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Title: Advanced *in vitro* experimental models for tissue engineering-based reconstruction of a 3D dentin/pulp complex: a literature review.

Hadjichristou Christina¹, About Imad², Koidis Petros³, Bakopoulou Athina⁴

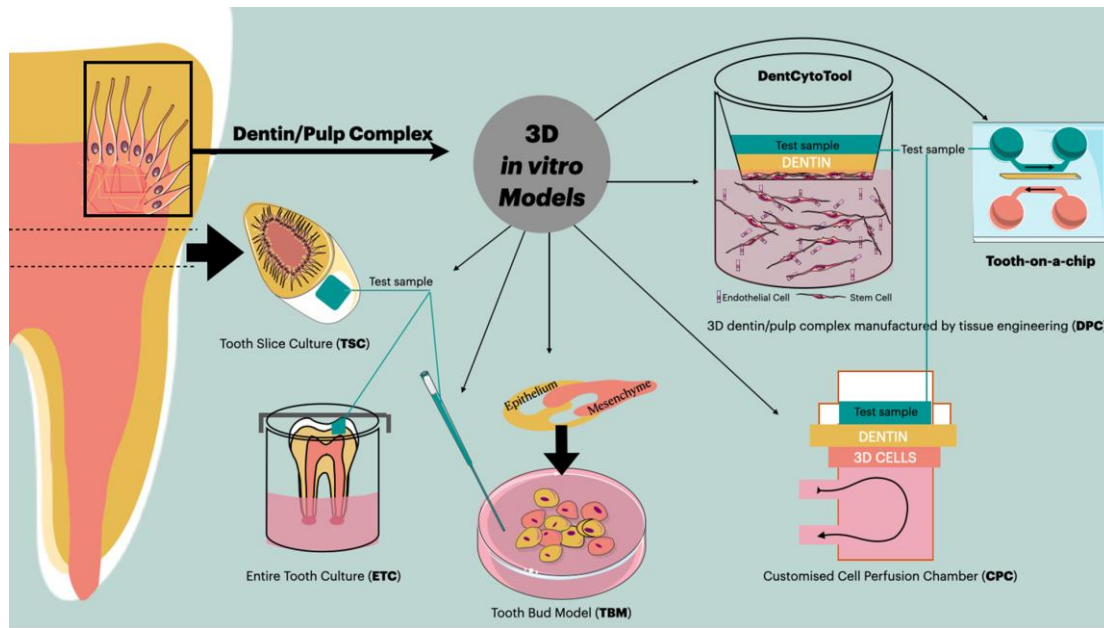
¹ Department of Prosthodontics, School of Dentistry, Faculty of Health Sciences, Aristotle University of Thessaloniki (A.U.Th), GR-54124 Thessaloniki, Greece. Electronic address: cchatzic@dent.auth.gr

² Aix Marseille University, Centre National de la Recherche Scientifique, Institute of Movement Sciences, Marseille, France. Electronic address: imad.about@univ-amu.fr

³ Department of Prosthodontics, School of Dentistry, Faculty of Health Sciences, Aristotle University of Thessaloniki (A.U.Th), GR-54124 Thessaloniki, Greece. Electronic address: pkoidis@dent.auth.gr

⁴ Department of Prosthodontics, School of Dentistry, Faculty of Health Sciences, Aristotle University of Thessaloniki (A.U.Th), GR-54124 Thessaloniki, Greece. Electronic address: abakopoulou@dent.auth.gr.

Abstract



Objective: Experimental procedures have been used to monitor cellular responses at the dentin/pulp interface. Aiming to divert from *in vivo* studies and oversimplified two-dimensional assays, three-dimensional (3D) models have been developed. This review provides an overview of existing literature, regarding 3D *in vitro* dentin/pulp reconstruction.

Material & Methods: PubMed, Scopus, Cochrane Library and Web of Science- were systematically searched for attributes between 1998-2020. The search focused on articles on the development of three-dimensional tools for the reconstruction of a dentin/pulp complex under *in vitro* conditions, which were then screened and qualitatively assessed. Article grouping according to mode of implementation, resulted in five categories: the customised cell perfusion chamber (CPC) (n=8), the tooth bud model (TBM) (n=3), the 3D dentin/pulp complex manufactured by tissue engineering (DPC) (n=6), the entire tooth culture (ETC) (n=4) and the tooth slice culture model (TSC) (n=5).

Results: A total of 26 publications, applying nine and eight substances for pulp and dentin representation respectively, were included. Natural materials and dentin components were the most widely utilized. The most diverse category was the DPC, while the CPC group was the test with the highest longevity. The most consistent categories were the ETC and TSC models, while the TBM presented as the most complete *de novo* approach.

Conclusion: All studies presented with experimental protocols with potential upgrades. Solving the limitations of each category will provide a complete *in vitro* testing and monitoring tool of dental responses to exogenous inputs.

Clinical Relevance: The 3D dentin/pulp complexes are valid supplementary tools for *in vivo* studies and clinical testing.

Keywords: dentin pulp complex, three-dimensional, tissue engineering, tissue manufacturing, regeneration, bioengineering.

Introduction

Dentin and the dental pulp are two components of the dental tissue, which function as a complex. Odontoblasts, lining the circumference of the pulp, give rise to predentin and eventually to dentin and in turn dentin -being more robust- protects the dental pulp [1]. Stimuli from the external environment are transferred from dentin to the pulp and in return the pulp responds accordingly [2]. This sensitive system, which is a structure lying within the dental cavity, is constantly exposed to acids, sugars and thermal shocks [3]. The scientific community has been occupied with experimental setups in order to recapitulate cellular mechanisms underlying certain functions or to monitor reactions to external stimuli and materials[4] . These efforts were addressed to solely pulp components, neglecting the synergistic effect of dentin and probably altering the output. With this in mind, investigations were directed in 2-dimensional experiments incorporating slices of dentin [5–8]. However, this was also troublesome, as it did not recapitulate the complexity or the structural anatomy of the natural 3D tissue. The factor of dimensionality is also degraded as cells are organized in complex three dimensional (3D) extracellular matrices (ECM) interacting with surrounding cells [9]. For these reasons, researchers have focused on developing 3D matrices to host cells, in a way that better mimics their physical environment. Three-dimensional analogues of the dentin/pulp complex include the customised cell perfusion chamber test with particular reference to the artificial pulp chamber and cell seeding in 3D polyamide meshes, instead of the initial 2D cell cultures on coverslips [10–13].

Another example is “the entire tooth culture” presented in 2005, in which extracted teeth with undeveloped roots were suspended in polystyrene well plates using metal wires to keep the roots in culture medium [14]. The

ability of these teeth to behave similarly to the clinical situation when placed in culture, (i.e. the ability of the cells to proliferate and migrate to injury sites after pulp exposure) was evaluated [14–17]. Another example is the tooth bud model illustrating organ germs representing a revival of the natal structure of the human tooth [18–20]. This attempt has been brought to life by culturing distinct mesenchymal and epithelial cell populations as representatives of pulp and dentin, respectively, in the form of *de novo* synthesis based on the embryological derivative of the complex tooth tissue.

More recently, a new advancement has been proposed in the medical field, the microfluidic devices, lined with living human cells. These small devices -microchips- that fit between the two fingers are constructed by PDMS and they replicate the microarchitecture and basic functions of human organs such as kidney [21], blood-brain barrier [22], lung [23], liver [24]. Dentistry followed this trend and devised the “tooth on a chip” [25]. This model would serve as a biocompatibility assessment tool by combining a pulp and a cavity chamber in two compartments separated by a dentin fragment. Other examples utilising 3D tissue engineering for this purpose are met through stiffness modulation, bioprinting, hanging culture inserts, and will be discussed later in detail.

