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Extracting, quantifying, and comparing dynamical and biomechanical properties of living matter through single particle tracking

This Perspective article describes soft and living matter from a physical stance, highlighting experimental and theoretical techniques relevant for the investigation of forces, materials properties, transport and emergent organisation phenomena. The Perspective addresses scientists in physics and chemistry but is also meant to serve as a starting point for researchers in the life sciences interested in the implementation of biophysical methods.







### **PERSPECTIVE**

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# Extracting, quantifying, and comparing dynamical and biomechanical properties of living matter through single particle tracking

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A panoply of new tools for tracking single particles and molecules has led to an explosion of experimental data, leading to novel insights into physical properties of living matter governing cellular development and function, health and disease. In this Perspective, we present tools to investigate the dynamics and mechanics of living systems from the molecular to cellular scale via single-particle techniques. In particular, we focus on methods to measure, interpret, and analyse complex data sets that are associated with forces, materials properties, transport, and emergent organisation phenomena within biological and soft-matter systems. Current approaches, challenges, and existing solutions in the associated fields are outlined in order to support the growing community of researchers at the interface of physics and the life sciences. Each section focuses not only on the general physical principles and the potential for understanding living matter, but also on details of practical data extraction and analysis, discussing limitations, interpretation, and comparison across different experimental realisations and theoretical frameworks. Particularly relevant results are introduced as examples. While this Perspective describes living matter from a physical perspective, highlighting experimental and theoretical physics techniques relevant for such systems, it is also meant to serve as a solid starting point for researchers in the life sciences interested in the implementation of biophysical methods.

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### I. Introduction

Living organisms are incredibly complex systems, with different aspects being well described by physics, chemistry, or biology; however, no single classical discipline is sufficient to successfully describe the dynamics and function of the entire organism, including health and disease. To truly understand how living creatures function, a thorough investigation combining tools and insights from different disciplines is crucial. Stressing this point is the fact that despite their biochemical composition, the behaviour of all living beings is in large part governed by the laws of physics and important insight into how biological systems function can be gained through the analysis of their physical properties.

Beyond advancing our understanding and treatment of disease, biophysics has been instrumental in advancing our understanding of biological systems, in particular the physical phenomena governing certain biological processes. Significant progress has been made over the past thirty years in studying cells and cell compartments, either at the individual molecule, cell, or tissue levels. These advances have demonstrated that the behaviour of individual biological systems may differ from the average collective behaviour of these same individuals in a group. One example is the structure of the cells cytoskeleton, an interconnected network of proteins and filaments that link a cell's nucleus to its membrane.1 This structure's mechanical characteristics are not only a consequence of the mechanical properties of the individual biopolymers, but also react to extracellular stresses, demonstrating a collective cellular response to the properties and signals of the external environment. Recent studies2 have clearly shown that biophysical

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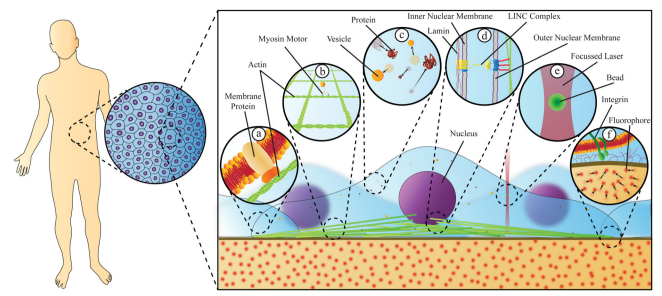


Fig. 1 Physical properties of cells originate from their molecular components and are responsive to both intracellular and extracellular events. Living organisms are composed of tissues, which themselves are composed of cells in close proximity to one another. Cells living in such tissues transmit forces intracellularly via a number of mechanisms. (a) Forces from the extracellular environment are sensed by cells through membrane proteins which relay the force to the cytoskeleton through attachments, triggering biochemical responses. (b) The cytoskeleton, composed of protein filaments such as actin polymers and/or microtubules, can transmit physical forces throughout the cell, serve to correctly position the nucleus in the cytoplasm, including during cell division, and allows cells to adjust their material properties, potentially in response to their exterior environment. Protein motors, such as myosin V, actively transport materials along the cytoskeleton filaments. (c) Cargo, such as vesicles or proteins, can diffuse or be actively transported by molecular motors through the crowded cytoplasm. (d) Physical forces can be transmitted through the nuclear membrane via the LINC complex. This complex is attached to the cytoskeleton and pierces both nuclear membranes, thus transmitting forces to the lamin proteins that act as nuclear structural components. (e and f) Force measurement techniques tracking the movement of micro- or nano-scopic particles can be used to quantify material properties inside cells or measure forces exerted by cells. Examples of such techniques include optical tweezers (e) and traction force microscopy (f).

methods, based on the observation of single particles, can be used to measure not only the forces exerted on cells, but also how cells respond to these forces, or even exert microforces within the cells, thus uncovering differences in disease states with clinical implications. Diseases themselves have a strong impact on the biomechanical properties of cells<sup>3</sup> and on intracellular material properties and trafficking.4 Fig. 1 gives an overview of relevant physical phenomena within cells, where dynamics and material properties play a tremendous role, and it also depicts two biophysical methods capable of quantifying such properties.

It is challenging to detect and quantify the complex physics occurring in and around living cells. An emerging and promising way to address this challenge is by using single particle techniques to directly measure biomechanical properties of living cells, e.g., their microrheological properties. This branch of rheology applies microscopic means of measuring the mechanics both within and without the cellular environment, thus providing critical information on biophysical phenomena affecting cell behaviour and function. As the wider field of single molecule tracking in biology and biophysics has already uncovered significant amounts of important information, as exemplified by 3D single-molecule active real-time tracking (3D-SMART) to measure DNA or protein diffusion,<sup>5</sup> here we narrow the discussion to single and multiple particle tracking and their relation to rheology and particle diffusion. A plethora of particle-based methods, experimental and theoretical, exist

to measure, analyse, and quantify the dynamics and mechanics of living systems, each with its own strengths and limitations. This range of choice can be overwhelming, the terminology imprecise, and, in the worst case, interpretations may be flawed if one is not aware of the limitations of the methods used. Here, we review particle-based methods, experimental and theoretical, relevant for studying soft- and living matter with the aim of providing enough information to enable both physical and life scientists to critically apply and evaluate the methods. We not only introduce passive particle tracking and microrheology methods, but also methods which actively disturb biological systems by mechanical forces and, through the system's response, provide information about the biomechanical properties of the system. In this way, we aim to contribute to the fields of biology and physics, and encourage further quantitative exploration and understanding of the living organism.

# II. Measuring biomechanical forces

Cells and tissues are subject to biomechanical forces, both from external sources and from within. These biomechanical forces play an immense role in controlling cell adhesion, tissue morphogenesis, and development,6 as well as in disease, such as the spreading of cancer. Cells can sense the stiffness of their surrounding environment through a signalling pathway called mechanotransduction.7 This mechanical sensing can be

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mediated by integrins, found in the membrane of cells.8 In addition, cells adapt their mechanical properties to their environment9 and the differentiation of stem cells is directed by the mechanical properties of the extracellular environment.<sup>10</sup> The mechancotransductive signalling pathway is not only sensitive to force amplitude, but also to direction: for instance, it was demonstrated that fibroblast cells could detect an oscillating force applied to the scaffold on which they were grown, orienting themselves perpendicular to the direction of oscillation.<sup>11</sup> Hence, force is a decisive factor in biological decision-making, and presumably the impact of mechanobiology has expanded during evolution.12 Despite the progress made in understanding the link between external mechanical forces and cellular processes, there remain many open questions to be answered.<sup>13</sup> Therefore, the measurement of biomechanical forces, preferably in a non-invasive manner and inside living organisms, is an important step forward in understanding proper cell function and control mechanisms.

To understand the nature and role of biomechanical forces, specific methods are needed that optimally allow for: (i) force quantification, (ii) application in vivo in a nearly non-invasive manner, and (iii) probing across a wide range of time scales. One can distinguish between forces measured at the surface of,

or within a cell, tissue or organism, necessitating penetrating or observing through a barrier such as the cellular membrane.

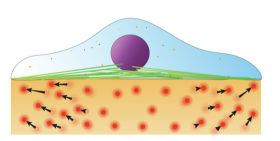
Advances in technology have led to the creation of numerous means of measuring the microrheology of living cells and tissues using single particle techniques.

### A. Traction force microscopy

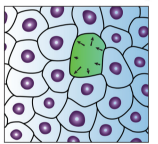
Traction force microscopy (TFM), see Fig. 2(a), is a useful method to quantify cell-generated forces. 14-16 TFM allows for measurement of the forces that cells exert on a substrate. This is done by tracking the displacement of fluorescent marker beads in the substrate, most often in the form of a soft, elastic matrix, on which cells are grown, thus allowing direct observation of cellular forces exerted on the matrix. The matrix must be optically transparent enough to allow for the observation and tracking of the embedded fluorescent markers. Common scaffolds include polyacrylamide (PAA) or polydimethylsiloxane (PDMS), though a wide variety of biocompatible materials, such as hydrogels, are used.

