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Novel Biodegradable Drug Delivery Systems for the Controlled Release of Growth Factors in Bone Healing and Tissue Engineering

ABSTRACT

Tissue engineering has emerged recently as a promising alternative in cases of tissue loss or organ failure by overcoming the problems of rejection and donor scarcity. Tissue engineering attempts to mimetize the body mechanisms of healing. By combining cells from the patient, growth factors and supportive scaffolds, it seeks to achieve the total regeneration of tissues. Bone morphogenetic proteins (BMPs) are cytokines with important roles during embryogenesis, and with strong ability to induce formation of new bone and cartilage. These growth factors have been used as powerful osteoinductive components in several tissue engineering products for bone grafting.

In this context, the main objectives proposed for this thesis were:

i. To develop a strategy to produce, purify and characterize high amounts of recombinant BMPs.

ii. To evaluate the potential of new natural-origin biodegradable polymers for delivery of BMPs.

Currently, most BMPs are obtained from mammalian cell cultures in low yields or from bacteria inclusion bodies after time-consuming refolding steps. Aiming to circumvent the disadvantages of the previously reported methods, we have developed a novel approach for the production of high amounts of pure recombinant BMPs. The mature domains of human BMPs were cloned in pET-25b vector in *Escherichia coli*. This vector contained a leader sequence for secretion of the recombinant protein into the periplasm, where conditions are more adequate for the formation of the dimer conformation of BMPs. The method was applied for producing human BMP-2, BMP-4, BMP-9, BMP-10, BMP-11 and BMP-14. Overexpression of the BMPs was achieved and scaled-up in a bioreactor. The BMPs were purified with affinity chromatography by tagging the protein with an histidine tag, resulting in significantly large amounts of pure protein. In murine myoblast C2C12 cells and in primary cultures of human fat-derived adult stem cells, these growth factors showed bioactivity. This was observed by the induction of alkaline phosphatase activity and the upregulation of the
expression of different genes, stimulated during osteogenic differentiation. No cytotoxicity was observed as inferred by cell viability assays.

Different materials have been proposed as carriers for BMPs. Carriers are used to increase the lifetime, stability and bioactivity of the BMPs and release these growth factors in both timely and site-specific ways. Silk fibroin and an elastin-like polypeptide were investigated as potential microparticle systems for the sustained delivery of different recombinant BMPs.

Silk fibroin microparticles were produced by dropwise addition of ethanol. BMP-2, BMP-9 or BMP-14 were loaded into the particles and could be released in a sustained way in a period of two weeks, as evaluated by ELISA and quantitative dot-blot. The fibroin particles did not show any significant in vitro cytotoxicity. The fibroin particles loaded with BMP-2 were able to induce high levels of ALP activity and osteogenic mineralization in C2C12 cells. In rats, the BMP-2 loaded particles were implanted subcutaneously and ectopic bone formation was observed by live microCT and histology, four weeks post-implantation.

Elastin-like nanoparticles, created by thermo-responsive self-assembly, were also explored for a combined release of BMP-2 and BMP-14. Both growth factors were encapsulated with a high efficiency, and could be delivered in a combined and sustained way, retaining their bioactivity, as observed in C2C12 cells.

The results obtained in the thesis demonstrate that the methodologies that were developed may result in novel therapeutical applications in the biomaterial and regenerative medicine fields.
Novos Sistemas Biodegradáveis para Libertação Controlada de Factores de Crescimento em Medicina Regenerativa e Engenharia de Tecidos do Osso

RESUMO

A Engenharia de Tecidos emergiu recentemente como uma alternativa promissora nos casos de perda ou falha da função de um órgão ou tecido, ultrapassando os problemas de rejeição imunitária e da escassez de dadores. Através da combinação do uso de células do paciente, de factores de crescimento e de estruturas de suporte, espera-se a total regeneração dos tecidos. As proteínas morfogenéticas do osso (BMPs) são citocinas com funções importantes durante a embriogénese, possuindo uma forte capacidade de induzir a formação e regeneração de osso e de cartilagem. Estes factores de crescimento têm sido usados como potentes componentes osteoindutores de diversos produtos de engenharia de tecidos para aplicações ósseas.

Neste contexto, os objectivos principais propostos nesta tese são:

i. Desenvolver uma estratégia para produzir, purificar e caracterizar quantidades significativas de BMPs recombinantes.

ii. Avaliar o potential de novos sistemas naturais de polímeros biodegradáveis para a libertação de BMPs.

De momento, a maioria das BMPs são obtidas a partir de culturas de células de mamífero, produzidas em pequenas quantidades, ou ainda a partir de corpos de inclusão em bactéria, após protocolos morosos de renaturação. Com o objectivo de ultrapassar estas desvantagens, foi desenvolvida uma nova metodologia para a produção de largas quantidades de BMPs puras. O domínio funcional das BMPs humanas foi clonado no vector pET-25b em Escherichia coli. Este vector contém uma sequência sinal que permite a secreção da proteína recombinante para o periplasma, onde o ambiente favorece a formação dos dímeros de BMPs. Este método foi aplicado para a produção de BMP-2, BMP-4, BMP-9, BMP-10, BMP-11 e BMP-14 humanas. A sobreexpressão de BMPs foi atingida e aplicada à larga escala, em bioreactor. As BMPs foram purificadas por cromatografia de afinidade para uma cauda de histidinas, resultando em quantidades significativamente elevadas de proteína pura. Estes factores de crescimento demonstraram bioactividade em células mioblásticas C2C12 e em culturas primárias de células estaminais adultas obtidas de tecido adiposo, observada através da indução da actividade da fosfatase alcalina e da expressão de genes relacionados com a
diferenciação osteogénica. Não foi observada citotoxicidade, como demonstraram testes de viabilidade celular.

Diferentes materiais têm sido propostos como suportes de libertação de BMPs. Estes auxiliam o aumento da estabilidade e bioactividade das BMPs, libertando estes factores de crescimento de maneira específica espacial e temporalmente. A fibroína da seda e um polipéptido baseado em elastina foram investigados como sistemas microparticulados para libertação controlada de diferentes BMPs recombinantes.

Micropartículas de fibroína da seda foram produzidas por adição gota-a-gota com etanol. BMP-2, BMP-9 ou BMP-14 foram imobilizadas nas partículas e poderam ser libertadas de modo contínuo durante um período de duas semanas, como demonstrou a análise por ELISA e dot-blot semi-quantitativo. As partículas de fibroína não demonstraram citotoxicidade in vitro. Partículas carregadas com BMP-2 foram capazes de induzir elevados níveis de actividade de ALP e mineralização osteogénica na linha celular C2C12. Em ratos, as partículas com BMP-2 foram implantadas subcutaneamente e foi observada formação ectópica de osso, através de microCT e análise histológica, quatro semanas após implantação.

Nanopartículas baseadas em elastina foram também estudadas para a libertação combinada de BMP-2 e BMP-14, criadas por auto-organização termosensível. Ambos os factores de crescimento foram encapsulados com eficiências elevadas, e poderam ser libertados de maneira controlada, retendo a sua bioactividade, como observado em células C2C12.

Os resultados descritos nesta tese demonstram que as metodologias que foram desenvolvidas podem resultar em novas aplicações terapêuticas nas áreas dos biomateriais e da medicina regenerativa.
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LIST OF ABBREVIATIONS

AEC (3-amino-9-ethylcarbazole)
ALP (Alkaline phosphatase)
BMP (Bone morphogenetic protein)
BSA (Bovine serum albumin)
BSP (Bone sialoprotein)
BW (Body weight)
Ca-P (Calcium phosphate)
CHO (Chinese Hamster Ovary)
C2C12 (Mouse myoblast cell line)
DNA (Desoxyribonucleic Acid)
cDNA (complementary DNA)
Dlx5 (Distal-less homebox 5)
DLS (Dynamic light scattering)
DMEM (Dulbecco's modified Eagle's medium)
E. coli (Escherichia coli)
E.E. (Encapsulation efficiency)
ELISA (Enzyme Linked Immunosorbent Assay)
ELPs (Elastin-like polymers)
FDA (Food and Drug Administration)
GDF (Growth and differentiation factor)
H & E (Hematoxylin and eosin)
HA (Hydroxyapatite)
HRP (Horseradish peroxidase)
HSV (Herpes simplex virus)
IPTG (Isopropyl-1-thio-β-galactopyranoside)
ITT (Inverse Temperature Transition)
LB (Luria-Bertani broth)
L.C. (Loading capacity)
MTS(3-(4,5)-dimethylthiazol-2-yl)-5-(3-carboxy-methoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium)
MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide)
Mw (Molecular weight)
MwCO (Mw cut-off, in membrane dialysis)
n (release exponent of Korsmeyer model of kinetics)
OC (Osteocalcin)
OP-1 (Osteogenic protein-1)
OSX (Osterix)
p (Probability value, in statistic significance tests)
PBS (Phosphate buffer saline)
PCR (Polymerase chain reaction)
pl (Isoelectric point)
PLA (Polylactic acid)
PLA-DX-PEG (Polylactic acid p-dioxanone polyethylene glycol)
PLGA (Polylactic-co-glycolic acid)
PEG (Polyethylene glycol)
PGA (Polyglycolic acid)
POLII (Polymerase II)
PVDF (Polyvinylidene Fluoride)
R (Released percentage)
$r^2$ (Correlation value)
rhBMP (Recominant human bone morphogenetic protein)
RNA (Ribonucleic Acid)
mRNA (Messenger RNA)
rpm (Rotations per minute)
RT-PCR (Reverse transcription polymerase chain reaction)
Runx2/Cbfa1 (Runt-related transcription factor 2/ core binding factor alpha-1)
SaOS-2 (Human osteosarcoma cell line)
SD (Standard deviation)
SDS-PAGE (Sodium dodecyl sulphate polyacrylamide gel electrophoresis)
SEM (Scanning electron microscopy)
SOC (Super Optimal broth with Catabolite repression) (added glucose)
$T_t$ (Transition temperature) (of elastin-like polypeptides)
TGF-β (Transforming growth factor β)
UR (Unreleased percentage)
v/v (Volume per volume)
w/v (Weight per volume)
$W_d$ (Dry weight)
$W_w$ (Wet weight)
WU (Water uptake)
μCT (Micro computer tomography)
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Figure 4. SDS-PAGE of samples of crude lysates (pellet fraction) with different time periods for expression of rhBMP-2. Respectively, A-H correspond to the following times of production: A (0h), B (4h), C (8h), D (16h), E (22h), F (24h), G (2 days) and H (4 days). Temperature of expression was 25°C. BMP-2
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**Figure 9.** Silk fibroin microparticles, for loading and delivery of rhBMPs, were formed after dropwise addition of ethanol and overnight incubation at - 20 °C.

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**CHAPTER IV**

*Osteoinduction in human fat-derived stem cells by recombinant human BMP-2 (bone morphogenetic protein-2) produced in Escherichia coli*

**Figure 1.** Strategy used for cloning the mature domain of human bone morphogenetic protein-2. A) BMP-2 gene contains three exons represented by boxes and two introns corresponding to the lines. Coding region is represented by black boxes located in exon 2 and 3. The mature domain of human BMP-2 protein is entirely located on exons 3 correspondent to amino acids 283 to 396. B) Construction of vector pET-25b/rhBMP-2 for expression of human BMP-2 in *E. coli*. The mature BMP-2 was cloned into the vector pET-25b together with its own leader sequence and 6-histidine tag for purification.  

**Figure 2.** A) Coomassie-blue stained SDS-PAGE of crude lysates, pellet fraction, from batch cultivation of transformed *E. coli*. Lanes 1) pET-25b clone; 2) rhBMP-2/pET-25b clone. Monomer appears as a well-defined band around 17 kDa. Molecular weight is in kDa (Broad Range, BioRad). B) rhBMP-2 purified by affinity chromatography: 1) Silver stained reduced SDS-PAGE, showing the monomer of BMP-2 with significant purity; 2) Western-blot using anti-his tag antibody; 3) Western-blot using anti-BMP-2 antibody. BMP-2 is visible in monomer, dimer and polymer.  

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standardized with Polymerase II expression. OSX: osterix; OC: osteocalcin; ALP: alkaline phosphatase. Conditions used: 1) No rhBMP-2; 2) 250 ng/ml; 3) 500 ng/ml; 4) 1000 ng/ml; 5) 250 ng/ml commercial preparation; 6) 500 ng/ml commercial preparation; 7) 1000 ng/ml commercial preparation. Data presented are mean values of two independent experiments. (* bioassay was not performed)

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**CHAPTER V**

*Expression, purification and osteogenic bioactivity of recombinant human BMP-4, -9, -10, -11 and -14*

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**Figure 4.** Cell morphology of C2C12 cells after 5 days of cell culture with A) no growth factor (control), B) 500 ng/ml rhBMP-4, C) 500 ng/ml rhBMP-9, D) 500 ng/ml rhBMP-10, E) 500 ng/ml rhBMP-11, and F) 500 ng/ml rhBMP-14. Purified BMP was used. x400. Osteoblast-like morphology (o) was observed in C2C12 cells stimulated with BMP-4, BMP-9 and BMP-14 and myoblast-like morphology (m) in control and cells stimulated with BMP-10.

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Silk fibroin microparticles as carriers for delivery of human recombinant BMPs.

Physical characterization and drug release

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**Figure 4.** Evolution of weight loss of unloaded elastin particles in phosphate buffer saline as a function of immersion time (t). (mean ± SD, n = 3). The remaining mass decreased to 97.1 ± 3.1 % after day 5, 88.9 ± 6.3 % after day 14, and to 64.5 ± 12.5 % after day 30.

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ATTACH I
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SHORT CURRICULUM VITAE

Paulo C. Bessa was born in 1981 in Porto, Portugal. He presently lives in Vienna, Austria and works, as a researcher, in the 3B’s Research Group (Biomaterials, Biodegradables and Biomimetics), under the supervision of Prof. Rui L. Reis, in the Ludwig Boltzmann Institute for Experimental and Clinical Traumatology, in Vienna Austria, under the supervision of Prof. Heinz Redl and of Prof. Martijn van Griensven, and in the Center for Molecular and Environmental Biology (CMBA), under the supervision of Prof. Margarida Casal.

His background includes a four-year graduation in Applied Biology, by the School of Sciences, University of Minho. He has just submitted his PhD thesis entitled “Novel Biodegradable Drug Delivery Systems for the Controlled Release of Growth Factors in Bone Healing and Tissue Engineering” to the University of Minho, which was prepared in cooperation with the Ludwig Boltzmann Institute for Experimental and Clinical Traumatology, in Vienna, Austria.

During the last year of his graduation, Paulo C. Bessa worked in neurobiology and molecular biology fields, in Leiden, The Netherlands. This work was the subject of his senior research project.

In 2004 he joined the 3B’s Research Group, where he has been working ever since on the production of recombinant growth factors and in the development of new drug delivery systems, for both controlled release and tissue engineering applications. During this phase he worked for different time periods at the Ludwig Boltzmann Institute for Experimental and Clinical Traumatology, in Vienna, Austria, under the supervision of Prof. Heinz Redl and of Prof. Martijn van Griensven, under the same scope of study.

As a researcher of the 3B’s Research Group he has been involved in the preparation of several grant proposals, both at the National (Portuguese Foundation for Science and Technology) and European levels (frameworks VI and VII), including the only Tissue Engineering Network of Excellence of Europe, EXPERTISSUES and the EU-funded project HIPPOCRATES.

Paulo C. Bessa has been also involved in the organization of some meetings and conferences such as in the first meeting of the SPCE-TC (Sociedade Portuguesa de Células Estaminais e Terapia Celular) in 2005, Lisboa, Portugal, and in the ESF-EMBO Symposium on “Stem Cells in Tissue Engineering: Isolation, Culture, Characterization and Applications”, in 2006, in Sant Feliu
de Guixols, Spain. In the group of Prof. Margarida Casal, he has also been involved in assisting in several practical classes of both Genetics and Genetic Engineering (Applied Biology BSc) and in the Master in Biotechnology, during 2004, 2005 and 2008. Finally, he has also assisted in the invited peer-revision of several manuscripts in the field.

As a result of his research work he attended several important international and national meetings in the present field of research. Presently he is the author of 7 papers in international peer-reviewed journals (4 published, 3 submitted), 1 published book chapter and 11 abstracts published in international conferences.
The work performed during this PhD resulted in the following publications:

- PAPERS IN REFEREED JOURNALS


**Bessa PC**, Casal M and Reis RL, 2008, Bone morphogenetic proteins in tissue engineering: the road from laboratory to the clinic, part I, basic concepts, Journal of Tissue Engineering and Regenerative Medicine 2, 1-13


**Bessa PC**, Cerqueira MT, Rada T, Gomes ME, Neves NM, Nobre A, Reis RL and Casal M, 2009, Expression, purification and osteogenic bioactivity of recombinant human BMP-4, -9, -10, -11 and -14, Protein Expression and Purification 63, 89-94.


- **BOOK CHAPTERS**


- **CONFERENCES PROCEEDINGS IN SCIENTIFIC JOURNALS:**


- **COMMUNICATIONS IN NATIONAL AND INTERNATIONAL CONFERENCES:**

**Bessa PC**, Pedro AJ, Casal M and Reis RL, Production of biologically active recombinant human BMP-2 for Tissue Engineering applications, European Society for Artificial Organs Congress (ESAO2005), Bologna, Italy, October 2005, oral
Bessa PC, Pedro AJ, Reis RL and Casal M, Production of recombinant human BMP-2 for bone biomedical applications, Congresso Nacional Micro’05 e Biotec’05, Póvoa de Varzim, Portugal, November 2005, poster


Bessa PC, Gomes S, Leonor IB, Klösch B, Redl H, Casal M and Reis RL, Natural origin biomimetic scaffolds for the delivery of human bone morphogenetic protein-2 produced by recombinant technology, InVents 4th Marie Curie Cutting Edge Conference (Biocompatibility evaluation and biological behaviour of polymeric biomaterials), Algarve, Portugal, October 2007, oral

Balmayor ER, Bessa PC, Tuzlakoglu K, Azevedo HS, Casal M and Reis RL, Novel biodegradable polymeric microparticles for the localized delivery of differentiation agents in bone tissue engineering applications, TERMIS-EU 2008 Annual meeting, Porto, Portugal, June 2008, oral

Bessa PC, Balmayor ER, Cerqueira MT, Rada T, Gomes ME, Neves NM, Azevedo HS, Casal M and Reis RL, Silk nanoparticles for delivery of human BMP-2 in bone regenerative medicine applications, TERMIS-EU 2008 Annual meeting, Porto, Portugal, June 2008, oral

Bessa PC, Cerqueira MT, Rada T, Gomes ME, Neves NM, Casal M and Reis RL, Expression, purification and bioactivity of recombinant human bone morphogenetic protein-4, -9, -10, -11 and -14 produced in Escherichia coli for Tissue Engineering applications, TERMIS-EU 2008 Annual meeting, Porto, Portugal, June 2008, poster
INTRODUCTION TO THE THESIS FORMAT

This thesis is divided in five sections containing nine different chapters, with five of them being experimental research. The contents of each are summarized below.

Section 1 (Chapter I, II) presents an extensive overview on the several aspects behind the field of bone morphogenetic proteins and their importance for the research in bone regenerative medicine. Chapter I starts by presenting an historical background on the discovery and main advances with the use of BMPs in the biomedical field, revealing the importance of these growth factors in inducing new bone and cartilage formation. At the same time, the progress with the expression of recombinant BMPs is discussed. Chapter II introduces the emerging challenges of drug delivery and biomaterial science, from bench to clinical approval. It covers the literature with the use of synthetic polymers, natural polymers, microparticle systems, and ceramics for a controlled release of BMPs, and main clinical applications in bone healing.

Section 2 (Chapter III) presents in detail the materials and experimental procedures used for cloning, expression, purification and bioactivity characterization of the proposed BMPs, and for the fabrication of a silk fibroin microparticle delivery system. In Section 3 (Chapters IV, V) different studies are described that explore the use of recombinant technology in bacteria for the production of human BMPs and their bioactive potential in selected cell lines and in primary cultures of human adult stem cells. In Chapter IV, a novel methodology was proposed for the production of human recombinant BMP-2 that showed bioactivity in human adult stem cells. In Chapter V the same methodology was used for the production of human BMP-4, -9, -10, -11 and -14. In Section 4 (Chapter VI, VII, VIII) novel particulate-based delivery carriers are proposed for the controlled release of human BMPs. In Chapter VI, the silk microparticles were characterized using SEM, DLS, swelling ability, and the release profile of BMP-2, -9 and -14. In Chapter VII the particles were studied for the bioactivity of released BMP-2, using C2C12 cell line, and a rat ectopic bone formation model. In Chapter VIII, elastin-like nanoparticles were produced by thermoresponsive self-assembly, for a combined delivery of BMP-2 and BMP-14.

Section 5 (Chapter IX) contains the general conclusions regarding the overall work carried out under the scope of this thesis, as well as some final remarks and future directions. In the end, Attach I provides supplementary data on the refolding and bioactivity of recombinant BMPs.
Pedras no Caminho?
Guardo todas, um dia vou construir um castelo.
Fernando Pessoa (1888-1935)
Chapter I

General Introduction

Bone morphogenetic proteins in tissue engineering: the road from the laboratory to the clinic, part I (basic concepts)

This Chapter is based on the following publication:

Bessa PC, Casal M and Reis RL, 2008, Bone morphogenetic proteins in tissue engineering: the road from laboratory to the clinic, part I, basic concepts, Journal of Tissue Engineering and Regenerative Medicine 2, 1-13
Chapter I

General Introduction

Bone morphogenetic proteins in tissue engineering: the road from the laboratory to the clinic, part I (basic concepts)

Abstract

Discovered in 1965, bone morphogenetic proteins (BMPs) are a group of cytokines from the transforming growth factor-β (TGF-β) superfamily with significant roles in bone and cartilage formation. BMPs are used as powerful osteoinductive components of diverse tissue engineering products for the healing of bone. Several BMPs with different physiological roles have been identified in humans. The purpose of this review is to cover the biological function of the main members of BMP family, the latest research on BMPs signaling pathways and advances in the production of recombinant BMPs for tissue engineering purposes.

1. Body morphogenetic proteins

Research on bone regeneration began decades ago as a result of intensive studies on bone growth and healing. Bone has been recognized, among the many tissues in human body, as one with the highest potential for regeneration. As early as 1889 Senn noticed that decalcified bone could induce healing of bone defects (Senn, 1889). Later Lavender provided the first evidence of ectopic bone formation after injecting bone crude extracts into muscle (Levander, 1934, 1938). In 1965 the pioneering work by Urist marked a landmark on the research in bone regeneration. Urist discovered that the active compound responsible for bone regeneration was a mix of proteins and named these as bone morphogenetic proteins (BMPs; Urist, 1965). In the years that followed Sampath and Reddi created a crude but highly reproducible bioassay for BMP for ectopic bone formation (Sampath and Reddi, 1981). The assay was based on the activity of the enzyme alkaline phosphatase enzyme and the content of calcium in the newly formed bone. Reddi proposed that BMPs are responsible for the initiation of a cascade of developmental events, in which progenitor cells in the bone marrow were induced by these factors to produce bone cells leading to bone regeneration (Reddi and Huggins, 1972; Reddi, 1981).

During the decades of 80s and 90s the BMP genes were cloned and the recombinant proteins were shown to be biologically potent (Wozney et al., 1988). Much work followed with
the use of recombinant BMPs (rhBMPs) for clinical applications such as spinal fusion, fracture healing and dental tissue engineering (Nakashima and Reddi, 2003; Seeherman and Wozney, 2005). Human BMPs are now produced in larger amounts by recombinant technology. In 2002 FDA gave approval of two products containing rhBMP-2 and rhBMP-7 in absorbable collagen carriers for spinal fusion and long bone non-unions (McKay et al., 2007; White et al., 2007). In a recent review Reddi proposed naming BMPs also as body morphogenetic proteins, due to their extensive roles in various tissues and organs beyond the bone (Reddi, 2005).

2. Members of the BMP family

In humans several members are under the designation of BMPs, from BMP-2 to BMP-18 (Table 1). BMP-1 is not a member of the BMP family. It is a misidentified protein with chordinase and procollagen proteinase activities, implicated in embryonic patterning (Kessler et al., 1996). In fact, since BMPs were named by homology-based cDNA cloning, there is confusion over BMP designations and there is a need to develop a more appropriate nomenclature for the BMP family.

The biological functions of BMPs are mainly related to bone and cartilage formation (Reddi, 2005), although BMP-8b, -10 and -15 do not have known roles in bone or cartilage. BMP-10 is involved in cardiac development and BMP-15 in ovarian physiology (Chen et al., 2004; Knight and Glister, 2006). BMP-8b is involved in reproductive cells (Zhao et al., 1996). Usually only BMP-2 to BMP-11 are considered to be BMPs. BMP-12, -13 and -14 are named cartilage-derived morphogenetic proteins (CDMPs), as these induce chondrogenic phenotypes rather than osteogenesis (Reddi, 2003). BMPs are also involved in embryonic patterning (Kishigami and Mishina, 2005), in skeletal formation (Tsumaki and Yoshikawa, 2005) and in organogenesis of other tissues behind bone. For example, BMP-2 plays a role in heart morphogenesis (Callis et al., 2005) and in neural stem cells (White et al., 2001), BMP-7 in kidney formation (Simic and Vukicevic, 2005) and diverse BMPs have unique roles in reproductive organs (Shimasaki et al., 2004; Tsumaki and Yoshikawa, 2005). During the events of early embryogenesis, BMPs are involved in the dorsal–ventral patterning with different roles, e.g. BMP-2 and -4 induce embryo ventral differentiation, while BMP-3 and -3b oppose the ventralizing effect by inducing tail and head formation, respectively (Hino et al., 2004). During gastrulation a BMP gradient in signaling established by BMP antagonists is responsible for directing cells into forming organs such as bone, cartilage, kidney or heart, depending on the levels of BMP activity and of other cytokines (Yamamoto and Oelgeschlager, 2004). The pleiotropic effect of the different BMPs is of important consideration for tissue engineering. For instance, besides being used for
differentiation of cartilage and bone BMP-4 is being studied for a role in keeping the undifferentiated state in embryonic stem cells (Qi et al., 2004). BMPs clearly possess pivotal roles in controlling the proliferation and differentiation fates of cells. In bone formation a clear bone-inducing role is observed for BMP-2, -4, -6, -7 and -9. These BMPs have been shown to induce bone mineralization and increase in osteocalcin levels in C2C12 cells (Chen et al., 2003) and orthotopic ossification in mice (Kang et al., 2004). BMP-3 and -3b are possibly inhibitors or negative regulators of osteogenesis, as these downregulate the expression of ALP in bone cells (Hino et al., 2004). During fracture healing different BMPs follow specific defined temporal sequences. BMP-2 appears to be an early factor peaking at day 1 after fracture while BMP-14 peaks at day 7 during cartilage formation and BMP-3, -4, -7 and -8 are expressed mainly after 2 weeks (Cho et al., 2002). In analogy different BMPs are also expressed in site-specific patterns during the formation of bone (Zoricic et al., 2003).

2.1. Subgroups within the BMP family

BMPs belong to TGF-β superfamily, which includes several other growth factors, such as activins, inhibins or TGF-βs. The members of the BMP family may be subdivided into different subgroups based in their gene homology and similarity in protein structure (Wozney and Rosen, 1998; Kishigami and Mishina, 2005; Reddi, 2005; see Table 2). The proteins within the BMP-2/4 group, osteogenic protein-1 (OP-1) group, BMP-9/10 group and BMP-12/13/14 group share sequence similarities of more than 50%. BMP-11 and BMP-15 are more distant members, similar to growth differentiation factors -8 and -9, respectively. Interestingly, the morphogens named as growth and differentiation factors (GDFs) have similarity to some BMPs and could therefore be included in the BMP family. However, these are not described in this review since their biological roles are beyond bone induction, except for GDF-3 a BMP inhibitor, and GDF-8, a negative regulator of bone and muscle mass (Levine and Brivanlou, 2006; Hamrick et al., 2007). Additionally, under the term BMP are BMP-16 to -18. These were given disclosure in patent applications (Celeste and Murray, 1999, 2000). BMP-16 is a human homologue to murine Nodal and BMP-17/18 are related to Lefty. Both have important roles during embryonic patterning, mainly by antagonizing the effect of BMP signaling in mesoderm formation (Nodal) or for the establishment of left-right embryonic asymmetry (Lefty) (Meno et al., 1997; Thisse et al., 2000). Since BMP-16 to -18 are more distantly related to BMPs than to other TGF-β superfamily members these should conceivably form a different group in the TGF-β superfamily, distinct from BMPs.
Table 1. BMP members in humans and their main physiological roles. BMPs known to induce complete bone morphogenesis are underlined.

<table>
<thead>
<tr>
<th>BMP</th>
<th>Nomenclature</th>
<th>Main physiological roles</th>
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<tbody>
<tr>
<td><strong>BMP-2</strong></td>
<td>BMP-2a</td>
<td>Cartilage and bone morphogenesis / heart formation</td>
<td>(Wang et al., 1990a; Kang et al., 2004; Callis et al., 2005)</td>
</tr>
<tr>
<td><strong>BMP-3</strong></td>
<td>Osteogenin</td>
<td>Negative regulator of bone morphogenesis</td>
<td>(Hino et al., 2004)</td>
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<td><strong>BMP-3b</strong></td>
<td>GDF-10</td>
<td>Negative regulator of bone morphogenesis</td>
<td>(Hino et al., 2004)</td>
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<td><strong>BMP-5</strong></td>
<td>-</td>
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<td>Bone morphogenesis ? / spermatogenesis</td>
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<td>(Chen et al., 2003; Kang et al., 2004; Lopez-Coviella et al., 2006)</td>
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<td><strong>BMP-13</strong></td>
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<td>Cartilage development and hypertrophy</td>
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<td><strong>BMP-14</strong></td>
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<td>Chondrogenesis / Angiogenesis</td>
<td>(Yamashita et al., 1997; Reddi, 2003; Zeng et al., 2007)</td>
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### Others

<table>
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<tr>
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<td>-</td>
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<td>(Chen et al., 2004)</td>
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<td>GDF-9b</td>
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<td>(Knight and Glister, 2006)</td>
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<td>Lefty</td>
<td>Embryonic patterning</td>
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### Table 2. Suggested subgroups within BMP family (grouped by similarity to OP-1)

<table>
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<tr>
<td>OP-1 (BMP-7)</td>
<td>BMP-5, -6, -7, -8, -8b</td>
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<td>BMP-2/4</td>
<td>BMP-2, -4</td>
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<tr>
<td>CDMP1/2/3</td>
<td>BMP-12, -13, -14 (GDF-5, 6, 7)</td>
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<td>BMP-9/10</td>
<td>BMP-9 (GDF-2), BMP-10</td>
</tr>
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<td>BMP-3/3b</td>
<td>BMP-3, BMP-3b (GDF-10)</td>
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<td>BMP-11/GDF-8</td>
<td>BMP-11 (GDF-11), GDF-8</td>
</tr>
<tr>
<td>BMP-15/GDF-9</td>
<td>BMP-15 (GDF-9b), GDF-9</td>
</tr>
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</table>

### 3. The structure of BMPs

BMPs as all members of the TGF-β superfamily, are homo or heterodimers linked via disulphide bridges. These proteins are expressed as large precursor polypeptide chains containing a hydrophobic signal sequence, a long and poorly conserved N-terminal pro-region sequence, a mature domain with a highly conserved C-terminal region and an N-terminal region that varies among the different BMPs.

BMPs are biologically active in homodimer and heterodimer conformation (Israel et al., 1996). BMPs form a conserved motif of seven cysteines, which is involved in the formation of six intrachain disulphide bonds and a single interchain bond, necessary to dimer formation. Heterodimers in cell cultures have been observed to induce much higher yields of osteogenic marker alkaline phosphatase (ALP) (Aono et al., 1995) than homodimeric BMP. Moreover, BMP-4/7 heterodimers have also been reported for their very potent mesoderm inducing activity in *Xenopus* (Suzuki et al., 1997). A role has been attributed to heterodimers of BMP-
4/8b in primordial germ cell formation (Ying et al., 2001). In humans, there is evidence that BMP heterodimers may be also involved in specific biological processes (Butler and Dodd, 2003), challenging researchers to look for working with heterodimers besides the widely documented investigation with BMP homodimers.

Intracellularly BMPs are produced as precursors that dissociate after proteolytical cleavage by subtilisin-like proprotein convertases to form mature proteins (Ge et al., 2005). However, in some cases the pro-region remains associated after secretion to the extracellular space, inhibiting the binding of BMPs to cell receptors. Importantly, propeptide forms of mature BMPs have been linked to specific physiological roles, such as in synovial rheumatoid arthritis for proBMP-2 and proBMP-6 (Lories et al., 2007), and in the stabilisation of BMPs in the cases of proBMP-4 (Degnin et al., 2004) and proBMP-9 (Brown et al., 2005). It is possible that the half-life of mature BMPs may be influenced by the identity of their prosequences, and that prodomains serve for other cellular effects (Gregory et al., 2005) which by itself is an important consideration for tissue-engineering approaches involving the use of these morphogens.

3.1. Wrist and knuckle epitopes

During the last years crystallographic studies have provided insights into the structure of BMPs and their interaction with receptors upon binding. Crystallographic studies have been reported for BMP-7 (Griffith et al., 1996), BMP-2 (Scheufler et al., 1999) BMP-9 (Scheufler et al., 1999), BMP-9 (Brown et al., 2005) and BMP-14 (Schreuder et al., 2005). These studies have revealed a common polypeptide core for BMPs and TGF-β. The differences in hydrophobic core amongst these BMPs explain the different affinities for the various cell receptors and possibly for its different physiological roles (Allendorph et al., 2006). In BMP-2 two symmetrical pairs of juxtaposed epitopes were described: the wrist epitope, with affinity to BMP receptor I (BmprI), containing residues from both monomers; and the knuckle epitope, which includes residues from only one monomer, with low affinity for BMP receptor II (BmprII; Nickel et al., 2001). Mutations in epitope II lead to loss of activation of both BMP/Smad pathway and induction of ALP, or even complete inhibition of its function (Kirsch et al., 2000). Variants with antagonistic properties are exclusively generated by mutations in the knuckle epitope of BMP-2. A synthetic knuckle peptide alone was also sufficient to induce ALP, osteocalcin and ectopic bone formation in a rat model (Saito et al., 2003). Alterations in epitope I lead only to a reduction of ALP activity while activation of the BMP–Smad pathway is maintained (Knaus and Sebald, 2001). From these findings it was possible to conclude that the activation of the BMP–Smad pathway and the induction of early osteogenic marker ALP were triggered by distinct
BMP–receptor complexes. Different pathways are probably triggered in consequence to the interaction between the different epitopes of BMPs and cell receptors (Hassel et al., 2003).

3.2. N-terminal of BMP

Another particular feature of the structure of BMP-2 is a heparin-binding domain in the N-terminal region of the mature polypeptide, which modifies its biological activity (Ruppert et al., 1996). Ruppert and colleagues demonstrated that the presence of the N-terminus of BMP-2 reduced its specific activity by interacting with heparinic sites in the extracellular matrix. A variant of recombinant BMP-2 was produced without this domain (EHBMP-2), and biological activity increased five-fold in a limb bud assay when compared to normal BMP-2 and was not affected by the presence of heparin. However, other studies have demonstrated that the presence of heparin binding site increased region increased retention times at the site of injury, and that this was necessary for higher osteoinductive effects (Wurzler et al., 2004). In a recent work a basic core of only three amino acids in the N-terminal region of BMP-4 was demonstrated to be required for its site restriction to the non-neural ectoderm (Ohkawara et al., 2002). The authors suggested that heparan sulphate proteoglycans bind to this basic core and thus play a role in trapping BMP-4. This study was the first to identify a critical domain responsible for the interaction between the BMP and the extracellular environment that restricts its diffusion in vivo. Interestingly, heparan sulphates were also observed to be required for BMP-7 signaling (Irie et al., 2003) and recently these have been observed to greatly influence the chondrogenic activity of BMPs (Fisher et al., 2006) and the transcription of several osteogenic genes in response to BMP (Manton et al., 2007). For these reasons heparin has been included in several different tissue engineering constructs using BMPs (Jeon et al., 2007; Lin et al., 2008).

3.3. BMP antagonists

There are diverse BMP-binding proteins that act as modulators of BMP activity or as antagonists, such as Noggin (Zimmerman et al., 1996), Chordin (Blader et al., 1997), Follistatin (Fainsod et al., 1997) and Gremlin (Hsu et al., 1998). These are extracellular regulator proteins that prevent the BMP assembly to cell receptors and in this way modulate the effect of BMP signaling during the formation of tissues. Crystallographic studies showed that the binding of antagonists, such as Noggin, to BMPs inhibited its signaling through blocking the binding of BMP epitopes for type I and type II receptors to which BMPs bind (Groppe et al., 2002). Recently, two new cofactors that regulate BMP signaling, Betaglycan and crypto, were described (Dale et al., 2004). Betaglycan formed a complex with inhibins which binds to activin
type II receptors, thus preventing BMP signaling (Wiater and Vale, 2003). Because of the association of these proteins with various diseases and with additional biological roles, the generation of antagonists of TGF-β superfamily members might generate potent tools for basic research and therapeutic approaches (Gazzerro and Canalis, 2006).

