

1 **Novel advances in the design of three-dimensional Bio-**
2 **scaffolds to control cell fate: translation from 2D to 3D**

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4 **Edorta Santos^{a, b}, Rosa M^a Hernández^{a, b}, Jose Luis Pedraz^{a, b}, Gorka**
5 **Orive^{a, b, *}**

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8 ^a NanoBioCel Group, Laboratory of Pharmaceutics, University of the Basque Country,
9 School of Pharmacy, Vitoria, Spain

10
11 ^b Biomedical Research Networking Center in Bioengineering, Biomaterials and
12 Nanomedicine (CIBER-BBN), Vitoria, Spain

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14 *Corresponding Author: Gorka Orive, PhD
15 Laboratory of Pharmaceutics, University of the Basque Country, School of Pharmacy,
16 Vitoria, Spain
17 gorka.orive@ehu.es

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1 **ABSTRACT**

2

3 Recreating the most critical aspects of the native extracellular matrix is fundamental to
4 understand and control the processes regulating cell fate and cell function. From the ill-
5 defined complexity to the controlled simplicity, we discuss the different strategies that
6 are being carried out by scientists worldwide to achieve the latest advances in the
7 sophistication of three-dimensional scaffolds, stressing their impact on cell biology,
8 tissue engineering and regenerative medicine. Synthetic and naturally derived polymers
9 like polyethylene glycol, alginate, agarose, etc., together with micro- and
10 nanofabrication techniques are allowing the creation of three-dimensional models where
11 biophysical and biochemical variables can be modified with high precision,
12 orthogonality and even in real-time.

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15 **Cellular scaffolds: more than a platform**

16 Three-dimensional biosystems are evolving rapidly. Their continuous sophistication is
17 accelerating our nascent understanding of cellular microenvironment and how the basic
18 building blocks of biological systems are integrated into the dynamic landscape of
19 tissue physiology. Traditionally, cellular scaffolds, from the typical 2D polystyrene
20 surfaces to the first 3D constructs (natural or artificial), were intended as inert platforms
21 that merely served as support for the cultured cells. Since then, more emphasis was
22 given to provide these matrices with suitable physical (e.g. stiffness and mass transfer)

1 and chemical (e.g. employed material type and degradation rate) properties for tissue
2 engineering and cell transplantation [1]. More recently, the biology of the scaffolds is
3 gaining the attention of scientists, including signals that cells receive via adhesion to the
4 material or directly from soluble factors in the microenvironment [2,3].

5 It is demonstrated that cells are able to sense and interpret the information coming from
6 the extracellular matrix (ECM) responding and reorganizing in function of topography
7 [4,5], mechanical properties (e.g. stiffness, viscosity and elasticity) [6-8], molecules
8 presented by the ECM [9] and concentration gradients of both soluble and tethered
9 growth factors [10]. Thus, cells receive and process a multiple combination of
10 physicochemical and biological cues always within a spatio-temporal context and in
11 three main ways: cell-cell contacts [11], cell-ECM interactions and cell-soluble/tethered
12 factor interactions [12].

13 Interestingly, the inspiration that guide the design of new biomaterial approaches is
14 always drawn from the observation on various length scales of the materials arranged
15 naturally by the cells in the tissues [13]. Thus, gaining insight into so far unknown
16 questions motivates the design of new models that allow for investigating more
17 thoroughly the cell-ECM interaction and its effects in a feedback manner.

18 Biomaterial strategies are bridging the gap in many scientific fields, as they have
19 become a necessary tool in tissue engineering or regenerative medicine among others.
20 In fact, microfabrication, and more recently nanofabrication [14] are allowing the
21 creation of suitable models where key factors may be studied from the nanometer to the
22 supramillimeter length scale [15,16]. Moreover, the ability of the new bioinspired
23 materials to be tuned in a wide range of biophysical and biochemical features is also
24 optimizing the way scaffolds control the different biological properties of the cells.

1 Here, we present the latest advances in the sophistication of 3D scaffolds and its impact
2 on cell biology, tissue engineering and regenerative medicine.

3

4 **Translation from 2D to 3D**

5 Recent findings suggest that cells often show a non-natural behavior when they are
6 moved away from their natural niches and seeded onto flat substrates. Only to mention
7 some examples, breast epithelial cells exhibited a tumoral trend when assayed in 2D
8 (likely due to the unnatural extreme stiffness provided by the plate), while regressed to
9 normal state upon transferal to 3D models resembling their natural niche [17]. In the
10 same way, increased chondrogenesis have been noticed in embryonic stem cells (ESCs)
11 cultured as 3D embryoid bodies when compared to the monolayer conformation [18].
12 Therefore, while 2D experiments represent a versatile and accurate way to screen the
13 effects of isolated compounds of the ECM on cells (Box 1), 3D experiments are
14 designed to direct a progressive and steady reconstruction of the complexity that entails
15 the native ECM.

16 The disparities in cellular function described between 2D and 3D approaches are mainly
17 given by the manner in which cells perceive their surrounding microenvironment. Cells
18 plated onto 2D substrates are polarized, maintaining only part of their surface anchored
19 and exposing remaining parts to the culture media. Moreover, the contact with
20 neighboring cells is also limited to the flat edges that share each-others. This is in sharp
21 contrast with the natural environment of the tissues, where each cell closely interacts
22 with the nearby cells and the ECM [19]. Hence, 3D environment-based interplay
23 reflects a more distributed integrin usage and enhanced biological activity [20]. Mass

1 transport physics is also absolutely altered. Growth factors, morphogens, cytokines and
2 so forth quickly diffuse in the media of 2D cultures, reaching cells uniformly, whereas
3 native ECM produces chemical and biological diffusion gradients that play a key role in
4 signaling and tissue development [21]. In addition, cell shape also has its influence on
5 cell commitment. Once again, cells on 2D cultures are limited to a planar and spread
6 morphology and do not experience the more complex morphologies found in vivo [18].
7 Furthermore, 2D surfaces offer almost undetectable resistance to cell migration, which
8 contrasts notably with the mechanical interactions provided in vivo.

9 Therefore, the design of 3D models that resemble more or less accurately the native
10 ECM is crucial in order to obtain reliable results that approximate to reality.
11 Nonetheless, mimicking the ECM in the lab is not simple, especially because there is
12 much we do not know yet about the cell-ECM cross-talk that occurs in vivo. As a
13 consequence, the most frequently used models so far have been hydrogel scaffolds
14 formed by animal ECM-derived proteins, Matrigel[®] or Vitrogen[®] among others [22].
15 Even if these biosystems have provided seminal understanding for cell biology field in
16 the past few decades, they are far from being ideal. Limitations include: (i) reduced
17 flexibility to modulate a cell's biophysical and biochemical properties (and furthermore,
18 to control such variables independently), (ii) immunogenicity, (iii) batch-to-batch
19 variability and (iv) ill-defined complexity that provides too little mechanistic
20 information to the cell [12,22].

