

Published DOI:10.1103/PhysRevFluids.5.074002

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# Numerical simulation of droplet impact on wettability-patterned surfaces

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(Received 11 December 2019; accepted 16 June 2020; published 16 July 2020)

Numerical simulations have unexplored potential in the study of droplet impact on nonuniform wettability surfaces. In this paper, we compare numerical and experimental results to investigate the application potential of a volume-of-fluid method utilized in OPENFOAM. The approach implements the Kistler model for the dynamic contact angle of impacting droplets. We begin with an investigation into the influence of the most important solver parameters to optimize the computational setup and reach the best compromise between computational cost and solution errors, as assessed in comparison to experimental results. Next, we verify the accuracy of the predictions for droplet impact on uniformly hydrophilic or superhydrophobic surfaces. Benchmarking the maximal spreading factor, contact, and spreading times, as well as contact-line behavior, we show strong agreement between the present numerical results and the models of Pasandideh-Fard et al., Phys. Fluids 8, 650 (1996) and Clanét et al., J. Fluid Mech. 517, 199 (2004). Lastly, we demonstrate the capability of the model to accurately predict outcome behaviors of droplets striking distributed-wettability surfaces, which introduce 3D outcome characteristics, even in orthogonal impact. The model successfully predicts droplet splitting and vectoring, as reported in the experiments of Schutzius et al., Sci. Rep. 4, 7029 (2014). Finally, we demonstrate a configuration wherein a droplet centrally strikes a circular disk of different wettability than its surrounding domain. The main contribution of the present paper is a numerical model capable of accurately simulating droplet impact on spatially nonuniform wettability patterns of any foreseeable design.

DOI: 10.1103/PhysRevFluids.5.074002

#### I. INTRODUCTION

Droplet impact on solid surfaces, which is frequently encountered in everyday life, has been an important target of human curiosity and scientific studies dating back hundreds of years. The complexity of the problem lies in the pursuit for deeper understanding of the underlying physical mechanisms e.g., surface tension, viscous forces, friction, and inertia) affecting the dynamics and the manner in which they are interwoven [1-3]. In addition to the impact problem producing fundamental insight into crucial physical phenomena, it is also relevant to a plethora of applications, for example, ink-jet printing [4,5], spraying [6,7], anti-icing [8], self-cleaning [9], and anticorrosion [10].

Over the past several decades, a large number of studies has been conducted on droplets striking solid surfaces. These studies have ranged from experimental to analytical or entirely computational

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efforts. Several experimental investigations have been performed to evaluate the effect of common parameters and discern some empirical relationships [11–13]. These parameters include but are not limited to the Weber number, defined as the ratio between inertial and surface tension forces  $We = \frac{\rho U_o^2 D_o}{\sigma}$ , where  $\rho$ ,  $U_o$ ,  $D_o$  are the density, impact velocity, and droplet diameter, respectively, and  $\sigma$  is the liquid surface tension; the Reynolds number, which is the ratio of inertial and viscous forces  $Re = \frac{\rho U_o D_o}{\mu}$ , where  $\mu$  is the dynamic viscosity; the Ohnesorge number, which relates viscous, inertial, and surface tension forces  $Oh = \frac{\mu}{\sqrt{\rho\sigma D_o}} = \frac{We}{\sqrt{Re}}$ ; the Eotvos Number,  $Eo = \frac{\Delta\rho g D_o^2}{\sigma}$ , which compares gravitational and surface-tension forces,  $\Delta\rho$  is the difference in density between the two phases, and g the gravitational constant; the Capillary number,  $Ca = \frac{\mu U_o}{\sigma}$ , or ratio of viscous and surface tension forces. The interaction between the solid surface and the two fluids (liquid, gas) is commonly described by the so-called contact angle  $\theta_o$ , a parameter derived from the equilibrium among the liquid, solid, and vapor interfaces.

Several analytical models have been put forth in attempts to predict droplet-impact outcome. These models generally attempted to describe the spreading and receding behavior of impacting droplets depending on the aforementioned dimensionless parameters. The majority of these models were derived by considering the energy changes of the system, with several geometric assumptions made for the shape of the spreading droplets. Some of the most successful and well-known models are those of Pasandideh-Fard *et al.* [14], Scheller and Bousfield [15], Roisman [16], and Clanét *et al.* [17].

As insightful as prior reports have been, the complexity of droplet impact cannot be fully comprehended only by experimental and analytical techniques. A need still exists for the use of high-performance computational tools in unraveling the physics of droplet impact. In fact, a large body of work has reported computational and numerical approaches of droplet impact on spatially-uniform solid surfaces. For comprehensive reviews of experimental, analytical, and computational studies in this area, the reader is referred to Yarin [18], and Josserand and Thoroddsen [3].

The vast majority of prior studies reported in the literature primarily concerned droplet impact on surfaces with spatially-uniform wettability. Little experimental or analytical work concerned droplet impact on nonuniform wettability surfaces (also known as wettability-patterned surfaces) and even less so on computational/numerical approaches addressing the matter. Nonuniform wettability surfaces are primarily sorted in two categories: multiregion wettability surfaces (MRS), and gradient-wettability surfaces (GWS). MRS [19-27] comprise solid substrates with spatial domains of distinct wettabilities placed adjacent to one another with clear boundaries. On the other hand, GWS [28–30] are surfaces with a gradient in wettability properties due, for instance, to a continuous variation in roughness. These two kinds of nonhomogeneous surfaces allow for passive control of the droplet impact dynamics and can be designed to cause specific outcomes. As important and relevant as these surfaces are, studies of droplet impact on such surfaces in the computational realm [31] are rare, with very few numerical reports [32–34] addressing such nonuniform surfaces. For instance, in a recent work, Xu et al. [35] studied numerically the droplet impact on a hydrophobic surface patterned with hydrophilic dots. A volume of fluid (VOF) method has also been adopted to study hydrophilic surfaces decorated with a hydrophobic strip in Ref. [36]. In addition, recent reports in the area of impact on nonuniform surfaces have demonstrated the limitless opportunity of complex physics that can be extracted from these types of surfaces. Li et al. [37] showed that the translational motion of an impacting droplet can be converted to gyration with a proper wettability-patterned surface. Such complex physics cannot intuitively be derived. The physical opportunities that these surfaces provide are limitless and require proper investigation. Thus, there is a need to develop a comprehensive, holistic method capable of studying these surfaces.

