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Denis Aubry, Hawa Thiam, Matthieu Piel, Rachele Allena. A computational mechanics approach to assess the link between cell morphology and forces during confined migration. Biomechanics and Modeling in Mechanobiology, Springer Verlag, 2015, 14 (1), pp.143-157. <10.1007/s10237-014-0595-3>. <hal-01083871>

## HAL Id: hal-01083871 https://hal.archives-ouvertes.fr/hal-01083871

Submitted on 18 Nov 2014

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Denis AUBRY, Hawa THIAM, Matthieu PIEL, Rachele ALLENA - A computational mechanics approach to assess the link between cell morphology and forces during confined migration - Biomechanics and Modeling in Mechanobiology p.nn - 2014

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## A computational mechanics approach to assess the link between cell morphology and forces during confined migration

D. Aubry · H. Thiam · M. Piel · R. Allena

Abstract Confined migration plays a fundamental role during several biological phenomena such as embryogenesis, immunity and tumorogenesis. Here, we propose a twodimensional mechanical model to simulate the migration of a HeLa cell through a micro-channel. As in our previous works, the cell is modelled as a continuum and a standard Maxwell model is used to describe the mechanical behaviour of both the cytoplasm (including active strains) and the nucleus. The cell cyclically protrudes and contracts and develops viscous forces to adhere to the substrate. The micro-channel is represented by two rigid walls, and it exerts an additional viscous force on the cell boundaries. We test four channels whose dimensions in terms of width are i) larger than the cell diameter, ii) sub-cellular, ii) sub-nuclear and iv) much smaller than the nucleus diameter. The main objective of the work is to assess the necessary conditions for the cell to enter into the channel and migrate through it. Therefore, we evaluate both the evolution of the cell morphology and the cell-channel and cell-substrate surface forces, and we show that there exists a link between the two, which is the essential parameter deter-

**Electronic supplementary material** The online version of this article (doi:10.1007/s10237-014-0595-3) contains supplementary material, which is available to authorized users.

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R. Allena (⊠) Arts et Metiers ParisTech, LBM, 151 Bd de l'hôpital, 75013 Paris, France e-mail: Rachele.allena@ensam.eu mining whether the cell is permeative, invasive or penetrating.

**Keywords** Confined cell migration · Continuum mechanics · Computational mechanics · Forces

#### **1** Introduction

In our previous works (Allena and Aubry 2012; Allena 2013), we have presented numerical models which helped to understand the mechanisms controlling cell motility on twodimensional (2D) flat surfaces. Nevertheless, during many biological processes such as embryogenesis, immunity and tumorogenesis, cell migration takes place in confined environments of tissues (Friedl and Wolf 2010). In these cases, cell locomotion is influenced by the presence of attractant molecules, but also by the morphology of the extracellular matrix (ECM). In fact, the surrounding tissues may vary in terms of heterogeneity, fibres density and organization. As shown both experimentally (Erler and Weaver 2009; Wolf et al. 2009; Egeblad et al. 2010; Friedl and Wolf 2010) and theoretically (Zaman et al. 2005, 2006, 2007; Scianna et al. 2013), the width of the ECM pores, the degree of ECM alignment as well as the ECM stiffness are fundamental parameters, which determine how and how much the ECM steers or inhibits the cell movement. Therefore, the cell needs to continuously adapt its shape and consequently its migratory behaviour. In tumorogenesis for instance, cancer cells develop an invasive behaviour, which allows them to enter and progressively invade healthy tissue as they are constantly exposed to biomechanical and biophysical stimuli. Such adaptation requires an internal reorganization of both the cytoskeleton and the embedded organelles, among which the nucleus is the stiffest and the most voluminous. Consequently, it has become essential to quantitatively assess the cell ability to deform as well as which mechanical forces the cell has to develop in order to move forward within a confined micro-structure.

In the last few years, several experimental studies have tried to provide such data. Systems like collagen gels or lattices are commonly used to simulate cell migration in confined connective tissues (Wolf et al. 2009). Although very simplified, such systems are highly complex and difficult to control since many physical parameters (i.e. gel density and elasticity, local constrictions) may affect the global mobility of the cell and furthermore fail to reproduce spatial tracks or obstacles (Provenzano et al. 2008; Wolf et al. 2009; Egeblad et al. 2010). More recently, it has been possible to better control, vary and tune the geometrical characteristics of the patterned micro-structure using micro-laser techniques (Ilina et al. 2011) or photolithography (Heuzé et al. 2011). In the latter work, cells migrate through straight micro-channels made of silicone rubber (i.e. polydimethylsiloxane, PDMS), whose sub-cellular dimensions vary between 2 and  $10 \mu m$ in width and highly depend on cell type. Such an approach has provided interesting results for cancer cells (Irimia and Toner 2009; Ronot et al. 2000), immune cells (Irimia et al. 2007; Faure-André et al. 2008) and neurons (Taylor et al. 2005). Micro-channels may be modulated in order to investigate specific biological problems such as trans-migration ability within a well-defined geometry or the influence of the substrate stiffness by letting channel material vary. Additionally, more complex geometries can be obtained to force the cell to take turns and explore its 2D confined environment.

From a numerical point of view, many models have been proposed to simulate single cell migration on 2D flat surfaces or in three-dimensional (3D) environment (Rangarajan and Zaman 2008). Such models have used different approaches resulting in force-based dynamics models (Zaman et al. 2005, 2006), stochastic models to simulate persistent random walks (Tranquillo and Lauffenburger 1987; Tranquillo et al. 1988; Stokes et al. 1991; Stokes and Lauffenburger 1991), models reproducing the movement of cancer cell spheroids (McElwain and Ponzo 1977; McElwain 1978; McElwain et al. 1979), Monte Carlo models (Zaman et al. 2007; Scianna and Preziosi 2013; Scianna et al. 2013) or purely mechanical models (Allena and Aubry 2012; Allena 2013). Active gel layers submitted to external forces have been used to represent acto-myosin cells migrating in a free (Recho and Truskinovsky 2013; Recho et al. 2013) or confined (Hawkins et al. 2009; Hawkins and Voituriez 2010) environment. Scianna and Preziosi (Scianna and Preziosi 2013) have presented a cellular potts model (CPM), which reproduces an experimental assay very similar to those used in (Taylor et al. 2005; Irimia et al. 2007; Faure-André et al. 2008; Irimia and Toner 2009; Rolli et al. 2010; Heuzé et al. 2011). In this model, the cell is modelled as a discrete physical unit, including the cytosol and the nucleus, while channels of different

widths constitute the migration chamber. The authors have investigated the invasiveness of tumour cells by evaluating their displacement and velocity as well as their deformability, which seems to be strongly linked to the deformability of the nucleus. In (Tozluoğlu et al. 2013), a hybrid agent-based finite element model is proposed to evaluate the migration strategy of the cell in different environments such as confinement. The model is able to simulate both the protrusion-contraction and the membrane blebbing modes of migration. Therefore, the authors estimate the effects of the ECM geometry on the relationship between cell velocity, contractility and adhesion, and they also find interesting effects of membrane blebbing on cell velocity and morphology. Finally, in (Giverso et al.), an energetic continuum approach is employed to investigate the necessary condition for which a cell migrate through a cylindrical channel. They consider the nucleus either as an elastic membrane surrounding a liquid droplet or as an incompressible elastic material. By taking into account and balancing different forces exerted by and on the cell during confined movement, they are able to determine the minimal size of the cylindrical structure and they observe that cell ability to migrate through it depends on both nucleus stiffness and adhesion to ECM.

