

In vitro three-dimensional cell cultures for bone sarcomas

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In vitro three-dimensional cell cultures for bone sarcomas Javier Munoz-Garcia^{1,2,#,*}, Camille Jubelin^{1,2,3,#}, Aurélie Loussouarn¹, Matisse Goumard^{1,2}, Laurent Griscom⁴, Axelle Renodon-Cornière¹, Marie-Françoise Heymann^{1,2}, Dominique Heymann^{1,2,5,*} ¹ Université de Nantes, INSERM, Nantes, France ² Institut de Cancérologie de l'Ouest, Saint-Herblain, France ³ Atlantic Bone Screen, Saint-Herblain, France ⁴ BIOSIT CNRS UMS3480, Université de Rennes-1, Rennes, France ⁵ University of Sheffield, Department of Oncology and Metabolism, Medical School, Sheffield, UK # JMG and CJ contributed equally to the work Running title: 3D cultures of bone sarcomas **Corresponding Author:** Dr Javier Munoz-Garcia Email: javier.munoz@ico.unicancer.fr Prof. Dominique Heymann Université de Nantes Institut de Cancérologie de l'Ouest Blvd Jacques Monod, 44805 Saint-Herblain, France Email: dominique.heymann@univ-nantes.fr Tel: +33 (0) 240 679 841

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Abstract

Bone sarcomas are rare tumour entities that arise from the mesenchyme most of which are highly heterogeneous at the cellular, genetic and epigenetic levels. The three main types are osteosarcoma, Ewing sarcoma, and chondrosarcoma. These oncological entities are characterized by high morbidity and mortality and an absence of significant therapeutic improvement in the last four decades. In the field of oncology, in *vitro* cultures of cancer cells have been extensively used for drug screening unfortunately with limited success. Indeed, despite the massive knowledge acquired from conventional 2D culture methods, scientific community has been challenged by the loss of efficacy of drugs when moved to clinical trials. The recent explosion of new 3D culture methods is paving the way to more relevant *in vitro* models mimicking the *in vivo* tumour environment (e.g. bone structure) with biological responses close to the *in vivo* context. The present review gives a brief overview of the latest advances of the 3D culture methods used for studying primary bone sarcomas.

Key words

Osteosarcoma; Ewing sarcoma; Chondrosarcoma; Extracellular matrix; 3D culture; Multicellular tumour spheroid; Scaffold-based 3D culture; Microfluidics; Bioprinting.

1-. **Introduction**

Bone sarcomas correspond to < 0.2% of diagnosed cancers registered in the EUROCARE database and then they are considered as orphan tumours. The three main types of bone sarcomas are: osteosarcoma (OS), Ewing sarcoma (ES) and Chondrosarcoma (CS). Despite theirs low incidence, bone tumours are associated to high morbidity and mortality, with an important impact in children and young adult population (e.g., 80% of ES patients are under 20 years of age at diagnosis). The absence of specific symptoms at early stages of the disease leads to late diagnosis that frequently corresponds to aggressive phases that include cancer cells spreading and establishment of bone and lung metastases. Unfortunately, whether the 5-year survival rate is 50-70% for OS and ES, there is a drop around 20-30% when lung metastases are detected at the time of the initial diagnostic [1]. CS are associated with high local recurrence associated with high morbidity [2].

OS constitutes the main entity with an incidence of two-thirds of primary bone tumours and affects preferentially children and adolescents. In most of the cases, clinical treatment includes surgical procedure combined with adjuvant and neo-adjuvant polychemotherapies with or without radiotherapy. Unfortunately, distant recurrences (with a high predilection for the lung) frequently occur and are associated with drug resistance [2]. OS normally germinates from malignant mesenchymal cells of long bones committed in osteoblastic differentiation and are characterised by the production of an osteoid matrix by tumour cells [3]. The aetiology of the disease is explained by initial somatic mutations of p53, Rb and a BRACness signature that lead to chromosomal instabilities, complex genomic profile and high cellular heterogeneity [2,4]. Cancer stem like cells [5], tumour microenvironment (TME) including immune infiltrated cells and extracellular matrix (ECM) that modulates tumour cell adhesion and migration are also suspected to contribute to this high heterogeneity and to the acquisition of drug resistance [6,7]. Thus, due to their highly complex pathobiology and the limited access to patient samples, a better understanding of OS growth and drug development require the generation of new cell culture methods that mimic native TME of OS [8].

ES is characterized by its high aggressiveness, fatal malignancy developed in bone and extra skeletal sites and with a rapid metastatic expansion mainly in lung. ES is the second most

common paediatric bone tumour affecting 3 children per million [9]. ES principally affects Caucasian patients with a slight prevalence in men than women. ES cell classically presents a round morphology, with common expression of the CD99 (MIC2) antigen and chromosomal translocation of the *EWSR1* gene to *ETS* family genes [10]. Experimental evidence suggests that ES cells may originate from undifferentiated mesenchymal stem cells (MSCs) characterized by neuroendocrine features and acquisition of *EWSR1* translocation [11,12]. In addition to conventional chemotherapies, clinical developments are focused on downstream partners of the EWSR1/FLI1 signalling pathways [13,14]. An important feature of ES is the high resistance to chemotherapy agents, in part due to their particular MSCs origin but also to its complex TME. Reproduction of TME by 3D culture techniques results in a key progress to better understand the behaviour and drug resistance of ES.

CS compose a heterogeneous group of primary malignant tumours characterised by relative low growth ratio and the formation of hyaline cartilaginous neoplastic tissue. Depending on their malignancy, CS are classified in three grades: low-metastatic grade I, intermediate grade II, and high metastatic grade III [15]. CS are characterised by a high chemo- and radio-resistance mainly due to the presence of large amount of ECM and poor vascularity that restrict the diffusion of anticancer agents and slow down their effectiveness [16]. The importance of these features has motivated the scientific community to switch to 3D culture systems that can reproduce the native CS condition and have the potential to be a great tool for developing new therapies against CS.

Beside the genetic charge present in tumour initiation, TME has emerged as a key factor for tumour development and malignancy. For a better understanding of tumour biology, the scientific community has to reproduce, as close as possible, natural cell growth conditions [17]. During last decades, many technological progresses have been proposed to mimic native tumour biology. Whereas the first documented cell culture methods date from 1885 by Wilhelm Roux [18], the establishment of a true two-dimensional (2D) laboratory tissue culture system has been described by Ross Harrison at the beginning of the 20th century [19]. This event led to a scientific revolution in the understanding of cell behaviour during healthy and pathogenic situations. While 2D cell culture techniques became standards in research laboratories for a wide window of studies and fields, they do not reproduce the dynamic evolution of tumour growth and failed to

mimic cell-to-cell or cell-to-microenvironment interactions. To overcome these 2D culture issues, during last decades, a variety of 3D cell culture techniques has been developed including liquid-based scaffold free methods, scaffold 3D systems and the emerging organ-on-a-chip platforms: microfluidics and bioprinting systems [20-23].

