Biocompatible elastomers for 3D biomaterials by additive manufacturing


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INTRODUCTION: The future vision of implants comprises individually tailored prostheses and the generation of artificial tissue and organs generated from the patient’s own cells. In order to develop artificial, biomimetic structures which perform as well as natural ones, we need fabrication processes that do not set any limits to the generation of shapes, and materials that allow for tailoring of their physical, chemical, and biological properties. We introduce new biocompatible materials for the manufacturing of flexible structures by freeform fabrication.

METHODS: We develop printable and photo-crosslinkable material systems, either based on bio-polymers derived from the native extracellular matrix, e.g. gelatin or glycosaminoglycans, or fully synthetic resins [1-3]. We use (meth)acrylation of the precursor polymers to achieve photocrosslinkability. Further chemical modifications and additives are applied in order to achieve inkjet-printable resin viscosities. Inkjet-printing combined with UV laser or two-photon-polymerization (TPP) in inert atmosphere is developed for 3D material structuring. Computational fluid dynamics are used to find optimal geometries for specific functions. Using bio-based materials inkjet-printing and 3D encapsulation of cell-laden bio-ink is performed.

RESULTS: The synthesis of polymers suitable for additive manufacturing processes was successful. Different types of non-degradable synthetic polymers were tailored to be non-cytotoxic and surface functionalization of the polymers with RGD peptide modified heparin-derivatives resulted in cell adhesion and proliferation to confluent monolayers. The material was crosslinked by UV laser and TPP and tubular systems were prepared e.g. with respect to future nutrient supply of large in vitro tissue constructs or blood vessel substitutes. The E-modulus of the crosslinked synthetic materials was adjustable. Modelling and computational fluid dynamics allowed for prediction of biomimetic bifurcations with optimized wall shear stress.

The new gelatin-based biomaterials constitute both, printable non-gelling precursor solutions and crosslinked hydrogels with tunable physico-chemical properties. Such bioinks were used for cell printing and 3D encapsulation of porcine chondrocytes.

DISCUSSION AND CONCLUSION: Our results constitute that so far individual tailoring of implants and generation of tissue substitutes by additive manufacturing remain sensitive hypotheses.


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Development of an atrial septal occluder using a biodegradable framework

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INTRODUCTION: The Solysafe® Septal Occluder has been used for closure of atrial septal defects, including patent foramen ovale, with high closure rates, excellent tissue response and few major complications. Design, and over-the-wire delivery technique provide the ability to retrieve and reposition [1]. It was voluntarily taken from the market due to isolated cases of irregularities with the metal framework. The development of a septal occluder based on the design of the established Solysafe® occluder but with a biodegradable framework presents significant new challenges compared to an occluder using a metal framework. This article outlines the challenges during development, production, sterilisation and implantation of such an occluder.

METHODS: The aim of the project was the replacement of the metal wires by a degradable material. Due to the mechanism of the device during implantation and deployment, the mechanical properties of the biodegradable material had to be equivalent or superior to those of the original material and eliminate the irregularities noted with the metal wire design. The degradable material has to keep the implant in position until it is ingrown completely. Different materials such as magnesium, degradable glass, and pure iron wires were tested without success because of the insufficient mechanical properties. First prototypes using a degradable polymer showed very promising results.

Based on the lower bending stiffness of the degradable polymers the diameter of the monofilaments had to be increased to reach the same behaviour the metal wires showed. Also, the filaments needed to be kept as thin as possible to allow the placement using a 12 F transseptal sheath.

The biodegradable material that is used for the framework degrades on the effect of hydrolysis [2] which causes a drop of inherent viscosity. Therefore, the biodegradable polymer is sensitive to water, moisture and high temperature. For this reason the monofilaments are stored under reduced pressure and the mounted implants are packaged and delivered in a sealed aluminium peel pouch filled with inert gas to prevent early degradation.

This moisture and temperature sensitivity impacts as well the process of sterilisation. Damage on the material that is caused by the sterilisation process has to be as low as possible. Thus, a major challenge was to minimize the impact on the material caused by sterilisation. Autoclaving is not possible as temperature and air humidity are too high and would destroy the material. Radiation also has a negative impact on the material since it causes an uncontrolled drop of inherent viscosity.3 Ethylene oxide and H2O2-sterilisation are proper methods for sterilisation of the degradable polymer. For this device H2O2-sterilisation was chosen as the temperature and the effect on the material are the lowest.

The continuous movement of the beating heart and the removal of the residuals by the blood flow influence the biodegradation rate. These effects have to be taken into consideration for the determination of the biodegradation time.

RESULTS: An occluder with non degradable frame was transformed into an occluder with a biodegradable frame by critical adjustments on dimensions and design. The design allows a fast and easy placement followed by an immediate closure of an artificially created defect in an animal model.

DISCUSSION & CONCLUSIONS: A septal closure device that occludes and then ‘goes away’ has long been a desirable attribute of septal devices. Our results demonstrate that:
1. a biodegradable framework can be successfully incorporated in to an intracardiac device
2. the device can be safely, securely and consistently deployed in an ASD model
3. a fully biodegradable occluder seems feasible and development of such a device should be intensified

Is increased occurrence of peri-implanted osteosarcoma associated with cast stainless steel implants?

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INTRODUCTION: Peri-implant osteosarcoma (OS) occurrence was found in dogs treated with tibial leveling osteotomy (TPLO) and the Slocum® cast stainless steel TPLO plate [1-3]. Recently, it was assumed that the metallurgical inhomogeneity of the plate surface, or the corrosion resistance of the cast stainless steel material are related factors [1-2]. Therefore, the aim of the current study was to investigate the corrosion behavior of TPLO plate surfaces from animals with and without osteosarcoma. Special attention was paid to the influence of plastic surface deformation which is the result of contouring required for plate application to the bone.

METHODS: Eighteen retrieved Slocum® TPLO plates made of 316L cast stainless steel, 9 from dogs with (CwOS) and 9 from dogs without proximal tibial osteosarcoma (CnOS) were investigated. Three retrieved plates made of forged stainless steel (Synthes) from dogs without proximal tibial osteosarcoma (FnOS) were included as examples for a different manufacturing process. On all plates visual inspections with a stereomicroscope and local micro electrochemical corrosion measurements were performed in 1M NaCl on various local spots (d = 100 µm) and corrosion resistance factors (CRF) were calculated [4]. All measurements were performed on non-deformed and on plastic deformed surface areas.

RESULTS: The implant time in-situ was comparable for both cast groups (CwOS 59±19 months vs. CnOS 52±14 months) and was shorter for FnOS with 35±5 months. Microscopic inspections of all retrievals on the cast groups showed rough surfaces, residues of sharp edged material and signs of local corrosion attacks. Furthermore, local notches and more severe tool marks were found next to the contoured regions of the plate. On the forged plates only a few marks, but no residues or microscopic signs of corrosion could be distinguished. The CRF values determined on different local surface spots showed a wide variation for the cast plates and did not vary to that extent in forged plates (Fig. 1).

Consistently, the CRF values obtained from cast plates at non-deformed locations showed higher values compared to the CRF values from plastic deformed locations. Interestingly, this effect was not observed for the surfaces of forged plates.

DISCUSSION & CONCLUSIONS: The micro electrochemical corrosion investigation showed no clear difference between local corrosion behaviors of the two groups of cast plates due to a large standard deviation. In comparison, the forged plates clearly exhibited higher corrosion resistance values throughout the various surface locations that were investigated. However, the overall number of measurements was smaller than for the cast plates. For plate adaptation to the bone shape the straight cast plates has to be bended in contrast to the anatomically pre-shaped forged plates. Our results clearly show that bending reduces the corrosion resistance of cast plates while forged plates are more unsusceptible to this alteration.

Currently, there is no clear causal connection between the surface properties of cast plates, the local CRF and the occurrence of peri-implant osteosarcoma. Nevertheless, the present results add to the speculation that osteosarcoma development might be related to local surface corrosion and the influence of that condition to neighboring bone cells.

Development of a novel glenoid implant

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INTRODUCTION: Today, the revision rate of total shoulder arthroplasty is still higher than for hip and knee arthroplasty. According to literature data, up to a third of the complications are linked to the glenoid component [1]. Cemented glenoid implants mostly fail because of wear of the bearing surface or because of cement failure. Uncemented metal-backed glenoids additionally show severe backside wear and screw- or tray breakage. To solve these problems a new glenoid implant has been developed, combining the new cross-linked, vitamin E stabilised vitamys® polyethylene for shoulder implants with the proven RM particle coating. This work introduces the concept of the new implant and summarizes the pre-clinical testing results.

METHODS: Wear behaviour of the new implant material was assessed using hip and shoulder simulators under different conditions. Furthermore push-out testing in PUR and bovine bone was used to test primary fixation, which is a prerequisite for osseointegration and long term fixation.

RESULTS: Shoulder simulator studies on vitamys® for shoulder implants showed wear reduction by a factor of three compared to standard UHMWPE (see figure 1).

Fig. 1: Shoulder simulator wear rate of UHMWPE and vitamys® glenoids versus Al2O3 heads.