#### 1.1 Objective of the present work

In the present paper, a finite element model that reproduces the experimental set-up used in (Heuzé et al. 2011) for HeLa cells is proposed, which is based on the following hypotheses:

- the 2D geometry represents a top view of the global structure, and a plane stress hypothesis has been made;
- as in (Giverso et al.), a purely mechanical approach is used to describe the cell behaviour. However, a different mathematical method is applied. In fact, the decomposition of the deformation gradient is employed to consider both the active (i.e. protrusion and contraction) and the elastic (i.e. strains generated by the interaction with the environment) strains undergone by the cell;
- contrary to previous works (Scianna and Preziosi 2013; Scianna et al. 2013; Tozluoğlu et al. 2013), the cell is modelled as a continuum. Nonetheless, both the cytoplasm and the nucleus have been originally represented through two characteristic functions, and a standard Maxwell model has been used to describe their viscoelastic behaviour (Allena and Aubry 2012; Allena 2013; Tozluoğlu et al. 2013);
- the cell is able to cyclically protrude and contract. Such active strains are triggered respectively by the polymerization and depolymerization of the actin filaments and are synchronized with the viscous adhesion forces between the cell and the substrate (Allena and Aubry 2012; Allena 2013);





 the micro-channel is represented by two rigid walls, which are described by two characteristic functions, and exerts an additional normal viscous force on the cell boundaries when contact condition is fulfilled.

The main objective of our work is to assess the necessary conditions for the cell to enter into the channel and migrate through it. In order to do so, we test four different channels whose dimensions in width are i) larger than the cell diameter, ii) sub-cellular, iii) sub-nuclear and iv) much smaller than the nucleus diameter. We analyse the evolution of the cell morphology by consistently comparing it with experimental observations, and we classify the cell behaviour according to the covered distance inside the channel. Additionally, we evaluate both the cell-substrate and the cell-channel surface forces during migration, and we find that there exists a link between such forces and the changes in cell shape, which may be essential in determining the invasive behaviour of the cell.

The paper is organized as follows: in Sect. 2, the geometry of the cell, the constitutive model and the mechanical approach are described. In Sect. 3, the results of the numerical simulations are presented. First, we analyse the cell behaviour (Sect. 3.1). Second, we evaluate the mechanical cellsubstrate and cell-channel surface forces (Sect. 3.2), and we find the necessary conditions determining whether the cell is penetrating, invasive or permeative.

#### 2 The model

In this section, we provide the general framework of the model. First, we describe the geometry of the cell including the cytoplasm and the nucleus. Second, we focus on the mechanics of the system. Specifically, we introduce the standard Maxwell models, which are used to reproduce the viscoelastic behaviour of the nucleus and the cytoplasm. Third, the intra-synchronization is presented. As in our previous works (Allena and Aubry 2012; Allena 2013), this represents the key ingredient of the cell movement. In fact, we show how the cyclic active strains (i.e. protrusion and contraction) are strongly coupled with the viscous forces generated by the cell to adhere to the substrate and necessary to efficiently move forward. Finally, we describe the geometry of the micro-channel and the associated viscous force exerted on the cell boundaries, which allows the cell to squeeze and pass (or not) through it.

#### 2.1 Cell geometry

HeLa cells are human cells with a rather rounded initial shape and a diameter of about 15  $\mu$ m (Ronot et al. 2000; Ngalim et al. 2013). For the numerical model, the geometry of the cell has been simplified by a circular domain  $\Omega_{cell}$  of radius  $r_{cell}$  (Fig. 1a). Here, we consider two main components of the cell: the cytoplasm ( $\Omega_{cytoplasm}$ ) and the nucleus ( $\Omega_{nucleus}$ ) (Fig. 1a, Sect. 5.1). Additionally, the cell cyclically generates a frontal ( $\Omega_f$ ) and a rear ( $\Omega_r$ ) adhesion region in order to move forward (Allena and Aubry 2012; Allena 2013) (Fig. 1b, Sect. 5.1).

#### 2.2 Constitutive model and mechanics of the cell

Both the nucleus and the cytoplasm are assumed to be viscoelastic materials, and their behaviour is described by two standard Maxwell models (Larson 1998) (Sect. 5.2). On one hand, the nucleus is composed by the nuclear lamina  $\Omega_{\text{lamina}}$ (the solid phase, Sects. 5.1 and 5.2), which surrounds the viscoelastic nucleoplasm  $\Omega_{nucleoplasm}$  (the fluid phase, Sects. 5.1 and 5.2). On the other hand, the cytoplasm is essentially made of a solid phase represented by the cell cortex  $\Omega_{cortex}$ (Sects. 5.1 and 5.3) and a fluid-like phase, the cytosol  $\Omega_{cytosol}$ (Sects. 5.1 and 5.3) in which the organelles such as the actin filaments are embedded. As in our previous works (Allena and Aubry 2012; Allena 2013), we assume that the polymerization of the actin filaments inside the cytosol, which mostly occurs at the front of the cell (Schaub et al. 2007), generates the protrusive force at the leading edge, and their contraction due to binding of myosin generates the contractile stress at the rear of the cell (Mogilner 2009). Such active strains triggering the deformability of the cell are then described through a deformation tensor  $F_{\text{cytosol},a}$  (Sects. 2.3 and 5.3) in the fluid-like branch of the symbolic standard Maxwell model of the cytoplasm.

As described in (Allena and Aubry 2012; Allena 2013), the global equilibrium of the system is expressed as

$$\rho a = Div_p \left( J \sigma F^{-T} \right) + f_{adh} + f_{channel}$$
(1)

where  $\rho$  is the cell density, *a* is the acceleration,  $Div_p$  is the divergence with respect to the initial position *p*, *J* is the determinant of the deformation gradient *F* and  $A^{-T}$  denotes the inverse transpose of the matrix *A* (Holzapfel 2000; Taber 2004).  $f_{adh}$  and  $f_{channel}$  indicate, respectively, the viscous adhesion forces between the cell and the substrate (Sect. 2.3) and the viscous force exerted by the channel on the cell boundaries (Sect. 2.4). Here, all the body forces but the inertial effects are neglected (Gracheva and Othmer 2004; Allena and Aubry 2012; Allena 2013).

#### 2.3 Intra-synchronization

To describe the oscillating movement of the cell, two main assumptions have been made:

- the active strains of protrusion and contraction are only applied in the cytosol. In fact, as in our previous works (Allena and Aubry 2012; Allena 2013), we assume that the oscillatory movement of the cell is triggered by the periodic polymerization and depolymerization of the actin filaments, which are embedded in the cytosol. The former only occurs at the front of the cell, while the latter takes place from the front towards the rear of the cell. Therefore, although the nucleus does not undergo any active strain, it will interact with the surrounding cytosol apart from the protrusion phase (Friedl et al. 2011);
- although the cell may form multiple pseudopodia (Allena 2013), here only one is generated in the direction of

migration, which, to reproduce the experimental set-up where the cell is constrained into a micro-channel (Heuzé et al. 2011), corresponds to the horizontal axis  $i_x$ .