To fill this gap, the main objective of the present paper is to develop a numerical capability to analyze droplet impact on solid surfaces, with special attention to wettability-patterned domains. This capability is developed using OPENFOAM, an open-source computational fluid dynamics code [38–40]. In particular, the present paper comprises an investigation on the influence of the most important OPENFOAM solver parameters toward achieving the best compromise between

computational power and solution errors, as assessed in comparison to experimental results. In addition, a study of the so-called spurious currents, i.e., nonphysical phenomena often plaguing numerical simulations, is performed to limit their adverse effects on the model predictions. First, we use previously-reported models to benchmark our numerical predictions for impact on uniform surfaces. Upon demonstrating proper agreement, we expand the method to accommodate a multitude of wettability-patterned surfaces. Specifically, the model is shown to accurately reproduce two important physical phenomena: (1) droplet splitting [31,41], and (2) droplet vectoring [42,43]. Splitting consists of dividing an impacting liquid droplet into multiple volumes through a combination of hydrophilic and superhydrophobic target regions [25]. Vectoring is a directed rebound of the droplet after orthogonal impact, passively guided by the nonuniform wettability of the patterned surface [25,28]. Using available results, we show good agreement between experimental droplet impact on nonuniform wettability surfaces and the present numerical simulations. We conclude that even though the present study is confined to a limited number of wettability designs, the techniques are amenable to any conceivable wettability-patterned surface.

## **II. NUMERICAL MODEL**

#### A. Algebraic volume of fluid method

Due to the multiphase nature of droplet impact, numerical simulations require an interface tracking method, such as marker and cell [44,45], VOF [46–50], or level set [51,52], with some studies making use of finite elements applied to moving grids [53] and lattice Boltzmann methods [54,55]. In the present paper, a modified version of the VOF method, initially developed by Hirt and Nicholson [56], is employed considering its ability to handle severe interface deformations. The method, based on a fully-Eulerian formulation, features the transport equation of an indicator function called phase fraction,  $\alpha$ , together with the continuity and momentum equations. The transport equations for incompressible fluids are considered in the following forms:

$$\frac{\partial \alpha}{\partial t} + \nabla \cdot (\mathbf{U}\alpha) = 0, \tag{1}$$

$$\boldsymbol{\nabla} \cdot \mathbf{U} = \mathbf{0},\tag{2}$$

$$\frac{\partial \rho \mathbf{U}}{\partial t} + \boldsymbol{\nabla} \cdot (\rho \mathbf{U} \otimes \mathbf{U}) = -\boldsymbol{\nabla} p + \boldsymbol{\nabla} \cdot \left[ \mu (\boldsymbol{\nabla} \mathbf{U} + (\boldsymbol{\nabla} \mathbf{U})^T) - \frac{2}{3} \mu (\boldsymbol{\nabla} \cdot \mathbf{U}) \mathbf{I} \right] + \rho \mathbf{a}, \quad (3)$$

where **U** is the velocity vector shared by the fluids (different fluids occupying the same volume element are assumed to have the same velocity), p is the pressure, **I** the identity matrix, **a** the external force per unit mass, such as gravitational acceleration (source term), and  $\alpha$  within each cell of the mesh designates how much of the volume in this cell is occupied by each fluid. Consequently,  $\alpha$  is a scalar function defined inside the simulation domain and varying between zero (i.e., all vapor) and one (i.e., all liquid). With this quantity and the assumption of immiscible fluids, the numerical model considers the two fluids in each cell as a single medium with physical properties computed by an  $\alpha$ -weighted average of the constituent properties. Thus, density and viscosity of the equivalent fluid are computed by  $\rho = \rho_l \alpha + \rho_v (1 - \alpha)$  and  $\mu = \mu_l \alpha + \mu_v (1 - \alpha)$ , where the subscript *l* refers to liquid and *v* to vapor.

The OPENFOAM-based solver INTERFOAM implements a modified two-phase formulation of the VOF method. This new formulation gives rise to an additional convective term, which is due to the representation of the velocity terms as weighted averages. The full derivation of this convective term begins by solving the phase fraction equations separately,

$$\frac{\partial \alpha}{\partial t} + \nabla \cdot (\mathbf{U}_l \alpha) = 0, \tag{4}$$

$$\frac{\partial (1-\alpha)}{\partial t} + \nabla \cdot (\mathbf{U}_{v}(1-\alpha)) = 0, \tag{5}$$

where l, v signify the liquid and vapor phases, respectively. The velocity vector of the equivalent fluid is considered to be a weighted average of the two velocities,  $\mathbf{U} = \mathbf{U}_l \alpha + \mathbf{U}_v (1 - \alpha)$ . Combining these terms together yields

$$\frac{\partial \alpha}{\partial t} + \nabla \cdot (\mathbf{U}\alpha) + \nabla \cdot (\mathbf{U}_r \alpha (1 - \alpha)) = 0, \tag{6}$$

where  $\mathbf{U}_r = \mathbf{U}_l - \mathbf{U}_v$ , is the relative velocity between the two fluids.

In regard to the surface tension, its effect is considered inside the momentum equation by adding the source term  $\mathbf{f}_{\sigma}$  evaluated according to the continuum surface model introduced by Brackbill *et al.* [57]. Thus  $\mathbf{f}_{\sigma} = \sigma \kappa \nabla \alpha$ , where  $\kappa$  is the mean curvature, evaluated as  $\kappa = -\nabla \cdot \mathbf{n}$ . Far from the wall, the unit vector normal to the free interface is computed by  $\mathbf{n} = \nabla \alpha / |\nabla \alpha|$ .