For the scope of this review, studies on the *in vitro* 3D dentin/pulp complex reconstruction were identified and analysed regarding the recapitulation of the pulp component, the dentin component, the modes of the assembly of the two components, the interventions and intentions of use of these devices, as well as the biological endpoints and the actual efficacy of each method.

There is still much to learn from the *in vitro* testing, although it is accepted that this will not totally replace the *in vivo* and clinical tests prior to material market release. There are several reasons that enforce seeking alternatives to *in vivo* animal models and clinical trials. These are the limited number of test subjects as well as the limited availability of donor tissue, the high cost of purchase and maintenance as well as the ethical concerns of pain and discomfort to living subjects. Adding to this notion, the predictive value of animal models and the projection of the results to humans and donor specific factors -such as disease or genetics- may still influence the outcome [26]. As in other medical fields -such as cancer research- a bridging vehicle is sought, halfway between unrealistic and oversimplified 2D *in vitro* models and animal models[27]. Nonetheless, *in vitro* methods should be used for initial screening purposes offering high throughput information, aiding in the apprehension of cellular mechanism responses through an *in vitro* environment that closely mimics the natural tissue. The aim of this review is to reveal and analyze the studies and experimental setups which have been

implemented with a 3D *in vitro* rational with the ultimate purpose to mimic the dentin/pulp complex as closely as possible and therefore to be used as monitoring tools for the biological processes and reactions of this complex tissue to external stimuli.

Material & Methods

Study Design

All authors involved in this study developed a protocol for a qualitative review of the literature based on recommended methods. The main research question of this study was: Do *in vitro* 3D studies, regenerating the dentin/pulp complex, resemble the native tissue successfully? The review followed the PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses)-statement.

Inclusion and exclusion criteria

Literature research was performed by applying specific search terms, aiming to find publications with the purpose to device a 3D analogue of the dentin/pulp complex. Eligibility criteria included journal articles published until February 2020 in peer-reviewed journals regarding *in vitro* studies. The attributes which did not provide information for both the pulp and the dentin component as well as their assembly method were eliminated. Case reports, finite element analysis studies, review articles, and *in vivo* studies were also excluded.

Search strategy

Four scientific databases - Pubmed, Scopus and Cochrane Library and Web of Science- were used for the search. The key words for this search were: "(dentin pulp complex OR pulp dentine OR tooth) AND (3D OR three dimensional) AND (tissue engineering OR tissue manufacturing OR regeneration OR bioengineering)". The search was limited to articles written in English, published until February 2020. Additional articles from paper references were added manually.

Study selection

After removing duplicates, the full content of the resulting papers was screened manually. Inclusion and exclusion criteria were applied for the final selection. The included studies were subdivided into studies performing customised cell perfusion chamber tests, tooth bud models, 3D dentin/pulp complex manufactured by tissue engineering studies, the entire tooth culture models and the tooth slice organ culture models.

Data extraction

Two independent reviewers assessed the titles and abstracts of the collected studies and data were extracted from the full texts of the finally included studies. The data collected were: author(s) and year of study, method for pulp representation, method for dentin regeneration, assembly manner for dentin/pulp complex, intervention - aim(s), methods, biological end-points, and efficacy measurement (outcomes or main findings). Disagreements during study selection, data extraction, and quality assessment were resolved by discussion between the two reviewers.

Risk of bias and quality assessment

Studies were not evaluated for risk of bias as there is no published risk – of – bias approach for mechanistic studies or framework for rating confidence in bodies of evidence for *in vitro* studies.

Results

Search results and study characteristics

The initial search procedure using PubMed (n=812), Web of Science (n=385), Scopus (n=40), and Cochrane (n=41). Another six records were added through other sources, leading to an overall of 1284 search results. After screening titles and abstracts, the total findings remaining were 88. The excluded studies involved evaluation of tissues other than dentin/pulp complex (n= 154), review articles (n=51), *in vivo* (n=64) or clinical testing (n=54), case reports (n=31) not written in English (n=82) or irrelevant to the subject (n=763). Out of the 88 articles remaining, 42 were duplicates, and as a result, the total findings were reduced to 46 unique journal articles. Following the inclusion and exclusion criteria, 20 studies had to be excluded leading to a total of 26, which were further probed and later synthesized in this review (Figure 1).

The retrieved studies were first found in 1998. In the past 22 years, the interest in the subject has been steady as the timeline reveals, on average 4 publications per 5 years, until 2018. In the last two years, 2019-2020, 6 relevant studies have already been published, indicating an increase in the interest towards this subject.

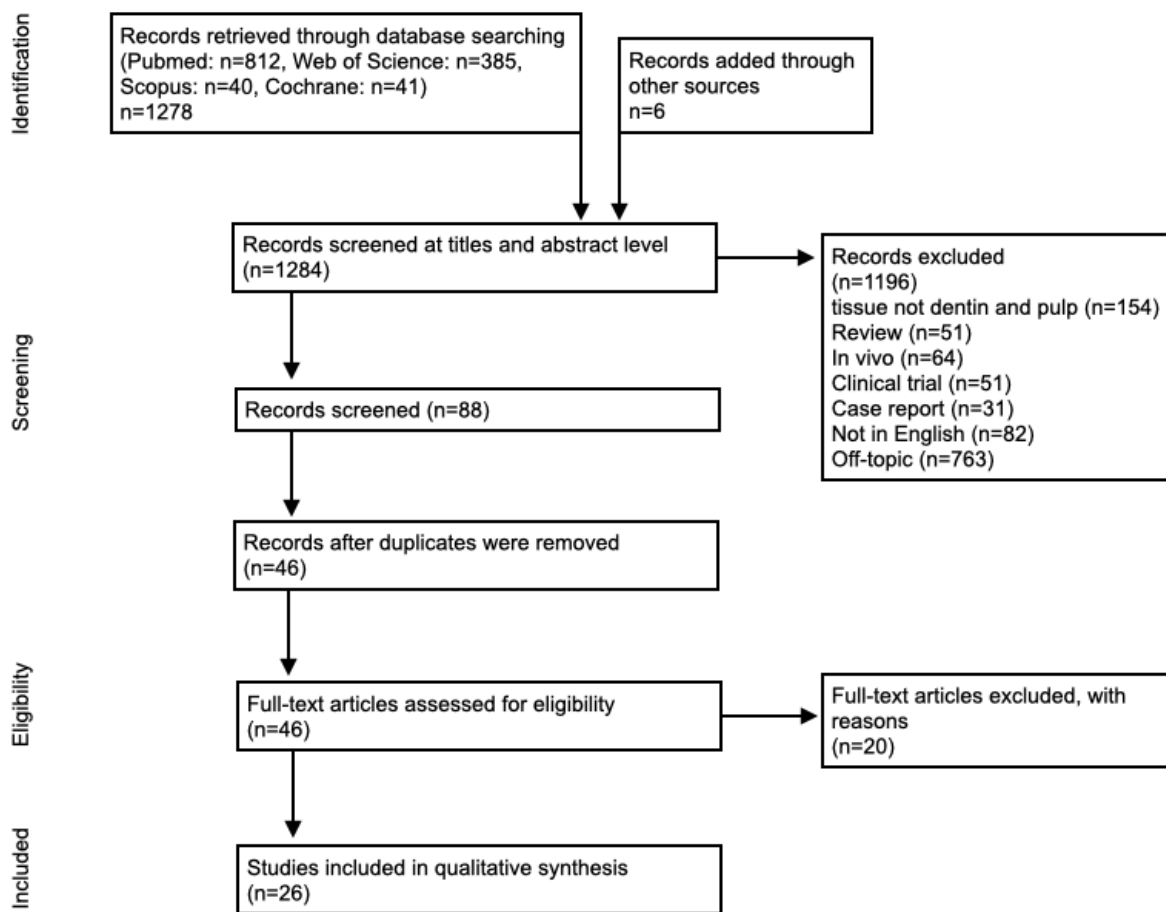


Figure 1: Flow diagram of the review process according to “Preferred Reporting items for systematic reviews and meta-Analyses: The Prisma Statement” [28]

Types of Three-dimensional dentin/pulp complexes

All of the 26 included studies were assigned to subgroups according to their defined category: customised cell perfusion chamber (n=8), tooth bud model (n=3), 3D dentin/pulp complex manufactured by tissue engineering (n=6), entire tooth culture (n=4) and the tooth slice organ culture model (n=5).

