Tailoring relaxation dynamics and mechanical memory of crumpled materials by friction and ductility

Eric van Bruggen, Erik van der Linden and Mehdi Habibi

Crumpled sheets show slow mechanical relaxation and long lasting memory of previous mechanical states. By using uniaxial compression tests, the role of friction and ductility on the stress relaxation dynamics of crumpled systems is investigated. We find a material dependent relaxation constant that can be tuned by changing ductility and adhesive properties of the sheet. After a two-step compression protocol, nonmonotonic aging is reported for polymeric, elastomeric and metal sheets, with relaxation dynamics that are dependent on the material’s properties. These findings can contribute to tailoring and programming of crumpled materials to get desirable mechanical properties.

1 Introduction

Crumpled structures are found in many areas of research as well as in a large range of length scales: from DNA packed in viral capsids1–3 and crumpled graphene4 to crumpled paper balls, car wrecks, or even the geological stratum.5 Crumpled systems (CSs) can combine low density structures with surprising mechanical strength and the ability to absorb mechanical energy.6–9 This combination of properties opens doors to use CSs for a variety of applications such as shock absorbers or light weight sandwich panels.10,11 However, in order to rationally design disordered materials, a thorough physical understanding of their unique features is needed. One remarkable physical property observed in CSs is their slow mechanical relaxation and their ability to carry long lasting memory of previous mechanical states. Research on conductors, glasses and polymers12 suggests that slow relaxation dynamics are a generic feature of amorphous materials and ubiquitous. In spite of a significant amount of literature on crumpled materials, a lot of questions about the mechanisms that govern slow stress relaxation are still unanswered. Hitherto, several theoretical models have been suggested to predict force recovery dynamics in CS. The two most popular descriptions being the Kohlrausch function ('stretched exponential'):13–15

\[ F(t) = \frac{a_1}{(t/t_f)^{a_2}} \]  

(1)

and the logarithmic decay function:8,16–18

\[ F(t) = a_2 + b_2 \log(t/t_f) \]  

(2)

Both models are capable of describing a broad range of behaviours in the development of force over time \( F(t) \) over many orders of magnitude19,20 with two free variables \((a_1, b_1)\) and a characteristic time scale \( t/\tau \) further denoted as \( t^* \). The stretched exponential function has been used to model luminescence decay of colloidal quantum dots,21 remnant magnetization in spin glasses22 or structural relaxation in glasses and polymers,23 while the logarithmic function has been widely applied to model slow processes such as decay of current in superconductors,24 conductance relaxation in electron glasses25 and mechanical relaxation of plant roots.12,26

Extensive effort has been put into developing microscopic models that are compatible with eqn (1) or (2). For example Palmer et al.27 show how a stretched exponential function can naturally arise from hierarchically constrained dynamics. Amir et al.,12 provide an interesting discussion on how logarithmic relaxation might be explained from a \( P(\lambda) \sim 1/\lambda \) probability distribution \( P \) of relaxation rates \( \lambda \) that can emerge from various random multiplicative processes. More discussion on the topic can be found in ref. 28 and 29. However, most models still lack a clear physical basis that links the material’s microstructure to the observed long time scale relaxations.33

A remarkable observation is that despite the similarities in stress relaxation for various crumpled systems, the stretched exponential model is preferred to describe crumpled stress relaxation in aluminum systems,13–15 whereas the logarithmic model seems to be more capable of fitting mechanical relaxation of crumpled paper and Mylar.8,16–18

The fact that material properties have a profound effect on the mechanical response of crumpled structures has already been convincingly shown. Many numerical30,31 and experimental6,10,32,33
studies have been dealing with the effect of elastoplasticity in the crumpling process. Matan et al.\(^8\) showed that the force needed to crumple a sheet (\(F\)) obeys a power law that depends on the size (\(D\)) of the crumpled object (\(F \sim D^{-\beta}\)), where the exponent \(\beta\) is material dependent. This was further quantified by Habibi et al.\(^10\) who showed that \(\beta\) is plasticity dependent and can be related to a dimensionless parameter called a foldability index, which is a measure for the ductility in the material. The effect of friction, adhesion and other surface interactions is an often ignored part of the crumpling process.\(^{15,17,34}\) Structural relaxation can also be influenced by the material properties and friction, but to what extent they can determine a crumpled material’s relaxation behaviour is an open question.