4. BMP signaling – from cell receptor to gene activation

BMPs, and all members of the TGF-β superfamily, bind to serine–threonine kinase receptors on the cell surface, triggering specific intracellular pathways that activate and influence gene transcription, having precise effects in cell proliferation and differentiation (Figure 1). There are types I, II and III receptors for TGF-β superfamily members. Only types I and II appear to play significant roles in BMP binding and signaling, both of which are required for signal transduction (Heldin et al., 1997; Shi and Massague, 2003). For the BMP family, significantly more ligands than receptors are known. To the best of our knowledge the main receptors to which most BMPs bind are type I receptors activin receptor Ia (ActR-I or Alk2), Bmprla (Alk3) and Bmprlb (Alk6) and the type II receptors BMP receptor II (BmprII), activin type II receptor (ActR-II) and activin type IIB receptor (ActR-IIB) (Derynck and Zhang, 2003). However, type III TGF-β receptors have been shown to serve as cell receptors for BMP signaling, mediating epithelial to mesenchymal cell conversion (Kirkbride et al. 2008).

Upon BMP binding low-affinity type II receptors are constitutively activated and type I receptors are activated by the transphosphorylation of the glycine–serine (GS)-rich domain of these receptors (Heldin et al., 1997). Therefore, the specificity of intracellular signals is mainly determined by type I receptors (Miyazono et al., 2005). BMP2-binding to preformed receptor complexes induces the SMAD pathway whereas signaling complexes sequentially assembled by BMP2 result in the activation of p38-mitogen-activated protein kinase (MAPK; Hassel et al., 2003). Different combinations of type I and type II receptors are the key to providing different and specific signals which result in different cell effects (Sebald et al., 2004). In fact, a recent study showed that Alk3 (type Ia) and Alk6 (type Ib) BMP receptors may transmit different signals during the specification and differentiation of mesenchymal lineages (Kaps et al., 2004). BMP-2 and BMP-4 bind preferentially to Alk3 and Alk6, whereas the BMP-7 group binds more to Alk2 and Alk6. BMP-14 (GDF5) binds to Alk6, but not efficiently to other receptors (Miyazono and Miyazawa, 2002), and the same is noted for BMP-15 (Shimasaki et al., 2004; Tsumaki and Yoshikawa, 2005). BMP-13 (GDF6) binds to Alk3 (Kishigami and Mishina, 2005). All these BMPs activate Smad-1, -5 and -8 signaling, whereas for TGF-β, Nodal and activins, Smad-2 and -3 are mainly involved (Shimasaki et al., 2004; Tsumaki and Yoshikawa, 2005). Differently,
BMP-11 binds to Alk4, -5 and -7 to activate Smad-3 signaling (Andersson et al., 2006). It has been shown that BMP-3 binds to Alk4 type I receptor and that it may block BMP signaling by competing with ActR-IIB and reducing Smad-1, -5, -8 or Smad-2, -3 phosphorylation (Gamer et al., 2005). Finally, activin receptor-like kinase 1 (Alk1) was identified as a potential receptor for BMP-9 (Brown et al., 2005) acting through Smad-1 and -5 signaling (Lopez-Coviella et al., 2006). Mutations in the BMP receptors have been described to cause diseases. Mutations in the BmprII gene have been found in patients with primary pulmonary hypertension (PPH; Lane et al., 2000); those in the BmprIA gene have been found in some patients with juvenile polyposis (Howe et al., 2001). Furthermore, proteomic studies have revealed that the C-terminal tail of BMP receptors type II is also involved in the regulation of cytoskeletal proteins (Hassel et al., 2004). Targeting BMP cell receptors may then be an alternative way to mediate bone-inducing stimuli in tissue-engineering approaches.

4.1. BMP/Smad signaling

Smads are the main signal transducers of serine–threonine receptors (Derynck and Zhang, 2003). Upon BMP binding, type I receptors phosphorylate receptor-regulated Smads (R-Smads). Different R-Smads were identified in mammals: Smad1, Smad5, Smad8 and possibly Smad9 (Xu et al., 2003). Activated R-Smads form a complex with common-partner Smad (Co-Smad(Xu et al., 2003)). There is only one described Co-Smad, Smad4. Two phosphorylated R-Smads form a heterotrimeric complex with one Co-Smad, which is translocated into the nucleus and modulates gene transcription with cooperation with other transcription factors (Massague and Wotton, 2000; Derynck and Zhang, 2003; Vincent et al., 2003). All R-Smads are activated by BMP-2 or -4, whereas BMP-6, -7 and -9 only efficiently induce Smad1 and -5 but not Smad8 (Aoki et al., 2001; Brown et al., 2005). Negative regulation of BMP-induced Smad signal transduction is modulated by inhibitory Smads (I-Smads). I-Smads act by inhibiting the activation of type I receptors upon BMP binding (Hanyu et al., 2001). Smad6 and -7 act as I-Smads. Smad6 preferentially inhibits BMP signaling while Smad7 inhibits both TGF-β –activin and BMP signaling. I-Smads also interact with R-Smads, preventing the formation of the complex with Smad4 (Murakami et al., 2003) inhibiting the transcription of specific genes (Edlund et al., 2003). BMP signaling is also modulated by ubiquitin-dependent protein degradation. Smad ubiquitin regulatory factors (Smurfs) induce the ubiquitination and degradation of Smads thus controlling the signal of BMP. Smurfs form a complex with inhibitory Smad7, which associates with type I receptors, promoting the ubiquitination and degradation of these receptors (Murakami et al., 2003). Interestingly, Smurfs also enhance the responsiveness to Smad2, which mediates TGF-β signaling and has as opposite effect to that of
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BMPs (Zhu *et al*., 1999; Zhang *et al*., 2001). The Smad signaling pathway has been also investigated in cancer treatment (Leivonen and Kahari, 2007; ten Dijke and Arthur, 2007; Katoh, 2007). By controlling the different Smad signaling and thus specific cell responses, researchers could in the near future not only achieve a better knowledge of how BMPs but also modulate the subtle biological roles inherent to the diverse TGF-β and BMP members (Schmierer and Hill, 2007).

4.2. BMP/MAPK signaling

Mitogen-activated protein kinase (MAPK) pathways are alternative pathways that are also implicated in the signal transduction of BMPs (Derynck and Zhang, 2003; Massague, 2003). There are three characterized MAPK pathways in mammalian cells: the extracellular signal regulated kinases (ERKs); the c-Jun-NH2-terminal kinases (JNKs); and the p38 MAPK pathways. These pathways are triggered either by cytokines such as BMPs or by environmental stress (Tibbles and Woodgett, 1999). Thus it may be of importance for biomedical purposes that MAPK pathways are considered since these are connected to a wide range of cellular responses. Cooperative interactions have been reported between Smads and transcription factors activated by MAPK pathways. The X-linked inhibitor of apoptosis links the BMP receptor signaling to TGF-β1-activated tyrosine kinase 1 (TAK1) and activates p38 and JNK kinases and nuclear factor-kappaB (NF-κB) (Silverman *et al*., 2003; Takaesu *et al*., 2003). In osteoprogenitor cells TAK1 interacts with Smads, thus interfering with the BMP signals and osteogenic differentiation (Hoffmann *et al*., 2005). MAPK pathways interact with other pathways mediated by different cytokines such as epithelial growth factor (EGF), insulin-like growth factor (IGF) or fibroblast growth factor (FGF) (Massague, 2003). These act by extracellular signal-regulated kinases which inhibit the BMP–Smad signaling through Ras/Raf/Mek (Aubin *et al*., 2004). It was reported that during embryonic differentiation activation of these pathways inhibit the BMP signaling thus favoring neural differentiation (Pera *et al*., 2003). It appears that MAPK pathways modulate BMP–Smad signaling through interaction with two conserved globular domains present in Smads, the MH1 and MH2 domains. The MH1 domain is involved in DNA binding and the MH2 domain in binding to cytoplasmic retention factors, activated receptors, nucleoporins in the nuclear pore, and DNA-binding cofactors, coactivators and corepressors in the nucleus (Yamamoto *et al*., 2003). Recently, it has been shown that MAPK pathways promoted the termination of Smad mediated BMP signaling, in *Xenopus* (Fuentealba *et al*., 2007). The simultaneously application of BMPs and other cytokines in therapeutical situations may therefore have specific
antagonistic or synergistic effects depending on the interactions occurring between the signaling pathways triggered by each growth factor.

4.3. BMP gene modulation

When translocated to the nucleus, Smads regulate gene transcription by interacting and binding to specific DNA sequences and DNA-binding proteins, such as transcriptional factors (Miyazono et al., 2000). BMP activated Smads bind preferentially to the GCCGnCGC sequence of target genes and only weakly to AGAC or GTCT gene sequences, which are preferentially bound by TGF-β- or activin-activated Smads (Kusanagi et al., 2000). Smads also interact with transcriptional factors and transcriptional coactivators and corepressors.

Runt-related transcription Factor (Runx) is one the most studied transcription factors for BMP signaling, regulating processes such as bone formation and haematopoiesis (Ito and Miyazono, 2003). Three isoforms have been identified in mammals: Runx1, Runx2 (also known as Cbfa1, core binding factor A1) and Runx3. The three are reported to interact with R-Smads. Runx2 cooperatively regulates transcription of genes leading to the differentiation of mesenchymal progenitor cells into osteoblasts (Zhang et al., 2000; Miyazono et al., 2004), and for this reason is widely screened as a marker for commitment of cells into the osteochondral lineage and differentiation into bone. Expression levels of Runx2 are low in mesenchymal cells and are induced upon BMP signaling (Maeda et al., 2004). The induction of Runx2 is mediated by a balance between two transcription factors, Dlx5 andMsx2 that responding to BMP signaling, triggering the activation of Runx2 (Miyama et al., 1999; Lee et al., 2003, 2005). Diverse osteogenic markers are then induced by Runx2, such as ALP, osteocalcin, osteopontin and others (Kumar et al., 2005; Takahashi et al., 2005), which are closely related to the process of endochondral calcification.

Osterix (OSX) is another transcription factor that is mediated by BMP/Smad signals and probably by MAPK signaling and other pathways (Celil et al., 2005). Osterix, together with Runx2, are the most studied transcription factors specific for BMP signaling, involved in the differentiation of mesenchymal stem cells into bone cells (Satija et al., 2007). Smad signaling also interacts with other proteins such as ATF4, TAZ, and NFATc1 transcriptional factors (Deng et al, 2008), Hoxc-8 (Shi et al., 1999), MyoD (Liu et al., 2001), OAZ (Hata et al., 2000), Msx1 (Yamamoto et al., 2000), and SIP1 (Verschueren et al., 1999). Menin is another required factor that regulates Runx2-induced transcription of genes during the early commitment phase of osteoblast differentiation (Sowa et al., 2004). Among the several BMP target genes, Id (inhibitor of differentiation or inhibitor of DNA binding) proteins are some of the most important ones (Ogata et al., 1993). Id proteins function as negative regulators of cell
differentiation and as positive regulators of cell proliferation (Yokota, 2001). Recent advances in microarray analysis enable the identification of many BMP target genes in diverse cell types (Clancy et al., 2003; Garcia et al., 2002; Korchynskyi et al., 2003; Kowanetz et al., 2004) and allow a better understanding of the physiological effects of BMPs.

Figure 1. BMP signaling pathways: schematic representation of Smad-dependent and -independent pathways and their main mechanisms of modulation

4.4. Cross-talk with other pathways

BMP signaling is modulated by cross-talk with several other pathways in a very elaborate network of interacting molecules. BMP and TGF-β–activin pathways compete with each other, as they share similar signaling molecules, such as Smad-4 (Candia et al., 1997). Inhibitory Smads are also modulated by several other cytokines and the presence of shear stress in cells (Miyazono et al., 2000). Expression of endogenous TGF-β in preosteoblastic differentiation cells induces I-Smads, which regulate a faster or slower BMP-induced differentiating effect (Maeda et al., 2004). The BMP–Smad pathway was also observed to interact in either synergistic or antagonistic ways with other signaling pathways, such as the Notch pathway (Dahlqvist et al., 2003), the EGF pathway (Kretzschmar et al., 1997), the STAT/LIF pathway (Nakashima et al., 1999), the Wnt pathway (Nishita et al., 2000), IGF and FGF signaling (Pera et al., 2003) and in convergence/antagonistic effect with the MAP kinase pathway (Aubin et al., 2004). MAPK pathway has also been shown to enhance BMP signaling in response to collagen
stimulus through activation of the Ras–ERK pathway (Suzawa et al., 2002). These molecular pathways have been involved in modulating BMP signaling in diverse cellular effects and processes, such as osteoblast maturation (Zamurovic et al., 2004), migration of endothelial cells (Itoh et al., 2004), epithelial-to-mesenchymal cell transition (Zavadil et al., 2004) and inhibition of myogenic differentiation (Dahlqvist et al., 2003).

5. Recombinant BMPs for tissue engineering

After Urist’s pioneering experiments in 1965, BMPs were extracted and purified from bone of several different species including rabbit (Urist et al., 1979), cow (Wang et al., 1988) and human (Urist et al., 1983). Isolation of native BMP from bone results in very low yields around 1–2 μg/kg cortical bone. The need for obtaining larger quantities of BMPs for tissue engineering prompted researchers to produce and purify these growth factors by DNA recombinant technology (Table 3).

Nowadays, recombinant BMPs are produced mainly by two expression systems: in mammalian cells or in bacteria. With mammalian cells the recombinant protein is obtained active but yields are usually very low. With bacteria the protein is obtained in much larger yields but usually in non-soluble inactive inclusion bodies which are later solubilized and refolded to yield bioactive protein. A common disadvantage is that the protein is not glycosylated as is normal in the human body and thus may present reduced stability or biological activity.

Recombinant human BMP-2 was first obtained in mammalian cell cultures from Chinese hamster ovary (CHO) cells (Wang et al., 1990a). The rhBMP-2 induced ectopic bone and cartilage formation when implanted in rats after 2 weeks. Characterization of rhBMP-2 produced in CHO cells led to the conclusion that the protein was secreted in three forms, a propeptide of 40–45 kDa, a mature homodimer of 30 kDa and a small portion of uncleaved precursor of 60 kDa (Israel et al., 1992). The mature peptide possessed N-glycosylation. Sampath et al. confirmed the bioactivity of rhBMP-7 from CHO cells by analysis of bone formation and specific osteogenic markers, such as osteocalcin (Sampath et al., 1992). Koenig et al. (1994) noticed a strong tendency of recombinant BMP-2 to adhere non-specifically to culture dishes or to the negatively charged extracellular matrix via its positively charged N-terminal. They modified the BMP by a mild trypsin digestion, removing the first seven to nine residues, and named it digit-removed BMP-2, that still binds to the BMP receptor. Nowadays, commercial recombinant human BMP-2 lack this N-terminal heparin binding end, since the region decreases the specific bioactivity of the BMP (Ruppert et al., 1996).
In 1996, various combinations of rhBMPs were coexpressed in CHO cells to test for possible heterodimers formation (Israel et al., 1996). Transient co-expression between the different rhBMPs resulted in more bioactivity than expression of any single rhBMP. The rhBMP-2/7 heterodimer yielded 20-fold higher activity in the in vivo ALP assay. In 1998 Kubler and colleagues produced rhBMP-2 and -4 in *Escherichia coli* and rhBMPs were tested in a collagen carrier and induced bone formation in rats after 4 weeks. Recombinant human BMP-6 was later expressed and purified in *E. coli* (Yang et al., 2003), using pET-15b vector. The protein was obtained in inclusion bodies, refolded and induced ALP activity in C3H10T1/2 cells and other osteogenic markers, as observed by RT–PCR.

During the last years, Vallejo and colleagues reported interesting works regarding the production of rhBMP-2 in inclusion bodies, with optimized protein refolding procedures (Vallejo et al., 2002; Vallejo and Rinas, 2004). The group found that refolding of BMP-2 was particularly sensitive to pH, temperature and concentration, but not so much to ionic strength or redox conditions. The rhBMP-2 was refolded in buffer with L-arginine or CHES at pH 8.5 to a yield of 750 mg dimer per litre of bacteria growing media.

In 2005, mature rhBMP-2, unprocessed rhBMP-2 and propeptide of rhBMP-2 were produced in *E. coli* (Hillger et al., 2005), using a pET-11a expression vector. This group reported bioactivity from both the propeptide and an N-terminal truncated BMP-2 form. Curiously, in 2004, Wurzler and co-workers used mutants of rhBMP-2 with enhanced heparin binding sites (in the N-terminus) and a variant with no heparin binding site. The variant with no binding site showed great reduction in osteoinduction, while the mutants with additional heparin binding sites led to enhanced bone formation, due to longer retention times of the BMP-2 in the extracellular matrix (Wurzler et al., 2004; Depprich et al., 2005). This fact could be of great interest for drug-delivery approaches providing recombinant proteins with increased affinity for binding to the extracellular matrix itself in vivo. In 2005, rat BMP-4 has been expressed and purified as inclusion bodies in *Escherichia coli* that are stable in a physiological compatible buffer showing bioactivity in MC3T3-E1 cells (Klösch et al., 2005).

A report on the production of rhBMP-2 was published in 2006, using pET-21a in *E. coli*. The protein, obtained in inclusion bodies, was refolded in different folding buffers (Long et al., 2006). RhBMP-2 was best refolded at low concentrations (0.1 mg/ml), 4 °C and in the presence of L-arginine. Recently, soluble rhBMP-2 was obtained upon production with the use of pET25b *E. coli* vector. The rhBMP-2 showed bioactivity in primary cultures of human fat-derived stem cells and an increase in levels of osteogenic markers Runx2, Osterix and Smad-1 and -5 in the C2C12 cell line (Bessa et al., 2008).
Table 3. Selected works reporting the production of recombinant BMPs in diverse expression systems.

<table>
<thead>
<tr>
<th>BMP</th>
<th>Expression system</th>
<th>Novelty</th>
<th>References</th>
</tr>
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<tbody>
<tr>
<td>BMP2</td>
<td>CHO cells</td>
<td>Bone formation, characterization of expressed BMP-2</td>
<td>(Wang et al., 1990b; Israel et al., 1992)</td>
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<tr>
<td>BMP-2</td>
<td>Insect cells</td>
<td>Alternative expression system</td>
<td>(Maruoka et al., 1995)</td>
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<tr>
<td>BMP2</td>
<td>E. coli</td>
<td>The heparin binding domain reduces ALP and specific in vitro biological activity</td>
<td>(Ruppert et al., 1996)</td>
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<td>BMP2</td>
<td>E. coli</td>
<td>High density expression in bacteria</td>
<td>(Li et al., 1998)</td>
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<td></td>
<td>E. coli, pCYTEXP3</td>
<td>Optimization of refolding conditions for BMP-2</td>
<td>(Vallejo et al., 2002; Vallejo and Rinas, 2004)</td>
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<tr>
<td>BMP2</td>
<td>E. coli</td>
<td>Additional heparin binding domains enhance in vivo bone formation</td>
<td>(Wurzler et al., 2004).</td>
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<td>BMP2</td>
<td>E. coli, pET-11a</td>
<td>Comparison with BMP-2 propeptide</td>
<td>(Hiliger et al., 2005)</td>
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<td>BMP2</td>
<td>E. coli, pET-21a</td>
<td>Use of different refolding buffers</td>
<td>(Long et al., 2006)</td>
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<td>BMP2</td>
<td>E. coli, pET-25b</td>
<td>Bioactivity in human stem cells</td>
<td>(Bessa et al., 2007)</td>
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<td>BMP4</td>
<td>E. coli</td>
<td>Bone formation in mice</td>
<td>(Kubler et al., 1998a)</td>
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<td>BMP4</td>
<td>E. coli, pET-11a</td>
<td>Stability in physiological buffer</td>
<td>(Klösch et al., 2005)</td>
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<tr>
<td>BMP6</td>
<td>E. coli, pET-15b</td>
<td>Induction of several osteogenic markers</td>
<td>(Yang et al., 2003)</td>
</tr>
<tr>
<td>BMP7</td>
<td>CHO cells</td>
<td>Bioactivity in rats</td>
<td>(Sampath et al., 1992)</td>
</tr>
<tr>
<td>BMP2/7</td>
<td>CHO cells</td>
<td>Heterodimer expression</td>
<td>(Israel et al., 1996)</td>
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</tbody>
</table>

5.1. From manufacture of BMPs to tissue-engineering products

Since their discovery in 1965, BMPs have revealed a promising future in the field of tissue engineering as powerful components of biomedical products for the regeneration of body parts of human patients, namely bone and cartilage. At present, the state of the art of regenerative medicine envisages the use of growth factors, loaded into scaffolding materials and using precursor or stem cells, possibly from the patient him/herself. It is expected that future advances in fundamental biology will allow researchers to design novel tissue engineering strategies that make use of combinations of different BMPs and other growth factors. The use of these growth factor cocktails will certainly present a challenging approach.
in tissue engineering since much is still poorly understood about the orchestration of molecular cascades underlying the regeneration of tissues.

Currently, there are two products approved by the Food and Drug Administration (FDA) consisting of recombinant BMP-2 and BMP-7 loaded into absorbable collagen sponges for clinical use for fusion of spinal vertebrae and treatment of long bone fractures (McKay et al., 2007; White et al., 2007). However, a whole set of other less understood BMPs will probably make its way into the clinic, such as with the case of cartilage derived morphogenetic proteins (BMP-12, -13 and -14) for cartilage or tendon reconstruction, or BMP-9 that has stronger osteogenic potencies compared to other BMPs (Kang et al. 2004). Recombinant BMPs used in clinics are derived from mammalian expression cells, which elevates much of the cost of clinical products containing rhBMPs. It is expected that advances in recombinant technology namely those involving the large-scale production of BMPs in bacteria will allow researchers to obtain and use these proteins at a much lower cost.

At present, recombinant technology is also a useful tool to achieve molecular modifications of BMPs to improve their bioactivity, stability and affinity to both the tissue-engineering scaffold or the extracellular matrix and cells (see Table 4). For example, by adding additional heparin binding domains, there is a significantly different bioactivity and bioavailability of the rhBMP, decreasing its specific activity in vitro but increasing bone formation in vivo (Depprich et al., 2005). Currently, most recombinant BMP-2 and BMP-7 are produced with no heparin binding domain and thus may present reduced bioactivity when used for clinical uses. Another approach may consist of producing recombinant peptides containing only the binding sites to the cell receptors thus achieving specific biological effects with no problems derived from loss of bioactivity or stability which occur during the incorporation into the scaffold (Suzuki et al., 2000; Saito et al., 2003, 2004, 2005, 2006). Designing chimeric proteins is a further powerful approach which may include the addition of other domains of biological importance such as immune modulatory agents, RGD for increased cell attachment, GFP for in vivo localization, or domains for specific binding to biomaterials such as for example to collagen (Han et al., 2002) or to fibrin (Schmoekel et al., 2005). Naturally, the chimeric rhBMPs would have different refolding characteristics as compared to native proteins. These could have altered stability, solubility, surface binding and bioactivity/biospecificity thus making an appealing approach of producing specific tissue target proteins and with increased bioactivity (Oppermann et al., 2005). Chemical modification also improves binding, stability, bioactivity and bioavailability of BMPs (Luginbuehl et al., 2004). Thus, by chemically or recombinantly modifying the isoelectric point of the BMP the affinity of the BMPs to the delivery matrices could be greatly enhanced that allows the achievement of a
desired release profile. Chemically modified rhBMP-2 with enhanced affinity to carriers has been reported by succinylation (Hollinger et al., 1998), acetylation (Uludag et al., 2000) and biotinylation (Uludag et al., 1999b). Finally, the expression of recombinant BMP heterodimers is another way of obtaining these growth factors with much increased bioactivity (Aono et al., 1995; Israel et al., 1996).

Despite being promising recombinant technology is still at an early stage, since there is a need to bypass some fundamental limitations, such as optimizing the stability and bioactivity of recombinant BMPs, obtaining glycosylation patterns identical to that of native BMP, and reducing the possibility of triggering immunogenic responses when used in clinical situations. In the next years, the combination of advances of the BMP field with that of biomaterial science (Mano et al., 2007) will surely herald a revealing future for the use of these morphogens for biomedical and tissue-engineering applications.
Table 4. Chemical or genetic engineering modifications for improving particular characteristics of BMPs for use in tissue engineering applications.

<table>
<thead>
<tr>
<th>Modification process</th>
<th>Advantages</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Enzymatic cleavage of BMP heparin binding domain</td>
<td>Reduces non-specific interactions with extracellular matrix, leading to increased bioactivity <em>in vitro</em></td>
<td>(Koenig <em>et al.</em>, 1994; Uludag <em>et al.</em>, 2001)</td>
</tr>
<tr>
<td>Plasmin enzymatic cleavage of BMP</td>
<td>Reduces non-specific interactions with extracellular matrix, leading to increased bioactivity <em>in vitro</em></td>
<td>(Israel <em>et al.</em>, 1992; Hollinger <em>et al.</em>, 1998)</td>
</tr>
<tr>
<td>Adding additional heparin binding domains</td>
<td>Increases retention to extracellular matrix or affinity to biomaterial, leading to increased activity <em>in vivo</em></td>
<td>(Wurzler <em>et al.</em>, 2004; Depprich <em>et al.</em>, 2005)</td>
</tr>
<tr>
<td>Immobilization of synthetic peptides of partial BMP sequences</td>
<td>Improves the stability and specific activity of the BMP and reduces the initial burst release</td>
<td>(Suzuki <em>et al.</em>, 2000; Saito <em>et al.</em>, 2003, 2004, 2005; Saito <em>et al.</em>, 2006; Seol <em>et al.</em>, 2006; Bergeron <em>et al.</em>, 2007; Park <em>et al.</em>, 2007)</td>
</tr>
<tr>
<td>Fusion protein BMP with domain to bind a specific biomaterial</td>
<td>Increases the retention of BMP on carrier, accelerating tissue regeneration; increases stability and activity of the BMP</td>
<td>(Han <em>et al.</em>, 2002; Schmoekel <em>et al.</em>, 2005)</td>
</tr>
<tr>
<td>Fusion protein BMP with RGD for enhanced cell attachment or domain for other specific biological function</td>
<td>Enhances attachment of cells, tissue regeneration and possibly the biocompatibility of the material</td>
<td>Not yet reported</td>
</tr>
<tr>
<td>Chemical modification of BMP such as with succinylation, acetylation or biotinylation, changes the isoelectric point</td>
<td>Enhances the affinity for binding to the biomaterial by changing the peptide solubility, to suit a desired release profile</td>
<td>(Hollinger <em>et al.</em>, 1998; Uludag <em>et al.</em>, 1999a; Uludag <em>et al.</em>, 1999b; Uludag <em>et al.</em>, 2000; Uludag <em>et al.</em>, 2001)</td>
</tr>
<tr>
<td>Formation of heterodimers between different BMPs</td>
<td>The increased bioactivity of BMP heterodimers leads to lower dose</td>
<td>(Aono <em>et al.</em>, 1995; Israel <em>et al.</em>, 1996; Zhu <em>et al.</em>, 2006)</td>
</tr>
</tbody>
</table>
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Chapter II

General Introduction

Bone morphogenetic proteins in tissue engineering: the road from the laboratory to the clinic, part II (BMP delivery)

This Chapter is based on the following publication:
Chapter II

General Introduction

Bone morphogenetic proteins in tissue engineering: the road from the laboratory to the clinic, part II (BMP delivery)

Abstract

Bone morphogenetic proteins (BMPs) are cytokines with a strong effect on bone and cartilage growth and with important roles during embryonic patterning and early skeletal forming. BMPs have promising potential for clinical bone and cartilage repair, working as powerful bone-inducing components in diverse tissue engineering products. Synthetic polymers, natural origin polymers, inorganic materials and composites may be used as carriers for delivery of BMPs. Carriers range from nanoparticles to complex 3D scaffolds, membranes for tissue-guided regeneration, biomimetic surfaces and smart thermo-sensitive hydrogels. Current clinical uses include spinal fusion, healing of long bone defects, craniofacial and periodontal applications, amongst others. BMP-2 and BMP-7 have recently received approval by Food and Drug Administration (FDA) for specific clinical cases, delivered in absorbable collagen sponges. Considering the expanding number of publications in the field of BMPs, there are prospects of a brilliant future in the field of regenerative medicine of bone and cartilage with the use of BMPs.

1. Introduction

Every year millions of surgical operations are performed for the healing or repair of an organ. In the past two decades, tissue engineering has emerged as a very promising alternative that circumvents several of the limitations of the existing options of autografting and allografting for the treatment of a malfunctioning or lost body part. Tissue engineering combines precursor cells from the patient with scaffolding matrices and the stimulus of growth factors. Since the advent of tissue engineering, bone has received particular interest since it is one of the tissues with most regenerative abilities in the human body.

Bone morphogenetic proteins (BMPs) are probably the most important growth factors in bone formation and healing (Reddi, 1998, 2005). These cytokines have been extensively studied during the past decades and, nowadays, recombinant human BMPs (rhBMPs) are widely used several tissue engineering products that might serve for the complete
regeneration of bone or cartilage. Current applications include loaded rhBMPs in delivery systems made of synthetic or natural polymers and the differentiation of transplanted stem cells from the patient with rhBMPs for later body implantation. The purpose of this review is to cover the latest developments in the research for a BMP delivery carrier involving the use of biomaterials science, particularly with the use of natural origin polymers, to the recent preclinical trials and approved products for clinical applications.

2. Delivering BMPs

2.1. BMP carriers– from bench to clinical approval

The main role of a delivery system for BMPs is to retain these growth factors at the site of injury for a prolonged time frame, providing an initial support for cells to attach and form regenerated tissue (Seeherman and Wozney, 2005). The carrier should provoke optimal inflammatory responses, be biodegradable to allow the formation of an interface with the surrounding biological tissue or complete biodegradability for complete invasion of healed tissues, it should present adequate porosity to allow the infiltration of cells and formation of blood vessels at the new bone. Furthermore, the carrier should protect the BMPs from degradation and maintain its bioactivity whilst releasing the protein on a time and space-controlled way for promoting the formation of new bone at the treatment site. Finally, carriers should be conveniently sterilized, easy to handle, stable over time with well-defined storage procedures, as well as suitable for commercial production, allowing scale-up production and approval by regulatory agencies. Having in consideration, the type of tissue to be regenerated is also of critical importance as different mechanical requirements differ for repair of bone, cartilage or tendon. For example, bone carriers are simplified by the fact that, upon fracture, bone is immobilized but carriers should allow vascular ingrowth due to the highly vascularised nature of bone. In cartilage, defects are subject to high compressive and shear stresses thus making healing more challenging. In tendon the regenerative ability appears to be intermediate to those of bone and cartilage, so tendons are very difficult to immobilize, needing a carrier that is able to withstand considerable tensile forces. The geometry of the carrier also affects significantly the biophysical process of osteoinduction and capillary penetration (Jin et al., 2000). All this in consideration, a research may have to keep in mind that the carrier is evidently aimed for common usage by surgeons and physicians.
2.2. BMP retention at the orthopedic site

The delivery of BMP shall last for a sufficient period of time to induce a specific amount of bone mass to treat the defect. Retention of BMP at the orthopedic site of injury is affected by many parameters such as interaction between the biomaterial and the BMP, influence of pH, temperature, porosity, influence of salts, concentration. Evidently, retention of the growth factor depends on whether the BMP is immobilized on the carrier during its manufacture or absorbed into the surface of the device.

Immobilization of the BMPs in a delivery system may be performed by different methodologies: via adsorption, entrapment or immobilization or by covalent binding (Luginbuehl et al., 2004). In case of adsorption, impregnation of the delivery matrix with the BMP is simpler but conformational changes might occur and the release of the protein is less sustained. Furthermore, delivery by adsorbed growth factors often results in initial burst release. With entrapment methodology, hydrophobic polymeric matrices are well known and described to immobilize and release bioactive agents over extended period of times (Langer and Folkman, 1976). However, difficulty arises over the fact that during processing of certain materials into carriers, pH conditions or temperature conditions often result in denaturation of the protein. Much research nowadays aims to develop specific methods of producing delivery carriers for BMPs that do not cause their loss of activity. Lastly, the BMP may become immobilized by covalent binding to the carrier. This may be performed by production of a fusion BMP protein with a domain of specific binding to a biomaterial (Suzuki et al., 2000). In this regard, recombinant technology offers great versatility for expression of a BMP capable of binding to most natural polymers. Other interesting approaches include exploring the strong affinity of BMPs to the extracellular matrix heparan sulfate/heparin proteoglycans (Blanquaert et al., 1999), ion-complexation by binding to charged polymers such as the cases of chitosan, alginate, hyaluronans or synthetic polyelectrolytes (Yamamoto et al., 1998) and crystallization of growth factors (Jen et al., 2002).

2.3. Pharmacokinetic profiles of released BMP

It is crucial to consider that release may preferably be sustained over time. Extremes in release profiles such as long low-amount release of BMP or initial burst of BMP are known for not being beneficial to bone healing. A delicate balance in concentration of BMPs helps to prevent either insufficient binding to carrier due to low concentration or to precipitation due to high BMP concentration. The time of release may be dependent on the type of fracture or the application in question. It is clear that there is more than one desirable pharmacokinetic profile. The pharmacokinetic profile varies according to the material in consideration, its
formulation, and the type and amount of BMP in use. By chemical modification of the carrier or the BMP, we may achieve a specific release profile, which is of interest since different BMPs may present different release profiles due to their different amino acid sequence, different species have different optimal release profiles (Li and Wozney, 2001) and chances are that the optimal profile may be also site-specific. Depending on the site of injury or on a particular application, various formulations of delivery systems may be designed, from simple nano/microparticles to scaffolds of increased 3D complexity, such as those that mimic the physical properties of the extracellular matrix or hydrogels that respond to physiological shifts such as pH or temperature.

3. Carriers for BMPs

3.1. Synthetic biodegradable polymers

Synthetic polymers have been widely used in tissue engineering applications (Saito and Takaoka, 2003) (Table 1). Initially, poly(lactic acid) (PLA) was investigated as a carrier for BMP delivery (Miyamoto et al., 1992) but the material was considered ineffective due the release of acidic degradation sub-products. However, novel biodegradable synthetic polymers have attracted attention since these are free of the risk of disease transmission that occurs with other materials used for bone applications such as collagen. Biodegradable polymers like polylactic acid p-dioxanone polyethylene glycol (PLA-DX-PEG) allow percutaneous injection after heating, for use as a scaffold and a delivery carrier for BMPs, due to its versatile temperature-dependent liquid–semisolid transition. This plasticity allows synchronizing the biodegradation of the polymer with the induction of new bone by BMP (Saito et al., 2001) and this type of injectable polymeric delivery system, polymerizing in situ, enables a less invasive approach to bone surgeries (Saito et al., 2003b). These scaffolds were tested, as carriers for BMPs, in a variety of models, such as a canine spinal fusion model and in the formation of artificial joint (Saito et al., 2005b), for long bone defects in rabbits (Yoneda et al., 2005) and in dogs (Murakami et al., 2003), and in healing of rat cranial defects (Suzuki et al., 2006). These works showed that PLA-DX-PEG delivered rhBMP-2 successfully, inducing the repair of bone defects several weeks after implantation. In other reports, composites of PLA-DX-PEG with calcium phosphate were shown to require less rhBMP to induce new bone formation in mice (Matsushita et al., 2004) and in healing femur defects of rabbits (Matsushita et al., 2006). Composites of PLA-PEG with hydroxyapatite were also evaluated for articular cartilage repair in rabbits (Tamai et al., 2005) and in a rabbit radii model (Kaito et al., 2005), showing enhanced tissue repair in the animals treated with rhBMP-2 and hydroxyapatite composites.
Poly (lactic-co-glycolic acid) (PLGA) combines the adsorptive stability of PLA with the mechanical strength of polyglycolic acid (PGA) and has received particular attention (Winet and Hollinger, 1993). Biodegradation of the synthetic composite is achieved by varying the proportion of each of the two component materials (Miller et al., 1977; Grayson et al., 2004). PLGA as a carrier for rhBMP-2 delivery was reported in alveolar cleft repair on dogs (Mayer et al., 1996), in gelatin sponge composites in a rabbit ulna model (Kokubo et al., 2003) and in tooth defects of dogs (Kawamoto et al., 2003), and in combination with bone marrow cells in a rabbit segmental bone defect model (Hu et al., 2005). These studies confirm the good results that are usually obtained with PLGA scaffolds; bone formation was observed successfully when the scaffolds delivered rhBMP, as compared to controls. The dosage of rhBMP was also observed to significantly affect the repair of bone defects. Recently, PLGA scaffolds have been also tested in rats (Shimazu et al., 2006), a canine model (Jones et al., 2006) and sheep (Zheng et al., 2006), showing that delivered BMP induced much higher bone formation than the scaffold alone, over the several weeks following implantation. Another report, which involved a PLGA scaffold conjugated to heparin, showed that a much longer sustained release of rhBMP-2 and significantly increased in vivo new formation of bone were achieved (Jeon et al., 2007), indicating the promising potential that heparin has as a stabilizing agent for BMP bioactivity.