In total, nine materials were used as a 3D substrate for the representation of the dental pulp, with polyamide meshes and the natural pulp being the most frequently utilized. Other natural materials were gelatin, collagen, fibrinogen, and their combinations. Synthetic materials consisted of nylon, polyamide, polystyrene meshes and GelMa.

The dentin analogue was mostly represented by natural dentin discs, dentin fragments or cavities in dentin, of either human or bovine origin. Others employed the same gels used for manufacturing the pulp, but at higher

concentrations (e.g. higher stiffnesses of GelMa or Gelatin). A few studies used cell sheet layers of appropriate cell lines or single-cell suspensions in a 3D arrangement (Table 1). The most popular assembly method for the development of dentin/pulp complex was through a customized cell perfusion chamber. Other methods were 3D printing of the two tissues, hanging culture inserts, compartmentalized PDMS molds and direct cell sheet layering. In the case of the entire tooth culture and the tooth slice organ culture, the two tissues (dentin and the dental pulp) were naturally in contact beforehand.

Pulp analogue material	Number of studies (reference)	Dentin analogue materials	Number of studies (reference)
Nylon mesh	1 [10]	Dentin disc	16 [10,12,29–42]
Polyamide mesh	6 [12,39–43]	Dentin fragments	1 [25]
Polystyrene scaffold	1 [38]	Tooth cavities	4 [14,15,17,44]
Cell matrix	1 [18]	Single cell suspension	1 [18]
GelMa	2 [19,20]	Cell sheet layers	1 [19]
Gelatin	1 [45]	GelMA	1 [20]
Natural pulp	10 [14,15,46,17,30–34,37,44]	Gelatin	1 [45]
Fibrin/fibrinogen	2 [36,47]	Fibrinogen	1 [35]
Collagen and Fibrin	1 [35]		

Table 1: Overview of the materials used for manufacturing the three-dimensional dentin/pulp complexes.

Customised Cell perfusion chamber.

The included studies in this sector used customized cell perfusion chambers. Animal-derived cells were seeded in three-dimensional scaffolds, such as nylon meshes [10], polyamide meshes [11,12,29,40,41] or polystyrene [38]. In all studies, cells were bovine-derived fibroblasts except from Jiang et al. who used mouse fibroblasts. All studies used dentin discs as dentin analogues to separate the pulp compartment from the exposed dentin side, and the most commonly found dentin thickness was 500 μm [10,11,39,40]. Two studies tested variations of dentin thicknesses ranging from 100-1000 μm [12,38]. All dentin discs followed a similar protocol for smear removal (50% citric acid for 30'') and sterilization by autoclave (121C/25') except from Jiang et al., who used 70% ethanol for disinfection purposes. All experimental designs consisted of dynamic cultures, as perfusion was

incorporated in the setup, within the limit of 0.3-5ml/h. Lower flow values (0.3ml/h) were used for the first 24h of culture. The perfusion was then arrested during the incorporation of the test specimens and the perfusion was re-initiated for another 24h period at higher flow values. The aim of these studies was to test the cytocompatibility of dental materials via MTT assay, 24h post-exposure to: dental adhesives, resin cements, primers, resin-modified glass ionomer cements, zinc oxide eugenol, zinc phosphate cements. Outcomes of these studies indicated the importance of incorporating the dentin barrier. Indeed, the lower dentin thicknesses were insufficient in terms of protecting the cells against cytotoxic effects by various bonding agents [12,38]. The parameter of perfusion, also simulating the natural condition in the natural tooth, gave controversial results. This parameter was material-dependent as some studies revealed an increase in the cytotoxicity caused by glass ionomer cements (Vitrebond, Ketac-Fil) by increasing the perfusion rate from 0.2 to 3 or 5ml/h [10,48]. Conversely, high perfusion rates (2ml/h) caused a decrease in cytotoxicity caused by a bonding agent, Syntac, where the authors noted that this open-test system may have simulated the natural tooth more closely as these materials have not been found to exert adverse reactions *in vivo* where the material is being washed away by the bloodstream [41]. Among different bonding agents, Sengun et al. reported that the contents of each material had different cytotoxic effects. However, they were introduced in identical conditions in the cell perfusion chambers, stating that bis-GMA and UDMA containing bonding agents were more cytotoxic than those containing TEGDMA and HEMA [39]. Recently, Jiang et al. mentioned that the type of adhesive, whether total-etch or self-etch, also had an effect, with the former being more cytotoxic than the latter [38]. These parameters: dentin thickness, perfusion, the material under investigation, reinforced the importance of employing a three-dimensional system to mimic the natural situation better.

Author, year	Pulp analogue (Scaffold/cells)	Dentin analogue	Assembly	Intervention	Methods/ Biological endpoints	Efficacy measurement / outcomes
Schmalz 1999	nylon mesh/ fibroblasts	Dentin discs -(500 µm) - bovine incisors. -treatment: etching 50%	Customised Cell perfusion chamber	Insertion of dental filling materials (cements): Zinc phosphate	MTT assay 24h after exposure to sample ±perfusion	Vitrebond was cytotoxic w/o perfusion. 0.3 ml/h perfusion: no

		citric 30'' for smear removal, autoclaved (121°C, 25 min)		cement (Harvard) Zinc oxide- eugenol cement Glass ionomer cement (Ketac-fil) Resin modified glass ionomer cement (Vitrebond)	(0.3 or 5 ml/h)	significant difference from w/o perfusion. 0.3→5 ml/h significant difference (except ketac silver) but still not cytotoxic.
Schuster 2001	Polyamide meshes/bovine pulp-derived cells	Dentin discs -(500 µm) - bovine incisors. -treatment: etching 50% citric 30'' for smear removal, autoclaved (121°C, 25 min)	Customised Cell perfusion chamber	Insertion of dental filling materials (cements): Zinc phosphate cement (Harvard) Zinc oxide- eugenol cement Glass ionomer	MTT assay 24h after exposure to sample ±perfusion (0.3 or 2 ml/h)	lower survival rates at perfusion conditions, compared with static experiments. However, not statistically significant. Only Vitrebond was cytotoxic

				<p>cement (Ketac-fil, ketac-silver)</p> <p>Resin modified glass ionomer cement light cured (Vitrebond)</p>		
Schmalz 2002	Polyamide meshes/bovine pulp-derived cells	Dentin discs -(500 μm) - bovine incisors. -treatment: etching 50% citric 30'' for smear removal, autoclaved (121°C, 25 min)	Customised Cell perfusion chamber	Insertion of dental materials: All- Bond 2, Prime & Bond NT, Syntac SC, Syntac Classic, and Prompt L-Pop	MTT assay 24h after exposure to sample ±perfusion (0.3 or 2 ml/h)	Only Syntac Classic significantly decreased the cell activities. 0.3ml/h perfusion increased cytotoxicity, but 2ml/h perfusion reduced the cytotoxicity low pH dentin- bonding agents have no effect when with a