Furthermore, for convenience in the setup of the boundary conditions, the hydrostatic component of the pressure is subtracted so a modified quantity,  $p_d = p - \rho \mathbf{g} \cdot \mathbf{h}$ , where p is the pressure of the droplet and **h** the depth, is adopted. Consequently, the momentum equation becomes

$$\frac{\partial \rho \mathbf{U}}{\partial t} + \nabla (\rho \mathbf{U} \otimes \mathbf{U}) = -\nabla p_d - \mathbf{g} \cdot \mathbf{h} \nabla \rho + \nabla \cdot [\mu (\nabla \mathbf{U} + (\nabla \mathbf{U})^T)] + \mathbf{f}_{\sigma}.$$
 (7)

The last relationship completes the system and allows us to solve for  $U_{r,f}$ , i.e.,  $U_r$  evaluated at each computational cell face from the following relation [58]:

$$\mathbf{U}_{\mathrm{r,f}} = \min\left[C_{\alpha} \left| \frac{\phi}{|\mathbf{S}_{f}|} \right|, \left| \frac{\phi}{|\mathbf{S}_{f}|} \right|_{\mathrm{max}} \right], \tag{8}$$

where  $\phi$  is the face flux,  $\mathbf{S}_f$  is the face area vector,  $C_{\alpha}$  a constant ( $C_{\alpha} = 1$  in this case), and  $\left|\frac{\phi}{|\mathbf{S}_f|}\right|_{\max}$  is the maximum phase flux velocity in the flow field. It is worth noticing that the compression term contributes only for  $0 < \alpha < 1$ , i.e., only at the interface, and limits the error due to interfacial smearing, enhancing the convergence of the scheme for fine meshes [58]. Moreover, in line with previous works, a value of the parameter  $C_{\alpha}$  equal to unity is adopted, since increasing or decreasing it was found to exacerbate errors in interfacial curvature and interfacial smearing (the reader is referred to Refs. [39,58] and references therein for a comprehensive discussion on this term).

# **B.** Discretization and MULES/PISO parameters

The discretization approach is the well-established finite volume method [59–61], which subdivides the flow domain into a finite number of smaller nonoverlapping control volumes that completely fill the domain. In particular, a colocated arrangement of the variables, stabilized by the Rhie and Chow velocity interpolation method [62], is implemented in OPENFOAM [63]. The discretization of Eqs. (6) and (7) requires specific schemes for the transient and spatial terms. All spatial derivative terms are converted to integrals over the cell surface, bounding the volume using Gauss's theorem [39], while an Euler implicit scheme is used for the transient terms. Following this, interpolation between neighboring cell-center quantities is required to find values at the face centers.

The advection terms in Eq. (6) require special treatment to ensure that  $\alpha$  stays between 0 and 1. This task is made by the INTERFOAM solver MULES, a flux-corrected transport technique that maintains boundedness of  $\alpha$  regardless of underlying numerical scheme and mesh structure [39,40]. A prudential approach was adopted, choosing a bounded Van Leer interpolation scheme [64,65] for the linear term in Eq. (6) and a linear scheme for the quadratic term in the same equation.

The pressure implicit with splitting of operators (PISO) algorithm [66] was adopted to handle the pressure-velocity coupling. In this approach, the equations of the system are solved one by one in a loop, seeking convergence of the solution. The PISO algorithm is widely used and detailed descriptions of its implementation can be found in Versteeg and Malalasekera [60] and Ferziger and Peric [67].

Property	Value	
Water density Air density Surface tension Water kinematic viscosity Air kinematic viscosity	$1000 \text{ kg m}^{-3}$ 1 kg m <sup>-3</sup> 0.07 N m <sup>-1</sup> 10 <sup>-6</sup> m <sup>2</sup> s <sup>-1</sup> 1.48 × 10 <sup>-5</sup> m <sup>2</sup> s <sup>-1</sup>	

TABLE I. Fluid physical properties.

# C. Boundary conditions

The most commonly used parameter to characterize the wettability of a surface by a specific liquid is the equilibrium contact angle,  $\theta_0$ , which is defined as the angle between the liquid/solid and liquid/gas interfaces at the three-phase sessile contact line. For ideal surfaces under equilibrium conditions, the value of  $\theta_0$  can be found using Young's equation [68]. However, for real (nonideal) surfaces, it has been verified that the contact angle varies over a certain range. The upper and lower limits of this range are known as the advancing contact angle ( $\theta_a$ ) and receding contact angle ( $\theta_r$ ) [69,70], respectively. In addition, it has been observed experimentally that  $\theta_a$  and  $\theta_r$  are functions of the Capillary number at the contact line [69]. For this reason, the so-called dynamic contact angle,  $\theta_d$ , has been introduced. To impose a boundary condition that yields an accurate approximation of the dynamic behavior of the contact angle, we have tested several dynamic contact angle models and adopted the one by Kistler [71]. In this model, the dependence of the generic contact angle upon the contact-line speed  $U_{cl}$ , can be evaluated through the following expressions:

$$\begin{aligned} \theta_d &= f_h \Big[ \operatorname{Ca} + f_h^{-1}(\theta_a) \Big] & \text{for } \operatorname{Ca} > 0 \\ \theta_d &= \theta_0 & \text{for } \operatorname{Ca} = 0 \\ \theta_d &= f_h \Big[ \operatorname{Ca} + f_h^{-1}(\theta_r) \Big] & \text{for } \operatorname{Ca} < 0, \end{aligned}$$
(9)

where Ca is evaluated at the contact line and  $f_h$  is Hoffman's function [72] defined by

$$f_h(x) = \cos^{-1} \left\{ 1 - 2 \tanh \left[ 5.16 \left( \frac{x}{1 + 1.31 x^{0.99}} \right)^{0.706} \right] \right\}.$$
 (10)

While the above choice of the functional form of the dynamic contact angle may not be universally accommodating, it is made here as one that produced reliable results.

The implementation of the contact angle as a boundary condition follows the approach in Brackbill *et al.* [57]. The unit vector normal to the interface n is evaluated as

$$\boldsymbol{n} = \boldsymbol{n}_w \cos\theta_d + \boldsymbol{n}_t \sin\theta_d \tag{11}$$

with  $n_w$  and  $n_t$  being the unit vectors respectively normal and tangential to the wall. The unit vector n is then used to compute the curvature and the surface tension force  $\mathbf{f}_{\sigma}$  in the cells at the boundaries, as discussed in the previous sections.

#### **III. COMPUTATIONAL AND PHYSICAL PARAMETERS**

The physical properties for water and air (room temperature and 1 atm) used in the simulations are listed in Table I. The release height necessary to reach a drop impact velocity of 1.18 ms<sup>-1</sup> is  $\sqrt{\frac{v^2}{2g}} \approx 71$  mm, implying the need for a large computational domain and a high computational cost. Thus, we explored alternative avenues to simulate an impact event. We compared the results of various techniques to select the most physically realistic method for modeling the impact.

