2D and 3D cell cultures – a comparison of different types of cancer cell cultures

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Abstract

Cell culture is a widely used in vitro tool for improving our understanding of cell biology, tissue morphology, and mechanisms of diseases, drug action, protein production and the development of tissue engineering. Most research regarding cancer biology is based on experiments using two-dimensional (2D) cell cultures in vitro. However, 2D cultures have many limitations, such as the disturbance of interactions between the cellular and extracellular environments, changes in cell morphology, polarity, and method of division. These disadvantages led to the creation of models which are more closely able to mimic conditions in vivo. One such method is three-dimensional culture (3D). Optimisation of the culture conditions may allow for a better understanding of cancer biology and facilitate the study of biomarkers and targeting therapies. In this review, we compare 2D and 3D cultures in vitro as well as different versions of 3D cultures.

Key words: co-culture, cell culture methods, 3D culture, 2D culture, cancer research.

Cell cultures as a research model

Studies on the mechanisms underlying the formation, function and pathology of tissues and organs are manageable largely due to the use of cell culture systems and animal models [1]. Harrison carried out the first cell cultures in 1907 during research into the origin of nerve fibres [2]. Since then, the method has been improved and used to observe the growth and differentiation of cells outside the body [3, 4]. Nowadays, experiments can be conducted using primary cells isolated directly from the donors’ material or using established cultures deposited in cell banks [5]. Primary cultures are isolated from living organisms and usually contain populations of different cell types present in the source tissue. In this case, it is important to isolate the correct cell type [5]. Characteristic features of primary cell lines are: i) difficulties with isolation and ii) short life span. On the other hand, they closely mimic the in vivo genetic features of tumours and thus make it possible to perform some functional experiments. An alternative option is the use of an established cell line. Bioresource centres, such as
the ATCC (American Type Culture Collection), offer characterized models of various types of cancer cell lines that are routinely used in research [6].

Cell cultures make it possible to understand cell biology, tissue morphology, mechanisms of diseases, drug action, protein production and the development of tissue engineering [7]. They are often used in the preclinical research of many drugs, in cancer research, and in studies on gene function [5].

The choice of the most appropriate cell culture methods in the area of cancer research may allow us to better understand tumour biology, and hence to optimize radio- and chemotherapy, or even to find new treatment strategies [8].

The cultures can be carried out under adherent conditions wherein the cells are attached to a glass or plastic dish or in a suspension, which in some cases (e.g. cultures of lymphocytes) corresponds more faithfully to the natural environment [6]. The most commonly used type of cell culture is the 2D model, but recently the 3D culture method has been gaining in popularity (Figure 1) [9]. Depending on the type of culture chosen, cell behaviour differs in many aspects [7].

2D cultures

In adherent 2D cultures, cells grow as a monolayer in a culture flask or in a flat petri dish, attached to a plastic surface [10].

The advantages of 2D cultures are associated with simple and low-cost maintenance of the cell culture and with the performance of functional tests. Unfortunately, adherent cultures also have numerous disadvantages. First, 2D cultured cells do not mimic the natural structures of tissues or tumours (Figure 2 A). In this culture method, cell-cell and cell-extracellular environment interactions are not represented as they would be in the tumour mass. These interactions are responsible for cell differentiation, proliferation, vitality, expression of genes and proteins, responsiveness to stimuli, drug metabolism and other cellular functions [9, 11–13]. After isolation from the tissue and transfer to the 2D conditions, the morphology of the cells is altered, as is the mode of cell division. The loss of diverse phenotype is also a result of 2D culturing [14, 15]. The changed morphology of the cells can affect their function [16, 17], the organization of the structures inside the cell, secretion and cell signalling [18, 19]. Due to disturbances in interactions with the external environment, cells growing adherently lose their polarity [20], which changes the response of those cells to various phenomena, such as to apoptosis [21, 22]. Another drawback of 2D culture is that the cells in the monolayer have unlimited access to the ingredients of the medium such as oxygen, nutrients, metabolites and signal molecules. For cancer cells in vivo, the availability of nutrients, oxygen, and so forth, is more variable because of the natural architecture of the tumour mass [9]. Furthermore, it has been observed that the 2D system changes the gene expression and splicing, topology and biochemistry of the cell [23–26]. In addition, adherent cultures are usually monocultures and allow for the study of only one cell type [27], which results in a lack of tumour

![Figure 1. Types of cell culture methods commonly used in research studies. A – Cells flattened in a monolayer on the bottom of the culture vessel. They are in contact with the culture vessel, neighbouring cells, and the culture medium. B – Cells attached to a scaffold are in contact with the scaffold, neighbouring cells, and the culture medium. C – A group of cells suspended in the culture medium or cultivated in gel-like substance; the cells are in contact with neighbouring cells and with the culture medium.](image-url)
microenvironment, or niches, which in vivo are required by cancer-initiating cells [28, 29].

