Draft for the thesis

Francisco Vazquez-Tavares
July 11, 2025

Summary Document for the draft of the Thesis.

Introduction

Acknowledge

Objectives

Justification

Theoretical framework

Description of the microgel

Description of the microgel as a colloid to introduce the langevin methodology.

"The dynamics of a macromolecular system is entirely determined by the potential U(rN) associated with the process. For computational and practical reasons, this potential is virtually always an approximation of the real physical potential¹." [Paquet and Viktor, 2015]

Brownian dynamics

2

From a general point of view there are two types of methods to make a quatitative description of systems: one focused on simulating dynamics at the microscale, and the other dedicated to deriving or establishing evolutionary equations at the macroscale[Wang et al., 2025]. Since we assume that the a microgel's mechanical response derives from its internal structure³ we choose to simulate the dynamics at the microscale. Additionally, by treating the microgel as a colloid, permits applying Brownian motion theory to model its response under shear deformation. Finally, there are two commonly used mathematical frameworks to model the Brownian motion, the continuous time random walk (CTRW) model and the Langevin equation[Wang et al., 2025], in this work we decided⁴ to use the langevin dynamics mathematical framework.

This is because, the solid phase of the colloid has a large mass and will change their momenta after many collisions with the solvent molecules and the picture which emerges is that of the heavy particles forming a system with a much longer time scale than the solvent molecules[Thijssen, 2007] and Langevin theory takes advantage of this difference in time scale to eliminate the details of the degrees of freedom of the solvent particles and represent their effect by stochastic and dissipative forces allowing longer simulations that would be impossible if the solvent were explicitly included[Pastor, 1994]. However, the representation of the solvent by a

¹ Those potentials are explained in the following section?
Would be better to describe the parameters in the implementation and the model here?

² The problem is characterised by two very different timescales, one associated with the slow relaxation of the initial velocity of the brownian particle and another linked to the frequent collisions that the brownian particle suffers with particles of the bath. (Problema de implementación o representación matemática, eso del tiempo de relajación por condiciones iniciales es por que matemáticamente se reuieren condiciones iniciales para resolver el sistema ecuaciones diferenciales.) Langevin assumed that the force acting on the brownian particle consists of two parts: a systematic, frictional force proportional to the velocity u(t), but acting in the opposite sense, and a randomly fluctuating force, R(t), which arises from collisions with surrounding particles[Hansen and McDonald, 2006].

³ Poner citas que desmuestrén que no es hipótesis, si no que se sabe

⁴ Supongo que eventualmente justificaré la desición.

stochastic and dissipative force, introduce the problem of characterize two very different timescales, one associated with the slow relaxation of the initial velocity of the brownian particle and another linked to the frequent collisions that the brownian particle suffers with particles of the bath[Hansen and McDonald, 2006]⁵. Therefore, two terms are used to create a mathematical representation of the solvent: a frictional force proportional to the velocity of the brownian particle and a fluctuating force. Hence,

$$m\frac{\mathrm{d}\vec{v}(t)}{\mathrm{d}t} = \vec{F}(t) - m\gamma\vec{v}(t) + \vec{R}(t). \tag{1}$$

The friction constant γ^6 parametrises the effect of solvent damping and activation and is commonly referred to as the collision frequency in the simulation literature, even though formally a Langevin description implies that the solute suffers an infinite number of collisions with infInitesimally small momentum transfer. Also, the fact that the second term is not a function of the position of any of the particles involves the neglect of involves the neglect of hydrodynamic interaction or spatial correlation in the friction kernel spatial correlation in the friction kernel [Pastor, 1994]. On the other hand, $\vec{R}(t)^7$ is a "random force" subject to the following conditions

$$\left\langle \vec{R}(t) \right\rangle = 0$$

 $\left\langle \vec{R}(t)\vec{R}(t') \right\rangle = 2k_B T \gamma \delta(t - t')$

The no time correlation is equivalent to assuming that the viscoelastic relaxation of the solvent is very rapid with respect to solute motions⁸.