In this paper, we present a comparative study where we look into a set of materials with different ductilities and friction coefficients and compare their relaxation behaviour after uniaxial compression. Moreover, we make use of a compaction protocol of Lahini et al.\(^18\) to further characterize and compare the material’s aging behaviour. Our findings can help elucidate the origins of slow stress relaxation in disordered crumpled structures and will provide insight in rational design of future crumpled materials.

2 Experimental procedure

2.1 Sample preparation

A variety of widely available sheet materials was used. Regular printing paper, BOPP (biaxially oriented polypropylene) used as transparent wrapping sheet, rubber, aluminum and brass sheets offered an economic means for studying crumpling phenomena while at the same time representing a wide range in elastoplastic response and frictional coefficients. In addition to using different materials, elastomers were powdered with a very fine corn starch flour, creating a minimal layer of starch particles as to reduce friction and adhesion without altering its bulk properties.

2.2 Friction measurement

To give a measure for both static and dynamic friction coefficients of our sheet materials, a standard rheometer (Anton Paar MCR 302) was transformed into a tribometer with a few simple adjustments.\(^15\) First a glass sphere was attached to a plate. Both glass sphere and bottom plate were covered carefully with the sheet material (Fig. 1a). To start a measurement, the probe is brought into contact by lowering the measuring tool. The normal force is controlled by vertical displacement of the glass sphere. The probe follows a circular trajectory on the bottom plate. Torque \(\tau\), angle of deflection \(\phi\) and normal force \(F_N\) are measured during rotation. Dividing the torque by length of the arm \(r \sim 6\) mm gives us a frictional force \(F = \tau/r\). Following Amonton’s second law of friction, the frictional force is proportional to the magnitude of the normal force. A frictional coefficient (\(\mu\)) can be determined by dividing frictional force by the normal force \(F_F/F_N = \mu\). At small deformations, a peak in frictional force can be observed, that corresponds to the static friction coefficient \(\mu_s\). At larger deformations frictional force reaches a plateau where we find the kinetic friction coefficient \(\mu_k\) (Fig. 1b).

2.3 Stress relaxation procedure

All samples were subjected to uniaxial compression tests in a closed die on a TTC TAXT plus texture analyser. Crumpled samples were prepared by confining a sheet of roughly A4 size by hand in a hollow plexiglass cylinder of diameter and height of 50 mm. Sheets were compressed by a cylindrical plexiglass probe under a constant strain rate of 2 mm s\(^{-1}\) until a pre-set force was reached. The force exerted on the probe was measured over time under constant strain. For every material, a number of samples were taken, varying the initial compaction force within range \(F = 1–200\) N.

2.4 Two-step compression protocol

To look further into the fundamentals of stress relaxation, a procedure, proposed by Lahini et al.\(^18\) was followed, i.e. a two-step compression protocol. After initial compression from height \(h_1\) to \(h_2\) (with force \(F_1\) to \(F_2\)), strain was kept constant for a specific waiting time \(\tau_w\) after which the probe was lifted to \(h_3\) with a strain rate of 5 mm s\(^{-1}\) towards a new force \(F_3\) for which \(F_1 < F_3 < F_2\) (Fig. 2). The normal force was continuously measured over time. \(h_1\) was set in a way that \(F_1 \sim 0.1\) N
was obtained, while $F_2$ varied in between 20–50 N and $F_3$ between 1–45 N.

3 Results and discussion

Typical stress relaxation curves for the tested materials are shown in Fig. 3 that show relaxation after initial compaction at $F \sim 25$ N. All studied samples show slow logarithmic-like stress relaxation with a steep decrease in magnitude of stress for short times $<20$ s and a slow but persistent decrease in the magnitude of stress for greater time scales. Relaxation data were fitted with eqn (1) and (2). All data sets show (at least slight) non-stochastic deviations from both the logarithmic and exponential model. Note that for aluminium (Fig. 3d) first a slower relaxation regime is observed, followed by a faster (logarithmic) relaxation, while for the BOPP, printing paper and brass (Fig. 3a–c) relaxation seems to slightly slow down for larger time scales. In line with published literature, it can be seen that relaxation of crumpled aluminium is very well described by a stretched exponential model, while paper and BOPP show a trend that closely resembles logarithmic decay.