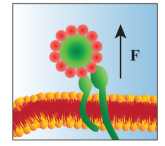
Traction forces exerted by the cells can be derived mathematically by first expressing the force-displacement relation via a Green's function, in the most general case expressed as a tensor. 14 While valid only for small forces, this regime is typically

### a) Traction force microscopy



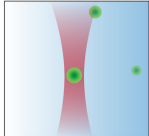
### c) Biofunctionalised microdroplets e) Magnetic Tweezers





b) Traction force microscopy - Pillars





f) Atomic force microscopy

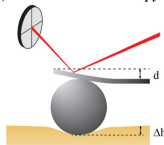


Fig. 2 Single-particle tools to guantify forces in biological systems. (a and b) Traction force microscopy (TFM) measures cellular forces exerted on a substrate either by observing the movement of fluorescent molecules embedded in the substrate (a), or by the bending of columns under the cells (b). (c) Biofunctionalised microdroplets quantify forces exerted on them by surrounding cells by measuring their deformation. Using mathematical models and computational methods, the strains on individual cells or cellular components can be calculated. (d) Optical tweezers (OT) are formed by tightly focusing a laser beam, thus being able to trap a particle, an organelle, or a molecule. OT are an excellent and nearly non-invasive tool to measure forces and probe material properties at the nano- or micron-scale, even inside living organisms. (e) Magnetic tweezers (MT) apply a magnetic field to magnetic particles inserted into cells. The resultant motion of the particle can be used to either exert or measure forces, or to apply a twisting motion. (f) Atomic force microscopy (AFM) brings a microscopic cantilever close to a material surface. Using a photodiode to measure the movement of a laser deflected off the back of the cantilever, it is possible to quantify nanoscopic movements of the cantilever. Affixing a bead on the cantilever allows for force measurements in soft matter.

suitable for the forces exerted by cells on their substrate. This approach is commonly referred to as an inverse problem: while the resultant marker displacement on the substrate is measurable, the applied cellular forces are unknown. Mathematically, the displacement field  $\mathbf{u}(\mathbf{r})$  is related to the traction force field  $\mathbf{T}(\mathbf{r}')$  through the Green's function tensor  $\mathbf{G}(\mathbf{r},\mathbf{r}')$ 

$$u(r) = \int \underline{\underline{\underline{G}}}(r,r') T(r') \text{d}r'. \tag{1}$$

1. Boussinesq approximation. The simplest way to tackle the inverse problem and solve for the Green's function is to use a thick substrate. Under these conditions, cellular traction forces decay before reaching the substrate edge, allowing for the use of the Boussinesq solution. Alternatively, if a thick substrate is not experimentally feasible, the Green's function for finite substrate thickness can be used. It is a substrate thickness can be used.

In many cases, TFM is performed in 2-dimensional systems where the Green's function can be expressed as a convolution:  $\underline{\underline{G}}(\mathbf{r},\mathbf{r}') = \underline{\underline{G}}(\mathbf{r}-\mathbf{r}')$ . For a sufficiently thick isotropic elastic material with Young's modulus E and Poisson ratio  $\nu$ , the Boussinesq solution for such a system gives

$$\underline{\underline{\mathbf{G}}}(\mathbf{r}) = \frac{(1+\nu)}{\pi E \|\mathbf{r}\|^3} \begin{pmatrix} \nu x^2 + (1-\nu) \|\mathbf{r}\|^2 & \nu xy \\ \nu xy & \nu y^2 + (1-\nu) \|\mathbf{r}\|^2 \end{pmatrix}, \quad (2)$$

where  $\|\mathbf{r}\| = \sqrt{x^2 + y^2}$ . The Green's function given in eqn (2) is invertible, permitting the inversion of eqn (1) to solve for the traction forces applied by cells on the substrate.

Historically, to measure the cellular forces using TFM, two images of the substrate are made: one with the cell attached, showing bead displacement caused by the cells, then a second of the same area where the cell has been detached chemically, usually via trypsin which degrades cell adhesions. 19 Computer software is then used to compare the two images and determine the bead displacement  $\mathbf{u}(\mathbf{r})$ . Modern techniques avoid taking two images by printing a grid pattern of beads into the substrate, allowing bead displacement to be directly determined using only one image.  $^{20,21}$  As the parameters E and  $\nu$  are already known for the substrate used, T(r') can be readily calculated using computational software. TFM has been successfully used to measure stresses/forces in: rat cardiac cells grown on substrates mimicking normal and diseased tissue, 22 human induced pluripotent stem cells under a variety of conditions, 23 and mouse embryonic fibroblasts (MEFs) under applied shear stresses.<sup>24</sup>

Alternatively, traction forces can be calculated from the bending of micro-scale $^{25,26}$  or nano-scale $^{27}$  pillars grown on cell culture substrates, see Fig. 2(b). Similar to TFM, described above, the positions of the fluorescent pillar tips on which cells are grown are tracked over time. These pillars are assumed to be elastic, and deformations assumed to be small, allowing for the application of linear elastic theory. In this case, the magnitude of cellular forces applied at each tip may be calculated from the tip displacement via

$$F = \frac{3EI}{L^3} \Delta r,\tag{3}$$

where E is the material's Young's modulus, I is the moment of inertia of the pillar, L is the length of the pillar, and  $\Delta r$  is the measured displacement of the pillar tip. Different pillar geometries have been used, from circular<sup>25,26</sup> to hexagonal,<sup>27</sup> though other geometries could be employed, requiring only a change in the equation for the geometry-dependent I. Using these pillars, both the forces exerted by cells<sup>28</sup> and cell layers can be detected.<sup>29</sup> Micropillar studies have also been used to demonstrate which nucleo-cytoskeletal proteins are responsible for nuclear deformation.<sup>30</sup> Additionally, as the tip's radius a scales with E via  $E \propto a^4$ , the sensitivity range of this technique is quite vast, permitting the detection of a wide-range of forces.

### B. Deformation microscopy

A similar technique called deformation microscopy has been used to quantify forces at the tissue level, which is also particularly suited for investigations at longer time scales. Here, the deformation of biofunctionalised microdroplets, see Fig. 2(c), can be recorded and used to measure forces. <sup>31</sup> In this technique, the curvature of a droplet surrounded by cells can be related to the forces these cells apply to the droplet. Specifically, the stress  $\sigma$  applied by a cell in the direction normal to a droplet with spherical coordinates  $\theta$  and  $\phi$  is

$$\sigma(\theta, \phi) = p_e^{\prime} - p_i^{\prime} + 2\gamma H(\theta, \phi), \tag{4}$$

where  $p_{\rm e}^{'}$  and  $p_{\rm i}^{'}$  are the external and internal hydrostatic pressures of the droplet,  $\gamma$  is the droplet's interfacial tension, and H is the droplet's local mean curvature at the surface. 31 These microdoplets must be of a similar size to cells, possess ligands to which cells can adhere, and be fluorescently labelled in order for their shape to be measured. Furthermore, only partially embedded microdroplets can be used to calculate both isotropic and anisotropic forces exerted by cells as this is the sole means of measuring the tissue pressure: fully embedded microdroplets only permit anisotropic force measurements to be made.<sup>31</sup> Thus, deformation microscopy is most useful when applied to measure forces in cultured cells or epithelial tissues, though in certain cases, anisotropic force measurements in cells can provide some information, as is the case for spatial inhomogeneities caused by cell movement during embryonic development. Microdroplets have been useful in spatiotemporal studies of developing tissues, specifically to measure forces in tissues during embryogenesis. 32 For more information on this technique, we refer the interested reader to Gómez-González, et al.,16 or Campás, et al.31

Deformation microscopy has shown potential for intracellular and intranuclear force mapping, <sup>31,33</sup> shedding light on both the biomechanical processes in and around cells, and on local mechanical stresses. Combining deformation microscopy with high-resolution microscopy techniques has allowed for dynamic strain mapping in individual cells, such as for contracting cardiomyocytes placed on either soft or stiff polydimethylsiloxane (PDMS) substrates.<sup>33</sup> Such a combination is particularly useful when seeking to measure forces during cell migration. However, the possibility of interplay between these

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detection methods and observed cell behaviour cannot be ignored. TFM, for example, employs very soft hydrogel matrices with Youngs moduli in the kPa regime; at this stiffness, the substrate material itself might interfere with the cell due to mechanosensory processes.<sup>34</sup> Another nearly non-invasive means of investigating local mechanical disturbances in biological systems is by focused-light-induced cytoplasmic streaming (FLUCS).35 FLUCS can be used to observe active microrheology, similar to intracellular viscoelasticity characterisation methods.

### C. Optical tweezers

Optical tweezers (OT), see Fig. 2(d), are a technology based on a tightly focused laser beam and with the capacity to reach inside living cells, <sup>36</sup> or even inside living organisms. <sup>37</sup> OT can actively apply controlled forces or perform quantitative force measurements in a nearly non-invasive manner. To minimise physiological damage, however, the laser power should be kept as low as possible and the wavelength in the biological transparency window. An optical trap exerts a harmonic force on the trapped object:

$$\mathbf{F}_{\text{trap}} = -\kappa_i \mathbf{x},\tag{5}$$

where **x** is the distance from the equilibrium position and  $\kappa_i$  is the spring constant characterising the optical trap in the idirection, typically determined through a calibration measurement. Notice that eqn (5) holds in each translational direction and the elements of the tensor  $\kappa$  usually differ from one another.

Optical tweezers based on a tightly focused near-infrared Gaussian laser beam can trap particles with a larger index of refraction than the surrounding media, for instance, micronsized polystyrene spheres (to which molecules can be specifically attached), metallic nanoparticles, or endogenously occurring organelles such as lipid granules.<sup>38</sup> The positions visited by the trapped particle, x, can be found by video microscopy or by focusing the back-scattered light from the laser beam onto a quadrant photodiode, with the latter method having the great advantage of higher time resolution (up to MHz) and easier data acquisition.

One can also estimate the forces without prior calibration of the trap stiffness using the so-called momentum method<sup>39</sup> by collecting and analysing the forward scattered light; this has the advantage that, e.g., the geometry of the particle does not need to be known. Recently, optical tweezers were combined with light-sheet microscopy to measure 100 pN-range tension at cell-cell junctions.40 The pN forces exerted by standard optical tweezers can even induce large-scale cellular reactions, such as calcium transients in neuroblastoma cells.<sup>41</sup>

In practice, optical tweezers are often implemented by tightly focusing a laser beam through a high numerical aperture objective. 42 Silica or polystyrene are among the most commonly used materials for optically trapped particles with typical sizes ranging between the 0.2-5 µm.43 However, also even smaller metallic nanoparticles can be optically trapped. 44 A small linker, such as antibody-antigen or streptavidin-biotin pairs, can be used to chemically affix the bead to the researcher's molecule of interest. 45 As the tracer bead may interact with the molecule of interest, in particular proteins, a double-stranded

DNA linker can be added between the small linker and molecule of interest to increase the distance between them, reducing possible interactions. If the spring constant,  $\kappa$ , is known, the force exerted by the optical trap can be determined by measuring the tracer's displacement with respect to the centre of the trap. Such measurements are useful to quantify the mechanical properties of living systems, including at the single molecule level.