Synthetic polymers have been also formulated as hydrogels for delivery of BMPs. Since hydrogels contain large amounts of water, they are interesting devices for the delivery of therapeutic proteins. Lutolf and colleagues reported using synthetic PEG-based hydrogels that mimic the invasive characteristics of extracellular matrices, with integrin-binding sites for cell attachment and substrates for matrix metalloproteinases (Lutolf et al., 2003a; Lutolf et al., 2003b), in a rat model, for rhBMP-2 delivery. The authors demonstrated that cells were able to fully penetrate the hydrogels and bone tissue was formed within 3-4 weeks in the gels that delivered rhBMP-2. Similarly, PEG-based hydrogels were reported by Pratt and co-workers showing that cells were able to fully invade the gel networks that were conjugated with peptides that mimic characteristics from extracellular matrix such as plasmin and a heparin molecule to improve the rhBMP-2 stability (Pratt et al., 2004). In another work, Fisher and colleagues evaluated thermoreversible hydrogels of poly(propylene fumarate-co-ethylene glycol), that mimicked properties of cartilage matrix hydrophilic proteoglycans, for cartilage tissue engineering, using rhBMP-7 (Fisher et al., 2004). The solutions of this polymer were aqueous at 25 °C but readily polymerized into gel above 35 °C. The group proposed the use of these hydrogels for articular cartilage repair. Identically, Gao and Uludag also reported using rhBMP-2 in N-isopropylacrylamide-based thermoreversible hydrogels in a rat model (Gao and
Uludag, 2001). The authors studied the effect of different hydrogel compositions on the *in vivo* retention of rhBMP and conclude that these polymers were much versatile for delivering proteins such as BMPs in more effective and controlled ways. A major disadvantage of the use of synthetic polymers is the risk of inflammatory response due to acidic sub-products of degradation (Winet and Hollinger, 1993), which may be also detrimental to the stability of the incorporated BMPs. This has lead researchers to look forward to other materials such as collagen and other natural polymers, as alternatives for BMP delivery.

Table 1. Synthetic-polymer-based matrices/scaffolds for drug delivery of BMPs for tissue engineering applications.

<table>
<thead>
<tr>
<th>Polymer(s)/Carrier/ scaffold structure</th>
<th>Formulation</th>
<th>Biological model</th>
<th>References</th>
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<tbody>
<tr>
<td>PLA</td>
<td>Scaffolds</td>
<td>Rabbit ulna</td>
<td>(He et al., 2003)</td>
</tr>
<tr>
<td></td>
<td>Scaffolds</td>
<td><em>In vitro</em> differentiation of chondrocytes</td>
<td>(Yang et al., 2006)</td>
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<td></td>
<td>Scaffolds</td>
<td>Rat ectopic bone formation</td>
<td>(Chang et al., 2007)</td>
</tr>
<tr>
<td>PLA-collagen</td>
<td>Membrane</td>
<td>Rabbit ectopic bone formation</td>
<td>(Tian et al., 2004)</td>
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<td>PLA-collagen-HA</td>
<td>Composites</td>
<td>Radius defects in dogs</td>
<td>(Hu et al., 2003)</td>
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<td></td>
<td>Composites</td>
<td>Mice ectopic bone formation</td>
<td>(Zhang et al., 2005)</td>
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<td>PLA-PEG-HA</td>
<td>Composites</td>
<td>Rabbit radius model</td>
<td>(Kaito et al., 2005)</td>
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<td></td>
<td>Composites</td>
<td>Articular cartilage repair rabbits</td>
<td>(Tamai et al., 2005)</td>
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<tr>
<td>PLA-DX-PEG</td>
<td>Scaffolds</td>
<td>Femoral canine model</td>
<td>(Murakami et al., 2003)</td>
</tr>
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<td></td>
<td>Scaffolds</td>
<td>Rat cranial defects</td>
<td>(Suzuki et al., 2006)</td>
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<td></td>
<td>Scaffolds</td>
<td>Mice ectopic bone formation</td>
<td>(Kato et al., 2006)</td>
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<td>PLA-DX-PEG-CaP</td>
<td>Composites</td>
<td>Ectopic bone formation in mice</td>
<td>(Matsushita et al., 2004)</td>
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<td>Composites</td>
<td>Spinal fusion in rabbits</td>
<td>(Namikawa et al., 2005)</td>
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<td>Composites</td>
<td>Femur defects in rabbits</td>
<td>(Yoneda et al., 2005)</td>
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<td></td>
<td>Composites</td>
<td>Femur defects in rabbits</td>
<td>(Matsushita et al., 2006)</td>
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### Chapter II. Introduction. Bone morphogenetic proteins in tissue engineering: BMP delivery

<table>
<thead>
<tr>
<th>Protein</th>
<th>Type</th>
<th>Application</th>
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<tr>
<td>PGA</td>
<td>Membrane</td>
<td>Periodontal repair in dogs</td>
<td>(Wikesjo et al., 2003)</td>
</tr>
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<td>Scaffolds</td>
<td>In vitro cartilage formation</td>
<td>(Blunk et al., 2003)</td>
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<td>Scaffolds</td>
<td>Cervical spinal fusion in goats</td>
<td>(Lippman et al., 2004)</td>
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<td>PLGA</td>
<td>Scaffolds</td>
<td>Alveolar cleft repair in dogs</td>
<td>(Mayer et al., 1996)</td>
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<td>Scaffolds</td>
<td>Rabbit radius defects</td>
<td>(Hu et al., 2005)</td>
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<tr>
<td></td>
<td>Scaffolds</td>
<td>Alveolar ridge defects in rats</td>
<td>(Shimazu et al., 2006)</td>
</tr>
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<td></td>
<td>Scaffolds</td>
<td>Canine mandible defects</td>
<td>(Jones et al., 2006)</td>
</tr>
<tr>
<td></td>
<td>Scaffolds</td>
<td>Reconstruction of orbital floor defects in sheep</td>
<td>(Zheng et al., 2006)</td>
</tr>
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<td>PLGA-heparin</td>
<td>Composite</td>
<td>Rat ectopic model</td>
<td>(Jeon et al., 2007)</td>
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<td>PLGA-gelatin</td>
<td>Composites</td>
<td>Rabbit ulna defects</td>
<td>(Kokubo et al., 2003)</td>
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<td>Tooth defects in dogs</td>
<td>(Kawamoto et al., 2003)</td>
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<td>Composites</td>
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<td>Hydrogels</td>
<td>Rat cranial defects</td>
<td>(Lutolf et al., 2003a; Lutolf et al., 2003b)</td>
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<td>Hydrogels</td>
<td>Rat critical size calvarial defects</td>
<td>(Pratt et al., 2004)</td>
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<tr>
<td>PEG-based</td>
<td>Hydrogels</td>
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<td>(Rizzi et al., 2006)</td>
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<tr>
<td>Polypropylene fumarate</td>
<td>Hydrogels</td>
<td>Proliferation of chondrocytes</td>
<td>(Fisher et al., 2004)</td>
</tr>
<tr>
<td>Isopropylacrylamide</td>
<td>Hydrogels</td>
<td>Ectopic bone formation</td>
<td>(Gao and Uludag, 2001)</td>
</tr>
</tbody>
</table>

### 3.2. Collagen

Collagen is the major non-mineral component of bone and also the most abundant protein in connective tissues of mammals. Collagen has received lots of attention due to having good biocompatibility, degrading into physiological compatible products and being suitable for interaction with cells and other macromolecules. The large variety of collagen formulations includes collagen gels, demineralized bone matrix, fibril collagen, collagen strips, membranes, absorbable collagen sponges and composites (Kirker-Head, 2000; Geiger et al., 2003). Another advantage is that collagen can be processed in aqueous form. Collagen has also a favorable influence on cell infiltration and wound healing. During the last years, most researchers have
Chapter II. Introduction. Bone morphogenetic proteins in tissue engineering: BMP delivery

focused on the use of absorbable collagen sponges although several other formulations have been investigated (Kirker-Head, 2000). Collagen sponges are very versatile, easily manipulated and wettable. Manufacture of collagen sponge carrier depends in several factors that include sponge mass, cross-linking methods, sterilization methods, soaking time, protein concentration and buffer composition (Geiger et al., 2003). These steps impact the interaction of the BMP with the collagen carrier and therefore the profile and the efficacy of released protein. For collagen sponges, binding of rhBMP is highly dependent on the pH. Studies using modified versions of recombinant BMP led to conclude that modification of the isoelectric point could bring up to 100-fold differences in the retention of protein to the collagen carrier (Uludag et al., 1999b). Binding of rhBMP-2 is therefore dependable on the isoelectric point of the two proteins and other factors such as ionic strength. Collagen sponges have since then been tested and evaluated in several animal models and clinical trials for cases of fracture repair, critical size defects, spinal fusion and dental and craniofacial reconstruction (Geiger et al., 2003). The collagen sponge consists of lyophilized rhBMP which is reconstituted with water prior to injection and impregnates the collagen sponge for several minutes before implantation. Two models using collagen sponges delivering recombinant human BMP-2 or BMP-7 were approved by FDA for human use as an alternative to bone grafts, for spinal fusion and long bone fractures, after many pre-clinical trials that have been recently reviewed (Gautschi et al., 2007). The collagen sponge holds the BMP and releases it just in the local environment where the surgery was performed, eliminating the need to harvest autologous bone which causes post-operative pain. Based on the extensive preclinical and clinical trials, the use of collagen sponges delivering BMPs has revealed to be a safer and superior alternative to autogenous bone graft. However, although showing success, collagen sponges pose risks of immunogenic reactions since the collagen used on these applications is derived from animal tissues, creating concerns on the risks of transmission of infectious agents and of immunological reactions. For this reason, the development of a superior carrier material for BMP delivery based on other natural polymers is currently being investigated. Alternatively, other sources of collagen, namely from recombinant origin, provides means of obtaining reliable and chemically defined sources of purified human collagens that are free of animal components (Yang et al., 2004).

3.3. Natural origin polymers

The materials for tissue engineering applications should ideally mimic the natural environment of tissues and, in this regard, natural polymers can send signals to guide cells at the various stages of their development and thus accelerate healing (Mano and Reis, 2007).
There are several natural polymers that may be used as carriers for BMP delivery. These may include collagen, starch-based polymers, chitin and chitosan, hyaluronans, alginate, silk, agarose, soy and algae derived materials, and poly(hydroxyalkanoates) (Mano et al., 2007) (Table 2). Several of these polymers are derived from occurring substances in bone, cartilage or in the extracellular matrix. For this reason, these materials often present excellent properties for use in regenerative medicine applications, such as being biodegradable, bioreabsorbable and versatile, as these may be processed into different formulations (Malafaya et al., 2003; Gomes et al., 2004). Natural polymers may present risks of immunogenic reactions and disease transmission, and disadvantages such as the sourcing and processing of the materials. Nevertheless, researchers have been looking for materials from plant origin and produced by microorganisms and/or from recombinant technology which may overcome these concerns.

**Alginate** is a generally safe polysaccharide, known to support the proliferation of chondrocytes *in vitro* (Park et al., 2005b) Very interesting work has been developed by Saito and colleagues with small synthetic peptides corresponding to BMP-2 regions binding to cell receptors, incorporated in cross-linked alginate gels, showing *in vitro* osteogenic differentiation and success in repairing bone defects in rats (Saito et al., 2003a, 2004, 2005a) and in rabbit radial bone defects (Saito et al., 2006). The use of alginate seems to be particularly appealing for cartilage tissue engineering applications since alginate is a major component of cartilage tissue. Alginate hydrogels were also reported for delivery of rhBMP-2 in rats (Simmons et al., 2004) with use of bone marrow cells with RGD peptide for improving cell adhesion.

**Chitosan** is another natural degradable polymer, obtained by alkaline deacetylation of chitin, extracted from the exoskeletons of arthropods. Chitosan has been formulated in many forms, such as hydrogels (Baran et al., 2004) and fiber meshes (Tuzlakoglu et al., 2004), that showed potential for use in osteochondral tissue engineering, making it suitable for BMP delivery (Prabaharan and Mano, 2005). Several works have reported the use of chitosan for delivering BMPs, particularly in composites with synthetic polymers or with other natural polymers. A chitosan-alginate composite gel, loaded with mesenchymal stem cells and rhBMP-2, was evaluated as an injectable tissue engineering construct in mice and induced new trabecular bone formation over a period of 12 weeks (Park et al., 2005a). Liang and colleagues described a chitosan-gelatin scaffold with incorporated rhBMP-2 (Liang et al., 2005) which demonstrated increased expression of bone-marker osteocalcin in osteoblast and myoblast cell lines. In another report, a chitosan blend with PGA was studied as a novel delivery carrier for rhBMP-2 (Hsieh et al., 2006). Derivates of chitosan are also reported. Chemical
modifications of chitosan may enhance certain bioactive properties and increase its solubility in water, thus aiding in the incorporation of rhBMPs, such as in the case of carboxymethyl chitosan. Mattioli-Belmonte and co-workers reported the use of N,N-dicarboxymethyl chitosan, with delivery of rhBMP, for enhancing cell proliferation and healing in articular cartilage lesions (Mattioli-Belmonte et al., 1999a). Recently, rhBMP-2 was immobilized directly on a guided bone-regenerative membrane surface made of chitosan nanofibers that functioned as a bioactive surface to enhance bone-healing (Park et al., 2006). The BMP-2 conjugated membrane surface retained bioactivity for up to 4 weeks of incubation, as well as holding over 50% of the initial BMP-2 attached, promoting cell attachment, proliferation, ALP activity and calcification, when compared with BMP-2 absorbed to the membrane. In two other works, dextran/gelatin based microspheres, containing rhBMP-2, were adhered to chitosan films for guided-tissue regeneration (Chen et al., 2005a) and chitosan membranes activated with BMP-2 were also reported to successfully differentiate C2C12 cells (Lopez-Lacomba et al., 2006b).

Fibrin is derived from blood clots and can be formulated into an adhesive glue-like delivery system (Hattori, 1990). Fibrin has been used as a delivery system for BMPs in a variety of animal models, that includes the use of a fibrin-fibronectin sealing system for rat calvarial defects as a carrier for rhBMP-4 (Han et al., 2005) and for rhBMP-2 (Hong et al., 2006), and a fibrin sealant with rhBMP-2 in the healing of dental pulp of dogs (Ren et al., 2000). In these reports, bone formation was much higher when the fibrin carrier was loaded with the rhBMP, as compared to controls. Fibrin glue might be also a great aid in limiting the diffusion of BMPs into surrounding tissues which could cause undesirable biological effects. In a rat spinal model, fibrin glue significantly limited the diffusion of rhBMP-2 that was loaded into a collagen sponge, preventing the BMP from inducing bone growth in the surrounding spinal cord and nerves (Patel et al., 2006a). Interesting research has been developed by the group of Hubbell with the use of fibrin matrices for the delivery of rhBMPs (Schmoekel et al., 2004; Schmoekel et al., 2005a; Schmoekel et al., 2005b). The group studied the influence of a non-glycosylated form of rhBMP-2 (Schmoekel et al., 2004) in fibrin. Since non-glycosylated rhBMP-2 is less soluble, retention into the fibrin scaffold was enhanced. The fibrin matrices were used to treat critical-size defects and non-unions in rats, dogs and cats. In these studies, bridging of bone defects showed more successful percentages of tissue healing when compared to controls. The group has also reported the use of a fusion BMP protein with an affinity domain to fibrin to increase binding to the carrier (Schmoekel et al., 2005b). Recently, a work using fibrin constructs to deliver rhBMP-2, vascular endothelial growth factor (VEGF) and fibroblast growth factor-2 (FGF-2), combined with hyaluronic acid or collagen, dramatically improved the ability
of blood vessels to directly invade the fibrin-based scaffolds (Smith et al., 2007). Finally, a human trial was reported showing partial reconstruction of a frontal bone defect using heparin together with bovine collagen, hyaluronic acid, and fibrin as vehicles for rhBMP-2 (Arnander et al., 2006). Altogether, fibrin glue certainly seems to be a very useful addition to a bone tissue engineering scaffold using BMPs, having in consideration that aids in promoting osteoinduction (Schwarz et al., 1993) and retention of growth factors (Hubbell, 2006).

**Hyaluronans** are present in extracellular matrix and can be formulated into gels, sponges and pads. Hyaluronans have been used in a variety of trials as a delivery vehicle for rhBMPs including in sponge form in the treatment of alveolar ridge defects in dogs (Hunt et al., 2001), periodontal repair in dogs (Wikesjo et al., 2003), in tibial defects of rabbits (Eckardt et al., 2005), in sheep in combination with hydroxyapatite (Aebli et al., 2005), in the healing of critical size defect in rats in composites with polyactic acid (Vogelin et al., 2005) and in gel and paste forms in non human primates (Seeherman et al., 2004). Kim and Valentini evaluated the kinetics of hyaluronic acid as a delivery system for rhBMP-2 in vitro and demonstrated that hyaluronans-based carriers retained more BMP than collagen gels (Kim and Valentini, 2002). In two other studies, hyaluronic acid was used to deliver BMPS for treating mandibular defects of rats (Arosarena and Collins, 2005a; Arosarena and Collins, 2005b). Significantly more bone was formed in presence of rhBMP-2 and, although not significant, the volumes of new bone were larger for the hyaluronic acid carrier. Recently, a acrylated hyaluronic acid hydrogel was used with human mesenchymal stem cells and rhBMP-2, for healing of rat calvarial defects (Kim et al., 2007). Higher levels of osteocalcin expression and bone formation occurred when the BMP-2 and stem cells were tested. Diverse hydrogel formulations of hyaluronic acid were also evaluated by Bulpit and Aeschlimann, showing excellent cell infiltration and osteochondral differentiation when loaded with BMP-2, in combination with either insulin growth factor-1 or transforming growth factor beta, implanted into rats (Bulpitt and Aeschlimann, 1999). Hyaluronans are observed to interfere positively with BMP cascade (Zou et al., 2004) and since these are part of the extracellular matrix, these may well be priority choices as scaffolds for delivery of BMPs in regenerative medicine of bone.

**Gelatin** has been used mostly in form of hydrogels for delivery of BMPs. Gelatin is an irreversibly hydrolyzed form derived from collagen that is usually cross-linked or hardened through thermal treatment to reduce its high water solubility and enhance the retention of protein to achieve a long-term release. Gelatin hydrogels delivering rhBMP-2 were studied in rabbit skulls (Hong et al., 1998), in mice (Yamamoto et al., 2003b) and recently in skull of non-human primates (Takahashi et al., 2007). Gelatin hydrogels delivering rhBMP-2 were observed to show higher levels of ALP and osteocalcin in comparison with rhBMP-2 delivered...
in collagen sponges (Yamamoto et al., 2003a). Recently, thermomechanical hydrogels based on methacrylated dextran in combination with gelatin have been reported by Chen and colleagues (Chen et al., 2007a; Chen et al., 2007b). The group used rhBMP-2 encapsulated in microspheres of same materials, loaded into the hydrogels, which delivered the growth factor over a period of 18 to 28 days. Their work will be discussed later on the section referring to nanoparticles.

**Dextran** is another natural polysaccharide, synthesized by some bacteria, that has attracted attention for use as a BMP delivery system, because of its excellent hydrophilic nature and biocompatibility. Dextran has been particularly used in form of nanospheres for delivery of rhBMPs that will be detailed in a later chapter. Dextran hydrogels has been evaluated for rhBMP-2 delivery both *in vitro* and *in vivo*, in a rat ectopic model, showing formation of new bone (Maire et al., 2005). The possibility of using natural polymers for designing intelligent hydrogel systems for BMP delivery, is also an interesting and very attractive option. However, no studies have been reported with the use of these systems.

**Starch based polymers** are another interesting alternative for delivering BMPs, that was proposed by Reis as materials with high potential for tissue engineering of bone and cartilage due to their interesting mechanical properties (Malafaya et al., 2001; Elvira et al., 2002). These starch based polymers are used in composites with different synthetic polymers and have been formulated into a variety of forms such as hydrogels (Pereira et al., 1998), nanofibbers (Tuzlakoglu et al., 2005), microparticles (Silva et al., 2004b), or 3D scaffolds (Gomes et al., 2002). The wide variety of formulations and compositions make these polymers suitable scaffolds for bone tissue engineering and controlled release of BMPs. In general, composites of natural polymers with synthetic polymers may become the future option of choice for tissue engineering of bone, since these combine the own specificities of synthetic and natural polymers to produce superior materials.

**Silk fibroin** is a protein derived from cocoons made by the larvae of silkworms. Silk has been proposed and widely investigated as a delivery carrier for BMPs in some contributions reported by the group of Kaplan. In one work, rhBMP-2 was directly immobilized on silk fibroin films and the effect of the delivery system studied in human bone marrow stromal cells and in critical sized cranial defects in mice (Karageorgiou et al., 2004). The rhBMP retained its biological activity. In another report, silk scaffolds fibers, prepared by electrospinning, were used to deliver rhBMP-2 and hydroxyapatite nanoparticles, *for in vitro* bone formation (Li et al., 2006). The rhBMP-2 survived the aqueous-based electrospinning process in bioactive form and induced osteogenesis in cultures of human mesenchymal stem cells. The group also tested BMP-2 delivered via silk fibroin scaffolds in critical size defects in mice (Karageorgiou et al.,
2006). In both works, the delivered rhBMP-2 increased levels of ALP activity, calcium deposition, and transcript levels for bone sialoprotein, osteopontin, osteocalcin, and runx2. In the last years, Meinel and co-workers evaluated the use of silk for tissue engineering constructs with silk-RGD covalently bound matrices, in human mesenchymal cells (Meinel et al., 2004) but not with use of BMPs. In 2006, Meinel and colleagues tested human stem cells loaded in silk fibroin scaffolds, in combination with rhBMP-2, and compared stem cells transfected with BMP-2 via an adenovirus with exogenous protein (Meinel et al., 2006). The expression of osteogenic markers was induced but the BMP was not studied when delivered directly on the silk scaffolds. Recently, rhBMP-2 delivered via silk fibroin scaffolds, in combination with human mesenchymal stem cells, was reported with promising results, in the healing of critical size defects of femurs in rats (Kirker-Head et al., 2007). Compared with other protein-based materials such as collagen, silks have distinguishable mechanical properties, presenting slower degradation times thus allowing adequate time for proper bone remodelling. For this reason, silk is a feasible and potential option as a carrier for controlled delivery of BMPs and, in general, for generating diverse bone tissue engineering constructs for clinical applications (Meinel et al., 2005). Other possible sources of natural polymers for BMP delivery include soy, casein, polyhydroxyalkanoate, polyhydroxybutyrate, corals, carrageenan, gellan gum, agarose and other fibrous proteins such as keratin and elastin (Kirker-Head et al., 2007).
### Chapter II. Introduction. Bone morphogenetic proteins in tissue engineering: BMP delivery

Table 2. Natural origin polymer based matrices for delivery of BMPs for tissue engineering applications. Please refer to Table 3 for micro and nanoparticles formulations.

<table>
<thead>
<tr>
<th>Polymer(s)/Carrier</th>
<th>Formulation</th>
<th>Biological model</th>
<th>References</th>
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</thead>
<tbody>
<tr>
<td>Alginate</td>
<td>Hydrogels</td>
<td>Ectopic bone formation in mice</td>
<td>(Simmons et al., 2004)</td>
</tr>
<tr>
<td></td>
<td>Gels, synthetic BMP-2 oligopeptides</td>
<td>Ectopic bone formation and tibial defects in rats</td>
<td>(Suzuki et al., 2000; Saito et al., 2003a, 2004, 2005a)</td>
</tr>
<tr>
<td></td>
<td>Gels</td>
<td>Rabbit radial bone defects</td>
<td>(Saito et al., 2006)</td>
</tr>
<tr>
<td>Carboxymethylchitosan</td>
<td>-</td>
<td>* In vivo cartilage formation</td>
<td>(Mattioli-Belmonte et al., 1999b)</td>
</tr>
<tr>
<td>Chitosan</td>
<td>Nanofibre membranes</td>
<td>Differentiation of osteoblast cells</td>
<td>(Park et al., 2006)</td>
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<td></td>
<td>Chitosan films</td>
<td>Differentiation of C2C12 cell line</td>
<td>(Lopez-Lacomba et al., 2006a)</td>
</tr>
<tr>
<td>Chitosan-alginate</td>
<td>Gels</td>
<td>Trabecular bone formation in mice</td>
<td>(Park et al., 2005a)</td>
</tr>
<tr>
<td>Chitosan-gelatin</td>
<td>Composites</td>
<td>Differentiation of osteoblasts/myoblasts</td>
<td>(Liang et al., 2005)</td>
</tr>
<tr>
<td>Chitosan-PGA</td>
<td>Composites</td>
<td>*</td>
<td>(Hsieh et al., 2006)</td>
</tr>
<tr>
<td>Dextran</td>
<td>Hydrogels</td>
<td>Rat ectopic model</td>
<td>(Maire et al., 2005)</td>
</tr>
<tr>
<td>Fibrin</td>
<td>Sealant</td>
<td>Dental pulp of dogs</td>
<td>(Ren et al., 2000)</td>
</tr>
<tr>
<td></td>
<td>Gels</td>
<td>Rats, rabbits, dogs and cats; different types of bone defects</td>
<td>(Schmoekel et al., 2004; Schmoekel et al., 2005a; Schmoekel et al., 2005b)</td>
</tr>
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<td></td>
<td>Sealant</td>
<td>Rat calvarial defects</td>
<td>(Han et al., 2005)</td>
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<tr>
<td></td>
<td>Sealant in PCL scaffolds</td>
<td>*</td>
<td>(Rai et al., 2005)</td>
</tr>
<tr>
<td></td>
<td>Sealant</td>
<td>Ectopic bone formation in mice</td>
<td>(Zhu et al., 2006a; Zhu et al., 2006b)</td>
</tr>
<tr>
<td></td>
<td>Sealant</td>
<td>Humans, frontal bone defect</td>
<td>(Arnander et al., 2006)</td>
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<td></td>
<td>Sealant</td>
<td>Differentiation of rabbit bone marrow cells</td>
<td>(Cui et al., 2007)</td>
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<tr>
<td>Material</td>
<td>Form</td>
<td>Application</td>
<td>Study(s)</td>
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<td><strong>Fibrin-CaP</strong></td>
<td>Sealant</td>
<td>Rat calvarial defects</td>
<td>(Hong et al., 2006)</td>
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<td><strong>Fibrin-collagen</strong></td>
<td>Sealant in collagen sponge</td>
<td>Rat spinal model</td>
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<td>Rabbit skull defects</td>
<td>(Hong et al., 1998)</td>
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<td>Hydrogels</td>
<td>Ectopic bone formation in mice</td>
<td>(Yamamoto et al., 2001; Yamamoto et al., 2003b)</td>
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<tr>
<td></td>
<td>Hydrogels</td>
<td>Critical size defects in rabbit ulnas</td>
<td>(Yamamoto et al., 2006)</td>
</tr>
<tr>
<td><strong>Gelatin/dextran</strong></td>
<td>Hydrogels</td>
<td>Skull; non-human primates</td>
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<td><strong>Hyaluronic acid</strong></td>
<td>Hydrogels</td>
<td>Ectopic bone formation in rats</td>
<td>(Bulpitt and Aeschlimann, 1999)</td>
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<tr>
<td></td>
<td>Sponges</td>
<td>Alveolar ridge defects in dogs</td>
<td>(Hunt et al., 2001)</td>
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<td></td>
<td>Scaffolds</td>
<td>Differentiation of CH3H10T1/2 cells</td>
<td>(Kim and Valentini, 2002)</td>
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<tr>
<td></td>
<td>-</td>
<td>Periodontal repair in dogs</td>
<td>(Wikesjo et al., 2003)</td>
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<tr>
<td><strong>Gels</strong></td>
<td>Osteotomy in non-human primates</td>
<td>(Seeherman et al., 2004)</td>
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<td><strong>Gels</strong></td>
<td>Non-union tibial defects in rabbits</td>
<td>(Eckardt et al., 2005)</td>
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<td><strong>Sponges</strong></td>
<td>Rat mandibular defects</td>
<td>(Arosarena and Collins, 2005a; Arosarena and Collins, 2005b)</td>
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<td><strong>Hyaluronan acrylated</strong></td>
<td>Hydrogels</td>
<td>Rat calvarial defects</td>
<td>(Kim et al., 2007)</td>
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<td><strong>Hyaluronic acid- Ti</strong></td>
<td>Composites</td>
<td>Cranial defects in rats</td>
<td>(Itoh et al., 2001)</td>
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<td><strong>Hyaluronic acid- HA</strong></td>
<td>Scaffolds</td>
<td>Osteointegration in cancellous bone in sheeps</td>
<td>(Aebli et al., 2005)</td>
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<td><strong>Hyaluronic acid- PLA</strong></td>
<td>Composites</td>
<td>Critical size defect in rat femurs</td>
<td>(Vogelin et al., 2005)</td>
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<tr>
<td><strong>Silk Fibroin</strong></td>
<td>Films</td>
<td>Cranial defects in mice</td>
<td>(Karageorgiou et al., 2004)</td>
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<tr>
<td></td>
<td>Scaffolds, loaded with human stem cells</td>
<td>Cranial defects in mice</td>
<td>(Karageorgiou et al., 2006)</td>
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</tbody>
</table>
Electrospun nanofibers
Differentiation of human bone marrow cells (Li et al., 2006)
Scaffolds, loaded with human stem cells
Critical size femur defects in rats (Kirker-Head et al., 2007)

* These studies involved solely material testing and delivery kinetics, with no in vitro or in vivo bioactivity models

### 3.4. Ceramics

Many studies dedicated to the understanding of the processes of bone mineralization and it was concluded that ceramic materials such as hydroxyapatite (HA) and other types of calcium phosphates can promote, when implanted, the formation of a bone-like mineral surface layer that leads to increased interface between the materials and the surrounding bone. Calcium phosphate for tissue engineering of bone include use of calcium phosphate layers, films or coatings to promote bone ingrowth, and use of calcium phosphate fillers to replace fractured or damaged bone. Hydroxyapatite (HA) is a form of calcium phosphate mineral that comprises seventy percent of bone and can be formulated as powder, granule, disk or block (Tsuruga et al., 1997). However, for bone tissue engineering applications, specific formulations work better than others, dependant on the geometric structure of the carrier (Kuboki et al., 1998). Hydroxyapatite is a fairly osteoconductive material, and has been used for BMP delivery alone (Noshi et al., 2001) or in composites with natural or with synthetic polymers as previously detailed. Hydroxyapatite has been used in combination with collagen for rabbit spinal fusion (Krai wattanapong et al., 2005), with natural origin polymers (Aebli et al., 2005), with tricalcium phosphate in a rabbit calvarial model delivering rhBMP-2 (Schopper et al., 2007), for differentiating mesenchymal stem cells with BMP-14/GDF-5 (Shimaoka et al., 2004), or for lumbar spinal fusion in non-human primates (Boden et al., 1999). Based on these and other works, hydroxyapatite has proven to be a suitable carrier for BMP delivery, not only enhancing the delivery of the growth factor but also in aiding its retention to the carrier and the osteoconductivity of the scaffold (Uludag et al., 1999a).

Calcium phosphates for delivery of BMPs include calcium phosphates cements and ceramics and calcium phosphate coatings. Calcium phosphate (Ca-P) cements have been extensively investigated as these are osteoconductive, biocompatible and show fast deposition of new bone at the cement surface (Driessens et al., 1998). The BMP may be incorporated into low temperature Ca-P cements by adding the protein in lyophilized form or in aqueous phase.
prior to formation of cement without any risk of denaturation of the growth factor. In high temperature cements, the BMP is generally only adsorbed onto the surface. Porous structure can be fabricated to mimic the structure of trabecular bone (Dutta Roy et al., 2003). Trials for rhBMP-2 delivery have included studies in rabbit ulnas (He et al., 2003) and femurs (Cao et al., 2006), and a canine tibial defect model (Edwards et al., 2004). These studies have demonstrated that the use of Ca-P cements accelerates bone healing. Trials are have also demonstrated the efficacy of calcium phosphate matrices in some non-human primates trials, such as in alveolar ridge surgery using a composite of Ca-P, hydroxyapatite and a collagen sponge (Miranda et al., 2005), in osteotomy sites with a single percutaneous injection of rhBMP-2 loaded into Ca-P cements (Seeherman et al., 2006a), and in posterolateral fusion where Ca-P functioned as a bulking agent to improve osteogenic potential of rhBMP-2 loaded on an absorbable collagen sponge (Barnes et al., 2005). Seeherman also reported achieving bridging of critical size defects in rabbits using the same minimally invasive injectable Ca-P cements (Seeherman et al., 2006b). Ruhé and colleagues also reported several in vivo studies with the use calcium phosphate cements, loaded with rhBMP-2 (Ruhé et al., 2005; Ruhé et al., 2006). A main advantage of the use of calcium phosphates compared to other carriers is that, in general, high doses of rhBMPs are not required (Yuan et al., 2001).

Calcium phosphate coatings are another elegant approach for delivering BMPs by incorporating these growth factors into the latticework of these mineral layers that may serve to coat specific scaffold materials. The BMP is biomimetically deposited during the formation of the calcium phosphate film which is form when the material is immersed on a solution of simulated body fluid that mimics the human blood plasma (Liu et al., 2004). The group of de Groot has used calcium phosphate coated titanium disks for delivery of rhBMP-2 in a rat model, showing that much lower concentrations of BMP are required in comparison with collagen matrices (Liu et al., 2004; Liu et al., 2005). Alternatively, bioactive glass (45S5 - Bioglass), a synthetic surface reactive glass that is commonly used as a filler for damaged or fractured bone, may be also used to form biomimetic calcium phosphate coated scaffolds (Leonor et al., 2003). The biomimetic layers, similar to bone apatite, may be used in combination with BMPs to guide the attachment and differentiation of bone precursor cells, given that the coatings have been shown to promote osteointegration and osteoinduction. Silva and colleagues proposed using blends of starch with polylactic acid and bioglass microspheres for the delivery of BMPs (Silva et al., 2004a). Promising potential arises from the fact that bioglass is osteoconductive and osteoinductive, stimulating the recruitment and differentiation of osteoblasts which produce new bone and completely resorb the material.
Table 3. Micro and nanoscale drug delivery systems based on synthetic and natural-origin polymers. The average size or size range of the particles is noted on the formulation.

<table>
<thead>
<tr>
<th>Carrier</th>
<th>Formulation/ size</th>
<th>Biological models</th>
<th>References</th>
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<td>PLGA</td>
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<td>Rat calvarial bone defects</td>
<td>(Kenley et al., 1994)</td>
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<td>Microparticles (247-430 μm)</td>
<td>Rat femur defects</td>
<td>(Lee et al., 1994)</td>
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<td>Microparticles</td>
<td>In vitro differentiation of osteoblasts</td>
<td>(Oldham et al., 2000)</td>
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<td>Microparticles</td>
<td>Rabbit calvarial bone defects</td>
<td>(Schrier et al., 2001)</td>
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<td>Microparticles</td>
<td>Sheep vertebrae</td>
<td>(Phillips et al., 2006)</td>
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<td>Nanoparticles (300 nm)</td>
<td>Rat ectopic model</td>
<td>(Wei et al., 2007)</td>
</tr>
<tr>
<td>PLGA/Ca-P</td>
<td>Microparticles (66 μm)</td>
<td>Rat ectopic model/ cranial model</td>
<td>(Ruhe et al., 2005)</td>
</tr>
<tr>
<td>PLA</td>
<td>Microparticles</td>
<td>Rat ectopic bone formation</td>
<td>(Saitoh et al., 1994)</td>
</tr>
<tr>
<td>Collagen/HA</td>
<td>Microparticles</td>
<td>Rabbit femoral bone defects</td>
<td>(Wang et al., 2003)</td>
</tr>
<tr>
<td></td>
<td>Nanocrystals/fibbers</td>
<td>Dogs, spinal fusion/ tibia fractures</td>
<td>(Itoh et al., 2004)</td>
</tr>
<tr>
<td>Chitosan/Alginate</td>
<td>Microparticles</td>
<td>In vitro differentiation of rabbit bone marrow stem cells</td>
<td>(Qin et al., 2003)</td>
</tr>
<tr>
<td>Dextran</td>
<td>Nanoparticles (20 nm)</td>
<td>In vitro differentiation of rabbit bone marrow stem cells</td>
<td>(Chen et al., 2005b)</td>
</tr>
<tr>
<td>Dextran/PEG</td>
<td>Microparticles (20-40 μm)</td>
<td>In vitro differentiation of human periodontal ligament cells</td>
<td>(Chen et al., 2006)</td>
</tr>
<tr>
<td>Dextran/Gelatin</td>
<td>Microparticles (20-40 μm)</td>
<td>Canine defects</td>
<td>(Chen et al., 2005a)</td>
</tr>
<tr>
<td></td>
<td>Microparticles (0.5-1.5 μm)</td>
<td>Periodontal regeneration in dogs</td>
<td>(Chen et al., 2007b)</td>
</tr>
</tbody>
</table>

3.5. Microparticles and nanoparticles for BMP delivery

The search for efficient, simple and cheap delivery systems for drug targeting has lead to a great investment on the area of nanoparticles and microparticles for drug delivery. Most common materials for the design of nanodevices to deliver BMPs include synthetic materials,
natural polymers and hydroxyapatite-based particles. Both in the scale of nano (up to 100 nm) and microspheres are reported (Table 3).

Polyactic acid and polyactic-co-glycolic acid have been used as materials for nanoparticles-based delivery system for BMPs. PLA was initially studied as a carrier for BMPs in a rat ectopic bone formation model (Saitoh et al., 1994), showing formation of new bone at 4 weeks after implantation and mature bone after 24 weeks. However, by blending PLA with polyglycolic acid in copolymer polyactic-co-glycolic acid (PLGA), biodegradation is controlled by changing the proportions of each of the two materials, since PLA degrades much slower than PGA. Microspheres of PLGA have been since then evaluated in diverse animal models such as on rat calvarial bone defects (Kenley et al., 1994), rat femur defects (Lee et al., 1994), and in calvarial defects in rabbits (Schrier et al., 2001), forming much more bone when BMPs were delivered via the PLGA particles. Interesting work has been developed by Ruhé et al., using microspheres of PLGA in combination with Ca-P cement, as carriers for rhBMP-2 delivery (Ruhé et al., 2005). The release of rhBMP-2 was observed to be dependent in the composite composition and the nanostructure, as well as of the pH of the release medium. Sustained slow release was observed possibly due to interaction of rhBMP-2 with the calcium phosphate cement. Delivery of rhBMP-7 was evaluated in PLGA nanospheres encapsulated in PLA scaffolds, with interconnected macroporous and nano-fibrous architectures (Wei et al., 2007). The group concluded that the carrier delivered rhBMP-7 in a time-controlled manner and induced significant bone formation.