21 Assuming some of these limitations, hydrogels formed by synthetic polymers like poly
22 ethyleneglycol (PEG), and naturally derived polymers including alginate, agarose,
23 chitosan etc. have become the biomaterial of choice for artificial ECM reconstruction.
24 Hydrogels are able to resemble the nature of most tissues due to their high water

1 content, and the presence of pores that facilitate the free diffusion of oxygen, nutrients
2 and growth factors, morphogens, etc. [18,23]. Most importantly, many hydrogels offer
3 the possibility to encapsulate cells under gentle and cytocompatible conditions, and
4 furthermore, their physicochemical properties can be easily tuned [24]. Indeed,
5 nonfouling polymers like PEG present the additional advantage of minimizing
6 uncontrolled variables such as protein adsorption and their indirect effects, which makes
7 PEG one of the most attractive biomaterials for 3D scaffold designs. On the contrary,
8 one of the reasons alginate has been so widely employed so far is that it can be further
9 injected in vivo in a minimally invasive way, undergoing gelation in situ.

10

11 No single model can recapitulate the whole complexity of every tissue type ECM. Many
12 authors agree on the fact that high level of complexity is not necessary for many
13 applications, and indeed simpler and practical models are enough to solve some specific
14 questions [12,16,25]. In fact, cells enclosed within 3D matrices rapidly remodel their
15 microenvironment depositing their own ECM molecules [22,26]. For that reason, it is
16 possible to compensate the lack of such complexity with artificial systems capable of
17 inducing desired effects to the hosted cells in a more efficient and rational way [27]. For
18 example, presenting cells with tethered small-molecule chemical functional groups was
19 enough to recreate unique chemical environments and induce multiple MSC
20 differentiated lineages [28]. Nonetheless, if the goal is tissue-like structures for
21 regenerative medicine for example, higher complexity levels in time and space are
22 absolutely justified [22].

23

1 **Sophistication of 3D biosystems**

2 *Adhesion ligand presentation*

3 Cells may be provided with adhesion surfaces by using a variety of naturally derived
4 ECM molecules such as collagen or fibrin, or using these molecules to decorate
5 synthetic polymers to which adhesion is regulated by adsorbed proteins (Box 2).
6 However, protein engineering allows us to isolate functional domains within large ECM
7 molecules and incorporate them into otherwise inert substrates. Thus, epitopes that
8 mediate cell-adhesion can be mimicked using synthetic peptides. Among them, perhaps
9 the most known are arginine-glycine-aspartic acid (RGD), derived from fibronectin, and
10 tyrosine-isoleucine-glycine-serine-arginine (YIGSR), derived from laminin. PEG
11 hydrogels can also be modified by novel polymerization mechanisms such as thiol-ene
12 [29] and thiol-acrylate chemistries [30], while other polymers like alginate are usually
13 modified by means of carbodiimide chemistry [31].

14 It has been observed that not only the adhesion moieties themselves, but also their
15 density and spatial distribution on micrometer and nanometer scales influence cell fate
16 [32]. By manipulating the way adhesion moieties are presented to the cells, it is possible
17 to induce major cellular processes such as migration, proliferation and differentiation
18 [33]. With this idea, nanoscale patterns of RGD islands in hydrogels have been varied
19 without altering the final ligand density. For instance, hydrogels with reduced island
20 spacing were produced by uniformly distributing alginate chains containing a single
21 ligand, while more increased island spacing was achieved by mixing unmodified chains
22 and chains coupled with multiple peptides (Fig. 1A). Thus, more closely spaced island
23 favored cell spreading, while more widely spaced islands supported differentiation [34].

1 Beyond these approaches, much attention has been lately paid to the patterning of
2 adhesive moieties. Biomolecules can also be immobilized in micropatterned volumes
3 within agarose gels with a multi photon laser [35]. When agarose or hyaluronan is
4 covalently modified with a derivative of cysteine protected with a photocleavable group,
5 the protection groups can be removed upon exposure with a laser beam (micrometric
6 resolution). As a result, desired oligopeptides are covalently immobilized in patterned
7 sites via Michael-type addition (Fig. 1B). Using this procedure and by means of
8 orthogonal physical binding pairs (barnasa-barstar and streptavidin-biotin),
9 simultaneous patterning of multiple growth factors has been achieved to direct neural
10 precursor cell differentiation [36]. Two-photon laser scanning (TPLS) photolithography
11 in PEG diacrylate (PEGDA) hydrogels, can also be used to guide encapsulated dermal
12 fibroblasts with precisely patterned RGD moieties [37].

13

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15 *Mechanical properties*

16 The most common way to control scaffold stiffness is by using polymers and cross-
17 linkers at different concentrations or varying the molecular weight of the polymers. In
18 this way, hydrogels formed by macromers of PEG and poly(lactic acid) (PLA),
19 modifying the initial macromer concentration from 10% to 20% result in gels with
20 elastic moduli increased from 60 to 500 kPa. The latter is used to restore initial function
21 in chondrocytes and facilitate the production of cartilaginous production [38]. The
22 compressive modulus of hydrogels comprising HA-tyramine conjugates can be
23 controlled by varying the concentration of hydrogen peroxidase (H_2O_2) employed to

1 produce the covalent cross-linkage. Thus, lower cross-linked matrices give as a result
2 enhanced chondrogenesis in encapsulated MSCs, while increasing cross-linking degree,
3 and thereby matrix stiffness, MSC differentiation towards fibrous phenotypes is induced
4 [39].

5

6 A blend of high MW and low MW alginates gives rise to highly cross-linked hydrogels;
7 the mixture has a pre-gelled viscosity similar to that of pure high MW at low
8 concentrations. Hence, rheological and mechanical properties can be decoupled:
9 scaffolds can have a high elastic modulus but cells are not sheared during encapsulation
10 [31]. Employing this procedure it has been demonstrated how MSCs are able to
11 reorganize the adhesion ligands on the nanoscale in function of the stiffness offered by
12 alginate matrix where they were encapsulated. This adhesion ligand reorganization
13 process may play an important role in MCS commitment [6].

14

15 *Chemical signaling*

16 The regulation of soluble molecule distribution within 3D scaffolds is difficult because
17 the availability of biomolecules depends on (i) the total concentration in the medium,
18 (ii) diffusion rate within the gel, and (iii) cellular metabolic activity [18]. In addition,
19 artificial ECMs may also require the presence of growth factors and morphogens in a
20 pharmacokinetic manner that resembles the natural cell niche For example, during
21 angiogenesis vascular endothelial growth factor (VEGF) promotes the proliferation of
22 endothelial cells at the first steps of the process, while platelet derived growth factor
23 (PDGF) regulates the maturation of these new created vessels a posteriori. Therefore,

1 similar pharmacokinetics must be reproduced to succeed in the attempt to induce
2 angiogenesis (Fig. 2A). In this sense, different approaches have been carried out in the
3 attempt to regulate the kinetics and distribution of soluble factors. In an attempt to
4 mimic the native ECM, where glycosaminoglycans act as depots for growth factors,
5 heparin was incorporated into the scaffold backbone for posterior sequestering and
6 controlled release of growth factors [40]. Similarly, HA hydrogels have been designed
7 to ongoing different degradation rates and further provide controlled release of cationic
8 proteins such as bone morphogenetic protein-2 (BMP-2) and VEGF anchored to the
9 matrix via electrostatic interactions [41]. Other approaches propose covalent linkage of
10 specific ligands from the desired molecules to the scaffold (also known as phage
11 display) [42]. By including multiple soluble factors within different encapsulation levels
12 (e.g. PLG spheres within alginate hydrogels), it is possible to sustain a simultaneous or
13 sequential factor delivery. The significance of exerting control over growth factors
14 availability in time and space has been probed, for instance, in stem cell differentiation
15 or functional repair of segmental bone defects [16,43,44] or therapeutic approaches to
16 induce angiogenesis [45].