Therefore, the solid active deformation tensor  $F_{\text{cytosol},a}$  reads

$$\mathbf{F}_{\text{cytosol},a} = \begin{cases} e_{a0} \sin\left(2 \pi \frac{t}{T}\right) h_{\text{cytosol},\text{front}} \mathbf{i}_{x} \otimes \mathbf{i}_{x} & \text{if } \sin\left(2 \pi \frac{t}{T}\right) > 0\\ \frac{e_{a0}}{2} \sin\left(2 \pi \frac{t}{T}\right) h_{\text{cytosol}} \mathbf{i}_{x} \otimes \mathbf{i}_{x} & \text{if } \sin\left(2 \pi \frac{t}{T}\right) < 0 \end{cases}$$
(2)

where  $e_{a0}$  is the amplitude of the active strain, *t* is time, *T* is the migration period,  $h_{\text{cytosol}}$  and  $h_{\text{cytosol,front}}$  are two characteristic functions (Sect. 5.1) and  $\otimes$  indicates the tensorial product.

As shown in (Allena and Aubry 2012), in order to be able to effectively migrate, the cell must adhere on the substrate otherwise it would only deform on place. Thus, an intra-synchronization is required which coordinates the cyclic protrusion–contraction deformations with the adhesion forces  $f_{adh}$  (Eq. 1) generated between the cell frontal and rear adhesion surfaces and the underneath substrate. As in previous works (Phillipson et al. 2006; Sakamoto et al. 2011; Allena and Aubry 2012; Allena 2013), such forces are assumed to be viscous and may be distinguished into a frontal  $(f_{adh, f})$  and a rear  $(f_{adh, r})$  force as follows

$$f_{adh,f}(\mathbf{n}_{cell}) = -\mu_{adh}h_{sync} \left(-\frac{\partial e_a}{\partial t} \mathbf{v} \quad \text{on}\Omega_f\right)$$
$$f_{adh,r}(\mathbf{n}_{cell}) = -\mu_{adh}h_{sync} \left(\frac{\partial e_a}{\partial t} \mathbf{v} \quad \text{on}\Omega_r\right)$$
(3)

with  $n_{cell}$  the outward normal to the cell boundary,  $\mu_{adh}$  the friction coefficient and v the velocity. The characteristic function  $h_{sync}$  is the key ingredient of the preceding equations since it couples the adhesion forces with the active strains, which results in the intra-synchronization mentioned above. Thus, we observe two main phases during the migratory movement of the cell: i) the protrusion and the adhesion at the rear edge; ii) the contraction and the adhesion at the frontal edge.

#### 2.4 Micro-channel

Here, we want to reproduce the micro-channel-based assay presented in (Heuzé et al. 2011). Thus, the micro-channel domain  $\Omega_{\text{channel}}$  is represented by two pseudo-elliptical rigid walls with no top roof (Sect. 5.4).

When the cell enters into the micro-channel, it is then submitted to a viscous force  $f_{\text{channel}}$  (Eq. 1), which can be distinguished into an upper ( $f_{\text{channel},uw_i}$ ) and a lower ( $f_{\text{channel},lw_i}$ ) force as follows  $f_{\text{channel},uw_{i}}(\boldsymbol{n}_{uw,i})$   $= -\mu_{\text{channel}} \frac{1}{(l_{uw,i}+1)^{8} + \alpha} \left( \frac{\partial \boldsymbol{u}}{\partial t}, \boldsymbol{n}_{uw,i} \quad n_{uw,i} \quad \text{on}\partial\Omega_{uw,i} \right)$   $f_{\text{channel},lw_{i}}(\boldsymbol{n}_{lw,i})$   $= -\mu_{\text{channel}} \frac{1}{(l_{lw,i}+1)^{8} + \alpha} \left( \frac{\partial \boldsymbol{u}}{\partial t}, \boldsymbol{n}_{lw,i} \quad \boldsymbol{n}_{lw,i} \quad \text{on}\partial\Omega_{lw,i} \right)$ (4)

where  $\mu_{channel}$  is the viscosity of the micro-channel,  $l_{uw,i}$  and  $l_{lw,i}$  are two level set functions (Sect. 5.4),  $\alpha$  is a constant and  $n_{uw,i}$  and  $n_{lw,i}$  are the outward normal to the boundaries  $\partial \Omega_{uw,i}$  and  $\partial \Omega_{lw,i}$  of the upper and lower wall, respectively, which are here originally calculated (Sect. 5.4) (the subscript 'i' indicates the channel number as explained in Sect. 3.1 and 5.4). Finally, (a,b) defines the scalar product between two vectors.

#### **3 Results**

The numerical simulations have been run using the finite element software COMSOL Multiphysics<sup>®</sup> 3.5a. As described in Sect. 2.2, the viscoelastic behaviour of the cell has been taken into account. The components of the cytoplasm and the nucleus have been implicitly described by specific characteristic functions (Sect. 5.1) in order to be able to define the parameters of the standard Maxwell models. The radius r<sub>cortex</sub>, r<sub>cytosol</sub>, r<sub>lamina</sub> and r<sub>nucleoplasm</sub> of the HeLa cell have been fixed to 7.5, 7.25, 4.5 and 4.4 µm, respectively. Then, the cell cortex and the nuclear lamina have a thickness  $t_{cortex}$ and  $t_{\text{lamina}}$  of 0.25  $\mu$ m (Pesen and Hoh 2005; Tinevez et al. 2009; Jiang and Sun 2013) and  $0.1 \,\mu$ m (Righolt et al. 2010), respectively. The nominal values of the Young moduli  $E_{\text{cortex},0}$  of the cell cortex and  $E_{\text{cytosol},0}$  of the cytosol have been chosen equal to 100 and 10 Pa (Crick and Hughes 1950). For the nucleus, assuming that its stiffness is mostly provided by the nuclear lamina, we have set  $E_{\text{lamina},0}$  and  $E_{\text{nucleoplasm},0}$  to 3,000 Pa (Caille et al. 2002; Dahl et al. 2008) and 25 Pa (Vaziri et al. 2006), respectively. According to a simple spatial homogenization approach (Christensen 1991; Larson 1998), such moduli have then been recalculated according to the surface occupied by each component in the cell to obtain  $E_{\text{cortex}}$ ,  $E_{\text{cytosol}}$ ,  $E_{\text{lamina}}$  and  $E_{\text{nucleoplasm}}$ (Table 1). Since we consider here that the cell cortex and the nuclear lamina are rather elastic, while the cytosol and the nucleoplasm are rather viscoelastic, the Poisson's ratios  $v_{\text{cortex}}$  and  $v_{\text{lamina}}$  have been set to 0.3, while  $v_{\text{cytosol}}$  and  $\nu_{\text{nucleoplasm}}$  to 0.4. The viscosities  $\mu_{\text{cvtosol}}$  and  $\mu_{\text{nucleoplasm}}$  are equal to  $3 \times 10^5$  Pa-s (Bausch et al. 1999; Drury and Dembo 2001). The cell density  $\rho$  has been set to 1,000 kg/m<sup>3</sup> (Fukui et al. 2000), and the viscous friction coefficient  $\mu_{adh}$  is equal  $10^{8}$ Pa-s/m. Finally, the intensity of the active strain  $e_{a0}$  and the migration period T have been chosen equal to 0.2 and 600 s, respectively, in order to obtain an average migration velocity of the order of magnitude of the one experimentally observed for HeLa cells (Ronot et al. 2000; Ngalim et al. 2013).