In comparing the results of Langevin dynamics with those of other stochastic methods [28-31], the relevant variable is the velocity relaxation time, τ_{ν} which equals γ^{-1} [Pastor, 1994] The Langevin equation improves conformational sampling over standard molecular dynamics[Paquet and Viktor, 2015].

- Hablar acerca de que la fuerza aleatoria puede tener distribución gaussiana, pero no necesariamente.
- hablar de la ecuación de Green-Kubo:

$$\eta = rac{V}{k_B T} \int_0^\infty \left\langle \sigma_{xy}(t) \sigma_{xy}(0)
ight
angle \mathrm{d}t$$

 No se que tanto hablar de la idea de correlación y su aplicación en estos temas.

Stress

Introductory paragraph To characterize the behaviour of materials, constitutive relations serve as an input to the continuum theory...⁹

This derivation can be found in the apendix of [Admal and Tadmor, 2010]¹⁰.¹¹ Consider a system of N interacting particles with each particle position given by

$$\vec{r}_{\alpha} = \vec{r} + \vec{s}_{\alpha},\tag{2}$$

⁵ Para traer a colación la sensibilidad de la respuesta mecánica al parámetro de damp.

⁶ Cuidado con las unidades. Hacer análisis dimensional, porque por la condición de correlación en R, γ ocupa tener unidades de masa entre tiempo, pero en la ecuación, solo ocupa unidades de 1/s.

⁷ No me acuerdo en donde está que se puede asumir que tiene distribución gaussiana.

⁸ Grote land Hynes [26] have investigated this assumption for motions involving barrier crossing and have found that while it is seriously in error for passage over sharp barriers (such as 12 recombination); it is quite adequate for conformational transitions such as might be found in polymer motions.[Pastor, 1994]

⁹ Capaz e ir introduciendo ideas del Clausius[Clausius, 1870]

¹⁰ Describe more if what is done in this article

¹¹ (Eventualmente pondré esto en párrafo) Notation: σ Tensor, $\vec{\sigma}$ vector, $\sigma_{i,j}$ tensor, $\overline{\sigma}$ time average.

where \vec{r} is the position of the center of mass of the system and \vec{s}_{α} is the position of each point relative to the center of mass. Hence, we can express the momentum of each particle as

$$\vec{p}_{\alpha} = m_{\alpha} (\dot{\vec{r}} + \dot{\vec{s}}_{\alpha}) = m_{\alpha} (\dot{\vec{r}} + \vec{v}_{\alpha}^{\text{rel}}).$$
 (3)

Before starting the procedure, lets take into account that the center of mass of the system is given by

 $\vec{r} = \frac{\sum_{\alpha} m_{\alpha} \vec{s}_{\alpha}}{\sum_{\alpha} m_{\alpha}},\tag{4}$

and by replacing (2) in (3) we get the following relations, which will be used later,

$$\sum_{\alpha} m_{\alpha} \vec{r}_{\alpha} = \vec{0}, \quad \sum_{\alpha} m_{\alpha} \vec{v}_{\alpha}^{\text{rel}} = \vec{0}. \tag{5}$$

Now we can start by computing the time derivative of tensorial product $\vec{r}_{\alpha} \otimes \vec{p}_{\alpha}^{12}$,

$$\frac{\mathrm{d}}{\mathrm{d}t}(\vec{r}_{\alpha} \otimes \vec{p}_{\alpha}) = \underbrace{\vec{v}_{\alpha}^{\mathrm{rel}} \otimes \vec{p}_{\alpha}}_{\text{Kinetic term}} + \underbrace{\vec{r}_{\alpha} \otimes \vec{f}_{\alpha}}_{\text{Virial term}}, \tag{6}$$

which is known as the *dynamical tensor virial theorem* and it is simply an alternative form to express the balance of linear momentum. This theorem becomes useful after making the assumption that there exists a time scale τ , which is short relative to macroscopic processes but long relative to the characteristic time of the particles in the system, over which the particles remain close to their original positions with bounded positions and velocities. Taking advantage of this property we can compute the time average of (6),

$$\frac{1}{\tau} (\vec{r}_{\alpha} \otimes \vec{p}_{\alpha}) \Big|_{0}^{\tau} = \overline{\vec{v}_{\alpha}^{\text{rel}} \otimes \vec{p}_{\alpha}} + \overline{\vec{r}_{\alpha} \otimes \vec{f}_{\alpha}}. \tag{7}$$