![Stress relaxation curves](image)

We can expand the logarithmic model by including physical cutoff rates $\lambda_{\text{max}}$ and $\lambda_{\text{min}}$. Eqn (2), was based on the assumption that during our experiments time scales far from the inverse of the extreme relaxation rates are considered: $1/\lambda_{\text{max}} \ll t < 1/\lambda_{\text{min}}$. If this is not the case, we can revert to the more general equation:36

$$F(t) = F_0/[E_1(\lambda_{\text{min}}t) - E_1(\lambda_{\text{max}}t)]$$  \hspace{1cm} (3)

That is the sum of two exponential integrals $E_1$ which enables us to incorporate a minimum and maximum rate of relaxation that can account for a plateau at very small or very large time scales. In Fig. 3 fits to eqn (3) are plotted. Especially for the brass sample (Fig. 3c), eqn (3) is very able to describe the observed relaxation behaviour, indicating that the examined time scales are not far from $1/\lambda_{\text{min}}$ and thus responsible for the flattening of the curve near $t \sim 10^5$. For the aluminium sample flattening near $t \sim 10^9$ occurs (Fig. 3c), indicating that examined time scales are not far from $1/\lambda_{\text{min}}$.

However, applying finite cutoff rates cannot account for the bended shape that is apparent for the rubber sheets (Fig. 3e) and to lesser extent can also be observed for both BOPP and paper samples (Fig. 3a and b). Instead, these apparent curved lines bare remarkable similarities to the results obtained by Lahini et al. for elastic foams18 i.e. two logarithmic regimes can be distinguished and in a similar fashion, we can use a superposed logarithm to fit our data:

$$F = a + b \log(t^*) + c \log(t^* - t_o^*)$$  \hspace{1cm} (4)

Eqn (4) provides a solid fit to all our data and plots of the residuals show a stochastic spread in a large time window. Non-stochastic deviation can still be seen at short time scales $t < 20$ s which we attribute to the finite strain rate during compaction, the associated premature relaxation and the earlier discussed finite maximum cutoff rate.

3.1 Relaxation rate

The applicability of eqn (4) to all our tested samples, allows us to do an inter material comparison of its plotting parameters set out against a range of initial compaction forces $F_{(t=1s)}$ (the force measured 1 second after compaction) to which our samples were subjected (see Fig. 4a).

![Stress relaxation over time](image)

We report a linear dependence of fitting parameter $b$ on the initial compaction force $F_{(t=1s)}$. Notice that $b/F_{(t=1s)}$ is constant for all initial packing forces and protocol independent and thus reflects an intrinsic material property. Our results are comparable to the findings for relaxation of a single fold in polymeric sheets by Thiria et al.37 who reported an intrinsic material constant in the single folding of sheets, but to our knowledge, it is the first time such a constant is proposed for crumpled materials.

To systematically investigate how the material dependent relaxation constant for CSs $b/F_{(t=1s)}$ is affected by the sheet's material properties, a dimensionless foldability index $i_i$ is employed. $i_i$ introduced by Habibi et al. gives a measure for the material's ductility or behaviour beyond yielding.10 In Fig. 4b constant $b/F_{(t=1s)}$ is plotted as a function of the foldability index. For most tested materials, an increase in ductility is accompanied
by a higher relaxation constant. However, this observation fails for the starch powdered rubber material that has an $i$ comparable to the untreated rubber sheet, but a distinctly different response, indicating that surface phenomena are at play.

### 3.2 Effect of friction

Using the setup of Fig. 1, friction coefficients for all sheet materials were measured. Results are shown in Table 1. The determination of the friction coefficient for rubber was complicated by stick-slip and adhesive phenomena. For adhesive materials, frictional force depends nonlinearly on the applied load and the restriction $F_t = 0$ at $F_N = 0$ is not valid, where $F_t$ and $F_N = 0$ are friction and normal forces respectively. Since there is no linear dependence, the friction coefficient is ill-defined. Nevertheless, we included the calculated value in Table 1 as it provides insight in the rubber's friction-like behaviour.

After a coating of corn starch flour is applied to the rubber elastomer, stick-slip phenomena are no longer observed. Moreover, the apparent friction coefficient is considerably lower, confirming that indeed, addition of starch is an effective way to reduce adhesion and lower the friction coefficient, thus allowing us to control the surface properties of our rubber material.

Returning to the simple relaxation experiment: in Fig. 3 we observed that addition of the very fine corn starch powder has a profound impact on the stress relaxation of rubber. Reduction of friction through addition of corn starch changes the shape of the relaxation data from a bend line in the semi-log plot to a straight line that can be very well described by a single logarithmic model.