17 Soluble biomolecules often show improved bioactivity when they directly attached to
18 the hydrogel network [46]. In addition to improved stability, covalently immobilized
19 growth factors can be used to spatially direct cell behavior (e.g. chemotaxis or
20 differentiation) [47]. However, it is important to ensure that active domains of the
21 molecules are accessible upon covalent linkage.

22 On the other hand, cells are exposed to gradients of morphogens, growth factors and
23 cytokines progressively in physiological tissue. They play a key role not only in
24 morphogenesis, chemotaxis and axogenesis [52], but also during processes like wound

1 healing or tissue homeostasis. Such gradients can be introduced into 3D models, for
2 instance, using the same micropatterning techniques described above to attach ligand
3 moieties [48]. In this way, endothelial cells (EC) tubule-like formation was guided
4 through VEGF gradients patterned within RGD-modified agarose hydrogels [49].

5 Microfluidics-based systems are also increasingly being used to generate gradients
6 within 3D models [15][50]. These platforms represent one of the most accurate and
7 robust ways to reproduce morphogen gradients given in vivo, as they allow small
8 amounts of expensive factors to be patterned into scaffolds with tight control [47].
9 Some approaches have already been carried out. For example, microfluidic channels
10 have been embedded directly in cell enclosing alginate scaffolds; the channels control
11 the distribution and flux of solutes in the total volume by means of convective mass
12 transfer (Fig. 2B) [51]. Moreover, since biomolecules can also be tethered to the
13 backbone of artificial ECMs, applying microfluidics technology with anchored proteins
14 would give rise to more comprehensive and realistic ECM surrogates [50].

15

16 *Mass transport and matrix permissiveness*

17 One of the prime points that concerns scientists when it comes to leap to 3D is the fact
18 that cells may suffer the lack of gases and nutrients. This is especially evident in vivo,
19 where encapsulated and transplanted cells rely on diffusion for oxygen and nutrients
20 from surrounding blood vessels. The maximum thickness for these biosystems has been
21 reported to be limited to approximately 100-200 μm [52]. Moreover, 3D culture designs
22 present physical constraints that hamper cell proliferation, migration and morphogenesis.
23 In general, pore sizes of less than 1 μm are able to support free diffusion of molecules,

1 but not cellular migration, whereas pores in the range of $\approx 10\text{-}100\ \mu\text{m}$ readily allow host
2 cells to migrate through the entire volume of the scaffold [53]. Most chemically cross-
3 linked polymer hydrogels form mesh-like structures with pores on the order of tens of
4 nanometers, which means that they are small enough even to prevent cellular events
5 such as the formation of filopodia [22]. Thus, cells remain literally trapped within their
6 microvoids, showing round morphology. Nonetheless, using smart engineering tricks
7 researchers have managed to improve mass transport conditions of their scaffolds and
8 increase the functionality of the enclosed cells.

9 By assembling a PEG hydrogel in the presence of crystal colloidal templates that could
10 be further removed by solvent extraction (“leaching”) scaffolds with a pore range of 20-
11 $60\ \mu\text{m}$ were formed [54]. Another alternative approach is the use of CO_2 as porogen in
12 the production of PEG scaffolds with interconnected pores ranging in size from 100 to
13 $600\ \mu\text{m}$, which were used to promote osteogenesis in MSCs [55]. Similarly, two-photon
14 lasers can be used to direct the patterned patternalized polymerization of multifunctional
15 acrylate monomers. This technique uniform 12 to $110\ \mu\text{m}$ pores and can be used to
16 study cell migration on basis of pore size [56]. A more recent work showed that
17 permeability can be readily improved in PEG hydrogels incorporating hydrophobic
18 nanoparticles that induced partially looser cross-linking density. By these means
19 viability and functionality of encapsulated cells was improved without altering scaffold
20 mechanical properties [57].

21 Matrix permissiveness also drastically influences tissue morphogenesis. Matrix
22 degradation is fundamental, so that encapsulated cells can remodel the scaffold by
23 secreting their own ECM molecules.. In addition, degradation allows cell migration and
24 regulates the release of matrix-tethered biomolecules that induce different cellular

1 functions [58]. Apart from the scaffolds formed by ECM derived molecules, which
2 present inherent degradability, it is possible to design inert matrices which can be
3 degraded according to different strategies. For example, synthetic hydrogels can be
4 designed to include degradable polymers within their network. Some studies describe
5 the use of poly(lactic acid) [59] or poly(caprolactone) [60] blocks in combination with
6 PEG backbone. Similarly, the scaffolds can be built by co-polymerization of different
7 ratios of degradable and non-degradable macromers [61]. For all these types of designs,
8 the degradation rate is governed by the number of hydrolytically labile bonds in the
9 hydrogel, although in general, normal cellular processes take place in less time than that
10 required for these labile bonds to be degraded. are on another scale faster than
11 mentioned rate [18].

12 In alginate, a well known strategy to control the degradation rate of the scaffolds is
13 partial oxidation of the main chains to create controllable numbers of functional groups
14 in the backbone susceptible to hydrolysis [62]. Hydrolytically labile hydrogels have
15 predictable degradation profiles, but the properties cannot be altered after gelling and
16 the degradation rate is both uniform and independent of cellular interactions. In order to
17 permit cellularly driven matrix degradation, synthetic hydrogels, such as those formed
18 by PEG acrylate, can be modified by Michael addition and photoinitiated reactions to
19 include specific sequences that are recognized and cleaved by proteases like matrix
20 metalloproteinases (MMPs) secreted by cells [18]. This design enables cells to locally
21 remodel their surrounding matrix and deposit their own ECM proteins on the matrix,
22 mimicking more realistically what occurs in vivo during wound healing, regeneration or
23 tumor metastasis [22].

