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Bioink properties before, during and after 3D bioprinting

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TOPICAL REVIEW

Bioink properties before, during and after 3D bioprinting

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Abstract

Bioprinting is a process based on additive manufacturing from materials containing living cells. These materials, often referred to as bioink, are based on cytocompatible hydrogel precursor formulations, which gel in a manner compatible with different bioprinting approaches. The bioink properties before, during and after gelation are essential for its printability, comprising such features as achievable structural resolution, shape fidelity and cell survival. However, it is the final properties of the matured bioprinted tissue construct that are crucial for the end application. During tissue formation these properties are influenced by the amount of cells present in the construct, their proliferation, migration and interaction with the material. A calibrated computational framework is able to predict the tissue development and maturation and to optimize the bioprinting input parameters such as the starting material, the initial cell loading and the construct geometry. In this contribution relevant bioink properties are reviewed and discussed on the example of most popular bioprinting approaches. The effect of cells on hydrogel processing and vice versa is highlighted. Furthermore, numerical approaches were reviewed and implemented for depicting the cellular mechanics within the hydrogel as well as for prediction of mechanical properties to achieve the desired hydrogel construct considering cell density, distribution and material–cell interaction.

1. Introduction

Fabrication of scaffolds by employing additive manufacturing technologies (AMT) also referred to as three-dimensional (3D) printing, has been widely used in tissue engineering to restore, replace or regenerate defective tissues \cite{1, 2}. Bioprinting can be considered an additive manufacturing technique during which cells and biomaterials, often referred to as ‘bioink’, are deposited simultaneously \cite{3, 4}. Bioprinting allows to skip the cell seeding procedure, which often proved to be challenging for classical scaffold-based tissue engineering. Moreover, it provides a possibility to distribute different cell types at desired locations within the bioprinted construct and achieve high initial cell densities. A number of comprehensive reviews, book chapters and books, discussing the relevant technologies and materials have been published over the years \cite{3, 5–21}. In order, to achieve the desired tissue construct, it is essential to understand the properties of bioprinted hydrogel matrix and identify its key parameters (cell density, geometry, stiffness, etc) influencing tissue development and maturation.

In this review, recent progress in bioprinting and relevant bioink properties with focus on the interaction between hydrogel materials and cells is summarized. The properties of the hydrogel, which are required for the printing process as well as to ensure cell survival are discussed along with the influence of cells on the hydrogel properties itself.

We also describe a numerical approach allowing the estimation of the mechanical properties of a cell-containing hydrogel. The effect of cell densities and
distributions within the hydrogel was discussed, which might help to design the bioprinted constructs in a way to achieve the desired properties.

2. Overview of bioprinting methodologies and suitable bioink materials

The physico-chemical parameters of a hydrogel precursor including the rheological behavior, the swelling properties, the surface tension and the gelation kinetics are important factors for its printability. This especially applies for biofabrication techniques that rely on bioink dispensing. Therefore, different hydrogel properties are essential depending on the particular bioprinting technique to be applied. With regard to that, such processing methodologies can be subdivided into three groups including extrusion bioprinting (pneumatic and mechanical), orifice-free bioprinting (laser-induced forward transfer (LIFT) and printing by surface acoustic waves) and inkjet bioprinting (piezoelectric and thermal) (see figure 1).

2.1. Inkjet bioprinting
An inkjet bioprinter delivers small droplets of bioink (1–100 picoliters; 10–50 μm diameter) [16, 22] on predefined locations of a substrate. The two most commonly used methods for inkjet printing of cells are piezoelectric and thermal inkjet bioprinting [6, 23, 24]. The piezoelectric inkjet printer uses piezoelectric crystals to produce acoustic waves to force the liquid in small amounts through the nozzle [16, 25–28]. The thermal inkjet system produces pulses of pressure by vaporizing the bioink around the heating element expelling the droplets out from the printing head. Several studies have already indicated that cells are not affected by the local high temperature of the heating element up to 300 °C due to the short period of exposure (2 μs) during the printing process [29–32].

In inkjet bioprinting the surface tension is an important parameter that determines to what extent the processing technology will result in the formation of droplets or a jet. Surface tension is the result of the cohesive forces existing between the compounds present in the liquid. The charges on the surface of the bioink are weaker than the surface tension, droplets are formed. Conversely, a jet is produced. The surface tension decreases with increasing cell concentration in the bioink, because more cells are adsorbed to the liquid-gas interface. Therefore, the total free energy is reduced, resulting in a smaller surface tension [33].

Gelation methods including physical [27, 34], chemical and photo-crosslinking [35] are used to ensure the stability of bioprinted constructs. Gelation of the bioink should occur in situ after the material exits the nozzle and simultaneously with the printing process (e.g. by photopolymerization) [32], because when it already takes place inside the printing head, blockages are created in the nozzle [23]. When hydrogel formation does not occur rapidly in situ the bioprinted construct might be compromised due to possible spreading of non crosslinked bioink solution. Furthermore, the shear stress characteristic to this process can negatively influence the cell viability [36]. As a result, the bioink must exhibit low viscosities (<10 mPa s) and cell densities (<10⁶ cells ml⁻¹) (see figure 1 and table 1) [29, 37, 38]. These conditions result in limitations for the printing process. Despite the disadvantages, inkjet bioprinters are successfully applied with a micrometer resolution (10–50 μm) [23, 28, 39] for the deposition of cells and are compatible with various bioinks [23, 29, 40].