### A. Computational domain

The computational domain is simplified by taking advantage of the symmetries (when present) of the event; such simplification is necessary due to limits in computational power. Axial-symmetric impact cases are modeled with a wedgelike shape domain with a single cell along one direction. For more complex cases that lack axisymmetry, other symmetry planes are exploited to reduce the computational grids. Preliminary tests showed that when the expanding liquid rim got close to the border of the computational domain, assuming that symmetric boundary conditions were met, the interactions with the boundary affected the results. For this reason, the domain size was chosen to be, depending on the simulated configuration, large enough to prevent any influences from the outer boundaries.

#### **B.** Initial conditions

The main parameter that affects the impact most drastically is the impact velocity, defined as the velocity of the center of mass of the droplet upon first contact with the stationary solid. However, the local state of the fluid and the interface could be significantly different even with the same impact velocity. In a real-world scenario, this state would be dependent on the motion within the droplet before impact [73]. Most of the computational studies on droplet impact have assumed an initial condition (extremely close to the surface) with a predetermined impact velocity imposed uniformly on the liquid phase. Here, instead, an attempt was made to include the droplet internal motion before impact by releasing the droplet from different heights and always with zero velocity.

To investigate the height-release effect, we first considered droplet impact with  $U_{\circ} = 1.18 \text{ ms}^{-1}$ , as in Rioboo et al. [74]. Under earth's gravity, the droplet has to be released from a height  $h \sim 71$  mm, which translates into a prohibitively large simulation domain. To avoid this issue and analyze preimpact effects, we split the simulation in two stages: a first stage with increased gravitational acceleration during the falling phase and a second stage with real gravity acting during the impact phase. To preserve the physics of the phenomenon in the modified configuration, we maintain the nondimensional numbers involving gravity, constant. Under these circumstances, we explored impact events in two alternative configurations: (a) with constant Eotvos number, and (b) with constant Weber and Capillary numbers. In the first case, both g and  $\sigma$  were multiplied by a factor K (K = 17 in this case), while in the second case g,  $\sigma$ ,  $\rho$ , and  $\mu$  were multiplied by the same factor K. The comparison between the two cases in Figs. 1(a) and 1(b) shows the presence of nonphysical deformation at the end of the falling stage in the case of constant Eo. This is due to the appearance of the so-called spurious currents, numerical artifacts that appear at the fluid interface when the discrete balance is not verified and/or the interface curvature is not computed accurately [75–77]. These currents are continuously supplied by the numerical imbalance and are bounded only by viscous dissipation. If viscous dissipation is low and surface tension is high, as in the case of constant Eo, the currents can reach severely high values, affecting the predicted shape of the droplet (see Supplemental Material [78] and Harvie et al. [79] for more details). In relation to the two configurations tested, constant We and Ca yield more physically sound results.

Subsequently, a simpler approach was tested, with the impacting droplet positioned a few millimeters above the solid with the specified impact velocity. The results were compared against the previous two-stage approach with constant We and Ca. It is a foreseeable consequence that the shear (from the surrounding air) during the fall could produce internal water circulation [73,80] that may affect the shape evolution of the droplet at impact. Comparing these two circumstances, however, as Fig. 1(c) shows, no relevant effects on the shape, impact, or spreading behavior of the impacting droplets were observed. Thus, to reduce the computational intensity, for all simulations reported below, all impact events were simulated with droplets released from a few millimetres above the solid floor with a specified and uniform impact velocity.



FIG. 1. Effects of the "spurious currents" at the end of the falling stage for droplet impact with  $D_{\circ} = 2.75 \text{ mm}$  and  $U_{\circ} = 1.18 \text{ ms}^{-1}$  in two cases: (a) when the Eotvos number is kept constant and (b) when the Weber and Capillary numbers are constant. In (c), we compare the shapes of a droplet falling at constant Weber and Capillary numbers (purple) and a droplet released from a few millimeters above the solid, i.e., without falling stage (green). There is little to no difference between the two shapes.



FIG. 2. Total error  $\varepsilon_{tot}$  (blue line) computed as the sum in time of the absolute differences between the spreading radii obtained for each mesh and the most refined one versus the total number of mesh cells for (a) wedge-shaped meshed domain (inset), and (b) full-3D meshed domain (inset).  $\varepsilon_{tot}$  is larger for the wedge-shaped domain but its order of convergence (*n*) is higher compared to the full-3D meshed domain. The black and red lines in each graph correspond to the first (*n* = 1) and second (*n* = 2) orders of convergence, respectively.

#### C. Solution convergence

To reach a compromise between solution accuracy and execution time, a preliminary study on the main solver parameter was performed and the convergence of the solution was verified by refining the adopted computational meshes. Specifically, we focused on the parameters of the MULES and PISO algorithms introduced earlier and the convergence of the pressure-velocity coupling algorithm PIMPLE, which iterates over the MULES and the PISO solvers (more details on the meaning of the parameters and the setup are provided in the Supplemental Material [81] and Ubbink [82]).

In addition, convergence of the numerical results with respect to global mesh refinement was studied. The simulation requires the transient analysis to be performed at every time step, each of the order of  $10^{-6}$  seconds, while the entire event lasts for about  $10^{-2}$  seconds. Due to the extensive resources needed to model these phenomena, a compromise had to be reached between accuracy and computational time. Figure 2 plots the total error for each mesh with a number of cells *i* as  $\varepsilon_{tot} = \sum_t |r_t^i - r_t^{i_{max}}|$ , where  $i_{max}$  corresponds to the most refined mesh, as a function of the number of mesh cells. The orders of convergence *n* observed for the wedge shape and the full-3D domain are about 1.9 and 1.4, respectively. In the following, we adopted meshes with  $\approx 10^5$  cells for the wedge-shaped domains, and between  $\approx 10^6 - 10^7$  for full-3D domains, depending on the impact surface.

# **D.** Computational resources

Because of the large number of cells required, the simulation was carried out using the openMPI implementation of Message Passing Interface. The mesh and fields are decomposed using the DECOMPOSEPAR utility, employing the Scotch decomposition, which requires no geometric input from the user and attempts to minimize the number of processor boundaries [39]. Specifically, numerical computations have been performed on a supercomputer with 17 nodes, each with 16 cores characterized by 2x AMD Opteron 6276/6376 (Bulldozer) 2.3 GHz (turbo 3.0 GHz). The jobs have been parallelized among 16 and 128 cores with processing time ranging between 10 to 300 processor hours.