A double logarithm can be used to describe the bend relaxation curves, reminiscent of the shapes obtained by Lahini et al. for relaxation of the polymeric foams. That crumpled systems showing foam-like properties, has already been convincingly pleaded by Bouaziz et al., but relaxation in polymeric foams has shown markedly different behaviour from crumpled polymeric sheets. Moreover, it has been argued that friction has only a small effect on the crumpling process. However, in present literature only materials with friction coefficients smaller than one and frictional forces that vanish near zero load have been considered. The examined rubber sample now gives us a means to investigate samples with higher friction coefficients and a finite friction near zero load due to adhesive contacts. One can hypothesize that due to these adhesive contacts, inter-layer movements during compaction and relaxation are reduced to a minimum, or even are completely absent resulting in the foam-like relaxation behaviour of the adhesive sheet.

Furthermore, under an equal compaction force, untreated rubber has a considerably faster relaxation rate than the starch powdered samples (Fig. 4a). A similar observation was made by Cottrino et al.: applying a higher strain rate resulted in a faster relaxation rate, and it has been explained by the inability of the internal micro-structure to properly rearrange itself.

![Fig. 4](a) Material dependent relaxation rates, with rate $b$ from equation $F = a + b \log(t^*) + c \log(t^* + t_0^*)$. Dashed lines are least square fits to the data forced through the origin. Increased initial compaction force $F_{(t=1s)}$ leads to an increased relaxation rate. (b) The material parameter $b/F_{(t=1s)}$ is plotted against a dimensionless foldability index (results measured using the procedure of Habibi et al.).

### Table 1 Friction coefficients for material samples

| Sheet material | Dynamic friction $\mu_k$ | Static friction $\mu_s$ |
|----------------|--------------------------|-------------------------|
| Aluminium      | 0.17 ± 0.01              | 0.21 ± 0.03             |
| Paper          | 0.29 ± 0.02              | 0.37 ± 0.03             |
| Rubber         | 2.71 ± 0.99              | 3.36 ± 1.65             |
| Rubber (coated)| 0.36 ± 0.04              | 0.41 ± 0.04             |
| BOPP           | 0.50 ± 0.03              | 0.53 ± 0.05             |

* No ‘true’ friction coefficient (see discussion).

![Fig. 5](Examples of nonmonotonic stress relaxation – depending on the compaction protocol, peak time and peak force change. Shown are coated rubber samples with $F_1 = 0.1$ N, $F_2 = 40$ N, while $F_3 = 12$ N (red hexagons), $F_3 = 18$ N (blue circles) and $F_3 = 22$ N (yellow diamonds).
In the same train of thought, we can assume that due to increased surface interactions, crumpled structures are not able to rearrange properly during compaction. This leads to an increased storage of energy in the form of ridges and vertices during the crumpling process.\textsuperscript{41} Consequently, after compaction a system high in friction is further away from its equilibrium state than its frictionless counterpart, which can explain the larger relaxation rate of the adhesive rubber sample at similar initial compaction forces (Fig. 4).

### 3.3 Non monotonic aging

In Fig. 5 stress evolution in time after the two step compression protocol is plotted. After decompression from $F_2$ to $F_3$, non-monotonic stress relaxation is apparent for all tested materials:

In the short time towards the peak, contributions of the fast relaxation modes are prevalent, thus leading to the observed peak in normal force.

If the waiting time $t_{w}$ at $F_2$ is varied, while other conditions are kept constant, it is found that peak time $t_p$ depends linearly with $t_w$ and a slope $t_p/t_w$ can be extracted. The steepness of the normal force shows a steep increase for the first seconds to minutes, then reaches a well-defined peak at a time $t_p$, and after, resumes slow decay. This means that before and after the peak, sets of data-points exist for which the strain, force and all other macroscopic observable are identical, however the direction of the force evolution is qualitatively different, indicating that force evolution cannot be predicted based on these observable parameters on its own. Instead, one needs to take into account the history of the system or deal with the system’s ‘memory’.