						0.5-mm dentin barrier
Galler 2005	Polyamide meshes/calf-bovine dental papilla derived cells	Dentin discs -(100, 200, 300, and 500 µm) -bovine incisors. -treatment: etching 50% citric 30'' for smear removal, autoclaved (121°C, 25 min)	Customised Cell perfusion chamber	Insertion of dentin contacting materials: Syntac Classic (Primer), Prompt L-Pop (adhesive), Vitrebond (Resin modified glass ionomer cement)	MTT assay 24h after exposure to sample, ±perfusion 0.3-2ml/h	Cell survival rates for Syntac Classic were significantly lower under perfusion conditions, no sign diff regarding dentin thickness. Vitrebond and Prompt L-Pop no statistical difference between static and perfusion conditions or different dentin thicknesses. Cytotoxicity Syntac> Vitrebond> L-Pop
Ulker 2009	Polyamide meshes/calf-bovine dental	Dentin discs -(500 µm) -bovine	Customised Cell perfusion	Insertion of composite resin cements:	MTT assay 24h after exposure to	Maxcem was similar to the negative control

	papilla-derived cells	incisors. -treatment: etching 50% citric 30'' for smear removal, autoclaved (121°C, 25 min)	chamber	Rely X, Clicker, MaxCem, Panavia F 2.0, BisCem, Bistite II DC	sample 0.3-2ml/h perfusion	group (P>.05) all other tested materials were cytotoxic
Sengun 2010	Polyamide meshes/bovine fibroblast pulp- derived cells	Dentin discs -(500 µm) - bovine incisors. -treatment: etching 50% citric 30'' for smear removal, autoclaved (121°C, 25 min)	Customised Cell perfusion chamber	Insertion of dentin bonding agents: G-Bond, Adper Prompt Self-Etch, Clearfil DC Bond System, and Quadrant University-1-Bond.	MTT assay 24h after exposure to sample 0.3-2ml/h perfusion	UB and CDCB were similar to the control group (P<0 .05). All other tested materials were cytotoxic
Schmalz 2014	Collagen hydrogel on polyamide meshes/human fibroblast - pulp derived (SV40 large T antigen transfected)	Dentin discs -(200 µm)- bovine lower anterior teeth.	Dentin barrier test	Insertion of resin-based composites and adhesive: Tetric EvoCeram, N'Durance, Clearfil SE Bond.	WST-1 cell proliferation assay. 0.3ml/h perfusion	The resin-based composites ±dental adhesive were non-toxic with values similar to the non-toxic control.

Jiang 2016	3D polystyrene scaffolds /L929 mouse fibroblasts	Dentin discs -(300, 500, 1000µm) - human molars. -treatment: 70% EtoH	Customised Cell perfusion chamber	Insertion of dental adhesives: Vitrebond, Gluma bond 5, Gluma Self Etch, Single bond Universal (self-etch)	MTT assay 24h after exposure to sample 0.3ml/h perfusion	Self-etch materials are non-toxic. Vitrebond and gluma bond 5 (for 300 µm dentin thickness) are more cytotoxic than the positive control.
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Table 2: Customised Cell perfusion chamber studies.

Tooth bud model.

For this category of experiments, a model for the developing tooth was assembled de novo using mesenchymal cells for the pulp and epithelial cells for dentin. The three articles retrieved used different techniques for the assembly. This was achieved either by encapsulating the cells in a gel drop (CellMatrix) and injecting epithelial cells adjacent to the mesenchymal cells within the CellMatrix [18], or by using GelMa and layering cell sheets of epithelial cells over cells sheets of epithelial cells [19] or by superimposing different concentrations of GelMa: 5% for the pulp analogue containing mesenchymal cells, and 3% for the dentin analogue containing epithelial cells layered on top of the 5% GelMa [20]. Most studies used osteogenic medium for the tooth bud model culture [19,20], the observation period varied from 1 day to 6 weeks, and the cells utilised were all of animal origin. The evaluation of the success of this attempt was performed mostly by histological analysis via Hematoxyline & Eosin (H&E) staining, Immunohistochemistry (IHC), immunofluorescence (IF), Polarised Light imaging (Pol) and Atomic Force Microscopy (AFM). The primary outcomes were the successful cell to cell compaction and interactions between epithelial and mesenchymal cells [18,19]. Expression of angiogenic markers, such as CD31, mesenchymal marker Vimentin, dentin markers, such as DSPP, and cell-cell marker E cadherin, were expressed by the interacting cellular populations [20]. The ultimate purpose of these attempts was to present a viable 3D method for tooth replacement through biomimetic tooth formation.

Author, year	Pulp analogue (Scaffold/cells)	Dentin analogue	Assembly	Intervention	Methods/ Biological endpoints	Efficacy measurement
Nakao 2007	Cellmatrix type I-A / Mice mesenchymal cells	Epithelial single cells injected adjacent to the mesenchymal cell aggregate	Pipetting cells- direct cell- cell contact	Organ culture for 2+14d	H&E	1 d of organ culture: -formation of a tooth germ -appropriate compartmentalization between epithelial and mesenchymal cells -cell-to-cell compaction
Monteiro 2016	5% GelMA /Porcine dental mesenchymal (DM) progenitor cells	The Dental Epithelial (DE) Cell sheet layers	Layering cell sheets (CS) of DM-DE over polymerised DM 5% GelMA	Osteogenic medium 1,4,7,12sd	H&E, IHC and IF analyses IHC (H&E, FAK, TEN and SYN4 Pol (polarized light imaging) IF (Sonic Hedgehog, BMP-2, RunX2)	<i>in vitro</i> cultured multilayered DE-DM CSs expressed appropriate tooth marker expression patterns including SHH, BMP2, RUNX2, tenascin and syndecan, which normally direct -DE-DM interactions, -DM cell condensation, and -dental cell differentiation.

Smith 2017	5% GelMA / Porcine Dental mesenchymal- HUVECs (1:1)	pDE- HUVECs (1:1) 3% GelMA	Layer ing GelM A on top of each other and photo crossl inking	preculture d <i>in vitro</i> in osteogenic media for 2 weeks→ evaluated after 1 or 6weeks	AFM (atomic Force Microscopy , H&E, IF, IHC (CD31, E-cadherin, OC, DSPP, Amelogenin)	H&E: ECM increase over time (cell differentiation) IHC: -(Ecad)-expressing pDE cells, vimentin (Vm)-expressing pDM cells, and CD31-expressing HUVEC cell populations present throughout the constructs -robust expression of the DM (DSPP) after both 1 week and 6 weeks of culture -(OC) and AM were faintly detected in 6-week.
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Table 3: Tooth bud model studies.

3D dentin/pulp complex manufactured by tissue engineering.

This group is the most diverse, with the most recent studies in terms of year of publication. The dentin/pulp complexes of this category are comprised of: spheres within rings with different stiffness values [45], slices of dentin with the preservation of the dental pulp [37], cells encapsulated in fibrin including Growth factors enhancing angiogenesis (*VEGF*) accompanied with *BMP-2* coated dentin discs [36], stacking fibrinogen - of various concentrations - in a bioprinting process within a polycaprolactone frame for dentin and the pulp [47], hanging culture inserts dividing two compartments - a dentin analogue and a pulp analogue - through a membrane with a dentin slice [35] and lastly, a three-dimensional tooth on a chip assembly with two perfusable compartments representing dentin and the dental pulp. In this category cellular populations were all of human origin, mostly DPSCs [35,36,45,47] and SCAP [25], while Hadjichristou et al. used a combination of HUVEC and SCAP for the pulp compartment and DPSCs for the dentin compartment [35]. The observation period of these experiments varied from 3 days [37] up to 4 weeks [45] and the main outcome of these studies was to evaluate the feasibility of a dentin/pulp complex implementation via mineral deposition quantification [45,47],

marker gene expression for angiogenic (*vWF*, *VEGF*, *PECAM-1*, *VEGFR-2*, *ANGPT-1*, *TIE-2*) [35–37] as well as odontogenic markers (*BSP*, *DMP-1*, *OCN* and *CBFA*, *DSPP*, *RunX2*, *BMP-2*) [35,36,47] and thyrotropin-releasing hormone degrading enzyme (*TRHDE*) and syndecan3 which are highly expressed in the natural pulp [45]. Some of the constructs were further utilised to deduce the cytotoxicity of dental materials such as eluates of resin monomers (*HEMA*, *TEGDMA*) [35] or of resin monomer (*HEMA*) and phosphoric acid [25] using techniques such as MTT, LDH and Live cell imaging. All results seemed promising in their utilisation as 3D biocompatibility investigation tools, all with potential improvements and high output capabilities.