24

1 *Nanofabrication*

2 A typical cell size ($\approx 7-15 \mu\text{m}$) is similar to or smaller than the so far described hydrogel
3 microstructures. Thus, some authors argue that the range of microporosities ($\approx 10-100$
4 μm) will effectively act as 2D surfaces with curvature for cell attachment [18,19]. One
5 possibility to address this problem is the fabrication of nanofibrillar nanofibilar
6 architectures. In fact, the necessity to understand in detail the nature of the native ECM
7 has fueled new paths towards the fabrication of biomimetic scaffolds with nanoscale
8 properties. Starting with the natural fibrous mesh of the ECM, it is possible to construct
9 novel scaffolds with interconnected and porous structures formed by interwoven fibers
10 with similar diameters to those presented by collagen fibers [14]. Thus, the forces
11 exerted by cells in the scaffold influence further material structural reorganization [19].
12 In this sense, electrospinning [63] and molecular self-assembly [64] (Box 3) are
13 increasingly growing nanofabrication techniques used to create 3D scaffolds of
14 interwoven fibers that resemble collagen structures of the native ECM [14].

15

16 *External control of spatio-temporal signal presentation*

17 The native ECM is highly dynamic. Therefore, the temporal and spatial variability
18 typical of ECM properties must also be introduced into 3D models in order to simulate
19 contextually meaningful and realistic microenvironments. Local modifications of the
20 environment at certain times can force a few cells to adopt decisions and develop new
21 functionalities, which may give rise to start a hierarchical reorganization at the
22 multicellular scale, reproducing those processes that take place in the nature (Fig. 3A-B)
23 [65]. For example, it has been reported that cells tend to invade stiffer areas guided by a

1 process known as “durotaxis” [66-68]. It has been observed a noticeable increase in the
2 elastic modulus of fibrotic tissues formed as a result of processes like acute myocardial
3 infarction [66]. To shed light on whether “durotaxis” is the mechanism promoting MSC
4 homing to these injured zones, an ideal 3D in vitro model would allow certain in situ
5 manipulation to recreate such physiological situation. Therefore, the creation of models
6 that can be externally manipulated in time and space results very advantageous to study
7 cell-ECM dynamic interplay.

8 With this aim in mind, a photodegradable PEG-based hydrogel model has been
9 developed which has predictable degradation rate patterns and stiffness gradients in
10 real-time under cytocompatible conditions (long-wavelength UV light). The gel can be
11 further manipulated at the micrometer-scale resolution with light guided gel patterning
12 [69]. As a result, cell behavior can be conditioned in situ within a 3D environment, for
13 example, by creating elastic modulus microgradients with well defined structures at
14 desired times [65]. This technology can be also employed to dynamically alter other
15 biophysical and biochemical properties. For instance, chondrocytes show an enhanced
16 differentiation in a scaffold with photolabile RGD moieties when the moieties are
17 removed at certain time points during 3D cell culturing [70].

18 “Click” reactions, can also be used to attach varying concentrations of biomolecules
19 (adhesion ligands in this case) to a scaffold backbone by means of cytocompatible
20 photolithographic patterning (Fig. 3D) and/or focused laser light guided gel patterning
21 (micrometer resolution) (Fig. 3E) after cell encapsulation [71]. Taking into account that
22 photoreactive groups for patterning are coupled with enzymatically degradable
23 sequences, this approach represents a valuable strategy to build artificial ECMs in vitro
24 with the possibility to modulate a wide number of variables in a spatio-temporal way.

1 More recently, it has been described the development of a novel light-based strategy
2 that enables the combination of mutually exclusive technologies to date. Thus,
3 biophysical and biochemical properties of the hydrogel can be controlled in a
4 independent way with orthogonal photoreactions in real-time (Fig. 3F-G) [72].

5

6

7 **Concluding remarks: challenges and future directions**

8 There is a wide range of possibilities to build 3D scaffolds. Design considerations are
9 varied according to the intended use and pursued goal. For instance, researchers
10 interested in the study of cell migration through given biomolecule gradient in vitro,
11 will possibly prefer the use of synthetic hydrogels like PEG to create their own patterns.
12 On the contrary, those more interested in forming bone-like tissue within scaffolds in
13 vivo will probably choose polymers such as alginate that can be easily injected to form
14 hydrogels once implanted. In any case, the ideal model should offer wide possibilities to
15 tune and modulate structural and mechanical properties such as elastic modulus, pore
16 size or topography. Moreover, all biophysical and biochemical properties should allow
17 independent manipulability (orthogonality) from each other. For example, increasing
18 polymer concentration to achieve a higher elastic modulus should neither affect
19 adhesion-ligand density nor mode of presentation.

20

21 It should be considered if simplifying is the best strategy for intended scaffold utility.
22 Lately for example, the type of cell attachment that is provided to promote cell-substrate
23 interactions is under debate, since it is not clear whether this biofunctionalization is

1 better to be accomplished by short peptides like the so far well-known RGD or, on the
2 contrary, by full ECM proteins like fibronectin or collagen [73]. Those who root for the
3 use of short synthetic peptides base their arguments on the fact that these short peptides
4 are chemically well defined and, thereby, much easier to isolate and understand the
5 effects caused by their use. Indeed, this precise composition avoids the uncertainty of
6 possible adverse effects providing a favorable ending through regulatory pathways.
7 Finally, the possibility to alter the ligand type, density or presentation patterns results in
8 a more interesting model. On the other hand, researchers against this motion state that
9 the integrin-mediated signaling mechanisms are much more complex and cannot be
10 completely reproduced by isolated RGD moieties. In fact, it has also been demonstrated
11 the relevance of the synergistic sites found in natural ECM proteins to regulate cell fate.

12

13 It has to be realized that understanding the complexity of the whole tissue physiology
14 by deconstructing its building blocks and studying their effects in an isolated way is
15 challenging by itself, since contextual meaning is lost. Perhaps, the real challenge lays
16 on finding a balance that would allow us to study well defined and controllable variants
17 while taking advantage of the biological mechanisms that we do not understand yet.

18 Likely, in the future, 3D models will be replacing much of those routine procedures so
19 far performed on 2D flat surfaces. Indeed, as the technology advances and we gain new
20 insights into the mechanisms that regulate cell-ECM interactions, we will be able to
21 design more sophisticated and tailor-made 3D scaffolds for the study of particular tissue
22 physiologies, always having in mind that the only true results are those validated in
23 vivo.

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15

16

1 GLOSSARY BOX

2

3 **Anoikis:** Cell apoptosis induced by lack of correct cell-ECM attachment.

4

5 **Carbodiimide chemistry:** Compounds containing the carbodiimide functionality
6 (RN=C=NR) are usually used to activate [carboxylic acids](#) towards [amide](#) or [ester](#)
7 formation. Additives, such as [N-hydroxybenzotriazole](#) or [N-hydroxysuccinimide](#), are
8 often added to increase yields and decrease side reactions. Thus, it is a common way to
9 attach peptides to the hydrocarbonated backbone of certain biomaterials.

10

11 **“Click” reactions:** simple orthogonal reactions starting from small molecular groups
12 that do not yield side products and that give heteroatom-linked molecular systems with
13 high efficiency under a variety of mild conditions.

14

15 **Focused laser light guided gel patterning (single or multi-photon):** Patterning
16 technique with micrometric resolution that enables full three-dimensional control over
17 the whole volume of the hydrogel. A laser beam (either single or multi-photon) is used
18 along with a confocal microscope to direct photoreactions at well defined points within
19 the hydrogel matrix.