Hip simulator studies on vitamys® for hip implants showed no significant change in wear behaviour after artificial ageing, even after 60 days of ageing which corresponds to 40 years in-vivo [2]. The same behaviour is expected for vitamys® for shoulder implants as the amount of vitamin E is the same for both materials.

Push-out tests in PUR foam and bovine tibial trabecular bone showed that the dovetail-like peg design can provide stable primary fixation. Push-out force in bovine bone was highly dependent on bone density and preparation method (see Fig. 2). In bovine bone with bone volume fractions similar to glenoid trabecular bone (20 – 30 %), push-out forces were between 250 and 730 N. Secondary stability will be attained due to osseointegration which is likely to start at the stable fixation pegs and move to the backside of the glenoid implant.

Fig. 2: Push-out forces of glenoid implants from primary fixation test in bovine bone blocks. Bone blocks prepared by hand or machine.

DISCUSSION & CONCLUSIONS: Combining the new vitamys® polyethylene for shoulder implants and the proven RM coating technology, the new glenoid implant was designed to reduce the amount of potential complications in total shoulder arthroplasty. Because the wear of the bearing surface is considerably lower and backside wear is eliminated, problems related to osteolysis might be avoided. Primary stability is provided by the fixation pegs and secondary stability is expected after osseointegration.


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Antibiotic loaded plaster of Paris: Wear of artificial hip joints in the presence of gypsum particles

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INTRODUCTION: Prosthetic joint infections usually require an implant removal and a treatment with antibiotics. After recovery, a new joint is implanted. This two-stage-procedure is an enormous stress, especially for elderly patients. Alternatively, beads made of plaster of Paris (gypsum) loaded with antibiotics can be implanted at the infection site [1] without removing the prosthesis. There they are slowly dissolved and antibiotics are locally released to fight the bacteria [1]. There is however concern regarding third-body-wear caused by small gypsum particles.

METHODS: Inlays made of ultra-high-molecular-weight polyethylene (UHMWPE) and cross-linked polyethylene (XLPE: vitamys®) against 28 mm CoCrMo heads and alumina pairings (36 mm, Bionit®, all n=3, from Mathys Ltd. Bettlach, CH) were tested using a hip simulator according to ISO 14242-1:2012 (Endolab, Germany). 10 g/L calcium sulphate hemihydrate (VWR, 1-100 µm particle size) was added to the standard test liquid, where the so called plaster of Paris forms the more stable dihydrate called gypsum.

RESULTS: In both cases, with and without gypsum particles in the test liquid, the wear of the inlays increased continuously (Fig. 1). In presence of the gypsum particles, the wear rates of the polymer inlays (red bars) were slightly higher than without gypsum (grey bars) (Fig. 2). The wear rates of the alumina inlays were 0.3±0.1 mg/mill. cycles both with and without gypsum. When no more gypsum was added to the metal-on-polymer articulations, the wear rates decreased (green bars). For the UHMWPE it was in the range of the reference samples while for the XLPE inlays it was still higher compared to the references. All heads and inlays showed few scratches, but there was no obvious difference between the articulations with and without gypsum particles. On the polymer inlays, additionally some pitting was observed.

Fig. 1: Wear of the hip inlays in presence of gypsum particles in the test liquid. The polymer inlays were subsequently tested without gypsum particles.

DISCUSSION & CONCLUSIONS: Neither the alumina articulations nor the CoCrMo heads were affected by the gypsum particles since gypsum is a relatively soft material. Only the much softer polymers were worn 50-70 % more, but they recovered at least partially when no more particles were added. For ceramic-on-polymer articulations and mixed ceramics even less effects by the gypsum is expected, since these heads/materials are more scratch resistant.

Thus, if the infection is treated successfully with the antibiotic loaded plaster of Paris, this is a good alternative to the 2-stage-procedure in elderly patients.


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Employing biological and mechanical cues for bone regeneration

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ABSTRACT: Bone is one of the few tissues in the human body that has the capacity for scarless regeneration. Nevertheless, complication rates of up to 10-20% are likely by means of delayed healing or non-unions [1].

Failed regeneration may be caused by biological deficits or mechanical demanding situations that impair endogenous cascades of healing. Approaches to locally deliver stimulatory factors to bone have been realized during the last decades. However, in most cases these concepts are not patient specific; they do not account for the deficits in the individual nor the specific trauma location. To be able to provide a platform for implant customization and to enable a “freedom of choice” for drug type, combinations thereof, and the applied amount, we developed intraoperatively applicable drug delivery systems (DDS). One example that is already available in the clinics is a poly(D,L-lactide) coated tibia nail (Expert Tibial Nail PROtect, DePuy Synthes) with incorporated antibiotics for a continuous release [2-5]. However, the drug itself and the location cannot be customized by such pre-coated approach. Thus intraoperative strategies are needed: Drug loaded patches that are glued to implant surfaces by rapid polymerization represent an additional strategy for implants customization (Fig. 1A) [6]. The gluing process does not impair the biological activity of the drugs within the patch, does not interfere with cellular activity or harm the mechanical stability after gluing to implant surfaces such as endoprosthetic devices.

But even if adequate local biological stimulation is available, mechanical conditions should support the healing cascade and not interfere with it. Surprisingly, a strong crosstalk between bone morphogenetic protein (BMP) signalling and mechanical tissue straining have been described (Fig. 1B). Mechanical conditions alter the effectivity of BMP proteins [7-9]. To include the mechano-sensitivity of healing in therapy concepts, a customized implant strategy has been developed to treat large bone defects. In this approach, rapid prototyping techniques are used to develop custom made scaffolds consisting of Ti beams, which meet at different angles, allowing tuning of their mechanical properties. Using computational algorithms, scaffolds were optimized to provide sufficient mechanical stimulus to tissue regeneration while avoiding implant fractures [10].

Implantable ceramic MEMS electrodes for cardiac pacemakers

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ABSTRACT: The conductive interface between the myocardial electrode and the tissue allows for transferring electron current from the pacemaker into the heart tissue. Unfortunately state of the art metallic electrodes are known to produce inflammations leading to fibrosis. The fibrosis results in increased interface resistance over time, which may eventually require an early replacement of the battery and in extreme cases the electrode. To limit the problem, the common solution is to use a steroid elution system [1]. Myocardial electrodes made by novel improved biocompatible materials like ceramics will therefore contribute to overcome these major drawbacks.

In this project the development of novel conductive ceramics are combined with advanced micromoulding techniques to study novel 3D implantable electrodes for low power, long term pacemaker applications. Most ceramics are electrically insulating in nature and produced via traditional powder route. The challenge is to develop an electrically conductive ceramic which can be moulded using today’s MEMS technologies [2].

Polymer derived ceramics (e.g. Ceraset) that exist as liquid precursors are a potential candidate material for shaping by micromoulding. The MEMS process is summarised in Fig. 1.

The first developed materials from PDCs with a modified microstructure for increased electric conductivity have been produced and are currently being evaluated by in-vitro biocompatibility tests, e.g. for cytotoxicity by the MTT test (see Fig. 2). We are currently investigating within a SNF funded project if these ceramic materials are similar in biocompatibility to zirconia and alumina ceramics.


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Evaluation of process capability of implant manufacturing

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INTRODUCTION: Many features of medical implants cannot be adequately described by the normal distribution. Consequently, capability indices based on parameter estimates assuming a normal distribution may be biased and not reflecting true process capability. Examples of features best described by distributions other than the normal include shape, surface quality, sphericity and angles, or position, coaxiality and concentricity.

METHODS: The calculation of capability indices (e.g., Cp and Cpk) is based on the location and dispersion of characteristic values with respect to a specified tolerance [1]. For example, Cp is expressed as the value of the specified tolerance divided by a measure of the length of the reference interval, usually 99.865% distribution quantile – 0.135% distribution quantile, which is equal to six standard deviations for a normal distribution.

Features where the deviation from a specified target value is expressed as an absolute value, i.e., ignoring the direction, are best described by fitting a truncated normal distribution [B1 in 2, 3] (Figure 1). Composite features are best described by fitting a Rayleigh distribution [B2 in 2].

The estimation of the parameters for the aforementioned features directly from the sample rather than from the implied distribution has recently been challenged [3]. We used packages “mam” and “VGAM” of the R software [3] to fit truncated normal and Rayleigh distributions to selected implant characteristics and compared the resulting capability indices to those obtained with a normal distribution.

RESULTS: Table 1 shows the difference between Cpk value estimates for two implant characteristics (angle and position) based on no specific distribution, on the normal distribution and on a distribution suggested to best fit the selected character [2]. Cpk values ≥ 1.33 are acceptable according to the SOP of the manufacturer who supplied the data.

Table 1. Minimum process capability index Cpk estimated for two features of an orthopedic implant with no specific distribution implied and based on two different implied distributions.

<table>
<thead>
<tr>
<th></th>
<th>Angle</th>
<th>Position</th>
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<tbody>
<tr>
<td>No distribution implied</td>
<td>0.707</td>
<td>0.860</td>
</tr>
<tr>
<td>Normal</td>
<td>0.790</td>
<td>0.833</td>
</tr>
<tr>
<td>Truncated normal</td>
<td>1.246</td>
<td>-</td>
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<tr>
<td>Rayleigh</td>
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<td>1.493</td>
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DISCUSSION & CONCLUSIONS: The capability of manufacturing processes of implants with shape, position and surface quality as key features should be estimated by using parameters from an appropriate distribution.