2.2. Orifice-free bioprinting
LIFT is also known as laser-assisted bioprinting and biological laser printing. In LIFT a pulsed laser beam is focused and scanned over a donor substrate that is
coated with an absorbing layer (e.g. gold or titanium) and a layer of bioink [41]. The focal point of the laser causes local evaporation of the absorbing layer thereby creating a high-pressure bubble that propels small portions of bioink towards a collector platform. The bioink jet extends towards the collector before separating from the donor substrate, by this way creating a temporary connection between both substrates [42–45]. This bioprinting technique is nozzle-free and is therefore not affected by clogging problems. Another substantial advantage is that the shear stress caused by the material passing through a nozzle (inkjet) or a needle (extrusion) is avoided. The resolution of LIFT is in the range of 10–100 μm [41, 45]. It is influenced by various factors, such as the laser parameters, the air gap between the donor substrate and the collector platform, the thickness and viscosity of the bioink layer [44]. LIFT is suitable for bioinks with a viscosity ranging from 1 to 300 mPa s and medium cell densities of \( \sim 10^6 \text{cells ml}^{-1} \) (table 1) [40, 41, 46, 47]. Understandably, bioprinting of well-defined 3D structures from low viscosity bioinks might be quite challenging. For the fabrication of the predesigned 3D constructs at high spatial resolution the bioink must exhibit fast crosslinking. Among the suitable cross-linking mechanisms, ionic crosslinking of sodium alginate containing bioink is frequently used. Also the temperature dependent gelation of Matrigel or enzymatic driven polymerization of fibrinogen were demonstrated [41, 42, 46].

Another elegant orifice-free bioprinting technique is relying on surface acoustic waves [19, 48]. The latter are produced by an acoustic ejector, which uses a surface acoustic wave piezoelectric substrate (e.g. lithium niobate, quartz, etc) with interdigitated gold rings placed on top of the substrate. Due to the circular geometry of the waves, an acoustic focal plane is generated at the air-liquid interface in the microfluidic channel. As a result, the bioink droplets are ejected from the microfluidic channel. The diameter of the droplets is uniform and can be set between 3 and 200 μm by changing the wavelength of the acoustic ejector. The embedded cells are not exposed to nozzle geometry, heat or high pressure, which results in a high cell viability (>89.8%). Furthermore, bioinks with various surface tensions and viscosities can be ejected [19, 48].

### 2.3. Extrusion bioprinting

Perhaps the most widespread method for the fabrication of 3D cell-laden constructs is extrusion bioprinting [41, 49]. For extrusion bioprinting, the bioink is generally inserted in disposable plastic syringes and dispensed either pneumatically or mechanically (piston- or screw-driven) on the receiving substrate [15]. In contrast with a LIFT or an inkjet bioprinter, an extrusion bioprinter does not dispense small bioink droplets but rather larger hydrogel filaments (approximately 150–300 μm in diameter) [16, 50–54]. A piston-driven system may provide more direct control over the flow of the bioink, when compared to pneumatic-based systems, prone to delays associated with the compressed gas volume. Screw-based deposition provides more spatial control and is capable of dispensing bioinks exhibiting higher viscosities [12]. However, the larger pressure drops generated by this extrusion method can be harmful for the suspended cells due to possible disruption of the cell membranes which results in cell death [55]. Because of the possibility to adjust the air pressure, pneumatic deposition can be used for a broad range of bioink types and viscosities. Advantages of extrusion bioprinting include the ability to print viscous bioinks (\(30–6 \times 10^7\text{ mPa s}\)) with very high cell densities, and even cell spheroids, into 3D scaffolds (see figure 1 and table 1) [40, 56]. The drawbacks related to this approach are its inferior resolution (200–1000 μm), potential nozzle clogging and the decreased cell viability due to shear stress [12, 40, 57]. The cross-linking pathways for fixation include physical (shear thinning and thermally induced), chemical (e.g. Michael addition reactions, click chemistry, etc) and photo-induced crosslinking [16, 40].

Another common aspect is that bioink formulations having adequate mechanical properties for

<table>
<thead>
<tr>
<th>Table 1. Overview of crucial bioink parameters, which are characteristic for the discussed bioprinting approaches. Adapted from [40].</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Orifice-free bioprinting</strong></td>
</tr>
<tr>
<td>LIFT [41, 67, 68]</td>
</tr>
<tr>
<td>Acoustic [19, 48, 69]</td>
</tr>
<tr>
<td><strong>Inkjet bioprinting</strong></td>
</tr>
<tr>
<td>[23, 29, 30, 37, 38, 61]</td>
</tr>
<tr>
<td><strong>Extrusion bioprinting</strong></td>
</tr>
<tr>
<td>[7, 36, 53, 54, 57, 62–66]</td>
</tr>
<tr>
<td>Viscosity</td>
</tr>
<tr>
<td>Bioink</td>
</tr>
<tr>
<td>1–300 mPa s</td>
</tr>
<tr>
<td>1–18 mPa s</td>
</tr>
<tr>
<td>&lt;10 mPa s</td>
</tr>
<tr>
<td>30–6 ( \times 10^7 ) mPa s</td>
</tr>
<tr>
<td>Cell density</td>
</tr>
<tr>
<td>Medium (10^6 cells ml^{-1})</td>
</tr>
<tr>
<td>Low (&lt;16 ( \times 10^6 ) cells ml^{-1})</td>
</tr>
<tr>
<td>Low &lt;10^6 cells ml^{-1}</td>
</tr>
<tr>
<td>High, cell spheroids</td>
</tr>
<tr>
<td>Resolution</td>
</tr>
<tr>
<td>10–100 μm</td>
</tr>
<tr>
<td>3–200 μm</td>
</tr>
<tr>
<td>10–50 μm</td>
</tr>
<tr>
<td>High</td>
</tr>
<tr>
<td>Medium</td>
</tr>
<tr>
<td>Single cell control</td>
</tr>
<tr>
<td>Control</td>
</tr>
<tr>
<td>Fabrication speed</td>
</tr>
<tr>
<td>Medium</td>
</tr>
<tr>
<td>Fast</td>
</tr>
<tr>
<td>Fast</td>
</tr>
<tr>
<td>Slow</td>
</tr>
<tr>
<td>(200–1600 mm s^{-1})</td>
</tr>
<tr>
<td>1–10 000 droplets s^{-1}</td>
</tr>
<tr>
<td>(100 000 droplets s^{-1})</td>
</tr>
<tr>
<td>(700 mm s^{-1}–10 μm s^{-1})</td>
</tr>
<tr>
<td>Cell viability</td>
</tr>
<tr>
<td>&gt;95%</td>
</tr>
<tr>
<td>89.8%</td>
</tr>
<tr>
<td>&gt;85%</td>
</tr>
<tr>
<td>80%–90%</td>
</tr>
</tbody>
</table>
fabrication of stable 3D constructs at good bioprinting accuracy often present suboptimal environment for cell migration and spreading [12]. A new bioprinting approach, which might overcome some of these drawbacks was reported recently by [58, 59]. In this gel-in-gel bioprinting method bioink is extruded into a volume of self-healing hydrogel acting as a support material. The support hydrogel deforms upon the injection of bioink and heals immediately after deposition enclosing the printed structure inside (see figure 2a). By using photocrosslinking as secondary stabilization step the mechanical properties of the printed construct can further be improved. Moreover, in combination with photocrosslinking, where either the bioink or the support hydrogel is photosensitive, freestanding 3D structures or structures with voids can be generated, by washing away the unstabilized hydrogel. This gel-in-gel printing method also opens up the possibility to print multiple materials and as the hydrogels are shear thinning also printing of cells results in a high viability (>90%).