All the parameters of the model have been reported in Table 1.

#### 3.1 Cell behaviour and morphology

As described in Sect. 2.4, the channel is represented by two pseudo-elliptical walls  $(l_{uw,i} \text{ and } l_{lw,i})$ , whose semi-axes *a* and *b* are 30 and 2 µm long, respectively.

For the simulations, only two-thirds of the total length of the channel are considered, which corresponds to  $40 \,\mu$ m.

By letting the position of the upper and lower walls centres  $c_{uw,i}$  and  $c_{lw,i}$  vary, we have tested four channels with different width as follows:

- *channel 16* has a width  $W_{c,1}$  of 16 µm, which is larger than the cell diameter with  $c_{uw,16}$  (42.5, 10 µm) and  $c_{lw,16}$  (42.5 µm, -10 µm);
- *channel 12* has an intermediate width  $W_{c,2}$  of 12 µm, which is smaller than the cell diameter and bigger than the nucleus diameter, with  $c_{uw,12}$  (42.5, 8 µm) and  $c_{lw,12}$  (42.5, -8 µm);
- *channel* 7 has a width  $W_{c,3}$  of 7µm, which is slightly smaller than the nucleus diameter with  $c_{uw,7}$  (42.5, 5.5µm) and  $c_{lw,7}$  (42.5, -5.5µm);
- *channel 4* has a width  $W_{c,4}$  of 4µm, which is much smaller than the nucleus diameter with  $c_{uw,4}$  (42.5, 4µm) and  $c_{lw,4}$  (42, -4µm).

For the first set of simulations, the viscous friction coefficient  $\mu_{\text{channel}}$  and the constant  $\alpha$  have been set equal to  $10^{10}$  Pa-s/m and 0.1, respectively.

We have studied the cell behaviour for each of the previous configurations by analysing specific aspects of the confined movement, and the main results are listed in Table 2.

First, we have evaluated the efficiency of the migration in terms of covered distance. In Fig. 3, the total displacement of the frontal edge of the cell is reported for the four simulations. Then, as previously proposed by (Rolli et al. 2010; Scianna et al. 2013), we can classify the cell as permeative, invasive or penetrating. The permeative behaviour is observable for *channel 16* and *channel 12* (Fig. 2a, b) where the cell reaches the other side of the channel by covering a distance of 38  $\mu$ m in 9,000 s (blue and red lines in Fig. 3, and Movie 1 and Movie 2, respectively). The invasive behaviour occurs when the cell enters into the channel, but it is not able to achieve the other side (Fig. 2c). This is the case of *channel 7* where the cell only migrates over 25  $\mu$ m in 6,000 s (green line in Fig. 3 and Movie 3). Finally, the cell is penetrating (Fig. 2d)

Table 1Main geometrical andmaterial parameters of themodel

Parameter	Description	Value (unit)	References
r <sub>cell</sub>	Cell radius	7.5 µm	
r <sub>cortex</sub>	Cortex radius	7.5 µm	
r <sub>cytosol</sub>	Cytosol radius	$7.25\mu m$	
rlamina	Lamina radius	4.5 µm	
rnucleoplasm	Nucleoplasm radius	4.4 µm	
t <sub>cortex</sub>	Cortex thickness	$0.25\mu m$	Posen and Hob (2005)
			Tinevez et al. (2009), Jiang and Sun (2013)
t <sub>lamina</sub>	Lamina thickness	0.1 µm	Righolt et al. (2010)
$l_f$	Distance cell centre— boundary of frontal adhesion region	4µm	
lr	Distance cell centre— boundary of rear adhesion region	4μm	
$\Omega_{cell}$	Initial cell area	$176.6\mu m^2$	
$\Omega_{\rm cortex}$	Initial cortex area	$11.6\mu m^2$	
$\Omega_{\rm cytosol}$	Initial cytosol area	$101.4\mu m^2$	
$\Omega_{\rm cytoplasm}$	Initial cytoplasm area	$113\mu m^2$	
$\Omega_{\text{lamina}}$	Initial lamina area	$2.8\mu m^2$	
$\Omega_{nucleoplasm}$	Initial nucleoplasm area	$60.8\mu m^2$	
$\Omega_{nucleus}$	Initial nucleus area	$63.6\mu m^2$	
$\Omega_f$	Initial frontal adhesion region area	$31 \mu m^2$	
$\Omega_r$	Initial rear adhesion region area	$31 \mu m^2$	
$E_{\rm cortex,0}$	Nominal cortex Young modulus	100 Pa	
$E_{\rm cytosol,0}$	Nominal cytosol Young modulus	10 Pa	Crick and Hughes (1950)
$E_{\text{lamina},0}$	Nominal lamina Young modulus	3,000 Pa	Caille et al. (2002),
Enucleoplasm 0	Nominal nucleoplasm Young modulus	25 Pa	Vaziri et al. (2008)
E <sub>cortex</sub>	Equivalent cortex Young modulus	15 Pa	
Ecvtosol	Equivalent cytosol Young modulus	8 Pa	
$E_{\text{lamina}}$	Equivalent lamina Young modulus	196 Pa	
Enucleoplasm	Equivalent nucleoplasm Young modulus	23 Pa	
V <sub>cortex</sub>	Cortex Poisson ratio	0.3	
Vcytosol	Cytosol Poisson ratio	0.4	
Vlamina	Lamina Poisson ratio	0.3	
Vnucleoplasm	Nucleoplasm Poisson ratio	0.4	
$\mu_{ m cytosol}$	Cytosol viscosity	$3 \times 10^5$ Pa-s	<b>D</b> 1 (1000)
		a 405 b	Drury and Dembo (2001)
$\mu_{ m nucleoplasm}$	Nucleoplasm viscosity	$3 \times 10^{9}$ Pa-s	Bausch et al. (1999), Drury and Dembo (2001)
ρ	Cell density	$1,000  \text{kg/m}^3$	Fukui et al. (2000)
$e_{a0}$	Amplitude of the active strain	0.8	
Т	Migration period	600 s	
$\mu_{ m adh}$	Cell friction coefficient	10 <sup>8</sup> Pa-s/m	
a		20	