Briefly, liquid-based scaffold-free methods are based on the prevention of cell adhesion to the cell culture container surfaces (e.g., vessels, plates) by coating them with non-adherent materials such as agar or poly-hydroxyethyl methacrylate [24]. The absence of adherent surfaces promotes cell-to-cell adhesion and formation of spontaneous spheroids. Wide variety of low/non-adherent supports are commercially available nowadays. Hanging drop technique is another liquid-based scaffold free methods that allows the production of spheroids using mono- or multi-cellular (co-cultures) approach thanks to the effect of the gravity (Figure 1) [25,26]. Low adherent supports and hanging drop methods have been also widely used to study cell organisation, embryonic development, tumour biology and tissue formation [25,27-32]. However, one of the main drawbacks of liquid-based scaffold methods is the lower reproducibility and control of the surrounding cell microenvironment.

Scaffold 3D systems consist in a structural support that favours cell adhesion, proliferation, migration, cell-to-cell interaction and signalling [33,34]. Natural scaffolds are based on molecules that are present in the ECM such as collagen, gelatine and derivatives [35,36], complex matrix (e.g., commercial MatrigelTM) and hydrogels [37,38], and polysaccharides as alginate, chitosan or hyaluronic acid [39-43]. Whereas the main advantage of these natural scaffolds is their biocompatibility, their production and inter-batches variability are the main issues of these materials. To solve these problems, synthetic scaffolds characterised by high stability, reproducibility and biocompatibility have been developed [44]. The most used synthetic scaffolds are based on polyethylene glycol (PEG) polymer hydrogels [45-48].

Finally, organ-on-a-chip platforms are based on microfluidic devices or 3D bio-printed systems [49-53]. Both techniques allow a precise control of the TME by applying a tuneable perfusion of media that mimics blood flow and facilitates a continuous access of nutrients, oxygen and drugs [54-56]. In addition, these systems can reproduce the complexity of tissue by adding layers or

compartments by co-culture of different various sets of cells in the presence of various ECM components [57-59].

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In the present review, we will focus on the different 3D culture techniques recently developed for the three main bone sarcoma entities.

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2-3D culture methods of primary bone tumours

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2-1. 3D Osteosarcoma culture models

With a low 5-years survival rate and no improvement in the last 4 decades, OS is a rare and devastating oncological entity that affects mainly children and young adults. The complexity of the bone structure and surrounding TME imply that 2D monolayer culture stays far away from the organisation of natural tumour tissue and impairs the study OS development [60].

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2-1.1. 3D culture and drug resistance in Osteosarcoma

196 Low attachment and hanging-drop cell culture are the most frequently employed liquid-based 197 scaffold-free method and consist in an easy 3D approach for generation of OS spheroids to study 198 cell behaviour and drug resistance (Figure 1 and Figure 2) (Table 1). By using hanging-drop 199 methodology, Rimann et al. demonstrated that spheroids produced from established cell lines, 200 like SaOS2 or HOS, exhibited a totally different pattern of resistance to a panel of antitumor 201 drugs compared to 2D (monolayer) culture methods [61]. Indeed, the IC50 values for 202 doxorubicin, cisplatin, taxol, and taurolidine significantly increase in 3D culture, meaning that 203 3D cells are more resistant to those drugs than monolayer culture. Those data were consistent 204 with the observation done in patients where treatment based in 2D dose concentrations showed a 205 decrease of drug effectiveness compared to 2D culture [61]. Many papers appeared reinforcing 206 the concept that 3D spheroids are really closed to real tumour behaviour to drug treatment [62-207 65]. Similarly, U2OS spheroids generated by ultra-low attachment methods were used to mimic 208 tumour structures and demonstrated the potential use of the nuclear NAD synthesis enzyme 209 nicotinamide mononucleotide adenylyltransferase-1 (NMNAT1) as a target for anti-tumour drugs 210 [66]. The expression of this enzyme increased in several tumour cell lines after exposure to DNA damaging agents as cisplatin and doxorubicin, suggesting an important role of this enzyme in tumour resistance [66].

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2-1.2. Impact of the tumour microenvironment structure in drug resistance