REFERENCES:  

ACKNOWLEDGEMENTS: The data set was kindly supplied by a manufacturer of orthopaedic implants.

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INTRODUCTION: Ceramic materials and especially oxide ceramics are well-known for their biocompatible properties. Beyond this aspect, these kinds of materials can also offer a lot of supplementary advantages when used as implants. Since several decades alumina is used as implant material for tribological pairings in Total Hip Arthroplasty. For dentistry, the use of zirconia as privileged material for the CAD/CAM applications is well recognized. More recently, composite ceramic material is also used as dental implant. METOXIT, as supplier for both the dental and orthopaedic community, has the potential to develop new ceramics which could potentially combine the advantages from both materials.

METHODS: A few years ago, METOXIT developed a new technology to obtain a ceramic coating onto ATZ Ziraldent® dental implants. This patented technology allows the production of a thin layer of porous ceramic structure for bony ingrowth, even onto complicated design such as screws. Figure 1 shows an enlarged view of this superficial porous microstructure. This successful technology called Zircapore® is documented up to 5 years in a prospective clinical survey.

The requests for a successful porous structure on orthopaedic implants are well described for metallic coatings, principally made of titanium alloy. Also, several ISO/ASTM standards are proposed to characterize the coating from a physical and mechanical point of view. The question is now: Can this Zircapore® technology be applied for a safe fixation of orthopaedic components?

RESULTS: In collaboration with EMPA in the frame of a CTI project, a new superficial porous structure is developed for an application in resurfacing implants [1].

The osteoconductive potential of the Zircapore®-like structure has been analysed in-vitro using human bone cells. Cell adhesion is assessed by scanning electron microscopy (SEM), as shown in Figure 2.

DISCUSSION & CONCLUSIONS: Oxide ceramics have optimal properties to be used as implants, e.g. for dental applications. For an application as full-ceramic, metal-free orthopaedic resurfacing implant, there is still multiple testing to be performed. However, at this stage, the manufacturing feasibility has been demonstrated.


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Phase contrast and X-ray dark field imaging: New possibilities for non-destructive testing of materials and components

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ABSTRACT: Plastics and fibre composite materials have been attracting increasing attention as materials in engineering because of their outstanding thermo-mechanical properties. In particular in medical technology plastic engineering offers a wide range of opportunities. Specific advantages of plastics and composite materials are their robustness, biocompatibility and, in particular for implants, their X-ray transparency. For plastics engineering, however, not only new manufacturing processes but also appropriate non-destructive testing and characterization tools are required.

The presentation concentrates on a novel technique that has demonstrated great potential for non-destructive testing (NDT) and non-destructive evaluation (NDE). This method uses the Talbot-Lau grating interferometer principle [1]. It enables X-ray insights extended by two additional contrast mechanisms: X-ray Phase Contrast Imaging (XPCI) and Scatter Dark Field Imaging (SDFI) - two new imaging modalities that have been developed at CSEM for NDT applications [2-4]. Remarkably, a single measurement delivers also the conventional X-ray attenuation contrast image in other words it combines three contrast mechanisms simultaneously. In Fig.1, the X-ray grating interferometer set-up at CSEM Zurich is shown.

The interferometer detects minor deflection and scattering that the X-ray beam encounters when penetrating the sample. These phenomena, though unobserved in conventional X-ray imaging, are related to relevant material properties. On the one hand, this enables improved contrast in weakly absorbing materials, such as plastics or soft tissue, and, on the other hand, it offers spatial sensitivity to the materials micro-morphology, such as porosity. Moreover, the technique allows for single projection images as well as tomographic acquisition and computed reconstruction.

Due to its sensitivity to the microscopic structure, SDFI enables the detection of a variety of microscopic properties, such as fibre orientation in fibre-reinforced materials. In addition, related defects such as cracks, delamination or fibre wrinkles can be detected. An example is shown in Fig.2 where the tomographic cross-section of a fibre-reinforced PEEK specimen is shown: The comparison of the equivalent tomographic slices reveals a domain of enhanced porosity in the SDFI which remains undetectable in the conventional CT.

The presentation will illustrate various application opportunities and examples will be given for the superior detection of characteristics and defects. In addition, a benchmark with existing NDT methods is presented.

REFERENCES:

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Development of a miniaturized hand-held navigation system for medical applications: recent results

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INTRODUCTION: A novel approach of an optical navigation and measurement system for computer-assisted surgery (CAS) is presented. The main features of the miniaturized hand-held system include clip-on abilities to a handle, instruments or a tool, an integrated optical surface scanner, and miniaturized tags attachable to the patient.

METHODS: State-of-the-art optical navigation systems for supporting CAS incorporate more than 20 years of experience. Nevertheless, there are many limitations like line-of-sight problems, clumsy optical locators, a substantial number of the required instrumentations and its logistics, time consuming registration procedures, or related complex workflows with the CAS.

The novel approach reduces or avoids many of those limitations do to a miniaturized hand-held system (Fig. 1). The system includes a stereo camera and at least one patterned tag (Fig. 1, insert). The tags are attachable to the patient. The identifications, the positions and the orientations of the tags are measured (6-dof, degrees of freedom). A clip-on mechanism permits attaching the camera to a handle, an instrument or a tool. An integrated optical surface scanner measures the surface topology of implants or body parts (Fig. 2). Presently, the prototype is wired for power and data processing. The measuring volume depends on the used optics design. For the system as shown in Fig. 1 the designed field view is about 60° and the measuring distance ranges from about 180 mm to 350 mm.

RESULTS: The position and angular accuracies (RMS) of the prototype is smaller than 50 µm and smaller than 0.5° within the measuring volume, respectively.

The miniaturized hand-held system has the potential for more economic workflows of the CAS, for quality management of the surgical procedure, for its usage for orthopaedics surgery, for new workflows for surgery of soft tissue like the liver using modified tags, or for new workflows for its usage in the field of laparoscopy.

DISCUSSION & CONCLUSIONS: The novel approach of the miniaturized optical navigation and measurement system is widely accepted by surgeons and has the potential for more economic or new CAS workflows especially towards soft tissue surgery.
Colorimetry on implant surfaces for esthetically demanding applications

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INTRODUCTION: The surface colour of implants is rarely the focus of attention. However, in certain aesthetically demanding applications, e.g. fracture plates for hand surgery or dental implants and abutments, the surface colour does become important. In patients with thin soft tissue or after bone resorption, the dark grey colour of metals, in particular titanium, results in a dark shadow leading to unsatisfactory aesthetic results. Different solutions have been proposed and are in clinical use. It was the goal of this study to compare the different surfaces and their effect on the colour of thin soft tissue.

METHODS: 12mm cpTi or ZrO\textsubscript{2} discs were used in this study. The discs were surface treated according to standard commercial processes to represent surfaces currently in clinical use. The surfaces were as follows: (1) turned, (2) polished, (3) sandblasted and acid etched, (4) pink anodisation, (5) TiNbN coated (golden) (6) ZrO\textsubscript{2}. Additionally, an experimental surface for Ti was included (7). For colour measurements (CIELAB) a commercially available camera was used (color-guide, BYK-Gardner, Germany). First, measurements were taken on the surfaces after undergoing a commercial cleaning (5 discs per surface). After that, the surfaces were covered by thin gingival sheets (0.5-1.0 mm) \cite{1} obtained from fresh pig jaws and the colour measurements repeated. Each specimen was tested on every surface (total 8 specimens). The colour of the jaws prior to resection of the tissue specimen was determined as reference.

RESULTS: The results are illustrated in Figures 1 and 2.

DISCUSSION & CONCLUSIONS: The colour values measured correlated generally very well with the optical impression. The results demonstrate clear differences between the surfaces. However, the colour value of the surface in the dry state did not always translate into a lighter or darker colour when covered with tissue; e.g. (3).

As seen clinically, ZrO\textsubscript{2} appeared lighter than the natural tissue. The pink (5) and golden (6) surfaces shifted the visible tissue colour towards the respective colour spectrum (magenta (5), yellow (6)). Further the anodised surface (5) appeared rather dark both when dry and when covered by tissue.

Our data suggest that colour measurements for aesthetically demanding implants surfaces should be conducted simulating the clinical conditions through appropriate in-vivo modelling.

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Fig. 1: CIELAB values of experimental surfaces as received.

Fig. 2: CIELAB values of experimental surfaces after coverage with gingival sheets in comparison to native gingiva.

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CVD coating technology: A coating alternative for CoCr alloys

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INTRODUCTION: One solution to avoid the CoCr particles released from the implants due to wear and corrosion is to coat a metal joint with a more wear resistant low corrosion ceramic coating. This has been proposed as for example titanium and zirconium nitride coatings on medical implants. But for the material systems such as CoCr, the problem with ion release from the substrate causing long term problems such as inflammation and aseptic loosening requires mostly dense multilayer films with higher fracture toughness. Ionbond has been pioneer of PVD multilayer coatings applied on knee systems. The relatively soft CoCr substrates theoretically require a thick hard coating in order to avoid as much as possible the egg-shell effect. Physical Vapour Deposition (PVD) coatings, evidenced to be good technology for multilayer coatings however layer interfaces would, theoretically, require higher coating deposition temperature to avoid as much as possible coherency stresses between the coating layers.