#### Table 1 continued

Parameter	Description	Value (unit)	References
b	Semi-axis of the pseudo-elliptical walls	2 µ m	
<i>x</i> <sub>0</sub>	x-coordinate of the pseudo-elliptical walls centre	42.5 µm	
Yuw0,i	y-coordinate of the upper pseudo-elliptical wall centre	$y_{uw0,1}$ : 10 $y_{uw0,2}$ : 8 $y_{uw0,3}$ : 6 $y_{uw0,4}$ : 4 $\mu$ m	
Ylw0,i	y-coordinate of the lower pseudo-elliptical wall centre	$y_{lw0,1}$ : -10 $y_{lw0,2}$ : -8 $y_{lw0,3}$ : -6 $y_{lw0,4}$ : -4 $\mu$ m	
$\mu_{\text{channel}}$	Channel viscous friction coefficient	10 <sup>10</sup> Pa-s/m	
α		0.1	
$W_{c,16}$	Width of channel 1	16 µm	
$W_{c,12}$	Width of channel 3	12 µm	
$W_{c,7}$	Width of channel 3	8 µ m	
$W_{c,4}$	Width of channel 4	4 µ m	

## Table 2 Main numerical results

	Channel 16	Channel 12	Channel 7	Channel 4
Displacement (µm)	38	38	25	7.5
Protrusion average velocity (µm/s)	0.0055	0.0051	0.0055	0.0053
Contraction average velocity (µm/s)	0.0102	0.0122	0.0118	0.0115
t <sub>contact</sub> (s)	_	1,950	1,250	1,220
t <sub>penetration</sub> (s)	3,900	4,600	4,610	-
T <sub>entry</sub> (s)	_	2,650	3,360	-
Maximal ratio cell area/nucleus area	3.29	2.89	2.25	3.29
Minimal ratio cell area/nucleus area	2.11	1.93	1.35	2.11
t <sub>regime1</sub> (s)	_	1,800	1,250	1,230
$t_{\text{regime2}}(s)$	_	2,450	1,350	1,250
$t_{\text{regime3}}$ (s)	-	2,600	1,850	-

when only part of the body (or nothing) penetrates within the channel as it takes place for channel 4 (purple line in Fig. 3 and Movie 4) where the total displacement is only equal to 7.5 μm.

In Fig. 4, the trend of the cell average velocity is represented. As a general remark, the velocity during the contraction phase is slightly higher than during the contraction phase, since the former only involves the frontal portion of the cytoplasm (see Sect. 2.3). While the average protrusion velocity remains rather constant for all the channels (roughly  $5 \cdot 10^{-3} \,\mu m/s$ ), the average contraction velocity varies between a minimal value of about  $10^{-2} \mu m/s$ for channel 16 (blue line Fig. 4) and a maximal value of  $1.2 \cdot 10^{-2} \,\mu$ m/s for channel 12 (red line Fig. 4). Additionally, for channel 7 (green line Fig. 4), we observe a peak of the velocity up to  $1.3 \cdot 10^{-2} \,\mu\text{m/s}$  at the entrance of the channel, while afterwards the cell acquires again a constant velocity. Such values are of the same order of magnitude of those experimentally observed for HeLa cells (Ronot et al. 2000; Ngalim et al. 2013).

Second, for each configuration, we have quantified the entry time  $(T_{entry})$ , which has been defined by Lautenschläger et al. (Lautenschlager et al. 2009) as the time interval between the first contact of the cell with the channel walls ( $t_{contact}$ ) and the complete penetration of the cell body within the channel (t<sub>penetration</sub>). For channel 16 and channel 4, such a parameter cannot be evaluated since the cell either does not enter in contact with the channel (channel 16) or does not migrate through it (channel 4). For channel 12 and channel 7, we found 2,650 and 3,360 s respectively, which confirms that the smaller the channel, the more the difficult is for the cell to get in. In fact, the contact cell channel occurs earlier for channel 7 than for channel 12 ( $t_{contact} = 1,250$  s versus  $t_{\text{contact}} = 1,950 \text{ s}$ , while  $t_{\text{penetration}}$  is almost the same



Fig. 2 Snaphots of the permeative (a and b), invasive (c) and penetrating (d) cell



Fig. 3 Total displacement of the cell for *channel 16* (blue line), *channel 12* (red line), *channel 7* (green line) and *channel 4* (purple line)

for both channels ( $t_{\text{penetration}} = 4,610 \, s$  versus  $t_{\text{penetration}} = 4,600 \, s$ ).

Third, we have evaluated the ratio between the total cell area and the nucleus area. At the initial configuration, such ratio is equal to 2.8, but it undergoes an oscillatory variation due to the protrusion–contraction movement of the cell. In the case of *channel 16* (Fig. 5, blue line), it varies between a maximal value of 3.3 during protrusion and a minimal value of 2.1 during contraction. Here, such values are the same at the end of each phase during the whole simulation since the cell overall deformation is not perturbed by the contact with channel. For *channel 12* instead, we observe a decrease of the maximal value of the ratio to 2.9 once the cell has completely



**Fig. 4** Migration average velocity of the cell for *channel 16* (*blue line*), *channel 12* (*red line*), *channel 7* (*green line*) and *channel 4* (*purple line*)

entered the channel ( $t_{penetration} = 4,600 \text{ s}$ , Fig. 5, red line), while the minimal value decreases to 1.9. Such drop is mainly due to a bigger shrinkage of the cell cytoplasm rather than of the nucleus due to the subcellular dimensions of the channel. However, in the case of *channel* 7 (Fig. 5, green line), both cytoplasm and nucleus contribute to the progressive decrease of the ratio. In fact, the nucleus must squeeze too to move forward since the channel has sub-nuclear dimensions. Then, the maximal and minimal values of the ratio at  $t_{penetration} = 4,610 \text{ s}$  decrease down to 2.25 and 1.35, respectively. For *channel* 4 (Fig. 5, purple line), the ratio evolution is the same as for *channel* 16 since the cell is not able to penetrate the channel and neither cytoplasm nor nucleus do not undergo large deformation.



**Fig. 5** Ratio between the cell area and the nucleus area for *channel 16* (*blue line*), *channel 12* (*red line*), *channel 7* (*green line*) and *channel 4* (*purple line*)

Finally, we have analysed the morphology of the cell relative to the channel, which can actually be divided into three regimes. The first regime is observed when the length  $L_p$ of the cell protrusion into the channel is smaller than half the width of the channel  $W_{c,i} (2L_p/W_{c,i} < 1)$ , and it has been indicated as  $t_{\text{regime1}}$ . The second regime occurs when  $2L_p/W_{c,i} = 1$ , and the protrusion is hemicircular with radius equal to  $W_{c,I}(t_{\text{regime2}})$ . Finally, the third regime is obtained when  $2L_p/W_{c,i} > 1$  ( $t_{\text{regime3}}$ ). At this point, the first half of the protrusion is rectangular of length  $W_{c,i}$  and the second half is hemicircular of radius  $L_c$ .