Author , year	Pulp analogue	Dentin analogue	Assembly	Intervention	Methods/ Biological endpoints	Efficacy measurement
Qu 2015	low stiffness nanofibrous (NF) gelatin scaffold /DPSCs (sphere)	high stiffness (NF) gelatin scaffold /DPSCs (ring shape)	Mechanica l seamless assembly of the sphere within the ring	differential medium for 4 weeks	H&E, Von kossa, thyrotropin- releasing hormone degrading enzyme (TRHDE), syndecan3 (SDC3)	cell density in the central area was higher than in the peripheral area of the construct. von Kossa: mineralization in the high-stiffness area the expression of (TRHDE) and (SDC3), (highly expressed in natural pulp) were significantly higher in the central area than in the peripheral area.

Seang 2018	Natural pulp	Dentin discs (2mm thick) from human 3 rd molars	Natural slices from parts of tooth containing both dentin and pulp.	slices were cultured without (control) or with iloprost for 1 or 3 days. Serum-free DMEM was used as the culture medium in all experimenta l conditions	Confocal microscop y (live/dead staining) IHC and IF (vWF, VEGF, Col-1), H&E counterstai ning	Cells viability was unaffected at 1 and 3 days Iloprost promoted angiogenesis vWF and vEGF demonstrated a significant increase at both 1 and 3 days. Iloprost Stimulated Collagen Synthesis shown by masson's trichrome staining and Col-1 IF.
Aksel 2018	Fibrin gel +VEGF/ Human DPSCs and VEGF	BMP-2 coated demineraliz ed human dentine discs (dd- BMP-2). Dentin treatment: betadine for 30 min and 1.5% sodium	Fibrin gel- VEGF inserted into dd- BMP-2	DPSCs in the absence or presence of VEGF and BMP-2 for 28d	MTT and Live&Dea d assays ELISA for release profiles of VEGF and BMP-2 RT-qPCR analysis: angiogenic and	Higher expressions of PECAM as an angiogenic factor, and BSP, DMP-1, OCN and CBFA as odontogenic factors were observed in 3D culture model as compared to the other fg/dd combinations and the monolayer control group

		<p>hypochlorite for 10'</p> <p>Soaked in 17% EDTA/15'.</p>			<p>odontogenic differentiation markers</p>	
<p>Han 2019</p>	<p>Fibrinogen (5mg/ml) - bio-ink/human DPSCs</p>	<p>fibrinogen (20mg/ml)- bio-ink (Polycaprolactone for 3D overall shape of the tooth)</p>	<p>3D construct produced by repeated printing process with stacking.</p>	<p>cultured with odontogenic differentiation medium for 15 days</p>	<p>Live & dead staining Alamar blue Alizarin red S mRNA expression of (DMP-1) and (DSPP)</p>	<p>mineral deposition was locally observed only in the outer region, and not in the central region of the pulp tissue. This result confirmed that the designed 3D bioprinting process could induce localized odontogenic differentiation in a designed 3D space, specifically patient-specific shaped 3D dentin-pulp complexes.</p>

					SEM	
Hadjic hristou 2019	Collagen I/Fibrin hydrogel /human HUVEC/S CAP co- cultured cells	Human DPSCs on membranes of hanging culture inserts covered by a disc-shaped human treated dentin matrix (hTDM)	Mechanica l placement of the hanging culture insert (dentin analogue) in 24-well plates containing pulp analogue	resin monomers (TEGDMA/ HEMA) and Bacterial endotoxin (LPS) inserted through the hanging culture insert	MTT LDH Real time PCR for angiogene sis-related genes (<i>VEGFα</i> , <i>VEGFR-2</i> , <i>ANGPT-1</i> , <i>TIE-2</i> and <i>PECAM-1</i>)	MTT: time- and concentration-dependent increase of cell cytotoxicity was observed after application of monomers HEMA or TEGDMA \pm LPS. LDH: time- and concentration-dependent decrease of cellular viability was observed after application of monomers HEMA or TEGDMA \pm LPS. PCR: resin monomers and LPS caused significant decrease in angiogenesis related genes at 7 days of observation.

França 2019	Perfusable chamber in PDMS mold/ SCAP	500µm human dentin fragments (17% EDTA for 45'' to remove the smear layer, thoroughly rinsed with water)	Separation of pulp side and cavity side by the dentin fragment within the PDMS mold	a. 10 mM, (b) 37% phosphoric acid gel (PA) used to etch the dentin for 15 s, and (c) 35% PA plus Adper Single Bond 2 (SB) applied through the cavity side of the device.	Live-cell imaging to test cytotoxicit y (incubatio n with 50 nM of Helix NP NIR). Gelatinoly tic activity by fluoresce	Live-cell imaging: Each tested material elicited apparent cellular injury with as early as 24 h, visible reduction in cell number relative to the untreated controls Green fluorescence, indicative of MMP activity, was visible after 24 h and peaked after 48 h.
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Table 4: 3D dentin/pulp complex manufactured by tissue engineering.

Entire tooth culture.

The four studies utilising the entire tooth culture have followed a similar implementation protocol based on the first reported study by Tévelès et al. [14]. In this case, freshly extracted teeth with open apical foramens were fixed and cultured in tissue plates with their apical parts immersed in culture medium, allowing the diffusion of the medium to the dental tissues for up to 4 weeks [14]. After the efficacy of the method was sufficiently studied through histological analyses and IHC [14], additional studies followed, aiming to investigate the reaction of the pulp after the application of dental materials within cavities of the suspended teeth. Among the tested materials, the retrieved articles included Mineral Trioxide (MTA) [15,17], Calcium hydroxide [15], and Biodentine [44]. The methods of evaluation for the pulp response after the application of dental materials comprised a histological (H&E staining, Gram staining) [17,44] and an IHC evaluation for *collagen I*, *Osteonectin*, dentine

sialoprotein (*DSPP*), *nestin* [15,44]. Through the results, it was possible to show that this method -the entire tooth culture- was able to recapitulate some of the responses of the natural teeth to the same materials when applied *in vivo* [15]. It could be recognised that this is an easily reproducible method without the need to regenerate the natural tooth *de novo*, as is the case with most of the other categories in this review. It also comprises a smart tool to test dental materials in direct contact with the three-dimensional native tissue for the amount of time that the tooth can uptake nutrients from the surrounding culture medium.