Owing to the many disadvantages of 2D systems, there was a need to find alternative models, better able to mimic a natural tumour mass, such as 3D culture systems (Table I).

**3D cultures**

One of the first three-dimensional cultures was made in soft agar solution, and was carried out by Hamburg and Salmon in the 1970s [30]. Since then, striking similarities between the morphology and behaviour of cells growing in a tumour mass and in cells cultured under 3D conditions have been well described and documented [9, 31].

Due to the method of preparation, 3D models can be divided into: i) suspension cultures on non-adherent plates (Figure 2 C); ii) cultures in concentrated medium or in gel-like substances (Fig-
ure 2 B) and iii) cultures on a scaffold (Figure 2 E). All of these models are characterized in Table II.

The concept of 3D spheres is based on the creation of spheroid structures in which cells form various layers. This structure mimics the physical and biochemical features of a solid tumour mass. Morphological analysis of 40 tumour cell lines (originating from: glioblastoma, astrocytoma, Wilms' tumour, neuroblastoma, head and neck squamous cell carcinoma, melanoma, lung, breast, colon, prostate, ovarian, hepatocellular and pancreatic cancers) cultured in 3D spheroid conditions led to the identification of three distinct groups according to the architecture of spheroid shapes: i) tight spheroids, ii) compact aggregates and iii) loose aggregates [32, 33]. Some cells under non-adhesive conditions display reduced cell-cell and cell-matrix interactions, lose their anchorage, escape from anoikis, divide and create spheres [34].

Cells from the donor's tissues are cultured in a culture medium, which is refreshed every 2-3 days [11, 34, 57]. Moreover, in 3D cultures, the morphology and polarity of the cells are maintained, and they can be restored to cells previously cultivated in 2D [1, 15, 38]. Furthermore, in some 3D systems, e.g. acinar-like spheroids, specific internal architecture with lumen formation is observed. This is the result of cell apoptosis in the central part of the spheroids. Cell proliferation depends on cell location and is higher in the peripheral part of the 3D structures [39–41]. Another important attribute of 3D culture is its similarity to cells growing in vivo in terms of cellular topology, gene expression, signalling and metabolism [42–47].

All these features create a specific platform which can be used for the study of the biology of cancer-initiating cells, invasion and metastatic processes, as well as for drug testing or for testing the response of cells to irradiation.

3D suspension culture systems are widely used as a model in studies, e.g. for increasing the population of cancer-initiating cells. This method allows for simple and low cost biological research [37, 48]. The spheroids obtained from oral cancer cell lines show an increased proportion of cancer-initiating cells, probably due to the epithelial-mesenchymal transition process occurring under 3D conditions. The spheres exhibit loss of E-cadherin expression, and overexpression of fibronectin, Sox2, Oct4 and Nanog. Expression of putative stem cell markers such as CD133 and ALDH also occurs. Sphere-related enrichment of the cancer-initiating cell population is also why a lower number of cells, derived from the spheres, is needed to generate a tumour in xenograft mice, compared to parental cells [37]. It has been observed that the number of spheres is reduced with