20

21 **Ligand island spacing:** Distance between adhesion ligand clusters within patterned
22 hydrogels.

23

24 **Michael-type addition:** 1,4-addition of a doubly stabilized carbon nucleophile to an
25 α,β -unsaturated carbonyl compound. This reaction is one of the most employed methods

1 for the mild formation of C-C bonds. It belongs to the larger class of [conjugate](#)
2 [additions](#).

3

4 **Photolithographic patterning:** Masked light is directly focused onto the hydrogel,
5 allowing photoreactions in precise regions within the hydrogel. These regions are
6 defined by two-dimensional patterns included in the photomask.

7

8 **RGD:** Specific adhesion ligand epitope formed by arginine, glycine, aspartic acid,
9 which is originally found in fibronectin, vitronectin, fibrinogen and osteopontin among
10 other natural ECM proteins. It is used to promote integrin mediated cell-ECM
11 attachment.

12

13 **Thiol-ene chemistry:** The thiol-ene reaction is an [organic reaction](#) between a [thiol](#) (-
14 SH) and an [alkene](#) (-C=C) forming a [thioether](#). One of its main advantages Thiol-ene
15 reactions can be readily induced by a photoinitiator. This reaction is considered to be
16 cytocompatible, bio-orthogonal and is included into the group of "click" reactions.

17

18 **YIGSR:** It is found in the laminin of natural ECMs and represents an integrin mediated
19 binding domain for cells. It is composed by tyrosine-isoleucine-glycine-serine-arginine.

20

21

22

23

1 TEXT BOXES

2 BOX 1. 2D approaches, overview

3 2D cell culture approaches have given rise to important advances exciting results, many
4 of which have been pivotal in the understanding of cell-ECM interaction [66,74].
5 Deconstructing 3D complexity into 2D simple models is a smart way to perform
6 univariable experiments to parse out the effects of isolated factors (e.g. matrix elasticity
7 or adhesion ligand density)– either natural or synthetic – in cells [50]. Microarray
8 technology and combinatorial and high-throughput screening (CTHS) approaches are
9 powerful tools to make infinite combinations of structural, biophysical and biochemical
10 parameters and thus elucidate some of the mechanisms that dictate cell biology (Fig. I)
11 [75,76]. Other advantages of 2D models include the facility to exert a precise control
12 over chemical and topographical properties even at nanometer scale, the overall
13 straightforward processing and the possibility to harvest the cells effortlessly.

14

15 **Box 1. Fig. I.** Schematic summary of 2D microarrays systems to test the impact of
16 multiple biophysical and biochemical variables on cell fate. Each panel is subdivided in
17 two parts. The top part (side view) reflects cells seeded onto uniquely engineered
18 substrates (grey). Below, color blocks represent the different signals that are presented
19 (view from above). (a) Substrate displaying individual molecular signals. (b) Different
20 signal mixtures can also be displayed to explore their combinatorial effects. (c)
21 Substrate stiffness is varied, for example, to study the influence of mechanic properties
22 on cell fate. (d) Adhesion ligand peptides have been printed forming spots of different

1 sizes to control cell shape and, consequently, cell fate. Reprinted and adapted from [50]
2 with permission from Macmillan Publishers Ltd: Nature, copyright 2009.

3

4

5

6 BOX 2. Cell-ECM interaction through focal adhesions and mechanosensing

7 **Focal Adhesion**

8 One of the most relevant ways to establish cell-ECM interaction is given through
9 integrin-mediated adhesions, which cells use to connect cell cytoskeleton to adhesion
10 molecules, such as fibronectin or laminin, located on the fibers [77]. This phenomenon
11 is known as focal adhesion (FA), which constitute specific types of large
12 macromolecular assemblies through which both mechanical force and regulatory signals
13 are transmitted. FAs serve to guide the cell through the ECM; these linkages induce the
14 arrangement and polarization of cell cytoskeleton. Furthermore, FA is absolutely
15 necessary to prevent anoikis in anchorage dependent cells [33].

16 **Mechanosensing**

17 Mechanical properties of biomaterials can also influence cell commitment behavior and
18 lineage differentiation [78]. Mechanosensing is an active cellular process that entails a
19 dynamic and reciprocal interaction between the ECM and the motor proteins that are
20 connected to the cytoskeleton [13]. Cells do not only exert forces, but also respond to
21 the resistance sensed through cytoskeleton organization/tension. These external forces

1 trigger a series of intracellular signaling pathways that activate or inhibit gene
2 expression [79]. In this context, elastic substrates with variable matrix rigidities can be
3 used to study the traction forces exerted by cells and to establish correlations with
4 triggered effects (Fig. II).

5

6 **Box 2. Fig. II.** The influence of matrix mechanical properties on cell fate behavior.
7 Scheme illustrating the elastic modulus scale of different tissues ranging from the
8 softest (brain) to the stiffest (bone). Reproduced with permission from ref [66].

9

10

11 BOX 3. Electrospinning and self-assembly, general concepts

12 **Electrospinning**

13 Electrospinning is a technique in which different polymer fibers (natural and synthetic)
14 are deposited on a defined substrate by means of an electric field [80]. Resulting
15 scaffolds present continuous fibers with high porosity. The nanofibers can be orientated
16 to recreate more or less arranged tissues [14]. Moreover, the structure can be designed
17 to incorporate delivery systems, for controlled release of cytokines, growth factors and
18 drugs among others [81-83]. One important limitation of electrospinning is the harsh of
19 the fabrication process: cells cannot be encapsulated in situ [19], the resulting scaffolds
20 are weak, and the fiber diameters only emulate the thickest ranges found in the native
21 ECM (50-500 nm) [14].

1 **Molecular self-assembly**

2 Molecular self-assembly is based on the spontaneous arrangement of individual
3 building-blocks into ordered and stable architectures by means of non-covalent bonds
4 [14]. For example, one of the most broadly described nanofiber is formed by the
5 amphiphile peptide (Fig. III) [84]. These nanofibrillar matrices are very close in
6 architecture to those composed of collagen in the native ECM. They have 10 nm
7 oscillating fiber diameters, pores ranging 5 to 200 nm, and high water content , >99.5%
8 [64] (the ability to retain water is fundamental to mimic the features of a real ECM,
9 where water represents the highest percentage of the total weight). The amphiphile
10 peptides can form hydrogels at near-physiological conditions, and in many cases the
11 fiber morphology can also be controlled [85,86]. Furthermore, they can be designed to
12 be sensitive to (easily degraded by) the actions of proteases and include adhesion
13 moieties in their backbone structure to support cell migration or induce lineage
14 differentiation [87]. Scaffolds presenting the laminin epitope IKVAV can prompt neural
15 progenitor cells to differentiate into neurons [88]. Remarkably, some of these
16 nanofibilar constructs such as PuraMatrix™, are now commercial products intended to
17 be used in the fields of cell biology or tissue engineering [22]. Unfortunately, the nature
18 of the cross-linkages (noncovalent bonds such as hydrogen bonds, electrostatic
19 interactions, hydrophobic interactions, van der Waals interactions etc) does not offer
20 flexibility in tuning the scaffold's mechanical properties or, at least, they have not been
21 described yet [89].

