

New Composite Materials for Bone Tissue Engineering

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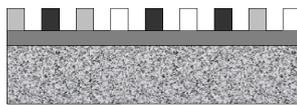
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1 Introduction

The main idea of this work was to develop new multifunctional composite scaffolds for bone cell growth and differentiation. For this purpose, a mineral macroporous support, based on ZrO₂ was used as scaffold. Water soluble aldehyde-containing copolymers of N-vinylpyrrolidone were covalently coupled to amino containing biospecific ligands and the complex was adsorbed on the matrix surface. The composite materials were tested initially in cell culture experiments.

2 Schematic approach

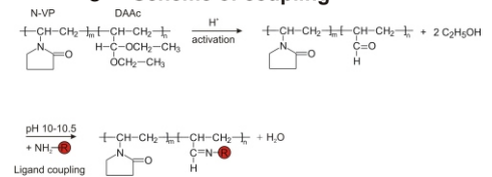


- Growth factor (BMP-2)
- Non-specific ligands (lysine derivatives)
- ▨ Biospecific ligands (RGD-peptide)

The main idea is to develop polymer-mineral composite materials with different surface ligands.



3 Scheme of coupling



Scheme of covalent coupling of amino-containing ligands to N-VP-aldehyde bearing polymer. The main advantage of chosen chemistry is the absence of toxic or non-metabolic byproducts. Moreover, the resulting Schiff's bonds are stable at physiological conditions which provide the stability of the ligand in its immobilized form.

4a Material and methods

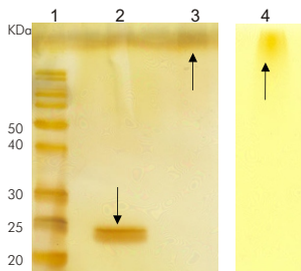
Copolymer of N-vinylpyrrolidone with acrolein diethyl acetal (N-VP-DAAc) was used as functional vector. The activation was carried out in aqueous solution at pH 2, 70-80°C, 60 min. The solution was then adjusted to pH 7. For **covalent binding**, BMP-2 or RNase (as model protein) + polymer were dissolved in sodium borate buffer, pH 10. The reaction took place at room temperature for 1 h at slight stirring. The total binding of BMP-2 to polymer was investigated by SDS-PAGE. For **adsorption**, Sponceram scaffolds were incubated in polymer solution at room temperature for 2 h at slight stirring. The concentration of adsorbed polymer was determined by analysis of the supernatant in comparison to the same non adsorbed solution. The concentration of polymer was determined using a specific iodine test.

4b Material and methods

Scaffold: monolithic macroporous Sponceram®

Cell seeding: After adsorption of polymer, the scaffolds were washed with PBS and incubated for 24 h in cell culture medium (DMEM containing 10% FCS, L-glutamine, antibiotics) at 37°C, 5% CO₂. An excess of osteoblast-like SAOS-2 cells were seeded on each scaffold in 96-well plates for 2 h at gentle stirring. Before each of the following tests the scaffolds were replaced into a new 96-well plate. Since it was not possible to estimate the attached number of cells on the scaffolds, the first measurements were performed directly after cell seeding. Cell metabolism was assayed using **MTT-test**.

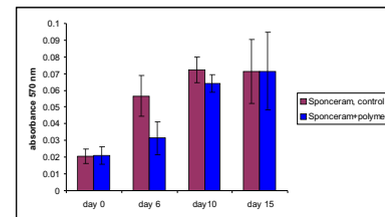
5 Gel electrophoresis



15% SDS gel, silver staining (left); iodine staining (right)
Lane 1: Marker; lane 2: BMP-2; lane 3: BMP-2 + polymer
lane 4: BMP-2 + polymer

The quality control of covalent binding was based on detection and comparison of migration distance of coupled and non-coupled BMP-2. While the charged protein migrates across the gel (lane 2), the uncharged complex does not move (lane 3). Lane 4 shows the iodine staining of the polymer (same sample as in lane 3).

6 Cell culture



Cell metabolism of SAOS-2 cells on Sponceram, Sponceram covered with polymer over a time period of 15 days. Values represent the mean of 5 samples of cultured scaffolds ± SD.

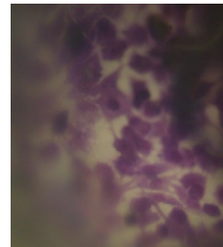
One goal was to test the modified scaffolds for their cytotoxicity. It is obvious that covering TCP and Sponceram by polymer did not influence the amount of vital cells.

7 Adsorption

Copolymer	Adsorption µg/g sorb.	Desorption		
		2 days, µg/g sorb.	6 days, µg/g sorb.	14 days µg/g sorb.
Sponceram				
N-VP-DAAc	205	10	5	0
N-VP-Ac - RNase (5:1)	240	0	0	0
N-VP-Ac - RNase (10:1)	205	0	0	0

A sonication step was added to the washing procedure to extract non-adsorbed polymer from the inner porous space of the monolithic material. There was no significant desorption of polymer measured within 14 days at physiological conditions.

8 Microscopy



Toluidin-Blue staining of SAOS-2 cells cultured on Sponceram® for 7 days.

Conclusion

The results obtained in this work clearly showed that water soluble biocompatible polymers based on N-vinylpyrrolidone are worthwhile perspectives for the development of composite scaffolds for bone tissue engineering. The copolymer of N-VP with aldehyde bearing monomer seems to be appropriate for the purpose of generating multifunctional solid surfaces triggering intensive bone cell growth.

Acknowledgement

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