Assuming that $\vec{r}_{\alpha} \otimes \vec{p}_{\alpha}$ is bounded, and the time scales between microscopic and continuum processes are large enough, the term on the left-hand side can be as small as desired by tacking τ sufficiently large and by summing over all particles we achieve the *tensor virial theorem*:

$$\overline{\mathbf{W}} = -2\overline{\mathbf{T}},\tag{8}$$

where

$$\overline{\mathbf{W}} = \sum_{\alpha} \overline{\vec{r}_{\alpha} \otimes \vec{f}_{\alpha}} \tag{9}$$

is the time-average virial tensor and

$$\overline{\mathbf{T}} = \frac{1}{2} \sum_{\alpha} \overline{\vec{v}_{\alpha}^{\text{rel}} \otimes \vec{p}_{\alpha}} \tag{10}$$

is the time-average kinetic tensor. This expression for the tensor virial theorem applies equally to continuum systems that are not in macroscopic equilibrium as well as those that are at rest.

The assumption of the difference between the time scales allow us to simplify the relation by replacing (3) in (10), so that,

$$\overline{\mathbf{T}} = \frac{1}{2} \sum_{\alpha} m_{\alpha} \overline{\vec{v}_{\alpha}^{\text{rel}} \otimes \vec{v}_{\alpha}^{\text{rel}}} + \frac{1}{2} \left[\overline{\sum_{\alpha} m_{\alpha} \vec{v}_{\alpha}^{\text{rel}}} \right] \otimes \dot{\vec{r}}, \tag{11}$$

which is not the simplification we expected, however, by the relations from (5), equation (11) simplifies to (5)

 $^{^{12}}$ It is interesting to note that the tensorial product $\vec{r}_{\alpha} \otimes \vec{p}_{\alpha}$ has units of action and by tacking the time derivative we are dealing with terms that has units of energy.

¹³ No estoy muy seguro si incluir una discusión acerca del término cinético en la expresión del virial. Posiblemente un párrafo... posiblemente lo ponga en la interpretación del teorema. También, no se si ir metiendo interpretación durante la derivación o no, pero bueno.

$$\overline{\mathbf{T}} = \frac{1}{2} \sum_{\alpha} m_{\alpha} \overline{\vec{v}_{\alpha}^{\text{rel}} \otimes \vec{v}_{\alpha}^{\text{rel}}}.$$
 (12)

On the other hand, instead of reducing the expression, we start to create the conection with the Cauchy stress tensor by distributing (9) into an internal and external contributions,

$$\overline{\mathbf{W}} = \underbrace{\sum_{\alpha} \overrightarrow{r}_{\alpha} \otimes \overrightarrow{f}_{\alpha}^{\text{int}}}_{\overline{\mathbf{W}}_{\text{int}}} + \underbrace{\sum_{\alpha} \overrightarrow{r}_{\alpha} \otimes \overrightarrow{f}_{\alpha}^{\text{ext}}}_{\overline{\mathbf{W}}_{\text{ext}}}.$$
 (13)

The time-average internal virial tensor takes into account the interaction between particle α with the other particles in the system, meanwhile, the time-average external virial tensor considers the interaction with atoms outside the system, via a traction vector \vec{t} and external fields acting on the system represented by $\rho \vec{b}$, where ρ is the mass density of it and \vec{b} is the body force per unit mass applied by the external field. Therefore we can express the following,

$$\sum_{\alpha} \overline{\vec{r}_{\alpha} \otimes \vec{f}_{\alpha}^{\text{ext}}} := \int_{\delta\Omega} \vec{\xi} \otimes \vec{t} dA + \int_{\Omega} \vec{\xi} \otimes \rho \vec{b} dV. \tag{14}$$

Where $\vec{\xi}$ is a position vector within the domain Ω occupied by the system of particles with a continuous closed surface $\delta\Omega$. Assuming that Ω is large enough to express the external forces acting on it in the form of the continuum traction vector \vec{t} .