METHODS: CoCr alloys can be coated by using Chemical Vapour Deposition (CVD) technology at temperatures higher than 800°C. At such temperatures the CVD coating applied on implants with complex geometries exhibit, in all its 3D shape, rather uniform coating thickness distribution which guarantee a good ion release blockage. For the articulating surfaces the density of CVD multilayer coatings showed advantage in mechanical and wear behavior. Such advantage is due to the CVD’s higher coating thickness combined with a dual or multi dual layer deposited at high temperature leading to virtually no risk for delamination whether if in between the substrate or in between layers.

RESULTS: Femoral components are compared in terms of implantation feedback and simulator results. The coatings used for comparison are:
- CVD bilayer (top layer ceramic Al2O3)
- PVD multilayer - AS coating from Aesculap, with top layer made of a ZrN compound,
- PVD-Medthin™ 01 TiN

The coatings associated with their technologies are compared through their structure, thickness distribution, roughness and adhesion.

DISCUSSION & CONCLUSIONS: The CVD bilayer (TiCN +Al2O3) coating with its higher thickness and better thickness distribution on 3D geometry consists in a good alternative coating when compared with the PVD deposition coatings. The CVD coating is limited for application in CoCrMo since TAV alloys become brittle during coating process attributed to the high reactivity of the reaction gases with the TAV. The Ionbond CVD bilayer (TiCN +Al2O3) coating evidences interesting properties compared to other ceramics in the market such as Oxinium (ZrO2) because:
- CVD preserves the fracture toughness of the CoCrMo substrate as CVD is not a diffusion process but a coating (adding material) process.
- CVD has a superior hardness on the top layer(> 20 GPa)

Regarding the PVD coatings the feedback from the market after over one decade of implantations is excellent and suggests an increase in volume of coatings on implants.

ACKNOWLEDGEMENTS: We would like to thank for the collaboration in the characterization:
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- b. Institut de Physique de la Matière Condensée (ICMP), École polytechnique fédérale de Lausanne, CH.

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Silver as a lubricant additive for implant manufacturing

B. M. Keller 1, M. A. Gehri 1, S. Maric 2, H. Fisch 2

1 preenTec AG, Marly, CH. 2 Motorex AG, Langenthal, CH

INTRODUCTION: Implants are manufactured from a broad variety of metals e.g. titanium alloys. During the manufacturing process, metalworking fluids, so-called lubricants, are used. Today, lubricants are highly complex systems which also contain human incompatible compounds. Water based lubricants are preserved with biocides. Without proper conservation, lubricants are infected by micro-organisms within a short time. This leads to increased production costs, and potentially endangers the health of employees. Unfortunately, the use of biocides also poses problems. Employees who come into direct contact with biocides, either by inhalation of aerosols or through skin contact at their working place, may experience adverse health effects. The use of biocides results in potential endotoxin release that triggers pro-inflammatory reactions and fever. Here we demonstrate the potential of stabilizing metalworking fluids with silver components, for use in the metalworking industry.

METHODS: Through the oligodynamic effect, silver shows antibacterial and antifungal properties, even at low ppm concentrations. In the past, the incorporation of silver compounds into lubricants was limited due to the reactivity of silver e.g. oxidation and unselective reactions with sulphur containing molecules. This led to reduced long term activity of the silver and reduced stability of the lubricant. Encapsulating silver compounds into an organic matrix makes it possible to incorporate them, with good stability, into lubricants [1]. The particle size must be within the submicron scale: small enough to avoid clogging filter systems and big enough to avoid any atypical size effects characteristic of nanoparticles. The biological evaluation of medical devices is an essential step in their development. Therefore any potential risk of the devices in humans should be investigated thoroughly. Medical devices meant for use in the human body have to be tested for haemocompatibility and pyrogenic activity, since any contamination can induce pro-inflammatory reactions [2]. Residues of biocide products are known to be potential pyrogenic contaminants. Therefore the absence of biocides in the implant production chain is preferable.

RESULTS: Encapsulating silver compounds into organic matrices and then incorporating these into lubricants, results in a technically and biologically stable, nontoxic, biocompatible metalworking fluid. By varying the silver concentration into the lower ppm range, the neutralizing effect can be shifted towards a biostatic state, i.e. the working fluid bacteria fall into a so called “dormant state”. The bacterial activity, as determined by gene probes and fluorescence measurements, was far below 10% of normal state bacterial activity [3].

DISCUSSION & CONCLUSIONS: MOTOREX AG Langenthal has recently developed silver containing lubricants, which are available under the trade name of PMC “precious metal catalyst”. This new technology keeps the lubricant bio-stable without the use of common biocides. Moreover, these lubricants are non-toxic and therefore very useful for applications in the medical sector.

New opportunities for using tantalum for implants with Additive Manufacturing

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1 LayerWise NV, Heverlee, BE. 2 KU Leuven, Department of Mechanical Engineering, Leuven, BE. 3 KU Leuven, Department of Metallurgy and Materials Engineering, Leuven, BE

INTRODUCTION: Tantalum is generally known as a dense, ductile, very hard material and its high resistance to corrosion of acids. Besides a highly biocompatible material, tantalum also has a very good apposition to human bone. Bone tends to grow very close to the tantalum surface, making tantalum a perfect material for bone implants. However, due to the high density, high melting point, high cost and difficulty to machine it using conventional machining, the use of tantalum for implants is currently limited to surface coatings and coated carbon porous structures. Selective Laser Melting (SLM) is an Additive Manufacturing technology which uses a focused laser beam to subsequently melt thin layers of metal powder to create functional full dense metal parts. AM allows for almost complete freedom of design and efficient material consumption. Given the unique properties of tantalum as a material combined with the advantages of AM to overcome current issues in using tantalum, this could open up a lot of new opportunities for the use of tantalum for medical implants.

METHODS: All parts were manufactured using the Selective Laser Melting technology. In order to characterize the solid properties of tantalum processed with SLM, test samples with almost full density were tested at the test facilities of KU Leuven Department of Metallurgy and Materials Engineering.

RESULTS: Figure 1 illustrates the nearly full (99.6 ± 1 %) achieved density with a cross section view using Light Optical Microscopy (LOM) and a top view indicating the melt tracks under secondary electron microscopy (SEM).

Table 1 summarizes the mechanical properties of SLM processed test samples and compares it to the properties required by ISO 13782 ‘Unalloyed Tantalum for Surgical Applications’.

Table 1: Comparison of mechanical properties of Ta required by ISO 13782 and processed by SLM.

<table>
<thead>
<tr>
<th>Property</th>
<th>ISO 13782</th>
<th>SLM Ta</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tensile strength [MPa]</td>
<td>170-520</td>
<td>513-540</td>
</tr>
<tr>
<td>Yield strength [MPa]</td>
<td>140-345</td>
<td>463±9</td>
</tr>
<tr>
<td>Elongation [%]</td>
<td>2-30</td>
<td>29±1</td>
</tr>
<tr>
<td>Young’s modulus [GPa]</td>
<td>n.a.</td>
<td>168±8</td>
</tr>
<tr>
<td>Hardness [HV]</td>
<td>n.a.</td>
<td>207±2</td>
</tr>
</tbody>
</table>

DISCUSSION & CONCLUSIONS: From the results presented above it can be stated that the mechanical properties of tantalum parts processed using Selective Laser Melting achieve the requirements of ISO 13782. Furthermore, processing conditions and their influence on the compression yield strength have been investigated. SLM therefore seems to be a novel method to manufacture tantalum implants. In a next step, porous tantalum scaffolds are being investigated. SLM allows manufacturing complex porous structures which enforces bone ingrowth and prevents stress shielding. In this continued research the static and dynamic mechanical properties of a defined lattice structure will be evaluated under compression. Also, the in vitro and in vivo behaviour of the SLM tantalum scaffolds will be investigated.


ACKNOWLEDGEMENTS: Research funded by the Agency of Innovation by Science and Technology (IWT).

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Breakthroughs in patient matched medicine through advancements in 3D image analysis

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INTRODUCTION: Today, with the increasing use of 3D medical imaging (e.g. CT or MRI), it is possible to analyse in detail 3D patient anatomy to extract personalized features and measurements for designing patient matched devices, but also to quantify trends and population specific shape information for standard implant design.

METHODS: In order to create a patient matched implant or surgical instrument, a medical image based workflow is performed (Fig. 1). Starting from a CT or MRI scan of the patient, a group of voxels that belongs to the anatomy of interest is selected in a process called segmentation. From the segmented anatomy an accurate 3D triangulated surface model is created. On this anatomical model, complex 3D measurements or a virtual surgery can be performed. Based on this analysis and planning, a patient matching implant and/or surgical instrument is then designed. Finite element analysis (FEA) can be performed to simulate the loading conditions on the implant during the patient’s daily activities and to optimize the design. Patient specific implants today are manufactured via milling or to an increasing extent via additive manufacturing (3D printing).