#### D. Magnetic tweezers

Magnetic tweezers (MT), see Fig. 2(d) and (e), can exert or measure forces, both intra-cellularly 46,47 or on the surface of cells. 48 Magnetic tweezers allow for application and measurement of larger forces (up to nN) than optical tweezers (hundreds of pN regime). Magnetic tweezers measurements are made by introducing magnetic particles (either superparamagnetic beads or ferromagnetic nanowires) into a target region and applying magnetic field gradients to either exert or measure forces within cells, or by twisting magnetic particles specifically attached to molecules. Such beads experience a force determined by49

$$\mathbf{F} = \begin{cases} \frac{1}{2} \nabla (\mathbf{m}_{\text{max}} \cdot \mathbf{B}), & \text{for strong magnetic fields} \\ \frac{V \chi}{2\mu_0} \nabla |\mathbf{B}|^2, & \text{for weak magnetic fields} \end{cases}$$
 (6)

Here, V is the volume of the magnetic particle,  $\mu_0$  is the magnetic permeability, **B** is the magnetic field,  $\chi$  is the magnetic susceptibility of the particle, and mmax is the maximum possible magnetic moment induced by the magnetic field on the particle. The magnetic particle's susceptibility can be calculated from its relative permeability  $\mu_r$  via  $\chi = 3\frac{\mu_r - 1}{\mu_r + 2}$ , assuming that it is scalar in this first estimation. The equations to determine the forces for strong and weak magnetic fields are equivalent when  $\mathbf{m}_{\mathrm{max}} = \left(\frac{V\chi}{\mu_0}\right)\mathbf{B}$ . As magnetic particles normally used in such experiments are composed of aggregated magnetic nanoparticles their shape may not be isotropic and they can also be subject to a torque  $\Gamma = \mathbf{m} \times \mathbf{B}$ , where  $\mathbf{m}$  is the magnetic moment of the particle.

Similarly to optical tweezers, to use magnetic tweezers, the magnetic particle must be chemically bonded to the molecule of interest, whose opposite side is chemically bonded to a surface, typically a glass coverslip in a flow cell placed onto an inverted microscope. 50 Historically, the magnetic field was generated by placing two magnets on opposite sides of the bead, though multiple magnets can be used to create stronger magnetic fields, and thus apply stronger forces. Rotating these magnets induced rotation of the bead and the molecule to which it is attached. Typically, the magnetic field must be dynamically adjusted, requiring the use of electromagnets. As the magnetic field B may be determined from the geometry, number, and positions of the magnets, and  $\mathbf{m}_{\text{max}}$ , V, and  $\chi$  for the bead are known, the force exerted on the bead may be calculated. Other implementations of magnetic tweezers using

a cylindrical magnet have also been used, which has the advantage of allowing stretching forces to be applied, while also uncoupling stretching from rotational torsion.<sup>50</sup> Magnetic tweezers can be used to measure forces associated with protein unfolding, and have shown that ligand binding mechanically stabilises this process.<sup>51</sup> DNA supercoiling, describing the under- or over-twisting of DNA helices has been shown to drive transitions to uncommon DNA secondary structures, such as Z-DNA, DNA cruciforms, or DNA unwinding.<sup>52</sup> The possibility of both stretching and rotating DNA using magnetic tweezers makes them highly suited to study such phenomena.<sup>50</sup> Recent advances in magnetic tweezers technology have also allowed for the precise application of piconewton forces to beads implanted in cells, allowing for the determination that the nucleus stiffens upon force application, and that the actin filaments are principally aligned along the major nuclear axis.<sup>46</sup>

A similar method to magnetic tweezers is 3D-magnetic twisting cytometry (3D-MTC), which applies local mechanical stresses to living cells by twisting magnetic particles in a rotating magnetic field.<sup>53</sup> This method is suitable for investigating, *e.g.*, the mechanical response of cells to specific receptors.

### E. Structured illumination light sheet microscopy

Structured illumination microscopy is a type of super-resolution microscopy which uses different excitation patterns to illuminate a sample, then recombines them computationally to produce an image.<sup>54</sup> Light sheet microscopy (also called Selective Plane Illumination Microscopy, or SPIM) is a similar technique to widefield inverted microscopy, save that the sample is illuminated from the side. This geometry is advantageous: standard widefield microscopy illuminates not just the part of a sample in focus, but also fluorophores not in the focal plane, increasing the background in the observed sample. Light sheet microscopy uses a cylinder lens in the illumination path to stretch the laser light into a thin line, which is then used to illuminate the sample.<sup>55</sup> Combining the illumination patterns used in structured illumination microscopy with a light sheet microscope allows for very high resolution in even widefield images, allowing for highly accurate tracking of quantum dots.<sup>56</sup>

The wide variety of techniques presented here present several means of measuring tissue, cellular, or subcellular, forces and/or rheological properties. The information gleaned from these techniques can provide valuable insight into how cells interpret and transmit extracellular signals, or into subcellular structures such as DNA or proteins. In the following section, we will discuss in-depth material properties measured by these techniques.

### III. Determining materials properties

Living organisms are made of tissues, cells, or molecules, with highly varying properties. Tissues or cells can be designed to withstand forces (*e.g.*, in bones) or to be compliant, allowing them to easily pass through narrow spaces, such as a blood cell through a blood vessel. A tissue's material properties govern its

ability to deform and determine how the tissue reacts to external stresses. Materials properties also determine the fluidity and stiffness of a biomaterial, as well as its reaction towards externally or internally generated forces.

Certain materials have a texture which changes when stress is applied and this change depends on the frequency with which the stress is applied. One example is ketchup which becomes 'runnier' when shaken. Living organisms are to a large extent composed of such matter and it is thus crucial to take into account the frequency-dependent response when describing the properties of living matter. For cellular movement accompanying organ development, the relevant time scales may be on the order of hours, or even days. On the other hand, the time scales relevant for the dynamics of biopolymers constituting the cytoskeleton is on the order of milliseconds.<sup>57</sup> Hence, the time scales relevant for biological function vary dramatically, and the materials properties change accordingly; a cell may appear rather rigid on short time scales but quite compliant on longer time scales.

#### A. Review of basic mechanics

For a detailed discussion of materials properties in a biological context, it is helpful to recall basic mechanics. Determination of a material's properties typically involves applying a force F and measuring the resultant deformation, or vice versa. As samples may vary geometrically from one another, and forces are usually only applied to one surface of a sample, it is common to describe the applied force using the stress  $\sigma = \frac{\mathbf{F}}{A}$ . where A is the surface area onto which a force is applied. As the force may be applied longitudinally (through either stretching or compression), or transversally (e.g., shear),  $\underline{\sigma}$  itself is a tensor. Depending on how the stress is applied geometrically, and to control for different sample sizes, sample deformation is characterised by the strain & Like stress, strain is a tensor with each component defined as  $\varepsilon_{ij} = \varepsilon_{ji} = \partial u_i/\partial j + \partial u_j/\partial i$ , where **u** describes the displacement field of the sample, including both displacement of the object and its deformation.

1. Isotropic elastic material: Young's modulus and Poisson ratio. One of the simplest ways of describing a materials properties is by Hookes law, relating the stress,  $\underline{\sigma}$ , linearly to the strain,  $\underline{\varepsilon}$ , via the 4th-order elasticity tensor. For most practical purposes in soft-condensed matter physics, this can be reduced to a formulation based on Youngs modulus, E:  $\underline{\sigma} = E_{\underline{\varepsilon}}$ . This equation assumes an isotropic material, and that E is scalar. The tensorial nature of the stress-strain relation can be readily seen in practice by applying pressure, or stress, to a small area of an elastic substance. While the elastic material compresses in the direction of the applied stress, there are also longitudinal strains on the material, pulling it from the sides towards the applied pressure. For experiments where the sample is uniformly compressed, or stretched, along one axis across all of the material, this equation becomes onedimensional, giving  $\sigma = E\varepsilon$ . Youngs modulus describes a materials response in the direction of the applied stress, a

relation that is only valid in the regime where the response is

elastic and linear, which is typically true for small strains or

Materials properties can also be determined by applying forces in directions other than perpendicular to a given material's surface. In particular, the orthogonal responses to a stress that maintains an isotropic material's volume, termed shear, are described by the shear modulus G. Experiments to determine G apply stresses by deforming one surface in a direction perpendicular to that of its normal vector, while maintaining a constant volume. As with E, measurements of G are only practically possible in the regime where the material's response is linear and elastic.

While the direction of the material's response may differ between G and E, both these properties measure the fundamental "stiffness" of the material, and are thus related. Assuming the material is isotropic, this can be seen through

$$E = 2G(1 + \nu) = 3K(1 - 2\nu), \tag{7}$$

where K is the bulk modulus, a measure of a material's resistance to compression, and  $\nu$  is Poisson's ratio. <sup>58</sup> Poisson's ratio describes a material's response in directions orthogonal to that of an applied stress. Eqn (7) provides a convenient way to calculate any of the three materials properties E, G, or K for a sample from another one, given that the sample's  $\nu$  is known. The latter can be determined for a material by stretching it, or compressing it, along a certain axis, and calculating

$$\nu = -\frac{\mathrm{d}\varepsilon_{\mathrm{trans}}}{\mathrm{d}\varepsilon_{\mathrm{long}}},\tag{8}$$

where  $d\epsilon_{trans}$  and  $d\epsilon_{long}$  are the changes in the material's size transverse and longitudinal to the applied deformation, respectively. Unlike a shearing test to determine G for a material, a material could undergo a change in volume during an experiment to determine  $\nu$ .