Author, Year	Dentin/Pulp complex preparation	Intervention	Methods/ Biological endpoints	Efficacy measurement
Téclès 2005	Teeth were cultured in four-well-tissue plates. The crowns were fixed to the cover with sterile wax and the apical part of the teeth was dipped in the culture medium.	± cavities	Histology: H&E staining IHC: BrdU	1 day, the BrdU was localised to the nuclei of cells in the perivascular area and decreased in those away from the cavity. 2 weeks: labelled cells were seen in the vicinity of the cavity. 4 weeks: the immunolabelling was localised to the cavity area only. Control teeth without cavities or with shallow dentine cavities did not show any perivascular labelling after culture.
Téclès 2008	After cavities preparations and restorations, the teeth were fixed by the crowns to a wire and were suspended in the culture medium in 12-	Cavity was gently dried and immediately with: -Mineral Trioxide	Histology: H&E staining IHC: Collagene type I, Dentin sialoprotein, and Nestin	Histological staining after direct pulp capping with Calcium Hydroxide XR1 or MTA revealed early and progressive mineralized foci formation containing BrdU-labeled sequestered cells.

	well tissue plates, permitting the diffusion of the medium through the apex without any direct contact between the biomaterial and the medium.	Aggregate (MTA) -calcium hydroxide	BrDU	The molecular characterization of the matrix and the sequestered cells by IHC (Collagene type I, Dentin sialoprotein, and Nestin) clearly demonstrates that these areas share common characteristics of the mineralized matrix of reparative dentin formed by odontoblast-like cells. This reproduces some features of the pulp responses after applying these materials <i>in vivo</i> .
Laurent 2012	A cavity was performed ex vivo with a truncated diamond bur mounted on a high-speed hand-piece and under sterile saline cooling until the pulp exposure was obtained	Cavity was gently dried and immediately restored with Biodentine	Histology: H&E staining IHC: collagen I, Osteonectin, dentine sialoprotein, nestin	Biodentine induced mineralized foci formation early after its application. The mineralization appeared under the form of osteodentine and expressed markers of odontoblasts
Pedano 2019	The tooth crown was gently dried with a sterile cotton swab, upon which the occlusal surface was locally etched for 30'' with phosphoric acid, rinsed with distilled water	Cavity was gently dried and restored with ProRoot MTA	Histology: H&E staining and Gram-Twort bacteria staining	Histology of the teeth subjected to the pulp-capping assay showed the formation of mineralized tissue after 4-week exposure to ProRoot MTA and normal histological features in the control teeth.

	and gently dried. The crowns were hanged in the wells of a 24 plate, using steel wire and flowable composite, containing tooth-culture medium.			
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Table 5: Entire tooth culture studies.

Tooth slice organ culture.

The tooth slice organ culture, has been implemented by a research group at the university of Birmingham. This group has come up with a protocol where teeth of rat or human origin, were embedded in a semisolid medium, firstly mentioned by Sloan et al. in 1998[30]. This embedded tissue was then transferred to a Millipore membrane and was let to float on culture medium. Following a certain time period, the embedded tissue was fixed in paraffin and histologically examined. After the investigation of the performance of this method in terms of cell viability and maintenance of the morphological characteristics of the dentin/pulp complex without any exogenous additions, this model received further additions. This development consisted of the introduction of growth factors, dental materials or even an award winning tensile/compressive force testing assembly. The endpoints of evaluation were mainly through histomorphometric analyses and H&E staining [30–32,34,46] and in one occasion, through Semi-quantitative reverse transcriptase polymerase chain reaction (Sq-RT-PCR) analysis [34]. Through the results, it was possible to show that this method -the tooth slice organ culture - was able to maintain the histological characteristics of the natural dentin/pulp tissue for up to 14 days when interventions were excluded [30]. This model has proved to be of multiple uses. By the introduction of different exogenous stimuli, it was possible to monitor tissue repair responses after the introduction of growth factors [31]. The same model was also useful as an *in vitro* cytotoxicity screening tool for dental materials [46] or even as a dentin/pulp response model to orthodontic force application [34]. In this experimental setup, similar to the entire tooth culture model, the natural tissues are utilized eliminating the need to recapitulate the dentin/pulp complex from scratch.

Author, year	Dentin/Pulp complex preparation	Intervention	Methods/ Biological	Efficacy measurement / outcomes
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			endpoints	
Sloan 1998	<p>Tooth slice -(2mm)- from maxillary and mandibular rat incisors.</p> <p>The tooth slices were embedded in low melting point agar, in 96-well plates.</p> <p>When the agar was semisolid, the slices were transferred on Millipore filters, floating in DMEM in Trowel-type cultures in Petri Dishes.</p> <p>2-14 days incubation 37C/5% CO2 followed by: paraffin embedment</p>		<p>Histological examination.</p> <p>Viability assessment with acridine orange.</p> <p>Radiolabelling with proline for ECM synthesis.</p> <p>TEM (transmission electron microscopy)</p>	<p>The integrity of the dentin/pulp complex was maintained for the total experimental period (14d).</p> <p>Pulp cells and odontoblasts were vital after all culture periods.</p> <p>At 14d in culture, radiolabelling was greater than at 7d within predentin.</p> <p>TEM: cell architecture appeared to correlate well with that seen in control uncultured tissues.</p>
Sloan 1999	<p>Tooth slice -(2mm)- from rat incisors.</p> <p>The tooth slices were embedded in low melting point agar, in 96-well plates.</p>	<p>Introduction of: Agarose beads with isoforms of TGF-β 1, 2 or 3 were placed on the edge of the mineralized tissue</p>	<p>Histological examination: hematoxylin and eosin (H&E) staining.</p> <p>Morphometrical analysis:</p>	<p>At areas without agarose beads, the normal architecture of dentin/pulp complex was maintained.</p> <p>Similar to controls, TGF-β₂, caused no significant differences.</p>

	<p>When the agar was semisolid, the slices were transferred on Millipore filters, floating in DMEM in ‘Trowel-type’ cultures in Petri Dishes.</p> <p>7 days incubation 37C/5% CO2 followed by: paraffin embedment</p>	<p>corresponding to the odontoblast layer.</p>	<p>subodontoblast layer cell counting in areas with or without the agarose beads.</p>	<p>TGF-β_1 and TGF-β_3 caused similar differences such as an increased ECM secretion and subodontoblast cell density at the site of application which was observed for almost 70% of the samples. These two growth factors can stimulate dentinogenic effects in odontoblasts <i>in vitro</i>.</p>
<p>Murray 2000</p>	<p>Tooth slice -(2mm)- from maxillary and mandibular rat incisors.</p> <p>The tooth slices were embedded in low melting point agar, in 96-well plates.</p> <p>When the agar was semisolid, the slices were transferred on Millipore filters, floating in DMEM in Trowel-type cultures in Petri Dishes.</p> <p>2 and 10 days of incubation 37C/5% CO2</p>	<p>Introduction of dental materials: Salicylic acid, Calcium hydroxide, Kalzinol zinc oxide eugenol, high-mercury Amalgam, Prime&Bond, Dycal, Barium sulphate, Hypocal, Scotchbond, Calasept, Life and One-step.</p>	<p>Histological examination: H&E staining.</p> <p>Histomorphometric analysis: cell number counting within the dentin/pulp complex.</p>	<p>For samples without dental materials, the normal architecture of dentin/pulp complex was maintained for the total study period.</p> <p>The dental materials caused cytotoxicity in the following order (from highest to the lowest):</p> <p>Salicylic acid, Calcium hydroxide, Kalzinol zinc oxide eugenol, high-mercury Amalgam, Prime&Bond, Dycal, Barium sulphate, Hypocal, Scotchbond, Calasept, Life and One-step.</p>

	followed by: paraffin embedment.			These effects were only detectable on the 10 th day of experimentation.
Dobie 2002	<p>Tooth slice -(0.5mm)- from human 3rd molars.</p> <p>The tooth slices were cultured on the base of a petri dish, with the application of low melting point agar.</p> <p>7 days of incubation 37C/5% CO2 followed by: paraffin embedment.</p>	<p>Introduction of: alginate hydrogels with different concentrations of TGF-β1, \pmacid treatment, \pm TGF-β1 antibodies.</p>	<p>Histological examination: H&E staining.</p> <p>Predentin width measurement in each tooth slice.</p>	<p>For samples without TGF-β or antibodies, the normal architecture of dentin/pulp complex was maintained for the total study period.</p> <p>Acid treatment and TGF-β caused reactionary and reparative dentinogenesis.</p>
Dhopatkar 2005	<p>Specimens from mandibles of rat incisors -2mm thick sections cultured in Petri Dishes.</p> <p>5 days of incubation 37C/5% CO2 followed by: paraffin embedment.</p> <p>3 days of incubation 37C/5% CO2 followed by: RNA isolation</p>	<p>Introduction of tensile or compressive forces (50g) to specimens through springs.</p>	<p>Histological examination: H&E staining followed by cell number counting.</p> <p>Semi-quantitative reverse transcriptase PCR for gene expression analysis.</p>	<p>When no forces were applied, dental and supporting tissues maintained a healthy appearance.</p> <p>Force application induced an increase in pulp fibroblast density.</p> <p>Force application also induced upregulation of Proliferating Cell Nuclear Antigen (PCNA), c-Myc, Collagen 1α, TGF-β1 and alkaline phosphatase and</p>

				<p>downregulation of osteocalcin.</p> <p>The nature of force (tension or compression) did not alter the result.</p>
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Table 6: Tooth slice organ culture studies.