2.4. Methods for hydrogel gelation
Hydrogel fixation (i.e. ‘gelation’) is an important aspect in preserving the shape of a bioprinted constructs thereby minimizing structure collapse [12]. The different gelation mechanisms can be subdivided into two categories being physical and chemical crosslinking. The network formation of a physical hydrogel is reversible and is the result of the occurrence of ionic interactions, high molecular chain entanglements, hydrogen bonds and/or hydrophobic interactions [70, 71]. Physically crosslinked hydrogels are usually associated with poor mechanical stability [55]. To overcome this limitation, chemical functionalities can be introduced to improve the mechanical strength of the hydrogel by creating covalent crosslinks, thereby resulting in an irreversibly crosslinked network [72, 73]. An irreversible network can be achieved by Michael-type addition reactions [74], click chemistry [75], enzymatic reactions [76] and photo-induced polymerization [77]. Li et al recently reported the development of a two-component bioink based on a supramolecular polypeptide-DNA hydrogel
A combination of both physical as well as chemical crosslinking can also be pursued [72, 79]. Physical crosslinking is generally used for biofabrication processes, since chemical crosslinking is often associated with stringent control over the crosslinking kinetics to avoid blocking of the nozzle [12]. Therefore, chemical crosslinking is frequently used as post-processing fixation and stabilization of the printed 3D constructs [55]. For example, Billiet et al. already reported on the application of methacrylamide-modified gelatin which was exposed to post-processing photo-induced crosslinking to produce mechanically stable 3D-constructs [63]. Sometimes a two-step photopolymerization approach is used to create a viscous, yet printable bioink and then the printed construct is fully photopolymerized to obtain the final shape of crosslinked scaffold. Skardal et al. used this two-step photopolymerization method to create 3D scaffolds based on a methacrylated ethanola- mide derivative of gelatin and methacrylated hyaluronic acid for tissue engineering applications. First, the gelatin and hyaluronic acid derivatives were partially photocrosslinked to obtain a gel-like bioink. Then the desired constructs were printed and photocrosslinked completely to fix their shape [80].

Colosi et al. has recently reported the use of low-viscosity bioink blend of alginate and gelatin metha- croyl (GelMA) with a coaxial dispensing system [81]. GelMA at low concentrations (<5% w/v) exhibits favourable properties for cells, but is not printable. Combining it with alginate results in a bioink mechanically stabilized by physically cross-linked fibers. Coaxial needle system (figure 2c) allows to precisely tune the gelation kinetics of this bioink by adjusting concentrations of alginate and CaCl$_2$. After bioprinting the hydrogel construct is further reinforced by UV cross-linking of GelMA.

### 2.5. Commercialization of bioprinting

The recent progress along with the increased attention to the field of bioprinting lead to intensified commercialization of devices and materials. Table 2 provides an overview of some commercially available bioprinters with their specifications and typical printing materials. Currently there is no standard way for defining the resolution of the printing process. Hence, manufacturers provide different parameters to describe it. For future evaluation, standardized parameters have to be defined in order to reasonably compare different printing methods.

While most of these printing devices rely on a single biofabrication method, RegenHU offers the selection between different fabrication technologies or combinations thereof. Very often bioprinters are also equipped with an additional light source enabling photo-induced polymerization, also referred to as curing, of the specialized bioinks.

As bioprinting has been commercialized, companies also start to offer their own bioinks. Among these, materials based on various cross-linking technologies are provided, including water-based, hydrogels, alginate, polyethylene glycol and water-based, alginate, polyethylene glycol (PLA).