### Viscoelasticity

(a) Frequency dependent viscoelastic modulus. Soft-matter and biological materials, however, typically possess both elastic and viscous, or viscoelastic, properties; the response of such systems is thus highly dependent upon the time scale of the applied stress. The viscoelastic properties of cells originate from their molecular constituents where the cytoskeleton and nuclear matrix act as elastic components, while the dense packing of the cytoplasm creates a highly viscous environment.<sup>13</sup> To characterise viscoelastic systems it is convenient to use the time-dependent shear modulus G(t) which is defined such that:

$$G = \lim_{t \to \infty} G(t). \tag{9}$$

This equation summarises the idea that the long-time shear response G of a material may be different from that on shorter time scales. This can more readily be seen when observing the flow of certain viscoelastic materials, such as whipped cream, after they are deformed: on short time scales, these materials can maintain a certain shape created through a deformation, while at long time scales they relax, losing their shape. One way of sampling how a material behaves across different time scales is

to apply an oscillatory shear  $\sigma(f)$  to it at different frequencies f(or angular frequencies  $\omega$ ), and measure the compliance of the medium (how much it deforms under stress),  $\gamma(f)$ , or *vice versa*. These frequency-dependent viscoelastic properties can be conveniently expressed through the complex shear modulus, also called the dynamic or complex shear modulus:

$$G^*(f) = G'(f) + iG''(f).$$
 (10)

 $G^*(f)$  is a sum of two parts: (i) a real part, G', termed the storage modulus and quantifying the material's elastic response, (ii) an imaginary part, G'', termed the loss modulus and describing the dissipated energy, or viscous response, of the material. These moduli also do not oscillate in phase with one another; this time lag is characterised by  $tan(\delta) = \frac{G''}{G'}$ . This time lag characterises how elastic or viscous a material is: for  $\delta \approx 0$ , the stress and strain are in phase with one another, such as for perfectly elastic materials, while for  $\delta \approx \pi/2$ , the stress and strain are the furthest out of phase with one another, a characteristic of viscous materials.  $G^*(f)$  and its two components are commonly measured in bulk for a material using a rheometer. For a detailed derivation of these properties and how to measure these, we refer to the book by D. Boal for more details on the complex shear modulus and viscoelastic properties of living organisms.59

(b) Linear response theory. The frequency-dependent response of a material at thermal equilibrium is equally well described by the positional power spectrum,  $\mathcal{P}(f)$ , which can be calculated through a Fourier transformation of the thermally driven positional time series of a tracer particle, x(t). There is a 1-to-1 relation between  $\mathcal{P}(f)$  and  $G^*(f)$  because the Fourier transform of the stochastic thermal force, F(f), and the Fourier transform of the position of the particle, x(f), are related through linear response theory:60

$$x(f) = \gamma(f)F(f), \tag{11}$$

where  $\gamma(f)$  is the compliance of the medium, describing its deformation under a certain, not too large stress.61

(c) Generalised Stokes-Einstein equation. To relate this compliance to the viscoelastic properties of the medium, consider the classical generalised Stokes-Einstein relation for a tracer particle of radius r in a fluid with viscosity  $\eta(t)$  at any given time t. As previous forces affecting the system determine the state of the material at all future times, the memory function  $\zeta(t)$  is commonly used. To simplify calculations of viscoelastic properties, Mason and Weitz assumed that the Laplace transform to the viscosity was related to the memory

function by  $\widetilde{\eta}(s) = \frac{\widetilde{\zeta}(s)}{6\pi r}$ , where s is the Laplace frequency.<sup>62</sup> For such a system, the dynamic shear modulus in Laplace space is

$$\tilde{G}(s) = s\tilde{\eta}(s) = \frac{k_{\rm B}T}{\pi rs\langle \tilde{r}^2(s)\rangle},$$
 (12)

where  $\langle \hat{r}^2(s) \rangle$  is the mean squared displacement.<sup>63</sup> Applying the Fourier transform to eqn (12), and assuming that the angular **Perspective** 

frequency  $\omega = 2\pi f$  is related to *s* by  $s = i\omega$ , the Generalised Stokes-Einstein relation is obtained,

$$G^*(f) = \frac{1}{6\pi r v(f)},\tag{13}$$

where  $\gamma(f)$  is the compliance of the medium.<sup>64</sup> As the system can be described through linear response theory, the Kramers–Kronig relation, coupling a complex system's real and imaginary components, may be applied, thereby showing that the scaling properties of G' and G'' are identical to the scaling properties of  $\mathscr{P}(f)$ ,  $^{60,64}$  as described below. There exist many different versions of such mechanistic models to describe more complex materials  $^{65,66}$  and viscoelastic properties for materials with long-range memory effects were successfully modelled by fractional-order models.  $^{67-69}$ 

Physical models of viscoelasticity employ viscous or elastic components placed in series and/or parallel with one another. These models use a 'dashpot' to illustrate viscous components, and a spring for the elastic components. Of these models, the most basic include a dashpot in series with a spring (the Maxwell model), or a dashpot in parallel with a spring (the Kelvin-Voigt model). Applying a stress to a material that follows the Maxwell model (Maxwell material) initially stores the applied energy in its elastic component, which is slowly dissipated by its viscous component. For materials that follow the Kelvin-Voigt model (Kelvin-Voigt material), a suddenly applied stress is initially greatly resisted by the viscous component, a resistance which decreases over time as its elastic component undergoes a slow compression.

In practice, a convenient way to quantify the viscoelastic properties of a material at thermal equilibrium is by tracking an inert tracer particle placed within it, as illustrated in Fig. 3a. First, the time series of the positions visited by the tracer particle should be measured, potentially by video microscopy or by tracking using optical tweezers. Then the positional time series is Fourier transformed and from this the positional power spectrum calculated.

For biological viscoelastic materials, there exists a frequency regime, typically 300 Hz < f < 6000 Hz, where thermal fluctuations dominate over active processes  $^{57}$  and which is still below the filtering effect of a typical photodiode.  $^{70}$  In this frequency interval, the experimentally obtained power spectrum is observed to scale with frequency and can be fitted by:  $^{71}$ 

$$\mathscr{P}(f) \propto f^{-(\alpha+1)}.\tag{14}$$

In such regimes, the scaling exponent,  $\alpha$ , carries information on the materials properties of the viscoelastic system and hence also the properties of the surrounding medium (see also Section IV):

 $\alpha$  = 0 The particle is restrained or even immobilised by the surrounding medium.

 $0<\alpha<1$  The subdiffusive regime, where the particle diffuses more slowly than in a purely viscous medium. Here, lower values of  $\alpha$  indicate a more elastic medium and values closer to unity indicate a more viscous medium (if there are no active processes in the observed frequency window).

 $\alpha$  = 1 The particle exhibits Brownian motion, and the surrounding medium is purely viscous.

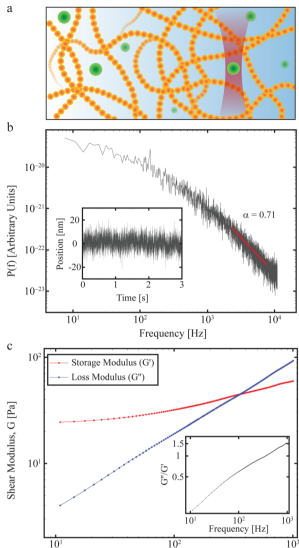


Fig. 3 Materials properties such as viscoelasticity can be measured using tracer particles. (a) The positions visited by a tracer particle (green ball) can be measured using a focused laser beam to form an optical trap (see also Section II), and used to quantify the materials properties of a sample, such as the polymer solution shown here. (b) Power spectrum as a function of frequency of an optically trapped tracer particle in a 50% Matrigel solution. The position of the particle as a function of time. (c) Extracted complex shear moduli for a tracer particle in a 50% Matrigel solution calculated on the basis of the same data as shown in (b). Inset: The loss tangent,  $\tan(\delta) = G''/G'$ , a measure of how solid- (G''/G' < 1) or fluid-like (G''/G' > 1) the viscoelastic material is.

Frequency [Hz]

 $\alpha > 1$  The particle undergoes superdiffusion, moving faster than it would by thermal diffusion in a viscous medium, *i.e.*, active processes propel the particle.

The real and imaginary parts of the complex shear modulus  $G^*(f)$  scale with frequency in the following form:

$$G'(f) \sim f^{\alpha} \tag{15a}$$

$$G''(f) \sim f^{\alpha} \tag{15b}$$

**PCCP** Perspective

permitting the materials properties to be calculated from the power spectrum through  $f^{\alpha}$ . Note that both G' and G'' scale with  $f^{\alpha}$  only for polymer networks in the frequency range below the molecular high frequency cut-off and above the characteristic mesh relaxation time.<sup>60</sup>

An example of how any material's viscoelastic properties can be quantified, both living or non-living, is given in Fig. 3. In this example, a tracer particle (a micron-sized polystyrene bead) is embedded in a Matrigel possessing viscoelastic properties. The experiment is described in detail in ref. 73 which outlines the properties of the Matrigel as a function of polymer concentration and how this affects organoids growing in the matrix. Fig. 3a demonstrates the experiment with the green dots signifying the polystyrene tracer particles, of which one is optically trapped. The forward scattered laser light is picked up by a quadrant photodiode operating at 22 kHz and located close to the back focal plane. The inset in Fig. 3b shows the positions visited by the tracer and the main figure shows the power spectrum as a function of frequency calculated from the positional time series on a double-log plot. Consistent with eqn (14), there exists a frequency regime within which  $\mathcal{P}(f)$ scales with frequency (red line). At frequencies below this regime the tracer particle feels the restoring force from the optical trap and at frequencies larger than this regime the photodiode exerts a filtering effect.<sup>70</sup> Due to these experimental limitations, a scaling relation can only be confirmed in a certain frequency window, this frequency window being relevant for describing polymer dynamics.<sup>57</sup> Within this frequency window, a fit (red line in Fig. 3b) returns  $\alpha = 0.71$ , indicating that the tracer exhibits subdiffusive motion (see above and Section IV), as expected for a polymeric matrix. Fig. 3c shows the storage and loss moduli as a function of frequency, calculated from the data in Fig. 3b and using the relations described in this section.

As living systems are dynamic in nature, their non-equilibrium materials properties are of importance. Non-equilibrium stress properties within cells can also been measured via force spectra,74 demonstrating that the changing stress properties within the cell can be separated from thermal fluctuations. Using this technique, it has been shown that cells can adapt to changing external stresses, adjusting their intracellular stress. Further use of force spectra could provide more much needed insight into intracellular non-equilibrium dynamics.

(d) Microrheology. Using a microrheological methodology as illustrated in Fig. 3 and in combination with endogenously occurring lipid granules as tracer particles, it is possible to map out the viscoelastic properties of living biological systems, such as cells, using optical tweezers. By doing so, anomalous diffusion<sup>38</sup> as well as weak ergodicity breaking<sup>75</sup> (see below) has been demonstrated within living yeast cells. And recently, using this technique invasive cancer cells have been shown more capable of adjusting to the stiffness of their environment than non-invasive cells.<sup>2</sup> Also, microrheological quantification of the materials properties of the basement membrane in combination with mathematical simulations have shown that Net4 softens the mechanical properties of native basement

membranes, thereby decreasing cancer cell potential to transmigrate this barrier, and thus finding that the stiffness of the basement membrane is a key determinant for metastases formation.76

(e) Crossover frequency and solid/liquid transition. One interesting property of many viscoelastic materials is that, depending on the time scale/frequency at which they are observed, they may behave more like a viscous liquid than an elastic solid, or vice versa. The point at which a viscoelastic material transitions from viscous liquid to an elastic solid is called the crossover frequency, and is typically defined as the frequency at which the values of G' and G'' are equal to one another. At frequencies below this crossover frequency (long time scales), these viscoelastic materials behave more as viscous liquids. As the frequency is increased, the material begins to have more elastic properties, until the crossover frequency is reached where the elasticity of the material makes it act more as a solid than a viscous liquid (shorter time scales).