Medical image based engineering is also used to speed-up the design process of standard implants. When personalized measurements are analysed on a large number of patient cases, population analyses can be performed in order to extract trends and population-specific shape information. By using statistical shape modelling [1], it is possible to determine the average shape and the variation across the population (Fig. 2).

RESULTS: Personalized analysis and surgical simulation allow for the design of custom implants that have shown to result in better surgical outcomes, and sometimes to be the only solution for extreme patient cases, such as for selected hip revision cases. A medical image based approach also allows for the design of patient-specific guides and is nowadays provided by the biggest implant companies, such as Biomet or Zimmer, for total knee replacement.

Image based population analysis have been used successfully to design standard implants that better fit the population. Using the average anatomy as the starting point, it is now possible to reduce the number of cadaver trails, thereby reducing R&D cost and reducing the time to market for a new device.

DISCUSSION & CONCLUSIONS:

Advanced anatomical image processing, analysis, design and modelling software enables efficient production of high quality patient matched implants and surgical instruments.

In many cases, a standard implant still provides a sufficient solution. Advanced statistical modelling of medical image data makes it possible to faster and cheaper develop implants that better fit the population.

Patient specific implants in orthopaedics – Efficient production with software

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INTRODUCTION: Patient specific instruments and patient specific implants are used more frequently in Orthopedics. The advantage of patient specific technology is shorter operation time and reduced overall instruments use [1].

The process for the production of patient specific instruments or implants requires the precise segmentation of the obtained image data. This segmentation causes significant problems to orthopedic manufacturers when the amount of cases increases: It is a slow and human intense work and therefore not well scalable. Furthermore it is an error prone and quality wise not well controllable work.

Medivation in conjunction with the Institute of Medical and Analytical Technologies in Muttenz has developed an own software technology called Auto-3D to segment the image data mostly automated and with as least human interaction as possible. The aim of this study was to analyze the efficiency and precision of the newly developed segmentation tools.

METHODS: The software engine Auto-3D segments the CT data by first automatically initializing an anatomical shape based model. Then the software applies a fine-segmentation algorithm in defined regions.

Once this automatic process is finished, the software then guides the operator through a manual process of checking the automatic work in a step by step process. Segmentation errors can be manually corrected.

Then the software exports the surface models with accurate precision for further processing.

In this study we used 20 reference models that were manually segmented with a standard image processing software (Mimics, Materialise, Belgium) by an experienced operator. The output of the Auto-3D software was checked against these 20 reference models in terms of a) overall precision of the anatomical shape based model, b) automatic segmentation process, c) manual fine-adjustments. Furthermore, the time was compared between the segmentation of the reference model and the final result with the software engine.

RESULTS: Table 1 summarizes the time measurements. The mean time used to create the reference models was 38.0 minutes with manual segmentation (min 28...max 60) versus the mean time of 17.6 minutes (min 8...max 30) with the Auto-3D segmentation software which is highly significant (p<0.001).

Table 2 summarizes the difference between the reference models vs the different stages of the Auto-3D segmentation process. It can be seen, that after applying the automatic segmentation process, the precision (mean surface deviation) is already within 0.8mm.

Table 1. Time comparison between manual segmentation and the software Auto-3D.

<table>
<thead>
<tr>
<th></th>
<th>Conventional Software [minutes]</th>
<th>Auto-3D Engine [minutes]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time</td>
<td>38.0</td>
<td>17.6</td>
</tr>
<tr>
<td>Min</td>
<td>28</td>
<td>8</td>
</tr>
<tr>
<td>Max</td>
<td>60</td>
<td>30</td>
</tr>
</tbody>
</table>

Table 2. Precision comparison between manual segmentation and the software Auto-3D. Stage 1 is the anatomic shape based model, stage 2 is after automatic segmentation and stage 3 is after manual fine-editing. All values in mm.

<table>
<thead>
<tr>
<th></th>
<th>Stage 1</th>
<th>Stage 2</th>
<th>Stage 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Femur</td>
<td>1.5</td>
<td>0.8</td>
<td>0.25</td>
</tr>
<tr>
<td>Tibia</td>
<td>1.4</td>
<td>0.8</td>
<td>0.24</td>
</tr>
</tbody>
</table>

DISCUSSION & CONCLUSION: The Auto-3D software improves the time significantly. The precision reached was similar to manual editing. The overall process quality due to the standardized process was not measured in this study but is another obvious advantage of this approach.

The software is today in regular use and is recognized by more and more manufacturers.


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Regulatory approval of implants produced with Additive Manufacturing

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INTRODUCTION: Additive Manufacturing (AM) is already in full use for the production of CE-certified and FDA-cleared orthopaedic implants, with more than 30,000 EBM-manufactured acetabular cups with integrated trabecular structures for improved osseointegration implanted to date. This rapid development has caused a heightened interest for AM within the implant manufacturing industry in general, and subsequently an increased focus on how to obtain regulatory approval for implants produced with additive technologies.

In this paper we will discuss Arcam’s experiences from how the implant manufacturers who have implemented AM in their production have gone about obtaining CE certification and FDA clearance for their EBM-manufactured orthopaedic and spinal implants.

BACKGROUND: Electron Beam Melting (EBM) technology manufactures parts by melting thin layers of metal powder. The energy source is an electron beam gun, and the process takes place in a vacuum chamber. The technology’s additive, layer-based nature also enables the production of implants with the integrated trabecular structures that enhance the osseointegration.

For most implant applications AM is considered a special process – a process for which the product cannot be fully verified and consequently the process needs to be validated. Validating an AM-based manufacturing process requires extra vigilance on three levels.

Firstly, we need to understand the technical differences between additive and conventional manufacturing. Conventional manufacturing is based on removing material until the final part shape has been obtained. In AM material is added layer-by-layer to obtain the final shape which means that the material properties of the part are generated as the part is being built.

Secondly, AM technologies generically rely on a stable process for each layer. For EBM specifically, achieving a stable process means obtaining a thermal steady-state. As the heat balance depends on the cross-section of your part the parameters controlling the process may have to be adjusted to handle different geometries – they become geometry-dependent.

The third point of vigilance concerns the supply of raw material. With AM only the material that is actually needed to produce the implant is used. The remaining powder metal is recycled and reused. Hence, powder handling procedures must ensure rigorous quality control and full traceability of each part back to its source powder.

VALIDATION GUIDELINES: AM in general and the Arcam EBM process in particular (although widely used for production of implants and despite a large number of already certified medical devices) are still relatively new to most implant producers. To facilitate the validation process, Arcam has developed a validation template with Arcam’s view on each aspect of FDA and ISO validation guidelines. Each of these guidelines is commented in detail and links are provided to relevant Arcam documentation, such as drawings, manuals, calibration reports and software documentation.

RESULTS: In 2007, the two Italian orthopaedic OEMs Adler Ortho and Lima were awarded CE-certificate for their acetabular cups featuring integrated porous structures. Under the trademarks of Fixa Ti-Por™ and DELTA TT™ these cups are now in volume production at Adler Ortho and Lima-Lto. In 2010 Exactech introduced the InteGrip™ cup and became the first U.S. manufacturer to offer FDA-cleared orthopaedic implants.

To date over 30,000 EBM-manufactured cups with integrated trabecular structures have been implanted, and approximately 2% of the global production of acetabular cups is now manufactured with EBM.
Fixation concept for a metalback/ceramic connection

M. Schmidt

Jossi Orthopedics Ltd, Islikon, CH

INTRODUCTION: "However, the metal-on-metal system generated adverse reactions in patients that are typical of all devices manufactured in this way."1 This type of generalising statements is often communicated since two market leading orthopaedic companies recalled their metal-on-metal (MoM) hip resurfacing implants in 2008 and 2010. Instead of enquiring on the reasons for increased clinical failure, no distinction seems to be made between design and material, and MoM devices are generally dismissed. Nevertheless, the concept of hip resurfacing2 is appreciated by surgeons and patients, thus creating a need for alternatives. From all accepted materials, only ceramic-on-ceramic (CoC) combinations are suited. Jossi Orthopedics’ interest is to produce the titanium alloy metalback by its unique HybridManufacturing™ process, combining deep-drawing and precision machining.

EVALUATION: When looking for an alternative to MoM, a thorough evaluation is necessary to decide whether concept, material, design, or clinical handling led to failure of resurfacing devices. It turned out that both withdrawn products had design weaknesses, one acetabular component badly osseointegrating3, the other MoM combination having too little clearance between head and cup, leading to increased friction and ultimately component loosening4. Since resurfacing needs large-diameter ball-heads and thin-walled acetabular cups, polyethylene is to be excluded for tribological reasons. Thus, only CoC is left to create the artificial joint components. Accepted combinations are Al-Al, ZTA-ZTA (zircona-toughened alumina) and ATZ-ATZ (alumina-toughened zirconia). On the acetabular side, the ceramic sliding surface is embedded in a titanium alloy metalback (Fig. 1) enabling cementless fixation and osseointegration, whereas a thin-walled ceramic ball-head is either cemented directly onto the femoral neck or a standard ball-head is fixed to a femoral stem.