Diverse natural origin materials were also proposed as carriers at a nano and micro scale for delivering BMPs. Collagen-hydroxyapatite microspheres were evaluated for rhBMP-4 delivery in femoral defects of rabbits (Wang et al., 2003). Regeneration occurred in the animal group treated with BMP-4 particles while with the carrier alone the defects were filled with fibrous tissue and inflammatory cells. Microspheres based in blends of chitosan with sodium alginate were reported in vitro in bone marrow derived cells, showing an increased on the levels of ALP (Qin et al., 2003). During the last years, dextran-based microspheres and nanospheres were extensively evaluated by Chen et al. for the delivery of BMPs. In 2005, the group reported delivering BMP-2 with dextran-based microparticles (20-40 μm) in canine defects (Chen et al., 2005a) and nanoparticles (20 nm) in the differentiation of rabbit bone marrow cells (Chen et al., 2005b) One year later, the authors studied a novel class of methacrylate dextran/PEG microspheres in periodontal ligament cells (Chen et al., 2006). Recently, the group reported dextran-gelatin microspheres loaded into thermomechanical dextran/gelatin hydrogels to deliver rhBMP-2 for periodontal regeneration in dogs (Chen et al., 2007b). The group studied the kinetics of release and demonstrated that by changing the ratio
of components, the rhBMP release could be varied from 18 to more than 28 days (Chen et al., 2007a).

Nanoparticle technology seems definitely one of the most promising approaches for the future of bone tissue engineering, by overcoming some fundamental issues in the methods applied for tissue regeneration such as the insufficient mechanical strength of scaffolds and the lack of stability or bioactivity of growth factors such as BMPs at the defect site (Kim and Fisher, 2007). Major nanotechnology areas of research such as the fabrication of scaffold-nanoparticle composites and the design of nano-patterned materials are some of the areas we found with greatest potential for the delivery of BMP in orthopedic regenerative science.

4. Human clinics and the future of bone tissue engineering

In the western world, an estimated 5-10% of all bone fractures show deficient healing, leading to delayed union or non-union, causing significant morbidity and psychological stress to the patients and bringing elevated costs to society (Westerhuis et al., 2005b). Fortunately, the current advances in bone tissue engineering have led researchers to find new strategies and devices with use of BMPs for accelerate healing of bone tissues on orthopedic field. In fact, by the end of 2007, nearly one million patients worldwide will have been treated with BMPs for diverse bone related problems and diseases (Pecina and Vukicevic, 2007). The clinical uses of BMPs include spinal fusion, treatment of long bone defects and non-unions, dental and periodontal tissue engineering, craniofacial defects and diseases, fracture repair, the improvement of osteointegration with metallic implants, musculoskeletal reconstructive surgery and tendon and ligament reconstruction. There are currently two main collagen-based products containing BMP-2 or BMP-7 that were approved by the Food and Drug Administration (FDA) in the last years for human clinical use: Infuse™ Bone Graft (Medtronic, US; Wyeth, UK) containing rhBMP-2 and Osigraft™ (Stryker Biotech) containing rhBMP-7, known by the designation of OP-1 (osteogenic protein-1). BMP-2 Infuse™ bone graft was approved for certain intervertebral body fusion procedures in 2002, for open tibial fractures in 2004, and for alveolar ridge and sinus augmentations in 2007 (McKay et al., 2007). BMP-7 Osigraft™ was approved for long bone fractures and as alternative to autografts in patients requiring posterolateral lumbar spinal fusion. There has been also an increasing number of trials that provide supporting evidence for the use of rhBMP-7/OP-1 in treatment of open tibial fractures, distal tibial fractures, tibial nonunions, scaphoid nonunions and atrophic long bone nonunions (White et al., 2007).
4.1. Spinal fusion

Spine fusion applications are an important part of clinical trials currently ongoing (Carlisle and Fischgrund, 2005). Spinal fusions consist of nearly half of all grafting surgery. Furthermore, failure rates up to 35% have been reported. The interest is in the use of rhBMPs to accelerate healing in patients with disk degenerative disease, removing the need for autograft harvesting and reducing morbidity. Degenerative disc disease is defined as back pain caused by degeneration of the disc as confirmed by clinical data and symptoms. The common approach is to use a collagen or other carriers soaked with rhBMP and place these within titanium spacers called cages which will be implanted in the spine. There are two types of fusion approaches: posterolateral fusion involving placing the bone graft between the transverse processes in the back of spine and intervertebral body fusion which involves placing the bone graft between the vertebrae in the area occupied by the intervertebral disk. In intervertebral body fusion of lumbar vertebrae, based in the success of previous trials, a prospective study for rhBMP-2 was then performed by McKay and Sandhu involving 279 patients with disk degenerative disease, from which 143 patients received lumbar tapered cages filled with rhBMP-2 and 136 patients received the device filled with autologous bone from iliac crest (McKay and Sandhu, 2002). Since at the conclusion of the study, higher rates of spinal fusion were observed for the rhBMP-2 group and less operative time and morbidity were reported, FDA granted approval for use of rhBMP-2 in treatment of single-level lumbar degenerative disc disease. Posterolateral spinal fusions are common for treating spondylolisthesis but require distinct mechanic and biological properties of the carrier. Boden et al. using rhBMP-2 delivered via a biphasic carrier of tricalcium phosphate and hydroxyapatite, reported achieving complete fusion in all patients treated with rhBMP-2 as compared with the control group (Boden et al., 2002). Although the carrier did not obtained approval from FDA, there are several other trials currently ongoing in humans, such as with collagen sponges (Glassman et al., 2007).

The efficacy of use of rhBMP-7/OP-1 as a replacement for iliac crest autograft was first evaluated by Johnsson et al. showing higher rates of fusion in posterolateral spinal fusions with application of rhBMP-7 (Johnsson et al., 2002), which were confirmed by subsequent work (Vaccaro et al., 2004; Kanayama et al., 2006). With no adverse effects, OP-1 was considered as a viable alternative to autograft and, as a result, FDA gave approval of rhBMP-7 for patients who have failed a posterolateral fusion and are at risk for repeated pseudarthrosis. At the present, there is little focus on human trials involving either the use of other BMPs or with the use of natural polymers as delivery carriers, but one is to expect that these options would be soon evaluated clinically, in the next years, considering the current state of biomaterials research.
4.2. Long bone fractures

In the most cases where rhBMPs are applied to fractures, these consist in non-unions of long bones. It is estimated as an example that in the UK, 42% of these fractures are of tibias, 20% are femurs and the rest are of other bones (Giannoudis and Tzioupis, 2005). RhBMP-2 has received approval of FDA for use in treating open tibial fractures. Initially, Govender et al. performed a randomized trial with 450 patients having open tibial fractures. Patients were randomized to receive different doses of rhBMP-2: 0.75 mg/ml (total dose of 6 mg), 1.50 mg/ml (total dose of 12 mg) or no rhBMP-2, in collagen sponges. After 12 months, analysis showed accelerated healing and reduced infection with increasing dosing of rhBMP-2 (Govender et al., 2002). There are numerous studies in the literature suggesting that rhBMP-7/OP-1 is also a safe and effective alternative for treatment of diverse long-bone fractures and non-unions. In tibial non-unions, a trial that led to multiple regulatory approvals worldwide, concluded that OP-1 delivered in a collagen sponge, was a safe and effective alternative to bone grafting (Friedlaender et al., 2001). In scaphoid nonunions, a study led to conclude that OP-1 could allow successful use of allograft, eliminating the donor site morbidity of an iliac crest autograft (Bilic et al., 2006). In another curious trial, McKee and colleagues assessed the efficacy of OP-1 on treating diverse long bone non-unions on 62 patients that failed previous autograft operations. The involved bones included 16 tibiae, 18 claviculae, 11 humeri, 10 femora, 4 ulnae and 3 radii. At the end of the study, 54 of 61 non-unions (89%) had healed, pointing that rhBMP-7 was an effective treatment (White et al., 2007).

4.3. Dental Tissue Engineering

In periodontal and dental tissue engineering, rhBMPs find their place in inducing pulp stem cells to differentiate into odontoblasts and promoting regeneration of pulp and teeth. Since the pulp, is an organ known to have tremendous regenerative abilities, during the last years tissue engineering has been considered as a promising approach for diverse clinical cases such as caris, pulpitis, and apical periodontitis (Nakashima and Akamine, 2005). Particularly interesting is the fact that human pulp stem cells have self-renewal ability and that tubular dentil is formed after the transplantation of these stem cells with hydroxyapatite power in mice (Gronthos et al., 2002). Recombinant BMPs have been noted to induce dentin formation in vivo when delivered with a collagen scaffold (Nakashima, 1994). The ultimate goal in dental tissue engineering using BMPs is in achieving a complete restoration of physiological, structural and mechanical integrity of the native dentil-pulp complex, including nerve and vascular regeneration (Nakashima and Akamine, 2005).
4.4. Future challenges, a global perspective

Bone repair and regeneration with BMPs are ushering in a new era in orthopedics. The past 10 years have seen practical demonstration of bone repair in a series of animal studies and subsequently in clinical trials. The expected value of BMPs in the treatment of bone defects, spinal fusion applications and other types of related applications is enormous. Extensive research in preclinical models has led to the approval of restricted use for human trials. However, despite the significant evidence of potential for bone healing demonstrated in animal models, future clinical investigations will be needed to better define variables such as dose, scaffold and route of administration. The impressive results of animal models are difficult to replicate in humans. It is unclear why these differences occur. Some insight is provided by the clear species-specific dose response ranging from 25 μg/ml in rodents, to 50 μg/ml in dogs, 100 μg/ml in non-human primates and 800 μg/ml in humans (Luginbuehl et al., 2004), in the case of BMP-2. The recruitment of bone precursor cells and bone turnover may occur differently in rodents, small animals and large mammals. Likewise, the dosing may not be optimal yet. In fact, the concentrations of BMP in use are supraphysiological and of a million times factor higher (milligrams in assays as compared to nanogram range in vivo). BMP inhibitors such as noggin or sclerotin which are upregulated by BMP presence may be interfering and providing a negative feedback effect on the body healing mechanisms (Westerhuis et al., 2005a). Understanding of the regulation between BMPs and BMP-inhibitors might be a key issue. Moreover, different fractures may require different dosage (Schmitt et al., 1999). Critical issues to consider include the potential risk of BMPs inducing heterotopic bone formation, especially when implanted adjacent to neural tissues (Paramore et al., 1999), and the serious issue of reported antibody formation noted in up to 38% of patients in some trials with BMPs (Walker and Wright, 2002).

Clearly the use of BMPs in orthopedics is still in its early days, but the latest trials in humans suggest that an exciting and promising future will unfold in the development of novel tissue engineering products for a wide range of clinical situations, with the use of BMPs. To date, clinical trials have focused mostly in rhBMP-2 and -7 and with the use of collagen as delivery materials. However, given the intricate network of molecules interplaying during bone regeneration, it is possible that a “cocktail” of different BMPs with simultaneous or sequential release, would be the most desirable approach to clinical uses instead of a single stimulus or molecule (Hadjiargyrou et al., 2002). Raiche and Puleo have already explored the sequential release of rhBMP-2 in combination with insulin-like growth factor-1 (IGF-1) (Raiche and Puleo, 2004a, b). However, the development of such cocktail for clinical cases may come across with difficulties since the commercial rights of the two currently approved rhBMPs are for restricted
use and owned by different companies (Westerhuis et al., 2005b). Nevertheless, in near future, the emergent advances with recombinant production of BMPs (Klösch et al., 2005; Schmoekel et al., 2005b; Bessa et al., 2008) will aid researchers in obtaining larger amounts of bioactive rhBMPs which could be used for tissue engineering research and development of novel products.

With the excitement over the potential of other natural origin polymers as novel delivery systems for BMPs, there is little doubt that these will also find relevant places in regenerative medicine of bone and traumatology and may be soon approaching clinical trials in humans. Diverse natural origin polymers have shown promising success for bone tissue engineering such as fibrin, hyaluronic acid, chitosan, silk fibroin and starch-based composites. Furthermore, these overcome limitations and disadvantages from the use of synthetic polymers and the risks of disease transmission inherent to the use of collagen from bovine sources. The recent advances in biomaterials science will certainly boost the number of tissue engineering approaches for the healing of bone with the use of BMPs. Novel strategies will possibly involve the specific targeting of BMPs, in injectable systems and stimulus-responsible hydrogels, the use of nano-scale patterning or encapsulated particles or with the use of molecules combined with the BMP mimicking the extracellular matrix, all of which allow restricted and site-specific delivery of these growth factors. Additionally, the design of 3D specific-architecture scaffolds by methods such as rapid prototyping or design of bilayered scaffolds surely ensures that the carrier for delivering the BMP will mimic closely the bone structure. Guided-tissue engineering delivery systems, which would not only deliver BMPs but angiogenic factors, would, for instance, prompt the recruitment and distribution of blood-vessels precursor cells which is necessary for formation of mature bone. Finally, the use of Ca-P cements and biomimetic coatings is a very promising approach since it furthers mimics the bone mineral makeup and aids in retaining the BMP and improving the tissue-material integration. The expanding variety of options for biomedical usage of BMPs, gives the promise that the future of clinical regenerative medicine and that of BMPs, particularly for bone applications, will be certainly be a bright one in the coming decades for millions of people.
Chapter II. Introduction. Bone morphogenetic proteins in tissue engineering: BMP delivery

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Chapter III. Materials and Methods
Chapter III

Materials and Methods

This chapter intends to give an overview on the experimental work behind the studies presented in this dissertation. All the groundwork that support the achieved results (presented in Sections 3 and 4) will be herein described in detail. This is expected to leave the reader with a more clear vision of the purpose of each studied subject and of the way that these can be correlated.

1. Recombinant technology for cloning and expression of proteins

1.1. BL21 DE3 strain *Escherichia coli* expression system

Because of its long history of laboratory culture and ease of manipulation, *Escherichia coli* plays nowadays an important role in modern biological engineering and industrial microbiology. There are other expression systems for recombinant production of proteins such as mammalian cells or insect cells but these involved more sophisticated culturing conditions and significantly lower yields of expression of the recombinant protein. Using the recent advances in protein production in bacteria, it is possible to express proteins in large amounts and refold these into their native and bioactive structure.

BL21(DE3) strain is the most widely used host background for protein expression and has the advantage of being deficient in both *lon* and *ompT* proteases which could degrade protein during purification (Studier 1991). The BL21(DE3) strain has a high expression level of recombinant proteins, induced upon the addition of isopropyl-1-thio-β-galactopyranoside (IPTG). IPTG activates the T7 RNA polymerase *lac* promoter and consequently upregulates the expression of the cloned gene.

1.2. pET-25b expression system

The pET System is a powerful expression vector developed for the cloning and expression of recombinant proteins in *Escherichia coli*, based on the T7 promoter-driven system originally developed by Studier and colleagues (Studier and Moffatt 1986; Studier et al. 1990). The pET-25b vector carries an N-terminal pelB signal sequence for potential secretion of protein into the periplasm space, plus an optional C-terminal HSV-Tag and the 6x-histidine-Tag (Figure 1). The use of pET-25b vector provides a suitable strategy to express recombinant proteins that require disulfide-bond formation for full bioactivity, such in the case of BMPs.
The recombinant proteins are induced for export into the periplasm space, which is an environment favorable for disulfide bond formation, critical for the folding of dimeric BMPs.

In addition, pET-25b vector allows the purification of protein by affinity chromatography, due to the presence of a C-terminal polyhistidine-tag. The polyhistidine-tags are an amino acid motif in proteins that consists of at least six histidine (His) residues, often at the N- or the C-terminus of the protein, and commonly used for affinity purification of the recombinant protein. Please refer to section 3.3.1. of the present chapter for more information on the histidine-tag affinity purification.

2. Cloning of bone morphogenetic proteins

2.1 Obtaining bone morphogenetic cDNA (BMP mature domain)

Different cDNA sources were researched for cloning of bioactive domains of the different human BMPs. A common source is to clone the BMP corresponding DNA from an human cDNA library from cells or tissues that express BMPs. However, since it is an expensive approach, we opt to clone the DNA corresponding to the bioactive domain of BMPs directly from the human genome, since for the different BMPs used in this thesis, all their bioactive domains are located solely in one exon. Bacteria clones were acquired from Sanger Institute (Cambridge, US), which contained human DNA encoding the different human BMP genes. From this template, PCR was performed, as a way of cloning the DNA corresponding to the bioactive domain of the BMPs into the pET-25b vector and expressing the different recombinant proteins.
Figure 1. Vector pET-25b was used for the expression of different human BMPs in *E. coli*. In the representation above, the vector restriction sites (A) and multiple cloning site (coding strand transcribed by T7 RNA polymerase) (B), are shown.
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Figure 2. Schematic representation of cloning of human BMP-2 in pET-25b vector, showing the N-terminal histidine tag of vector, and pelB signal sequence for secretion into periplasm. The signalling sequence is proteolytically cleaved in periplasm of Escherichia coli. The (natural-occurring) heparin binding domain of BMP-2 is indicated in dark green color. The same methodology was performed for cloning of human BMP-4, BMP-9, BMP-10, BMP-11/GDF-11 and BMP-14/GDF-5. BamHI and XhoI restriction sites, used for cloning, are indicated.

2.2. Molecular biology tools used for cloning of BMPs

2.2.1. Designing primers for bone morphogenetic protein cloning

Primers were designed for amplifying the gene sequence coding for the mature part of the BMPs and flanked with restriction sites (BamHI and XhoI) allowing for insertion into the cloning and expression vectors (Figure 2). Primers were designed accordingly to a set of parameters well defined on the literature (such as nucleotide length, GC nucleotide percentage, and several other rules). Primer efficiency was first investigated by performing in silico PCR with Amplify and Primer 3 software tools.

2.2.2. Cloning of BMP cDNA by polymerase chain reaction (PCR)

Amplification by PCR of rhBMPs was performed upon the aforementioned cDNA, in section 2.1. In a PCR tube, 1 μg DNA was mixed with 0.5 μl dNTPs 10 mM, 2 μl MgCl₂ 50 mM, 1 μl of each primer (detailed information in chapters IV and V) of a 10 μM stock, 1 μl of proofreading accuzyme polymerase enzyme, 2.5 μl enzyme buffer (10x) and rest of ultrapure water for a total 25 μl.

Conditions regarding the (PCR) amplification cycles, were the following.

1) 94°C – 2 min
2) 30 cycles (94°C – 30 sec, Ta – 45 sec, 72°C – 1 min)
3) 72°C – 5min
4) 4°C – ad infinitium
The primer annealing temperature (Ta) was calculated in primer manufacturer instructions. After the PCR, a volume of 1-2 μl was run in agarose gel electrophoresis for confirmation of DNA integrity and size (1% agarose gel stained with ethidium bromide, 6 μl per 100 TAE buffer). Amplified BMP DNA was then purified with GenElute PCR DNA Purification Kit (Sigma), which is a critical step for efficient ligation to cloning/expression plasmids.

2.2.3. Restriction of DNA and ligation into expression vector

Restrictions of both amplified DNA and of vector are required for insertion of the gene of interest to the cloning vector. They allow the complementary between insert and plasmid, for ligation. The restrictions sites of BamHI and XhoI were selected for ligation, to discard the maximum of amino acids residues, upstream and downstream of BMP gene sequence, that could cause any interference in the folding and bioactivity of the protein. The restriction reaction was performed by incubating 10 μl purified DNA, 6 μl ultrapure water, 2 μl enzyme buffer and 1 μl of each restriction enzyme, at 37ºC (as indicated by the manufacturer) for 5 h. Following the restriction, the pET-25b-BMP2 vector was diphosphorylated to prevent the re-cycling of plasmid, by incubating the mixture with 1μl of SAP enzyme (30 min at 37ºC) and stopping the reaction (15 min, 65ºC). Before ligation, the DNA integrity was verified on gel electrophoresis, and DNA isolated using QIAquick Gel Extraction Kit (Qiagen). Ligation was performed by incubating 1 μl of plasmid, 7 μl of BMP DNA, 1 μl enzyme buffer and 1 μl ligation enzyme (Roche), overnight at 4ºC.

2.2.4. Transformation of XL1BLUE and BL21DE3 Escherichia coli strains

For cloning of plasmid, XL1BLUE or TOP10 Escherichia coli strains were used. For expression of protein, BL21DE3 E. coli strain was generally used (although Origami, Tuner and BLR strains were also investigated, data not shown). For transformation of XL1BLUE, bacteria stocks were thawed in ice (competent cells) and 10 μl ligation product were added and incubated for 30 min. Then, cells were incubated for 30 sec at 42ºC with agitation and immediately returned to ice, for more 10 min. After this, 0.8 ml of SOC medium (Super Optimal broth with Catabolite repression) was added and cells grown at 37ºC with vigorous agitation for 1h. Cells were then centrifuged, resuspended in a small volume (50 μl) and plated into LB culture plates which contained 50 μg/ ml ampicillin. Negative controls included bacteria transformed with a plasmid with no BMP gene and cells having no resistance to the antibiotic. E. coli was grown at 37ºC overnight. The following day, colonies were selected, grown overnight in new plates and
plasmids were isolated with use of GenElute Plasmid Minipreps Kit (Sigma, US) (according to manufacturer instructions).

The extracted plasmid was then digested with specific restriction enzymes and run in agarose gel electrophoresis for screening which colonies had the correct insertion of the BMP genes. The integrity of the gene sequences was verified by DNA sequencing (Sanger et al. 1977) using ABI PRISM 310 Genetic Analyzer. The pET-25b and pET-25b/BMP vectors were then used for transformation of BL21DE3 E. coli strain, for expressing the recombinant BMPs. Cryotubes were prepared using collected biomass in 1.5 ml 30% glycerol for long-term storage of bacteria at -80ºC. Plasmids were stored at -20ºC. All work was performed under sterile conditions.

3. Expression, folding and purification of bone morphogenetic proteins

3.1 Expression of rhBMPs

3.1.1. Expression of protein in BL21DE3 Escherichia coli strain

Expression of recombinant BMPs was performed by using BL21DE3 cells which are indicated for the expression of recombinant proteins from pET vectors. Transformed BL21DE3 bacteria with pET-25b/rhBMP and pET-25b were inoculated in 50-200 ml Luria Bertani (LB) medium with 50 μg ampicillin/ml and incubated at 37 ºC (180 rpm), until an OD_{600} of 0.6-1.0 was obtained. To induce the recombinant protein expression IPTG 1 mM was then added to the culture media, and the temperature lowered to 25 ºC during 24 h. Biomass was collected by centrifugation (4000 g, 20 min, 4 ºC), washed once with PBS and stored at -20 ºC.

3.1.2. Analysis of protein expression using SDS-PAGE

Protein was isolated by resuspending frozen bacteria in phosphate buffer saline (or other lyses buffer), with addition of a proteases inhibitor (Complete Mini EDTA-free, Roche), and performing cell lyses by incubating ultrasonication of samples for 4 times for 15 s with intervals of 1 min, in ice. Supernatant and pellet fractions were collected by centrifugation (4000 g, 30 min, 4ºC), for SDS-PAGE analysis. SDS-PAGE was applied for a qualitative and semi-quantitative analysis of the protein, at different stages of BMP production: protein expression, solubilization and folding, purification and storage.

For SDS-PAGE, protein concentrations of 0.1-1 mg/ml were used. Protein samples were mixed with SDS-PAGE loading buffer and heated at 95ºC for 5 min. Samples were separated by using hand-cast 10–12% acrylamide/bisacrylamide SDS-PAGE gels, at 15 mA, for 1 h, in
reducing or non-reducing conditions. Reducing conditions included the use of loading buffer with 5% (v/v) β-mercaptoethanol as a reducing agent. Coomassie Blue-R or silver staining were used for the visualization and staining of gels. Gels were analyzed with Gel Pro Analyzer software (Media Cybernetics), for the estimation of protein molecular weight and concentration.

3.1.3. Analysis of protein expression using Western-blot

For Western-blot, protein concentrations of 1-10 µg/ml were used. Protein samples were mixed with SDS-PAGE loading buffer and heated at 95 °C for 5 min. Samples were separated by using hand-cast 10-12 % SDS–PAGE at constant 15 mA, for 1 h. Gels were then fixed for 5 min in transfer buffer (20mM Tris-HCl, 20mM glycine, 20% ethanol) and samples were electro-transferred to a nitrocellulose membrane at 100 V for 60 min (wet Western-blot transfer) or 10V, 400 mA, for 30min (semi-dry Western-blot transfer). The membrane was washed once with PBS for 10 min and blocked with 2% (w/v) BSA in PBS-T buffer (PBS, 0.05% Tween 20) for 2h at room temperature. After this, the membrane was washed once with PBS-T buffer and incubated with peroxidase conjugated anti-His antibody (Sigma) diluted 1:2000 or rabbit anti-human BMP-2 polyclonal primary antibody (AbD Serotec) diluted 1:5000 followed by mouse anti-rabbit peroxidase conjugated secondary antibody (Amersham) diluted 1:30000. Prior to colorimetric development, the membrane was washed 3x with PBS and image detection was performed with ChemiDoc XRS and Quantity One software (BioRad).

3.1.4 Optimization of protein expression levels

Protein expression was optimized for different variables that affected the amounts of produced protein: temperature of induction and expression, time length of expression and the influence of codon usage. First, the expression of rhBMP-2 was performed for 24h at 25ºC, 18ºC and 4ºC. By analyzing crude lysates (pellet fraction) using SDS-PAGE, rhBMP-2 expression was found to be higher at 25ºC (Figure 3). Following this, the amount of expressed BMP-2 was tested along different times of production, and optimal recovery of protein was found after a period of 24-48 h of expression (Figure 4). Finally, the cloning of a codon-usage optimized BMP-2 gene sequence did not affected significantly the expression of protein, compared to the native sequence derived from the BMP-2 cDNA (Figure 5).
**Figure 3.** SDS-PAGE of samples of crude lysates (pellet fraction) with different temperature used for expression of rhBMP-2. Respectively, A-G correspond to the negative control, BMP-2 variant I (25°C), BMP-2 variant II (25°C), BMP-2 variant I (18°C), BMP-2 variant II (18°C), BMP-2 variant I (4°C), BMP-2 variant II (4°C). BMP-2 monomer and dimer are marked (18 kDa, 36 kDa). Marker: 116 kDa, 97 kDa, 66 kDa, 45 kDa, 22 kDa and 14 kDa. Expression of BMP-2 was significantly higher at 25°C. Expression was performed for 24 h. Please refer to section 3.2.5. for rhBMP-2 variants.

**Figure 4.** SDS-PAGE of samples of crude lysates (pellet fraction) with different time periods for expression of rhBMP-2. Respectively, A-H correspond to the following times of production: A (0h), B (4h), C (8h), D (16h), E (22h), F (24h), G (2 days) and H (4 days). Temperature of expression was 25°C. BMP-2 monomer and dimer are marked (18 kDa, 36 kDa). Marker: 200 kDa, 116 kDa, 97 kDa, 66 kDa, 45 kDa, 22 kDa and 14 kDa. Expression of BMP-2 peaked after 24 -48 h.

**Figure 5.** SDS-PAGE of samples of crude lysates (pellet fraction) with rhBMP-2 produced with native gene sequence (1), and a codon usage optimized sequence (2). The use of optimized codon use does not result in significant changes in the expression of rhBMP-2.
3.1.5. Bioreactor scale-up protein production

Scale-up expression of BMPs was performed using a 2 l Bioflo110 bioreactor (New Brunswick Scientific, USA) (Figure 6). Pre-cultures were transferred to the pre-autoclaved bioreactor which contained LB media with 50 μg ampicillin/ml. *E. coli* were grown at 37ºC until an OD<sub>600</sub> of 0.6-1.0. At this time point, 1 mM IPTG was added and temperature lowered to 25ºC to induce the expression of the recombinant BMPs. During the entire process of fermentation, aeration was kept at 2 l/min and an agitation cascade of 300-1000 rpm was selected in order to maintain minimum dissolved oxygen levels at 30% of saturation. The pH 7.4 was kept constant using NH<sub>4</sub>OH (30%, v/v) and H<sub>3</sub>PO<sub>4</sub> (5 M) which were automatically pumped into the bioreactor vessel. In the end, the biomass was collected by centrifugation (4000 g, 20 min, 4ºC), washed once with PBS and stored at -20ºC.

Figure 6. Use of a bioreactor allowed producing large-scale amounts of recombinant BMPs.

3.2. Recovery of rhBMPs

3.2.1. Protein folding and solubilization

The folding of proteins is dependent on several conditions, such as buffer composition, ionic strength, denaturant and protein concentration, pH and temperature (Vallejo and Rinas 2004). Since the recombinant protein was initially obtained in the pellet fraction of bacteria cell lysate, the protein solubilization was first analyzed in the presence of varied buffers and conditions (e.g. Triton, Tween, sodium lauryl sulphate, L-arginine, and urea, and different pH). The BMP-2 was found to be recovered best with phosphate buffer saline containing L-arginine (data not shown), a common aggregation-suppressor that enhances the solubilization and the refolding of proteins (Vallejo and Rinas 2004). Different concentrations of L-arginine were used,
with different pH, for optimizing the solubilization and folding of BMP-2 (Table 1, Table 2). A concentration of 0.5-0.7M and a pH 8.0-11.0 was found to be optimal and was generally used for this step.

**Table 1.** Effect of different L-arginine concentrations on the recovery of total protein of crude extracts, from which rhBMP-2 is a main fraction. Similar conditions of dilution of protein and pH 8.5 were used for these recovery assays. Protein was quantified using the Bradford method. The underlined conditions were selected.

<table>
<thead>
<tr>
<th>[L-Arginine] (M)</th>
<th>Recovery of total protein</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>32%</td>
</tr>
<tr>
<td>0.3</td>
<td>44%</td>
</tr>
<tr>
<td><strong>0.5</strong></td>
<td><strong>57%</strong></td>
</tr>
<tr>
<td>0.7</td>
<td>63%</td>
</tr>
<tr>
<td>1.0</td>
<td>58%</td>
</tr>
</tbody>
</table>

**Table 2.** Effect of different pH on the recovery of total protein of crude extracts, from which rhBMP-2 is a main fraction. Similar conditions of dilution of protein and a fixed concentration of 0.5 M L-arginine were used for these recovery assays. Protein was quantified using the Bradford method. The underlined conditions were selected.

<table>
<thead>
<tr>
<th>pH</th>
<th>Recovery of total protein</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>5%</td>
</tr>
<tr>
<td>5</td>
<td>6%</td>
</tr>
<tr>
<td>7</td>
<td>35%</td>
</tr>
<tr>
<td>8</td>
<td>48%</td>
</tr>
<tr>
<td><strong>8.5</strong></td>
<td><strong>57%</strong></td>
</tr>
<tr>
<td>9</td>
<td>68%</td>
</tr>
<tr>
<td>12</td>
<td>91%</td>
</tr>
</tbody>
</table>
3.2.2. Recovery of rhBMP-2 from periplasm

Recombinant proteins expressed with pET-25, are expected to be secreted to the periplasmatic space of bacteria where cellular environment allows for the formation of dimers (Messens and Collet 2006). Therefore, as a way of establishing an easier method for the recovery of BMPs, extraction of protein from the periplasm was performed by osmotic shock:

1) Unfrozen cells from bacteria fermentations were collected and washed with PBS.
2) Osmotic solution I (2mM CaCl$_2$, 20mM Tris-HCl, 20% sucrose w/v) was added and cells incubated 10min in ice.
3) The solution was centrifuged (10min, 4ºC, 5000 g) and supernatant discarded.
4) Add slowly osmotic solution II (2mM CaCl$_2$, 20mM Tris-HCl) and cells incubated 10min in ice.
5) The solution was centrifuged (10min, 4ºC, 5000 g) and supernatant containing the periplasm protein was collected and analyzed by SDS-PAGE.

However, no protein was recovered from the periplasm in significant amounts (data not shown), even when different conditions of expression were attempted (different temperatures, different time periods, or induction with different IPTG concentrations). As a consequence, the recovery of the recombinant BMPs was solely performed with the use of L-arginine as stated in 3.2.1.

3.3 Purification of rhBMPs

3.3.1. Purification by nickel affinity chromatography

Recombinant human BMPs were purified by nickel affinity chromatography. Briefly, up to 40 mg of filtered protein were loaded into a pre-equilibrated nickel HisTrap column (Amersham) (Figure 7), with phosphate buffer (40mM sodium phosphate, 0.5 M NaCl) containing 20 mM imidazole. Binding of protein to the column was optimal at pH 9.0 (data not shown). The column was then washed extensively with phosphate buffer containing 80 mM imidazole, with 60-100 ml. The recombinant BMPs were eluted with 15-20 ml of sodium phosphate buffer containing 400 mM imidazole, and stored at 4ºC for short-term use, or desalted and lyophilized.
Figure 7. Nickel affinity chromatography was used for the purification of histidine-tagged recombinant human BMPs.

3.3.2. Buffer exchange and storage

Desalting is an important step prior to bioactivity assays to remove buffer salts used during the folding and purification of recombinant proteins. The removal of salts or buffer exchange was first investigated using three different methods: dialysis tubing, ultracentrifugation columns or chromatographic desalting columns. The efficacy of each method was determined by assessing the amount of salts in solution using salt conductivity, and by measuring the loss of protein during buffer exchange (data not shown). It was found that with desalting columns (Amersham, UK), the recovery of BMP-2 was much higher than with ultracentrifugation columns (Vivascience, Germany) or dialysis cassettes (Pierce, US), while providing efficient salt removal. Therefore, this method was used for the desalting of rhBMPs.

For short-term storage, the BMPs were buffer exchanged to sodium phosphate buffer with no imidazole, prior to the bioactivity assays. These solutions of protein were stored at 4ºC for up to one month. For long-term storage, lyophilization or freeze storage were used. For lyophilization, the samples were desalted, quickly frozen in liquid N₂ and freeze dried. Lyophilized protein was resuspended in PBS, at concentrations up to 0.5 mg/ml. Optionally, bovine serum albumin (1%) was added to prevent degradation and L-arginine (0.5 M) was added to aid in the solubilization. Freeze-dried protein was stored at 4ºC or at room temperature, in dry conditions.
3.4 Stability of rhBMPs

The stability of rhBMP-2 during storage was studied in different conditions. Samples of purified BMP-2 were incubated in different buffers, concentrations and pH and analyzed after a period of two weeks, 4°C. Silver stained SDS-PAGE revealed that the protein was stable when stored in phosphate buffer, at pH 7.0-9.0 (Figure 8). At more acidic pH, the BMP-2 tended to precipitate, while at higher pH the protein showed degradation. In addition, the protein was also more stable when stored at concentration of 0.01-0.5 mg/ml (data not shown).

For additional information, please refer to Attach I – supplementary data.

![Figure 8. Silver stained SDS-PAGE showing the stability of BMP-2 in phosphate buffer, at different pH. The protein was stored for a period of two weeks at 4°C. Figure shows protein before and after centrifugation to remove precipitates. Lanes: A) protein stored in pH 5.0; B) protein stored in pH 7.0; C) protein stored in pH 9.0; D) protein stored in pH 11.0. BMP-2 seems to be more stable and soluble when stored at pH 7.0-9.0, in phosphate buffer (B and C).]

4. Assessing the bioactivity of BMPs

4.1. Bone differentiation and BMPs

During bone differentiation, bone morphogenetic proteins bind to serine-threonine cell receptors (Bessa et al. 2008), triggering the expression of genes involved in osteogenic differentiation.

One common assay to determine the biological activity of BMPs is to determine the enzymatic activity of alkaline phosphatase, which is a non-specific but highly upregulated gene during the differentiation to osteogenic phenotype, prior to bone mineralization (Gundberg 2000). Another method is to measure the induction of the expression of mRNA of BMP-target genes by semi-quantitative RT-PCR, having the advantage of providing more information about the extent and detail of osteogenic differentiation, at a genetic level. Different genes were assessed, including early markers such as transcriptional factors Osterix and Runx2, signaling molecules Smad-1, -5 or -3, and mineralization-related proteins such as Bone sialoprotein, Osteopontin, ALP and Osteocalcin. Finally, by performing mineralization calcium-specific
stainings such as Alizarin red or Von Kossa, it is possible to observe the the extent by which BMPs induce the formation of extracellular matrix of calcium phosphate, which is typical of bone formation. The morphological observation of cells was performed using a Axiovert10 (Zeiss) optical microscope with a coupled Coolpix950 (Nikon) camera.

4.2. Human adult stem cells

These cells have been shown to differentiate into osteoblasts in vitro when cultured with the appropriate supplements and factors such as BMPs (Skillington et al. 2002) To evaluate if this same behavior could still be observed when cultured with the recombinant BMPs, these cells were isolated and cultured according to the method described by Zuk et al. (2001). Human subcutaneous adipose tissue samples were obtained from lipoaspiration procedure. The tissue was firstly washed with PBS, 5 mg / 500 ml Ciprobay 200 (Bayer) and 10% Antibiotic-Antimycotic (Gibco, UK), and then digested with 0.2 % collagenase type I A (Sigma Aldrich, Germany) in PBS for 60 min at 37°C, in gentle stirring. The digested tissue was filtered using a 100 μm filter mesh (Sigma Aldrich, Germany) and centrifuged at 800 g for 7 min at 20°C and the supernatant eliminated. After performing the enzymatic digestion as above described, the cells were seeded in 75 cm² culture flasks with Basal Medium composed by DMEM (Sigma Aldrich, Germany), 10% FBS (Gibco, UK) and 1% Antibiotic-Antimycotic (Gibco, UK).