The occurrence of non-monotonic relaxation after decompression can be explained as a consequence of the non-equilibrium states of the system during relaxation.\textsuperscript{18} Starting point is the assumption that the system consists of an ensemble of relaxation modes, covering a wide range of relaxation rates that is at rest in an equilibrium state $E_1$ before compaction (at $F_1$). By moving the piston to $F_2$, a new equilibrium state $E_2$ is forced upon the system, to which all relaxation modes move towards while contributing to the total force opposite of the compression direction. Over time, this force is decreasing as more and more relaxation modes arrive in close vicinity of $E_2$. By introducing a decompression step to $F_3$, we force a third equilibrium state $E_3$ upon the system for which $E_1 > E_3 > E_2$. We now see that the fast relaxation modes which were already in close vicinity to $E_2$ and thus $E_3$, will now possess an sign compared to that of the slow relaxation modes that were still close to the original equilibrium state $E_1$ and for which $E_2$ is valid. In the short time towards the peak, contributions of the fast relaxation modes are prevalent, thus leading to the observed peak in normal force.

Fig. 6 Ratio $t_p/t_w$ against the ratio of forces $(F_2 - F_3)/(F_3 - F_1)$ for metal, polymeric and elastomeric crumpled systems. Dashed line with $(t_p/t_w) = 1$ provides a reasonable fit for all data.

![Graph showing ratio $t_p/t_w$ against the ratio of forces.](image)

Fig. 7 (a) Normalized peak force plotted against the peak time $t_p$ – the plastic aluminium and brass behave markedly different from the polymeric compounds. (b) Same as in (a) on linear scale. Dashed lines represent fits by robust bi-square fitting, restricted to the origin. (c) Slopes of (b) plotted against the material dependent relaxation $b/F_{t=1s}$. (d) Foldability index $[i_f]$ against $b/F_{t=1s}$. 

![Graph showing various data points and fitted lines for different materials.](image)
slope $t_p/t_w$ in its turn is dependent on the compaction protocol. If we assume that the compaction force is proportional to an intrinsic equilibrium state of the crumpled system, it can be derived that $t_p/t_w = (F_2 - F_1)/(F_3 - F_1)^3$. In Fig. 6, ratio $t_p/t_w$ is plotted against the force ratio $(F_2 - F_1)/(F_3 - F_1)$. Indeed we find that $[(F_2 - F_1)/(F_3 - F_1)][t_p/t_w] \approx 1$ independent of the used material and without the need for any fitting parameters. This implies that non-monotonic aging is a generic feature of crumpled systems, governed by universal relaxation dynamics.

In order to compare non-monotonic dynamics of different experiments, measured force is normalized by $F_3$ (at $t = 1$ s after decompression). Plotting the normalized forces at relaxation constant. With this paper, we contribute to an increased understanding of crumpled systems that can help in further tailoring and programming of these disordered materials.

**Conflicts of interest**

There are no conflicts of interest to declare.

**Acknowledgements**

We acknowledge the anonymous reviewers for the constructive comments. MH acknowledge funding from the Netherlands Organization for Scientific Research through NWO-VIDI grant no. 680-47-548/983.