Cancer cells receive multiple signals (autocrine, juxtacrine and endocrine messages) coming from their cellular neighbours, the extracellular matrix and distant organs. By integrating all of the information, cancer cells change their behaviour, modify their metabolism and migration properties and become quiescent, or highly proliferative or resistant to drugs. TME then plays a crucial role in drug resistance [67]. Whereas hanging drop techniques have been used with success for rapidly forming tumour spheroids, this approach is far from wholly reproducing the natural TME controlling the cancer cell behaviour. 3D cell culture based on scaffold methods using natural components identified in the natural TME is one of the strategies proposed to mimic the environment of cancer cells. Interestingly, 3D scaffold are also bivalent tools not only usable in *in vitro* assays but also as support of cancer cells in *in vivo* animal experiments. The use of alginate beads is a perfect example of a 3D scaffold frequently applied to the oncology field. Alginate is used firstly to encapsulate and proliferate OS cells in a 3D spheroid configuration, and secondly for studying the metastatic effect of OS cells in animal model after inoculation of encapsulated cells [68]. The drug sensitivity was compared between 2D and 3D cell culture conditions and revealed a significant higher drug resistance when cells were cultured in 3D scaffolds compared to monolayer (2D) cell cultures. The use of silk sponges was also described as 3D scaffold for the expansion of SaOS2 and U2OS cell lines, which appeared less sensitive to drug treatment (doxorubicin and cisplatin) than cell lines cultured in a 2D environment [69]. 3D SaOS2 and U2OS cell spheroids were also generated by using commercial culture plates coated with various matrix: hyaluronic acid, collagen and adhesion proteins (BiomimesysTM matrix, from HCS Pharma, France). By using such an approach, 3D cultures differentially modulated the ATP binding cassette transporter A1 (ABCA1) and B1 (ABCB1, also known as Multidrug resistance protein-1, MDR1) expressions associated with the drug efflux and resistance which has not been observed in 2D environments [70]. 3D culture cells exposed to doxorubicin were characterized by a higher expression of ABCB1 implicated in the intracellular drug efflux, induced by the ERK1/2/HIF-1 pathway. Surprisingly, the increase of expression of ABCB1 was accompanied by a reduction of ABCA1 and by T-cell inactivation. ABCA1/ABCB1 ratio was 242 then related to the chemo- and immune sensitivity of human OS. These resistances were reversed 243 when ABCA1/ABCB1 ratio was increased [70]. This observation suggested that the upregulation 244 of ABCB1 transporter may induce an anti-drug and anti-immune resistance properties in OS 245 tumours. 246 The physical characteristics (e.g. elasticity) of the scaffolds influence cancer cell properties. The 247 generation of a hydrogel with a tuneable network of PEGDA and Gelatin-Methacryloyl (GelMA) 248 enabled the control of the stiffness and adhesion properties of the substrate. The stiffness of the 249 substrate correlated with the proliferation and progression of SaOS2 OS cells which proliferated 250 much better when the rigidity of the substrate increased. This stiffness dependency relied in the 251 regulation of the integrin-mediated focal adhesion signalling pathway [71]. Similarly, a recent 252 study compared the viability of OS MG63 cells cultured in four different scaffolds (collagen, MatrigelTM, alginate and agarose) and demonstrated that their viability was also dependent on the 253 254 scaffold elasticity [72]. Whereas cell adherence was similar for the different cell types in 2D models, ranking collagen as the best substrate followed by MatrigelTM, alginate and agarose, 3D 255 256 cultures of OS cells were more dependent on substrate elasticity for an optimal proliferation. In 257 this case, robust gels such as collagen and agarose are more proliferative substrates than softer hydrogels, MatrigelTM and alginate. Interestingly, even if the four substrates were able to produce 258 259 in vivo tumour in animal model, tumour size and angiogenesis process also correlated to the 260 elasticity of the substrate and showed higher size and micro-vessel formation in collagen and agarose than in MatrigelTM or alginate [72]. Mechanical properties of the ECM were also related 261 262 to the drug resistance of cancer cells. Molina et al. developed a 3D culture system that allows 263 mechanical modulation of TME by changing substrate stiffness [73]. They observed that lower 264 stiffness induced the nuclear localisation of mechanotransduction pathways, contributing to 265 specific drug resistance to anti insulin-like growth factor-1 and mTOR drugs [73]. By using 266 collagen scaffold, Fallica et al. demonstrated that U2OS osteosarcoma cells exhibited increased 267 resistance to the anti-proliferative drug PI103 in 3D gels than in conventional 2D cultures [74]. 268 These authors observed that the increase of collagen concentrations augmented the resistance of 269 OS cells to the inhibitor. This observation was in agreement with many clinical cases in which 270 the increase of collagen levels in TME was associated to a poor patient survival [75]. Reinforcing 271 the importance of collagen scaffold composition, Charoen et al. demonstrated that concentration 272 of 3-4 mg/ml of type I-collagen gels was crucial for optimum development of OS spheroids, whereas the optimum concentration for MDA-MB-231 breast cancer spheroids was 2 mg/ml [76]. These data suggest that production of specific tumour niches depends on tissue ECM composition. MatrigelTM or agarose were replaced by methylcellulose for facilitating the development of cancer cell spheroids. Based in an *in vitro* methylcellulose scaffold model, Bai *et al.* generated spheroids from HOSS1 OS cell line and various soft-tissue sarcomas including HT1080 fibrosarcoma, RD rhabdosarcoma, SW872 liposarcoma cells. Spheroids formed in this 3D environment showed more resistant properties to doxorubicin, gemcitabine and docetaxel or X-ray radiation than those formed in 2D cultures [77].

Tumours are characterized by a high heterogeneity of cell distribution with a necrotic or apoptotic core surrounded by quiescent layer of cells followed by proliferative cells. This tumour stratification is associated with a different TME composition in each tumour region. The determination by mass spectrometry imaging (MSI) of the spatial distribution of metabolites in response to doxorubicin treatment on SaOS2 OS cells cultured in alginate compared to 2D underlined the role of the 3D environment. The combination of 3D culture and MSI techniques represent a new tool to better understand drug activities and design new therapeutic approaches [78]. In addition to its role in anti-tumour drug resistance, TME plays a role in cell accessibility for genetic manipulation. 3D mineralized alginate-chitosan cell encapsulation resulted in an efficient tool for gene transfection in human bone cells [79]. Polysaccharide beads facilitated gene uptake by SaOS2 cells when specific calcium phosphate and chitosan rate were used indicating that microcapsule environment composition is crucial for gene transfection in 3D bone model [79].

2-1.3. Osteosarcoma cancer stem cells and 3D culture methods

OS, and other tumours, are composed by highly heterogeneous cell populations that include "Cancer Stem Cells" (CSCs) or 'tumour-initiating cells" [6,80,81]. CSCs combine stem cell features with tumour characteristics as tumour initiation ability, dormancy, recurrence and metastasis [80,82]. Thus, CSCs respond differently to anti-tumour treatments than non-CSCs tumour cells by showing a more resistant drug phenotype and leading the role of treatment failure [81]. CD133 CSC spheroids were generated from the SaOS2 cell line by using a scaffold-free 3D model based [83]. The generated spheroids were viable, conserved their pluripotency, and

constituted an ideal model for drug screening. CSCs enrichment from MG-63 and SaOS2 spheroids by scaffold-free method was combined with two hybrid scaffolds that mimic ECM and used to analyse the impact of ECM in OS CSCs development [84]. Hybrid scaffold was constituted by Mg-doped hydroxyapatite coupled to collagen fibres and a porous hydroxyapatite substrate. Both hybrids scaffolds resulted in stable CSCs enriched OS spheroid growth without any loss of round morphology compared to 2D. Moreover, an increase of stemness markers including OCT-4, NANOG and SOX-2 was observed that indicated that both types of hybrids scaffolds were able to mimic native environment promoting CSC stimulation [84]. Hydroxyapatite nanoparticle 3D cultures had a strong impact on the survival of OS cells under anti-tumoral oxidative stress therapy [85]. Cold atmospheric plasma resulted in a potential therapy in OS by induction of oxidative stress and subsequently cell death in 2D cultures. However, when this therapy was tested in 3D, MG-63 OS cell cultures in hydroxyapatite nanoparticles were characterised by a significant decrease of cell death [85]. This property was related to 3D environment due to the nanoparticles that favoured cell scavenging and evasion from reactive oxygen and nitrogen particles. Moreover, the generated 3D TME enhanced CSCs subpopulation expansion [85]. These data suggest a relevant role of TME in the development and drug resistance of CSC on OS, and the advantage of the use of 3D culture techniques that mimic native 3D OS nature unlike 2D approaches.