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**Table I. Comparison of 2D and 3D cell culture methods**

| Type of culture                  | 2D                                           | 3D                                           | Ref. |
|----------------------------------|----------------------------------------------|----------------------------------------------|------|
| Time of culture formation        | Within minutes to a few hours                | From a few hours to a few days               | [11, 34, 57] |
| Culture quality                  | High performance, reproducibility,           | Worse performance and reproducibility,       | [12] |
|                                  | long-term culture, easy to interpret,         | difficult to interpret, cultures more        |      |
|                                  | simplicity of culture                         | difficult to carry out                        |      |
| In vivo imitation                 | Do not mimic the natural structure of the    | In vivo tissues and organs are in 3D form     | [35] |
|                                  | tissue or tumour mass                         |                                              |      |
| Cells interactions                | Deprived of cell-cell and cell-extracellular | Proper interactions of cell-cell and         | [13, 28, 29, 36, 37] |
|                                  | environment interactions, no in vivo-like     | cell-extracellular environment, environmental |      |
|                                  | microenvironment and no “niches”             | “niches” are created                          |      |
| Characteristics of cells          | Changed morphology and way of divisions;     | Preserved morphology and way of divisions,   | [1, 14–17, 20, 38] |
|                                  | loss of diverse phenotype and polarity        | diverse phenotype and polarity               |      |
| Access to essential compounds    | Unlimited access to oxygen, nutrients,       | Variable access to oxygen, nutrients,        | [10, 46] |
|                                  | metabolites and signalling molecules         | metabolites and signalling molecules         |      |
|                                  | (in contrast to in vivo)                      | (same as in vivo)                            |      |
| Molecular mechanisms             | Changes in gene expression, mRNA splicing,    | Expression of genes, splicing, topology      | [23–26, 42–45] |
|                                  | topology and biochemistry of cells           | and biochemistry of cells as in vivo         |      |
| Cost of maintaining a culture    | Cheap, commercially available tests and the  | More expensive, more time-consuming,         | [8, 48, 58, 75] |
|                                  | media                                        | fewer commercially available tests           |      |
the passage of time, and that the percentage of ALDH+/CD44+ cells decreases during the culture period, which may indicate an ongoing differentiation process [48]. Campos et al. observed that ALDH+/CD44+ cells create more orospheres, with higher proliferation rates and are more resistant to anoikis than ALDH-/CD44- cells growing in suspension conditions. This is probably caused by secreted endothelial factors which activate PI3K-Akt signalling, which in turn influences cells obtained from primary and metastatic cancers differently [49]. Sphere formation assays (in suspension or soft-agar cultures) are commonly used and inexpensive assays for evaluating the role of examined genes in self-renewal and maintenance of tumour stemness [50].

It is not surprising that tumour cells are less sensitive to drugs in 3D than in 2D cultures. This effect may be caused by reduced access to compounds in the medium or by pathophysiological differences due to hypoxia, or by changes in the cell cycle [35, 36]. Hsieh and colleagues showed that unstable culture conditions, which sometimes occur in vitro, and the type of culture method used can affect the results.

Table II. Characteristics of different 3D cell culture methods

| Type of 3D system | Description of cell culture | Advantages | Disadvantages | Ref. |
|-------------------|-----------------------------|------------|---------------|-----|
| Suspension cultures on non-adherent plates | • Single cells are seeded on non-adherent plates with medium | • Simplicity, easiness and speed of conducting culture | • Some cell lines need expensive plates coated with specific materials, for example polystyrene or covalently bound hydrogel, because of strong adhesion abilities of cells | [8, 48, 58, 59] |
| | • 3D structures can be observed after 3 days of culture | • Bacterial plates or non-adherent culture plates can be used but only for some cell lines | • Formation of aggregates of cells as a result of cells’ movement in medium | |
| Cultures in concentrated medium or in gel-like substances | • Single cells grow in medium containing substances with gelling properties: i) dissolved low-melting agarose with cell medium is poured on plate and incubated until solidifying to obtain the first, lower layer; the top layer consisting of agarose and the medium with single cells is added; ii) the cells are flooded in Matrigel (multiprotein hydrogel) | • Soft agar allows to study both the growth of a single cell regardless of attachment and the phenomenon of escape from anoikis | • Difficulty in obtaining spheres for certain cell lines, inconvenient and time-consuming preparation of the two layers of agar and requirement of long-term cultures | [7, 48, 58, 59, 75–81] |
| | • 3D structures can be observed after 7 days of culture | • Cells cultured in Matrigel can be easily recovered for further analysis | • Low repeatability of the results | |
| Cultures on scaffold | • The cells can migrate among fibres and attach to the scaffold, made of biodegradable material such as silk, collagen, laminin, alginate, and fill the space among fibres, grow and divide | • System is compatible with commercially available functional tests, as well as with DNA/RNA and protein isolation kits | • Cells attached to the scaffolds flatten and spread like the cells cultured under adherent conditions | [7, 8, 37, 82–90] |
| | | • Easy to prepare for immunohistochemical analysis | • Scale of scaffolds and topography of cell distribution may cause various behaviour of the cell | |
| | | | • Materials used to construct the scaffold may affect the adhesion, growth and cell behaviour | |
| | | | • Cell observation and cell extraction for some analyses are restricted | |
od used can significantly influence cellular metabolic activity, cell proliferation and, ultimately, changes in cell sensitivity to tested drugs. They also indicated that among 2D, 3D and spheroid models, only 3D cell culture, with the same cell density as natural tissue, shows a drug response comparable to that of a solid tumour [51].