22

1 **Box 3. Fig. III.** Schematic representation of peptide amphiphile and Cryo-TEM images
2 of resulting nanofibers. Reprinted with permission from [90]. Copyright 2009 American
3 Chemical Society

4

5

1 **FIGURE LEGENDS**

2 **Fig. 1.** Adhesion ligand patterning. (a) Commonly used strategy to achieve well-
3 characterized nanoscale patterns of RGD islands. Polymer chains modified with single
4 adhesion moieties gives rise to reduced island spacing (green dots). A blend of
5 unmodified polymer chains and polymer chains with multiple adhesion ligands results
6 in increased island spacing. (b) Schematized multiphoton chemical patterning in
7 hydrogels and resulting oblique and side views of fluorescence images taken from 3D
8 patterned squares and circle arrays (50 μm diameter). Agarose is covalently modified
9 with a derivative of cysteine protected with a photocleavable group. The protection
10 groups are removed in a patterned way by exposure with a laser beam. Thus, after
11 rinsing, free thiol groups are left in the irradiated areas. Finally, the hydrogel is rinsed in
12 a solution with desired oligopeptides (green fluorophore in this case), which are
13 covalently immobilized via Michael-type addition (reacting with free thiol groups). As
14 shown in fluorescence images, different biomolecules (represented with red and green
15 fluorophores) can be patterned in desired spaces. Scale bars represent 50 μm in both
16 fluorescence micrographs. Reprinted and adapted with permission from ref [35].
17 Copyright 2008 American Chemical Society.

18

19

20 **Fig. 2.** Spatio-temporal control of growth factor delivery. (a) Scheme illustrating a dual
21 delivery of growth factors as a way to regulate the kinetics. Delivery systems are
22 represented with capital letters indicating the growth factor encapsulated within their
23 matrices. VEGF (in red) and PDGF (in blue) contents are gradually released with
24 different kinetics. The size of the spots represents the amount of delivered growth factor

1 at each time point. Thus, there is a prompt VEGF burst, while PDGF burst is delayed in
2 time. (b) Cross-sectional schematic depiction of cellular microfluidic scaffolds showing
3 different manners to induce gradients of soluble factors. Encapsulated cells are shown
4 as double circles. Microchannels are depicted as squares. The pink shading represents
5 the gradients of soluble molecules at steady-state: On the top part, solutes are delivered
6 from the microchannels and are subsequently consumed by encapsulated cells as they
7 diffuse into the matrix. On the bottom part, solutes are delivered via the channels on the
8 left (Sources) and removed by the channels on the right (Sinks). λ_k (cm)= concentration
9 variation over particular distance (Krogh length); λ_c (cm)= interchannel distance; w_c
10 (cm)= microchannel width; h_c (cm)= microchannel height; K_c (cm s⁻¹)= mass transfer
11 coefficient of the flow in the microchannels; U_c (cm s⁻¹)= flow speed in the
12 microchannels. Reprinted and adapted by permission from Macmillan Publishers Ltd:
13 Nature Materials ref [51], copyright 2007. <http://www.nature.com>.

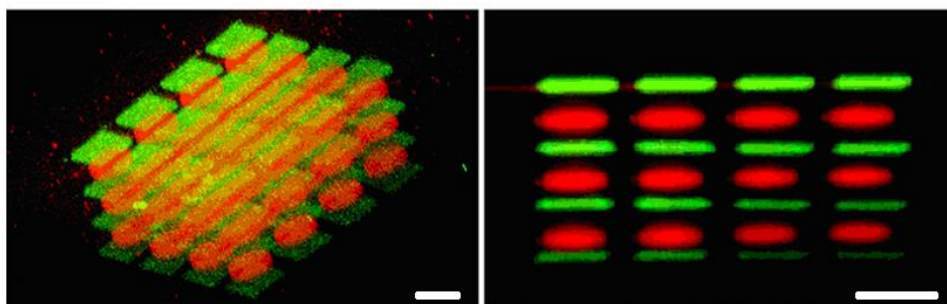
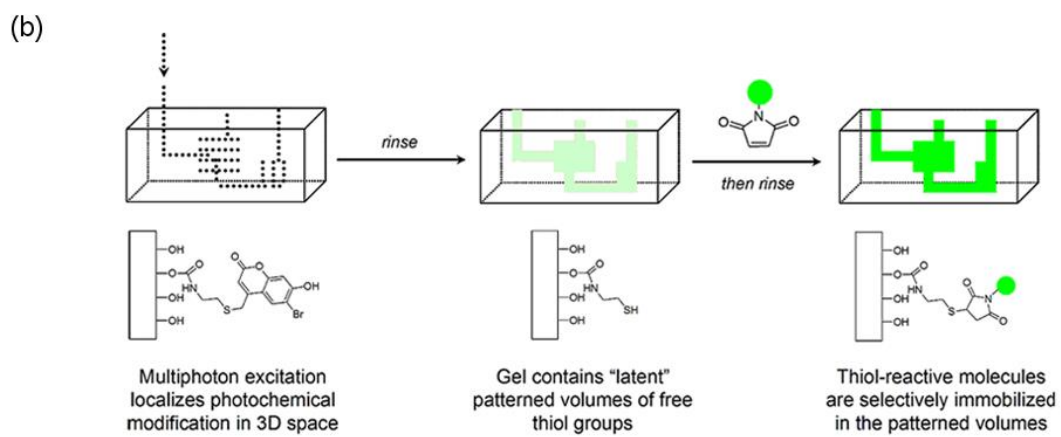
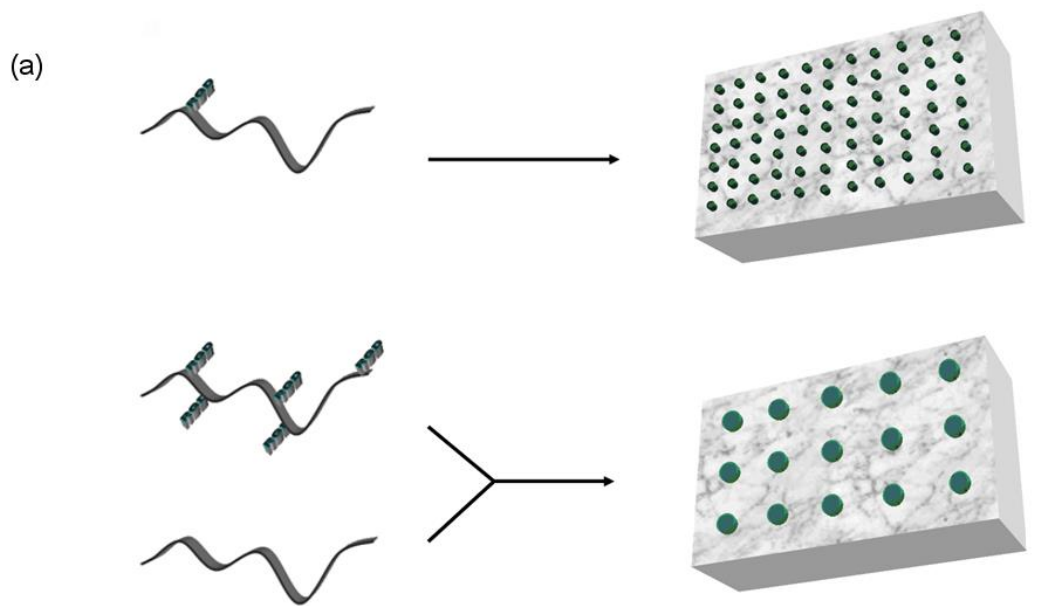
14
15 **Fig. 3.** Creating spatio-temporal signals by in situ modification of matrix properties. (a)
16 Cells within homogeneous hydrogels give rise to disorganized cellular structures with
17 no functionality. (b) Light-mediated in situ patterned hydrogels may possibly prompt
18 well defined structures and, ideally, tissue-like cell function. Photodegradable
19 hydrogels may be obtained introducing polymer chains that act as photolabile
20 crosslinkers. Upon irradiation, the photolabile groups are cleaved and hydrogel's
21 crosslinking density is decreased. Thus, large-scale stiffness gradients may be created,
22 for example, by flood irradiation with a highly absorbed wavelength, which gradually
23 penetrates into the hydrogel creating continuous and linear degradation gradients in the
24 z axis of the hydrogel. In addition, local patterns of degraded areas or three-dimensional
25 structures within the hydrogel may be formed using focused light directed by a single