### Table 2. Overview of commercially available bioprinters.

<table>
<thead>
<tr>
<th>Bioprinter and manufacturer</th>
<th>Fabrication technique</th>
<th>Specified resolution</th>
<th>Recommended materials</th>
</tr>
</thead>
<tbody>
<tr>
<td>3Dn300TE, NScrypt</td>
<td>Extrusion-based</td>
<td>Line widths 20–100 μm</td>
<td>Not specified (viscosity range: 0.001–1000 Pa s)</td>
</tr>
<tr>
<td>3D-Bioplotter$^a$, Envisiontec$^b$</td>
<td>Extrusion-based</td>
<td>Minimum strand diameter 100 μm</td>
<td>Hydrgels, ceramic, metal pastes, thermoplasts</td>
</tr>
<tr>
<td>Bioscaffold$^c$, Gesim$^c$</td>
<td>Extrusion-based</td>
<td>Not specified</td>
<td>Hydrgels, biopolymers (collagen, alginate) bone, cement paste, biocompatible silicons and melting polymers (CPL, PLA)</td>
</tr>
<tr>
<td>Biobot 1, Biobots$^d$</td>
<td>Extrusion-based</td>
<td>Layer resolution 100 μm</td>
<td>Hydrgels, biopolymers (viscosity range: 100–10$^4$ Pa s, see table 3 for more details)</td>
</tr>
<tr>
<td>Inkredible+, Cellink$^e$</td>
<td>Extrusion-based</td>
<td>Layer resolution 50–100 μm</td>
<td>Hydrgels (see table 3)</td>
</tr>
<tr>
<td>Biofactory$^f$, RegenHU$^f$</td>
<td>Extrusion-based Inkjet</td>
<td>Not specified</td>
<td>Bioink, Osteoink (see table 3 for more details)</td>
</tr>
<tr>
<td>Revolution, Ourobotics</td>
<td>Extrusion-based</td>
<td>Not specified</td>
<td>Collagen, gelatin, alginates, chitosan</td>
</tr>
<tr>
<td>Bio3D Explorers, Bio3D technologies$^{g}$</td>
<td>Extrusion-based</td>
<td>Not specified</td>
<td>Not specified</td>
</tr>
<tr>
<td>CellJet Cell Printer, Digilab</td>
<td>Extrusion-based</td>
<td>Droplet size 20 nl–4μl</td>
<td>Water-based, hydrogels, alginate, polylehne glycol</td>
</tr>
<tr>
<td>BioAssemblyBot, advanced solutions</td>
<td>Extrusion-based</td>
<td>Related to spheroid diameter</td>
<td>Not specified</td>
</tr>
<tr>
<td>Regenova, Cyfuse</td>
<td>Spheroid assembly</td>
<td>Related to spheroid diameter</td>
<td>Cells only (scaffold/biomaterial-free approach)</td>
</tr>
<tr>
<td>NovoGen MMX, Organovo$^{h}$</td>
<td>Injet</td>
<td>20 μm</td>
<td>Cellular hydrogels</td>
</tr>
<tr>
<td>Dimatix Materials Printer, Fujifilm</td>
<td>Injet</td>
<td>20 μm</td>
<td>Water-based, solvent, acidic or basic fluids</td>
</tr>
<tr>
<td>Poietis$^{i}$</td>
<td>LIFT</td>
<td>20 μm</td>
<td>Not specified</td>
</tr>
</tbody>
</table>

$^a$ Light curing system.  
$^b$ Not for sale, but utilized for bioprinting human tissue.
mechanisms and specified for different applications can be found (table 3). For example Bioink Solutions, Inc. offers gelatin-based bioinks containing growth factors that are specific for the printing of different tissue types.

Rheological properties of these bioinks are mostly not indicated by the manufacturer. Therefore, they might only be suitable in combination with the company’s own bioprinter and would have to be adapted for other printing techniques.

Generally, these bioinks are constituted from natural and/or synthetic polymers including collagen/gelatin, hyaluronic acid, PEG, etc. Collagen is the most abundant protein present in the extracellular matrix (ECM) of many tissues [82]. This protein forms a hydrogel at physiological conditions by triple helix formation. Collagen is a suitable material for cell encapsulation purposes because of the presence of cell-interactive RGD (Arginine-Glycine-Aspartic acid) sequences in their backbone, which stimulate cell adhesion. For example, Xu et al. have mixed rat embryonic hippocampal neurons with neutralized collagen and placed the cell-laden solution subsequently in the incubator at 37 °C to induce hydrogel formation [83]. The degradation of the triple helix of collagen by acidic or basic hydrolysis results in the production of gelatin [71]. Gelatin is a thermo-responsive protein with a sol-gel temperature of around 30 °C depending on the gelatin concentration applied. By cooling a gelatin solution below 30 °C, hydrogel formation is induced (cfr. upper critical solution temperature behavior) [84]. This protein is often employed for tissue engineering and regenerative medicine because various functional groups corresponding with constituting amino acids can be easily modified with (meth)acrylate groups to prevent

<table>
<thead>
<tr>
<th>Company</th>
<th>Bioink</th>
<th>Material</th>
<th>Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bioink Solutions, Inc.</td>
<td>Gel4Cell®</td>
<td>Gelatin-based</td>
<td>UV-crosslinkable</td>
</tr>
<tr>
<td></td>
<td>Gel4Cell®-BMP</td>
<td>Conjugated with different growth factors</td>
<td>Cell viability &gt; 90%</td>
</tr>
<tr>
<td></td>
<td>Gel4Cell®-VEGF</td>
<td></td>
<td>Osteoinductive</td>
</tr>
<tr>
<td></td>
<td>Gel4Cell®-TGF</td>
<td></td>
<td>Angiogenic</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Chondrogenic</td>
</tr>
<tr>
<td>CELLINK</td>
<td>CELLINK</td>
<td>Nano-cellulose/alginate mixture</td>
<td>Shear thinning</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Fast crosslinking</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>For soft tissue engineering</td>
</tr>
<tr>
<td>RegenHU</td>
<td>BioInk®</td>
<td>PEG/gelatin/hyaluronic acid-based</td>
<td>Good cell adhesion properties</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Biodegradable</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Mimics the natural ECM</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Possible combination with Osteoink®</td>
</tr>
<tr>
<td></td>
<td>Osteoink®</td>
<td>Calcium phosphate paste</td>
<td>Osteoconductive</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Chemical composition similar to human bone</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>For hard tissue engineering</td>
</tr>
<tr>
<td>Biobot</td>
<td>Bio127</td>
<td>Pluronic F127-based</td>
<td>Gels at room temperature</td>
</tr>
<tr>
<td></td>
<td>BioGel</td>
<td>Gelatin Methacrylate based</td>
<td>Dissolves when cooled</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>When combined with GelKey®</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Covalently crosslinks when exposed to light</td>
</tr>
</tbody>
</table>