For emulsions of two different kinds of liquids or materials in which one is not miscible or soluble in the other, the viscoelastic properties (G' and G'') are significantly different, and can be described by the liquid droplet model. As the materials are immiscible with one another, one will form droplets embedded in the other. Such a material's viscoelastic properties are a function of the droplet volume fraction  $\phi$  in the emulsion, with materials having low droplet  $\phi$  behaving as a viscous liquid, and those with high droplet  $\phi$  acting as an elastic solid.<sup>77</sup> In materials of moderately high  $\phi$ , G' is fairly constant for low frequencies, and G'' dropping with f until intermediate frequencies, with the behaviour resembling that of an elastic solid. 77,78 At higher frequencies, however, G' scales with  $f^{1/2}$ , while G'' rises with f, eventually surpassing G'. The other words, the liquid droplet model states that at low frequencies, as the droplets cannot be mixed with the surrounding material, the material acts as a solid. Once high frequencies are applied, equating to vigorous shaking, the droplets shrink in size or are destroyed altogether, and can be forced to mix in with the surrounding material, allowing it to behave as a viscoleastic liquid. As some biological components are immiscible with one another, as is the case with cell or nuclear membranes in living systems, the liquid droplet model can help describe their viscoelastic behaviour.

### B. Atomic force microscopy and other viscoelastic measurement techniques

Atomic force microscopy (AFM, see Fig. 2e) is another excellent tool for measuring materials properties. Using a sphere of radius R affixed to a cantilever lacking a tip, an AFM can measure Young's modulus,  $E_{1}^{79}$  or the complex shear moduli G' and  $G''^{80}$  of a viscoelastic material, such as cells. The cantilever is slowly lowered until the sphere indents the sample a distance  $\Delta h$ , bending the cantilever a distance  $d^{79,81}$  Assuming the measured samples contain an elastic component, forces F measured using such a system can be related to E and the

Poisson ratio,  $\nu$ , of the material or cell using the Hertz model<sup>81</sup>

$$F = \frac{4\sqrt{R}}{3} \frac{E}{1 - \nu^2} d^{3/2}.$$
 (16)

While seemingly straight-forward, for soft materials such as cells (0.1 kPa < E < 100 kPa) it is critical that an appropriate indentation depth  $\delta$  is used: for very small or very large values of  $\delta$ , measurements of E using eqn (16) are inaccurate. Recent advances in analytical methods have enabled the determination of the rheological properties G' and G'' directly from raw experimental data. This technique has been successfully integrated with AFM: by applying a step-strain indentation to samples using a bead affixed to a cantilever, Chim,  $et\ al.$ , were able to measure the local viscoelastic properties of gels, and even cells, using AFM. Results using this technique for a polydimethylsiloxane (PDMS) gel and polyacrylamide gel-like solution were compared to those obtained with a rheometer, showing reasonable agreement between values of G' and G'' between both methods.

Magnetic tweezers (see Section II) are another excellent tool with the capacity for measuring viscoelastic properties on a sub-cellular level: magnetic tweezers can be used to probe the materials properties within cells and tissues, such as the viscoelastic properties of individual fibroblasts, <sup>83</sup> the viscosity of fly embryos, <sup>84</sup> or the stiffness in mouse blastocysts. <sup>85</sup>

On larger length scales, viscoelastic properties of entire cells can be assessed by microfluidics-based real-time deformability cytometry. In this methodology, a cell is squeezed through a microscopic channel while measuring its deformation.86 This type of equipment has been successfully used, for instance, to measure how blood cells change their viscoelastic properties upon differentiation,87 and to demonstrate how neuronal cells change their mechanical properties through reprogramming and differentiation.88 Although in principle the response is only observed at one frequency (the inverse of the deformation time), this type of study does allow for comparisons between different materials. This is also true for studies based on optical stretchers where the so-called "compliance" is measured, which has been used to investigate the viscoelastic properties of glial cells<sup>89</sup> and cancer cells, relating viscoelastic properties to invasive potential.90

# C. Experimental consideration of sampling time vs. frequency response

For optical tweezer experiments seeking to measure viscoelastic properties, the frequency response can depend upon the sampling time. By calculating the Allan Variance, <sup>91</sup> the noise in optical tweezers setups can be quantified and one can determine the optimal measurement time, measurement frequency, and detection scheme, thereby providing a qualified choice of these parameters.

### D. Illustrative example

To provide a more concrete example of how to calculate the different materials properties, consider a sample of material with known Poisson's ratio  $\nu$ . One means of measuring its

materials properties is via AFM, as shown in Fig. 4. Here, a bead of known radius R is glued to an AFM cantilever with no tip via an epoxy, and its spring constant k is measured. The bead is then brought into contact with the material, and pushed in a distance d. This results in the bending of the cantilever a distance  $\Delta h$ , caused by a force of  $|\mathbf{F}| = k\Delta h$ . Applying the Hertz model (eqn (16)), the

Young's modulus is calculated as  $E = \frac{3F}{4\sqrt{R}} \frac{(1-\nu^2)}{d^{3/2}}$ . Note that for

softer materials, and biomaterials in particular, the measured E is a function of d until a certain indentation depth; to obtain the true value for Young's modulus of such a material, d must be increased until a plateau is observed in calculations of E. The shear modulus G can be calculated from the value of E using eqn (7),

resulting in  $G = \frac{3F}{8\sqrt{R}} \frac{(1-\nu^2)}{d^{3/2}(1+\nu)}$ . Alternatively, the bulk modulus

*K* can be calculated *via*  $K = \frac{F}{4\sqrt{R}} \frac{(1-\nu^2)}{d^{3/2}(1-2\nu)}$ .

# IV. Characterising transport and organisation

Self-organisation, in particular in living matter, emerges from an intimate interplay of transport processes, passive and active, on many length and time scales, along with a multitude of chemical reactions. Prominent examples of dynamic self-organisation phenomena range from dynamic protein gradients in single cells<sup>92–96</sup> and *in vitro* assays<sup>97,98</sup> over compartmentalisation of biochemical reactions<sup>99</sup> or active arrangements of cellular organelles, <sup>100–103</sup> up to organised fluid dynamics of tissues during development, *e.g.*, in zebrafish and fly embryos. <sup>104–107</sup> In the context of organismal development, mechanical cues, materials properties (see Section III), and the interplay with biochemical

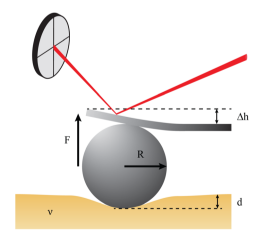


Fig. 4 Example measurement of elastic properties using AFM. A material of known Poisson's ratio  $\nu$  is indented by a sphere of radius R attached to an AFM cantilever. The cantilever has a spring constant of k (calculated before the experiment), and is bent by a distance  $\Delta h$ , indicating a force  $|\mathbf{F}| = k\Delta h$ . Applying the Hertz model from eqn (16), the Young's modulus is calculated as  $E = \frac{3F}{4\sqrt{R}} \frac{(1-\nu^2)}{d^{3/2}}$ .

gradients have been of prime importance (see ref. 108-113 for some examples).

Relating to the previous sections, we here focus on fundamental and generic aspects of transport in the self-organisation of living matter. Due to its global appearance and generic character, we concentrate on thermally driven or active diffusive motion as its epitomisation. In particular, the current section discusses known facts and current challenges of diffusional transport by first outlining relevant theory and second relevant experiments before highlighting some crucial aspects of the interplay between transport and (bio)chemical reactions in selforganisation processes.

#### The Gaussian probability density function

Ever since Robert Brown's vivid account of the jittery motion of microscopic particles derived from pollen and rocks, diffusive processes have been the focus of statistical physicists. Starting with the seminal works of Einstein<sup>114</sup> and Smoluchowski, <sup>115</sup> "Brownian motion" has become an overarching statistical description for non-equilibrium phenomena in all areas of physics, physical chemistry, and even financial mathematics. Langevin's formulation in terms of a stochastic differential equation connected diffusion to Newtonian mechanics in the presence of effective, random forces. 116 Today, massive advances in microscopic techniques and the ability to fluorescently tag and monitor submicron tracers or even single molecules, make it possible to garner single-particle trajectories at nanometer-resolution in complex environments such as living matter.117

The mathematical description of Brownian motion is often called "universal", as the mean squared displacement (MSD)  $\langle \mathbf{r}^2(t) \rangle = \int_{-\infty}^{\infty} \mathbf{r}^2 P(\mathbf{r}, t) dV = 2dK_1 t$  fully defines the Gaussian probability density function (PDF), i.e., the propagator of the diffusion equation, in d dimensions

$$P(\mathbf{r},t) = (4\pi K_1 t)^{-d/2} \exp\left(-\frac{\mathbf{r}^2}{4K_1 t}\right).$$
 (17)

Here,  $K_1$  is the diffusion coefficient with units m<sup>2</sup> s<sup>-1</sup>. The PDF is used as a measure of the likelihood of a particle being at position r at time t, while the particle's MSD is the expected squared distance travelled by a particle subject to Brownian motion relative to its initial position. For spheres with radius Rin a fluid of viscosity  $\eta$ , *i.e.*, d = 3, at thermal equilibrium the familiar Stokes-Einstein result  $K_1 = k_B T/(6\pi \eta R)$  holds. Similar relations  $K_1 \propto 1/R_{\rm eff}$  were also shown to hold for the momentary diffusivity of fluctuating proteins in water, 118 for drum-shaped proteins of radius  $R_{\rm eff}$  in crowded membranes, <sup>119</sup> and in general for sufficiently large membrane domains. 120,121 The Gaussian law, (eqn (17)), with time-linear MSD  $\langle \mathbf{r}^2(t) \rangle = 2dK_1t$  holds as long as the diffusing particle moves unbounded and interaction-free in unconstrained space. Allowing for interactions, the picture becomes already considerably more complex. When specialised proteins search for a specific DNA target sequence, for example, they switch intermittently between three-dimensional diffusion in the cell (or the nucleus) and one-dimensional diffusion along

the often randomly coiled DNA, 122-125 thereby significantly improving the search efficiency over a simple diffusive process. Similarly, particles may intermittently bind to a surface, e.g., a cellular membrane, which alters the overall diffusion statistic 126-128 and mediates the escape through a small opening in the surface. 129,130

On the other hand, restricting the available space for simple diffusion, e.g., by randomly placed (almost) immobile obstacles, may not only decrease the transport coefficient  $K_1$  but can result in strong deviations from the Gaussian law (eqn (17)), and even in a local confinement ("corralled diffusion"). Indeed, in more complex and/or non-equilibrium systems, such as in biological fluids, gels, or membranes, the simple laws of diffusion are no longer applicable, and are thus non-universal. 117,127,131-134 In the following paragraphs we discuss two specific showcases of nonuniversality of diffusion.