In the case of *channel 16*, the migration mode and the morphology of the cell do not change and are very similar to those observed for cell migrating over flat surfaces (Allena and Aubry 2012; Allena 2013). In fact, there is no contact

Fig. 6 The three regimes of the

cell morphology during the migration through *channel 7* 

between the cell cortex and the channel walls, and thus, the cell body is not perturbed during its movement. This is not the case for channel 12 and channel 7 where the cell needs to squeeze in order to enter the channel. For channel 12, regime 1 is observed at  $t_{\text{regime1}} = 1,800 \text{ s}$ , while  $L_p$  becomes equal to  $W_{c,2}/2$  at  $t_{\text{regime}2} = 2,450 \text{ s.}$  Starting from  $t_{\text{regime}3} =$ 2,600 s, regime 3 is achieved and the protrusion is clearly half rectangular and half hemicircular. For *channel* 7 (Fig. 6). steps occur earlier. In fact, regime 1 and regime 2 are reached at  $t_{\text{regime1}} = 1,250 \text{ s}$  and  $t_{\text{regime2}} = 1,350 \text{ s}$ , respectively, while regime 3 starts at  $t_{regime3} = 1,850$  s. For *channel 4* instead, only regime 1 and 2 observed at  $t_{regime1} = 1,230 s$ and  $t_{\text{regime2}} = 1,250 \text{ s}$ , respectively. The reason why regime 3 is not achieved is mainly due to the fact that, despite the cell tries to enter the channel by protruding and contracting, the force  $f_{\text{channel}}$  exerted by the channel walls on the cell boundaries is too high. This means that reaching regime 3 is a necessary, but not sufficient condition for the cell to be invasive. In fact, a second necessary condition need to be satisfied, that is, the cell-channel surface force  $f_{\text{channel}}$  at  $t_{\text{regime3}}$  must be low enough for the cell to enter.

We have also been able to experimentally observe such changes in morphology for two types of cells using a microchannel-based essay as proposed in (Heuzé et al. 2011). Figure 7a–d shows the successive steps (top view) of bone marrow-derived dendritic cells (BDMCs) migration through a 5 $\mu$ m (Fig. 7a–b) and 1.5 $\mu$ m (Fig. 7c, d) wide microchannel. It is possible to clearly distinguish the three regimes undergone by the whole cell body (Fig. 7b, d) and by the stained nucleus (Fig. 7a, c). Figure 7e shows instead a sagittal view of the successive steps of a HeLa cell migrating through a 20 $\mu$ m wide micro-channel. We observe the deformation undergone by the stained nucleus along the *z* axis. In fact, in this specific case, the cell is confined in the *x*–y plane,



(c) t = 1850 s



**Fig. 7** (a:d) Top view of successive steps of a bone marrow-derived dendritic cell (BMDC) migration (from the *left* to the *right*) through a  $5\mu$ m **a**-**b** and  $1.5\mu$ m **c**-**d** wide micro-channels. Nuclear staining with Hoechst (**a** and **c**) (*scale bar*:  $30\mu$ m) **e** Sagittal view of successive steps

of a HeLa cell entering (from the *left* to the *right*) a  $20\mu$ m wide microchannel (HeLa Histone2B-mcherry (nucleus), MyrPalm-GFP (plasma membrane), *scale bar*:  $15\mu$ m)

Table 3 Values of the mechanical forces for the different channels

	Channel 16	Channel 12	Channel 7	Channel 4
Maximal frontal cell-substrate surface force (Pa)	10	10	10	10
Maximal rear cell-substrate surface force (Pa)	4	4	4	4
Maximal cell-channel surface force at $t_{\text{contact}}$ (Pa)	-	3.3	3.1	7.2
Maximal cell-channel surface force at $t_{\text{penetration}}$ (Pa)	_	3.3	3.9	-
Average cell-substrate surface force between $t_{contact}$ and $t_{penetration}$ (Pa)	-	3.3	3.3	3.3
Average cell-channel surface force between $t_{contact}$ and $t_{penetration}$ (Pa)	-	2.5	3.2	$t_{\rm contact} - t_{\rm regime2}$ 4.5
Absolute maximal cell-channel surface force (Pa)	-	4.2	6.2	8.6

but also in the x-z plane. Although such an aspect has not been numerically considered so far, we are currently working to improve the model in order to have a three-dimensional representation of the cell and the micro-channel and therefore being able to implement this further confinement

#### 3.2 Mechanical forces

In this section, we try to evaluate the cell-substrate and cellchannel surface forces, in particular during the time interval between  $t_{\text{contact}}$  and  $t_{\text{penetration}}$  in which  $t_{\text{regime3}}$  is included. The main values are summarized in Table 3.

Some general remarks may be pointed out:

- given the asymmetry of the active strain (Sect. 2.3) and the equation expressing the cell-substrate surface forces (Eq. 3), we found 10 and 4 Pa, respectively, at the front and rear edge of the cell. Additionally, such values do not change from one configuration;
- the cell-channel surface force increases as the channel width  $W_{c,i}$  decreases (maximal absolute value of 4.25,

6.25 and 8.5 Pa for *channel 12, channel 7* and *channel 4*, respectively);

- as mentioned in Sect. 3.1, the necessary condition for the cell to be invasive is that the average cell-channel surface force at  $t_{regime3}$  must be lower than the average cell-substrate surface force at the same time point. This allows the cell to pull enough to penetrate within the channel without being obstructed by the channel. Since  $t_{\text{regime3}}$  is included in the time interval between  $t_{\text{contact}}$  and  $t_{\text{penetration}}$ , we have calculated the average cell-channel surface force during this period. For channel 12 and channel 7, we have found an average value for the cell-channel surface force of about 2.5 Pa and 3.2, respectively, which is lower than the average cell-substrate surface force of 3.3 Pa. As a result, the cell is able to enter the channel. For channel 4, since the cell-channel surface force between  $t_{\text{contact}}$  and  $t_{\text{penetration}}$  cannot be calculated, we have evaluated it between  $t_{contact}$  and  $t_{regime2}$  finding an average value of 4.5 Pa and a maximal value of 8.6 Pa, which is twice the cell-substrate surface force. Therefore, the cell is stuck at the entrance of the channel and shows a penetrating behaviour;
- once the cell has completely penetrated into the channel, the upper and lower central boundaries of the cell come very close or directly in contact with the nucleus, which is the stiffest component of the system. Then, a higher cellchannel surface force is necessary at this specific region to maintain the cell squeezing during the whole migration process and in order for the cell to be permeative. This is the case for *channel 12* for which the cell is able to reach the opposite end of the channel (Movie 5). However, for *channel 7*, the cell-channel surface force is slightly higher at the rear of the cell. Thus, the cell is slowed down and shows a penetrating behaviour (Movie 6).

For *channel 16*, only the cell-substrate surface force can be evaluated while the cell-channel surface force is null since no contact between the cell boundaries and the channel walls occurs.

#### 4 Conclusions

In this paper, we have proposed a 2D mechanical model to simulate the migration of HeLa cell under confinement. The model reproduces the set-up used in a micro-channel assay as presented in (Heuzé et al. 2011). As in our previous works (Allena and Aubry 2012; Allena 2013), the cell is modelled as continuum and a standard Maxwell model is used to describe the mechanical behaviour of the cytoplasm (including active strains) and the nucleus. The cell is able to cyclically develop protrusion–contraction strains, which are synchronized with

the adhesion forces between the cell and the substrate. By approaching the channel, which is represented here by two pseudo-elliptical rigid walls, the cell is submitted to an additional viscous force. We have tested four channels whose dimensions in terms of width are larger than the cell diameter (channel 16), sub-cellular (channel 12), sub-nuclear (channel 7) and much smaller than the nucleus diameter (channel 4). We have analysed the cell behaviour and classified it as permeative (channel 16 and channel 12), invasive (channel 7) or penetrating (channel 4) according to the distance covered by the cell inside the channel. From a morphological point of view, we have identified three different regimes in relation to the ratio between the cell protrusion length in the channel and the width of the channel. Additionally, we have evaluated the evolution of the cell shape and the cell-substrate and cellchannel surface forces between the first contact between the cell and the channel  $(t_{contact})$  and the complete penetration of the cell body within the channel ( $t_{penetration}$ ).