4.3. C2C12 cell line

C2C12 is a mice myoblast precursor cell line that differentiates into osteoblasts when cultured in presence of BMPs (Katagiri et al. 1994). C2C12 cells were originally obtained by Yaffe and Saxel (1977) through selective serial passage of myoblasts cultured from the thigh muscle of C3H mice 70 h after a crush injury. Due to its constant reliability, rapid growth and ease of manipulation, this was our standard choice for assessing the bioactivity of BMPs. In the present thesis, C2C12 were cultured in Dulbecco’s modified Eagle’s medium (DMEM) with 1% (v/v) fetal calf serum and 2mM L-glutamine, at 37ºC with 5% CO₂ in a humidified environment. Low serum concentration was used (1 %) to inhibit cellular proliferation. Since C2C12 cells became rapidly confluent, while differentiating into myoblasts, this occurrence was avoided, during expansion of the cell cultures. Thus, cells were generally seeded at lower densities, between 10⁴ to 10⁵ cells per ml, in a 24-well plate. Culture medium and BMPs were replaced every three to five days.
4.4. MTS cell viability assay

Short term cell viability screening, in response to purified BMPs was performed with the use of MTS test. The MTS test is a biochemical test widely used to assess cytotoxicity by measuring cell viability and proliferation in a qualitative way (Slater et al. 1963; Zeltinger et al. 2001). This biochemical test is based in the reduction of MTS salt (3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium), by cell dehydrogenase mitochondrial enzymes, into a formazan product that is soluble in tissue culture medium. The salt absorbs at a wavelength of 490 nm and since only living cells have the capability of metabolize the MTS, it gives a measurement of the viable cells, and an inverse relationship of toxicity to the cells can be assumed (Mosmann 1983). Briefly, cells were incubated with a tetrazolium salt solution for 2h at 37 °C and then absorbance read at 490 nm (Salgado et al. 2002). All the samples were tested in triplicate for at least two independent experiments. The results are expressed as a percentage of the control (scored as 100% viability) as mean ± standard deviation.

4.5. Alkaline phosphatase enzymatic activity

Alkaline phosphatase (ALP) hydrolyses phosphate esters and is involved in the initial processes of bone extracellular matrix mineralization. Consequently, it is widely regarded as an indicator of osteoblastic differentiation and analyzed by tests based on the hydrolysis of substrates containing phosphate. The enzymatic activity of ALP was measured by the conversion of p-nitrophenyl phosphate (pNPP) into p-nitrophenol (pNP) (Salgado et al. 2002). Cells were collected at each time point \( n = 3 \) and frozen in 50 μl phosphate buffer saline at -80 °C until further analysis. Then, cells were lysed by mixing the sample with 50 μl of PBS with 0.2 % Triton, for 1h at room temperature. The enzyme reaction was set up by mixing the sample with 50 μl of ALP-buffer containing 0.5 M 2-amino-2-methyl-1-propanol and 2 mM MgCl₂ (pH 10.3) and 7 mg/ml of pNPP. The solution was incubated at room temperature for 20 min and the reaction was then stopped by a solution containing 2 M NaOH and 0.2 mM EDTA in distilled water. The optical density was determined at 405 nm, with a reference filter at 620 nm. A standard curve was made using pNP solutions. The results were expressed in nmol of pNP/min/mg total protein. All samples were tested in triplicate, from at least two independent experiments. Assaying the membrane-bound ALP is much more accurate since it only quantifies the enzyme present in cells viable at the moment of the test and not the secreted enzyme which accumulates in the culture medium.
4.6. RNA extraction and RT-PCR analysis of osteoblastic markers

For RT-PCR, mRNA of cells was extracted after a specific time point in culture with TriReagent RNA Isolation Reagent (Sigma-Aldrich) from triplicate 24-well assays following the supplier instructions. Briefly, 500 μl of TriZol were added to each sample containing approx. 10^5 cells per ml. After an incubation of 5 minutes, additional 160 μl of chloroform (Sigma Aldrich, Germany) were added; the samples were then incubated for 15 min at 4 °C and centrifuged at the same temperature and 10000 g for 15 min. After the centrifugation, the aqueous part was collected and an equal part of isopropanol (Sigma Aldrich, Germany) was added. After an incubation of 2 hours at -20 °C the samples were washed in ethanol, centrifuged at 4 °C and 7000 g for 5 min and resuspended in 12 μl of water RNase/DNase free (Gibco, UK). The samples were then quantified using a NanoDrop ND-1000 Spectrophotometer (NanoDrop Technologies, USA) and cDNA was synthesized from 4 μg total RNA (A_{260nm} / A_{280nm} ratio between 1.7-2.0) with oligo dT primer (MWG Biotech AG) and AMV reverse transcriptase (Promega). The polymerase chain reaction (PCR) was performed with specific primers for osteogenic markers and with the use of HotStar Taq Polymerase (Qiagen). Agarose gels were imaged with Multimage Light Cabinet (Alpha Innotech, USA) and gene expression analyzed with ChemiImage 4400 AlphaEase FC Image Analysis Software (Alpha Innotech, USA) and using polymerase II for normalizing gene expression, as housekeeping marker.

4.7. Mineralization assay: Alizarin Red staining

For both mineralization assays, cell cultures were seeded at low seeding density (3 x 10^4 cells/ ml) and grown for a period of two weeks in the presence of BMPs and with 1% serum DMEM culture media, supplemented with 10 mM β-glycerophosphate and 100 μM ascorbic acid (Sigma, USA). The Alizarin Red mineralization test was performed using the method described by Bodine et al. (1996), with modifications. Cells were washed for three times with PBS (with no Ca^{2+}, Mg^{2+}), and subjected to a fixative 4% v/v solution of formaldehyde in PBS for 30 min. After removal of fixative, cells were washed twice with distilled water and covered with Alizarin Red (AR) 1% solution, followed by gentle agitation in an orbital shaker for 10 min. The AR solution was then removed and the cells washed four times with distilled water. The cells were observed in a Axiovert10 (Zeiss) optical microscope with a coupled Coolpix950 (Nikon) camera.

The mineralization staining was quantified by the method proposed by Gregory et al. (2004). Briefly, the calcium from cells was extracted by incubation of 800 μl acetic acid (10% v/v) to each well, for 30 min in a orbital shaker. Then, the suspensions were transferred to a clean 1.5 ml eppendorf and vortexed for 30 sec; 500 μl of mineral oil was added and the
suspensions heated at 85 °C for 10 min and then cooled in ice for 5 min. The suspensions were centrifuged (20 min, 10000 g), the supernatant carefully removed, and 200 μl ammonium hydroxide (10%) added (until pH ~4.1). The solutions were read for absorbance at 405 nm, in a Spectra III spectrophotometer (SLT, Austria).

5. Production of silk fibroin and elastin particles for BMP delivery

5.1. Development of fibroin microparticles

5.1.1. Isolation of silk fibroin

Silk fibroin was first isolated from Bombyx mori cocoons after a standard degumming protocol, with modifications (Zhang et al. 2007), for the removal of silk sericin. The cocoon shells of silkworm Bombyx mori were degummed twice in boiling solution of 0.5 % Na₂CO₃ for 0.5 h, at 90 °C, and the resulting degummed fiber was subsequently dissolved in a ternary system, i.e. a mixed solution of calcium chloride, ethanol and water (CaCl₂ / C₂H₅OH / H₂O: 1:2:8 mole ratio) at 90°C for 6 h. The solution was then filtered (0.22 μm, Rotilab, Germany), and the solution was extensively dialyzed, using a cellulose semi-permeable (molecular weight cut-off = 12,000 kDa) membrane, against distilled water to remove CaCl₂, smaller molecules and impurities. The silk fibroin was lyophilized and used for the preparation of microparticles.

5.1.2. Microparticle preparation and loading of BMPs

Silk fibroin microparticles were prepared by ethanol precipitation according to the method described by Cao and colleagues (2007), with modifications. A solution of approximately 1% (w/v) fibroin in phosphate buffer saline (PBS, pH 7.4) was prepared (5 ml) and filtered with a 0.22 μm PVDF membrane filter (Rotilab, Germany). Then, 5 μg/ml or 50 μg/ml BMP-2, BMP-9 or BMP-14 were added (approximately 5 or 0.5 μg BMP per mg of fibroin). The solution was incubated at room temperature with stirring (600 rpm). Absolute ethanol was added dropwise to a ratio of 1:2 to the initial volume of silk fibroin, and the solution incubated overnight at -20 °C. Different ratios were also tested for a comparative study of the particles size. The resulting microparticles (Figure 9) were collected by centrifugation at 4000 g, 4°C, for 5 min, washed, and stored at 4°C or lyophilized, after quick-freezing in liquid nitrogen, in a Christ Alpha 1-4 (Linder, Austria) lyophilizer. Sterile conditions were used during the entire procedure.
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5.2. Development of elastin-like nanoparticles

5.2.1. Nanoparticle preparation

The elastin-like polymer, \((VPAVG)_{220}\), consists of 220 repeats of the main monomer VPAVG and was produced by recombinant technology and purified as reported elsewhere (Machado et al. 2009). Particles were produced by resuspending lyophilized polymer in ice-cold phosphate buffer saline (PBS) to a final solution of 0.1%. The solution was then incubated at 37 °C for 30 min. Particles were collected by centrifugation (10000 g, 37 °C, 10 min), and washed once with pre-warmed PBS. Due to its hysteresis behavior, the particles are stable at room temperature and only solubilize when a strong undercooling is achieved (of about 6 to 8 °C). The particles were prepared under sterile conditions, and stored at room temperature.

5.2.2. BMP loading

Elastin nanoparticles were loaded with a mix or single BMP-2 and BMP-14, during the formation of the particles. Growth factor was added at a concentration of 20 µg/ml to the elastin solution. The nanoparticles were produced with the different combinations of the two growth factors:

I) Unloaded particles (no growth factor)

II) BMP-2 encapsulated

III) BMP-14 encapsulated

IV) BMP-2 and BMP-14 encapsulated

Figure 9. Silk fibroin microparticles, for loading and delivery of rhBMPs, were formed after dropwise addition of ethanol and overnight incubation at -20 °C.
5.3. Characterization of the developed particles

5.3.1. Morphological characterization

The particles were morphologically characterized by scanning electron microscopy (SEM). SEM analysis was performed on gold-coated air-dried samples (Agar Sputeroncoater 108, Essex, UK) and using a Philips XL20 microscope (Philips, Eindhoven, NL). Particle size measurements were obtained from different micrographs acquired in the SEM.

5.3.2. Size distribution

Measurements of particle size were performed on freshly prepared microparticle solution by dynamic light scattering (DLS), using a Zetasizer NanoZS (Malvern, UK). Each sample was diluted to an appropriate concentration with filtered (0.2 μm filters Millex-GN, Millipore) ultrapure water. Each analysis lasted 60 s and was performed at 25 ºC.

5.3.3. Water uptake

The hydration degree of silk fibroin and elastin particles was evaluated after immersion into PBS, pH 7.4. All experiments were conducted at 37 ºC, in triplicate. Percentage of water uptake (WU) after each time period of immersion (t) was calculated using the following equation:

\[
WU(\%) = \frac{(Ww-Wd)}{Wd} \times 100
\]

‘Wd’ and ‘Ww’ correspond to the weight of the particles in dry and wet state, respectively.

5.4. Release of BMPs

5.4.1. Determination of loading capacity and encapsulation efficiency

The particle encapsulation efficiency was determined upon their separation from the aqueous preparation medium containing the non-associated protein by centrifugation (10000 g, 10 min). The amount of free BMP-2 was determined in the supernatant by using a BMP-2 sandwich-type ELISA development kit (Eubio, Austria). The amount of free BMP-9 and BMP-14 was estimated by dot blot. The blot intensities were compared with standards of known concentration, using image software analysis (Chemilimage 4400, Alpha Innotech, US). Each sample was assayed in triplicate (n = 3). The particle encapsulation efficiency (E.E.) and BMP loading capacity (L.C) were calculated using the following equations:
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E.E.(%) = \frac{\text{Total BMP amount (μg)} - \text{Free BMP amount (μg)}}{\text{Total BMP amount}} \times 100

L.C.(μg BMP / mg particles) = \frac{\text{Total BMP amount (μg)} - \text{Free BMP amount (μg)}}{\text{Microparticles weight (mg)}}

5.4.2. In vitro release

In vitro release studies were conducted in phosphate buffer saline (PBS), pH 7.4. Particles were incubated in PBS, at 37 ºC, with 200 rpm agitation. At pre-determined time points, a sample from the solution was removed, centrifuged and stored at -20 ºC until measurement. The solution was replaced with a similar volume of fresh buffer. Sterile conditions were used to prevent any contamination of samples.

5.4.3. Quantification of released BMPs

For measurement of BMP-2 concentration, a BMP-2 sandwich-type ELISA kit (Eubio, Austria) was used, following the manufacturer’s instructions. A calibration curve was obtained using standard preparations of BMP-2 of known concentration. For detection of the concentration of BMP-9 and BMP-14, the samples were estimated by dot blot as described before. Each sample was assayed in triplicate \((n = 3)\) and mean ± SD values reported.

5.4.4. Release kinetics models

To study the release kinetics, the data obtained from the in vitro release was treated accordingly to zero order as cumulative amount of drug released vs. time (equation 1), first order as log cumulative percent drug remaining vs. time (equation 2), Higuchi kinetics as cumulative percent drug released vs. square root of time (equation 3) and Korsmeyer kinetics as log cumulative percent drug released vs. log time (equation 4) (Huguchi 1963, Chowdary and Ramesh 1993, Hadjioannou et al. 1993, Bourne 2002):

\begin{align*}
R &= k_1 t \\ 
\log UR &= k_2 t 2.303 \\ 
R &= k_3 t 0.5 \\ 
\log R &= \log k_4 + n \log t
\end{align*}
where R and UR are released and unreleased percentages, respectively, at time (t); \( k_1, k_2, k_3 \) and \( k_4 \) are the rate constants of zero-order, first order, Higuchi and Korsmeyer models, respectively; \( n \) is a exponent that characterizes the mechanism of release.

5.5. *In vitro* bioactivity

When a molecule is incorporated inside a release system, it is vital that its biological activity is maintained throughout loading, incorporation and release. For released BMPs, the maintenance of its bioactivity was measured by the induction of ALP activity in C2C12 cell line. C2C12 cells were incubated with the fibroin or elastin particles containing specific amounts of loaded BMPs. Growth factor was used to the culture media, as a positive control. Alkaline phosphatase (ALP) activity was measured accordingly to described in section 4.5, after 5 days of cell culture. Cell viability was evaluated, as described in section 4.4, by incubating specific amounts of particles into the cell cultures.

5.6. *In vivo* bioactivity (silk fibroin particles)

5.6.1. Rat ectopic bone formation model

The bioactivity of delivered BMP-2 in fibroin particles was assessed *in vivo* using a rat ectopic bone formation model. Male Sprague-Dawley rats (weighting 340-360 g), were purchased from the Institut für Labortiertechnik und -genetik (Himberg, Austria). They were housed in light and temperature-controlled facilities and given food and water *ad libitum*. In three animals, surgery was performed after general anesthesia with an intramuscular injection of 60 mg/kg of body weight (BW) ketamine (Ketavet®, Pharmacia, Erlangen, Germany) and 7.5 mg/kg BW xylazine (Rompun®, Bayer, Leverkusen, Germany). After shaving and disinfecting the dorsum of the animals, four midsagittal incisions were made dorsally in the proximal area of each limb and subcutaneous pockets were created by blunt dissection. The fibroin particles were implanted subcutaneously into each pocket. Three groups were defined: group I (unloaded particles), group II (particles + 5 μg BMP-2) and group III (particles + 12.5 μg BMP-2). The animals were randomized and were implanted with four samples \((n = 4)\) per each group. Only one group was assigned to each animal to avoid the potential interference of released BMP by blood circulation. After placement of the particles, the incisions were closed with sutures, and the wound covered with vapor-permeable spray dressing (Opsite®, Smith & Nephew, London, UK). Analgesia treatment was given post operation, of 0.01 mg/kg BW Buprenorphin. At the end of each implantation period (4 weeks), the animals were sacrificed.
and samples from the tissues surrounding the implantation site were harvested. The animal protocol was approved by the authority of the city government of Vienna.

5.6.2. *In vivo* microCT

Live microCT (vivaCT 75, Scanco Medical AG, Switzerland) was used to examine bone formation in individual rats. MicroCT imaging was performed at weeks 2 and 4 respectively. Rats were anesthetized for 5 min with 2% isoflurane during measurement. No X-ray contrast media was used in this study. Image reconstruction was performed using the built-in 3D visualization software (Scanco Medical AG, Switzerland). The evaluation of newly formed bone and bone density were performed using IPL version v5.06b (Scanco Medical AG, Switzerland).

5.6.3. Histological analysis

Harvested samples were fixed in neutral buffered 10% formalin and embedded in paraffin. Three-micrometer or four-micrometer sections were cut and stained with hematoxylin and eosin (H & E) or Alizarin Red S, respectively, for light microscopy observation. For a detailed analysis of the tissue differentiation, immunohistochemical staining was performed in selected samples. Briefly, paraffin sections were dewaxed with xylene and rehydrated in a graded ethanol series. After enzymatic antigen retrieval with 0.1% Proteinase (Sigma Aldrich, USA) sections were incubated with an Osteocalcin antibody (Santa Cruz Biotechnology, USA) in a humidified chamber at 4 ºC overnight. After rinsing, sections were incubated with the EnVision™ system (Dako, Denmark) at room temperature for 30 min. and immunoreactivity was visualized with AEC solution (Thermo Fisher Scientific, USA). Slides known to express Osteocalcin were used as positive controls, omitting of the primary antibody served as negative control.

6. Statistical analysis

In the scope of this thesis, experiments were performed at least in triplicate and expressed as means ± standard deviations. Student’s t test was used for statistical analysis. Statistical significance was defined as p < 0.01 or p < 0.05 for a 99% or a 95% confidence.
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References


Chapter IV

Osteoinduction in human fat derived stem cells by recombinant human bone morphogenetic protein-2 produced in *Escherichia coli*

This Chapter is based on the following publication:

Chapter IV

Osteoinduction in human fat derived stem cells by recombinant human bone morphogenetic protein-2 produced in *Escherichia coli*

Abstract

Bioactive recombinant human bone morphogenetic protein-2 (rhBMP-2) creating an alternative to the time-consuming methods of inclusion bodies preparation was obtained using *Escherichia coli* pET-25b expression system. 55 mg purified rhBMP-2 were achieved per gram cell dry weight, with up to 95% purity. In murine C2C12 cell line, rhBMP-2 induced an increase in the transcription of Smads and of osteogenic markers Runx2/Cbfa1 and Osterix, measured by semi-quantitative RT-PCR. Bioassays performed in human fat derived stem cells showed increase activity of early osteogenic marker alkaline phosphatase and absence of cytotoxicity.

1. Introduction

Bone morphogenetic proteins (BMPs) are a group of cytokines from the TGF-β superfamily (Wozney *et al.* 1988) with strong ability to induce bone and cartilage formation (Urish 1965, Reddi 1998). BMPs have been used as powerful osteoinductive components in several late-stage tissue engineering products for bone grafting (Westerhuis *et al.* 2005).

During the process of bone healing, signals from BMPs trigger the differentiation of stem cells into bone forming cells that are recruited to the site of injury (Reddi 1981). BMPs bind to serine-threonine kinase cell receptors and mediate their signals through Smad related cascades and Smad-independent pathways (Derynck 2001, Shi & Massague 2003). Signal is regulated by different factors such as binding to cell surface receptors (Kirsch *et al.* 2000, Sebald *et al.* 2004), formation of BMP-heterodimers (Israel *et al.* 1996), presence of antagonists such as Noggin (Groppe *et al.* 2002), cross-talk with other pathways (Miyazono *et al.* 2005) and the presence of a heparin binding domain in the N-terminal region of BMP-2 which provides anchorage of the cytokine to the extracellular matrix (Ruppert *et al.* 1996, Kubler *et al.* 1999, Wurzler *et al.* 2004).

Currently, most BMPs are obtained from mammalian cell cultures in low yields (Wang *et al.* 1990, Israel *et al.* 1992) or from bacteria inclusion bodies after time-consuming refolding procedures (Kubler *et al.* 1998, Vallejo *et al.* 2002, Long *et al.* 2006). Aiming to circumvent the...
disadvantages of the previously reported methods, we have developed a novel approach for
the production of high amounts of soluble and pure rhBMP-2, with bioactivity verified in
murine myoblast C2C12 cell line as well as in human fat derived adult stem cells.

2. Materials and methods

2.1. Cloning of rhBMP-2

The sequence coding for the mature (bioactive) domain of human BMP-2 was obtained
from a bacterial clone (Sanger Institute, UK), containing locus 20p12.1-13 of chromosome 20
of human genome, which includes the entire gene for human BMP-2 (clone ref. RPS-859D4).
According to the data in Human Genome Project (http://www.ensembl.org/homo_sapiens),
the entire human BMP-2 gene (Ensembl gene OTTHUMG00000031833) includes three exons
and two introns with the open reading frame located on exons 2 and 3. By consulting
UniProtKB/Swiss-Prot, human BMP-2 protein (P12643) mature domain consists of amino acids
283-396, which is obtained after cleavage from propeptide (Hillger et al. 2005). The region
corresponding to the mature domain is located solely in exon 3 and, for this reason, we have
cloned directly by PCR this sequence from the above mentioned clone (Figure 1A). Mature
domain was amplified by PCR using primers 5’ CGG GAT CCA CAA GCC AAA CAC AAA CAG C 3’
(forward primer) and 5’ CCC TCG AGG CGA SAC CCA CAA CCC TC 3’ (reverse primer). The
nucleotides for cleavage by BamH and XhoI in forward primer and reverse primers are
underlined, respectively; the codons corresponding to amino acids 283 and 396 are
represented in bold. PCR-product was first cloned into pGEM T-Easy vector (Promega, USA)
and then subcloned in a pET-25b vector (Novagen, USA) via the two restriction sites, BamH
I and XhoI (plasmid pET-25b/rhBMP-2), that was used to transform E. coli BL21(DE3) strain
(Invitrogen, UK). DNA cloning and manipulation were performed according to standard
protocols (Sambrook et al 1989). The integrity of cloned PCR products was verified by DNA
sequencing (Sanger et al. 1977) using ABI PRISM310 Genetic Analyzer.

2.2. Expression of rhBMP-2 in bioreactor

Transformed E. coli BL21(DE3) strain with pET-25b/rhBMP-2 or pET-25b vectors were
picked and transferred into 50 ml Luria Bertani (LB) media with 50 μg ampicillin/ml and
incubated at 37 °C for 1-2h. Pre-cultures were then transferred to a pre-autoclaved 2 L
Bioflo110 bioreactor (New Brunswick Scientific, USA) containing LB media with 50 μg
ampicillin/ml and cultured at 37 °C until an OD600 of 1.0. 1 mM IPTG was then added and
temperature lowered to 25 °C. During the entire process, aeration was kept at 2 L/min and an
agitation cascade between 300 and 1000 rpm was selected in order to maintain minimum dissolved oxygen levels at 30% of saturation. The pH 7.4 was kept constant using solutions of ammonium hydroxide (30%, v/v) and phosphoric acid (5 M) pumped into the bioreactor vessel. Biomass was collected by centrifugation (4000 g, 20 min, 4 ºC), washed once with PBS and stored at -20 ºC.

Figure 1. Strategy used for cloning the mature domain of human bone morphogenetic protein-2. A) BMP-2 gene contains three exons represented by boxes and two introns corresponding to the lines. Coding region is represented by black boxes located in exon 2 and 3. The mature domain of human BMP-2 protein is entirely located on exons 3 correspondent to amino acids 283 to 396. B) Construction of vector pET-25b/rhBMP-2 for expression of human BMP-2 in *E. coli*. The mature BMP-2 was cloned into the vector pET-25b together with its own leader sequence and 6-histidine tag for purification.
2.3. Purification of rhBMP-2

Frozen bacteria were resuspended in lyses buffer (20 mM sodium phosphate buffer, 0.5 M NaCl, 1 mg/ml lysozyme), with inhibitor of proteases (Complete Mini EDTA-free, Roche). Bacteria were ultrasonicated 4 times for 15 s with intervals of 1 min on ice and supernatant and pellet fractions collected by centrifugation (4000 g, 30 min, 4 °C). The pellet was then incubated overnight in solubilization buffer (phosphate saline buffer, 40 mM imidazole, 0.7 M L-arginine, pH 11.0) at 18 °C with gentle stirring and the supernatant containing soluble rhBMP-2 collected after centrifugation (4000 g, 4 °C, and 30 min). The supernatant was adjusted to pH 8.5 with 0.1M HCl and applied to a pre-equilibrated HisTrap chromatography column (Amersham). Briefly, column was equilibrated with 5 column volumes of sodium phosphate buffer containing 40 mM imidazole. The column was washed extensively with 60-80 ml of sodium phosphate buffer containing 80 mM imidazole and sample eluted in 15 ml of sodium phosphate buffer containing 500 mM imidazole. A flow rate of 5 ml/min was kept during the whole procedure by the use of a peristaltic pump. Samples were analyzed by SDS-PAGE and Western-blot. Imidazole was removed by buffer exchange to phosphate buffer saline with use of a HiTrap desalting column (Amersham) prior to bioactivity tests.

2.4. Analytical detection of rhBMP-2 by reducing and non-reducing SDS-PAGE and Western-blot

Protein samples were mixed with SDS-PAGE loading buffer and heated at 95 °C for 5 min. Samples were separated by using hand-cast 10-12 % SDS–PAGE gels in reducing or non-reducing conditions. Reducing conditions included the use of SDS-PAGE loading buffer with 5 % β-mercaptoethanol as reducing agent. Coomassie blue-R or silver staining was used for visualization and staining of gels. Samples were then electro-transferred (semi-dry Western-blot transfer) to a nitrocellulose membrane at 400 mA, 10 V for 30 min. The membranes were blocked with 2 % BSA in PBS-T buffer (PBS, 0.05 % Tween 20) and incubated either with peroxidase conjugated anti-His antibody (Sigma) diluted 1:5000 or rabbit anti-human BMP-2 polyclonal primary antibody (AbD Serotec) diluted 1:5000 followed by mouse anti-rabbit peroxidase conjugated secondary antibody (Amersham) diluted 1:30000. Image detection was performed with ChemiDoc XRS and Quantity One software (BioRad).

2.5. Bioactivity tests

C2C12 cells were seeded at 10⁵ cells/ml per well in a 24-well plate, attached overnight in Dulbecco’s modified Eagle’s medium (DMEM) with 1 % fetal calf serum and no antibiotics, at 37 °C with 5 % CO₂ in a humidified environment. Cells were incubated for 5 days with rhBMP-2
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purified by the method described above and a commercial rhBMP-2 (Wyeth, USA) used as control. Primary cultures of human fat tissue derived stem cells (Malafaya et al. 2005, Zuk et al. 2002) were seeded at $10^5$ cells/ml per well in a 24-well plate and attached overnight in DMEM with 50 μg/ml ascorbic acid, 10 mM β-glycerophosphate, 100 U/ml penicillin, 100 μg/ml streptomycin and 10 % fetal bovine serum at 37 ºC in 10 % CO₂. Cells were incubated for 14 days with rhBMP-2 purified by our method, with media replacement every two days. Alkaline phosphatase (ALP) enzymatic activity was determined accordingly to standard procedures (Salgado et al. 2002) after 5 days on C2C12 cell line cultures and after 3, 7 and 14 days on cell cultures of mesenchymal stem cells. MTS cell viability assay was determined after 3, 7 and 14 days of cell culture in human stem cells (Salgado et al. 2002). For RT-PCR, mRNA of C2C12 cells was extracted after 5 days of cell culture with TriReagent RNA Isolation Reagent (Sigma-Aldrich) and cDNA was synthesized from 4 μg total RNA with oligo dT primer (MWG Biotech AG) and AMV reverse transcriptase (Promega). The polymerase chain reaction (PCR) was performed with specific primers for osteogenic markers (Table 1) and with use of a Hot Star Taq Polymerase (Qiagen). Agarose gels were imaged with Multimage Light Cabinet (Alpha Innotech, USA) and gene expression analyzed with Chemilmage 4400 AlphaEase FC Image Analysis Software (Alpha Innotech, USA) and using polymerase II for normalizing expression.

Table 1. Oligonucleotide primers for semi-quantitative RT-PCR analysis in murine C2C12 cells.

<table>
<thead>
<tr>
<th>Gene marker</th>
<th>Primer sequence: Sense/ Antisense</th>
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<tbody>
<tr>
<td>Runx2/Cbfa1</td>
<td>5’ GCC GCA GTG CCC CGA TTG AGG 3’</td>
</tr>
<tr>
<td></td>
<td>5’ AGG GAG GGC CTG GGG TTC TGA GG 3’</td>
</tr>
<tr>
<td>Smad-1</td>
<td>5’ CCC ACC TGC TTA CCT GCC TCC TGA 3’</td>
</tr>
<tr>
<td></td>
<td>5’ TGG GGT GAA AGC CGT GGT GAG ATG 3’</td>
</tr>
<tr>
<td>Smad-5</td>
<td>5’ AAT GGG CAG AAA AGG CAG TGG ATG 3’</td>
</tr>
<tr>
<td></td>
<td>5’ AGC GTT GGT GGG TTG GTG GAA AGA 3’</td>
</tr>
<tr>
<td>Osterix</td>
<td>5’ GCA GTG GGG CAG GCC GTT CTA CC 3’</td>
</tr>
<tr>
<td></td>
<td>5’ GGG GCG GCC GCT GAT TGG CTT TCT 3’</td>
</tr>
<tr>
<td>Osteocalcin</td>
<td>5’ ATT CAT ATG AGG ACC CTC TCT CTG 3’</td>
</tr>
<tr>
<td></td>
<td>5’ ATT AGA TCT CTA ATG ATG ATG ATG AAT AGT GAT ACC GTA GAT 3’</td>
</tr>
<tr>
<td>ALP</td>
<td>5’ CAC GCG ATGCAA CAC CAC TCA GG 3’</td>
</tr>
<tr>
<td></td>
<td>5’ GCA TGT CCC CGG GCT CAA AGA 3’</td>
</tr>
<tr>
<td>Polymerase II</td>
<td>5’ TAC ACC CCA GCT TCT CCC AAA TAC 3’</td>
</tr>
<tr>
<td></td>
<td>5’ AGC TCT TCG CCC TGT TCG 3’</td>
</tr>
</tbody>
</table>
3. Results

3.1. Cloning, expression and purification of rhBMP-2

The PCR gave a single product of the coding region of human BMP-2, with expected size of 359 bp (data not shown). DNA sequencing revealed a total correspondence to the encoded amino acids (Ensembl gene OTTHUMG00000031833). Figure 1B shows the map of the expression vector pET-25b/rhBMP-2 used for transformation of bacteria. *E. coli* BL21 (DE3) strain transformants were grown and collected in conditions described in Material and Methods section. After bacteria lyses, rhBMP-2 was found mainly in the pellet (fraction corresponding to bacterial debris) (Figure 2A). Soluble protein was then recovered with phosphate buffer containing 0.7M L-arginine, pH 11.0. Purification procedures revealed that rhBMP-2 was expressed in monomer, dimer and polymer forms visible in Western-blot using antibodies anti-histidine tag and anti-BMP-2. Silver stained SDS-PAGE revealed up to 95 % purity (Figure 2B). RhBMP-2 stored in 20mM phosphate sodium, 0.5M NaCl was found to be stable over two months at pH range of 7-9 (data not shown).

![Figure 2](image.png)

*Figure 2.* A) Coomassie-blue stained SDS-PAGE of crude lysates, pellet fraction, from batch cultivation of transformed *E. coli*. Lanes 1) pET-25b clone; 2) rhBMP-2/pET-25b clone. Monomer appears as a well-defined band around 17 kDa. Molecular weight is in kDa (Broad Range, BioRad). B) rhBMP-2 purified by affinity chromatography: 1) Silver stained reduced SDS-PAGE, showing the monomer of BMP-2 with significant purity; 2) Western-blot using anti-his tag antibody; 3) Western-blot using anti-BMP-2 antibody. BMP-2 is visible in monomer, dimer and polymer.
3.2. Biological activity in C2C12 cells

Bioassays were performed in murine myoblast C2C12 cell line, by the administration of purified rhBMP-2. Semi-quantitative RT-PCR revealed an increase of expression of early osteoblast differentiation markers after 5 days of cell culture, such as the osteochondral transcription factor Runx2/Cbfa1 (by 2-3 fold), Smad-1 and -5 (up to 1.5 fold) and osteoblast transcription factor Osterix (up to 1.5 fold). In these tests, the levels of ALP mRNA did not change significantly. Commercial mammalian rhBMP-2 induced markedly Runx2/Cbfa1 by 3-7 fold, Smad-1 and Smad-5 by 2-5 fold, Osterix by 2-4 fold, Osteocalcin by 2-9 fold and high levels of ALP expression were detected (Figure 3).

![Figure 3](image-url)

**Figure.** 3. RT-PCR for genes involved during osteoblast differentiation of C2C12 cells after 5 days of culture with purified rhBMP-2 and commercial mammalian rhBMP-2 as control. Gene expression was analyzed with Chemilimage 4400 AlphaEase FC Image Analysis Software (Alpha Innotech) and standardized with Polymerase II expression. OSX: osterix; OC: osteocalcin; ALP: alkaline phosphatase. Conditions used: 1) No rhBMP-2; 2) 250 ng/ml; 3) 500 ng/ml; 4) 1000 ng/ml; 5) 250 ng/ml commercial preparation; 6) 500 ng/ml commercial preparation; 7) 1000 ng/ml commercial preparation. Data presented are mean values of two independent experiments. (* bioassay was not performed)
3.3. Biological activity in mesenchymal stem cells

RhBMP-2 did not show any significant evidence of cytotoxicity in human mesenchymal stem cells after two weeks of cell culture (Figure 4A). MTS is a viability/proliferation test and an inverse relationship of toxicity to cells can be assumed. In human primary cultures of fat derived stem cells, there was an increase in levels of ALP enzymatic activity (Figure 4B). Data correlates with morphological observations of the cells with optical microscopy (Figure 5).

**Figure 4.** A) Tetrazolium salt (MTS) test performed in fat tissue derived human stem cells as a function of rhBMP-2 concentration, after 3, 7 and 14 days of cell culture. Cell viability is expressed in percentage. B) Bioactivity of rhBMP-2 as measured by the induction of alkaline phosphatase activity in fat tissue derived human stem cells. ALP activity levels are expressed in μmol ALP per mg of total protein.

**Figure 5.** Cell morphology of fat derived human stem cells after 10 days of cell culture with A) no growth factor (control) and B) with 500 ng/ml purified rhBMP-2 (replacement of growth factor and culture media every two days). x400. Fibroblast-like (f) and osteoblast-like (o) morphologies were observed when rhBMP-2 was used on the mesenchymal stem cell primary culture.
4. Discussion

A novel and more simple approach for the production of readily soluble rhBMP-2 using \textit{E. coli} that, for the first time, is bioactive in human cell cultures. In previous reports, recombinant BMP-2 production was achieved in bacterial inclusion bodies in fairly large amounts (Li \textit{et al.} 1998, Vallejo \textit{et al.} 2002, Long \textit{et al.} 2005). However, inclusion bodies require time-consuming steps for solubilization and refolding. In our approach, we have achieved expression of rhBMP-2 directly folded in dimer by use of pET-25 expression system which directs production of soluble recombinant protein into periplasmatic space of \textit{E. coli}, thus allowing the formation of disulfide bridges.

Recovery with 0.7 M L-arginine buffer resulted in monomer, dimer and polymer forms adding value to what has been reported by Klösch \textit{et al.} (2005) and Long \textit{et al.} (2006). In fact, the formation of cysteine bridges is known to be favored at slightly alkaline pH, thus increasing amounts of dimeric rhBMP-2 (Vallejo & Rinas 2004). However, this contribution seems to be the first one to show this method to obtain soluble and dimeric BMP-2 directly upon production. The amount of purified rhBMP-2 was about 55 mg per gram cell dry weight, corresponding to 110 mg purified rhBMP-2 per litter of culture broth.

In previous studies, recombinant BMPs obtained from \textit{E. coli} were reported to be tested in murine cells lines C2C12 (Vallejo \textit{et al.} 2002, Long \textit{et al.} 2006), MC3T3-E1 (Klösch \textit{et al.} 2005) and CH310T1/2 (Yang \textit{et al.} 2003) or \textit{in vivo} rodent models (Kubler \textit{et al.} 1998, Bessho \textit{et al.} 2000, Hillger \textit{et al.} 2005). In our case, we report bioactivity on primary cultures of human stem cells. Binding of active BMP-2 to cell receptors results in an increase of early markers activated during osteogenic differentiation (i.e. Runx2, Osterix). In human stem cells, rhBMP-2 increased ALP enzymatic activity and differentiated cells into osteoblast-like morphology. In conclusion, the novel approach described herein shows steps for improving the production of high amounts of readily soluble rhBMP-2 that shows bioactivity in primary human adult stem cells, and can be use for diverse bone tissue engineering research applications.
References


Chapter IV. Osteoinduction by recombinant human BMP-2 produced in Escherichia coli


Chapter V

Expression, purification and osteogenic bioactivity of recombinant human BMP-4, -9, -10, -11 and -14

This Chapter is based on the following publication:

Bessa PC, Cerqueira MT, Rada T, Gomes ME, Neves NM, Nobre A, Reis RL and Casal M, 2009, Expression, purification and osteogenic bioactivity of recombinant human BMP-4, -9, -10, -11 and -14, Protein Expression and Purification 63, 89-94.
Chapter V
Expression, purification and osteogenic bioactivity of recombinant human BMP-4, -9, -10, -11 and -14

Abstract

Bone morphogenetic proteins (BMPs) are cytokines from the TGF-β superfamily, with important roles during embryonic development and in the induction of bone and cartilage tissue differentiation in the adult body. In this contribution, we report the expression of recombinant human BMP-4, BMP-9, BMP-10, BMP-11 (or growth differentiation factor-11, GDF-11) and BMP-14 (GDF-5), using *Escherichia coli* pET-25b vector. BMPs were overexpressed, purified by affinity his-tag chromatography and shown to induce the expression of early markers of bone differentiation (e.g. smad-1, smad-5, runx2/cbfa1, dlx5, osterix, osteopontin, bone sialoprotein and alkaline phosphatase) in C2C12 cells and in human adipose stem cells. The described approach is a promising method for producing large amounts of different recombinant BMPs that show potential for novel biomedical applications.