A frequent observation, especially in soft and living matter, is that of "Brownian yet non-Gaussian" diffusion. 135-137 Here, the MSD still grows linearly in time, but the PDF of step lengths deviates from the anticipated Gaussian law (eqn (17)), often displaying exponential tails. One possible explanation for this phenomenon employs a heterogeneous ensemble of trajectories, e.g., when monitored particles have different radii, as encountered for commercially available tracer beads. As a result, while each trajectory has its well-defined diffusivity  $K_1$ , the ensemble of trajectories of different particles is characterised by a distribution  $p(K_1)$  of diffusion coefficients. The non-Gaussian behaviour of the averaged PDF  $P(\mathbf{r},t)$  then emerges from a superposition of individual Gaussian contributions from each trajectory, with the particular features of  $P(\mathbf{r},t)$  depending on the exact shape of  $p(K_1)$ . 135,136 This "superstatistics" reproduces the widely observed exponential forms of  $P(\mathbf{r},t)$  for exponential shapes of  $p(K_1)$ . Thus, by simply lumping together all steps from all trajectories, deviations from the anticipated Gaussian behaviour may be created due to ensemble heterogeneity in an otherwise homogeneous environment.

Heterogeneity within individual trajectories can be emphasised by normalising each trajectory by its root-mean-squared step length, eliminating any impact of the ensemble of particles. If the resulting  $P(\mathbf{r},t)$  still deviates from a (standard) Gaussian (eqn (17)), then  $K_1$ -heterogeneity within individual trajectories can be deduced (as illustrated in Fig. 5a). Several reasons may account for such an observation, e.g., spatially and/or temporally varying viscosities or effective local mobilities may alter a particle's step length while it explores its surrounding. Alternatively, particles may switch stochastically between different mobility states, such as via a transient coupling to stationary structures or processive molecular motors. 140-142 Even the radius of the tracer particle itself may grow or shrink, e.g., by ongoing multimerisation, especially when using fluorescently-labelled biomolecules, 143-145 or via conformational fluctuations. 118 Such cases can be effectively modelled as "diffusing-diffusivity" processes, assuming that the diffusion coefficient becomes explicitly time-dependent,  $K_1(t)$ , and varies stochastically. <sup>139,146–148</sup> As a result, the MSD grows linearly in time with a constant effective diffusivity as prefactor, while the displacement PDF starts with

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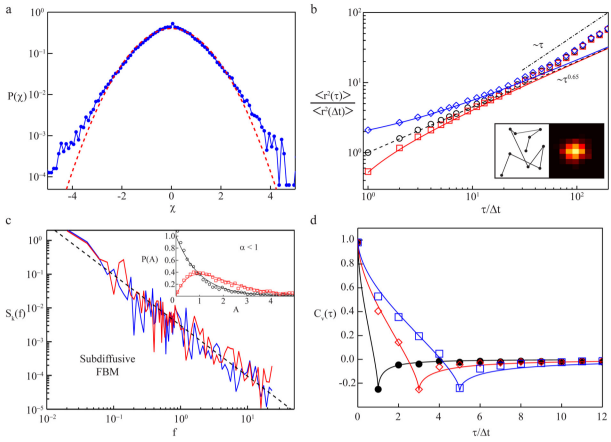


Fig. 5 Transport observables and measurement artifacts. (a) The PDF of normalised increments, P(y), from trajectories of quantum dots in the cytoplasm of living cells (blue symbols) deviates from the anticipated standard Gaussian (red dashed line), indicating significant diffusion heterogeneity of individual trajectories; data from ref. 142. (b) The TA-MSD of a single trajectory with exactly known positions follows the anticipated power-law scaling (black circles and dashed line). Both static and dynamic localisation errors (blue diamonds and red squares, respectively) induce considerable deviations for small lag times. Left inset: Visited positions during image acquisition, responsible for a dynamic localisation offset. Right inset: Camera image of a single immobile particle whose position can only be determined via static localisation error. Data from ref. 153. (c) The PSD of individual subdiffusive telomer trajectories (red and blue lines) follow the predicted FBM scaling (black dashed line). Inset: Normalised fluctuations, A, of individual PSDs from the ensemble mean feature universal distributions, P(A), depending upon dimension (black 1D and red 2D trajectories, respectively); data from ref. 154. (d) Velocity autocorrelation functions of subdiffusive quantum dots in the cytoplasm of living cells. The anti-correlation peak agrees with the FBM prediction with an anti-persistent memory. Instantaneous velocities were determined between frames separated by  $\Delta t$  (black circles),  $3\Delta t$  (red diamonds), and  $5\Delta t$ (blue squares). Data from ref. 142.

exponential tails (or more general shapes 149) and crosses over to an effective Gaussian at times longer than an intrinsic correlation time. 139,146-149 Alternatively, quenched spatial disorder may give rise to non-Gaussian behaviour.  $^{150-152}$ 

### B. Anomalous diffusion

Here, we discuss anomalous diffusion, as introduced in Section III and using the same scaling exponent,  $\alpha$ . Anomalous diffusion is conventionally defined by a nonlinear scaling of the MSD, often assuming a power-law form

$$\langle \mathbf{r}^2(t) \rangle \simeq K_{\alpha} t^{\alpha},$$
 (18)

where  $K_{\alpha}$  represents the generalised diffusion coefficient.  $K_{\alpha}$  has the physical dimension m s<sup>- $\alpha$ </sup>. The coefficient  $K_{\alpha}$  depends on the details of the specific physical process behind the observed motion. For instance, in long-range correlated dynamics, such as for viscoelastic anomalous diffusion, the exponent  $\alpha$  is

related to the power-law exponent of the noise autocovariance function. Or,  $\alpha$  emerges for the MSD of random walk processes with scale-free distributions  $\psi(\tau) \simeq t_0^{\alpha}/\tau^{1+\alpha}$  of waiting times  $\tau$ between successive jumps with  $0 < \alpha < 1$  and finite variance  ${\ell_0}^2$  of the jump lengths,  $K_\alpha = {\ell_0}^2/[2t_0^\alpha]$ . Generally  $K_\alpha$  is proportional to the strength of the driving noise. For instance, in the mentioned processes with waiting times  $K_{\alpha} \propto k_{\rm B}T$ , where  $k_{\rm B}$  is the Boltzmann factor and T temperature. 155

A prominent example is stochastic motion fuelled by a Gaussian yet power-law correlated noise, such as in viscoelastic systems. Depending on whether the system is at equilibrium and subject to a fluctuation-dissipation relation or not (intrinsically, biological cells are very far from equilibrated systems), the resulting motion is governed by the generalised Langevin equation 156,157 or fractional Brownian motion. 158 This behaviour is in fact closely related to the viscoelastic nature of many complex systems, e.g., in lipid bilayer membranes, 159 worm-like

**PCCP** micellar solutions, 160 or in crowded media like biomimetic fluids 161-163 or the cytoplasm of cells. 142,164-168 Notably, viscoelastic yet non-Gaussian processes have also been analysed. 140,167,169,170 Another class of systems is described by continuous-time random walks, composed of two, possibly heavy-tailed, PDFs governing the random jump lengths and waiting times of this process. 171 For instance, scale-free immobilisation times dominate the motion of membrane proteins<sup>172</sup> and insulin granules.<sup>173</sup> Numerous other anomalous diffusion processes exist. 132,150,174,175 As discussed in Section III, normal diffusion implies  $\alpha = 1$ . We mention that measured MSDs do not always follow a unique power-law trend across the entire measurement. Instead, cross-over behaviours occur, and normal diffusion with  $\alpha = 1$  may emerge as an intermediate-asymptotic behaviour. 128 Such a situation is demonstrated in Fig. 5b which shows the time-averaged MSD (TA-MSD) of a subdiffusive trajectory of a simulated FBM (black circles). The dynamics follows the anticipated subdiffusive power-law scaling  $\langle r^2(\tau)\rangle \sim \tau^{0.65}$ . Moreover, as demonstrated here, errors in the localisation of the tracer particle may spoil the power-law behaviour of the MSD. Thus experiment-inherent static or dynamic localisation errors induce significant perturbations (blue and red symbols, respectively): while the finite number of photons per frame introduces statistical position uncertainty in the associated raster image (cf. right inset), particle movement during this finite acquisition time smudges the visited loci (cf. left inset). For large lag times  $\tau$  in sufficiently large systems, a crossover to normal diffusion ( $\alpha = 1$ , dash-dotted line) is frequently observed at times beyond some

crossover time scale. Finally, finite systems such as biological cells

eventually enforce a crossover of the MSD to a constant plateau ( $\alpha$  = 0) due to an asymptotic confinement of tracer particles. Notably,

even at stationarity in a bias-free environment, the particle PDF

does not necessarily need to have the same amplitude everywhere,

as, e.g., for Brownian motion, but may increase or decrease close to boundaries. 176,177 Closing this brief overview, we want to emphasise

that the effects of additional system-inherent noise may also need

to be considered, e.g., created by the long-term motion of cells with

tracking particles, or by drift in the experimental apparatus. Such

noise may partially mask some of the actual physical processes

underlying the diffusive motion of tracked particles. 178-181

The existence of so many relevant physical processes, all characterised by the same scaling (eqn (18)) of the MSD, presents a major challenge for the analysis of (experimental) data: how can we infer the true physical process governing the observed systems from the measured data? Apart from simply characterising the system, this information is also vital to predict secondary and coupled processes, such as chemical reactions or diffusion-mediated relaxation dynamics, and eventually also emergent phenomena like pattern formation, a topic which is discussed further in the following section.