With this we can substitute the traction vector with $\vec{t} = \sigma \vec{n}$, where σ represent the Cauchy stress tensor and applying the divergence theorem in (14), we have

$$\overline{\mathbf{W}}_{\text{ext}} = \int_{\Omega} \left[\vec{\xi} \otimes \rho \vec{b} + \text{div}_{\vec{\xi}} \left(\vec{\xi} \otimes \boldsymbol{\sigma} \right) \right] dV = \int_{\Omega} \left[\boldsymbol{\sigma}^{\text{T}} + \vec{\xi} \otimes \left(\text{div}_{\vec{\xi}} \boldsymbol{\sigma} + \rho \vec{b} \right) \right] dV$$
(15)

Since we assume that we are under equilibrium conditions, the term ${\rm div}_{\vec{\xi}} {\bf \sigma} + \rho \vec{b}$ is zero (15) it simplifies to

$$\overline{\mathbf{W}}_{\text{ext}} = V \mathbf{\sigma}^{\text{T}}.\tag{16}$$

By tacking into account that we integrate over the domain Ω we can say that we compute the spatial average of the Cauchy stress tensor,

$$\mathbf{\sigma}_{\mathrm{av}} = \frac{1}{V} \int_{\Omega} \mathbf{\sigma} dV, \tag{17}$$

in which V is the volume of the domain Ω . Replacing (16) into (13), the tensor virial theorem (8) can be expressed as,

$$\sum_{\alpha} \overline{\vec{r}_{\alpha} \otimes \vec{f}_{\alpha}^{\text{int}}} + V \boldsymbol{\sigma}_{\text{av}}^{\text{T}} = -\sum_{\alpha} m_{\alpha} \overline{\vec{v}_{\alpha}^{\text{rel}} \otimes \vec{v}_{\alpha}^{\text{rel}}}.$$
 (18)

Finally, solving for the Cauchy Stress tensor we get,

$$\boldsymbol{\sigma}_{\rm av} = -\frac{1}{V} \left[\sum_{\alpha} \overline{\vec{f}_{\alpha}^{\rm int} \otimes \vec{r}_{\alpha}} + \sum_{\alpha} m_{\alpha} \overline{\vec{v}_{\alpha}^{\rm rel} \otimes \vec{v}_{\alpha}^{\rm rel}} \right], \tag{19}$$

an expression that describe the macroscopic stress tensor in terms of microscopic variables ¹⁴.

¹⁴ It is important to acknowledge that several mathematical subtleties were not taken into consideration, however all the mathematical formality is adressed by Nikhil Chandra Admal and E. B. Tadmor in [Admal and Tadmor, 2010]

To end the section it is important to show that (19) is symmetric. Therefore, we rewrite the internal force as the sum of forces between the particles

$$\vec{f}_{\alpha}^{\text{int}} = \sum_{\beta_{\beta \neq \alpha}} \vec{f}_{\alpha\beta},\tag{20}$$

and substituting (20) into (19), we have

$$\boldsymbol{\sigma}_{\rm av} = -\frac{1}{V} \left[\sum_{\alpha, \beta_{\beta \neq \alpha}} \overline{\vec{f}_{\alpha\beta} \otimes \vec{r}_{\alpha}} + \sum_{\alpha} m_{\alpha} \overline{\vec{v}_{\alpha}^{\rm rel} \otimes \vec{v}_{\alpha}^{\rm rel}} \right]. \tag{21}$$

Due to the property $\vec{f}_{\alpha\beta} = -\vec{f}_{\beta\alpha}$ we obtain the following identity

$$\sum_{\alpha,\beta_{\beta\neq\alpha}} \vec{f}_{\alpha\beta} \otimes \vec{r}_{\alpha} = \frac{1}{2} \sum_{\alpha,\beta_{\beta\neq\alpha}} \left(\vec{f}_{\alpha\beta} \otimes \vec{r}_{\alpha} + \vec{f}_{\beta\alpha} \otimes \vec{r}_{\beta} \right) = \frac{1}{2} \sum_{\alpha,\beta_{\beta\neq\alpha}} \vec{f}_{\alpha\beta} \otimes \left(\vec{r}_{\alpha} - \vec{r}_{\beta} \right).$$
(22)