Figure 3. (A) Structure of gelatin modified with methacrylamide groups (Gel-MOD). (B) Hepatocarcinoma cell were encapsulated in Gel-MOD constructs and cured using Irgacure 2959. Cell survival was evaluated with live/dead assay [63].
liquefying of gelatin at physiological temperature [60, 63, 77]. In addition, gelatin is bio-interactive due to the presence of RGD sequences (figure 3).

Not only proteins are present in the ECM, but also glycosaminoglycans including hyaluronic acid, which is a biodegradable, biocompatible and non-immunogenic biopolymer. The modification of hydroxyl and carboxylic acid functional groups of hyaluronic acid enable the introduction of photocrosslinkable moieties which can be photopolymerized in the presence of cells [85, 86]. Masters et al for example investigated the effect of photocrosslinkable methacrylated hyaluronic acid hydrogels on the cell response of encapsulated cardiac valvular interstitial cells (figure 4). Results showed that after 1 week, the embedded cells were still viable indicating the potential of these materials for tissue engineering purposes [87].

Also synthetic polymers such as PEG-based hydrogel precursors have already been frequently used for cell encapsulation because their mechanical, swelling and diffusion properties can be easily controlled by varying the crosslinking degree. On the other hand, these hydrogels lack inherent binding sequences, like the common RGD motif, for cell attachment and are not degradable. To make them more suitable for cells, these hydrogels have to be modified with the necessary binding peptides and enzymatically degradable groups. For example, Bryant et al developed PEG diacrylate hydrogels and incorporated RGD sequences in the backbone to enhance the cell-interactive properties of the hydrogel [88].

2.6. Summary
Each bioprinting technique has its own set of ‘ideal requirements’ with respect to the bioink properties, in order to achieve designed 3D geometries with the desired resolution and a high viability of the embedded cells (see table 1). In addition, process peculiarities might necessitate more specific material characteristics. A plethora of rheological parameters (i.e. viscosity, shear thinning and thixotropy) can be distinguished, which inherently affect the bioprinting process [12]. Viscosity is determined by temperature, the polymer concentration and its molecular weight. A bioink exhibiting a high viscosity is associated with an increased shear stress during the printing process, which can cause cell damage [89]. A previous study has already shown that the viscosity of a cell-laden hydrogel influences shape fidelity after deposition. Low-viscosity bioink forms strands which spread out on the receiving platform, while higher viscosity bioinks leads to the formation of filaments on the collecting substrate [90]. The viscosity of shear thinning materials decreases with increasing shear rate. This property results in high printing fidelity, because the applied pressure in the nozzle causes a decrease in viscosity thereby facilitating the deposition of the bioink. As the shear stress is removed after exiting the orifice, the viscosity increases sharply [18, 64]. This effect also takes place for thixotropic biomaterials, for which the decrease in viscosity is reversible and time-dependent [91]. Therefore, these phenomena are extremely useful for nozzle-based applications.

For example, inkjet bioprinting exhibits limitations regarding the material viscosity, while extrusion-based techniques may require shear-thinning properties to reduce the shear stress on the embedded cells to increase the cell viability [92]. The low viscosity bioinks for inkjet bioprinting require fast crosslinking mechanisms to facilitate the layering of the 3D-printed constructs. Conversely, crosslinking in extrusion bioprinting can be executed after fabrication, because the high viscosity bioinks maintain their 3D shape after deposition [12, 40]. However, the resolution of extrusion bioprinting is directly related to the diameter of the needle, which might result in some restrictions on material viscosity and affect the shear-stress induced during the dispensing process. Another interesting option is gel-in-gel bioprinting, allowing to extrude soft cell-friendly bioinks into the support gel. It remains to be seen how the supporting gel volume displaced by deposited bioink would affect the bioprinted construct in case when a large bioink quantity is used. By using a combination where one of the two

Figure 4. (A) Structure of methacrylated hyaluronic acid. (B) Live/dead staining results indicated that encapsulated cardiac valvular interstitial cells in methacrylated hyaluronic acid hydrogels remained viable after 1 week of cell culture [87].
hydrogels is photopolymerizable, gel-in-gel printing allows to create constructs with localized photosensitivity. As a result, a photopolymerizable part of the construct can be crosslinked, while the rest of the material is washed away to reveal the desired structure. A similar outcome is achieved by lithography-based 3D printing technologies. In this case it is not necessary to combine different properties, instead the same hydrogel is crosslinked selectively by controlling the material-light interaction volume. Somewhat higher spatial resolution and true 3D structuring, without the necessity to deposit material layer-by-layer, is possible with multi-photon processing.

For example, gelatin-based bioinks, already in their physical gel state, can be locally cross-linked by two-photon polymerization (figure 2c). In the case of LIFT and inkjet bioprinting techniques bioinks also encounter localized heating, which can further affect the viability of cells. Therefore, bioinks with a low thermal conductivity may be applied, to facilitate cell viability and superior cell function after the printing process.