#### **IV. RESULTS AND DISCUSSION**

#### A. Validation: Impact on uniform-wettability surfaces

To establish the validity of the numerical approach, simple homogeneous flat surfaces were used first to test the accuracy of the simulation results with respect to established relations found in the literature. We focused first on two types of surfaces, namely, all-hydrophilic and all-superhydrophobic. Due to the axially-symmetric nature of this problem, simulations were carried only along the x-y plane of the wedge domain configuration (x is the radial distance from the axis, y is along the symmetry axis).

Equilibrium, advancing, and receding contact angles of the surfaces are input parameters to the code and designate the wettability of each surface. In addition, droplets with different sizes but equivalent impact velocities are simulated. We maintain We values between 20 and 60 to adhere to conditions found in the literature. The main parameters reported in the literature and used to gauge our simulations are spreading contact line characteristics, maximum spreading diameter, contact time, and average receding velocity.

# 1. Impact on hydrophilic surfaces

A simple homogeneous surface was used first to test the agreement of the predicted results with relations found in the literature. Here, we focus on a uniformly-hydrophilic surface that, due to the axially symmetric nature of the problem, has been simulated by equations cast only along the x-y plane in the wedge configuration.

First, we perform a simulation of a droplet with  $D_{\circ} = 2.0$  mm, impacting with  $U_{\circ} = 0.2 \text{ ms}^{-1}$  on a surface with equilibrium, advancing and receding contact angles equal to 60°. As shown in Fig. 3(a), the simulation demonstrates the classical sequential features of impact, spreading, and receding. The predicted contact-line position as a function of time [Fig. 3(c)] was compared with both experimental and simulation data of Malgarinos *et al.* [46] for a similar impact scenario. Our numerical results are in agreement with the data in the literature. Only for the final portion of the receding phase, the simulation demonstrated a slight delay in the receding.

Simulations with different impact droplet sizes and same initial velocities were also conducted. The corresponding Weber numbers ranged between 20 and 51. We begin by comparing the numerically derived values of the spreading factor ( $\xi_{max} = \frac{D_{max}}{D_o}$ ) against those obtained by using the relations derived in Pasandideh-Fard *et al.* [14], Scheller and Bousfield [15], Roisman [16], Ukiwe and Kwok [83], and Wildeman *et al.* [84]. As seen in Fig. 4, the predictions are noticeably different from model to model. Our numerical results match quite well with the predicted  $\xi_{max}$  of Pasandideh-Fard *et al.* [14], expressed as  $\xi_{max} = \sqrt{\frac{We+12}{3(1-\cos\theta_a)+4WeRe^{-1/2}}}$ . This model not only takes into consideration We and Re, but also makes use of the advancing contact angle,  $\theta_a$  in predicting  $\xi_{max}$ . We also compared the maximum spreading time determined from the present numerical simulation to that predicted by Pasandideh-Fard *et al.* [14], namely,  $t_{sp} = \frac{8D_o}{3U_0}$ . As seen in Fig. 4(b), there is strong agreement between the two models. The present results not only confirm the accuracy and reliability of our technique, but also help reinforce the legitimacy of Pasandideh-Fard *et al.*'s model in its predictive capability of droplet impact on uniformly-hydrophilic surfaces.

#### 2. Droplet impact on superhydrophobic substrates

Next, we perform a similar set of runs to validate the numerical predictions of droplet impact on uniformly-superhydrophobic surfaces. We first considered a droplet with  $D_{\circ} = 2.1$  mm,  $U_{\circ} = 0.5 \text{ ms}^{-1}$  striking a surface with equilibrium, advancing and receding contact angles equal to 150°. As shown in Fig. 5(a), the simulation demonstrates the classical sequential behavior of impact, spreading, receding, and rebound. The predicted contact-line position as a function of time [Fig. 5(c)] was compared with the experimental data of Schutzius *et al.* [25] for a similar impact scenario. Besides a small asymmetry of the experimental droplet impact, not evident in the



FIG. 3. (a) Simulated dynamic droplet shapes with  $D_{\circ} = 2.0 \text{ mm}$ ,  $U_{\circ} = 0.2 \text{ ms}^{-1}$ , and  $\theta_0 = \theta_a = \theta_r = 60^{\circ}$  (We = 1.1 and Re = 400). The droplet shapes demonstrate the expected features of impact, namely, spreading and receding. (b) Velocity field near the contact line. (c) The simulation contact-line position (red line) versus time is compared to independently-reported experimental and simulation results under similar impact conditions. The graph shows good agreement of the data sets; the experimental and simulation data were obtained from Malgarinos *et al.* [46].



FIG. 4. Droplets impacting a hydrophilic surface with contact angles  $\theta_0 = \theta_a = \theta_r = 60^\circ$ . (a) Comparison of the numerically-predicted maximum spreading factor ( $\xi_{max}$ ) at an impact velocity  $U_o = 1.18 \text{ ms}^{-1}$  for droplets of varying initial diameter. The present simulation results (black squares) are compared to the models of Scheller and Bousfield [15] (red line), Roisman [16] (blue line), Wildeman *et al.* [84] (green line), Pasandideh-Fard *et al.* [14] (orange line), and Ukiwe and Kwok [83] (pink line). This graph shows strong agreement between the present simulation results and the model of Pasandideh-Fard *et al.* [14] and Wildeman *et al.* [84]. (b) Numerically predicted spreading times ( $t_{sp}$ ) compared to those of Pasandideh-Fard *et al.* [14].

simulation, numerical results are in accordance with the experiment. Only for the final portion of the receding phase, the simulation demonstrated a slight delay in the rebound.

Figure 5(d) plots the average droplet velocity  $U_{avr} = \frac{1}{V} \int_{V} \alpha \mathbf{U} dV$ , where V is the droplet volume and  $\alpha$  the liquid phase fraction. After the early spreading phase where a linear trend is observed,  $U_{avr} \propto t$ , the droplet velocity remains null (metastable equilibrium at maximum spreading) for about 1 ms. After this, the velocity starts increasing again in the rebound stage, exhibiting once again a linear trend  $U_{avr} \propto t$  but with a lower slope, ending with a plateau. After complete detachment from the solid substrate—where the maximum value of  $U_{avr}$  is attained—the droplet velocity starts decreasing again under the combined detracting effects of drag and gravity.