Therefore, by replacing the identity of (22) into (21), we have

$$\boldsymbol{\sigma}_{\text{av}} = -\frac{1}{V} \left[\frac{1}{2} \sum_{\alpha, \beta_{\beta \neq \alpha}} \overline{\vec{f}_{\alpha\beta} \otimes (\vec{r}_{\alpha} - \vec{r}_{\beta})} + \sum_{\alpha} m_{\alpha} \overline{\vec{v}_{\alpha}^{\text{rel}}} \otimes \overline{\vec{v}_{\alpha}^{\text{rel}}} \right], \quad (23)$$

expressed with indexical notation and using the eistein summation convention,

$$\sigma_{ij}^{\text{av}} = -\frac{1}{V} \left[\frac{1}{2} \sum_{\alpha, \beta_{\beta \neq \alpha}} \overline{f_i^{\alpha\beta} r_j^{\alpha} + f_i^{\beta\alpha} r_j^{\beta}} + \sum_{\alpha} m_{\alpha} \overline{v_i^{\alpha \text{ rel}} v_j^{\alpha \text{rel}}} \right], \quad (24)$$

which is the same expression implemented in LAMMPS[Thompson et al., 2022]. ¹⁵

Computational Implementation

General description

The simulation methodology is based on the work presented in [Gnan et al., 2017] and [Sorichetti et al., 2023], with the objective of create a representative polymer structure of a microgel and characterize the mechanical response under shear deformation. This methodology creates the structure by using a mixture of two types of patchy particles. The patchy particles are spheres of identical size and mass decorated by patches to represent interaction sites. One type represent a *Crosslinker* and is define by 1 central particle with 4 patches placed at the vertices of a circumscribed tetrahedron. The other one represent a *Monomer* define by 1 central particle and 2 patches placed at the poles.

The interaction between the central particles is modeled with a Weeks-Chandler-Andersen repulsive potential,

$$U_{WCA}(r_{i,j}) = \begin{cases} 4\varepsilon_{i,j} \left[\left(\frac{\sigma}{r_{i,j}} \right)^{12} - \left(\frac{\sigma}{r_{i,j}} \right)^{6} \right] + \varepsilon_{i,j}, & r_{i,j} \in [0, 2^{1/6}\sigma], \\ 0, & r_{i,j} > 2^{1/6}\sigma \end{cases}$$
(25)

where $r_{i,j}$ is the distance between the center of the central particles, σ is the diameter of the particles and $\varepsilon_{i,j}$ is the energy of the interacton. The

- \$\vec{r}_i\$: Position of Crosslinker central particle
- \vec{r}_j : Position of Monomer central particle
- {rⁱ_μ} Set of positions of the patches in the crosslinker patchy particle
- {\vec{r}_v^j} Set of positions of the patches in the monomer patchy particle

¹⁵ No se si poner la referencia a la pagina de documentacionhttps://docs.lammps.org/compute_stress_atom.html

patch-patch interaction is modeled with an attractive potential,

$$U_{\text{patchy}}(r_{\mu\nu}) = \begin{cases} 2\varepsilon_{\mu\nu} \left(\frac{\sigma_p^4}{2r_{\mu\nu}^4} - 1\right) \exp\left[\frac{\sigma_p}{\left(r_{\mu\nu} - r_c\right)} + 2\right], & r_{\mu\nu} \in [0, r_c], \\ 0, & r_{\mu,\nu} > r_c, \quad (26) \end{cases}$$

where $r_{\mu\nu}$ is the distance between two patches, σ_p is the diameter of the patches, r_c is the cut distance of interaction set to $1.5\sigma_p$ and $\varepsilon_{\mu,\nu}$ is the interaction energy between the patches. Moreover, the interaction between patches is complemented by a three-body repulsive potential, defined in terms of (26), that provides an efficient bond-swapping mechanism making possible to easily equilibrate the system at extremely low temperatures, while at the same time, reataining the single-bond-per-patch condition[Sciortino, 2017],

$$U_{\text{swap}}(r_{l,m}, r_{l,n}) = w \sum_{l,m,n} \varepsilon_{m,n} U_3(r_{l,m}) U_3(r_{l,n}), \quad r_{l,n} \in [0, r_c],$$
 (27)

where

$$U_3(r) = \begin{cases} 1 & r \in [0, r_{\min}], \\ -U_{\text{patchy}}(r)/\varepsilon_{m,n}, & r \in [r_{\min}, r_c] \end{cases}$$
 (28)