2-1.4. Proteomic profile in osteosarcoma 3D cultures

Protein expression and modification are highly impacted by nutrient availability and TME. Interestingly, the protein expression profiles significantly differ between 2D and 3D culture. A proteomic study using spheroids produced by ultra-low attachment supports from the dog OS cell line D17 demonstrated that the development in 3D culture induces an increase of glycolysis/glucogenesis pathways, biosynthesis of amino acids and changes in carbohydrate metabolism [86]. These data were in agreement with the metabolism observed during tumour development and the generation of a hypoxic local environment. Chaperon's family, which is composed by protein folders associated to cellular stress response and cytoskeletal organization, is similarly modulated by 3D context. On the opposite manner, general protein phosphorylation is upregulated in 2D cultures compared to 3D environment, probably due to the increase in the growth rate observed in monolayer cell cultures [86]. These data suggest that, in order to better

understand the proteomic profiles presented in tumours, all the previous information obtained by 2D studies must be re-evaluated in the light of the 3D culture methods close to the tumour behaviour.

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2-1.5. 3D osteosarcoma culture as a novel approach to study bone mineralisation

- 341 OS is characterised by the production of mineralised tissue. Osteoblastic-like OS cells show an 342 increase in the level of protein implicated in the mineralisation process (as TNAP, BMP-2 and 343 CaSR). The faster osteogenic properties of OS cells make them an interesting tool to better 344 understand the bone mineralisation process. While 3D culture techniques have been widely developed for mimicking TME and, subsequently, more reliable anti-cancer drugs screening, 345
- 346 bone 3D cultures were also used to study osteogenesis biology.

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Natural and synthetic scaffold gels generate a macro- and microstructural configuration similar to the trabecular bone identifying these materials as perfect supports to study bone mineralisation [87-89]. Type I collagen stands out as a great 3D scaffold for bone mineralisation in which SaOS2 cells can be expanded. Magnesium is a key cation involved in many biological activities such as metabolism, muscle contraction and bone cell function. Almost 50% of the magnesium present in the body is associated with bone tissue (hydroxyapatite crystals, HA) and influences 354 bone-remodelling processes. By using a 3D collagen scaffold approach, Picone et al. showed that intracellular magnesium was incorporated at the early phase of bio-mineralisation, a process 356 which may favour HA platelet formation and interfibrillar mineralisation [90]. The composition 357 of culture media appeared critical similarly to the 3D environment. Indeed, a Modified Eagle's Medium resulted in a better mineralisation induced by SaOS2 that conventional medias (e.g. 359 Dulbecco's Modified Eagle's Medium) used for this cell line [91].

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The morphology of mineralised matrix produced by HOS OS cells strongly differed according the cell culture condition used (2D to 3D) [92]. While 2D cell cultures produced spheroid particles in the surrounding cell layers, in 3D culture conditions (type I collagen and poly-ion complex hydrogels) amorphous mineralised particles were observed at the matrix layers. This phenomenon produced gel turbidity that could be used as an indicator of the level of mineralisation [93-95]. These data indicate that 3D gels are interesting approaches for osteogenesis studies associated with OS development. However, structure and mechanical properties of 3D gels are crucial parameters for investigating bone mineralisation. Similarly to tumour development, the pore size of the matrix used to promote osteogenesis is a critical factor. Using bioprinting approach, Vanderburgh et al. showed that a 300 µm pore size produced the optimum osteoblast differentiation and mineralisation. In addition, this pore size also favoured OS tumour growth and proliferation [96]. The distribution of pores also influences the structure of the extracellular matrix. The comparison of two types of 3D poly(D,L-lactic acid) scaffolds, one with regular pore distribution and the other with random distribution, showed that both types were adapted substrates for the attachment and proliferation of MG63 OS-derived osteoblasts. However, the random pattern, which is closer to real bone structure, induced a better distribution and organization of collagen fibres [97]. Polydimethylsiloxane (PDMS) is widely used as material to produce microfluidic devices but it can be also used as a scaffold for cell cultures [98]. A water-PDMS emulsion was used as a porous template for SaOS2 OS proliferation. Playing with different curing parameters for PDMS and pressure, Riesco et al. generated various grades of PDMS reticulation that allowed proper adhesion and proliferation of OS cells. Moreover, this system provides a fast and cheap way to produce scaffolds in mass [98]. As mentioned before, PDMS is the most commonly used material for microfluidic device fabrication (Figure 3). A microfluidic chip was developed for the production of OS spheroids in mass (up to 5000) [99]. Microfluidic device was treated with a surfactant (Synperonic®) to generate a nonadherent surface that favoured the generation of MG63 spheroids in a similar way as nonadherent plates. Massive production of spheroids was used to challenge spheroids to two different cellular stresses: nutrient deprivation (serum concentration) and hypoxia (HIF inhibition). 3D cultures obtained data confirmed in vivo observation where stress conditions favoured the increase of VEGF secretion and induction of malignancy processes [99]. This study confirmed the impact of ECM variation in tumour malignancy as well as demonstrated that microfluidic approaches represent an interesting tool for massive 3D cultures and analysis. However, it has been reported that the combination of commercial OS tumour cell lines and collagen and MatrigelTM scaffolds may not be the perfect model for 3D tissue bone engineering studies [100]. The hypoxia observed in 3D cultures using scaffolds like collagen and MatrigelTM generated less oxidative stress by tumour cells than in 2D. In addition, this oxidation can negatively impact the

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bone mineralisation [100]. By using non-tumour cell lines, Gamblin et al. generated 3D cultures to study osteoblastic and osteoclastic differentiation [101]. Based on biphasic calcium phosphate microbeads scaffolds, these authors induced the proliferation of MSCs that adhered and proliferated with abundant production of collagenous ECM. Interestingly, the system promoted the co-cultures of differentiated MSCs with osteoclasts generated from peripheral blood CD14positive monocytes. Altogether, the system was able to mimic bone precursors behaviour and established a valid non-tumour approach to study drugs for bone healing, osteoporosis and OS biology [101]. The production of hybrid scaffolds as the combination of layers of biodegradable polymer poly(\(\epsilon\)-caprolactone) (PCL) and layers of chitosan in combination with HA by electrospinning method resulted in continuous micro- and nano-fibers with high surface area and micropores that provided optimal attachment and proliferation of SaOS2 OS cells with high mineralisation activity [102]. Bioprinting methods have been also used to produce hybrid hydroxyapatite-chitosan-genipin hydrogels to analyse nanomechanical properties of generated bone tissue [103]. Bioprinter hybrid scaffolds resulted in a good structured TME that favoured 3D MG63 OS cell adhesion, culture and proliferation [103], indicating that bioprinting methods constitute interesting platforms to analyse how the composition and architecture of ECM impact bone mineralisation.