3. Hydrogel properties before and after bioprinting

The characteristics of the bioink should meet the mechanical requirements for the bioprinting process and at the same time ensure cell survival within the produced construct. Therefore, cytocompatibility of a bioink is another critical aspect concomitant with bioprinting. Hydrogels are commonly used for tissue engineering and biofabrication because of their high water content and low toxicity rendering them excellent mimics of the ECM. Several studies have already demonstrated that 3D hydrogels produced by a bioprinting technology, provide an excellent matrix for encapsulated cells. Malda et al gives a good overview of the cytocompatibility of different hydrogels when fabricated with different methods. For example, 3D printed cell-encapsulating methacrylamide-modified gelatin hydrogels with a substitution degree of 62% resulted in a cell survival of >97% and maintained cell expression of the liver-specific functions. Thus, the cell viability was not impaired due to the printing process (e.g. needle type, temperature, etc) and the exposure to increased fluid shear stresses.

In addition, Hsieh et al have shown that neural stem cells embedded in 20%–30% polyurethane hydrogels exhibit excellent proliferation and differentiation due to the low matrix stiffness. The developed hydrogel was anticipated to mimic the microenvironment of the brain, resulting in an excellent niche for neural stem cells. Markstedt et al studied the use of a bioink that combined the outstanding shear thinning properties of nanofibrillated cellulose with the fast crosslinking ability of alginate. The printed constructs were stable in their shape and size and the embedded human chondrocytes exhibited a cell viability of 86% 7 d after printing. They stated that the bioink was suitable for 3D printing in the presence of living cells for inducing the growth of cartilage tissue. Furthermore, Das et al assessed the differentiation potential of human nasal inferior turbinate tissue-derived mesenchymal progenitor cells embedded in silk fibroin-gelatin. In the latter, bioink gelation was induced via enzymatic crosslinking by mushroom tyrosinase and physical crosslinking via sonication (see figure 5). The results showed that the constructs...
supported multilineage differentiation of the encapsulated stem cells and specific tissue formation [101].

In clinical applications direct injection of cells within a carrier solution often leads to a low cell viability due to mechanical disruption of cell membrane. Mechanical properties of a hydrogel carrier can be designed in a way that protects cells during injection. Aguado et al shows the protective effect of crosslinked alginate hydrogel with different storage modulus on encapsulated cells. The cell viability after injection was significantly higher in all the crosslinked hydrogel carriers with \( G' \) ranging from 0.33 to 58.1 Pa compared to control samples where a cell suspension in PBS was injected. Cells encapsulated in the hydrogel with a storage modulus of 29.6 Pa showed the highest viability demonstrating the impact of cell carrier mechanics on the cell viability [89]. Also Yan et al investigated the flow profile of a \( \beta \)-hairpin peptide-based hydrogel and could show their big potential as cell carriers for injection due to their shear-thinning and self-healing properties [102]. Moreover, Burdick group designed a Dock-and-Lock mechanism to obtain a self-assembling, self-healing and shear-thinning hydrogel for needle injection. Encapsulated mesenchymal stem cells showed a viability of >90% and remained homogenously distributed in the gel after needle injection [103].

Another important physico-chemical parameter for the fidelity of bioprinted constructs is the swelling behavior of hydrogels, which is mainly determined by the crosslinking extent and the charge densities [104]. This characteristic influences the final shape and the size of the printed 3D construct [90].

Moreover, the use of high levels of crosslinking result in lower swelling ratios thereby reducing the diffusion of oxygen and nutrients required for the cells to survive as a result of the reduced pore sizes [105]. Diffusion of waste products away from the cells encapsulated in the hydrogel is also managed in this context [97]. In addition, highly cross-linked hydrogels,
providing good shape fidelity after bioprinting, might not always be optimal from biological perspective due to impairment of cell migration and proliferation [12]. The requirements towards the properties of a hydrogel with regard to bioprinting process and cell culture are often opposing. A possible solution to this issue is combining hydrogels optimized for cell culture with materials providing mechanical stability and facilitating shape fidelity [106].

In a method referred to as 4D-Printing, Lewis et al demonstrated that by controlling swelling anisotropies within a hydrogel construct it can deform in controllable manner. Swelling anisotropy was introduced during printing by spatial control over the orientation of the cellulose fibrils inside the hydrogel. Upon immersion in water the structure swells and acquires its final shape over time (see figure 7) [107]. In general every bioprinting process has a 4D printing character since, it is likely that the geometry and properties will change in long term, unless completely isotropic constructs are built. For example the change in the geometry induced by swelling of zonal hydrogel constructs with different mechanical properties in separate layers has to be taken into account if swelling would occur after bioprinting [108].

4. Effect of cell content on material processing

It is expected that high initial density of cells in the bioprinted construct will lead to faster tissue formation. However, the presence of cells significantly affects the printability of bioinks. Indeed, Billiet et al have compared the processing potential of methacrylamide-modified gelatin (Gel-MOD) with and without hepatocarcinoma cells. It was demonstrated that the incorporation of the cells altered the rheological properties. As such, this parameter will affect the printing process since the viscosity is altered. For temperatures above the gelation point, the viscosity was reduced by a factor of 2, up to a cell density of $1.5 \times 10^6$ cells ml$^{-1}$. Increasing the cell density further to $2.5 \times 10^6$ cells ml$^{-1}$ resulted in a further viscosity decrease up to a factor of 4 (see figure 8) [63]. Furthermore, Skardal et al have assessed the effect of the presence of human intestinal epithelial cells on hyaluronan-based hydrogels which were crosslinked using tetrahedral polyethylene glycol tetracrylate. Their report states that cell densities above a certain threshold, interfere with hydrogel formation. Bioink with cell densities up to $25 \times 10^6$ cells ml$^{-1}$ formed
hydrogels within 20 min. At higher cell densities the hydrogels did not form under the applied conditions [109].