To provide further validation, we also analyzed the maximum drop spreading factor. We compared the results from the simulation [Fig. 6(a)] of varying-diameter droplets impacting with the same velocity, to the models of Pasandideh-Fard *et al.* [14] and Clanét *et al.* [17]. The predictive equation of Pasandideh-Fard *et al.* is the same as previously stated for hydrophilic surfaces, while Clanét *et al.* derived an equation of the form of  $\xi_{max} \sim We^{\frac{1}{4}}$ . As seen from Fig. 6(a), the present simulation results bare a stronger agreement with the Clanét *et al.* model than the Pasandideh-Fard *et al.* prediction. On average, there is a 13% difference between Pasandideh-Fard *et al.* This is in agreement with Fig. 2 of Clanét *et al.* [17], where a similar behavior was observed for the experimental data. The model of Clanét *et al.* was specifically derived for impact on superhydrophobic surfaces and low-viscosity fluids (such as water). Thus, it is reasonable to expect that the present simulation is in better agreement with this model.

We also compared the predicted contact time  $t_c$ , i.e., the time during which droplet and surface remained in contact, with those predicted by Richard *et al.* [85]. The contact time,  $t_c$ , is a finite number only for droplet impacts having a rebound and, for this reason, it can be evaluated only in the analysis of droplet impact on superhydrophobic surfaces. According to Richard *et al.* [85], contact time of a bouncing drop can be approximately computed as  $t_c = k_c \sqrt{\frac{\rho D_0^3}{8\sigma}}$ , where  $k_c$  is a constant. Based on this equation,  $t_c$  is independent of the impact velocity and dependent on the



FIG. 5. (a) Simulated dynamic droplet shapes with  $D_{\circ} = 2.1$  mm,  $U_{\circ} = 0.5$  ms<sup>-1</sup>, and  $\theta_0 = \theta_a = \theta_r = 150^{\circ}$  (We = 7.5 and Re = 1050). The droplet shape demonstrates the expected behaviors of impact, spreading, receding, and rebound. (b) Velocity field near the contact line. (c) The contact line of the simulated droplet impact (red line) is compared to independently-reported experimental results with similar impact conditions. Apart from a slight asymmetry for the experimental data, there is fairly good agreement between the data sets; the experimental data were taken from Schutzius *et al.* [25] (squares) and Li *et al.* [88] (triangles). (d) Predicted average droplet velocity as a function of time after initial contact. This quantity is difficult to measure experimentally but it can be evaluated with relative ease numerically.



FIG. 6. (a) Numerically predicted maximum spreading factors for droplets with same initial velocity  $U_{\circ}$  and varying initial diameter  $D_{\circ}$  striking orthogonally a superhydrophobic surface. Numerical results on average show a 13% difference from the predictions of the model of Pasandideh-Fard *et al.* [14], and an average 6% difference from the predictions of Clanét *et al.* [17], the latter derived specifically for superhydrophobic surfaces. (b) Contact time (black triangles) and average receding velocity (blue squares) of droplets impacting with constant velocity,  $U_{\circ}$ , and varying diameter  $D_{\circ}$ . In agreement with the model of Richard *et al.* [85], the dependence of the contact time on  $D_{\circ}$  is clear. Moreover, as predicted by Bartolo *et al.* [87], the average receding velocity is proportional to the initial velocity (thus, in this case, the receding velocity remained almost constant). In the Supplemental Material, we report data on maximum spreading factor, contact time, and average receding velocity for impacts with the same initial diameter  $D_{\circ}$  and varying initial velocity  $U_{\circ}$ .

impact diameter. Thus, for constant-diameter droplets impacting at varying We,  $t_c$  is expected to stay constant. In contrast, for constant impact velocity but increasing diameter droplets,  $t_c$  is expected to increase. Both sets of circumstances have been studied. Figure 6(b) plots cases where the impact velocity was held constant but the diameter was varied. We observe the expected trend, with  $k_c = 2.7 \pm 0.2$  [86].

Another comparison was conducted on the droplet retraction velocity  $U_{rec}$  during the receding phase. According to Bartolo *et al.* [87], a proportionality exists between the average receding velocity and the impact velocity. Figure 6(b) also presents  $U_{rec}$  for simulations at constant  $U_0$  and varying initial radius  $R_0$ ; as expected,  $U_{rec}$  showed an almost flat trend with We, attaining values of 20–30% of the impact velocity  $U_0$ .

Based on the above agreement between the theoretical predictions of others and the present numerically-obtained trends, it is within reason to assume validity of our numerical simulation for droplet impact on superhydrophobic surfaces.

#### B. Impact on spatially-nonuniform surfaces

After validating the numerical model for impact on uniformly superhydrophobic and hydrophilic surfaces, we studied impact on nonuniform (in terms of wettability) surfaces. These consist of combinations of hydrophilic and superhydrophobic [89] domains of similar properties to those tested above. We consider designs previously reported and demonstrating droplet translation, vectoring, and volume splitting. In addition, we explore a wettability pattern that facilitates droplet entrapment.

# 1. Droplet impact on the boundary line between hydrophilic and superhydrophobic semi-infinite regions

This nonaxisymmetric case considers a droplet impacting a straight line separating two semiinfinite regions with different wettabilities. The regions feature hydrophilic and superhydrophobic properties, respectively, and the droplet impact occurs with  $D_0 = 2.1$  mm and  $U_0 = 1.41$  ms<sup>-1</sup>.



FIG. 7. (a) Droplet impact on a wettability step. The dark-grey region (hydrophilic) has contact angles of  $\theta_0 = 89^\circ$ ,  $\theta_a = 93^\circ$ ,  $\theta_r = 0^\circ$ , while the light-grey region (superhydrophobic) has  $\theta_0 = 159^\circ$ ,  $\theta_a = 166^\circ$ ,  $\theta_r = 155^\circ$ . As observed from the impact snapshots (a) and the contact-line graph (b), each portion of the impacting droplet spreads on its respective region, independent of what occurs over the opposite side. Once full spreading has been reached, the superhydrophobic (SHP) region allows the fluid to recede, while the fluid over the hydrophilic (HL) region remains pinned. The droplet portion on the superhydrophobic region gradually recedes until it is entirely drawn over the hydrophilic region, sometimes causing secondary-droplet departure.