Other types of scaffold methods for 3D culture of OS cells imply the use of stirred-tank bioreactors [104]. Base on this equipment, Chen *et al.* developed a fibrous bed bioreactor with a 3D polyester fibrous matrix that resulted in a better production of 143B OS spheroids compared to 2D cell cultures. Moreover, they gave evidence that a 3D scaffold favoured the retention of viable and non-apoptotic tumour cells together with a long-term stability [104].

2-1.6. 3D culture methods for deciphering osteosarcoma metastatic process

OS cells are characterized by their ability to spread to distant tissues forming metastases (lung and bone) as carcinomas. To understand the process of cell dissemination and metastasis development, 3D bone tissue cultures were produced using a microfluidics device to mimic and analyse the "metastatic" installation in bone [105]. Hao *et al.* [106] developed a bone-on-a-chip microfluidic device in which they generated mature osteoblastic tissue using the MC3T3osteoblast precursors cell line. Cells produced a layer of heavily mineralised collagen

fibres up to 85 µm in thickness. By using this system, they analysed similarly the capability of metastatic breast cancer cell line (MDA-MB-231) to invade bone tissue. After 14 days of coculture, cancer cells seeded and invaded the apical layer of mineralised bone tissue in an "Indian file" and formed "micro-metastases" [106]. Choudhary et al developed an interesting microfluidic PDMS device that contained culture chambers in which primary human osteocytes were cultured in the presence of collagen-coated biphasic calcium phosphate microbeads for producing bone tissue in hypoxic conditions. Co-culture of conditional reprogrammed prostate cancer cell line (PCa3) with 3D osteocyte culture induced an increase of fibroblast growth factor-23, RANKL mRNA expression levels and alkaline phosphate activity by osteocytes that was associated to an increase of the mineralisation process. These results suggested that 3D microfluidic devices can be useful to better understand the metastatic process induced by primary tumours in bone tissue [107]. 3D microfluidic devices were also used to analyse the cell traction force in confined TME [108]. This device consisted in deflectable PDMS microspots included in micro-channels with different wide cross-sections. Migration test using HOS OS cells demonstrated that, in contrast to what observed in non-confining microchannels, tumour cell traction forces did not depend on myosin-II. This result showed that migration mechanisms of tumour cells during metastasis can vary depending on tissue structure, which compromises antimetastatic drug approaches. Moreover, the traction force devices resulted in an appealing approach for new anti-metastatic drug selection screening [108].

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2-1.7. Combination of 2D and 3D culture methods for the study of new vessels during osteosarcoma development

An alternative to the development of complex 3D micro systems is the combination of 2D and 3D cultures. To study angiogenesis process during tumour development, 3D MG-63 OS spheroids were generated by hanging drops using Gravity PLUS plates, reaching a size of 400 µm in diameter, and co-cultured with a HUVEC endothelial monolayer [109]. MG-63 OS spheroids produced similar ECM compared to *in vivo* tumours and acquired similar tumour architecture with proliferation cells at the periphery and quiescent cells at the centre of the spheroids. The generation of a hypoxia compartment induced the production of VEGF factor by tumour cells promoting proliferation and differentiation of HUVEC to produce vascular tubule-like structures. Using dog OS cell lines (D22 and D17) and 3D collagen gels, Massimini *et al.*

demonstrated that a non-human OS model was associated with induced vasculogenic mimicry and that 17-AAg drug abolished tumour progression and micro vascular channel formation [110].

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2-2. 3D Ewing sarcoma culture models

ES is the second most common paediatric bone malignancy and the third most frequent primary bone sarcoma after OS and CS. In addition to the *EWS/ETS* fusion gene which is at the origin of the disease, numerous investigations highlighted the contribution of the TME in the progression and malignancy of ES [111]. Similarly to OS, scientific community has put their effort to set up new *in vitro* ES models (Table 1).

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2-2.1 Drug resistance in 3D Ewing cell cultures

Electrospun polymeric scaffolds based on the inert polymer PCL have become a promising 3D platform to study mechanistic and drug resistance processes in TC-71 ES cell line. TC-71 3D culture reproduced morphology, proliferation and protein expression similar to observations in human tumours. As remarked in OS models, a 3D configuration induces more resistance to drug treatment (doxorubicin) than 2D cultures [111,112]. Interestingly, the PCL 3D model revealed that the IGF-1R/mTOR signalling pathways were highly activated in ES 3D models and these pathways played a key role in the upmodulation of tumour cell adhesion, identifying IGF-1R/mTOR signalling pathways as new potential targets for drug treatment in ES [111]. Similarly, Santoro et al. emphasized the role of IGF1/IGF-1R pathway and biomechanical TME stimulation in drug resistance by using similar 3D models [113]. These data are in agreement with the recent study published by Molina et al. [114]. Indeed, these authors demonstrated that 3D TME favoured the downregulation of IGF-1R via mTOR pathway, which was accompanied by a reduction of the clathrin-dependent nuclear localisation and transcription activity of IGF-1R [114]. TC-71 3D culture was exposed to different shear stresses close to those observed in bone microenvironment in a flow perfusion bioreactor. Under shear stress, 3D ES cells enhanced cell tumour proliferation and induced an increase of IGF1 pathway compared to 2D cultures. Besides, the increase of IGF1 levels was associated to the resistance to dalotuzumab (an inhibitor of IGF1 receptor) and the downregulation of the c-KIT and HER2 oncoproteins [113]. These data suggested that biomechanical forces impacted the progression and malignancy of ES cells [113]. When ES spheroids were grown in a 3D mimic bone tissue, there was an increase in ERK1/2 phosphorylation and RUNX2 protein levels associated to drug resistance that was not observed when the same cells were cultured in 2D [115]. Interestingly, an increase of RUNX2 level was similarly observed in patients suffering from ES [115]. All those data suggest that ECM displays a key role in ES drug resistance by induction of specific mechanotransduction signalling pathways including RUNX2.