Discussion

The formation of a valid and easily reproducible 3D dentin/pulp complex holds a critical aspect in the tissue regeneration field in dentistry. Nine different materials have been devised for the representation of the dental pulp and eight for dentin tissue (Table 1). The timespan of the retrieved studies, more than 20 years, shows that this topic has continuously caught the interest of the research community. Well-established techniques that have received multiple publications, such as the perfusion chamber have been customized. New ideas have been further supported by other research groups; the methods were repeated and further validated, as was the case with the entire tooth culture. Furthermore, the field has been enriched with newly emerging ideas from different fields, e.g. 3D printing with bio-inks [47], and by incorporating state of the art with the latest updates in the research field, researchers have managed to include dentistry within these innovative ideas. Another example of this is the “tooth on a chip” which has borrowed the idea from the medical field through the “organ-on-a-chip”, finding place also in dentistry [25]. It is essential to mention that some studies sought ways to modify already existing knowledge from two-dimensional studies and bringing insight into the three-dimensional environment [35]. One of the reasons for the high demand in this subject might be the plethora of dental materials aiming to be released in the market with the need for validation and prediction of biohazards before animal testing, or even animal testing with the minimum number of sacrificed animals. The biocompatibility of dental materials and studying the reaction of dental tissues against external stimuli and the similarity to the native tissue is of utmost importance. Improving the resemblance of the *in vitro* methodology to the natural dentin/pulp complex by combining positive aspects from each of the collected experimental setups seems like a wise option to Replace the use of animal studies, Reduce the number of sacrificed animals and Refine methodology [49].

Literature search and excluded studies

After the initial search of the four databases, 1196 studies had to be eliminated. Apart from the standard exclusion criteria, such as the type of study - being a review article, clinical trials, *in vivo* studies or case reports,

not written in the English language, other significant reasons for exclusion were the following: some studies were attempting to regenerate a three-dimensional analogue of tissues other than the dentin/pulp complex, such as the periodontal bone-ligament, cementum, alveolar bone, periodontium, cartilage, salivary glands etc; other studies were focusing solely on either the dental pulp or dentin but not on the construct of both dentin and pulp; other studies, although matching the keywords for inclusion, they did not offer a detailed report on the manufacturing process of the two tissues and were therefore excluded. As stated initially, the studies that would be included should have a detailed information on the manufacturing of a three-dimensional dentin/pulp construct, therefore after removing the duplicates and the excluded studies, 21 manuscripts were finally included.

Customised cell perfusion chamber

This system has been developed in a way that many aspects mimic the natural tissue and the pulp responses to external stimuli. In defence of this statement, factors like the introduction of perfusion 0.2-5ml/h derive from literature data regarding the pulp flow, which is believed to be 20-82ml/min/100g [10]. The results, as stated by the authors are closer to those retrieved from animal experiments, agar diffusion tests and millipore filter tests and are dissimilar to those found by 2D protocols or direct contact tests when referring to resin modified Glass Ionomer Cements (RMGIC) [41,42]. Furthermore, when considering eugenol containing materials the results agree with animal model studies and studies on human teeth but not with other 2D culture tests such as the agar diffusion and the Millipore test. The simulation to the actual situation may resemble the *in vivo* tests [38-40] and authors state that the cell perfusion chamber may even be more sensitive than the *in vivo* tests [12] as the defence mechanisms exerted in the form of immune response are absent in the *in vitro* situation. The variation of dentin disc is a matter of controversy, as this affects the diffusion of monomers, finally reaching the pulp, the process of sterilisation alters the diffusion potential and the substance absorption by the dental tubules compared to the original tissue [39]. Other decisive factors regarding dentin are the thickness of this tissue [12,38,39], permeability [50], and location [39]. Meaning that thicker dentin discs provide higher protection to the underlying tissue, and number of tubules vary from the centre to the horns of the pulp and from areas proximal to the pulp to more distant locations, thus studies should focus on the consistency with which the methodology is followed [38]. Another important aspect is the timing of observation, as the early time of observation results in higher cytotoxicity values compared to later time points [39]. These studies have shown that the system is quite efficient, as many researchers conclude that there is material-dependent cytotoxicity. This depends on the

chemical nature of the introduced material [12], some constituents of the materials -as was Diphenyliodonium Chloride for Vitrebond- [38,42], some resin monomers which were more cytotoxic than others– as was bis-GMA> UDMA> TEGDMA> HEMA, and also the etching type for bonding agents - where total etch bonding agents were found to be more cytotoxic than self-etch systems [38,39]. For these reasons, this 3D dentin/pulp complex is under consideration as a strong candidate for replacement of animal models [40–42]. Worthy to mention is the fact that this test has been utilised so extensively, to a point that it has been included in the ISO 7405, for the evaluation of biocompatibility of medical devices used in dentistry [51].

Tooth bud model

The tooth bud model is a radical method proposed for the regeneration of a biomimetic organ, which will ultimately be able to function in the oral environment as an alternative to synthetic dental implants. All studies stated the importance of Dental epithelial and dental mesenchymal cell communication in the form of crosstalk between the cell lines, as well as the cellular compartmentalization in order to obtain all distinct cell tissues constituting the natural tooth, namely: enamel, dentin, root, pulp, blood vessels, bone and periodontal ligament [18–20]. The developmental stage, the cell seeding density and the substrate of encapsulation were equally important, as cells at cap stage seemed to perform better than cells at bell stage towards the development into teeth, and certain cell-scaffold combinations seemed to perform better for their intended use [18]. These models are promising as they were able to develop normal teeth with complete structures, and they could potentially serve as tools to study organogenesis and the underlying molecular mechanisms as well as the cell interactions at the Dental Epithelial-Dental Mesenchymal interface. Although these are innovative and inspirational attempts, there are still issues to solve, offering space for evolution.

3D dentin/pulp complex manufactured by tissue engineering

In this section, the implementation process was diverse, as mentioned earlier. Nonetheless, specific goals were matched for all studies. All studies mentioned the superiority of a 3D dentin/pulp complex over a two-dimensional counterpart, forming complete pulp-dentin complexes. This was vastly proven by PCR results showing an increase in dentin specific markers, such as *DSPP* and others such as *RunX2*, *ALP* [35], and angiogenic markers such as *VEGF*, *Angiopoietin-1*, and *PECAM* [35,37]. The stiffness of the scaffold forming the pulp analogue was an important factor as there was a delicate fine-tuning procedure to obtain a pulp-like analogue. Qu et al. attempted this by keeping the concentration of gelatin constant and varying the crosslinking

density [45], other by varying the concentration of fibrin and fibrinogen [35,47], while this offered control over the differentiation potential of cells into the intended cell populations. Another important characteristic was the utilisation of natural biomaterials for the pulp analogues. Gelatin (a derivative of collagen), collagen, and fibrin or fibrinogen were the major components used for this purpose. The reason for this selection was that fibrin is a biodegradable material with the ability of prevascularization [36], biocompatible with angiogenic by-products [35]. Collagen on the other hand, is a natural constituent of the dental pulp [35]. Dentin was another component of the 3D assemblies which was able to release growth factors sequestered within the matrix that were able to promote cell differentiation into odontoblasts [36] and was also able to host SCAP better than 2D monolayers [25,35] being a source of growth factors, limiting diffusion and acting as a semipermeable membrane for the introduced restorative materials [25]. Isolated points that have to be mentioned here are the real-time monitoring capability of the tooth on a chip device, which offered the potential to track cellular and subcellular responses in an *in vivo*-like environment [25], the patient-specific capability of 3D printing processes presented by Han et al.[47] offering the potential for personalised tooth - tissue engineering due to the excellent printability of fibrin and DPSCs as a bio-ink, as well as the serum free approach aided by the angiogenic incorporation of Iloprost, by Seang et al.[37], proposing a dentin/pulp slice as a viable 3D approach for the model to investigate therapeutic approaches towards dentin/pulp regeneration. All in all, these tools are introduced as candidates for risk assessment of restorative materials and monitoring tools for pulp reactions.