The sum in (27) runs over all triples of bonded patches (patch l bonded both with m and n). $r_{l,m}$ and $r_{l,n}$ are the distances between the reference patch and the other two patches. The parameter $\varepsilon_{m,n}$ is the energy of repulsion and w is used to tuned the swapping (w=1) and non-swapping bonds ($w\gg 1$). The cut off distance r_c is the same as in the potential of interaction between patches, meanwhile the minimum distance r_{\min} is the distance at the minimum of (26), i.e. $\varepsilon_{m,n} \equiv \left| U_{\text{patchy}}(r_{\min}) \right|$. Finally, the energy of interaction between crosslinker patches ($\varepsilon_{\mu^i,\mu^i}$) are set to 0 to allow only crosslinker-monomer and monomer-monomer bonding (figure 2).

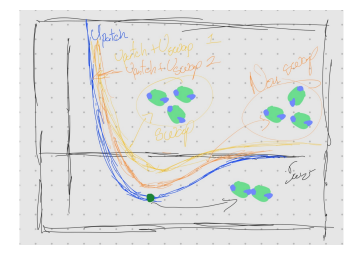


Figure 1: La idea de la figura es poner el pontecial de interacción entre paches y ver el efecto del pontecial de 3 cuerpos cuando w = 1 y cuando $w \gg 1$.

Assembly of the network

We perform molecular dynamics (MD) simulations at fixed temperature $T = kBT/\varepsilon = 0.05$, where kB is the Boltzmann constant. Thanks to such a low temperature, the system tends to maximize the number of bonds.

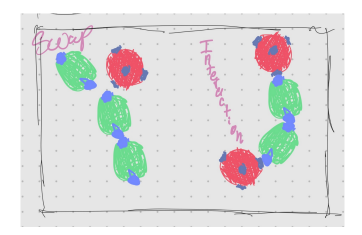


Figure 2: La idea de esta es mostrar las posibles configuraciones (monomero-monomero, monomero-crosslinker y un poco de pontecial de 3 cuerpos)

In addition, owing to the bondswapping mechanism, the system is able to continuously restructure itself, until the large majority of possible bonds are formed. It is important to said that the main difference between the articles cited and the implementation in this htesis are the absence of FENE bonds and the swelling potential.

References

Nikhil Chandra Admal and E. B. Tadmor. A unified interpretation of stress in molecular systems. *Journal of Elasticity*, 100(1):63–143, June 2010. ISSN 1573-2681. DOI: 10.1007/s10659-010-9249-6.

R. Clausius. Xvi on a mechanical theorem applicable to heat. *The London, Edinburgh, and Dublin Philosophical Magazine and Journal of Science*, 40(265):122–127, August 1870. ISSN 1941-5982, 1941-5990. DOI: 10.1080/14786447008640370.

Nicoletta Gnan, Lorenzo Rovigatti, Maxime Bergman, and Emanuela Zaccarelli. In silico synthesis of microgel particles. *Macromolecules*, 50 (21):8777–8786, November 2017. ISSN 0024-9297, 1520-5835. DOI: 10.1021/acs.macromol.7b01600.

Jean Pierre Hansen and Ian R. McDonald. *Theory of simple liquids*. Elsevier Academic Press, 2006.

Jing-Tao Lü, Bing-Zhong Hu, Per Hedegård, and Mads Brandbyge. Semi-classical generalized langevin equation for equilibrium and nonequilibrium molecular dynamics simulation. *Progress in Surface Science*, 94(1):21–40, February 2019. ISSN 0079-6816. DOI: 10.1016/j.progsurf.2018.07.002.

Eric Paquet and Herna L. Viktor. Molecular dynamics, monte carlo simulations, and langevin dynamics: A computational review. *BioMed Research International*, 2015:183918, 2015. ISSN 2314-6133. DOI: 10.1155/2015/183918.