Cell extracellular matrix (ECM) interactions seem to play an important role in the drug resistance of tumours. Cells growing in a 3D silk scaffold system, which have been found to be similar in fibre orientation and dimensions to native tumour ECM, are more resistant to paclitaxel. The use of artificial ECM is a good way to mimic the natural architecture of a tumour mass. It has been found that changes in ECM composition are associated with cancer progression and tumour features [52]. In this context, the use of 3D systems could avoid over- or underestimation of a specific drug in case of drug sensitivity and resistance, as well as its dosage [35, 36].

As mentioned above, the spheroids show different responses to drugs, but also the spatial structure of spheroids influences the irradiation response. Increased radiation survival is brought about by 3D architecture, which influences DNA heterochromatinization, characterized by deacetylation of histone H3 and high expression of heterochromatin protein 1α (HP1α). It should be noted that higher levels of heterochromatin partly protect DNA against radiation-dependent induction of double strand breaks in 3D structures. This phenomenon could be overcome by knockdown of histone deacetylase (HDAC) 1/2/4 or by application of the HDAC inhibitor LBH589. However, neither growth conditions nor HDAC modification affects ATM phosphorylation [53]. It has been shown that in some cases, such as head and neck cancers, integrins and their signalling cascades are critical for cell proliferation and survival. Use of the FAK/IGF-IR inhibitor TAE226 demonstrates strong radiosensitizing potential.
under in vitro 3D conditions, which strongly suggests that this inhibitor has potential in clinical practice [54]. Furthermore, the behaviour of cells cultured under 3D conditions shows that the combined targeting of FAK/IGF-IR by cetuximab and TAE226 induces cell death without the need for further irradiation [55].

Spheroids can be used to study the process of cell migration on ECM proteins, invasion into Matrigel, or simultaneously tissue invasion and angiogenesis. Characteristic features of migration are visible flattened cells surrounding spheroids (dispersed or radial migration). In the case of invasion the cells extend in visible invadopodia [35, 36]. Tissue invasion and angiogenesis assays can be performed by means of co-cultures of spheroids and embryoid bodies generated from mouse embryonic stem cells. This assay is designed to mimic xenograft tumour transplant systems [35, 36].

Moreover, 3D tissue culture system allows for the creation of imitation cancer tissue, with green fluorescent protein (GFP) expression and with the features of a solid tumour. The efficiency of transfection of anti-GFP oligonucleotides can be measured simply by fluorescence microscopy. Creation of systems with over- or down-regulated genes examined for usage in new treatment strategies is also possible. An example of application for 3D collagen matrix tissue structures could be in the establishment of an intracellular delivery system for oligonucleotides using the microneedle technique [56].

Apart from using 3D systems in the area of cancer research, they can also be applied in tissue engineering. For example, primary human salivary gland cells may be encapsulated in a 3D hyaluronic acid-based hydrogel scaffold, in order to obtain organized acinar-like spheroids with active protein secretion pathways. This approach might be used in the future to restore function to salivary glands damaged by radiation treatment [39–41].

A disadvantage of 3D cultures is that it requires the separation of single cells from spheroid structures by proteolytic degradation of single layers, which takes from several hours to a few days [57]. In many 3D methods, the efficiency, life-span, repeatability, and comfort of work are poorer than in the case of 2D systems [12]. It is often emphasized that a disadvantage of 3D structures is the fact that “spheres” can be formed, not from a single cell, but from a few cell clusters. However, even structures created as aggregates of several cells still have a three-dimensional form and seem to be a better model than flat, adherent cultures [58].