Figure 7(a) shows how each surface region interacts with the impacting liquid. More importantly, the maximum spreading on the hydrophilic side is equivalent to that for a uniformly-hydrophilic surface. Similarly, the maximum spreading on the superhydrophobic side is equivalent to that predicted for impact on a uniformly superhydrophobic surface. Intuitively, this outcome makes sense, even though the droplet is impacting a MRS; in this circumstance, the liquid over each region does not feel the corresponding effects of the other half region and thus the liquid spreading unravels as if each surface were on its own. After maximal lateral spreading, the superhydrophobic surface also facilitates receding. With the droplet pinned over the hydrophilic domain, the energy from the receding fluid is directed from the superhydrophobic to the hydrophilic side, and eventually brings the entire fluid into the hydrophilic region. This behavior can be seen in greater detail in the contact-line graph presented in Fig. 7(b). The images in Fig. 7(a) show that even though the droplet remains pinned as a whole, there is enough energy remaining in the receding fluid to induce a partial rebound [90].

In Fig. 8, we plot the positions of the contact line at the maximum spreading condition as function of the contact angle difference (CAD), namely, the difference between the contact angles of the two domains. For this impact, the results show that the values of the maximum spreading diameter are independent of CAD, but a shift of the droplet center toward the hydrophilic substrate is induced by the nonuniform wettability conditions.

#### 2. Droplet impact on a hydrophilic substrate with a superhydrophobic strip

Splitting droplets into prespecified volumes without external energy expenditure is an important capability that holds application possibilities in the realm of digital microfluidics. We use our numerical tools and wettability-patterned surfaces to induce volume splitting in impacting droplets. To this end, we simulate droplet impact on a surface with a superhydrophobic strip surrounded by semi-infinite hydrophilic regions. As previously reported in Schutzius *et al.* [25], two observed outcomes are possible: for wider strips, a partial droplet rebound occurs, while for narrower strips, the droplet splits into two parts, each lying entirely on the respective hydrophilic surface with no partial volume rebound.

Figures 9(a) and 9(b) show the simulated impact on the center of 3 mm and 0.2 mm-wide superhydrophobic strips, respectively. It can be seen that a partial rebound of the droplet occurs



FIG. 8. Maximum spreading of a droplet striking a wettability step with increasing contact angle difference (CAD) on either side of the step. Specifically, null CAD corresponds to the uniform-wettability case, i.e., when both sides have contact angles of  $\theta_0 = \theta_a = \theta_r = 90^\circ$ , while the nonzero CAD corresponds to hydrophilic (HL) and hydrophobic (SHP) sides with  $\theta_0 = \theta_a = \theta_r = 90^\circ - CAD/2$  and  $\theta_0 = \theta_a = \theta_r = 90^\circ + CAD/2$ , respectively.

on the wider (3 mm) strip, Fig. 9(a), but not on the 0.2 mm strip [Fig. 9(b)]. This observation is consistent with the experiments of Schutzius *et al.* [25]. The maximum spreading factor and the droplet shape during the impact are also evaluated numerically, and are in agreement with those observed experimentally. It is interesting to notice that the spreading factor evaluated for the direction parallel to the strip follows a trend typical of impacts on superhydrophobic surfaces. On the other hand, the spreading factor along the direction perpendicular to the strip, after maximum spreading, starts decreasing until it reaches a minimum value exactly at detachment of the rebounded droplet, and then it starts rising again during the relaxation phase [91]. Figure 9(c) shows that the maximum spreading radii evaluated at the superhydrophobic and the hydrophilic surfaces do not depend on the strip widths.

#### 3. Droplet impact on a superhydrophobic substrate with a hydrophilic arc

Proceeding to more complex configurations, we simulated droplet impact on a superhydrophobic surface with a thin hydrophilic circular arc, having external radius r, approximately equal to the maximum spreading droplet radius. Analogous to experiments performed by Schutzius *et al.* [25], the simulated impact occurred at We = 100. As seen in Fig. 10(a), the model predictions revealed nonorthogonal rebound, as also observed experimentally. The pinning of the liquid on the arc causes an asymmetric receding behavior, which results in a net transfer of momentum in the horizontal direction, yielding rebound in a non-purely-vertical direction. As seen in Fig. 10(b), the leftmost contact line remains pinned on the philic arc as the rightmost contact line continues to recede inward. Once the rightmost contact line traverses the arc center ( $R_{sp} = 0$ ), the droplet begins its rebound. Due to the pinning of the leftmost contact line, the receding contact line is forced to recede past the impact point ( $R_{sp} = 0$ ). This results in a prolonged contact time and increased energy losses (refer to Supplemental Material for a link to the full simulation video V3:Arc).



FIG. 9. Droplet impact (We = 60) on wettability-patterned surfaces (WPS) consisting of hydrophilic ( $\theta_0 = 89^\circ$ ,  $\theta_a = 93^\circ$  and  $\theta_r = 0^\circ$ ) areas (in dark grey) surrounding superhydrophobic ( $\theta_0 = 159^\circ$ ,  $\theta_a = 166^\circ$ ,  $\theta_r = 155^\circ$ ) strips (in light grey) of width: (a) 3 mm, and (b) 0.2 mm. The width of the superhydrophobic strip controls the ultimate shape of the impacting liquid, and residual liquid volume rebounded from the surface. The same dynamic phenomena were previously observed experimentally by Schutzius *et al.* [25]. (c) Maximum spreading factor for the superhydrophobic and hydrophilic surfaces for varying strip widths *w*.



FIG. 10. (a) Simulation of droplet impacting orthogonally with We = 100 ( $D_0 = 2.2 \text{ mm}$ ,  $U_0 = 1.83 \text{ ms}^{-1}$ ) on a superhydrophobic surface ( $\theta_0 = 159^\circ$ ,  $\theta_a = 166^\circ$ ,  $\theta_r = 155^\circ$ ) with a hydrophilic arc ( $\theta_0 = 89^\circ$ ,  $\theta_a = 93^\circ$ , and  $\theta_r = 0^\circ$ ) of external radius r = 3.29 mm and width w = 0.4 mm. As seen in the shapes in (a) and the temporal variation of the two contact-line radii on the symmetry axis in (b), the impacting droplet spreads, becomes pinned onto the hydrophilic arc (shown in dark grey), eventually recedes, and rebounds in a nonorthogonal fashion (event known as vectoring).