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2-2.2. Impact of tumour microenvironment in 3D Ewing cultures

ES are characterised by a rapid development of multidrug resistance due to the overexpression of Multidrug resistance associated protein-1 (MRP1) and ABCB1 [116]. Similarly to OS, the drug resistance of ES cells was also related to a set of stem cells with tumour-initiating properties with a development dependent of TME [116]. Supporting this idea, ES spheroids produced by ultralow attachment method under serum-free conditions failed to generate tumour-initiating cells [117]. Although ES cells (VH-64, TC-32, TC-71 and A4573) form spheroids in serum-free media with a diameter ratio of 200 µm and with phenotypes similar to 2D cultures, none of them was able to self-renew or expand in a clonogenic manner indicating that TME is a key factor for ES enrichment [117]. Reinforcing the role of TME in ES development, Villasante et al. developed an interesting 3D model based on the combination of an engineered bone tissue with spheroids produced from different ES cell types (RD-ES, SK-N-MC and EWS-GFP) [118]. Bone tissue was generated from induced osteogenic differentiation of human MSCs in a native bone ECM, whereas ES spheroids were formed thank to the intrinsic nature of ES cells to generate cell aggregates after long period of culture (one week at 37°C in Eagle's Minimum Essential Medium supplemented with 10% Hyclone FBV [119]). The co-culture of these ES spheroids in the tissue bone matrix recapitulated the tumour behaviour, including the re-expression of focal adhesion and related cancer genes, generation of a hypoxic and glycolytic phenotypes and development of angiogenesis potential [118]. This study pointed out the requirement of specific niche configuration for proper development of ES. Reproduction of hypoxic conditions is indeed necessary to better understand how tumours develop angiogenic mechanism. Agar coated plates were used to induce A673 ES spheroids that were moved to a hypoxia chamber for analysing the functional relationship between hypoxia, spheroid cell distribution and DNA damage response [120]. Under hypoxic conditions, A673 ES spheroids displayed a stratification of cellular population from necrotic cells at the nucleus of the spheroids to proliferating cells located at their surface. Moreover, cells localized at the nuclear and perinuclear zone of the spheroids were characterised by an increase of γ -H2AX via the ATM DNA repair pathway, indicating that this approach can be used for anti-ATM drug development in ES [120]. Recently, a new approach was described for encapsulation in alginate spheres of ES cells isolated from patient derived xenografts without losing their phenotype [121]. While ES primary cultures were maintained for at least one month, cells at the core of spheroids did not undergo to hypoxia which is a key step of tumour angiogenesis. That could be due to the limitations in spheroid size by the alginate beads tested (<200 μ m), as hypoxia has been observed when spheroids reach a size over 400 μ m [122]. Overall, this method has an interesting potential as drug screening platform and can be easily implemented for hypoxia studies. In addition, encapsulation of different cell types in alginate beads can be a useful tool for cell-to-cell interactions in ES.

2-3. 3D chondrosarcoma culture models

CS is the second most frequent bone cancer characterised by the production of malignant cartilaginous matrix [123,124]. Surgery is the only effective medical treatment as CS are characterised by a high resistance to chemo- and radiotherapy. However, the mechanisms that control and regulate CS differentiation are still not well defined. MSCs can undergo into chondrogenic differentiation and have been used to understand the gene expression that determine 3D chondrogenic mechanism [125]. Chondrogenic differentiation of MSCs was associated to an increase of *SERPINA1* and *SERPINA3* mRNA expression. Moreover, secretion of SERPINA-1 correlated with chondrogenesis and dedifferentiation during chondrocyte expansion, suggesting that SERPINA1 could be considered as a marker of chondrocyte differentiation [125]. Similarly, MSCs were used for studying chondrocyte differentiation in spheroids and simultaneous gene expression profiles to determine genes implied in prechondrogenic and chondrogenic phenotype compared to tumour samples [126]. Comparative gene analyses allowed the identification of two clusters that mainly include ECM components, remodelling matrix enzymes and few growth factors useful to predict the clinical behaviour of CS subtypes [126].

2-3.1. 3D models as tool to unravel drug resistance in chondrosarcoma

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As mentioned above, CS are resistant to conventional chemotherapies. CS cell resistance can be explained in part by the structure of the tissue and its dense hyalin ECM composition, indicating that 3D models will become crucial tools to better understand the mechanism underlying CS drug resistance (Table 1). Comparison study between 2D and 3D chondrogenic spheroids from different origins (SW1353, CAL78 and OUMS27) and new established CS cell line from primary tumour biopsy (CH03, CH34 and CH56) showed that 3D CS spheroids were more resistant to chemotherapeutic drugs (Doxorubicin and Mafosfamide) than 2D cultures [127]. As shown by RT-qPCR, these cell lines cultured in 2D cells lost expression of several genes (COL2A1, COMP, ACAN) implicated in cartilage development that was restored in 3D cultures. Moreover, the capacity of each cell type to produce cartilaginous matrix was directly related to its drug resistance [127]. These data suggested a direct functional relationship between cartilaginous matrix composition and chemoresistance. Similarly, spheroids produced by CH2879, OUMS27 and L835 cell lines were used to determine the mechanism involved in resistance to cisplatin and doxorubicin [128]. 3D spheroids were generated by differentiation of cell lines after long period of culture (6 weeks in chondrogenic medium, see [129]) and exhibited CS phenotype. Exposure to chemotherapy agents highly activated the multi-drug resistance pump (ABCB1) in all CS 3D spheroids. Inhibition of the anti-apoptotic BCL-2 family members by specific drug (ABT-737) resulted in a sensitization of 3D CS to doxorubicin. These results indicated that tumour drug resistance does not rely only on ECM composition and that other mechanisms must be implied and be considered as potential targets for development of new CS therapies. Taken together, these data demonstrated that the mimicking of the cell behaviour and ECM of CS is the added value of 3D cultures. As discussed for OS, non-adherent surface methods can be used for obtaining multicellular tumour spheroids. Combination of non-adherent plates and 0.5% methylcellulose generated CS spheroids from the HEMC-SS cell line [130]. HEMC-SS spheroids developed CS tumour features as proliferative cell population at the periphery of a hypoxic and apoptotic core, with ECM rich in glycosaminoglycans and VEGF excretion (figure 4). Moreover, this model recapitulated the drug resistance phenotype observed in CS tumour for classical chemotherapy agents [130]. While still far for native tumour environment, the absence of complexity to generate this 3D system and its close features to the in vivo tumours make it a convincing model for massive drug screening. In this way, HEMC-SS spheroid model was used to evaluate the effect of new hypoxia-activated pro-drugs that target the rich proteoglycan ECM of CS [131]. This study showed that quaternary ammonium, which is characterised by a positive charge that interacts strongly with the negative charges present in the proteoglycans [132], could be used as an adjuvant for CS drug targeting [131]. Generation of CS 3D cultures by similar hanging-drop methods showed the potential of the ionophore salinomycin (SAL) as a new anti-CS drug [133]. SAL resulted in a strong cytotoxic effect in both 2D and 3D SW1353 (grade II) CS model by inducing cellular apoptosis via caspase activation [133]. Another approach to produce 3D CS system consists in the use of natural or synthetic material that can serve as substrates for the formation of tumour mass cells in cartilage or chondrosarcoma niches. Alginate hydrogels were used to analyse the invasion and drug resistance of CS models [134]. Compared to other biomaterials as collagen and MatrigelTM, alginate is characterised by an inert and stable composition, which is translated in a more reproducible method to generate beads for encapsulation. Alginate encapsulated CH2879, JJ012 and SW1353 CS cell lines, compared to a 2D culture, were characterised by a long-term lifespan with generation of a hyaline-like cell matrix and demonstrated that 3D cultures recapitulated cell matrix gene expression. Interestingly, CH2879 cell line displayed an evasion phenotype from the beads compared to the other cell line models. This was somehow in coherence with the grade of malignancy III of this cell line whereas the other two are grade II. This data suggests that alginate beads are useful to analyse CS cell invasion properties. Thus, it can be combined with drug screening as alginate beads CS 3D model summed up the characteristic drug resistance phenotype of CS cell lines [134]. In agreement with this result and by using same methodology, Palubeckaite et al. showed that CS 3D spheroids reproduced similar phenotype that in vivo CS with production of an ECM enriched in collagen II and resistance to chemotherapeutic agents as doxorubicin, cisplatin, temozolomide and YM-155 [135].