Therefore, we have been able to define the necessary conditions in order for the cell to be invasive or permeative. In the first case, two main conditions must be satisfied: i) regime 3 (i.e. cell protrusion length in the channel larger than half the channel width) has to be achieved, and ii) simultaneously, the cell-substrate surface force must be higher than the cellchannel surface force so that the cell is able to pull on the substrate and enter into the channel. For the second behaviour to occur, a further condition must be satisfied, that is, the cell-channel surface force during the whole migration has to be maximal along the upper and lower central boundaries of the cell. Those boundaries may come very close or directly in contact with the cell nucleus, which is the stiffest component of the system. Then, a larger force is required to maintain the squeezed cell shape.

Despite the consistent results shown in the present paper, our model still presents some limitations. Firstly, the geometry is 2D, which does not allow considering a top-roofed micro-channel and the cell deformation in the third direction. Secondly, the active strains of protrusion and contraction have been defined through a sinusoidal function, which may lead to a rather stable periodic deformation of the cytoplasm and consequently of the nucleus. In order to control the effects of such a phenomenon, some stochastic active input close to cell perception may be introduced and improve the global movement. Finally, so far all the cell components have been considered as viscoelastic materials. However, the nucleus may be able to adapt its deformation to the forces exerted by the micro-channel on the cell boundaries. Therefore, a viscoplastic behaviour with restoration (Mandel 1972; Lubliner 2008) would probably be more appropriate. We are currently working on this aspect in order to be able to investigate the ability of the cell to penetrate micro-channels with significant sub-nuclear dimensions.

#### **5** Appendix

#### 5.1 Geometry of the cell

For any spatial point **p**, the four components of the cell body (the cortex  $\Omega_{cortex}$ , the cytosol  $\Omega_{cytosol}$ , the lamina  $\Omega_{lamina}$ and the nucleoplasm  $\Omega_{nucleoplasm}$ ) are described through characteristic functions (i.e. composition of a Heaviside and a level set function (Allena 2013) as follows

$$h_{\text{cortex}} \left( \boldsymbol{p} \right) = \begin{cases} 1 & \text{if} r_{\text{cytoplasm}}^2 < \| \boldsymbol{p} - \boldsymbol{c}_{\text{cell}} \| < r_{\text{cortex}}^2 \\ 0 & \text{otherwise} \end{cases}$$
(5)

$$h_{\text{cytosol}}(\boldsymbol{p}) = \begin{cases} 1 & \text{if } r_{\text{lamina}}^2 < \|\boldsymbol{p} - \boldsymbol{c}_{\text{cell}}\| < r_{\text{cytosol}}^2 \\ 0 & \text{otherwise} \end{cases}$$
(6)

$$h_{\text{lamina}}(\boldsymbol{p}) = \begin{cases} 1 & \text{if} r_{\text{nucleoplasm}}^2 < \|\boldsymbol{p} - \boldsymbol{c}_{\text{cell}}\| < r_{\text{lamina}}^2 \\ 0 & \text{otherwise} \end{cases}$$

$$c_{\text{oot}} \parallel < r^2$$
 .

(7)

$$h_{\text{nucleoplasm}}(\boldsymbol{p}) = \begin{cases} 1 & \text{if } \|\boldsymbol{p} - \boldsymbol{c}_{\text{cell}}\| < r_{\text{nucleoplasm}}^2 \\ 0 & \text{otherwise} \end{cases}$$
(8)

where p = x - u, with x and u being, respectively, the actual position and the displacement,  $c_{cell}$  is the cell centre and  $r_{\text{cortex}}$ ,  $r_{\text{cvtosol}}$ ,  $r_{\text{lamina}}$  and  $r_{\text{nucleoplasm}}$  are the external radius of the cell cortex, the cytosol, the nuclear lamina and nucleoplasm, respectively (Fig. 1a). Therefore, the cytoplasm  $\Omega_{cytoplasm}$  and the nucleus  $\Omega_{nucleus}$  domains are defined by the following characteristic functions

$$h_{\text{cytoplasm}}(\boldsymbol{p}) = h_{\text{cortex}}(\boldsymbol{p}) + h_{\text{cytosol}}(\boldsymbol{p})$$
$$h_{\text{nucleus}}(\boldsymbol{p}) = h_{\text{lamina}}(\boldsymbol{p}) + h_{\text{nucleoplasm}}(\boldsymbol{p})$$
(9)

The frontal portion of cytosol where the polymerization of the actin filaments takes place is described as follows

$$h_{\text{cytosol, front}} \left( \boldsymbol{p} \right) = \begin{array}{c} h_{\text{cytosol}} & if \, \boldsymbol{p} > \boldsymbol{c}_{\text{cell}} \\ 0 & \text{otherwise} \end{array}$$
(10)

The frontal  $(\Omega_f)$  and rear  $(\Omega_r)$  adhesion regions are also defined by two characteristic functions as

$$h_{f}(\mathbf{p}) = \begin{array}{c} 1 & (\mathbf{p} - \mathbf{c}_{\text{cell}}, \mathbf{i}_{x}) > l_{f} \\ 0 & \text{otherwise} \end{array}$$

$$h_{r}(\mathbf{p}) = \begin{array}{c} 1 & (\mathbf{p} - \mathbf{c}_{\text{cell}}, \mathbf{i}_{x}) < -l_{r} \\ 0 & \text{otherwise} \end{array}$$
(11)

with  $l_f$  and  $l_r$  the distances of  $c_{cell}$  from the boundaries of  $\Omega_f$ and  $\Omega_r$ , respectively, (Fig. 1b). As soon as the cell moves, the argument p is replaced by x-u, with x the actual spatial position and u the displacement.

#### 5.2 Nucleus constitutive law

As mentioned in Sect. 2.2, the nucleus is described through a viscoelastic constitutive equation based on a standard Maxwell model including a solid phase (i.e. the lamina) and a fluid phase (i.e. the nucleoplasm) (Fig. 8).