1. Introduction

Discovered in 1965, bone morphogenetic proteins (BMPs) are a group of several cytokines belonging to the transforming growth factor-β (TGF-β) superfamily. These proteins have strong inducing activity in bone and cartilage formation, and also have important roles during embryonic patterning and early skeletal development (Bessa et al. 2008a). During the last decades, BMPs have been used as powerful osteoinductive components in several late-stage tissue engineering products for bone grafting (Westerhuis et al. 2005). Since 2002, BMP-2 and BMP-7 received approval for specific clinical uses (Bessa et al. 2008b).

Despite being extensively investigated, to date, most research has been focused only in a few BMPs, particularly in BMP-2 and BMP-7, whilst little is known about the physiological roles and the biomedical relevance for the remaining BMPs. Both, BMP-4 and BMP-9 have been shown clearly bone-inducing roles, as shown by their ability to induce osteogenic differentiation in C2C12 cell line (Cheng et al., 2003) and orthotopic ossification in mice (Kang et al., 2004). BMP-4 is also implicated in diverse roles during embryonic development, and as a differentiation factor to haematopoietic and nerve cells (Li et al. 2001, Giacomini et al. 2006,
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Reissmann et al. 1996). BMP-14 (or growth and differentiation factor-5) induces mainly tendon and cartilage formation (Aspenberg and Forslund 1999; Dines et al. 2007) and has been also studied in healing of periodontal ligament (Nakamura et al. 2003). BMP-11 and BMP-10 have not been identified for any bone-inducing potential. BMP-10 has expression restricted solely to the heart tissue and is essential to cardiac morphogenesis (Neuhaus et al. 1999; Chen et al. 2004), while BMP-11 (or growth and differentiation factor-11) has roles in the establishment of axial patterning (Andersson et al. 2006). In former reports, human BMP-4 and BMP-14 have been expressed in E. coli and reported to induce alkaline phosphatase marker in murine cells but these were not tested in human cells (Kubler et al. 1998, Sieber et al. 2006). In addition, the expression of human BMP-9, -10 and -11 in Escherichia coli is not yet reported. In this work, we described the production, purification and bioactivity of recombinant human BMP-4, -9, -10, -11 and -14 by a novel method that was previously reported for the overexpression of human BMP-2 (Bessa et al. 2008c). The different BMPs were analyzed by their ability to induce the mRNA expression of several markers of bone differentiation in C2C12 cells and in human adult stem cells from adipose tissue. Our results suggest a promising and straightforward way for the production of different BMPs, all capable of inducing cell differentiation in the aforementioned models.

2. Materials and methods

2.1. Cloning and expression of rhBMP-4, -9, -10, -11 and -14

The sequences coding the mature (bioactive) domains of human BMP-4, BMP-9, BMP-10, BMP-11/GDF-11 and BMP-14/GDF-5 were obtained from different bacterial clones (Sanger Institute, UK), which contained the full genes encoding the human BMPs: clone reference RP11-808F04 for BMP-4, RP11-463P17 for BMP-9, RP11-88N13 for BMP-10, RP11-55N12 for BMP-11, and RP11-173P06 for BMP-14. According to the data from the UniProtKB/Swiss-Prot database (http://www.expasy.org) and from the Human Genome Project public database (http://www.ensembl.org/homo_sapiens), the sequences coding for the bioactive domains of these BMPs, are located in one sole exon, thus allowing direct PCR cloning of these BMPs from the above mentioned clones, as was previously described (Bessa et al. 2008c).

DNA coding for the mature/bioactive protein domains was amplified by PCR using the primers indicated in Table 1. PCR-products were cloned in a pET-25b vector (Novagen, USA) via the two restriction sites, BamHI and XhoI, that were used to transform E. coli BL21(DE3) strain (Invitrogen, UK). The restriction sites for these enzymes were specific.
Table 1. Oligonucleotide primers for cloning of human BMP-4, -9, -10, -11 and -14. The nucleotides for restriction by BamH1 and XhoI are underlined in the forward and reverse primers, respectively. The codons representing the beginning and the end of BMP sequences are represented in bold.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer sequence: Forward / Reverse</th>
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<tr>
<td>BMP-4</td>
<td>5’ CGG GAT CCA AGC CCT AAG CAT TCA C 3’</td>
</tr>
<tr>
<td></td>
<td>5’ CCC TCG AGG CGG CAC CCA CAT CC 3’</td>
</tr>
<tr>
<td>BMP-9</td>
<td>5’ CGG GAT CCA AGC GCC GGG GCT GGC AG 3’</td>
</tr>
<tr>
<td></td>
<td>5’ CCC TCG AGG CTG CAC CCA CAC TCT GGC AC 3’</td>
</tr>
<tr>
<td>BMP-10</td>
<td>5’ CGG GAT CCA AAC GCC AAA GGA AAC TAC 3’</td>
</tr>
<tr>
<td></td>
<td>5’ CCC TCG AGG CTG CCA CAT TCG GAG 3’</td>
</tr>
<tr>
<td>BMP-11</td>
<td>5’ CGG GAT CCA AAC CTG GGT CTG GAC TG 3’</td>
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<tr>
<td></td>
<td>5’ CCC TCG AGA GAG CAG CCA CAG CGA TC 3’</td>
</tr>
<tr>
<td>BMP-14</td>
<td>5’ CGG GAT CCA GCC CCA CTG GCC ACT C 3’</td>
</tr>
<tr>
<td></td>
<td>5’ CCC TCG AGG CTG CCA CAC GAC TC 3’</td>
</tr>
</tbody>
</table>

DNA cloning and manipulation were performed according to standard protocols (Sambrook et al. 1989). The integrity of the cloned PCR products was verified by DNA sequencing (Sanger et al. 1977) using an ABI PRISM310 Genetic Analyzer. Transformed E. coli BL21(DE3) strain with pET-25b/rhBMP were inoculated in 50 ml Luria-Bertani (LB) medium with 50 μg ampicillin/ml and incubated at 37 °C, until an OD600 of 0.8 was obtained. To induce recombinant protein expression IPTG 1 mM was then added to the culture media, and the temperature lowered to 25 °C during 24 h. Biomass was collected by centrifugation (4000 g, 20 min, 4 ºC), washed once with PBS and stored at -20 ºC.

2.2. Purification of rhBMPs

Frozen bacteria were resuspended in lyses buffer (20 mM sodium phosphate buffer, 0.5 M NaCl, 1 mg/ml lysozyme), with a protease inhibitors (Complete Mini EDTA-free, Roche). Bacteria cells were ultrasonicated 6 times for 30 s with intervals of 5 min on ice and supernatant and pellet fractions collected by centrifugation (4000 g, 30 min, 4 ºC). The pellet was then incubated overnight in solubilization buffer (phosphate saline buffer, 30 mM imidazole, 0.7 M L-arginine, pH 10.0) at 18 ºC with gentle stirring and the supernatant containing soluble rhBMP collected after centrifugation (4000 g, 20 min, 4 ºC). The pH of supernatant was adjusted to 7.5 with 0.5M HCl and applied to a pre-equilibrated HisTrap
chromatography column (Amersham). Briefly, the column was equilibrated with 5 column volumes of sodium phosphate buffer containing 30 mM imidazole, washed extensively with 100-120 ml of sodium phosphate buffer containing 60 mM imidazole and the sample eluted in 20 ml of phosphate buffer supplemented with 400 mM imidazole. A flow rate of 5 ml/min was kept during the whole procedure by the use of a peristaltic pump. Purified protein was desalted with the use of a HiTrap desalting column (Amersham) and freeze-dried. BMPs were resuspended in sterile phosphate buffer saline, prior to bioactivity tests. Protein concentration was measured by Bradford method (Bradford et al. 1996) using bovine serum albumin (BSA) standards of known concentration. Protein purity was estimated using SDS-PAGE image analysis (GelPro Analyzer 4.0, Media Cybernetics).

2.3. Western-blot detection of rhBMPs

Protein samples were mixed with SDS-PAGE loading buffer and heated at 95 °C for 5 min. Samples were separated by using hand-cast 10-12 % reduced SDS–PAGE gels and Coomassie blue-R was employed for visualization and staining of gels. Samples were then electro-transferred (wet Western-blot transfer) to a nitrocellulose membrane at 100 V for 60 min. The membrane was blocked with 2% (w/v) BSA in PBS-T buffer (PBS, 0.05% Tween 20) and incubated with peroxidase conjugated anti-His antibody (Sigma) diluted 1:2000. Image detection was performed with ChemiDoc XRS and Quantity One software (BioRad).

2.4. Bioactivity tests

C2C12 cells were seeded at 2.5 x 10⁴ cells/ml per well in a 24-well plate, attached in Dulbecco’s modified Eagle’s medium (DMEM) with 1% (v/v) fetal bovine serum, 100 U/ml penicillin and 100 μg/ml streptomycin, at 37 °C with 5% CO₂ in a humidified environment. Cells were incubated for 5 days with BMP-4, -9, -10, -11 and -14, purified by the method described above (500 ng/ml) and a control with no cytokine. MTS cell viability assay was performed in C2C12 cells after 3 days of culture (Salgado et al. 2002). Primary cultures of human adult stem cells isolated from adipose tissue (Malafaya et al. 2005) were cultured under similar conditions and incubated with BMP-4, -9 and -14 (500 ng/ml) for 3 days of cell culture. For RT-PCR, the mRNA of cells was extracted with TriZol Reagent (Invitrogen, USA) from triplicate 24-well assays following the procedure provide by the supplier. Samples were quantified using a NanoDrop ND-1000 Spectrophotometer (NanoDrop Technologies, USA) and cDNA synthesis performed with iScript cDNA synthesis Kit (BioRad, USA). PCR was performed with specific primers for osteogenic markers. Agarose gels were visualized using Eagle Eye imaging system.
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(Stratagene) and gene expression was analyzed with Gel-Pro Analyzer (Media Cybernetics), using polymerase II expression as a housekeeping marker.

2.5. Statistical analysis

Experiments were performed at least in triplicate and expressed as means ± standard deviations. Student’s t test was used for statistical analysis using a two-tailed paired test and analyzed by a paired analysis of variance. Statistical significance was defined as p < 0.05 for a 95% confidence.

3. Results

3.1. Cloning, expression and purification of rhBMPs

The PCR originated single products for the coding regions of human BMP-4, -9, -10, -11 and -14 (data not shown). DNA sequencing revealed a total correspondence to encoded nucleotides and corresponding amino acids. *Escherichia coli* BL21(DE3) strain transformants were then cultivated and collected as described in the Material and methods section. After cell lyses, rhBMPs were found mainly in the pellet (fraction corresponding to cell debris) (data not shown). Soluble protein was recovered using phosphate buffer containing 0.7M L-arginine, pH 10.0, as indicated in Materials and Methods. After purification by affinity chromatography, the rhBMPs were visible as monomer, dimer and polymer, with Western-blot, using an antibody against the six-histidine tag (Figure 1). Estimating from Coomassie blue-stained SDS-PAGE, the BMPs were pure up to 95% (Table 2).

![Figure 1. Western-blot of purified rhBMP-4, -9, -10, -11 and -14 and a negative control (respectively 1-6) using an anti 6-His tag antibody. The BMPs are visible in monomer, dimer and polymer.](image)

<table>
<thead>
<tr>
<th>Lanes</th>
<th>Size (kDa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
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</tr>
<tr>
<td>2</td>
<td>116</td>
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<td></td>
<td>14</td>
</tr>
</tbody>
</table>
Chapter V. *Expression, purification and bioactivity of recombinant human BMP-4, -9, -10, -11 and -14*

Table 2. Purification of recombinant human BMP-4, BMP-9, BMP-10, BMP-11 (GDF-11) and BMP-14 (GDF-5).

<table>
<thead>
<tr>
<th>Steps</th>
<th>Crude lysate</th>
<th>L-arginine solubilization</th>
<th>Ni-NTA chromatography</th>
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</thead>
<tbody>
<tr>
<td>Protein</td>
<td>Total protein (mg)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Purity (%)&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Total protein (mg)</td>
</tr>
<tr>
<td>hBMP-4</td>
<td>497.5</td>
<td>24.4</td>
<td>203.6</td>
</tr>
<tr>
<td></td>
<td>Yield 100 %</td>
<td></td>
<td>Yield 70.5 %</td>
</tr>
<tr>
<td>hBMP-9</td>
<td>531.5</td>
<td>17.2</td>
<td>220.9</td>
</tr>
<tr>
<td></td>
<td>Yield 100 %</td>
<td></td>
<td>Yield 66.5 %</td>
</tr>
<tr>
<td>hBMP-10</td>
<td>533.3</td>
<td>14.9</td>
<td>261.3</td>
</tr>
<tr>
<td></td>
<td>Yield 100 %</td>
<td></td>
<td>Yield 77.4 %</td>
</tr>
<tr>
<td>hBMP-11</td>
<td>506.4</td>
<td>13.5</td>
<td>278.5</td>
</tr>
<tr>
<td></td>
<td>Yield 100 %</td>
<td></td>
<td>Yield 80.1 %</td>
</tr>
<tr>
<td>hBMP-14</td>
<td>569.1</td>
<td>19.7</td>
<td>252.5</td>
</tr>
<tr>
<td></td>
<td>Yield 100 %</td>
<td></td>
<td>Yield 73.4 %</td>
</tr>
</tbody>
</table>

<sup>a</sup> The recombinant hBMPs were purified from 1 L *E. coli* culture.

<sup>b</sup> Total protein quantification was determined using Bradford method

<sup>c</sup> Estimated from SDS-PAGE image analysis

3.2. Cytotoxicity of rhBMPs

Bioassays were performed in murine myoblast C2C12 cell line, by the administration of purified rhBMP-2. The different proteins did not show cytotoxicity in C2C12 cells after three days of cell culture at 50 and 500 ng/ml (*Figure 2*). MTS is a viability/proliferation test and an inverse relationship of toxicity to cells can be assumed. BMP-4 (50 ng/ml), BMP-9 (50 ng/ml) and BMP-10 (500 ng/ml) significantly stimulated cell proliferation by approximately 10 to 25 % (*p* < 0.05).
3.3. Expression of osteogenic markers induced by rhBMPs

Semi-quantitative RT-PCR revealed an increase in expression of several early osteoblast differentiation markers in C2C12 cell line after five days of cell culture with the different BMPs (Figure 3). The increased mRNA levels were mean values of independent assays, and normalized against the expression of polymerase II. The expression of early osteochondral transcription factor runx2/cbfa1 was induced by 2 to 3-fold with BMP-4 and BMP-11, 4-fold with BMP-10 and BMP-14, and 6-fold with BMP-9 ($p < 0.05$ in all cases). Expression of smad-1 and smad-5, two signaling molecules for BMPs, were also induced but in a lower extent. The expression of smad-5 with induced 2-fold with BMP-4 and BMP-9 ($p < 0.05$). Expression of smad-1 was induced with BMP-9 (2-fold, $p < 0.05$). With BMP-10, BMP-11 and BMP-14, there was no significant increase in the expression of smad-1 or -5. Osterix, a transcription factor downstream of runx2/cbfa-1 and a marker of commitment to osteoblastic lineage, was induced both with BMP-4 (2-fold) and BMP-9 (2-fold, $p < 0.05$) and its expression maintained with the other BMPs. Osterix expression was slightly decreased by BMP-11 (not statistically significant). The expression of transcription factor dlx5 was nearly the same with all BMPs, with a slight increase with BMP-4 (1.5-fold) which is not significant. Alkaline phosphatase (ALP), bone sialoprotein and osteopontin, three markers of bone formation, were induced with some
of these BMPs. ALP was induced with BMP-4, BMP-9 (2-fold, \( p < 0.05 \)) and BMP-10 and BMP-14 (1.5-fold), and reduced by half (0.5-fold) with BMP-11 (\( p < 0.05 \)). Bone sialoprotein expression was up-regulated with BMP-14 (3-fold) and with BMP-4 (5-fold, \( p < 0.05 \)). Osteopontin expression was up-regulated mainly with BMP-9, BMP-14 and less with BMP-4 and BMP-10 (\( p < 0.05 \)), compared with basal expression in the control and with BMP-11. Osteocalcin, a marker specific of late-stages of bone mineralization, was not detected in any assay. The observations on cell morphology correlated well with the induction of the several osteogenic markers (Figure 4). BMP-9, and to a less degree BMP-4 and BMP-14, induced an osteoblast-like morphology.

In the primary cultures of fat-derived human stem cells, BMP-4, BMP-9 and BMP-14 were also able to induce an increase in the expression of the different early markers of osteogenic differentiation (Figure 5). BMP-4 induced the highest levels of osteogenic markers, particularly runx2 (7-fold, \( p < 0.05 \)), bone sialoprotein (7-fold, \( p < 0.05 \)) and osteopontin (8-fold, \( p < 0.05 \)). BMP-9 induced runx2 (2-fold) and alkaline phosphatase (2-fold), while BMP-14 induced the expression of alkaline phosphatase (5-fold, \( p < 0.05 \)) and bone sialoprotein (3-fold, \( p < 0.05 \)). Bacteria transformed with blank vector (e.g. no BMP) did not induce any expression of osteogenic genes (data not shown).
Figure 3. mRNA expression of signaling molecules and transcription factors (A) and osteogenic-specific markers (B) induced by BMPs, measured by semi-quantitative RT-PCR after 5 days of cell culture in C2C12 cell line. Bars represent mean values ± SD of gene expression fold variation of at least three independent determinations (* \( p < 0.05 \)). Smad-1 and Smad-5 were induced mostly for BMP-4, -9 and -14, Osterix for BMP-4 and -9, Runx2 was induced for all BMPs but higher for BMP-9. Bone sialoprotein was induced mostly for BMP-4 and -11; ALP was induced 1.5 to 2-fold for all BMPs except BMP-11; osteopontin was induced mainly for BMP-9, BMP-10 and BMP-14.
Figure 4. Cell morphology of C2C12 cells after 5 days of cell culture with A) no growth factor (control), B) 500 ng/ml rhBMP-4, C) 500 ng/ml rhBMP-9, D) 500 ng/ml rhBMP-10, E) 500 ng/ml rhBMP-11, and F) 500 ng/ml rhBMP-14. Purified BMP was used. x400. Osteoblast-like morphology (o) was observed in C2C12 cells stimulated with BMP-4, BMP-9 and BMP-14 and myoblast-like morphology (m) in control and cells stimulated with BMP-10.
Figure 5. mRNA expression of early markers of osteogenic differentiation induced by BMPs, measured by semi-quantitative RT-PCR after 3 days of human adult stem cell cultures. Bars represent mean values ± SD of gene expression fold variation of at least three independent determinations (* p < 0.05). The expression of osteogenic markers was induced mainly with BMP-4. BMP-9 induced runx2 and ALP (2-fold) and BMP-14 induced solely ALP (5-fold) and bone sialoprotein (3-fold).

4. Discussion

In this work we report a straightforward methodology for the expression of recombinant human BMP-4, BMP-9, BMP-10, BMP-11/GDF-11 and BMP-14/GDF-5 in *Escherichia coli*, that showed bioactivity in both C2C12 cell line and human adult stem cells.

Although the concept of overexpressing BMPs in *E. coli* is not novel, we have shown, for the first time, the overexpression of functional BMP-9, -10 and -11 using *E. coli* and characterized the bioactivity of BMP-4, BMP-14 and BMP-9 using primary cultures of human adult stem cells, which has not been shown before. Using a single-step method, which was previously reported for the solubilization of rhBMP-2 (Bessa et al. 2008c), the recovery with 0.7 M L-arginine buffer resulted in monomer, dimer and polymer, in a similar way to former reports (Klösch et al. 2005; Long et al. 2006), with up to 50 mg purified BMPs per liter of culture broth.
All tested BMPs have shown evidence of bioactivity in C2C12 cells. C2C12 cell line is a well-known model for studying the osteogenic differentiation induced by BMPs as has been reported by several investigators (Vallejo et al. 2002; Long et al. 2006; Bessa et al. 2008a). The different BMPs induced several early markers of BMP signaling (smad-1/-5, runx2/cbfa-1 or osterix) and of bone mineralization (bone sialoprotein, osteopontin or ALP).

Semi-quantitative RT-PCR analysis of mRNA expression was selected since it is well established that upon binding of the BMPs to their cell receptors, the expression of these genes is highly induced, triggering osteogenic differentiation (Bessa et al. 2008a). In C2C12 cells, BMP-4, -9 and -14 appeared to have had the most significant effects in the markers of bone formation. BMP-11 solely induced significantly the BMP-activated transcription factor runx2, showing that probably this BMP does not have any important role in bone formation and healing, besides those described in the literature during embryonic development (Andersson et al. 2006). In human stem cells, BMP-4, -9 and -14 also demonstrated bioactivity, despite the fact that the osteogenic potential was higher for BMP-4 than BMP-9 or -14, which could point for the stronger bone-inducing potential of this cytokine.

Despite the fact that these recombinant BMPs show bioactivity, further studies will be performed to assess the detailed response of cells to BMPs, including the induction of the different markers by Real-Time RT-PCR, and immunostaining analysis.

In conclusion, the approach we detailed herein shows a promising route for producing large amounts of several different bioactive recombinant BMPs. The results in human adult stem cell cultures suggest that some of these BMPs could be used with success for future clinical applications.
Chapter V. Expression, purification and bioactivity of recombinant human BMP-4, -9, -10, -11 and -14

References


Chapter VI

Silk fibroin microparticles as carriers for delivery of human recombinant BMPs. Physical characterization and drug release

This Chapter is based on the following publication:

Chapter VI
Silk fibroin microparticles as carriers for delivery of human recombinant BMPs. Physical characterization and drug release

Abstract

Bone morphogenetic proteins (BMPs) are cytokines with strong ability to promote new bone formation. Herein, we report the use of silk fibroin microparticles as carriers for the delivery of BMP-2, BMP-9 or BMP-14. BMP-containing fibroin microparticles were prepared by a mild methodology using dropwise addition of ethanol, exhibiting mean diameters of 2.7 ± 0.3 μm. Encapsulation efficiencies varied between 67.9 ± 6.1 % and 97.7 ± 2.0 % depending on the type and the amount of BMP loaded. Release kinetics showed that BMP-2, BMP-9 and BMP-14 were released in two phases profile, with a burst release in the first two days followed by a slower release, for a period of 14 days. The release data were best explained by Korsmeyer’s model and the Fickian model of drug diffusion. Silk fibroin microparticles can offer a promising approach for the sustained delivery of different BMPs in tissue engineering applications.

1. Introduction

Bone morphogenetic proteins (BMPs) have recently sparked great interest in the tissue engineering field due to their strong ability to promote new bone and cartilage formation (Reddi, 2005; Bessa et al., 2008a). Different materials have been proposed as carriers for BMPs (Bessa et al., 2008b). Carriers are used to increase the lifetime, stability and bioactivity of the BMPs and release these growth factors in both timely and site-specific sustained ways. Microparticles as carriers can offer the advantages of a controlled release, protecting the protein from loss of bioactivity and targeting the delivery into the specific injury site, with no dispersion into surrounding tissues.

Several different microparticulated systems have been investigated for the delivery of BMPs, most based on polylactic-co-glycolic acid (PLGA) (Lee et al., 1994; Phillips et al., 2006; Wei et al., 2007), on collagen (Wang et al., 2003, Itoh et al., 2004) or on dextran (Chen et al., 2006, 2007). Silk is an attractive choice as a material for bone tissue engineering, due to being biocompatible, slowly biodegradable and possessing excellent mechanical properties such as tensile strength and rigidity which are highly desirable for a potential use as a scaffold for bone tissue engineering (Scheibel 2006). Silk fibroin has been investigated as a carrier for BMPs in
several contributions by Kaplan and colleagues, in form of membranes for healing cranial
defects in mice (Karageorgiou et al., 2004), electrospun nanofibers (Li et al., 2006), and
scaffolds used for critical size defects in rats (Kirker-Head et al., 2007) and mice (Karageorgiou
et al., 2006).

Fibroin has been recently described to produce microparticles when precipitated with
ethanol at specific fibroin concentrations and freeze temperature (Nam and Park, 2001; Cao et
al., 2007; Zhang et al., 2007). Ethanol causes a shift in the random coil structure of fibroin to an
insoluble beta-sheet configuration (Nam and Park, 2001). The addition of ethanol to the fibroin
solution causes it to either precipitate to form a gel that can be used as a scaffold for tissue
engineering (Nazarov et al., 2004; Tamada, 2005) or to form a gel consisting of micrometric
particles (Nam and Park, 2001; Zhang et al., 2007). If these variables are controlled, a control
of particle size is possible and could be adapted to specific clinical applications. Therefore, we
were interested in using this system as a possible delivery micro-carrier for BMPs.

This study explores the use of silk fibroin microparticles as a potential delivery system for
the release of different BMPs. Fibroin microparticles were produced under very mild
conditions, with a concomitant loading of BMP-2, BMP-9 or BMP-14. Besides the fact that
BMP-2 has been widely researched as a osteogenic factor (Bessa et al., 2008b), we also
evaluated the potential of the fibroin particles to deliver BMP-9 and BMP-14 due to their
potential in novel therapeutical applications. BMP-9 is reported to be one of the most
osteogenic BMPs (Kang et al., 2004) and BMP-14 is used in tendon and ligament regeneration
(Nakamura et al., 2003; Dines et al., 2007), in spinal fusion (Spiro et al., 2001), bone formation
(Simank et al., 2004; Kadamatsu et al., 2008) and is critical for digit and limb development
(Merino et al., 1999). The particles were characterized for morphology, size distribution, water
uptake, encapsulation efficiency and the release kinetics of the three BMPs, with different
amounts of loaded growth factor.

2. Materials and methods

2.1. Materials

Silk containing Bombyx mori cocoons were purchased from Halcyon Yarn, US. Human
BMP-2, expressed in Chinese Hamster Ovary (CHO) cells, was purchased from Wyeth, UK.
Human BMP-9 and BMP-14 were cloned and expressed in Escherichia coli, purified by
histidine-tag affinity chromatography and freeze-dried, as described elsewhere (Bessa et al.,
2009). All chemicals were of analytical grade and used as received.
2.2. Methods

2.2.1. Preparation of BMP-loaded silk fibroin microparticles

Isolation of silk fibroin from Bombyx mori cocoons

Silk fibroin was isolated from *Bombyx mori* cocoons following a degumming protocol with modifications (Zhang et al., 2007). Briefly, the cocoon shells of silkworm *Bombyx mori* were degummed twice in a boiling solution of 0.5 % Na$_2$CO$_3$ for 0.5 h, at 90 ºC, and the resulting degummed fibers were subsequently dissolved, in a ternary system, i.e. a mixed solution of calcium chloride, ethanol and water (CaCl$_2$ / C$_2$H$_5$OH / H$_2$O: 1:2:8 mole ratio) at 90ºC for 6 h. The solution was then filtered (0.22 μm, Rotilab, Germany), and the solution was extensively dialyzed, using a cellulose semi-permeable (molecular weight cut-off = 12 000 kDa) membrane, against distilled water to remove CaCl$_2$, smaller molecules and impurities. The fibroin was lyophilized and used for the preparation of microparticles.

Microparticle preparation

Silk fibroin microparticles were prepared by ethanol precipitation according to the method described by Cao et al. (2007), with modifications. A solution of 1% (w/v) silk fibroin in phosphate buffer saline (PBS, pH 7.4) was prepared (5 ml) and filtered with a 0.22 μm PVDF membrane (Rotilab, Germany). For preparation of BMP loaded particles, 5 or 50 μg/ml of growth factor (either BMP-2, BMP-9 or BMP-14) were added to the fibroin suspension, corresponding to a smaller or a larger dose of loaded BMP (0.5 or 5 μg of BMP per mg of fibroin, respectively). The solution was incubated at room temperature with stirring (600 rpm) for 20 min. Absolute ethanol was added dropwise to a ratio of 1:2 to the initial volume of silk fibroin, and the solution incubated overnight at -20 ºC. Different ratios (ethanol to initial volume of fibroin suspension) were also tested to investigate the change in particle size. The resulting microparticles were collected by centrifugation at 4000 g, 4ºC, for 5 min, washed twice with distilled water and lyophiliized, after snap-freezing in liquid nitrogen, in a Christ Alpha 1-4 (Linder, Austria) lyophilizer. Sterile conditions were used during the entire procedure.

2.2.2. Physical characterization

*Scanning electron microscopy (SEM)*

Fibroin particles were morphologically characterized by scanning electron microscopy (SEM). SEM analysis was performed on gold-coated air-dried samples (Agar Sputercoater 108,
Essex, UK) and using a Philips XL20 microscope (Philips, Eindhoven, The Netherlands). Particle size measurements were obtained from different micrographs acquired in the SEM.

**Size distribution**

Measurements of particle size were performed on freshly prepared samples by dynamic light scattering (DLS), using a Zetasizer NanoZS (Malvern, UK). For the analysis, each sample was diluted to an appropriate concentration with ultrapure (filtered) water. Each analysis lasted 60 s and was performed at 25 °C.

**Water uptake and degradation studies**

The hydration degree of unloaded fibroin microparticles was evaluated after immersion into PBS, at 37 °C, for periods up to 30 days (30 min, 4 h, 2 d, 5 d, 8 d, 14 d, 21 d and 30 d). The weight loss of unloaded fibroin particles was assessed after immersion into PBS, at 37 °C, for periods up to 30 days (1 d, 2 d, 5 d, 8 d, 14 d, 21 d and 30 d). All experiments were performed in triplicate. Percentage of water uptake (WU) after each time period of immersion (t) was calculated using the following equation:

\[
WU(\%) = \left( \frac{W_w - W_i}{W_i} \right) \times 100
\]

The percentage of mass remaining (MR) after each time period (t) was calculated using the following equation: WL (\%) = \left( \frac{W_d}{W_i} \right) \times 100

‘\(W_w\)’ and ‘\(W_d\)’ correspond, respectively, to the weight of fibroin particles in wet state (after removal from solution, washing with distilled water and soft-blotting with filter paper) and dry state (after removal of solution, washing after distilled water and freeze-drying); ‘\(W_i\)’ correspond to the initial dry weight of particles, before immersion.

**2.2.3. In vitro release studies**

**Quantification of protein**

For measurement of BMP-2 concentration, a BMP-2 sandwich-type ELISA kit (Eubio, Austria) was used, following the manufacturer’s instructions. A calibration curve was obtained using standard preparations of BMP-2 of known concentration. For detection of BMP-9 and BMP-14, the sample concentrations were estimated by dot blot with the use of an anti-histidine tag antibody (Sigma, US). The blot intensities were compared with standards of known concentration, using image analysis (ChemiImage 4400, Alpha Innotech, US).
Determining the loading capacity and encapsulation efficiency

The microparticle encapsulation efficiency was determined upon their separation from the aqueous preparation medium containing the non-associated protein (free BMP) by centrifugation (10,000 g, 10 min). Theoretical drug content (total BMP) was calculated assuming that the entire amount of drug added to the fibroin solution was encapsulated and no drug loss occurred at any stage of the particle preparation. All experiments were performed in triplicate. The microparticle encapsulation efficiency (E.E.) and BMP loading capacity (L.C) were calculated using the following equations:

\[
E.E. (\%) = \frac{\text{Total BMP amount (μg)} - \text{Free BMP amount (μg)}}{\text{Total BMP amount}} \times 100
\]

\[
L.C. (\text{μg BMP / mg particles}) = \frac{\text{Total BMP amount (μg)} - \text{Free BMP amount (μg)}}{\text{Microparticles weight (mg)}}
\]

In vitro release

Silk fibroin particles loaded with BMP-2, BMP-9 or BMP-14 (initial loading of 0.5 or 5 μg /mg) were suspended in 5 ml PBS (pH 7.4), at 37 °C, with 200 rpm agitation, to a concentration of 2 mg/ml of microparticles. At pre-determined time points (1 h, 4 h, 1 d, 2 d, 4 d, 8 d and 14 d), 250 μl of solution was removed, centrifuged (10,000 g, 10 min) and filtered in 0.22 μm membranes and stored at -20 °C until quantification. The volume of solution removed was replaced with fresh buffer. Sterile conditions were applied to prevent any contamination of samples. All experiments were performed in triplicate. Percentage of release was obtained by comparing the measured growth factor content with the actual loading capacity.

Release kinetics models

To study the release kinetics, the data obtained from the in vitro release was treated accordingly to zero order as cumulative amount of drug released vs. time (equation 1), first order as log cumulative percent drug remaining vs. time (equation 2), Higuchi kinetics as cumulative percent drug released vs. square root of time (equation 3) and Korsmeyer kinetics as log cumulative percent drug released vs. log time (equation 4) (Huguchi, 1963; Chowdary and Ramesh, 1993; Hadjioannou et al., 1993; Bourne, 2002):

\[
R = k_1 t \\
\log UR = k_2 t 2.303 \\
R = k_3 t 0.5 \\
\log R = \log k_4 + n \log t
\]
where \( R \) and \( UR \) are released and unreleased percentages, respectively, at time \( (t) \); \( k_1, k_2, k_3 \) and \( k_4 \) are the rate constants of zero-order, first order, Higuchi and Korsmeyer models, respectively; \( n \) is an exponent that characterizes the mechanism of release.

2.2.4. Statistical analysis

Experiments were performed in triplicate \((n = 3)\) and mean ± standard deviations were reported. Student’s \( t \) test was used for statistical analysis using a two-tailed paired test. Statistical significance was defined as \( p < 0.01 \) for a 99% confidence.

3. Results

3.1. Morphology, size distribution and encapsulation efficiency

The morphology of fibroin particles observed by SEM is shown in Figure 1. Fibroin formed spherical microparticles with diameters increasing when the ratio of ethanol to silk solution (used during the preparation of particles) increased. The particles had mean sizes of 580.0 ± 120.6 nm using a 1:2 ratio of ethanol and silk, increasing up to 1.1 ± 0.3 μm with a 1:3 ratio \((p < 0.01)\), and to 1.2 ± 0.4 μm with 1:4 ratio (not significant). Using size distribution, the particles, using a 1:2 ratio, showed an average size of 2.7 ± 0.3 μm, in wet state (Figure 2). The largest part of microspheres (88.5 %) had sizes ranging from 1.5 to 3.0 μm in diameter. The encapsulation efficiency of BMP-2 into the fibroin particles was 97.7 ± 2.0 % per total amount of growth factor added during the preparation of the particles, when 0.5 μg BMP-2 (per mg fibroin) was loaded, and 76.8 ± 3.5 %, when 5.0 μg of BMP-2 were loaded (Table 1). The loading capacity was 0.69 ± 0.06 μg of BMP-2 per mg of fibroin particles, and 5.4 ± 0.5 μg of BMP-2 per mg of particles, when the different amounts were loaded. For BMP-9, the encapsulation efficiency was 90.2 ± 5.9 % and 72.4 ± 4.4 %, and the loading capacity was 0.63 ± 0.17 and 5.1 ± 0.5 μg/mg of particles, when 0.5 or 5.0 μg BMP-9 were loaded, respectively. For BMP-14, the encapsulation efficiency was 82.5 ± 6.0 % and 67.9 ± 6.1 %, and the loading capacity was 0.60 ± 0.17 and 4.8 ± 0.6 μg of BMP-14 per mg of particle weight, when 0.5 or 5.0 μg BMP-14 were loaded, respectively.
Figure 1. SEM micrographs of unloaded fibroin microparticles, produced with different ethanol : fibroin solution ratios: A) 1:2 ratio, B) 1:3 ratio, and C) 1:4 ratio. Magnification 10.000 x.
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Figure 2. Size distribution of unloaded silk fibroin particles. Almost all particles showed sizes ranging between 1.5 and 3.0 μm, with a mean diameter of 2.7 ± 0.3 μm. Size refers to particle diameter (nanometers).

Table 1. Loading and encapsulation efficiencies of BMP loaded fibroin microparticles (mean ± SD, n = 3).