FIG. 11. Maximum spreading factor evaluated with the present VOF numerical model for central and orthogonal droplet impact with We  $\approx 60$  on: (a) a superhydrophobic disk in hydrophilic surroundings, and (b) a hydrophilic disk in superhydrophobic surroundings. The superhydrophobic domains have contact angles  $\theta_0 = \theta_a = \theta_r = 150^\circ$ , while the hydrophilic ones have contact angles  $\theta_0 = \theta_a = \theta_r = 60^\circ$ . The impact diameter and the impact velocity are  $D_0 = 2.1$  mm and  $U_0 = 1 \text{ ms}^{-1}$ , respectively, and the disk diameter is w. We also report the maximum spreading diameters obtained with the models proposed by Pasandideh-Fard *et al.* [14] (blue line) and by Clanét *et al.* [17] (green line).

#### 4. Droplet impact on center of disk surrounded by a different-wettability area

We also simulated orthogonal impact at the center of a circular disk of specific wettability and size, and surrounded by a semi-infinite region of different wettability. The corresponding droplet behaviors during impact under two circumstances were studied: (1) Hydrophilic disk in superhydrophobic exterior, and (2) superhydrophobic disk in hydrophilic surroundings. The contact angles are  $\theta_0 = \theta_a = \theta_r = 150^\circ$  in the superhydrophobic regions and  $\theta_0 = \theta_a = \theta_r = 60^\circ$  in the hydrophilic domains. The impact conditions were kept constant and the diameter w of the disk was varied. The numerically calculated maximum spreading factors for each case are plotted in Fig. 11.

For the case of a superhydrophobic disk in hydrophilic surroundings, Fig. 11(a) shows that for smaller disk diameters, the maximum spread is comparable to that predicted for a droplet striking a uniformly hydrophilic area. This can been seen in the graph, where the blue line is the predicted spreading factor for a droplet impacting a uniform hydrophilic region of the same wettability and impact conditions. This implies that the spreading behavior is independent of the original small superhydrophobic disk and only dependent on the outside hydrophilic area. A reason for this outcome is that the superhydrophobic area is not large enough to have a significant impact on the liquid spreading behavior. In contrast, as the diameter of the disk increases, the model results deviate from the predictions of Pasandideh-Fard *et al.* [14]. Eventually, for large enough disks, the observed maximum spreading diameter is no longer predicted by Pasandideh-Fard *et al.* 's model, but rather the model of Clanét *et al.* [17] for a uniformly-superhydrophobic surface (green line). In the latter case, the disk is large enough to accommodate the entire spreading range, which now occurs entirely within the superhydrophobic area.

Next, we studied central impact on a hydrophilic disk in superhydrophobic surroundings. As seen from Fig. 11(b), for smaller disks, the observed spreading factor can be predicted using the Clanét model for a superhydrophobic surface. This observation can be explained using the same logic presented above; the area of the hydrophilic disk is small enough so it does not exert significant influence on the droplet spreading behavior when compared to the surrounding superhydrophobic region. As the disk diameter increases, an interesting situation occurs wherein the observed maximum spreading is equivalent to the disk diameter [linear portion of Fig. 11(b)]. In other words, for such conditions, a spreading droplet becomes entrapped within the hydrophilic disk and is unable to spread farther onto the superhydropobic surroundings. This physical entrapment is

interesting and intuitive. The outside superhydrophobic region acts as an energy barrier, preventing the spreading droplet from transitioning into it. This phenomenon should be observed not only for hydrophilic/superhydrophobic contrasts but also for other situations where the surrounding region is less wettable than the inner disk. As the diameter of the disk increases, the observed spreading factors begin to fall in with the predictions of Pasandideh-Fard *et al.*'s model for impact on a uniformly-hydrophilic surface. At this point, the diameter of the disk has become so large that the entire spreading is confined within the hydrophilic region.

# **V. CONCLUSIONS**

The realm of wettability-patterned surfaces is vast, with nearly limitless design combinations of different shapes, surface energies, and textures. Relying only on experimental investigations for testing such surfaces would be infeasible and expensive. In this paper, we have presented a numerical method capable of simulating droplet impact on such spatially-inhomogeneous planar surfaces. We began by benchmarking our numerical simulations against the commonly-accepted models of Pasandideh-Fard et al. [14] and Clanét et al. [17], and found strong agreement in select problems where the solid surface wettability is spatially uniform. Next, we examined droplet impact on wettability-patterned surfaces capable of producing droplet translation, splitting, and vectoring. In the case of translation, we studied a droplet impacting orthogonally a wettability step and moving laterally until equilibrating into the region of higher surface energy. For splitting, we showed how the width of a hydrophobic strip surrounded by hydrophilic areas dictates the size of the liquid-volume split and the residual volume that rebounds. Lastly, we simulated the nonorthogonal rebound (vectoring) of droplets impacting orthogonally on a superhydrophobic domain with a hydrophilic circular arc, which serves as an anchoring site. Our simulation results agreed with the data reported by Schutzius et al. [25], where the same patterns were experimentally tested and analyzed. Furthermore, we simulated cases where spreading droplets that impacted orthogonally a disk of specific wettability became entrapped by the surrounding region that had lower wettability. Although the present model has been vetted with a limited number of wettability designs, the techniques presented here are amenable to any foreseeable wettability-patterned surface. Moreover, following the recent experimental work by Jambon-Puillet et al. [92], the presented methodology can be extended to study numerically the deformation of elastic capsules filled with different liquids impacting rigid walls.

# ACKNOWLEDGMENTS

Computational resources were mainly provided by HPC@Polito, a project of academic computing within the Department of Control and Computer Engineering at the Politecnico di Torino (http://www.hpc.polito.it). M.I. and D.C. gratefully acknowledge the support of Raúl Tempone, the KAUST SRI-UQ center, and the KAUST HPC and Extreme Computing center. M.E. was supported by a National Science Foundation Graduate Research Fellowship under Grant No. 0907994.

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