Giorgio Parisi. Statistical field theory. Addison Wesley, 1988.

- R. W. Pastor. Techniques and applications of langevin dynamics simulations. In G. R. Luckhurst and C. A. Veracini, editors, *The Molecular Dynamics of Liquid Crystals*, pages 85–138. Springer Netherlands, Dordrecht, 1994. ISBN 978-94-011-1168-3. DOI: 10.1007/978-94-011-1168-3₅.
- Francesco Sciortino. Three-body potential for simulating bond swaps in molecular dynamics. *The European Physical Journal E*, 40(1):3, January 2017. ISSN 1292-8941, 1292-895X. DOI: 10.1140/epje/i2017-11496-5.
- Valerio Sorichetti, Andrea Ninarello, José M. Ruiz-Franco, Virginie Hugouvieux, Walter Kob, Emanuela Zaccarelli, and Lorenzo Rovigatti. Effect of chain polydispersity on the elasticity of disordered polymer networks. *Macromolecules*, 54(8):3769–3779, apr 2021. ISSN 0024-9297, 1520-5835. DOI: 10.1021/acs.macromol.1c00176.
- Valerio Sorichetti, Andrea Ninarello, José Ruiz-Franco, Virginie Hugouvieux, Emanuela Zaccarelli, Cristian Micheletti, Walter Kob, and Lorenzo Rovigatti. Structure and elasticity of model disordered, polydisperse, and defect-free polymer networks. *The Journal of Chemical Physics*, 158(7): 074905, feb 2023. ISSN 0021-9606, 1089-7690. DOI: 10.1063/5.0134271.
- Arun K. Subramaniyan and C.T. Sun. Continuum interpretation of virial stress in molecular simulations. *International Journal of Solids and Structures*, 45(14–15):4340–4346, July 2008. ISSN 00207683. DOI: 10.1016/j.ijsolstr.2008.03.016. URL https://linkinghub.elsevier.com/retrieve/pii/S0020768308001248.
- Robert J. Swenson. Comments on virial theorems for bounded systems. *American Journal of Physics*, 51(10):940–942, October 1983. ISSN 0002-9505, 1943-2909. DOI: 10.1119/1.13390. URL https://pubs.aip.org/ajp/article/51/10/940/1052035/Comments-on-virial-theorems-for-bounded-systems.
- Johannes M. M. H. Thijssen. *Computational physics*. Cambridge University Press, 2007.
- A. P. Thompson, H. M. Aktulga, R. Berger, D. S. Bolintineanu, W. M. Brown, P. S. Crozier, P. J. in 't Veld, A. Kohlmeyer, S. G. Moore, T. D. Nguyen, R. Shan, M. J. Stevens, J. Tranchida, C. Trott, and S. J. Plimpton. LAMMPS a flexible simulation tool for particle-based materials modeling at the atomic, meso, and continuum scales. *Comp. Phys. Comm.*, 271: 108171, 2022. DOI: 10.1016/j.cpc.2021.108171.
- Aidan P. Thompson, Steven J. Plimpton, and William Mattson. General formulation of pressure and stress tensor for arbitrary manybody interaction potentials under periodic boundary conditions. *The Journal of Chemical Physics*, 131(15):154107, October 2009. ISSN 0021-9606, 1089-7690. DOI: 10.1063/1.3245303. URL https://pubs.aip.org/jcp/article/131/15/154107/316893/General-formulation-of-pressure-and-stress-tensor.

D. H. Tsai. The virial theorem and stress calculation in molecular dynamics. *The Journal of Chemical Physics*, 70(3):1375–1382, February 1979. ISSN 0021-9606, 1089-7690. DOI: 10.1063/1.437577. URL https://pubs.aip.org/jcp/article/70/3/1375/89129/ The-virial-theorem-and-stress-calculation-in.

Heng Wang, Xuhao Li, Lijing Zhao, and Weihua Deng. Multiscale modeling and simulation for anomalous and nonergodic dynamics: From statistics to mathematics. *Fundamental Research*, January 2025. ISSN 2667-3258. DOI: 10.1016/j.fmre.2024.12.024.