The equilibrium and dynamic modulus at 1 Hz of agarose hydrogels with chondrocyte density at 0, 10, 40 million cells ml\(^{-1}\) were presented in the study by Buckley et al [110]. By maintaining the effective concentration of agarose as constant, both the equilibrium modulus and dynamics modulus at 40 million cells ml\(^{-1}\) were lower than acellular constructs. This result could be explained by the Young’s modulus of a chondrocyte, which has been reported as approximately 0.6 kPa [111], an order of magnitude lower than that of agarose. In this case, a larger cell-seeding density would reduce the modulus of cell seeded agarose.

5. Predicting properties of hydrogels containing living cells

There was a certain amount of effort dedicated to predicting properties of the 3D printed scaffolds [112–115]. The mechanical properties of scaffolds were mainly considered with regard to construct compliance, but also in terms of cell-material interaction.

In case of bioinks one has to consider the presence of living cells not only with regard to bioprinting, but also in terms of possible long-term changes of bioprinted construct properties due to cell proliferation, migration and interaction with hydrogel material (cell traction, enzymatic degradation matrix remodeling). As shown in figures 9(A)–(C) there are different ways for cells to proliferate inside the hydrogel construct. For example MC3T3-E1 cells were found to grow in clusters when encapsulated in alginate hydrogel. In contrast, fibroblasts encapsulated in methacrylated gelatin were able to spread through the hydrogel over time, whereas they were found to grow in aggregates in pullulan methacrylate [116–118]. This behavior is not only attributed to the cell type, but also to the properties of the hydrogels, such as porosity, stiffness and most important the presence of ligands facilitating cell attachment.

The relationship between the density of encapsulated cells and mechanical properties of the resulting hydrogels was investigated by Mauck et al [119]. In their work, chondrocyte-seeded agarose hydrogels were cultured in free-swelling and dynamic loading conditions over a 2 month culture period. Constructs containing 10 million cells ml\(^{-1}\) were initially twice stiffer than the ones seeded with 60 million cells ml\(^{-1}\). After culturing 56 d, the constructs seeded with more cells showed similar Young’s modulus with the lower cell density one under the condition of free-swelling (85.1 ± 15 kPa versus 78 ± 1.5 kPa). This indicated that higher cell-seeding density accelerates the hydrogel remodeling. This observation was confirmed by Chang et al [120], who demonstrated that a higher cell density led to a larger equilibrium modulus of tissue implants after 30 weeks of culture. In addition, the dynamic loading condition resulted in a much larger Young’s modulus for the hydrogel containing higher cell density (186.2 ± 11.3 kPa versus 91.5 ± 11.6 kPa). The loading condition induced dynamic modulus followed the same trend. The interesting observations were that both proteoglycan and collagen density remained the same regardless of free-swelling or dynamics loading. This was speculated that the loading condition promoted the production of linker
molecules and/or the aggregation of macromolecular proteoglycan. This work indicated that various cell densities encapsulated in the 3D hydrogel could promote material remodeling depending on external loading conditions. The experimental data reported in literature is generally obtained following different protocols, which makes it difficult to guide hydrogel design and optimization. Numerical models have the potential to predict mechanical properties of a construct with different cell density under various protocols. It is especially appealing considering the multitude of bioinks used by different groups and the diversity of their properties. Guilak et al [121] developed a finite element model of chondrocytes within an explant of cartilage to understand the interactions between cell and matrix, which differ by nearly three orders of magnitude in terms of their Young's moduli. The Gel-MOD loaded with different densities of MC3T3-E1 cells, ranging from 15 to 6.14 × 10^6 cells ml^-1, was characterized compared to acellular hydrogels using a photorheometer. The details of experimental procedure are similar to the ones reported previously by our group [123]. In short, the cells were directly resuspended in a 10% (w/w) Gel-MOD solution. Measurement of the storage modulus G’ and the loss modulus G” were taken with a photorheometer (Anton Paar MCR 302 WESP) during a dynamic time sweep at a frequency of 10 Hz and strain of 10% during 10 min of photopolymerization with a 320–500 nm light source.

Our results presented here (previously unpublished) show that the stiffness of hydrogel drops by 13% when the cell density is increased from 12 to 15 million cells ml^-1 (figure 10). Calibrated by the aforementioned experimental data, numerical models were developed to predict mechanical properties of hydrogels containing different cell densities and the corresponding cellular mechanics.

A representative volume element (RVE) of the hydrogel with side length of 150 μm was used to represent hydrogels containing different cell densities (figure 11). The encapsulated cells were modeled as solid sphere with 30 μm in diameter and 1.5 kPa in Young's modulus, which was adopted from the previous experimental observation on MC3T3-E1 cells [124]. Based on the aforementioned experimental measurements, the shear storage modulus and loss modulus of the acellular hydrogel was 7870 Pa and 11 Pa, respectively. In the model, the material behaviors of hydrogel were then defined by the magnitude of shear modulus and the Poisson’s ratio of 0.49. A 10% shear loading was applied on the RVE to mimic the testing condition. Nonlinear finite element models were solved using ABAQUS 6.12 (Simulia, Providence, RI, USA). Different cell densities varying from 6.14 to 15 million ml^-1, corresponding to the volume fraction of 8.68% to 21.2%, were considered in the RVE. The estimated shear modulus of the RVE was found to be in good agreement with experimental measurements (figure 10). Both experiment and simulation demonstrated that higher cell density led to reduced modulus of the hydrogel. These findings are also consistent with experimental observations reported by other groups [63, 110, 119].