<table>
<thead>
<tr>
<th>Growth factor</th>
<th>Amount of growth factor per mg of fibroin, prior to loading</th>
<th>Encapsulation efficiency (%)</th>
<th>Loading capacity (μg BMP / mg particles)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMP-2</td>
<td>0.5 μg</td>
<td>97.7 ± 2.0</td>
<td>0.69 ± 0.06</td>
</tr>
<tr>
<td>BMP-9</td>
<td>0.5 μg</td>
<td>90.2 ± 5.9</td>
<td>0.63 ± 0.17</td>
</tr>
<tr>
<td>BMP-14</td>
<td>0.5 μg</td>
<td>85.8 ± 6.0</td>
<td>0.60 ± 0.17</td>
</tr>
<tr>
<td>BMP-2</td>
<td>5.0 μg</td>
<td>76.8 ± 3.5</td>
<td>5.4 ± 0.5</td>
</tr>
<tr>
<td>BMP-9</td>
<td>5.0 μg</td>
<td>72.4 ± 4.4</td>
<td>5.1 ± 0.5</td>
</tr>
<tr>
<td>BMP-14</td>
<td>5.0 μg</td>
<td>67.9 ± 6.1</td>
<td>4.8 ± 0.6</td>
</tr>
</tbody>
</table>

Encapsulation efficiency (%) = \([\text{Total BMP amount (μg)} - \text{Free BMP (μg)}] / \text{Total BMP (μg)}\) x 100.
Loading capacity (μg BMP / mg particles) = \([\text{Total BMP amount (μg)} - \text{Free BMP (μg)}] / \text{Microparticles weight (mg)}\).
3.2. Water uptake and degradation studies

The swelling occurred rapidly during the first two days of immersion, up to 48.5 ± 6.7 % after 30 min and up to 167.7 ± 14.1 % after day 2 (p < 0.01), with water uptake reaching a peak at day 8 with 227.7 ± 26.6 % (p < 0.01) (Figure 3). This was followed by a slow and gradual decline in wet weight, with a swelling value of 203.4 ± 25.0 % (non-significant) after day 30, possibly due to weight loss. The remaining mass was 97.1 ± 1.5 % after day 5, 95.4 ± 2.5 % after day 14, and 93.5 ± 3.3 % after day 30 (Figure 4).

![Water uptake graph](image)

**Figure 3.** Evolution of water uptake behavior of unloaded silk fibroin microparticles in phosphate buffer saline as a function of immersion time (t). (mean ± SD, n = 3). Water uptake increased significantly up to 5 days, peaked after 8 days, and then slightly declined (non-significantly). *p < 0.01 related to the difference between consecutive time measurements.

3.3. *In vitro* release studies

Drug release studies were evaluated when fibroin particles loaded with BMP-2, BMP-9 and BMP-14 were immersed in PBS, at 37ºC, with agitation. The microparticles loaded with 0.5 μg of BMPs /mg fibroin, showed an initial burst release with more than half of protein being released during the first two days (64.1 ± 5.1 % for BMP-2, 78.4 ± 7.9 % for BMP-9 and 62.5 ± 7.3 % for BMP-14), followed by a second phase of a slower and more sustained release with 90.1 ± 6.9 % of BMP-2, 90.9 ± 5.2 % of BMP-9, and 80.4 ± 5.6% of BMP-14, released after 14 days (Figure 5A). This corresponds to a release rate, of 15 ng/day, 7 ng/day and 8 ng/day.
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(per mg of fibroin), between day 2 and 14, for each BMP, after a burst release of 442 ng, 499 ng and 377 ng of each BMP (in 2 days).

The particles loaded with 5 µg/mg of BMPs showed an initial release of about 28.6 ± 5.5 % of BMP-2, 22.7 ± 6.3 % of BMP-9 and 14.1 ± 6.7 % of BMP-14 being released after two days (Figure 5B), which corresponds to 1.5 µg, 1.1 µg and 0.7 µg of each growth factor. After 14 days, 47.3 ± 7.0 % BMP-2, 38.9 ± 8.9 % BMP-9, and 33.2 ± 8.8 % BMP-14 were released, corresponding to a release rate of 86 ng/day, 72 ng/day and 76 ng/day, for each BMP. At the end of this period, 2.5 µg, 2.0 µg and 1.5 µg of each growth factor were (cumulatively) released from the particles (per mg of fibroin).

The release data of the BMPs were treated accordingly to first order, Higuchi and Korsmeyer equations (Table 2) and zero order equation (data not shown). The release data of BMP-2, BMP-9 and BMP-14 were best explained by Korsmeyer’s equation ($r^2$ varying between 0.92 and 0.98), followed by Higuchi’s equation ($r^2$ varying between 0.75 and 0.95) and by first order kinetics ($r^2$ varying between 0.71 and 0.91). The release exponent “n” of Korsmeyer’s model, varied in the different experimental conditions between 0.30 and 0.42. Since $n < 0.43$ the release data fitted a Fickian mode of diffusion (Ritger and Peppas, 1987). The zero order modeling showed low linearity (data not shown).

Weight loss

![Weight loss graph](image)

Figure 4. Evolution of weight loss of unloaded fibroin particles in phosphate buffer saline as a function of immersion time (t). (mean ± SD, n = 3). The remaining mass decreased to 97.1 ± 1.5 % after day 5, 95.4 ± 2.5 % after day 14, and 93.5 ± 3.3 % after day 30.
Figure 5. Release kinetics of BMP-2, BMP-9 and BMP-14 immobilized in fibroin particles. ELISA was used for BMP-2 quantification and dot-blot for BMP-9 and BMP-14 quantification (Mean ± SD, n = 3). A) Release from particles containing 0.5 μg of BMP per mg of fibroin; B) release from particles containing 5.0 μg of BMP per mg of fibroin. Cumulative release is expressed in percentage of loaded protein.
**Table 2.** Release kinetics of BMPs immobilized in fibroin particles. First order (log cumulative percent drug remaining vs. time), Higuchi kinetics (cumulative percent drug released vs. square root of time) and Korsmeyer kinetics (log cumulative percent drug released vs. log time), with correlation values ($r^2$).

<table>
<thead>
<tr>
<th>Growth factor</th>
<th>Amount of growth factor per mg of fibroin, prior to loading</th>
<th>Equation modeling (First order (^a), Higuchi (^b), Korsmeyer (^c))</th>
</tr>
</thead>
</table>
| BMP-2         | 0.5 μg                                                   | \[ y = -0.057x + 1.7547 ; r^2 = 0.91 \(^a\) \]
|               |                                                          | \[ y = 17.926x + 26.183 ; r^2 = 0.85 \(^b\) \]
|               |                                                          | \[ y = 0.3029x + 1.6459 ; r^2 = 0.97, n = 0.30 \(^c\) \]          |
| BMP-9         | 0.5 μg                                                   | \[ y = -14.148x + 25.648 ; r^2 = 0.71 \(^a\) \]
|               |                                                          | \[ y = 19.509x + 25.325 ; r^2 = 0.75 \(^b\) \]
|               |                                                          | \[ y = 0.3674x + 1.6277 ; r^2 = 0.92, n = 0.37 \(^c\) \]          |
| BMP-14        | 0.5 μg                                                   | \[ y = -0.0344x + 1.8002 ; r^2 = 0.70 \(^a\) \]
|               |                                                          | \[ y = 18.232x + 17.785 ; r^2 = 0.80 \(^b\) \]
|               |                                                          | \[ y = 0.4195x + 1.519 ; r^2 = 0.92, n = 0.42 \(^c\) \]          |
| BMP-2         | 5.0 μg                                                   | \[ y = -0.0133x + 1.9131 ; r^2 = 0.71 \(^a\) \]
|               |                                                          | \[ y = 11.018x + 8.4402 ; r^2 = 0.88 \(^b\) \]
|               |                                                          | \[ y = 0.385x + 1.2845 ; r^2 = 0.98, n = 0.39 \(^c\) \]          |
| BMP-9         | 5.0 μg                                                   | \[ y = -77.226x + 150.34 ; r^2 = 0.77 \(^a\) \]
|               |                                                          | \[ y = 8.5682x + 7.496 ; r^2 = 0.92 \(^b\) \]
|               |                                                          | \[ y = 0.301x + 1.2361 ; r^2 = 0.96, n = 0.30 \(^c\) \]          |
| BMP-14        | 5.0 μg                                                   | \[ y = -0.0083x + 1.9499 ; r^2 = 0.83 \(^a\) \]
|               |                                                          | \[ y = 7.2919x + 5.3548 ; r^2 = 0.95 \(^b\) \]
|               |                                                          | \[ y = 0.2978x + 1.1429 ; r^2 = 0.97, n = 0.30 \(^c\) \]          |
4. Discussion

Microparticles as drug carriers have the advantages of sustained or controlled release, the possibility of drug targeting to specific tissues, thus reducing side effects of drugs and improving their bioavailability. Therefore, microparticles as delivery systems have drawn much attention in the pharmaceutical field and have been successfully applied in several clinical trials. The goal of this study was to assess the possibility of the use of silk-derived fibroin as a microparticle carrier for the release of different BMPs.

Fibroin microspheres were manufactured in a semi-aqueous system, avoiding the use of toxic organic solvents, harsh conditions, cross-linking agents or surfactants, as a way of minimizing the risk of a loss of BMP activity during the particle preparation. The group of Kaplan and colleagues have recently reported that fibroin could form either particles or films with methanol and conserving the activity of incorporated drugs (Hofmann et al., 2006; Wang et al., 2007). When compared with other methods of processing, such as spray drying (Hino et al., 2003), this approach not only avoids conditions unfavorable for the growth factor activity such as high temperature, but it also allows generating particles with smaller size.

The microparticles were easily prepared under sterile conditions and had a small size in a rather narrow size distribution. The size of particles was significantly smaller in the SEM counterpart images, probably due to the large swelling of the particles in the wet state. This was reported also in a former work (Cao et al., 2007). The small particle size presented a double advantage, first by potentially enhancing the drug availability in the particle surface, and also by providing a surgical material that may be suitable for injectable implantation in bone defects in a slurry form. The BMPs were successfully encapsulated with a rather high efficiency, which is comparable to other works (Patel et al., 2008; Wenk et al., 2008). The lower encapsulation efficiencies observed for BMP-14 could be due to the differences in the BMP charge and its interaction with the fibroin material. Interestingly, this BMP has a lower isoelectric point when compared to BMP-2 (calculated using Expasy Proteomics tool Compute pI/Mw, at www.expasy.org; pI = 6.3 for BMP-2, pI = 5.9 for BMP-9 and pI = 5.8 for BMP-14). It is possible that this growth factor interacted to a weaker extent with the fibroin carrier since fibroin has an acidic isoelectric point (pI = 5.5).

During the initial stages of swelling, the fibroin microspheres showed substantial water uptake, mostly during the first two days of immersion, which also corresponded to the initial burst release, as the drug was rapidly released from the particles. Then, as swelling became gradually counterbalanced, the BMP was released slower and the release was mainly determined by the diffusion of the growth factor from the particle to the medium. It is possible
that the erosion of the carrier contributed little to this, since we have found both that the release data best fitted with the Korsmeyer’s equations and slope values between 0.30 and 0.42, thus indicating that the release was of a Fickian type of diffusion, controlled by only one mechanism and thus not dependent on carrier erosion (this applies when $n < 0.43$, in the case of spherical particles) (Ritger and Peppas, 1987). In fact, as we have observed the silk fibroin microparticles exhibited a slow rate of degradation. The release amounts during this phase were such that they could meet the concentrations of growth factor required in most in vitro and in vivo applications (in the range of few hundreds of nanograms to a few micrograms), by choosing a specific amount of BMP during the loading process (Kenley et al., 1994; Hosseinkhani et al., 2007; Wei et al., 2007; Bessa et al., 2009).

The silk fibroin microspheres have shown to work as a microcarrier that provides a sustained release of the different BMPs in a timewise manner that could be used in bone regenerative applications. The bioactivity of the BMPs released from the silk fibroin microparticles will be explored in future works.
Chapter VI. Silk fibroin microparticles for delivery of BMPs. Physical characterization and drug release

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Chapter VII

Silk fibroin microparticles for delivery of BMP-2. In vitro and in vivo bioactivity

This Chapter is based on the following publication:

Chapter VII

Silk fibroin microparticles as carriers for delivery of human recombinant BMP-2. In vitro and in vivo bioactivity

Abstract

Bone morphogenetic proteins (BMPs) are growth factors with an important role in bone formation. In this study, we evaluated the use of silk fibroin microparticles as a novel delivery carrier for BMP-2, in vitro and in vivo in a rat model for ectopic bone formation. The particles did not exhibit significant cytotoxicity. In murine C2C12 cells, the fibroin particles with entrapped BMP-2 induced osteogenic mineralization and an increase in alkaline phosphatase activity. In the rat ectopic model, bone formation was observed by in vivo μCT after 2 and 4 weeks post-implantation, both with 5 and 12.5 μg BMP-2, with an increase in bone density over time. Histology revealed further evidence of ectopic bone formation, observed by strong Alizarin Red staining and osteocalcin immunostaining. Our findings showed that fibroin microparticles may be used for the sustained delivery of BMP-2, allowing the formation of new bone.

Introduction

Bone morphogenetic proteins (BMPs) are a group of cytokines from the transforming growth factor-beta (TGF-β) superfamily that have been used as powerful osteoinductive components in several late-stage tissue engineering products for bone grafting (Reddi et al. 2005, Bessa et al. 2008a, Bessa et al. 2008b). Currently, two devices using BMPs have been approved by the FDA for human clinical application based on the use of collagen sponges (Gautschi et al. 2007). However, as collagen poses several drawbacks such as sub-optimal handling conditions and the risk of disease transmission, there is still a need for an optimized BMP delivery system (Kirker-Head et al. 2007; Bessa et al. 2008).

The search for efficient, simple and cheap delivery systems for drug targeting has lead to a great investment in the area of nano and microparticles for drug delivery. In tissue engineering, these systems are excellent choices for growth factor delivery, since they can be easily prepared and sterilized. They can be processed into injectable systems allowing an easy and non-invasive implantation in the patient. Silk offers one versatile alternative as a natural biocompatible and biodegradable protein polymer that has been shown to efficiently deliver...
BMPs in a variety of works (Karageorgiou et al. 2004, 2006, Li et al. 2006, Kirker-Head et al. 2007).

We have been studying a fibroin-based microparticle system as a potential new carrier for loading and release of BMPs (Bessa et al. 2008c). In the present study, we examined the bioactivity of BMP-2 loaded fibroin microparticles in vitro in C2C12 cells, by quantification of the alkaline phosphatase (ALP) activity and calcium mineralization, and in a rat ectopic bone formation model, using in vivo μCT and histological analysis.

2. Materials and Methods

2.1. Materials

Silk-containing Bombyx mori cocoons were purchased from Halcyon Yarn, US. The silk fibroin was isolated from the cocoons following an adapted degumming protocol (Zhang et al. 2007). Fibroin microparticles were prepared using the method of by Cao and Colleagues (2007), with modifications. Briefly, a solution 1% (w/v) fibroin in phosphate buffer saline was prepared, to which 0.5 μg BMP-2 (Wyeth, UK) was loaded. This solution was incubated at room temperature with stirring (600 rpm), while absolute ethanol was added dropwise to a ratio of 1:2 to the initial volume of fibroin. The solution was incubated overnight at -20 ºC, washed with distilled water, collected by centrifugation and freeze-dried. The characterization of the fibroin microparticles by scanning electron microscopy (SEM), dynamic light scattering (DLS), swelling behavior, and the release profile of BMP-2, was described elsewhere (Bessa et al. 2008c). All other chemicals were of analytical grade.

2.2 Methods

2.2.1. Bioactivity in C2C12 cells

C2C12 cells have been used for screening the osteogenic activity of BMPs in a variety of works (Klösch et al. 2004, Long et al. 2006, Bessa et al. 2008d, Bessa et al. 2009). These cells do not express significant amounts of endogenous BMPs, thus making them an effective model for testing the activity of the released BMP-2 (Katagiri et al. 1994). C2C12 cells were seeded at $10^5$ cells/ml per well in a 24-well plate, attached in Dulbecco’s modified Eagle’s medium (DMEM), with 1% (v/v) fetal calf serum, at 37 ºC, with 5% CO₂ in a humidified environment. The cells were incubated with unloaded particles and with particles containing 0.5 and 2.5 μg loaded BMP-2 (Wyeth, UK). BMP-2 was added as a positive control, to the culture medium, at 0.1, 0.5 and 1.0 μg/ml. As another control, unloaded particles which were immersed for 30 min in 5 μg/ml BMP-2 and washed twice, were also added to the cells (adsorbed BMP control).
Alkaline phosphatase (ALP) enzymatic activity was measured accordingly to standard procedures (Salgado et al. 2002) after 5 days in C2C12 cultures. For the Alizarin Red mineralization assay, C2C12 cells were seeded at 3 x 10^4 cells/ml per well and cultured in DMEM supplemented with 100 μM ascorbic acid and 10 mM β-glycerophosphate, in the presence of both unloaded and BMP-2 loaded microparticles. Media and particles were replaced every 5 days. After 14 days of culture, the cells were washed for three times with PBS (without Ca^{2+}, Mg^{2+}) and fixed in 4% (v/v) formaldehyde, for 30 min. After removal of the fixative, cells were washed twice with distilled water and covered with Alizarin Red 1% solution, followed by gentle agitation in an orbital shaker for 10 min. The solution was then removed and the cells washed three more times with distilled water. The cells were observed using a Axiovert10 (Zeiss, Germany) optical microscope, and imaged with a coupled Coolpix950 (Nikon, Japan) camera.

2.2.2. MTS cell viability assay

Human osteosarcoma cells (SaOS-2 cell line; ECACC) were used for cell viability tests instead of murine C2C12 cell line, since these are a well-established human cell line of an osteoblast phenotype. Human osteosarcoma cells were seeded at 10^6 cells/ml in a 24-well plate, attached overnight in Dulbecco's modified Eagle's medium (DMEM) with 1 % fetal calf serum and no antibiotics, at 37 ºC, 5 % CO_2, in a humidified environment, and cultured for 1, 3 and 5 days. Unloaded fibroin particles were added to the cell culture at 0.5 mg/ml. MTS cell viability assay was determined after these time periods using a standard procedure (Salgado et al. 2002). Latex rubber (Velos-Perforex, Manchester, UK) and standard culture medium were used as positive and negative controls, respectively. Latex rubber is known to have a strong cytotoxic effect leading to extensive cell death and lysis, and is commonly used as a positive control for cell death. All the samples were tested in triplicate for at least two independent experiments with reproducible results. The results are expressed as a percentage of the control (scored as 100% viability) as mean ± SD.

2.2.3. Rat ectopic bone formation model

Male Sprague-Dawley rats (weighting 340-360 g), were purchased from the Institut für Labortiertechnik und -genetik (Himberg, Austria). They were housed in light and temperature-controlled facilities and given food and water ad libitum. In three animals, surgery was performed after general anesthesia with an intramuscular injection of 60 mg/kg of body weight (BW) ketamine (Ketavet®, Pharmacia, Erlangen, Germany) and 7.5 mg/kg BW xylazine (Rompun®, Bayer, Leverkusen, Germany). After shaving and disinfecting the dorsum of the
animals, four midsagittal incisions were made dorsally in the proximal area of each limb and subcutaneous pockets were created by blunt dissection. The fibroin particles were implanted subcutaneously into each pocket. Three groups were defined: group I (unloaded particles), group II (particles + 5 μg BMP-2) and group III (particles + 12.5 μg BMP-2). The animals were randomized and were implanted with four samples \((n = 4)\) per each group. Only one group was assigned to each animal to avoid the potential interference of released BMP by blood circulation. After placement of the particles, the incisions were closed with sutures, and the wound covered with vapor-permeable spray dressing (Opsite®, Smith & Nephew, London, UK). Analgesia treatment was given post operation, of 0.01 mg/kg BW Buprenorphin. At the end of each implantation period (4 weeks), the animals were sacrificed and samples from the tissues surrounding the implantation site were harvested. The animal protocol was approved by the authority of the city government of Vienna.

2.2.4. **In vivo μCT**

Live microCT (vivaCT 75, Scanco Medical AG, Switzerland) was used to examine bone formation in individual rats. MicroCT imaging was performed at weeks 2 and 4 respectively. Rats were anesthetized for 5 min with 2 % isoflurane during measurement. No X-ray contrast media was used in this study. Image reconstruction was performed using the built-in 3D visualization software (Scanco Medical AG, Switzerland). The evaluation of newly formed bone and bone density were performed using IPL version v5.06b (Scanco Medical AG, Switzerland).

2.2.5. **Histology**

Harvested samples were fixed in neutral buffered 10 % formalin and embedded in paraffin. Three-micrometer or four-micrometer sections were cut and stained with hematoxylin and eosin (H & E) or Alizarin Red S, respectively, for light microscopy observation. For a detailed analysis of the tissue differentiation, immunohistochemical staining was performed in selected samples. Briefly, paraffin sections were dewaxed with xylene and rehydrated in a graded ethanol series. After enzymatic antigen retrieval with 0.1 % Proteinase (Sigma Aldrich, USA) sections were incubated with an Osteocalcin antibody (Santa Cruz Biotechnology, USA) in a humidified chamber at 4 ºC overnight. After rinsing, sections were incubated with the EnVision™ system (Dako, Denmark) at room temperature for 30 min. and immunoreactivity was visualized with AEC solution (Thermo Fisher Scientific, USA). Slides known to express Osteocalcin were used as positive controls, omitting of the primary antibody served as negative control.
2.2.6. Statistical analysis

Experiments were performed at least in triplicate and expressed as means ± standard deviation. Student’s t test was used for statistical analysis using a two-tailed non-paired test. Statistical significance was defined as $p < 0.01$ for a 99% confidence.

3. Results

3.1. Alkaline phosphatase in C2C12 cells

The bioactivity of BMP-2 released from the silk fibroin microparticles was studied by determining the ability to induce ALP enzymatic activity over basal levels in C2C12 cells (negative controls) after 5 days of culture, and by comparing the cell response to a BMP-2 dose-response effect directly added to the culture medium (positive control). The BMP-2 released from the fibroin particles was able to induce a significant increase in ALP activity over basal levels after 5 days ($p < 0.01$, Figure 1). Particles containing 2.5 μg of BMP-2 could induce high levels of ALP activity comparable to the positive control. Cells were observed to have differentiated into osteoblast morphology (data not shown). The fibroin particles with only adsorbed BMP-2 was able to induce solely a small increase in ALP activity. Unloaded particles did not induce any increase in ALP activity or osteoblast morphology. After two weeks of cell culture, the BMP-2-loaded fibroin particles were able to induce mineralization as observed by Alizarin red staining (Figure 2), identically to positive control (BMP-2 added to culture medium).

3.2. Cell viability

The silk fibroin microparticles did not show any significant evidence of cytotoxicity in human osteosarcoma cells, after 1, 3 or 5 days of cell culture (Figure 3). The cells were able to proliferate normally along this time period. MTS is a viability/proliferation test and an inverse relationship of toxicity to cells can be assumed (Salgado et al. 2002). Data correlates with morphological observations of the cells with optical microscopy (data not shown). Similar data was also obtained when particles were tested in primary cultures of human adult adipose derived stem cells (data not shown).

3.3. In vivo μCT

All animals showed no complications in wound healing, during the 4-week follow up. After 2 and 4 weeks, all sites of fibroin particle implantation were easily identified and retrieved for analysis by μCT. The μCT reconstructions showed clearly ectopic calcifications in all BMP-2
loaded implanted particles, which varied from 1 to 3 mm in width. The bone formed as a solid shell on part of the surface and as having a trabecular structure inside (Figure 4 and 5). We were not able to image the implantation site of group III (fibroin particles + 12.5 μg BMP-2) after 2 weeks follow-up, as the animal breathing interfered with the imaging process. Comparing the reconstructions of group II (fibroin particles + 5 μg BMP-2) between 2 and 4 weeks follow-up, large differences in amount of ectopic bone formed could be observed (Table 1). There was an increase in both the bone density and bone volume in group II (fibroin particles + 5 μg BMP-2) between 2 and 4 weeks (178.3 to 288.2 mg HA/mm$^3$ bone and 1.9 mm$^3$ to 2.5 mm$^3$) and an increase in the ratio between bone and tissue (0.025, after 2 weeks, to 0.050, after 4 weeks). The dimensions of the newly formed bone was nearly similar between the two time periods. Furthermore, the μCT imaging at the 2 weeks measurement had to be performed with a lower threshold, thereby adding evidence that a significant increase in bone formation occurred in the period between 2 and 4 weeks, in group II. At 4 weeks post-implantation, group III (fibroin particles + 12.5 μg BMP-2) revealed that ectopically formed bone an higher density (433.0 mg HA/mm$^3$ bone) compared to group II (fibroin particles + 5 μg BMP-2) (288.2 mm HA/mm$^3$ bone). No ectopic bone formation was detected in group I, where unloaded fibroin particles were implanted (data not shown).

Figure 1. Alkaline phosphatase activity of C2C12 cell line after 5 days of culture. A-C) BMP-2 soluble protein, positive control, 0.1, 0.5 and 1.0 μg/ml; D) control cells (no growth factor or particles added); E and F) unloaded particles; G) particles with loaded BMP-2 (0.5 μg), H) fibroin particles with loaded BMP-2 (2.5 μg), I-J) fibroin particles with adsorbed BMP-2. ALP activity is reported in nmol /min /mg of total protein (Mean ± SD, n = 3), * p < 0.01 relative to negative control cells (D).
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Figure 2. Alizarin red mineralization staining of showing sites of calcium-phosphate deposits (orange color) in C2C12 cell line differentiated into osteoblast after 14 days of culture with particles loaded with 0.5 μg BMP-2 (A), and 0.5 μg/ml BMP-2 added to culture medium (B). Unloaded fibroin particles did not show any mineralization (C).

![Image of mineralization staining]

Figure 3. Cell viability tetrazolium salt (MTS) test performed in SaOS-2 cells as a function of the time of cells culture with 0.5 mg/ml silk fibroin microparticles. DMEM and Latex were used as a negative and a positive control for cell death, respectively. (mean ± SD, n = 3)

![Graph of cell viability test results]
Figure 4. Live microCT analysis of ectopic bone formation induced by fibroin particles + 2.5 μg BMP-2 after 2 weeks (A) and 4 weeks (B), and fibroin particles + 12.5 μg BMP-2 after 4 weeks (C). Scale is 5 mm.
Figure 5. Detailed live microCT analysis of ectopic bone formation induced by fibroin particles + 2.5 μg BMP-2 after 2 weeks (A and B) and 4 weeks (C and D), and fibroin particles + 12.5 μg BMP-2 after 4 weeks (E). Background signal in A and B is due to a lower threshold used for bone visualization. Scale is 1 mm.
Table 1. Analysis of in vivo μCT data, showing bone size, surface area, volume and mean density. Group II (fibroin particles + 5 μg BMP-2), group III (fibroin particles + 12.5 μg BMP-2). (mean ± standard error, \( n = 2 \))

<table>
<thead>
<tr>
<th>Group</th>
<th>Group III (4 weeks)</th>
<th>Group II (4 weeks)</th>
<th>Group II (2 weeks) *</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bone density (mg HA/mm(^3))</td>
<td>433.0</td>
<td>288.2 ± 20.8</td>
<td>178.3 ± 11.3</td>
</tr>
<tr>
<td>Bone volume (BV) (mm(^3))</td>
<td>1.2</td>
<td>2.5 ± 0.5</td>
<td>1.9 ± 0.6</td>
</tr>
<tr>
<td>Tissue volume (TV) (mm(^3))</td>
<td>89.6</td>
<td>47.9 ± 18.3</td>
<td>76.6 ± 18.9</td>
</tr>
<tr>
<td>Bone volume ratio (BV/TV)</td>
<td>0.013</td>
<td>0.050 ± 0.001</td>
<td>0.025 ± 0.002</td>
</tr>
</tbody>
</table>

* the threshold used for bone detection in μCT was lower than that used for the measurements at 4 weeks

3.4. Histology

H&E staining revealed no signs of vascularization, but in group II some parts showed calcified tissue. In this group, H&E staining revealed abundant presence of osteoblasts and osteocytes surrounded by an extracellular matrix (Figure 6A). These tissues showed intense Alizarin Red staining (Figure 6B). In group III, no Alizarin Red positive areas could be observed, despite the fact that in the μCT ectopic bone formation was observed. At the time of harvest it was not possible to localize where the new bone formation was occurring. No staining was detected in the control group as expected (data not shown). Immunohistological evaluation with an osteocalcin-specific antibody revealed moderate staining in sections of group II (Figure 6C), in the site where osteoblasts surrounding a mineralized matrix were detected. Staining was not detected in a negative control with no incubation with the osteocalcin antibody (data not shown).

4. Discussion

It is well recognized the need for an efficient delivery system for BMPs as a way of providing effective and sustained stimulation of bone formation, as shown in several experimental models (Seeherman and Wozney 2005). The main role of a carrier is to retain these growth factors at the site of injury for a prolonged time frame, protecting the
immobilized drugs from degradation and maintaining its bioactivity, whilst releasing the protein in a time and site-controlled way to promote the formation of new bone at the treatment site.

Currently, diverse synthetic and natural origin polymers have been investigated for the delivery of BMPs in bone tissue engineering. The most common includes collagen, polylactic-co-glycolic acid (PLGA), polyethylene glycol (PEG), hydroxyapatite, gelatin, chitosan, fibrin, hyaluronic acid, and silk fibroin, generally in hydrogels, scaffolds or microparticles form (Bessa et al. 2008b). Fibroin-based biomaterials have been explored as novel alternatives for drug delivery and in tissue engineering as a result of their biocompatibility, biodegradability, mechanical strength, and versatility of formats into which the material could be processed (Lammel et al. 2008). In the present study we examined the in vitro and in vivo delivery of BMP-2 from fibroin microparticles.

The basis for evaluating the activity of the BMP released by the fibroin particles was to test initially particles loaded with the growth factor in C2C12 cell line and then using a rat ectopic bone formation model. C2C12 murine pre-myoblast cells are well-defined by their ability to rapidly differentiate into osteoblasts when cultured in presence of BMPs (Vallejo et al. 2002, Klösch et al. 2004, Long et al. 2006). In these cells, fibroin particles loaded with BMP-2 (0.5 and 2.5 μg) were able to induce a significant increase in ALP activity (after 5 days of culture), osteoblast-like morphology and the formation of a mineralized matrix (after two weeks of culture), thereby confirming that the protein was loaded into the particles remaining in bioactive state. This adds value to the method formerly developed to produce fibroin particles by ethanol precipitation (Nam and Park 2001, Cao et al. 2007, Zhang et al. 2007), confirming that is retains the activity of loaded drugs. A similar case was reported for fibroin particles and films, obtained by methanol precipitation, that retained the activity of immobilized horseradish peroxidase and lysozyme (Hofmann et al. 2006, Wang et al. 2007). In addition, the levels of ALP activity obtained with BMP-2 loaded particles were similar to those of induced by BMP-2 added to culture medium, indicating that most of the growth factor was released over the timeframe of 5 days. Since particles with surface-adsorbed BMP-2 induced low levels of ALP, it is likely that most of the growth factor accountable for the significant increase in ALP levels is BMP-2 loaded into the particles during their fabrication. No cytotoxicity was detected by the use of fibroin particles in vitro, as inferred by the MTS cell viability assay. In general, silk fibroin has been regarded as a good biocompatible material, effecting low immune responses in vivo, and with an historical use as sutures in clinical applications (Altman et al. 2003, Meinel et al. 2005, Wang et al. 2008).
Ectopic bone formation models in rodents have been used to test the activity of BMPs in a wide variety of studies (Bulpitt and Aeschlimann, 1999, Maire et al. 2005, Zhang et al. 2005, Kato et al. 2006, Jeon et al. 2007). The microparticles could form a slurry that was easily implanted ectopically in rats. After two and four weeks, bone formation was observed in all sites of implantation of BMP-2 loaded particles and the extent of ectopically formed bone was measured by microCT. Mineralized areas of bone were found as early as after two weeks post-implantation, and increased both in bone volume and hydroxyapatite density, after four weeks. MicroCT is a reliable and non-invasive method for scan and evaluation along time of bone mass, structure, geometric and density parameters (Hangartner 2007). The use of microCT imaging maximizes the results from single animals, offering sequential data within the same animal, reducing this way the required number of animals per study (Cowan et al. 2007). Ectopic bone formation was also confirmed by histological analysis, performed at four weeks post-implantation. New bone formation was detected by the presence of a calcified extracellular matrix, which was stained by Alizarin Red, revealing only surface mineralization without quantitative data, as compared to the microCT data. In addition, osteocalcin, a calcium-binding protein and a marker of mature bone formation (Price 1989), was also detected in the cells surrounding the newly formed bone.

New bone formation occurs as released BMP-2 triggers the recruitment of stem cells that differentiate into osteoblasts and form a calcium-phosphate matrix (Cunningham et al. 1992). This can only occur if there is not only an initial stimulation of the site by BMP-2 but also a sustained released of the growth factor in the following weeks, that allows the formation of new bone to occur. In the lack of a sustained release the BMP-2 is usually cleared from the implant site within hours, supposedly terminating its local effect (Louis-Ugbo et al. 2002). This is demonstrated by studies where surface-adsorbed BMP-2 was used in the particles which failed to induce new bone formation probably due to either loss of the BMP bioactivity or insufficient duration of stimulus (Wei et al. 2007). Thus, it has been suggested that materials exhibiting an higher retention of BMPs yield increased osteoinductive activity (Uludag et al. 1999, 2000). In our contribution, the amounts of BMP-2 loaded into the fibroin microparticles (5 and 12.5 μg), are similar to the values reported in the literature to achieve ectopic bone formation in rats with BMP-2, using other delivery systems (Kenley et al. 1994, Hosseinkhani et al. 2007, Wei et al. 2007). The fibroin microparticles have clearly shown to be able to deliver BMP-2 in a sustained way, allowing therefore the formation of new bone.

In conclusion, the fibroin microparticles may constitute a promising delivery system of recombinant human BMPs, for future bone tissue engineering applications.
Figure 6. H&E staining for animals implanted with fibroin particles + 5 μg BMP-2 (group II) at 4 weeks post-implantation (A). Newly formed bone was indicated by the presence of a mineralized extracellular matrix (E.M.) which was stained with Alizarin Red (B), osteocytes (O.C.), and osteoblasts (O.B.) which were stained by osteocalcin immunostaining (C).
Chapter VII. Silk fibroin microparticles for delivery of BMP-2. In vitro and in vivo bioactivity

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Chapter VIII

Thermoresponsive self-assembled elastin-based nanoparticles for delivery of BMPs

This Chapter is based on the following publication:

Chapter VIII

Thermoresponsive self-assembled elastin-based nanoparticles for delivery of BMPs

Abstract

Elastin-like polymers are a new type of protein-based polymers that display interesting properties in the biomaterial field. Bone morphogenetic proteins (BMPs) are cytokines with a strong ability to promote new bone formation. In this work, we explored the use of elastin-like nanoparticles (237.5 ± 3.0 nm), created by thermoresponsive self-assembly, for the combined release of bone morphogenetic protein-2 (BMP-2) and bone morphogenetic protein-14 (BMP-14). These BMPs could be encapsulated at high efficiency into the elastin-like particles and delivered in a sustained way for 14 days. The activity of the growth factors was retained, as shown by the induction of ALP activity and osteogenic mineralization in C2C12 cells. Increased bioactivity was observed with a combined release of BMP-2 and BMP-14. This approach shows a significant potential for future tissue engineering applications in bone.

1. Introduction

Bone morphogenetic proteins (BMPs) have recently sparked great interest in the tissue engineering field due to its strong ability to promote bone and cartilage formation (Reddi 2005, Bessa et al. 2008a). During the process of bone formation, signals from BMPs trigger the differentiation of stem cells, which are recruited to the site of injury, into bone forming cells (Reddi 1981). In recent years, BMP-2 has demonstrated a successful regenerative potential in a variety of works in spinal disk regeneration, healing of long bone fractures, and periodontal repair (McKay et al. 2007). As part of the conceptual tissue engineering perspective, different materials have been proposed and researched as delivery carriers for BMPs (Bessa et al. 2008b). The materials for tissue engineering applications should ideally mimic the natural environment of tissues and, in this regard, natural polymers can exert signals to guide cells at the various stages of their development, accelerating healing, thus presenting excellent properties for clinical uses (Mano an Reis 2007).

Elastin-like polymers (ELPs) are repetitive polypeptides inspired on the mammalian elastin structure which consists of a pentapeptide repeat, VPGXG, where X may be any natural amino acid except proline (Chilkoti et al. 2006, Simnikc et al. 2007). The most interesting feature of
Chapter VIII. Thermoresponsive self-assembled elastin-based nanoparticles for delivery of BMPs

the ELPs is their Inverse Temperature Transition (ITT) behavior (Urry et al. 1992). Below a certain critical temperature, the transition temperature (T_t), and in the presence of water, the polymer remains soluble, with the chains relatively extended in a disordered state. Above the T_t, the polymer chains hydrophobically fold and self-assemble into a more ordered structure (Herrero-Vanrell et al. 2005, Machado et al. 2009). Poly(VPAVG), where the central glycine is replaced by an alanine, had been chemically synthesized and showed to form above its T_t, spherical microparticles, that are able to entrap active substances (Herrero-Vanrell et al. 2005, Rincón et al. 2006). The polymer self-assembling process occurs above a T_t of ~ 30 ºC, but the self-assembled structures only solubilize again when a strong cooling of ~10 ºC is achieved, a phenomenon called hysteresis (Herrero-Vanrell et al. 2005). The particles, once formed, are stable and can be used for drug delivery. In addition, ELPs demonstrate excellent biocompatibility (Urry et al. 1991). Apparently, the immune system ignores these polymers because it does not distinguish them from the natural elastin, and the degradation products are only natural amino acids.

Poly(VPAVG) can be produced by recombinant technology, in alternative to the typical chemical synthesis route (Machado et al. 2009). This has several advantages such as resulting in polymer with a defined molecular weight and, thus, a strictly monodisperse material. Moreover, the recombinant route allows the polymers to be designed accordingly for specific uses, by having a superior and easy control in the selection of its sequence, and hence, their physical-chemical properties. The recombinant polymer can be easily purified to high degrees by taking advantage of its thermal responsiveness (Meyer and Chilkoti 1999, Machado et al. 2009).