1 photon confocal laser scanner microscope. (c) Thiol-ene reaction employed to create in
2 situ patterning of hydrogels. (d) The thiol-ene reaction mediated incorporation of
3 biochemical cues is confined to user-defined spaces of the hydrogel by
4 photolithographic patterning technique. This process can be continuously repeated to
5 add new biochemical cues to the scaffold, and this is represented in the figure
6 introducing different fluorophores within the hydrogel. (e) Full three-dimensional
7 control can be obtained by using focused laser light guided gel patterning, with which
8 micrometric spatial resolution is achieved. (f) Orthogonality of photocoupling and
9 photodegradation reactions. Multiphoton visible light is first used to create user-defined
10 shapes by coupling a green fluorophore labeled peptide within the hydrogel (top part,
11 buffalo), and then desired regions are degraded with multiphoton ultraviolet light, thus
12 removing the fluorophore from the irradiated areas (bottom part, CU and horn). (g) The
13 techniques described above can be employed, for example, to direct cell outgrowth
14 through three-dimensional functionalized channels. The inset depicts the hydrogel
15 viewed from above. Scale bars represent 100 μm in all pictures. (a-b) Reprinted by
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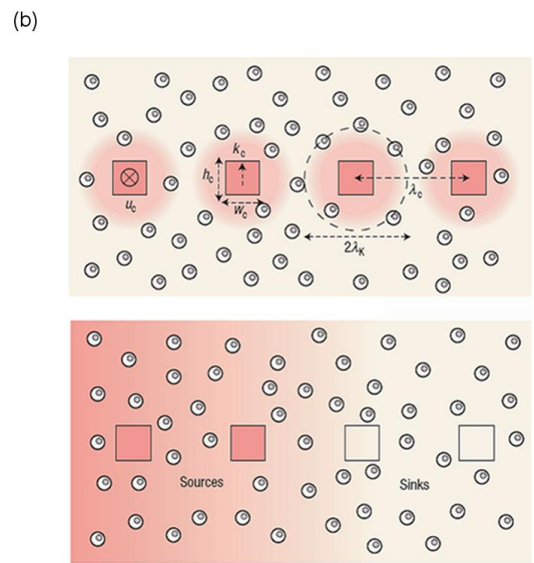
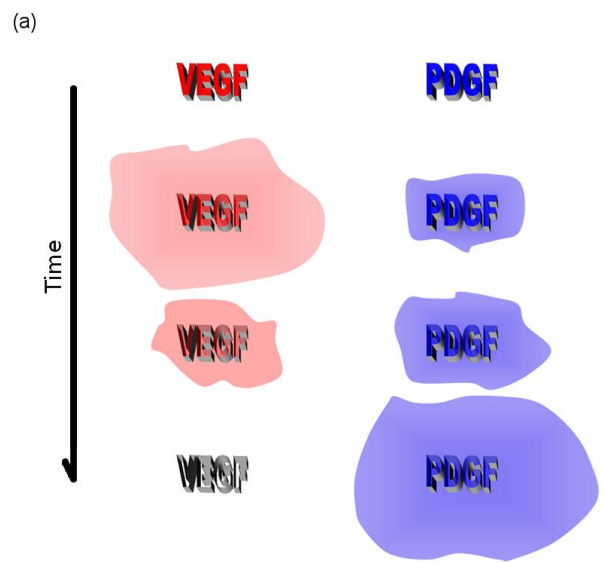