The Cauchy stress  $\sigma_{nucleus}$  and the deformation tensor  $F_{\text{nucleus}}$  in the nucleus are defined by

$$\sigma_{\text{nucleus}} = \sigma_{\text{lamina}} + \sigma_{\text{nucleoplasm}}$$

$$F_{\text{nucleus}} = D_p u + I = F_{\text{lamina}} = F_{\text{nucleoplasm}}$$
(12)

where  $D_p u = {3 \atop m=1} {\partial u \over \partial p_m} \otimes i_m$ , with u the displacement and I the identity matrix (Holzapfel 2000; Taber 2004), and  $F_{\text{nucleoplasm}} = F_{\text{nucleoplasm},e}F_{\text{nucleoplasm},v}$ . The solid part of the stress  $\sigma_{\text{lamina}}$  in the lamina reads

$$\boldsymbol{\sigma}_{\text{lamina}} = \frac{1}{J_{\text{lamina}}} \boldsymbol{F}_{\text{lamina}} \boldsymbol{S}_{\text{lamina}} \boldsymbol{F}_{\text{lamina}}^T$$
(13)

where  $J_{\text{lamina}}$  is the determinant of  $F_{\text{lamina}}$  and  $S_{\text{lamina}}$  is the second Piola-Kirchoff stress tensor, which is computed as an isotropic hyperelastic Saint Venant material as follows

$$S_{\text{lamina}} = \lambda_{\text{lamina}} Tr \left( E_{\text{lamina}} \right) I + 2\mu_{\text{lamina}} E_{\text{lamina}}$$
(14)

with  $\lambda_{\text{lamina}}$ ,  $\mu_{\text{lamina}}$  and  $E_{\text{lamina}}$  the Lame's coefficients and the Green-Lagrange strain tensor of the solid phase, respectively.

The fluid part of the stress  $\sigma_{\text{nucleoplasm}}$  in the nucleoplasm can be expressed as

$$\boldsymbol{\sigma}_{\text{nucleoplasm}} = 2\mu_{\text{nucleoplasm}}\boldsymbol{D}_{\text{nucleoplasm},v}$$
(15)

with  $\mu_{nucleoplasm}$ , the viscosity of the nucleoplasm and the eulerian strain rate  $D_{nucleoplasm,v}$  is computed from the strain gradient velocity as

$$2D_{\text{nucleoplasm},v} = \dot{F}_{\text{nucleoplasm},v} F_{\text{nucleoplasm},v}^{-1} + F_{\text{nucleoplasm},v}^{-T} \dot{F}_{\text{nucleoplasm},v}^{T}$$
(16)

#### 5.3 Cytoplasm constitutive law

The cytoplasm is composed by two phases: i) a solid phase represented by the cell cortex and ii) a fluid phase represented by the viscous cytosol with the embedded organelles such as the actin filaments that undergo the active strains (Fig. 8). It is assumed that the Cauchy stress  $\sigma_{cytoplasm}$  and the deformation tensor  $F_{\rm cytoplasm}$  read

$$\sigma_{\text{cytoplasm}} = \sigma_{\text{cortex}} + \sigma_{\text{cytosol}}$$

$$F_{\text{cytoplasm}} = F_{\text{cortex}} = F_{\text{cytosol}}$$
(17)

Additionally, the fluid deformation tensor  $F_{\text{cytosol}}$  is multiplicatively decomposed as follows

$$F_{\text{cytosol}} = F_{\text{cytosol},v} F_{\text{cytosol},e} F_{\text{cytosol},a}$$
(18)

where e and v stand for elastic and viscous, respectively.

The solid stress  $\sigma_{\text{cortex}}$  in the organelles can be written as

$$\boldsymbol{\sigma}_{\text{cortex}} = \frac{1}{J_{\text{cortex}}} \boldsymbol{F}_{\text{cortex}} \boldsymbol{F}_{\text{cortex}}^{T}$$
(19)

Fig. 8 Symbolic schemas for the standard Maxwell models used to describe the nucleus (*top*) and the cytoplasm (*bottom*) behaviours



with  $J_{\text{cortex}}$  the determinant of  $F_{\text{cortex}}$  and  $S_{\text{cortex}}$  the second Piola–Kirchoff stress tensor, which, similarly to the nucleus (Sect. 5.2), is defined as an isotropic hyperplastic Saint Venant material as follows

$$S_{\text{cortex}} = \lambda_{\text{cortex}} Tr \left( E_{\text{cortex}} \right) I + 2\mu_{\text{cortex}} E_{\text{cortex}}$$
(20)

where  $\lambda_{\text{cortex}}$ ,  $\mu_{\text{cortex}}$  and  $E_{\text{cortex}}$  the Lame's coefficients and the Green-Lagrange strain tensor of the solid phase, respectively.

Finally, the fluid stress  $\sigma_{\rm cytosol}$  in the cytosol reads

$$\boldsymbol{\sigma}_{\text{cytosol}} = 2\mu_{\text{cytosol}}\boldsymbol{D}_{\text{cytosol},v} \tag{21}$$

with  $\mu_{\text{cytosol}}$  the viscosity of the cytosol and  $D_{\text{cytosol},v}$  the eulerian strain rate expressed as follows

$$2\boldsymbol{D}_{\text{cytosol},v} = \dot{\boldsymbol{F}}_{\text{cytosol},v} \boldsymbol{F}_{\text{cytosol},v}^{-1} + \boldsymbol{F}_{\text{cytosol},v}^{-T} \dot{\boldsymbol{F}}_{\text{cytosol},v}^{T}$$
(22)

#### 5.4 Micro-channel geometry

As mentioned in Sect. 2.4, the micro-channel domain  $\Omega_{\text{channel}}$  is modelled as two pseudo-elliptical rigid walls: a upper one  $(\Omega_{uw})$  and a lower one  $(\Omega_{lw})$ . They are described through two characteristic functions as follows

$$h_{uw,i} (\mathbf{p}) = \begin{cases} 1 & \text{if} l_{uw,i} < 1 \\ 0 & \text{otherwise} \end{cases}$$

$$h_{lw,i} (\mathbf{p}) = \begin{cases} 1 & \text{if} l_{lw,i} < 1 \\ 0 & \text{otherwise} \end{cases}$$
(23)

where the subscript '*i*' indicates the number of the channel, and  $l_{uw,i}$  and  $l_{lw}$  are two level set functions expressed as

$$l_{uw,i} = \left(\frac{x - x_0}{a}\right)^4 + \left(\frac{y - y_{uw0,i}}{b}\right)^4$$
$$l_{lw,i} = \left(\frac{x - x_0}{a}\right)^4 + \left(\frac{y + y_{lw0,i}}{b}\right)^4$$
(24)

with *a* and *b* are the semi-axes of the pseudo-elliptical walls with centres  $c_{uw,i}(x_0, y_{uw0,i})$  and  $c_{lw,i}(x_0, y_{lw0,i})$ . Thus, the micro-channel is the composition of the two previous characteristic functions as follows

$$\Omega_{\text{channel}} = h_{uw,i} \left( \boldsymbol{p} \right) + h_{lw,i} \left( \boldsymbol{p} \right) \tag{25}$$

The c outward normals  $\boldsymbol{n}_{uw}$  and  $\boldsymbol{n}_{lw}$  to the boundary  $\partial \Omega_{uw}$ and  $\partial \Omega_{lw}$ , respectively, given by

$$\boldsymbol{n}_{uw,i} = h' \left( l_{uw,i} \right) \frac{\nabla l_{uw,i}}{\left\| \nabla l_{uw,i} \right\|}$$
$$\boldsymbol{n}_{lw,i} = h' \left( l_{lw,i} \right) \frac{\nabla l_{lw,i}}{\left\| \nabla l_{lw,i} \right\|}$$
(26)

where h' indicates the Dirac function.

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