RESULTS: Existing CoC resurfacing solutions are thoroughly protected by patents. The biggest obstacle therefore was not only to circumvent existing patents but to set an own IP protection. The crucial point was the connection between metalback and ceramic sliding surface. Having mastered this challenge, a setting instrument was developed, enabling a strong grip on the cup without interfering with the acetabular rim during impaction. Being solely a supplier to orthopaedic companies, i.e. without own branded products, the goal of Jossi Orthopedics always is to offer a complete solution to potential customers. The metalback/ceramic assembly then was presented to orthopaedic companies, however, in times of cost restrictions, it is rather preferred to buy a customised standard solution. Jossi's goal therefore is to collaborate with a ceramics producer.

Fig. 1: Metalback/ceramic assembly CeraGrip™, consisting of a thin-walled acetabular component (left), shown with femoral component for resurfacing (right). The ATZ ceramic components were provided by Metoxit, Thayngen, Switzerland.

DISCUSSION & CONCLUSIONS: Even being, as a supplier, only interested to produce a specific component, a complete solution should be targeted, and a strategic design feature should be IP protected. In the presented example, the crucial technological development is the connection between metal and ceramic, leading to an "invisible generic" solution. The titanium alloy metalback can be fully customised, e.g. by a proprietary porous coating, creating a distinct orthopaedic implant, however, with reduced time to market for the orthopaedic company.

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3www.orthoinfo.aaos.org/topic.cfm?topic=A00355

http://www.ecmjournal.org
Contemporary dental implant systems: Clinical, biological and mechanical aspects

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Zahnmedizin Zürich-Nord, Zurich, CH

ABSTRACT: Titanium has been established as the material of choice for endosseous implants resulting in a high degree of predictability. In recent decades, the clinical replacement of natural teeth by osseointegrated implants has represented one of the most significant advances in restorative dentistry. Since then, numerous studies of various clinical indications have documented high survival and success rates with respect to specific criteria [1]. Persisting osseointegration and restorations of good function have been key elements for the patient’s subjective satisfaction, which is one criterion. One should never stop to have a critical and objective look even at successful concepts.

Many types of implants require trans-mucosal abutments to retain implant restorations. The implant-abutment-interface (IAI) comes more and more in the focus of the present research. Modern IAI can be divided into three groups: The internal conical, the flat-to-flat and the tube-in-tube IAI. It is shown that this micro movement causes bacterial leakages, micro-pump effects in the sulcus and unintentional elastic deformations of the implant components. It is a clinical observation, that many IAI are not sealed and bacterial colonization is clearly visible in the implant and on the abutment screw after abutment removal. Unpublished data shows micro-particles as result of fretting in the IAI.

The data presented is indispensable in order to evaluate IAI, to optimize them and to bring the results in correlation with the clinical results. The avoidance of abutment loosening, surface abrasion, fractures and the formation of bacterial colonies has not really been adequately considered in the construction of most IAI.

Implant manufacturers should aim to reduce the micro-mobility by increasing the stability of the implant-abutment interface. Therefore, reducing the mobility of this connection by constructing physically tight connections with a high level of precision in the sub-micrometer range is considered to be an important precondition for microleakage prevention. With the current technology, micro-movements that are conditioned by chewing forces can only be avoided in IAI if they have a conical self-locking effect.

101 million cycle simulator wear characterization of diamond like carbon coated CoCrMo articulating implants

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1 Empa, Swiss Federal Laboratories for Materials Science and Technology, Dübendorf, CH.
2 DePuy Synthes, Dübendorf, CH

INTRODUCTION: Diamond like carbon (DLC) coatings have been proven to be an excellent choice for wear reduction in many technical applications. However, for successful adaption to the MedTech field, layer performance, stability and adhesion in realistic physiological setups are very important and not consistently investigated [1]. Simulator testing as well as corrosion tests are of great importance to verify the long term stability of such a DLC coated articulating implant in the human body. Commonly one million cycles of simulator testing correspond to 1 year of articulation in the human body.

![DLC layer on CoCrMo implant](image)

Fig. 1: Cross-section of a 4 µm thick DLC layer on a CoCrMo implant after 101 million cycles in a spinal disk simulator.

METHODS: Diamond like carbon coatings were deposited on CoCrMo biomedical implant alloy using a plasma-activated chemical vapor deposition (PACVD) process. As an adhesion promoting interlayer tantalum films were deposited using magnetron sputtering. The implants were mounted in a spinal disk simulator where they underwent more than 101 million cycles. Within this time these implants were characterized by high wear resistance, low friction coefficients, high corrosion resistance and low defect growth. These results were obtained by means of optical microscopy, SEM/EDX, FIB cross section and profilometry. The coatings were further analysed using XRD and XPS.

RESULTS: It is shown that metal-on-metal (MoM) pairs perform well up to 5 million loading cycles, after which they start to generate wear volumes in excess of 20 times those of DLC-coated implants [2]. This is attributed to the slight roughening observed on unprotected metal surfaces as usually also observed in-vivo. The DLC on DLC inlay pairs show comparable low volume losses throughout the full testing cycle (up to 101 million cycles over a period of three years and two months).

DISCUSSION & CONCLUSIONS: To our knowledge this is the first time a simulator test of a DLC-coated articulating implant running for more than 100 million cycles (corresponding to over 100 years of articulation in-vivo) is presented.


ACKNOWLEDGEMENTS: Financial support provided by the Swiss innovation promotion agency CTI, and CCMX is gratefully acknowledged.
Smart NiTi constructs for 3D cell culture applications

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INTRODUCTION: NiTi shape memory alloys have unique mechanical and physicochemical properties that are appealing for a wide variety of biomedical applications. Selective laser melting (SLM) is a versatile method to create porous scaffolds using computer aided design (CAD) [1]. With the ultimate goal of fabricating complex 3D NiTi implants for orthopaedic & dental applications, we validated the utilization of SLM-based NiTi constructs as scaffolds for Tissue Engineering applications. Targeting the beneficial properties, i.e. pseudoelasticity or the one- and two-way shape memory effect, NiTi scaffolds might be used as mechanically active implants stimulating the surrounding tissue and thereby assisting bone healing.

METHODS: We assessed the biocompatibility of NiTi scaffolds as well as the adhesion and proliferation of human bone marrow-derived mesenchymal stromal cells (hBMSC) and MG-63 osteosarcoma cells. The cells were cultured on rapid prototyped (RP) NiTi constructs both on two-dimensional disks and three-dimensional scaffolds. Cell adhesion on constructs was assessed both using SEM and confocal laser scanning microscopy (CLSM). Proliferation rates were assessed using the CyQUANT® Cell Proliferation Assay to determine cell numbers for several points in time.

RESULTS: Both cell types did not exhibit a cytotoxic effect cultured in extracts of NiTi constructs (data not shown). Following, MG-63 cells and hBMSC were seeded on 2D disks and 3D NiTi scaffolds revealing high colonization densities (Figure 1). Additionally, long-term cultures up to 21 days were performed in order to investigate hBMSC proliferation capacity cultured on NiTi constructs. hBMSC do proliferate on NiTi constructs with similar growth rates as on tissue culture plastic (TCP) (table 1) demonstrating that SLM-NiTi disks are permissive to cell proliferation. hBMSC indicate similar sprouting and cell adhesion behaviour as observed on TCP being the gold standard in vitro culture system. These findings indicate the high biocompatibility of NiTi constructs facilitating their further utilization as cell culture substrate.

DISCUSSION & CONCLUSIONS: The results demonstrate SLM-NiTi construct biocompatibility and underline their possible utilization as implant and/or scaffolding material exhibiting high colonization (adhesion and proliferation) capacities. Taking their unique shape memory properties into account, SLM-NiTi constructs could lead to personalized implants which allow for colonization and differentiation of host progenitors and at the same time might provide a unique platform to create active and thus smart implants.


ACKNOWLEDGEMENTS: The authors gratefully acknowledge the financial funding of the Swiss National Science Foundation within the research program NRP 62 “Smart Materials”.

Table 1. hBMSC growth rates [doublings/day].

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<thead>
<tr>
<th></th>
<th>NiTi 2D disk</th>
<th>TCP</th>
</tr>
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<tbody>
<tr>
<td>Growth Rates</td>
<td>0.170 ± 0.021</td>
<td>0.170 ± 0.033</td>
</tr>
</tbody>
</table>

Fig. 1: Left: CLSM image of MG-63 cells cultured on a 3D NiTi scaffold (scale 3.4 mm x 3.4 mm). Right: SEM image of hBMSC cultured on 2D NiTi disk for 11 days.
Effect of ultrasound on the electrochemical deposition of antibacterial copper particles on anodized titanium implant surfaces

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INTRODUCTION: Bacterial infections taking place immediately or years after orthopaedic, trauma or dental surgeries cause serious problems for the patients. Implant surfaces exhibiting antibacterial properties preventing infections are therefore highly desired. We have recently established an electrochemical method to deposit the antimicrobial agent copper on the rough, fine-porous surface of spark anodized titanium samples. The antibacterial effect and the copper release rate were demonstrated [1]. Copper was deposited as clusters of different sizes, forms and surface distribution [1]. In this study we demonstrate how the surface distribution of the copper deposits can be affected by ultrasound applied during the electrochemical process.