### C. Combining transport and reactions - crucial aspects of self-organisation

On the most fundamental level, transport impacts cellular organisation by bringing reactants closer together, such as for proteins, whose interaction triggers downstream events like gene expression or large-scale cellular motion. Yet, like the classical Michaelis-Menten scheme for enzymes and substrates, most cellular reactions, and even complex signalling networks, can be formulated and analysed in terms of concentrations as sets of ordinary differential equations (ODEs). 182,183 Spatial information might be considered here by restricting certain reaction steps to separated vessels that exchange material with some kinetic rate. Hence, transport events are lumped into effective (pseudo) rates and all particles are assumed to be well-mixed (in their respective vessel) at each instant of time. However, this approach fails miserably if the number of reactants is low. At this condition, the dissociation event in a Michaelis-Menten scheme does not, for example, lead back to a well-stirred mixture, but rather towards a diffusion-driven rebinding of the same substrate molecule to its corresponding enzyme, an interaction favoured over all competitors. 184 As a consequence, the very same enzyme can act on substrates in either a processive or distributive fashion, depending on the actual concentrations and transport coefficients. 185 In fact, considering diffusion-driven rebinding for the MAPK pathway (which communicates a signal from the cell surface to the nucleus), this rebinding erases the toggle-switch behaviour that had been predicted by ODEs. 184

Subdiffusion of the fractional Brownian motion (FBM) type can further enhance rebinding events, 186-188 and may even lead to fractal kinetics. 189,190 Notably, the crowded state of cells not only influences reactions by altering diffusive transport and rebinding, but it also stabilises associated states (e.g. protein adsorption to target membranes) in a very general way by rendering the immediate surrounding volumes inaccessible, i.e., hampering dissociation. 191

Given that transport significantly impacts information processing at the most fundamental level, it is mandatory to focus not only on ODEs and apparent kinetic rates, but also to thoroughly consider and quantify transport when aiming to understand cellular self-organisation. 192-194 Thus, considering both the experimental and theoretical limitations and challenges becomes even more important.

On larger length scales, diffusional transport is key for the emergence of spatial concentration patterns that facilitate intracellular self-organisation, tissue organisation, and embryonic development. 195 A prime example in this context are reaction-diffusion systems with an activator and an inhibitor, whose antagonistic and nonlinear reaction terms can lead to the emergence of stationary spatial patterns when diffusional mixing is sufficiently poor (see Fig. 6 for an example). 196 Such spatially non-uniform steady states are named "Turing patterns", in honour of A. M. Turing. By now, many similar pattern formation systems have been observed experimentally, and a multitude of pattern forming systems formulated mathematically. These include, but are not restricted to, scenarios that invoke depletion effects instead of inhibition, advection in addition to diffusion, or the conservation of particle numbers, showing temporal oscillations instead of a stationary spatial pattern. Unfortunately, a Babel-like state has emerged over the years in the literature: at times, any kind of gradient formation is simply named a Turing pattern, while at others already wellknown phenomena are coined with new and fancy names.

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Fig. 6 Transport involved in self-organisation. (a) One-dimensional reaction-diffusion system (known as Schnakenberg model) that develops a stationary Turing pattern over time (activator concentrations highlighted in colours). A necessary condition is that the activator and its antagonistic inhibitor possess vastly different diffusion constants. (b) If diffusional transport of activator and inhibitor are too similar, the pattern disappears, and a well-mixed homogeneous state emerges.

Given that a detailed but often neglected nomenclature on patterns was developed decades ago in the field of nonlinear dynamics (see, for example, ref. 197), it would facilitate comparisons between studies if the existing physical nomenclature was used instead of re-inventing (or renaming) the wheel.

An impressive amount of data has emerged over the years that clearly demonstrate the interplay between transport and reactions in living matter self-organisation on many length and time scales. Patterning in reaction-transport systems has been observed, for example, in the syncytium stage of developing flies, 198,199 for the division machinery of Escherichia coli bacteria in vivo<sup>92</sup> and in vitro, <sup>97,98</sup> and for morphogen patterns that govern the embryogenesis of vertebrates. 200-202 A particularly well-studied example for transport-induced pattern formation is the PAR protein gradient in the zygote of Caenorhabditis elegans. 94,96 Here, two antagonistic PAR protein species, supported by diffusive and advective intracellular transport, build a gradient that determines not only the distinct biochemical fate of the emerging daughter cells, but also sets the anteriorposterior body axis within the first cell division. The decisive role of directed transport in this example has recently been demonstrated in an elegant experiment<sup>203</sup> in which infrared light created a counteracting flow within the zygote that flipped the gradient. Subsequent patterns (e.g. concentration and mobility gradients for MEX-5 and PIE-1 proteins 93,95,204) are triggered by this initial patterning, including even the condensation of membraneless organelles into so-called p-granules in the posterior daughter cell. 205,206

Taken together, it appears fair to say that vital self-organisation phenomena in living matter rely crucially on transport processes: unambiguously revealing and naming them is of the utmost importance to obtain a quantitative understanding of living matter. The next subsections discuss important measures for this endeavour.

### D. Time-averaged moments and ergodic behaviour

The most informative way of monitoring transport in complex media is by acquiring single-particle trajectories, *i.e.*, time series of particle positions  $\mathbf{r}(t)$  from time t=0 to t=T (the "measurement time"); the numerous experimental challenges

associated with this approach are discussed below. Commonly, time series  $\mathbf{r}(t)$  are first evaluated in terms of moments,  $\langle |\mathbf{r}(t)|^q \rangle$  of (integer or fractional) order q, with each q providing information about the system; e.g., q=2 for the MSD. While ensemble-averaged moments (like eqn (18)) are typically easier to obtain in analytical calculations, the availability of individual particle traces in experiments advocates the use of time-averaged moments  $^{117,132,174}$ 

$$\overline{\delta^{q}(\tau)} = \frac{1}{N-k} \sum_{i=1}^{N-k} |\mathbf{r}((i+k)\Delta t) - \mathbf{r}(i\Delta t)|^{q},$$
(19)

for a time series of positions  $\mathbf{r}(i)$  labelled  $i=1,\ldots,N$ , where  $T=N\Delta t$  and  $\tau=k\Delta t$  is the lag time. This expression also highlights that averaging in  $\overline{\delta^q(\tau)}$  becomes increasingly poor as  $\tau\to T$ , where individual fluctuations of trajectories become dominant. We note that a widespread method—chopping up a long time series into short segments and then evaluating as an ensemble of trajectories—works well for stationary processes. Yet, for non-stationary dynamics, such as processes with scale-free immobilisation times, this segmentation introduces spurious correlations.

Using time- and ensemble-averaged moments, a central aspect of the system can already be analysed: the system is considered ergodic when  $\lim_{\tau/T\to 0} \overline{\delta^q(\tau)} = \langle |\mathbf{r}|^q(t) \rangle$ . A system is said

to be ergodic in a practical sense when a point (such as a particle) within the system will eventually explore all possible positions and states within it in a uniform, random way. While ergodicity is often tacitly assumed to hold, many stochastic processes do indeed violate this premise. Analytical expressions for ensemble- and time-averaged moments therefore need to be calculated and tested individually 117,132,207,208 for experimental data. This becomes even more important when considering that some processes eventually become ergodic, but only with different convergences of time- and ensemble-averaged quantities. 131,132,174 "Weak ergodicity breaking", 150 i.e., disparity between time and ensemble averages MSDs, reveals important clues about the underlying system, and hence must be thoroughly probed. An important indicator is also the magnitude of the fluctuations of the amplitude  $\delta^q(\tau)$  for finite T from one trajectory to another, i.e., to which degree a process is reproducible between different realisations. The PDF of these amplitude fluctuations and its variance, the "ergodicity breaking parameter", 132,207 have been demonstrated to be reliable ways to identify the underlying stochastic process. 117,132,207,208 A related feature of nonstationary dynamics, such as those based on scale-free waiting time processes, is so-called "aging", resulting in an explicit dependence of all observables on how long the process, or the experimental measurement itself, has been running. 117,132,207-209

### E. Power spectral density

Complementing the analysis of moments, the power spectral density (PSD) of individual position time series can also be exploited (see Section III). Fourier-transforming a *d*-dimensional

trajectory leads to the PSD's k-component (k = 1, 2, ..., d),

$$S_k(f,T) = \frac{1}{T} \sum_{j=1}^k \left| \int_0^T e^{ift} x_j(t) dt \right|^2,$$
 (20)

where f denotes the frequency. The sum takes into consideration the k-component contribution of the d-dimensional trajectory  $\mathbf{x}(t) = \{x_1(t), x_2(t), \dots, x_d(t)\}$ . The classical PSD  $\mu(f)$  is the long-time ensemble average  $\mu(f) = \lim_{T \to \infty} \langle S_d(f,T) \rangle$ . Given that singleparticle tracking experiments often cannot provide sufficiently many, and long, individual trajectories, it is pertinent to work with eqn (21). Detailed analyses for Brownian and fractional Brownian motion demonstrate that the power-law decay  $S_k(f,T) \sim 1/f^{\beta}$  is preserved in individual PSDs. In addition, similarly to the amplitude variations of the TA-MSD from one finite-time trajectory to the next, the PDF  $P_k(A)$  of fluctuations of the PSD's amplitude A for different trajectories from the same ensemble show a characteristic behaviour, which has also been verified with experimental data, 154,210-212 as shown in the inset of Fig. 5c. The two curves correspond to the case of k = 2 of the two-dimensional measurement (red curve and symbols) and the one-dimensional projection (black curve and symbols). For details on the mathematical form of  $P_k(A)$  see ref. 154, 210, 213 and 214. It is also possible to extend this analysis to random-diffusivity processes, and different processes, with and without ageing, can be distinguished by the specific form of  $P_k(A)$ , even if the scaling exponent  $\beta$  is the same. 154,210,213,214 Recently it was demonstrated that the coefficient of variation of the PSD is a robust measure for anomalous diffusion in the presence of static and dynamic errors.<sup>215</sup>

### F. First passage times

Another relevant piece of information allowing for close-up viewings of a system are first-passage times, for instance how long it takes a lipid molecule to escape a given radius around its initial position. <sup>159</sup> While valuable information is often already contained in mean first-passage times, <sup>216,217</sup> reconstructing their entire PDF unveils additional information, such as the distance between the initial position and the target, target reactivity, or the physical dimension of the accessible volume. <sup>192,194,218</sup> Notably, in many cases this PDF is wide, and individual realisations show a considerable scatter in their first-passage times. <sup>219</sup> Similar to this is the growing shell technique developed in ref. 220.