1

2 **Figure 1**

3

4



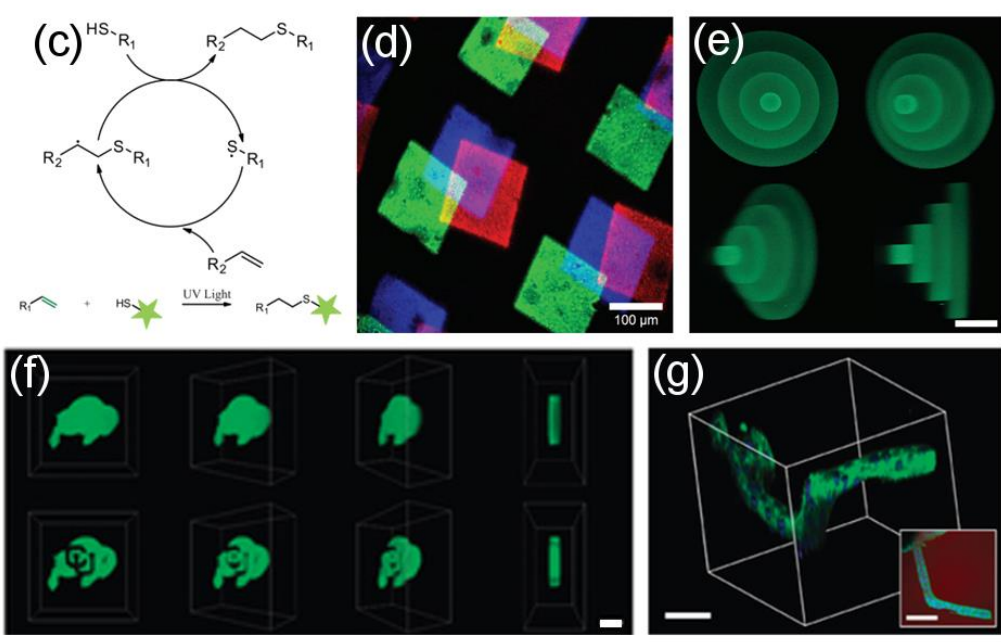
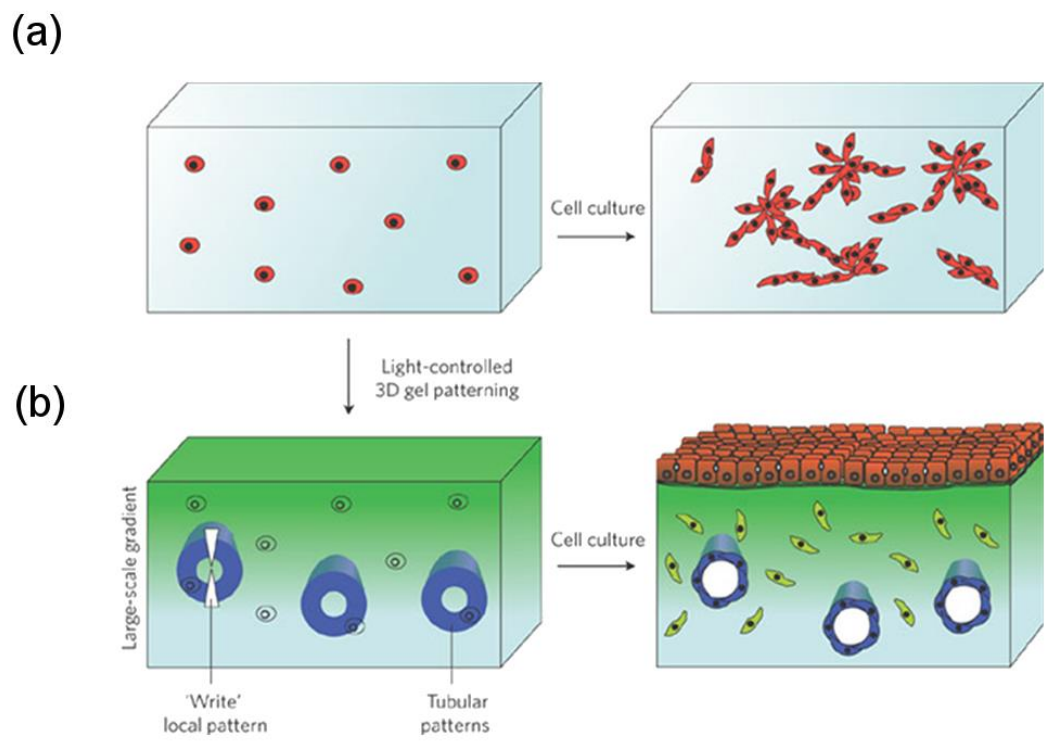
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2 **Figure 2**

3

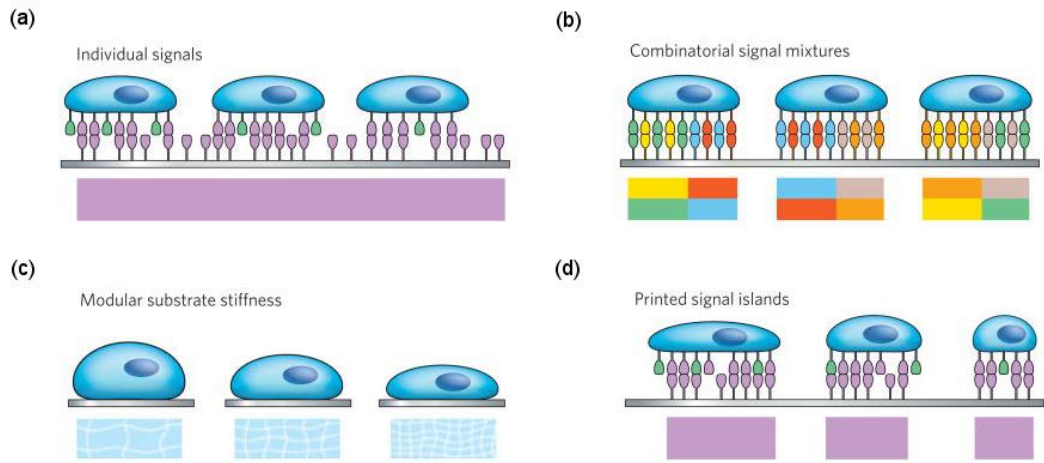
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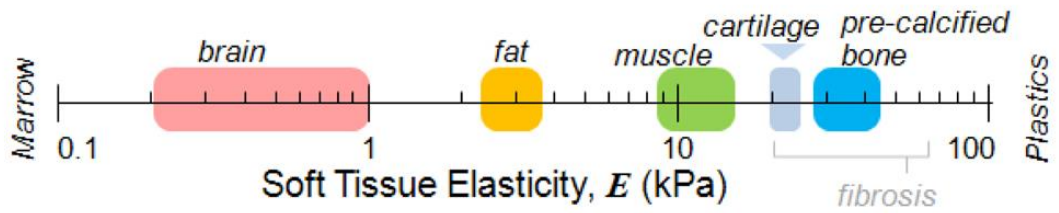
Figure 3



1

2 **Box 1. Figure 1**

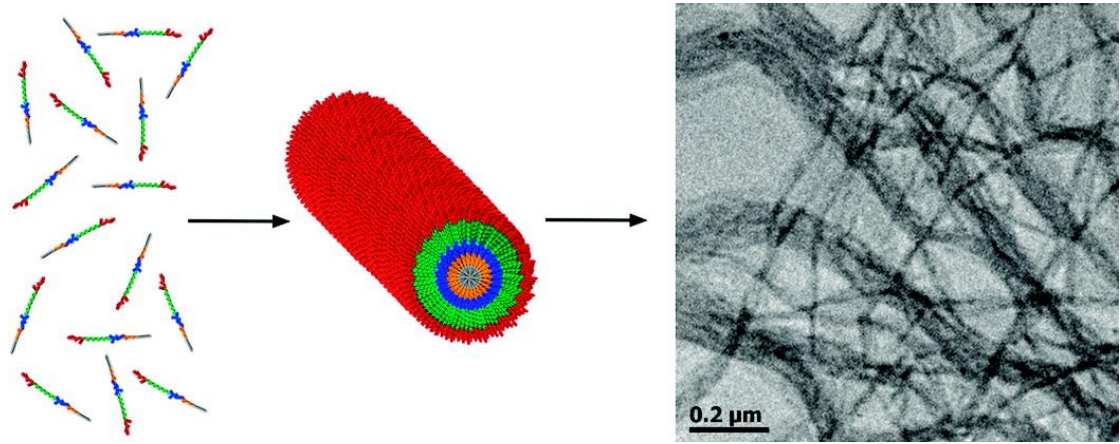
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4

5 **Box 2. Figure 1**

6



1

2 **Box 3. Figure 1**

3