METHODS: Mechanically pre-treated and ultrasonically cleaned discs of cp Ti (grade 4, Ø 14 mm, 1.5 mm thick) were anodized according to the spark-assisted anodizing (SAA) method [2] to produce a rough, fine-porous surface. Copper was electrochemically deposited using proprietary electrolytes and process parameters. During the deposition process ultrasound was applied with a frequency either of 27 kHz or 80 kHz and a power of 350 W each. Deposition studies were performed for different copper concentrations in the electrolyte and different deposition times. The copper deposits were characterized by SEM/EDX.

RESULTS: In the absence of ultrasound, copper is deposited on the fine-porous oxide layer of the anodized samples as large clusters of nanometer-sized copper particles and with inhomogeneous surface distribution (Fig. 1 left). When ultrasound is applied, the copper deposits are much smaller (size of few nanometers) and homogeneously distributed over the sample surface (Fig. 1 middle). Compared to the experiments without ultrasound the amount of deposited copper is significantly increased for all deposition times when 27 kHz is applied (Fig. 2). However, for 80 kHz, less copper is deposited after longer deposition time (120 s) and the copper deposits are slightly larger than for 27 kHz (Fig. 1 right).

DISCUSSION & CONCLUSIONS: Application of ultrasound during electro-chemical deposition of copper leads to a more homogeneous Cu allocation, to a frequency dependent deposit distribution and to an increased amount of deposited copper. It is suggested that the different size of the cavitation bubbles and acoustic streaming velocities for 27 kHz and 80 kHz may explain the different results [3].

Additive manufactured ceramic/polymer scaffolds

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INTRODUCTION: Tailor-made ceramic scaffolds in combination with the corresponding surface chemistry and biology is of great importance for successful implantation and guarantees a rapid osseointegration. This project investigates the fabrication of scaffolds with defined macro porosity by means of powder based 3D-printing. To reach mechanically stable structures, various post-processing methods like infiltration or thermal treatment have been applied.

METHODS: Two fabrication approaches were developed. On one hand cylindrical specimens (Ø 10 x 10 mm) were printed as a composite (hydroxyapatite/polymer). On the other hand printed hydroxyapatite (HA) specimens with open porosity were infiltrated with polymers after sintering. All structures were printed with acidic binder solution. Composites with HA as bulk material were blended either with 20 wt.-% or 30 wt.-% collagen, polycaprolactone (PCL) and chitosan. Specimens containing chitosan were post-hardened in acidic medium. Infiltration was performed with 4 wt.-% polyvinylalcohol (PVA), 3.6 wt.-% PCL and 10 wt.-% gelatine. The mechanical properties were determined by compressive strength (CS) measurements. Further analysis was performed by porosity measurement (Archimedes) and scanning electron microscope (SEM).

RESULTS: Cylinders with chitosan showed the highest CS values among all other composites compared to printed and sintered HA cylinders. Figure 1 illustrates the consistent distribution of a HA and polymer within the composite matrix. The porosity of all composite specimens decreased around 20 % compared to sintered HA samples.

DISCUSSION & CONCLUSIONS: In previous publications, CS of 3D-printed composites with other bio polymers are reported in a range of 0.5 - 5 MPa [1-2]. This emphasizes the potential of printed chitosan composites with their 16.3 MPa CS (Figure 2). In addition, the developed infiltration method led to a 4.5x higher compressive strength. However, the insolubility of chitosan, collagen and gelatine specimens needs to be further improved to ensure mechanical stability in body like environments.


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Macroporous titanium coating for challenging substrates

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INTRODUCTION: Macroporous metal structures obtained by thermal plasma spray (e.g. TiGrowth®) are becoming a reliable alternative strategy to reach a tough and time resistant bone fixation in metal joint replacement components [1]. Scope of this study is to highlight the possibility to extend the application of thermal sprayed titanium spongy surfaces to those non metal substrates that show limited osseointegrative potential [2].

MATERIALS & METHODS: By Modified Vacuum Plasma Spray (MVPS) a macroporous thick titanium coating (TiGrowth®) was applied either onto a Ti-6Al-4V or onto a polymeric substrate (PEEK Motis®, Invibio, UK). MVPS is a lower temperature process than conventional VPS thus more respectful of the substrate properties [3].

The morphology of macroporous surfaces was characterized by Scanning Electron Microscopy (SEM). Porosity and thickness were measured through optical micrographic analysis. Mechanical tests were performed for evaluation of coating to substrate adhesion according to ASTM F 1147.

In n=6 sheep, 36 rods of PEEK Motis®, with and without TiGrowth® coating, were randomly implanted in the right and left wing of the pelvis. Animals were allocated to 2 and 12 weeks observation periods (each n=3). 12 samples of each surface were used for a pull out test. 6 samples were used for a histologic analysis.

RESULTS: Mechanical tests: Adhesion test results are summarized in Table 1.

<table>
<thead>
<tr>
<th>Material</th>
<th>Adhesion (MPa)</th>
<th>Std. Dev. (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TiGrowth® on Ti6Al4V,</td>
<td>47.9</td>
<td>5.6</td>
</tr>
<tr>
<td>thickness @ 600 µm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TiGrowth® on PEEK,</td>
<td>43.8</td>
<td>2.4</td>
</tr>
<tr>
<td>thickness @ 400 µm</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Surface morphometric analysis: Micrographic sections showed titanium coatings with max thickness exceeding 900 µm on metals and 400 µm on PEEK. SEM analysis disclosed pore diameters distributed between 100 and 400 µm. The higher the thickness, the larger the pores obtained. Overall porosity was between 40 and 70 % with a continuity of interconnections.

In-vivo results: Pull out tests revealed a striking increase of the biomechanical implant-to-bone fixation from 2 to 12 weeks. (Figure 1).

CONCLUSIONS: Morphological analyses on TiGrowth® showed uncommon characteristics in terms of pores size, overall porosity and coating thickness for thermal sprayed coatings. Yet this had no major influence on coating adhesion to Ti6Al4V substrate, as required for orthopedic use. Adhesion was found adequate also for application on PEEK. In vivo results after 12 weeks clearly demonstrated significantly improved fixation values for titanium coated PEEK implants in contrast to uncoated PEEK specimens.


ACKNOWLEDGEMENTS: This study was cosponsored by Provincia Autonoma di Trento, IT.
Simulation of muscle and joint reaction forces as boundary conditions for implant design appropriate to the load

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INTRODUCTION: The design of implants appropriate to load is difficult since in vivo load cases are not known in all details. Muscle and joint reaction forces are very difficult to measure. Application of simulation software to determine these is one answer to this question. Virtual testing of implants generates new knowledge about mechanical behaviour and the design can be improved based on that.

METHODS: A validated three dimensional musculoskeletal gait model [1] of the AnyBody Modeling System included in the AnyBody Managed Model Repository Version 1.3.1 (AMMRV1.3.1) was used in this study. This model allows importing motion capture data as well as external forces measured by force plates in C3D format. The data was recorded at the Gaitlab of Orthopaedische Kinderklinik, Behandlungs-zentrum Aschau im Chiemgau. The female patient was 62 years old with normal weight and with a diagnosed coxarthrosis. For the musculoskeletal simulation the AnyBody Modeling System [2] version 5.0 including a polynomial muscle recruitment criterion of third order [3] has been used to calculate the muscle and joint reaction forces. In AnyBody the motion is divided in a discrete number of time steps defined by the user. For each step, muscles are recruited to gain mechanical equilibrium.

The results were applied to a finite element model in ANSYS 14 as boundary conditions. The model was meshed with tetrahedral elements. A rigid support fixed the femur at the condyles. Assuming an osseo-integrated implant allows using a bonded contact at bone-implant interface. A quasi-static analysis was performed for 10 time steps between heel strike and toe off.

RESULTS: Comparison of hip reaction forces of in vivo measurements [4] and simulation showed good correlation. The support reactions at the femur condyles determined with ANSYS correlated well with the knee reaction forces calculated by AnyBody. Maximum stress at the implant could be observed below the fatigue limit of approximately 600 MPa for medical titanium alloy Ti6Al4V.

DISCUSSION & CONCLUSIONS: Stress distribution in the implant is different if compared to simulated accreditation tests like ISO 7206 standard for hip implants and when only considering hip reaction forces. Contact conditions are unclear in vivo. Micro motion at the bone-implant interface could occur in reality. Much higher stress peaks are expected when higher load motions like running or stair climbing are investigated [4].

Hemi-resurfacing implants of the shoulder: short term osseous integration

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INTRODUCTION: Hemi-resurfacing shoulder implants restore the joint function with less trauma and bone loss compared to total shoulder arthroplasty. Good clinical short- and mid-term results are reported in the current literature [1]. However, due to the radiopaque cup, only little is known about the bone remodeling processes under the implants.

The aim of this study was to evaluate two different shoulder resurfacing designs regarding their osseous integration at the implant interface and the bone stock under the implant.