### G. Velocity autocorrelation function, VACF

Another method to analyse time series is through the correlation functions of successive steps, <sup>132</sup> often referred to as the velocity autocorrelation function (VACF):

$$\overline{\text{VACF}}(l) = \frac{1}{N - l - k} \times \sum_{i=1}^{N-l-k} \frac{\left[\mathbf{r}(i+l+k) - \mathbf{r}(i+l)\right] \cdot \left[\mathbf{r}(i+k) - \mathbf{r}(i)\right]}{k^2},$$
(21)

where the instantaneous velocity is the step length within the

considered time interval  $k\Delta t$ , while  $l\Delta t$  is the lag time. The VACF probes in particular the (anti-)persistent memory kernel in random processes, allowing, for example, a pronounced anticorrelation to be observed for subdiffusive fractional Brownian motion, <sup>132</sup> as visualised in Fig. 5d. Indeed, pronounced anticorrelations in the VACF are found for subdiffusive motion in crowded dextran and worm-like chain micellar solutions, <sup>160,163</sup> in intracellular fluids, <sup>142,167,181,221–224</sup> and in lipid bilayer membranes. <sup>159</sup> The autocorrelation function of squared increments, on the other hand, can reveal stochastic switching between different mobilities, with and without a non-Markovian memory kernel. <sup>142,225</sup>

More recently, automatic classification approaches based on Bayesian statistics<sup>180,226</sup> and deep-learning algorithms.<sup>227–231</sup> have been introduced to dissect possible transport models with a given set of trajectories. Another means of extending analysis tools is through the combination of physical data analysis with mathematical time series analysis.<sup>232,233</sup> The power of such analyses was highlighted in the recent AnDi (Anomalous Diffusion) community challenge, in which given time series from simulations and experiments had to be analysed. The performance for model identification and parameter estimation was presented in ref. 230. Future AnDi challenges are already planned.

In summary, a broad palette of analysis tools is already available to extract the transport model underlying experimentally acquired data. However, it should be kept in mind that in order to make a claim about the different key features of available models, it is not sufficient to measure just one quantity but rather a combination of several data-derived observables. Key information is often encoded in the fluctuations of observables such as the time-averaged MSD, the single trajectory power spectra, first-passage times, *etc.* Moreover, the different shortcomings and uncertainties of experimental data acquisition must be considered thoroughly in this context, as described below. Awareness of the non-universality of diffusion in complex systems such as living biological cells is necessary in order to meaningfully analyse data, and not simply assign some essentially meaningless parameters.

# H. Experimental approaches and challenges in quantifying transport

Quantification of diffusive transport, especially in living matter, often relies on light microscopy techniques with high spatiotemporal resolution. The most versatile approach is single-particle tracking (SPT), the rapid imaging of a sparse set of particles and determining their individual trajectories from a time series of images. <sup>234–236</sup>

Prior to data acquisition, the choice of imaging technique for SPT must be carefully considered in relation to the problem of interest: phase contrast *versus* fluorescence, widefield *versus* confocal, and so on. Each method has strengths and weaknesses in terms of acquisition time, potential phototoxicity, and other parameters. The most widespread imaging technique in the realm of living matter is fluorescence microscopy, for which a broad palette of genetically encoded or artificial fluorophores is available for *in vivo* imaging.<sup>237–243</sup> In most

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cases, experiments are reduced to rapid, two-dimensional fluorescence imaging, where particle positions are determined only in the imaging plane. To this end, epi-fluorescence, totalinternal reflection, confocal, or lightsheet microscopy can be used, all of which allow for an acquisition speed in the range of at least ten frames per second. While two-dimensional image acquisition is straightforward and rapid, gaining information about the third dimension requires considerably more effort due to its slow speed, reduced resolution and increased amounts of data to be processed. Neglecting motion in the third dimension to improve temporal resolution, however, requires that motion along the different spatial degrees of freedom are not coupled, i.e., that the two-dimensional trajectory contains all relevant information and is more than a lossy projection of the real motion. Probing the validity of this assumption, e.g., by testing whether the remaining two spatial degrees are uncorrelated, is rarely done, but would improve the grounds of the subsequent analysis.

Having recorded a potentially meaningful image series, a number of quality controls must be made before a more refined analysis in terms of MSDs, PSDs, etc. can be performed. First and foremost, drift and fluctuations of the microscopy stage and/or the entire specimen need to be removed. Correcting for this global centre-of-mass motion might require additional SPT experiments on fixed cells, or tracing either cell adhesion areas or even endomembranes to eventually reveal the relative motion of tracer particles. While fixation may be useful for some controls, it is important to notice that for experiments investigating cellular dynamics or transportation phenomena, fixed cells cannot be used as fixation destroys any biological activity.

A challenging next step involves identifying the labelled particles in each frame and combining their frame-wise positions to trajectories; failure to do so results in mixing up particle identities within a single trajectory, leading to an erroneous analysis. Often, the image contrast has to be enhanced locally, e.g., by bandpass filtering, to highlight local intensity maxima. Then, the maxima in each frame have to be assigned to a set of moving particles by some probabilistic measure. Many algorithms for SPT evaluation are available for the community, such as plugins for ImageJ/FIJI, or even as source codes for IDL/Matlab. Which to pick for data evaluation is a nontrivial decision, a choice which may be guided by a comprehensive comparative study.<sup>244</sup> It is evident, however, that all of these algorithms have internal degrees of freedom that potentially introduce artifacts to the extracted positions. For example, allowing for blanks in a trajectory and simply interpolating the missed position is not good practice, even though many tracking algorithms do just this. Using interpolated, basically diced positions might render trajectories more Markovian, i.e., existing memory effects within a trajectory are deliberately diminished, hence altering correlation functions like the VACF.

Even when carefully performing all of these steps, one ought to bear in mind that all particle positions inevitably have static and dynamic localisation errors<sup>245-248</sup> that require careful consideration in subsequent analyses. First, the position for

even an immobile particle can only be determined with an accuracy dependent upon the number of acquired photons  $(n_p)$ as  $\sim 1/\sqrt{n_{\rm p}}$ ; this static localisation error is basically the usual standard error of the mean. Second, recording an image to determine a particle's position takes a finite time during which it is in constant motion. As a result, the particle's position is smudged, creating a dynamic localisation error. Both localisation errors will affect the MSD and the VACF, 248 suggesting a different scaling exponent α and/or an altered (anti-) correlation in the VACF. Even worse, both localisation errors can induce opposing artifacts and it is a priori unclear which of the two will be dominant or if they even cancel each other out. For example, the static localisation error adds a positive offset to the MSD, 246,248 whereas the dynamic localisation error effectively adds a negative offset, 245,248 compare Fig. 5b. It turns out that it is possible to circumvent this issue in the MSD when using a resampling approach of the very same trajectories, 153 which typically allows for a more stable determination of the scaling exponent  $\alpha$ . At the same time, one needs to be aware of the finite length of experimentally determined trajectories, frequently in the range of N = 100 positions per trajectory. With short trajectories, for instance, the apparent scaling exponents derived from time-averaged MSDs feature a fairly wide distribution that hardly reveals the properties of a random walk. 153 Often, all averaging procedures discussed in the preceding section are hampered by finite-size effects when dealing with experimental data. Concurrently, mean-maximal distancebased analyses may be profitable.249

Taken together, the transportation measures discussed above can only be applied to experimental data with the caveat that they have been derived assuming long trajectories and perfectly known positions. In order to derive meaningful analysis tools for experiments, this uncertainty and the aforementioned superimposed noise need to be carefully taken into consideration. 178-180 We mention that additional challenges may be posed by the precise experimental protocol, e.g., when temperature or ATP concentrations are modified during the experiment, 250 or when the action of poisonous substances decay over time.<sup>251</sup>

Besides SPT, a number of ensemble-based measurement techniques for quantifying transport are available. These are good alternatives especially in less dilute systems in which individual particles may not be distinguishable from one another. It is important to note that they do not provide individual trajectories, instead reporting only on the MSD and assuming, more or less tacitly, spatial homogeneity. Frequently employed techniques are fluorescence recovery after photobleaching (FRAP)<sup>252,253</sup> and fluorescence correlation spectroscopy (FCS).<sup>254-258</sup> FRAP determines the half-time of diffusion-driven fluorescence recovery in a region in which dye molecules have been photobleached, whereas FCS relies on monitoring the fluctuations of stationary fluorescence signals to determine the mean residence time of particles in the focus. Complementary to both, a scattering technique called differential dynamic microscopy (basically small-angle dynamic light scattering) has been introduced and applied more recently. 259-263 If and to what extent these different methods provide comparable results to SPT, and

determining which is the most accurate in quantifying transport within a given system, is a nontrivial task (see, for example, 264-267 for some comparative studies). Compared to SPT, however, ensemble-averaged techniques provide significantly more limited information: inferring the actual nature of the transport process from these data is typically more than challenging.

### V. Conclusion

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Progress in quantifying and understanding living matter depends on novel experimental and theoretical methods to dissect the impact of different biological, physical, and chemical key factors and to uncover the underlying biological mechanisms. This Perspective focuses on how tracking single tracer particles enables quantification of materials properties, bio-mechanical forces and transportation mechanisms relevant for living organisms. A critical aspect when comparing such data obtained by different experimental platforms, or even from the same platforms but by different research groups, is to identify and use comparable experimental parameters, and similar data analysis strategies and to be critically aware of the limitations of the experimental methodology. Successful progress in quantifying and understanding dynamical and biomechanical properties of living systems is thus critically dependent on a mutual agreement between biologists and physicists to standardise their methods based on single particle tracking, on the use a common nomenclature, and on striving to achieve comparable data mapped in a universal language across disciplines. This Perspective is intended to provide a first and stimulative step towards harmonising experimental and theoretical approaches based on single particle tracking techniques at the interface of physics, chemistry, and the life sciences. Such inter-disciplinary harmonising effort is needed to exploit the emerging experimental platforms, including, e.g., super-resolution and light-sheet microscopy, and to advance our quantitative understanding of living matter.

### Conflicts of interest

There are no conflicts to declare.

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