A tumour is not a homogenous structure, but is built from tumour cells of various phenotypes. Furthermore, 2D cultures are, in fact, also a compound of various cell phenotypes. In spite of this, a homogenous structure can be achieved from culture of a single cell, with only one genetic background in the concentrated culture medium, as is the case, for example, in soft agar or Matrigel [59].

The problem of low reproducibility in 3D culture was solved by Vinci et al., who described a three-dimensional spheroid-based functional assay for tumour target validation and drug evaluation. They used 96-well ultra-low attachment plates to create just one spheroid per well. The size of the obtained spheroids was reproducible and showed Gaussian distribution [32, 33].

Owing to the large number of problems associated with 2D systems, 3D models would appear to be a good alternative, that could be an intermediate model between 2D and animal studies [1, 30]. The different technical approaches to obtaining 3D models possess their advantages and their limitations (Table I). The proper choice of 3D system mostly depends on the nature of the research. It must be emphasized that choosing the wrong model may influence the results. Clearly, the ideal 3D model does not exist. In some cases the use of a 2D culture system is enough, but 3D will be used more frequently in the future due to improvements to automation and cost reductions.

2D and 3D methods in co-culture systems

A tumour is a mass composed of multiple cell types [60]. In co-cultures, different cell types are grown together in the same environment [61]. This type of culture was described in the 1970s as a system by which to examine communication among cells [62]. Such communications include three types of intercellular interactions: cell-cell, cell-microenvironment and paracrine signalling by dissolved factors [61]. This allows us to observe interactions in functional structures that closely resemble interactions in vivo [63].

In co-culture we can distinguish between target cells and assistance cells that support their growth and development [61]. Studies show that both types of cells gain through co-culture [64].

Co-cultures can be divided into two types: direct and indirect [61]. In the first model, different types of cells are mixed and cultured together. In the second, cells are separated by a physical barrier [61]. Both types of co-culture can be carried out in 2D systems [65, 66], as well as in 3D models [67, 68]. In direct cultures, we can observe all three types of interactions described above, which would appear to be of great importance in the study of cellular behaviour [69, 70]. In contrast to the direct system, in the indirect model, the cells are deprived of interactions between the types of cells by the presence of a physical barrier [68, 71].

Our experience has shown that cells of different phenotypes do not grow with each other, even in direct models. Figure 4 shows mesenchy-
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Abnormal fibroblast cells (green) cultured with epithelial cancer cells (red). It can be seen that the cells grow within the limits of their own line both in 2D and 3D systems. Moreover, it was observed that under 3D conditions, the SCC-25 line (red) formed a 3D structure, while the MSU-1.1 line (green) surrounded the established structures as assistance cells.

The downside of co-cultures is the inclusion, by necessity, of many variables (i.e. the degree of similarity and separation of the population, the components of the medium, volume and duration of culture), which must all be optimized so that all types of cultured cells might thrive. All these factors make co-cultures difficult to conduct [72].

Conclusions

Cell cultures are commonly used in genetic and biological cancer research [6, 48]. They mimic in vivo conditions, to varying degrees, and may provide an alternative to animal models [73]. Currently, there are many forms of cell culture, which allows for the selection of a method well suited to the purpose of the study [74]. The most common research model is still the 2D culture system. However, owing to its limitations, 2D cultures are increasingly being seen as an inefficient model with which to study the processes associated with cellular responses to ionizing radiation or to exposure to chemotherapeutics. The 3D models are potentially a better approach in the search for new biomarkers and new treatment strategies, leading us closer to the goal of personalized medicine.

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Conflict of interest

The authors declare no conflict of interest.

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Figure 4. Co-culture of epithelial SCC-25 (red) and fibroblast MSU-1.1 (green) cell lines (scale bar represents 100 µm). A – cells cultured under 2D conditions are flattened and attached to the plate surface. The epithelial SCC-25 cells (red) have typical rhombus-like shape and MSU-1.1 cells (green) are spindle-like and surround SCC-25 cells; B – SCC-25 (red) and MSU-1.1 (green) cells cultured under 3D conditions changed their own morphology due to the lack of attachment. Cells lose their typical shape and aggregate, creating more (SCC-25) and less (MSU-1.1) compact structures.
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