METHODS: 10 uncemented hemi-resurfacing implants of the shoulder were retrieved from patients undergoing revision due to glenoidal erosion. 5 Epoca RH (Synthes, CH) and 5 Copeland (Biomet, USA) implants were analyzed. The implants including the bone were embedded in polymethylmethacrylate and a section was taken through the centre of the implant with a diamond cutting saw. Sections were stained with Giemsa-Eosin and digitalized.

The relative bone density (BD [%]) under the resurfacing implant and the relative bone implant contact (BIC [%]) at the interface were analyzed using digital imaging software (KS400, Zeiss, D). The interface between implant and tissue was further evaluated by scanning electron microscopy (SEM) and the chemical composition of the materials was analyzed by energy dispersive x-ray analysis (EDX).

An unpaired t-test was used to compare the two implant designs and a p<0.05 determined significance.

RESULTS: Qualitative histological evaluation revealed an inhomogeneous bone distribution with a reduced bone stock under the implant shell. The quantitative evaluation of the BD confirmed the reduced bone stock (Fig.1, Tab.1). No significant difference was observed between the two implant designs (p=0.43). The interface analysis in the SEM confirmed the good bony ingrowths.

DISCUSSION & CONCLUSIONS: Regardless of the implant design, the cementless shoulder resurfacing implants showed a good bone implant contact at the interface. This suggests a sufficient initial stability with a good ingrowth of the bone into the implant surface and coatings. However, the bone stock under the implant shell appeared reduced in most implants. This is probably related to changes in the load transfer and an unloading of the bone, similar as seen in hip resurfacing arthroplasty [2].

Further studies are needed to confirm these observations and also to evaluate the load transfer of cementless shoulder resurfacing implants into the humeral head.

Table 1: Quantitative evaluation results

<table>
<thead>
<tr>
<th></th>
<th>Bone density (BD [%])</th>
<th>Bone implant contact (BIC [%])</th>
</tr>
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<tbody>
<tr>
<td>Epoca RH</td>
<td>10.9±4.0</td>
<td>35.8±7.2</td>
</tr>
<tr>
<td>Copeland</td>
<td>8.7±4.4</td>
<td>36.8±12.2</td>
</tr>
</tbody>
</table>

**Biological safety testing on implant devices**

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**INTRODUCTION:** The biological evaluation of an implant device is an essential step in the progress of certification. Therefore, any potential risk for the use of the device in humans should be investigated thoroughly.

**COMPENDIUM:** The preferential aim of ISO 10993, is the protection of humans from biological risks. This includes risks from the biological compatibility of the device and also microbiological contamination.

For the assessment of the local effects after implantation, ISO 10993-6 has been established (currently under revision) including a new part for assessing the biological response of brain tissue to an implant.

At BSL BIOSERVICE studies for implants, such as orthopaedic implants, implants with contact to brain tissues, osteoinductive materials, drug delivery systems, tissue engineering products or cardiovascular implants are performed. Studies for functional implantation can be necessary to evaluate the functionality and biocompatibility under biological conditions in active mode. Possible endpoints could be the ingrowth behaviour of tissue into an implant, the mechanical resistance at the intended implantation location or the tissue reaction on electromagnetic radiation.

As the tissue configuration in the vicinity of an implant changes with time, ISO 10993-6 recommends to perform studies for short term as well as for long term implantation periods. The respective periods for each implantation device shall be determined by the intended clinical exposure. Long term studies are defined as studies exceeding 12-18 weeks. For these studies it is advisable to use larger species than rodents.

A hygienic concept in the production of an implant prevents microbiological contamination. The hygienic concept includes a set of microbiological studies on the finished product as well as within the production process. Monitoring of the air, surfaces, and staff and of the water quality, knowledge of the microbial quality of the raw materials and additives are basical components. If a cleaning step is implemented in the manufacturing process, the efficacy must be proven e.g. by a combination of a test for cytotoxicity, of scanning electron microscopy (SEM) and x-ray photoelectron spectroscopy (XPS).

![Fig. 1: Left: Orthopaedic implant. Right: Hygiene monitoring of surface by contact plates](image)

The packaged product before sterilization is examined for the bioburden (= microbial load on the product). The bioburden is the basis for the success of the sterilization procedure, which must be validated. A study for endotoxins/pyrogens is necessary for devices with in-/direct contact to cardiovascular blood, cerebrospinal and lymphatic tissue and ophthalmological products.

![Fig. 2: LAL-test for endotoxins](image)

**SUMMARY:** Evaluation of biological safety is mandatory for each implant device. It is necessary to create individual testing strategies for proving the biocompatibility as well as hygienic safety. These strategies need to be uniquely tailored considering the nature of the material as well as the intended clinical use of the device.

http://www.ecmjournal.org
**INTRODUCTION:** Experimental design is an excellent tool to explore the properties of products and materials depending on various parameters. However, the number of distinct parameters or nonlinearities often turns most design plans to unrealistic schedules and need to be simplified and reduced, sometimes until a level where no more useful information can be expected.

However, daily production is, up to a certain extent, an experimental design: production temperatures and speeds have variations, suppliers change, etc. There is often a lot of information to extract from these small changes, as long as the data is collected and treated adequately: without interrupting the production cycle the so-called data analytics cycle [2] is running offline, applying statistical and data mining methods [1] to find out which parameters are critical, which combination of settings bring the best production quality, and finally to suggest which parameters are the most interesting to look at in a future design of experiment.

**METHODS:** There are quite some advantages of analysing in detail the production data prior to any expensive and time consuming design of experiment. One major advantage is that this analysis can take place without interrupting the production. The source of information is the data collected during the daily production processes and the analysis takes place offline and possibly includes external environmental data as humidity, temperature, working plans, etc.

The results of a data analytics cycle on daily production data are multiple. First it is easy then to rank the importance of all recorded parameters. This is important when considering increasing or reducing the amount of stored data as well as when needing to know which parameter has the highest influence on target Key Performance Indicators (KPIs) as costs, speed, quality or a combination of them.

Secondly the effects of these important parameters can be estimated and analysed either as a single effect or in combination with other parameters. Non-linear effects can be extracted as well using non-linear methods in the analysis as decision trees or neuronal networks [1].

Finally all kind of interactions and multidimensional effects can be discovered: it can well be, for example, that the quality of the production is, in average, the same on every production line but that there are differences when considering separately the quality depending on the various supplier as well.

From that point of the data analytics cycle many actions can be taken: it is possible to make a numerical model of the production based on the most important effects, it is possible to make recommendations about optimal or critical combinations [2].

![Fig. 1: Design of experiment and Data Analytics, two complementary approaches to explore parameter variations and optimize KPIs as costs, rate and quality.](http://www.ecmjournal.org)

**DISCUSSION & CONCLUSIONS:** Finally, when considering extending the production to settings that have not yet been used, the data analytics cycle will be an important source of information to define more precisely the design of experiment and to increase the chances of finding the optimal production point that is needed (Fig. 1).

**REFERENCES:**  
Advanced laser ablation for the surface microstructuring of cardiovascular implants

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INTRODUCTION: Cardiovascular diseases account for more than half of all deaths in the developed world. With our aging society, there is a growing demand for cardiovascular implants of all kinds, including heart valves, ventricular assist devices, stents, vascular grafts and pacemakers.

DEVELOPMENT NEEDS: One of the most pressing needs in implant design is the increase of device biocompatibility. This may be achieved by functionalization of the implant surface, for example by application of micro-scale surface textures. Today, the processing of micro-structured surfaces is mostly achieved via physicochemical technologies such as surface roughening, electrochemical polishing or etching, and micro-patterning via imprint lithography. While these methods are well suited for creating either surface topographies with random feature orientation or for application on flat substrates, they show serious limitations when used on three-dimensional surfaces.

In this study we aimed at developing a laser processing technology for the microstructuring of 3D implants and at defining an optimal surface design to improve healing of the endothelium after implant placement.

APPROACH AND RESULTS: In order to produce a high quality microstructured surface on a macroscopic 3D object, a novel laser structuring setup was established, which met the requirements regarding process stability (>24 hrs. of continuous laser structuring) and precision (one micron structure size or less). In contrast to a conventional laser-structuring process, which is usually performed using a 3D scanner (movement of the laser beam achieved by mirrors) in combination with an F-theta lens (for beam focussing), a four-axis control system to move the workpiece and a special aspherical lens was employed to concentrate the laser energy. This led to focussing diameters around one micrometre combined with an elongated region with almost constant beam intensity distribution along the optical axis. Thus, the ablation of material even in the sub-micrometre scale could be achieved (Figure 1).

To test the effect of micro-structures on endothelial wound healing, in vitro experiments were performed using a custom-made flow chamber that reproduces physiological flow conditions. It could be verified that substrates featuring a defined anisotropic topography with one micron parallel ridges (Fig. 2) improved the endothelial wound healing time by approx. 50% compared to smooth, unstructured surfaces [1].

CONCLUSIONS: With the advanced laser ablation technique developed, it was possible to microstructure 3D surfaces in the micrometre and sub-micrometre range. Furthermore, it was shown that the laser processed prototypes meet the geometric requirements for improved endothelialisation. Currently, animal tests are on the way to validate the obtained results in vivo.


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