Entire tooth culture

Out of the five analysed categories of this paper, the entire tooth culture model is the most homogeneous in terms of implementation and the one with the maximum resemblance to the native tissue, as it uses the actual tooth in suspension from the cover of the culture dish, reflecting the *in vivo* situation. In support of this notion, the histological results, after the application of dental materials in cavities are similar to those observed in animal model studies or *in vivo* studies ([17,44]. The authors of the four retrieved studies propose that this is a valid model for the study of the early stages of dentinogenesis after inflammation, for the evaluation of responses to pulp capping material application and healing process monitoring [14,15,17,44]. So far, this has been evaluated for dental restorative materials such as MTA, calcium hydroxide, Biodentine and has proven that it is worth a spot in the *in vitro* experimental protocols. Nevertheless, it is not devoid of limitations, which are presented below.

Tooth slice organ culture

Similar to the previous category of this paper, the tooth slice organ model was also quite homogeneous in terms of implementation and resemblance to native tissue, since all attributes arose from the same research group. With time, each new publication included more refined information and details, adding valuable knowledge for other groups wishing to reproduce this experimental method. Indeed this method was able to sustain the native tissue morphology -as shown by the histomorphometric analyses- for an adequate time frame -up to 14 days- which is a valuable time span for experimentation [30]. This tool is said to be valuable for the injury and repair processes monitoring, as well as cell-matrix interactions and regulation of cell activity of the dentin/pulp tissue [30,31]. This is also the only tool that has evolved to the point that it may be used for the study of the effect of indirect external factors (tensile/compressive forces) to the dentin/pulp complex [34]. It is also mentioned as an alternative to in vivo animal cytotoxicity testing for some aspects of the exerted effects by dental materials (Murray), and generally similarities have been observed between the findings of their studies to in vitro and in vivo studies [32].

Limitations

The perfusion chamber has received multiple modifications through the years and still certain questions and concerns arise inevitably. One of these is the variation of dentin used in this model. This tissue has varying degrees of permeability, which is governed by the location from which the dentin discs have been harvested due to regional differences of the dental tubules. This means that it is so versatile that even within the same location (over the pulp chamber) there are variations between dentin covering the pulp horns and that over the area between the horns, where the former has higher permeability [38]. This could potentially influence the number of stimuli introduced in the system and, ultimately, the effect on the underlying tissue, namely the dental pulp. Another point of interest which could also alter the response compared to the natural tissue, concerning dentin, is the mode of sterilization before incorporation into cell cultures. Authors stated that autoclaving may cause protein denaturation, affecting mainly collagen, which is thermally denatured at 41 degrees Celsius and this may consequently negatively affect the organic components of the tissue [38]. Evidently, this alteration may influence the reactions that would typically occur in the natural tissue. Examples of this could be the hydroxyapatite in dentin that would naturally neutralize acids contained in dental materials under investigation, but this altered form may act differently. Also, ingredients that would normally cross-react with collagen, such as glutaraldehyde, and consequently decrease the cytotoxicity of the adhesive are able to surpass this defensive

barrier and alter the final response since collagen may be destroyed during preparation procedures [38]. Adding to the issue of defensive mechanisms, the immune reactions taking place in the living tissue are also absent in the *in vitro* arrangement, namely the inflammation and immune responses of mechanisms of repair [12]. For this reason, the *in vitro* perfusion chamber is considered to be a more sensitive system than *in vivo* models, and the results should be regarded with these limitations in mind.

As mentioned earlier, the tooth bud model is an up-and-coming and innovative model, and although progress has been achieved, authors stated that there are still issues to solve. One of these was that although Dental epithelium (DE) and Mesenchyme (DM) were seeded in a distinct manner, they appeared as mixed populations at one week; thus, at certain occasions, no distinct enamel/dentin layer was observed [20]. To address this, the authors suggested that in the future, they should apply sequential photocrosslinking so that the developing tissues will retain compartmentalization. Another suggestion was the addition of dental differentiation growth factors to engage the cellular populations into targeted differentiation directions.

Certain limitations were also stated for the category 3D dentin/pulp complex manufactured by tissue engineering. Han et al. [47] reported that in the future, their 3D printed model should be able to incorporate surrounding tissues (cementum, periodontium) for better representation of the whole tooth. The contraction of the scaffold was another limiting factor when researchers were trying to experiment with the stiffness of the scaffold. The medium and low stiffness scaffolds were presented with contractions, though this was solved by providing a supportive framework made of b-TCP (Qu 2015). On the other hand, this was beneficial as it increased the cellular differentiation potential and cell density providing maximum cell-cell contact [45]. Others stated that since this model was established to represent a cytotoxicity assessment tool, it should incorporate dental restorative materials as a whole, not only constituents of the product. Meaning that studying solely monomers contained within the resin products may neglect other confounding factors that could cause further reactions when reaching the dental pulp analogue [35]. Adding to this notion, França et al.[25] stressed the importance of adding the factor of perfusion as well as the inclusion of immune cells, functional capillaries and innervation, a microbiome and salivary flow [25]. This was not possible at the time of experimentation as it would be difficult to track all the aspects of their device at once, and this could be facilitated by built-in biosensors in further attempts [25].

Regarding the whole entire tooth culture model, although it uses the actual freshly extracted dental tissue and it closely simulates the *in vivo* situation, there are still certain limitations to it. Some of these are the limited duration of *in vitro* culture – up to four weeks - where the teeth seem to behave similarly to the living teeth

[17,44]. Further drawbacks of this model are the absence of clearance of noxious stimuli due to the absence of blood circulation, oxygen, and nutrient supply [15,17,44]. Lastly, as with many *in vitro* experiments, the inflammation reaction is not represented [44].

Limitations for the category tooth slice organ culture, were stated by the early studies [30]. This was referring to the observation of odontoblast de-polarization and detachment of odontoblasts from predentin, which was caused when mechanical trauma during the tooth sections to obtain the tooth slices.

Conclusions

To conclude, this paper gathers the evolution of 3D dentin/pulp complexes that are present in literature since 1998. This work aims to stress the increasing efforts through the years to devise a tool that may be controllable and handy for the risk assessment of newly introduced biomaterials, minimising the need for animal studies and *in vivo* tests. From the findings of these studies, it is evident that innovative ideas still arise, and certain limitations do exist, proving that there is still space for improvement and experimentation. As it is clear, it is not easy to conclude on the superiority of one category over the other as they all come with their strong points and weaknesses. The customised perfusion chamber, the tooth germ model, the different 3D dentin/pulp complexes manufactured by tissue engineering, the entire tooth culture and the tooth slice organ culture are all viable options, as long as their compromises are made clear and the developed protocols are followed closely to avoid pitfalls.

Compliance with ethical standards

Conflict of interest

The authors have no conflict of interest to declare.

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Ethical approval

This article does not contain any studies with human participants or animals.

Informed consent

For this type of study, formal consent is not required.

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