As cells are not always distributed homogeneously, but might be present in clusters inside the hydrogel,
the effect of different cell distributions on the hydrogel properties was also investigated. This was illustrated by comparison of three different cellular distributions (random distributed, corner clusters and central cluster) at the same cell density of 9.6 million ml$^{-1}$ as shown in figure 11. The obtained shear modulus from finite element models was summarized in table 4. It is clear that the samples with cell clusters are softer compared to the ones without randomly distributed cells.

**Figure 11.** Representative volume element (RVE) used for numerical model of hydrogel containing randomly distributed cells at density of 12 million ml$^{-1}$ (top). Numerical models of hydrogel containing living cells distributed (A) randomly, (B) in eight corner clusters and (C) in one central cluster at a density of 9.6 million ml$^{-1}$.

**Table 4.** Mechanical performance of hydrogel with different cell distribution.

<table>
<thead>
<tr>
<th>Shear modulus (Pa)</th>
<th>Random distributed</th>
<th>Edge clusters $(-7.3%)$</th>
<th>Central cluster $(-6.1%)$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>6628</td>
<td>6141</td>
<td>6221</td>
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**Figure 12.** Probability distribution of von-Mises stresses on cells.
The shear modulus for edge clusters and central cluster are 7.3% and 6.1% less than the Gel-MOD samples with random distributed cells. The minimal difference between two cluster distributions could be explained by the average distance between cells, which are close enough resulting in similar cell–cell interactions. The mechanical stresses sensed by cells embedded in Gel-MOD were depicted as probability curves in figure 12. It is obvious that cell clusters shifted probability distribution curve to a higher stress region. This implied that cells in cluster state are more prone to damage and therefore also more susceptible to mechanical stimulation. This behavior might be beneficial considering the matrix design for cartilage tissue engineering [125].

Moreover, multiple edge clusters shared a little larger loadings than the one central cluster, as indicated by the shapes of probability curves. However, in general the cluster distribution has minimal impact on the accumulated cellular loadings.

The formation of cell clusters inside the hydrogel might also result from proliferation of encapsulated cells (see figure 9). In order to estimate to what extent clusters might influence the temporal properties of the construct, the effect of cell proliferation was investigated by simulating four different situations related to cell division: initial distribution of single cells (8 cells per RVE), cell doubling (16 cells) etc (figure 13). Although this model might be not taking into account all the aspects of the cell–material interaction, it estimated that within the simulated range the shear modulus decreased from 7.425 to 6.141 kPa as a results of cell proliferation (see table 5). Since cell proliferation is imperative to most bioprinting methods it is important to be able to estimate the final mechanical properties of the construct based on the knowledge of initial material, cell density and proliferation rate. In addition, the appropriate numerical models should be capable of predicting the effect of material degradation, matrix remodeling etc [126].
The loadings sensed by cells are relevant to their behavior, including cellular viability, differentiation, and damage etc. We have delineated the maximum principal strain sensed by cells encapsulated in hydrogel as probability curves (figure 14a). It is clear that the cellular strain tends to be more inhomogeneous as the cell density increased. Higher strain regions were observed at the interface between the cell and hydrogel. This could be explained by the material mismatch between cell and hydrogel and the cell–cell interactions. Stress concentration was observed at the interface between cell and hydrogel, where there existed the material mismatch. Previous reports speculated, that this inhomogeneity is correlated with cellular damages as cells are more prone to be damaged at higher strain [122]. It is interesting to note though that mean stress and strain sensed by cells as a whole has no significant difference among various cell densities (figure 14b).

Another potentially important aspect of cell–material interaction, which might have an effect on the hydrogel construct, is cell contractraction. Our modeling results assuming the 5% cell contraction (implemented as eigenstrain, which was estimated from the traction force microscopy data) showed only minimal effect on the mechanical properties of the hydrogel. Specifically, for the case of hydrogel containing 9.6 million cells, the cell contraction altered the shear modulus of the hydrogel from 6.628 to 6.631 kPa.

6. Future perspectives

The dawn of bioprinting enabled the generation and transplantation of several tissues, including multilayered skin, bone, vascular grafts, tracheal splints, heart tissue and cartilaginous structures [40]. Bioprinting has come a long way with the portfolio of bioinks designed for different technologies and applications expanding rapidly. Hydrogel properties relevant to the process of bioprinting have been systematically investigated and adapted to match different technologies. The development of the bioprinted construct into a tissue is gathered by a multitude of factors such as cell proliferation, material degradation, matrix remodeling etc. Although modern computational approaches should be capable of predicting these processes and benefiting the field of bioprinting, their development is currently lacking attention. Availability of according modeling tools would be highly advantageous, especially taking into account the diversity of bioink properties, expenditures associated with experimental optimization of bioprinted tissue constructs and the possibility to apply these tools to different geometries, e.g. patient-specific cases, in the future. This work reviewed the recent efforts aiming at predicting the properties of cell containing materials and constructs. Furthermore, we present an own model allowing to estimate the mechanical properties of hydrogels containing different cell densities and distributions. This model provides a fundamental framework for designing bioprinted constructs considering the impact of cell density to achieve desired mechanical properties. The model could be extended to incorporate complex 3D construct architectures, which has demonstrated great influence on their mechanical environments [127]. The predicted mechanical response of cells in various printed hydrogel architectures, integrated with experimental data, could be used to determine the cellular loadings, its damage threshold, as well as the longitudinal behaviors. In addition, the bioprinting process-induced mechanical disturbances has also been found to affect the cell viability [36]. Numerical modeling could be also used to mimic the mechanics during the fabrication process. Optimized parameters such as printing speed or nozzle diameter might be obtained for certain mechanical properties of hydrogel containing cells. In perspective, such computational tools can be directly integrated with modeling of tissue and organ development [128].

Disclosure of interests

We declare no potential conflicts of interest relevant to this article.

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