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CS are resistant to radiotherapy. Hamdi *et al.* developed a 3D approach to analyse the impact of linear energy transfer ionizing radiation (LET) in CS [136]. Due to its relative photon radioresistance, metastatic potential and cartilage phenotype, intermediate-grade II SW1353 CS cell line was used as a model. 3D culture was performed by using a collagen scaffold and hypoxic conditions (2% of O₂) to mimic *in vivo* cartilage environment. Exposition to low and high LET radiation showed that 3D cultures resulted in a more resistant phenotype with a delayed response

of DNA repair mechanisms (evaluated by H2AX expression post radiation) than 2D cultures, where cell were grown in monolayer conditions and normoxia (20% oxygen). The main difference between culture methods relied in the microenvironment conditions, which implied that the microenvironment played a key role in the radiation resistance of CS and should be considered when determining the depth-dose profile in radiation therapy of SC [136].

2-3.2. Chondrosarcoma 3D culture approaches to investigate cell adhesion, migration and

cell-to-cell interactions

3D CS cultures were used to determine the impact of TME charges during cell adhesion [137].

Titanium beads from 400-500 µm were modified by deposition of polyelectrolyte multilayer film

that conferred a positive or negative surface charge. In the presence of negative charges, human

HCS-2/8 CS cells exhibited cytoplasmatic stress fibres that were totally absent when positive

charged TME were tested, indicating that anionic charges affected cytoskeleton organisation.

Moreover, anionic but not cationic surfaces promoted two modes of pseudopod formation by

HCS-2/8 cells, in a random progression and as a "cell recognition signal". This phenomenon was

associated with a cellular mechanism to optimize the anchoring process. Interestingly, cells

developed pseudopods on cationic surfaces when cells were cultured in the presence of

conditioned medium obtained from an anionic culture, suggesting that this process was regulated

by an exocytotic mechanism. This mechanism was linked to the MAPK ERK1/2 pathways as

phosphorylation levels were increased in the presence of anionic charges and reduced when

cationic surfaces were assessed [137]. Overall, these studies suggested that ECM has a relevant

role in CS cell adhesion and migration.

As described for OS, direct or indirect cell-cell interactions between CS cells and cells of the local microenvironment can be analysed by combination of 3D and 2D cultures. In a recent study, Minopoli *et al.* analysed the contribution of pro-tumoral M2 macrophages to CS development [138]. Primary CS cells were isolated from patient biopsies and cultured by hanging drop methods to produce 3D spheroids and then co-cultured in collagen/fibroblast matrix with blood isolated monocytes. The size of CS spheroids increased in the presence of monocytes, probably due to an increase of CS cell invasive capability induced by monocyte factors. And reciprocally, CS cells induced monocyte differentiation into a pro-tumoral M2 phenotype. These observations

indicated a crosstalk between CS cells and macrophages through soluble mediators [138]. The induction of CS proliferation by macrophages was inhibited by the addition of urokinase receptor (uPAR)-derived synthetic peptide RI-3 which was known to reduce the monocyte migration [139]. In this context, RI-3 could potentially avoid the recruitment of monocytes to CS niches and reduce proliferation and angiogenic properties of CS tumour [138].

Recently, an innovative method was developed to assemble 3D CS spheroids by levitational forces by using low doses of gadobutrol salt [140]. As biological tissues are considered to be diamagnetic, they can be levitated when paramagnetic medium is used. Paramagnetic ions of Gadolinium(III) (Gd³⁺) have been widely used as a contrast agent but are characterised by cytotoxicity at high concentrations and lower concentration of Gd³⁺ did not allow cell levitation. However, by using high magnetic fields, Parfeno *et al* induced SW1353 CS spheroid magnetic levitational bio assembly in the presence of lower doses of gadobutrol (0.8 mM). The study revealed minimal cytotoxicity by the magnetic field and opens a new area in the domain of microgravity and tumour biology [140]. In the meantime, further studies may be needed to determine if high magnetic fields have an impact in cell behaviour, genomic and proteomic expression pattern.