**Notes and references**

1. P. K. Purohit, J. Kondev and R. Phillips, *Proc. Natl. Acad. Sci. U. S. A.*, 2003, **100**, 3173–3178.
2. N. Stoop, J. Najafi, F. K. Wittel, M. Habibi and H. Herrmann, *Phys. Rev. Lett.*, 2011, **106**, 214102.
3. J. Najafi, N. Stoop, F. Wittel and M. Habibi, *Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys.*, 2012, **85**, 061108.
4. J. A. Baimova, E. A. Korznikova, S. V. Dmitriev, B. Liu and K. Zhou, *Rev. Adv. Mater. Sci.*, 2014, **39**, 69–83.
5. M. P. Golombok, F. S. Anderson and M. T. Zuber, *J. Geophys. Res.*, 2001, **106**B21, 811–823.
6. S. Deboeuf, E. Katzav, A. Boudaoud, D. Bonn and M. Adda-Bedia, *Phys. Rev. Lett.*, 2013, **110**, 104301.
7. M. B. Amar and Y. Pomeau, *Proc. R. Soc. London, Ser. A*, 1997, 729–755.
8. K. Matan, R. B. Williams, T. A. Witten and S. R. Nagel, *Phys. Rev. Lett.*, 2002, **88**, 7–10.
9. M. R. Shaebani, J. Najafi, A. Farnudi, D. Bonn and M. Habibi, *Nat. Commun.*, 2017, **8**, 15568.
10. M. Habibi, M. Adda-Bedia and D. Bonn, *Soft Matter*, 2017, **13**, 4029–4034.
11. M. Mirzaali, M. Habibi, S. Janbaz, L. Vergani and A. Zadpour, *Sci. Rep.*, 2017, **7**, 13028.
12. A. Amir, Y. Oreg and Y. Imry, *Proc. Natl. Acad. Sci. U. S. A.*, 2012, **109**, 1850–1855.
13. A. S. Balankin, M. A. Cruz, L. A. Caracheo, O. S. Huerta, C. D. Rivas, C. L. Martinez, D. S. Ochoa, L. M. Ruiz, S. M. Gutiérrez, J. P. Ortiz and M. P. Ortiz, *J. Mater. Sci.*, 2015, **50**, 4749–4761.
14. S. Cottrino, P. Vivies, D. Fabregue and E. Maire, *Acta Mater.*, 2014, **81**, 98–110.
15. R. F. Albuquerque and M. A. F. Gomes, *Phys. A*, 2002, **310**, 377–383.
16. A. S. Balankin, O. Susarrey Huerta, F. Hernández Méndez and J. Patiño Ortiz, *Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys.*, 2011, **84**, 1–10.
17. I. Dierking and P. Archer, *Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys.*, 2008, **77**, 1–5.
18. Y. Lahini, O. Gottesman, A. Amir and S. M. Rubinstein, *Phys. Rev. Lett.*, 2017, **118**, 1–6.
19 D. C. Elton, 2018, arXiv:1808.00881.
20 A. Amir, Y. Oreg and Y. Imry, Phys. Rev. B: Condens. Matter Mater. Phys., 2008, 77, 165207.
21 E. N. Bodunov, Y. A. Antonov and A. L. Simões Gamboa, J. Chem. Phys., 2017, 146, 114102.
22 R. V. Chamberlin, G. Mozurkewich and R. Orbach, Phys. Rev. Lett., 1984, 52, 867–870.
23 B. Sturman, E. Podivilov and M. Gorkunov, Phys. Rev. Lett., 2003, 91, 176602.
24 A. Gurevich and H. Kupfer, Phys. Rev. B: Condens. Matter Mater. Phys., 1993, 48, 6477–6487.
25 T. Grenet, Phys. Status Solidi C, 2004, 1, 9–12.
26 K. Büntemeyer, H. Lüthen and M. Böttger, Planta, 1998, 204, 515–519.
27 R. G. Palmer, D. L. Stein, E. Abrahams and P. W. Anderson, Phys. Rev. Lett., 1984, 53, 958–961.
28 J. C. Phillips, Rep. Prog. Phys., 1996, 59, 1133–1207.
29 B. Sturman, E. Podivilov and M. Gorkunov, Phys. Rev. Lett., 2003, 91, 1–4.
30 T. Tallinen, J. A. Åström and J. Timonen, Nat. Mater., 2009, 8, 25–29.
31 S. F. Liou, C. C. Lo, M. H. Chou, P. Y. Hsiao and T. M. Hong, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2014, 89, 1–5.
32 O. Bouaziz, J. P. Masse, S. Allain, L. Orgeas and P. Latil, Mater. Sci. Eng., A, 2013, 570, 1–7.
33 Y. C. Lin, J. M. Sun, H. W. Yang, Y. Hwu, C. L. Wang and T. M. Hong, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2009, 80, 066114.
34 K. P. Mota and P. M. C. de Oliveira, Phys. A, 2008, 387, 6095–6104.
35 B. Weber, Y. Nagata, S. Ketzetzi, F. Tang, W. J. Smit, H. J. Bakker, E. H. G. Backus, M. Bonn and D. Bonn, J. Phys., Lett., 2018, 9, 2838–2842.
36 A. Amir, S. Borini, Y. Oreg and Y. Imry, Phys. Rev. Lett., 2011, 107, 186407.
37 B. Thiria and M. Adda-Bedia, Phys. Rev. Lett., 2011, 107, 1–4.
38 G. A. D. Briggs and B. J. Briscoe, Wear, 1979, 57, 269–280.
39 Y. Fukahori, P. Gabriel and J. J. Busfield, Wear, 2010, 269, 854–866.
40 B. N. J. Persson, J. Phys.: Condens. Matter, 2008, 20, 312001.
41 T. Tallinen, J. A. Åström and J. Timonen, Comput. Phys. Commun., 2009, 180, 512–516.