3-. Conclusion and future perspectives

Primary bone tumour progression and metastasis rely on the particular combination of bone MSC differentiation and physiological, structural, biochemical TME interactions. 2D cultures were considered for a long time as a valid approach for improving the knowledge in tumour biology. Many studies gave evidence that 2D cultures did not reflect the real nature of tumours, as many treatments that were effective in 2D failed in clinical trials. In this context, development of 3D culture approaches that mimic TME is a new perspective. In particular, the use of natural materials already presents in the bone ECM or synthetic materials as scaffolds for 3D bone culture generation demonstrates to be promising approaches for the study of tumour invasion, metastasis, angiogenesis processes and anti-tumour drug development. Moreover, the development of novel technologies in 3D cultures as microfluidics and bioprinting that allow the generation of customizable 3D systems constitute a fully scientific revolution. These technologies can be combined with other 3D techniques and allow the creation of a fully controlled TME that

- 676 reproduces the native configuration of bone tumours, including cell-to-cell or tissue interactions,
- 677 cell adhesion, proliferation and migration, EMC structure and composition, physiology
- parameters as hypoxia, shear stress and mechanical forces. This 3D technological revolution will
- be an excellent opportunity for the identification of new bone therapeutic targets and drug
- discovery in bone sarcoma field.

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Table 1. 3D methods used in primary bone tumors

Bone tumor	Technique	Material	Cell line	Reference
	Scaffold-free	Hanging drop and Low-attachment	SaOS2 HOS U2OS D17 MG63	61-66,83,84,86,109
		Alginate beads	LM8 MG63	68,72,78,83
		Silk sponge	SaOS2 HOS	69
		PEG	SaOS2	71
		Collagen	MG63 U2OS	72,74,76,90-94,110
		Agarose	MG63	72
Osteosarcoma	Scaffold	PCL	MG63	73,102
	Scariola	Methylcellulose	HOS	77
		PLA	MG63	97
		НА	MG63	84,85,102,107
		ВСР	OS MSC	101,107
		Complex matrix	SaOS2 U2OS MG63	70,72,84,92
	Microfluidic/Bioprinting			96,110
		PDMS	SaOS2 MG63	98,99,106-109
	Scaffold-free	Hanging drop and Low attachment	RD-ES A673 SK-N-MC VH-64 T-32 TC-71 A4573	112,117-119
	wing Sarcoma Scaffold	PCL	TC-71	111,113,114
Ewing Sarcoma		Collagen	RD-ES A673 SK-N-MC	115
		НА	RD-ES A763 SK-N-MC	115
		Agar	A673	120
		Alginate	Primary cell lines	121,122
Chondrosarcoma	Scaffold-free	Hanging drop and Low attachment	SW1353 CAL78 OUMS27 CH03 CH34 CH56 CH2879	127-131,133

			L835	
		Levitation forces	SW1353	140
	Scaffold	Alginate	CH28979 JJ012 SW1353	134,135
		Collagen	SW1353	136,138
		Titanium beads	HCS-2/8	137

PEG: polyethylene glycol; PCL: poly(ε-caprolactone; PLA: poly(D,L-lactic acid; HA: Hydroxyapatite; BCP: biphasic calcium phosphate; PDMS: Polydimethylsiloxane

1270 Figure Legends

- Figure 1. Osteosarcoma spheroid. Representative images of OS spheroid. A) HES staining of MNNG/HOS spheroid. B) Wide-field MNNG/HOS spheroid using a Nikon Eclipse Ni microscope. C) MNNG/HOS spheroid at day 3 obtained from 20.000 cells in DMEM (Gibco®) supplemented with 1% L-glutamine plus 10% FVS using 96-well low adherent plate U-bottom (ThermoFisher), labelled with VybranTM DiO (ThermoFisher) and imaged in a Operetta CLS microscope (PerkinElmer).
 - **Figure 2. Sarcospheres from different OS cell lines.** Representative images of OS spheroid formed from different OS cell lines depicting differences in size and morphology. MNNG-HOS and MG63 OS cells were plated at 5,000 cells/well and 2,500 cell/well respectively in low adhesion plates (Corning Costar®) coated with DMEM (Gibco®) + 10% agarose and imaged on days 1-4 using the Celigo Imaging Cytometry System.

Figure 3. 3D Spheroid PDMS chip. PDMS microsystem for spheroid cell culture in a 60 x 22 mm slide. Microsystem is constituted by a reservoir for media (a 15 ml Falcon tube cut at desired size) glued to the PDMS microsystem. The reservoir is connected to the cell culture chamber by an 8 mm length channel (200 μ m wide and 70 μ m high). Cell culture chamber is 4 mm wide by 20 mm long (height 200 μ m). To slowdown media flow, a 2 cm long serpentine channel (200 μ m wide 70 μ m high) was placed after the cell chamber. Output through 1.5 mm Tygon tubing with 500 μ m internal diameter.

Figure 4. Cell subpopulation in an OS spheroid. Spheroids are characterised by a continuum subset of cells that goes from apoptotic or bone-like MSC non-dividing cells (in red) to a peripheral proliferative subset of cells (green cells). 10,000 GFP-MNNG/HOS cells were seeded into a 96-multiwells low-attachment plate and cultured for 13 days. Pictures showed population evolution from day 5 to day 13. GFP expressing MNNG/HOS osteosarcoma cells stained with DiD (ThermoFisher) to show the retention of DiD by a non-proliferating subpopulation of the

cells in the formed spheroid and imaged using fluorescent microscopy. Scale bar corresponds to $1299-50\ \mu m.$

Figure 1

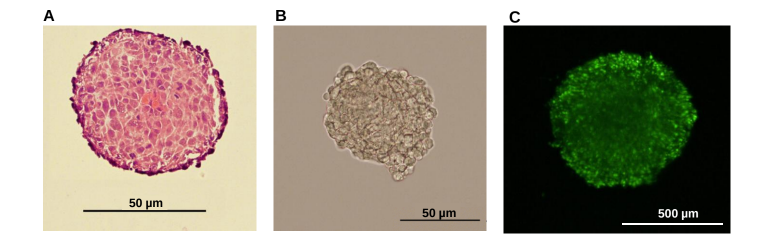


Figure 2

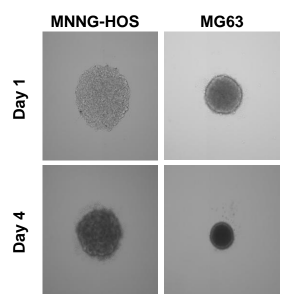


Figure 3

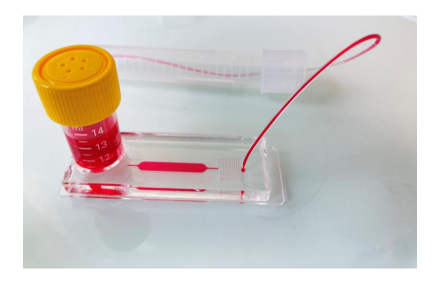


Figure 4