Herein, we report the production of elastin-like nanoparticles, by thermoresponsive self-assembly and by exploiting the inverse temperature transition of the biologically produced (VPAVG)220, for a sustained release of both BMP-2 and BMP-14. BMP-14 (also known as GDF-5, growth and differentiation factor) is involved in tendon, ligament, and bone regeneration (Nakamura et al. 2003, Simank et al. 2004, Dines et al. 2007), possessing significant angiogenic activity (Yamashita et al. 1997, Zeng et al. 2007, Kadomatsu et al. 2008). Thus, it makes an attractive choice as a therapeutical agent in addition to BMP-2, due to the highly vascularised nature of bone and the interest to promote angiogenesis during bone formation. BMP-14 is a key growth factor involved during fracture repair (Al-Aql et al. 2008). The elastin nanoparticles were characterized for swelling behavior, the release profile of BMP-2 and BMP-14, and biological activity of the released growth factors in C2C12 cells.
2. Materials and Methods

2.1. Materials

The elastin-like polymer, \((VPAVG)_{220}\), consists of 220 repeats of the main monomer VPAVG and was produced by recombinant technology and purified as reported elsewhere (Machado et al. 2009). The polymer has a molecular weight of 96 kDa and self-assembles at temperatures above 33 °C. More details in mass spectra (Maldi MS) verifying its amino acid composition, and self-assembling behavior, can be found in Machado et al. (2009). Human BMP-2, expressed in Chinese Hamster Ovary (CHO) cells, was purchased from Wyeth, United Kingdom. BMP-14, expressed in \(E. coli\) was purchased from PeProTech, United Kingdom. All other chemicals were of analytical grade and used as received.

2.2. Methods

2.2.1. Production of elastin-like nanoparticles

Particles were produced by resuspending lyophilized polymer in ice-cold phosphate buffer saline (PBS) to a final solution of 0.1%. The solution was then incubated at 37 °C for 30 min. The particles were then collected by centrifugation (10000 g, 37 °C, 10 min), and washed once with pre-warmed PBS. Due to its hysteresis behavior, the particles are stable at room temperature and only solubilize when a strong undercooling is achieved (of about 10 °C). The particles were prepared under sterile conditions, and stored at room temperature.

2.2.2. Physical characterization

**Scanning electron microscopy (SEM)**

Non-loaded elastin particles were morphologically characterized by scanning electron microscopy (SEM). SEM analysis was performed on gold-coated samples (Agar Sputtercoater 108, Essex, United Kingdom) and using a Philips XL20 microscope (Philips, Eindhoven, The Netherlands). Particle size measurements were obtained from several micrographs acquired in the SEM.

**Size distribution**

Measurements of particle size were performed on freshly prepared filtered samples by dynamic light scattering (DLS), using a Zetasizer Nano Series (Malvern, UK). The analysis was performed at 37 °C and by performing 10 readings.
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Water uptake and degradation studies

The hydration degree of unloaded elastin nanoparticles was evaluated after immersion into PBS, pH 7.4, for periods up to 72 hours (30 min, 4h, 24 h, 48 h and 72h). The weight loss of unloaded elastin particles was assessed after immersion into PBS, pH 7.4, for periods up to 30 days (1 d, 3 d, 7 d, 15 d and 30 d). All experiments were conducted at 37 ºC, in triplicate. Percentage of water uptake (WU) after each time period of immersion (t) was calculated using the following equation:

\[ \text{WU} (%) = \frac{W_w - W_d}{W_d} \times 100 \]

The percentage of mass remaining (MR) after each time period (t) was calculated using the following equation:

\[ \text{WL} (%) = \frac{W_d}{W_i} \times 100 \]

\(W_w\) and \(W_d\) correspond to the weight of elastin particles in wet and dry state, respectively. \(W_i\) corresponds to the initial dry weight of particles, before immersion.

2.2.3. In vitro delivery studies

Loading of BMP-2 and BMP-14

Elastin nanoparticles were loaded with BMP-2 or BMP-14 or a mix of the two, during the formation of the particles. Growth factor was added at a concentration of 20 µg/ml to the elastin-like polymer solution. The nanoparticles were produced with the different combinations of the two growth factors:

I) Unloaded particles (no growth factor)
II) encapsulated BMP-2
III) encapsulated BMP-14
IV) encapsulated BMP-2 and BMP-14

Determination of encapsulation efficiency

The nanoparticle encapsulation efficiency was determined upon their separation from the aqueous preparation medium containing the non-associated BMP by centrifugation (10000 g, 37 ºC, 10 min). The amount of free BMP-2 was determined in the supernatant by using a BMP-2 sandwich-type ELISA development kit (Eubio, Austria). The amount of free BMP-14 was estimated by dot blot using a rabbit anti-human BMP-14 antibody (BioVision, Germany) followed by an HRP-conjugated goat anti-rabbit antibody (Sigma, United States). The blot intensities were compared against BMP-14 standards of known concentration, with the use of image analysis software (Chemilimage 4400, Alpha Innotech, United States). Each sample was
assayed in triplicate \((n = 3)\). The nanoparticle encapsulation efficiency (E.E.) was calculated using the following equation:

\[
E.E.(\%) = \frac{\text{Total BMP amount (μg)} - \text{Free BMP amount (μg)}}{\text{Total BMP amount}} \times 100
\]

**In vitro release**

Pre-weighted particles (250 μg dry weight), corresponding to particles loaded with BMP-2 or BMP-14 (groups II and III), were incubated in 5 ml PBS (pH 7.4), at 37 °C, with 200 rpm agitation. At pre-determined time points (1 h, 4 h, 1 d, 2 d, 3 d, 7 d and 14 d), 100 μl of solution were removed, centrifuged (10000 g, 37 °C, 10 min) and stored at -20 °C until quantification. The solution was replaced with a similar volume of fresh buffer. Sterile conditions were applied to prevent any contamination of samples.

**Quantification of released protein**

For measurement of BMP-2 concentration, a BMP-2 sandwich-type ELISA kit (Eubio, Austria) was used, following the manufacturer’s instructions. A calibration curve was obtained using standard preparations of BMP-2 of known concentration. For detection of concentration of BMP-14, the samples were estimated by dot blot as described before. Each sample was assayed in triplicate \((n = 3)\) and mean ± SD values reported.

**Release kinetics models**

To study the release kinetics, the data obtained from the *in vitro* release was treated accordingly to zero order as cumulative amount of drug released vs. time (equation 1), first order as log cumulative percent drug remaining vs. time (equation 2), Higuchi kinetics as cumulative percent drug released vs. square root of time (equation 3) and Korsmeyer kinetics as log cumulative percent drug released vs. log time (equation 4) (Huguchi 1963, Chowdary and Ramesh 1993, Hadjiioannou *et al.* 1993, Bourne 2002):

\[
R = k_1 t \quad \text{(1)}
\]

\[
\log UR = k_2 t 2.303 \quad \text{(2)}
\]

\[
R = k_3 t 0.5 \quad \text{(3)}
\]

\[
\log R = \log k_4 + n \log t \quad \text{(4)}
\]
where R and UR are released and unreleased percentages, respectively, at time (t); $k_1$, $k_2$, $k_3$ and $k_4$ are the rate constants of zero-order, first order, Higuchi and Korsmeyer models, respectively; n is a exponent that characterizes the mechanism of release.

### 2.2.4. Bioactivity of released BMP

**MTT cell viability assay**

C2C12 cells were seeded at $5 \times 10^4$ cells/ml in a 24-well plate, attached overnight in Dulbecco’s modified Eagle’s medium (DMEM) with 1% fetal calf serum and no antibiotics, at 37 °C, 5% CO$_2$, in a humidified environment. Unloaded elastin particles were then added to the cell culture at 10 and 100 μg/ml. The MTT assay was determined after the 5 days of culture using a standard procedure (Salgado *et al.* 2002). Standard culture media, with no particles, was used as positive control of cell viability. All samples were tested in triplicate. The results are expressed as percentage of control (scored as 100% viability) as mean ± SD.

**Alkaline phosphatase (ALP) activity**

The bioactivity of the BMPs released from the elastin nanoparticles was studied by determining their ability to induce ALP enzymatic activity over basal levels in C2C12 cells (negative controls) after 5 days of culture, and by comparing the cell response to a BMP dose response effect when directly added to the culture medium (positive control). C2C12 cells were seeded at $5 \times 10^4$ cells/ml per well in a 24-well plate, attached in Dulbecco’s modified Eagle’s medium (DMEM), with 1% (v/v) fetal calf serum, at 37 °C, with 5% CO$_2$ in a humidified environment. The cells were incubated with either unloaded or loaded elastin nanoparticles, in the experimental groups described before (I to IV). Particles contained 0.1, 0.25, 0.5 and 2.5 μg of BMP-2, BMP-14 or a mix of both, in the particle preparation added to the cells. BMP-2 (Wyeth, UK) and BMP-14 (PeProTech, UK) were added as controls to the culture medium, at 0.1, 0.25, 0.5 and 2.5 μg/ml (corresponding to similar amounts of growth factor loaded in the particles). Alkaline phosphatase enzymatic activity was measured accordingly to standard procedures (Salgado *et al.* 2002) after 5 days in C2C12 cultures, and normalized to the total protein content. All samples were tested in triplicate, with two independent experiments, with reproducible results.

**Calcium mineralization assay**

For the Alizarin Red mineralization assay, C2C12 cells were seeded at $5 \times 10^4$ cells/ml per well, in a 24-well plate, and cultured in DMEM in the presence of both unloaded and loaded
BMP-2/ BMP-14 particles, in the experimental groups described before (I to IV). The particles contained 2.5 μg of single or of both of each BMPs in the particle preparation added to the cells. BMP-2 or BMP-14 were added as controls to the culture media, at 2.5 μg/ml. After 5 days of culture, the cells were washed three times with PBS (to ensure that few or no particles were retained in the cell monolayer), the culture media was replaced by DMEM supplemented with 100 μM ascorbic acid and 10 mM β-glycerophosphate, and fresh particles or BMPs were added. This was repeated after day 10. After 14 days of culture, the cells were washed three times with PBS (without Ca\(^{2+}\), Mg\(^{2+}\)) and fixed in 4% (v/v) formaldehyde, for 30 min. After removal of the fixative, cells were washed twice with distilled water and covered with Alizarin Red 1% solution, followed by gentle agitation in an orbital shaker for 10 min. The solution was then removed and the cells washed four more times with distilled water. The cells were observed using an Axiovert10 (Zeiss, Germany) optical microscope, and imaged with a coupled Coolpix950 (Nikon, Japan) camera. The mineralization was quantified by the method proposed by Gregory et al. (2004). Briefly, the calcium from cells was extracted by incubation of 800 μl acetic acid (10% v/v) to each well, for 30 min in orbital shaker. Then, the suspensions were transferred to a clean 1.5 ml eppendorf tube and vortexed for 30 sec; 500 μl of mineral oil was added and the suspensions heated at 85 °C for 10 min and then cooled in ice for 5 min. The suspensions were centrifuged (20 min, 10000 g), the supernatant was carefully removed, and 200 μl ammonium hydroxide (10%) were added (until pH ~4.1). The solutions were read for absorbance at 405 nm, in a Spectra III spectrophotometer (SLT, Austria).

2.2.5. Statistical analysis

Experiments were performed at least in triplicate and expressed as means ± standard deviation. Statistical analysis was performed using the Student t-test as well as one-way analysis of variance (ANOVA) followed by a Tukey HSD test for post-hoc comparison. Statistical significance was defined as \(p < 0.05\) for a 95% confidence. \(p < 0.01\) corresponds to a 99 % confidence, and \(p < 0.001\) to a 99.9 % confidence.

3. Results

3.1. Physical characterization

The elastin-like polymer was able to form spherical to slightly elongated nanoparticles, with average diameters of 115.5 ± 28.2 nm (Figure 1). Using dynamic light scattering, the particles showed an average size of 237.5 ± 3.0 nm, in wet state (Figure 2). Nearly all the particles (91.8 %) had sizes ranging from 190 nm to 295 nm in diameter. When incubated in
PBS, at 37 °C, the swelling of the particles occurred rapidly during the first hours of immersion, up to 261.9 ± 41.0 % after 30 min and up to 416.4 ± 37.1 % after 24h (p < 0.05), with water uptake stabilizing at 466.7 ± 33.0 % (Figure 3). The remaining mass of the particles was 97.1 ± 3.1 % after day 5, 88.9 ± 6.3 % after day 14, and 64.5 ± 12.5 % after day 30 (Figure 4).

Figure 1. SEM micrographs of unloaded elastin nanoparticles. Magnification 50.000 x. Scale is 1 μm.

Figure 2. Size distribution of unloaded elastin-like particles, obtained by Dynamic Light Scattering (DLS). Almost all particles showed sizes ranging between 190 nm and 295 nm, with a mean diameter of 237.5 ± 3.0 nm. Size refers to particle diameter (nanometers).
Figure 3. Evolution of water uptake behavior of unloaded elastin particles in phosphate buffer saline as a function of immersion time (t). (mean ± SD, n = 3). Water uptake increased significantly up to 24 hours, and then stabilized. * p < 0.05 is relative to the difference between consecutive time measurements.

Figure 4. Evolution of weight loss of unloaded elastin particles in phosphate buffer saline as a function of immersion time (t). (mean ± SD, n = 3). The remaining mass decreased to 97.1 ± 3.1 % after day 5, 88.9 ± 6.3 % after day 14, and to 64.5 ± 12.5 % after day 30.
3.2. Loading and release of BMP-2 and BMP-14

Encapsulation efficiency

The encapsulation efficiency (E.E.) of BMP-2 and BMP-14 into the elastin-like particles was respectively 94.5 ± 1.8 % and 99.2 ± 0.4 %, per total amount of growth factor added during the preparation of the particles.

In vitro release

The nanoparticles loaded with BMP-2 showed an initial burst release during the first 24 hours, of 20.0 ± 2.3 % released growth factor (Figure 5). This was followed by a gradual release, with up to 30.7 ± 1.9 % (0.8 % per day, 151 ng/day per mg polymer) of released protein after day 14. In the case of BMP-14, there was a burst release during the first 24 hours, of 19.2 ± 3.2 %. After this, there was a gradual release, with up to 36.1 ± 1.7 % (1.3 % per day, 258 ng/day) after day 14.

Figure 5. Release kinetics of BMP-2 and BMP-14 loaded in the elastin nanoparticles. ELISA was used for BMP-2 quantification and dot-blot for BMP-14 quantification (Mean ± SD, n = 3). Cumulative release is expressed in percentage of loaded protein.

Release kinetics models

The release profile fitted best the Korsmeyer model ($r^2$ of 0.97 for BMP-2 and $r^2$ of 0.96 for BMP-14) followed by Higuchi’s model ($r^2$ of 0.96 for BMP-2 and $r^2$ of 0.88 for BMP-14), while first and zero order models showed low correlations ($r^2$ less than 0.85). The release exponent
“n” of Korsmeyer model had values of n = 0.20 for BMP-2 and n = 0.14 for BMP-14 (Figure 6 and Figure 7). Since n < 0.43, in both cases, the release data fits a Fickian mode of diffusion (Ritger and Peppas 1987).

3.3. In vitro bioactivity

MTT cell viability

C2C12 incubated with unloaded elastin-like particles maintained their cell viability in a similar way to the positive control of cell viability (Figure 8). Unloaded particles have increased significantly cell viability to about 106.0 ± 0.3 % and 112.8 ± 1.2 % (p < 0.05 relative to control) when 10 or 100 μg/ml of elastin particles were added. Morphology observations corroborated the absence of cytotoxicity (data not shown).

![Graphs of release kinetics](image)

**Figure 6.** Release kinetics of encapsulated BMP-2. First order (log cumulative percent drug remaining vs. time), Higuchi kinetics (cumulative percent drug released vs. square root of time) and Korsmeyer kinetics (log cumulative percent drug released vs. log time), with correlation values (r²).
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Figure 7. Release kinetics of encapsulated BMP-14. First order (log cumulative percent drug remaining vs. time), Higuchi kinetics (cumulative percent drug released vs. square root of time) and Korsmeyer kinetics (log cumulative percent drug released vs. log time), with correlation values ($r^2$).

Figure 8. Cell viability tetrazolium salt (MTT) test performed in C2C12 cells, after 5 days of cell culture, with 10 or 100 µg/ml unloaded elastin-like nanoparticles. DMEM was used as a positive control for cell viability. Cell viability is expressed as a percentage of the control (scored as 100% viability) as mean ± SD ($n = 3$). * $p < 0.05$ is relative to differences to the control.
**ALP activity**

Cells incubated with BMP-2 loaded nanoparticles showed morphological changes into osteoblast-like shapes (data not shown) and a significant increase in ALP activity, after 5 days of culture (Figure 9). BMP-2 loaded particles induced 37-fold, 139-fold, 345-fold and 735-fold, corresponding to particles which contained 0.1, 0.25, 0.5 and 2.5 µg, respectively. The dose-wise increase in ALP activity was highly significant ($p < 0.001$). Particles loaded with the combination of BMP-14 and BMP-2, induced 40-fold, 296-fold, 369-fold and 783-fold, at the same respective amounts of loaded growth factor. Comparing particles with BMP-2 alone with BMP-2 and BMP-14, significant pairwise differences were found at specific doses (C with G and D with H, $p < 0.01$, E with I, $p < 0.05$). The overall additional effect of BMP-14 was significant ($p < 0.01$). Particles loaded with only BMP-14 induced ALP levels by 1.2-fold, 5-fold, 7-fold and 8-fold (not significant). Similar results were obtained with the control of BMP-14 added to medium. Differentiation into the osteoblast-like phenotype was also observed in these cases, although to a smaller extent as compared to BMP-2 loaded particles (data not shown).

**Figure 9.** Alkaline phosphatase activity of C2C12 cell line after 5 days of culture, incubated with loaded and unloaded elastin nanoparticles: A) negative control with no added particles or growth factor; Elastin particles were loaded with: B-E) BMP-2 (0.1, 0.25, 0.5, 2.5 µg); F-I) BMP-2 and BMP-14 (0.1, 0.25, 0.5, 2.5 µg each); N-Q) BMP-14 (0.1, 0.25, 0.5, 2.5 µg); J-M) BMP-2 added to culture medium (0.1, 0.25, 0.5, 2.5 µg) as positive control; R-U) BMP-14 added to culture medium (0.1, 0.25, 0.5, 2.5 µg) as positive control. ALP activity is reported in nmol/min/mg of total protein (Mean ± SD, $n = 3$). * $p < 0.05$ and ** $p < 0.01$ are relative to the differences indicated.
Figure 10. Alizarin Red mineralization staining of showing sites of calcium-phosphate deposits (orange color) in C2C12 cell line differentiated into osteoblast after 14 days of culture with particles loaded with BMP-2 (A), BMP-2 and BMP-14 (B), and the positive controls of BMP-2 (C) and BMP-2 + BMP-14 (D) added to culture media, at 2.5 μg/ml. Unloaded particles did not showed any mineralization (E). Unstained cells treated with particles loaded with BMP-2 + BMP-14 showed osteoblast morphology (F).
Alizarin Red mineralization

After two weeks of culture, BMP-loaded particles were able to induce osteogenic mineralization, as observed by Alizarin Red staining (Figure 10). Increased mineralization was observed in cells cultured with particles loaded with either BMP-2 or combined BMP-2 and BMP-14, than in cells cultured with corresponding doses of the growth factor added to culture media (at 2.5 μg/ml). No mineralization was observed for cells cultured with particles loaded with BMP-14 or with unloaded particles (data not shown). The amount of calcium was significantly higher \( p < 0.001 \) for particles loaded with BMP-2 \( (1463.0 \pm 11.2 \%) \) or with combined BMP-2 and BMP-14 \( (1563.5 \pm 15.0 \%) \), compared to BMP-2 added to media \( (956.2 \pm 29.3 \%) \) or to combined BMP-2 and BMP-14 added to the media \( (1053.0 \pm 3.5 \%) \). In addition, the amount of calcium was also significantly higher \( p < 0.001 \) with particles loaded with the combination BMP-2 and BMP-14 than with particles loaded only with BMP-2.

4. Discussion

Elastin-like polymers (ELPs) have been used in a wide variety of applications, ranging from protein purification to biosensors and surface-engineering, drug delivery and as biomaterials such as hydrogels (Chow et al. 2008). Recently, chemically synthesized poly(VPAVG) particles have been reported to deliver in a sustained way the model drug dexamethasone (Herrero-Vanrell et al. 2005). The self-assembly and smart behavior of ELPs make these a potential choice as a delivery system of growth factors in tissue engineering. Genetic engineering has been applied to obtain these polymers with a precise amino acid composition, sequence and length, resulting in the absolute control of its molecular weight, stereochemistry and thermoresponsive behavior (Chilkoti et al. 2006, Rodríguez-Cabello et al. 2007, Machado et al. 2009).

In this work, the self-assembly capacities of a recombinant ELP, \((VPAVG)_{220}\), has proven to be a potential way to obtain stable nanoparticle systems for delivery of BMPs. The repeat number of 220 was chosen to obtain a polypeptide considered as representative of an high molecular weight polymer (Machado et al. 2009). The particles showed average diameters of 237.5 nm, in DLS, which is of relevant interest for tissue engineering approaches, considering that the particles could be loaded into a specific scaffold or hydrogel. The size of the particles was smaller in the SEM counterpart images, mainly due to shrinkage of the particles caused by the dehydration process involved in SEM imaging, and also due to the initial high water content of the particles. The size in dynamic light scattering was significant smaller when
compared to that reported in a former work (Herrero-Vanrell et al. 2005). This is mainly due to the monodisperse nature of the (VPAVG)$_{220}$ polymer.

The particles were able to encapsulate significant amounts of BMP-2 or BMP-14, at high efficiencies. It is possible that hydrophobic interactions between the polymer and the growth factors could contribute to such efficiency. In fact, there is a significant difference in charge between the polymer and the loaded drugs; while (VPAVG)$_{220}$ has an isoelectric point of 5.5, BMP-2 and BMP-14 have isoelectric points around 9 and 8, respectively (Geiger et al. 2003, Plöger et al. 2008). Thus, this potential advantage could be applied for loading drugs that show low E.E. in other delivery systems due to low affinity to the material. Since the nanoparticles are manufactured in an aqueous-system, in mild conditions, with no harsh temperature, pH or organic solvents, this may also allow the incorporation of labile drugs; and may therefore be suitable for many therapeutical applications beyond those of bone tissue engineering by presenting an advantage due to its high E.E and easiness of preparation and handling. No cytotoxicity could be inferred by the cell viability assays, reinforcing the idea of the high biocompatibility of elastin-like polymers for clinical applications, observed before (Urry et al. 1991). The unloaded elastin particles had a small but significant stimulating effect in the cell proliferation. It is has been described that elastin peptides can induce the proliferation of diverse cell types, such as vascular smooth cells or fibroblasts, by interaction with specific cell signaling pathways (Tajima et al. 1997, Mochizuki et al. 2002).

The BMPs were successfully released over a sustained time period. The release of both BMP-2 and BMP-14 was associated with a two-phase delivery profile, consisting of an initial burst release during the first 24 h, followed by a slower and more gradual release for 14 days. During the initial stages of swelling, the elastin particles showed substantial water uptake (swelling ratio ~4) thus, the drug is rapidly released from the particle, correlating with the initial burst release observed during the first hours of immersion. The higher swelling ratio indicates the possibility that the system may be formulated also as a hydrogel-based material. It is probably that the diffusion is the only mechanism responsive for the release of the drugs in this second stage, since the release data matched Korsmeyer models of diffusion with Fickian correlation values, meaning that the release is mainly determined by the diffusion of the growth factor from the particle to medium and no degradation mechanisms. Since the temperature required for re-dissolution is well below the body or room temperature as observed previously (Herrero-Vanrell et al. 2005, Machado et al. 2009), due to the phenomena of hysteresis, the particles would remain in solid state in the body. This is also supported by the degradation data obtained. Stable particle formation is a peculiar behavior of the (VPAVG) pentapeptide. All ELPs based on the pentapeptide (VPGXG) (X representing any amino acid)
tend to form, upon heating above their $T_t$, unstable particles that rapidly sediment and coalesce to form a viscous continuous phase called coacervate. However, the polymer based on (VPAVG) is a peculiar member of the ELP family. It is obtained by substitution of the first Gly by Ala in the representing pentapeptide (VPGVG). This position for amino acid substitution is particularly sensitive. In fact this Gly can be substituted only by Ala. Any other amino acid yields a polypeptide that does not show inverse temperature transition and, therefore, is not smart and self-assembling.

In this work, we have seen that the elastin-like particles were capable of delivering, in a bioactive way, both BMP-2 and BMP-14. The sustained release profile resulted in enhanced mineralization induced by particles containing the growth factors, when compared to the corresponding doses of BMPs added to the culture media. This shows that their slow release may present a potential for triggering bone formation in vivo. Despite the effort of recent tissue engineering advances to promote the combination of a release of two growth factors to enhance bone healing (Raiche and Puleo 2004, Basmanav et al. 2008, Patel et al. 2008, Chen et al. 2009, Wang et al. 2009), it is still unclear which combination may result in a better effect and this may well depend in the type of clinical application (e.g. long bone fractures, spinal fusion or tooth regeneration). In this study, we have found an increase in the bioactivity (both ALP and osteogenic mineralization) with the combined release of BMP-2 and BMP-14 compared to both growth factors alone. This shows that the combination of these growth factors might be an interesting one. Nevertheless, further studies are required, probably by screening the induction in vivo of neo-vascularization by BMP-14, to evaluate a potential synergistic impact in bone formation.

This work provides the first report on the potential use of an elastin-like, VPAVG polymer, as a new carrier for the delivery of BMPs, offering a range of exciting properties, such as the thermoresponsive behavior, high encapsulation efficiency, and sustained release.
Chapter VIII. Thermoresponsive self-assembled elastin-based nanoparticles for delivery of BMPs

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Chapter IX. General Conclusions and Final Remarks
Chapter IX

General Conclusions and Final Remarks

This chapter intends to give a brief outlook in the main goals and results achieved in the present thesis. General conclusions are indicated, with an overall review of the steps taken throughout the research process. The main challenges encountered are drawn to attention, as well as the solutions applied. The advantages and disadvantages of the methods used are stated, as well as the reasons why specific choices were made. The main findings and contributions from this thesis are identified, relating these to the context of the tissue engineering, clinical applications and the scientific community. Finally, a few recommendations are made on possible further studies to overcome specific challenges and future directions to broaden the horizons of the current field of investigation.

GENERAL CONCLUSIONS

The main goals of the present thesis were to clone, express, purify and characterize human recombinant BMPs, in primary cultures of human stem cells and in C2C12 cells, and investigate novel delivery systems for the sustained delivery of these BMPs.

The overall work was divided into the following parts:

- Cloning, expression, folding, purification and bioactivity characterization of different human recombinant BMPs (BMP-2, BMP-4, BMP-9, BMP-10, BMP-11 and BMP-14);
- Production of silk fibroin microparticles and elastin-like polypeptide thermoresponsive self-assembled nanoparticles for delivery of different human BMPs with physical characterization and in vitro or in vivo bioactivity characterization.
1. A METHOD FOR THE PRODUCTION OF RECOMBINANT BMPs

The production of growth factors by recombinant technology was originally proposed in order to meet the requirements of tissue engineering field to obtain large amounts of these molecules for research purposes. Bone morphogenetic proteins were specifically selected since these are the main signaling growth factors that promote the differentiation of progenitor cells to bone-forming cells, as described in section 1, chapter I.

Bone morphogenetic proteins were produced by recombinant technology in *Escherichia coli*, in an attempt to overcome the challenges of former methods, such as the low yields of native BMP extraction from bone, or from mammalian expression cells (as stated in Chapter I and IV).

This process involves a stepwise approach on which different issues had to be overcome:

- **Source of genetic material for cloning of human BMPs.** As detailed in chapter IV, the DNA was obtained from the Human Genome Project, due to the mature domain of these BMPs being located in only one exon. However, this method is not be applicable to every gene, which in those cases, cloning from a cDNA library would have to be used.

- **Finding an ideal optimal plasmid as expression vector.** The innovative approach is based on the pET25 vector, using *Escherichia coli* as a host for recombinant expression. This had main advantages: i) the fact that this vector facilitated the secretion of recombinant protein to the periplasm, promoting thus the formation of BMPs dimers, ii) allowing easy overexpression of the protein, and iii) allowing easy purification by affinity chromatography due to its integrated six-histidine-tag.

- **Optimization of expression conditions.** These steps were performed as detailed in chapter III. The overexpression was then scaled-up in a bioreactor, to achieve very large amounts of BMPs, after several fermentation variables were optimized (codon usage, temperature, expression period, induction method).

- **Isolation of the recombinant growth factors and proper folding.** Since initially the protein was detected in insoluble fraction, the protein was easily solubilized in a buffer containing L-arginine, which acts as a suppressor of protein aggregation and a facilitator of folding, i.e. a chaotrophic agent. The BMPs were present in monomer, dimer and polymer forms. Purification by affinity chromatography resulted in isolation of the BMPs, to purities up to 90-95%. The yields achieved were 25 to 45 % from total, or 23 to 54 mg
per Liter of fermentation batch, or 110 mg for BMP-2 in bioreactor (the only growth factor which had a scale-up).

- Stability and bioactivity of protein. BMP-2 was stable in physiological conditions such as phosphate saline buffer, pH 7.4. The purified BMPs were able to induce several markers of osteogenic differentiations and did not possess significant cell cytotoxicity, as observed by the cell viability assays. C2C12 cells were used as a standard and reproducible model for testing the \textit{in vitro} bioactivity of the recombinant BMPs, while human fat-derived stem cells provides a human-based model to test the biological activities of the different BMPs.

Several main advantages were found with the overall methodology:

- The method developed is an easy way to obtain large amounts of recombinant BMPs of significantly purity, by a very straightforward, low-cost, and easy protocols.
- The proteins are stable in physiological conditions and easily stored.
- Recombinant BMPs are able to induce several markers of osteogenic differentiation.
- The expression and purification method can be easily adaptable to other growth factors.

And a few drawbacks:

- Bioactivity is lower when compared to commercial BMPs, produced in mammalian cells, requiring thus trial-and-error and time-consuming experimentation for the optimization of the protein folding and activity. Future studies are required to optimize the bioactivity to the levels obtained with BMPs expressed in mammalian cells.
- The protein is obtained in monomer, dimer and polymer forms, so further isolation of dimers is required.
- The protein, expressed in \textit{E. coli}, does not have its natural glycosylation pattern, hence its bioactivity and stability may be reduced, compared to mammalian-derived BMPs.
- Further purification from residual endotoxins is also required.
As investigated in Chapter V, different recombinant BMPs were produced. This can provide a number of important contributions:

- To show that the novel strategy can be applied to several growth factors.
- Since different BMPs participate in a timely sequence of growth factors involved in the differentiation of osteoblast and new bone formation, these may be required for novel therapeutical applications, involving a combination of BMPs. In fact it is still poorly understood the entire sequential activation and physiological roles of the different BMPs in different clinical situations, such as spinal fusion, fracture healing or dental healing.

2. INNOVATIVE MICROPARTICULATE DELIVERY SYSTEMS FOR BMPs

The choice for the carrier was based on the assumption that materials for tissue engineering should ideally mimic the natural environment of tissues. We have chosen silk fibroin (chapter VI and VII) and elastin-like polymers (chapter VIII) due to a series of potential advantages for bone tissue engineering, which were demonstrated in this thesis:

- The ability to form nano- or microparticles, with a rather homogeneous distribution of size, by a very easy methodology.
- Both systems have very easy method of drug loading, with no use of harsh conditions, allowing the loading of more labile molecules
- Both systems have high efficiency of loading, applicable for the different BMPs
- Both systems did not show significant cytotoxicity as inferred by in vitro viability assays.
- Both systems showed the ability to deliver the BMPs, retaining their bioactivity in vitro. Fibroin particles loaded with BMP-2 have demonstrated to be able to trigger new bone formation in vivo, as demonstrated in the ectopic bone formation model, in chapter VII.
- Both particulate systems showed relatively small burst release, followed by gradual and sustained release of the BMPs for up to 14 days.
- Both systems possess slow degradability, having small weight loss over the short-term, which make these materials very suitable for bone regeneration. Silks have interesting mechanical properties, since by having slow degradation times, may allow adequate time for proper bone remodeling.
Elastin-like polypeptides presented further specific advantages:

- The thermoresponsive and hysteresis behavior allows the preparation of multi-loading particles for the release of combinations of different growth factors, with a very easy and fast method. The optimization of a system for a potential sequential release of growth factors requires further studies.

A few drawbacks may be noted for these particulate systems, namely:

- Particles alone disperse in vivo, and thus require a scaffold material as carrier.
- The significant swelling in both particle systems may be a disadvantage in vivo, with risk of increased inflammatory in the implantation site. This can be avoided by pre-soaking the carrier prior to implantation in PBS, or by incorporation into a scaffold material.

The rat ectopic bone formation model (Chapter VII) also proved an efficient model for accessing the activities of recombinant BMPs delivered via a carrier in vivo. The microCT and histological analysis provide two powerful techniques to analyze the formation of bone both on a histochemical level, by screening the expression of important mineralization proteins such as osteocalcin, but also a three-dimensional assessment of the quality of bone formed, its density and length, and how the doses of the BMP affect these parameters. However, one main drawback is that the ectopic model only provides an idea whether the carrier is delivering the BMP in a sufficiently bioactive and sustained way to trigger bone formation. Further in vivo orthotopic studies, from a clinical perspective, are required, using, for example, a long bone fracture model.
FUTURE DIRECTIONS

In this section, recommendations are made for further studies to improve the methodologies previously described. These include:

- Improvement of refolding methodologies to obtain recombinant BMPs, with increased bioactivity to levels similar to the ones induced by BMPs expressed in mammalian cells.
- Isolation of dimeric BMPs by using size exclusion chromatography or heparin affinity chromatography. Endotoxins should be also removed prior to clinical assays.
- Alternative expression systems may be explored, for instance in yeast, thereby avoiding the issues of protein misfolding due to overexpression.
- Novel delivery systems based in a combination of micro- or nanoparticles and a scaffold or hydrogel, for a sequential delivery of two or more growth factors. The scaffold would provide as carrier for the particles, while supporting also cells. Sequential delivery would be achieved by one growth factor being released first from this carrier, and a second growth factor released from the particles.

We also highlight other directions of future work that might be fruitful and interesting to exploit:

- Formation of heterodimeric recombinant BMPs, possibly yielding stronger bioactivity;
- Fusion proteins, between growth factors or BMPs and domains of interest, such as domains of binding to biomaterials, to possibly increase bioactivity. One example would be the cloning and expression of fusion proteins of BMPs to silk fibroin and elastin sequences, to achieve, in theory, a polypeptide that would simultaneous work as a carrier and as a growth factor, and exhibiting thermoresponsive behavior.
- Biomimetic coatings in the particulate systems since these were previously reported to increase the binding and retention of BMPs, and therefore increase their bioavailability.
A more attentive look at folding is required, as the folding structure of the recombinant BMPs is critical to its bioactivity. It is known, and detailed in chapter II, that BMPs bind to combinations of cell surface receptors, and these combinations can explain different bioactive effects. Mutations in one epitope affect ALP activity, and in another the triggering of Smads pathway. Therefore, the critical correct folding and subsequent bioactivity of recombinant BMPs has to be studied both with assays of the induction of ALP, induction of Smads pathway and transcription factors specific to osteogenic pathway, and the induction of mineralization proteins. Correct refolding of recombinant BMPs is achieved by finding the optimal conditions for a renaturation of monomer to dimer (e.g. protein concentration, ionic strength and buffer composition, the presence of chaotrophic agents and aggregation inhibitors (e.g. L-arginine, urea), temperature and duration of reaction).
Attach I

Supplementary data

This section intends to give additional methodologies and results obtained during the current thesis, which are of not of critical importance to the overall conclusions, but still of relevance.

REFOLDING STUDIES WITH RECOMBINANT BMPs

When the bioactivity of recombinant BMP-2 was compared to mammalian-derived BMP-2 (Wyeth, UK), the activity was significantly lower, with no induction of alkaline phosphatase (ALP) activity and no osteogenic mineralization in C2C12 cells. For this reason, we aimed at optimizing the refolding steps of the recombinant BMP, in order to achieve the full bioactivity. Below are briefly described these procedures, after extensive trial-and-error, which nevertheless failed in resulting in reproducible batches of fully bioactive recombinant BMPs.

A stepwise optimization of BMP refolding

The new method involves a stepwise methodology:

1. **Lyses of bacteria.** Briefly, cells are lysed by incubation with PBS supplemented with 1 mg/ml lysozyme at 37 °C for 20 min, followed by extensive ultrasound treatment in ice. DNAse is added at a concentration of 20 mg/ml to promote the removal of nucleic acids and reduce the viscosity of the lysate. The lysate is resuspended at a concentration of 200 mg wet weight per ml.

2. **Wash and isolation of inclusion bodies.** The lysate is centrifuged and the pellet, where most BMP was found (chapter IV), is subsequently washed for several times with Tris buffer 0.1 M, 1 mM EDTA, and 2 % Triton, pH 8.0, to ensure the isolation of the inclusion bodies to high degree. Ultrasound treatment is used for the resuspending of inclusion bodies. These steps are performed at 4 °C. The purification of inclusion bodies is important as the impurities in the lysate may trigger the precipitation of protein during the refolding.

3. **Solubilization of inclusion bodies.** The purified inclusion bodies are solubilized using 4 M guanidine-HCl, 0.1 M Tris, pH 8.5, 2 mM EDTA and 50 mM TCEP or DTT (reducing agent), to a concentration of 5 to 15 mg/ml. This step is performed at room
temperature for 1h, or overnight, at 4 °C. The solubilized protein is centrifuged and the precipitate is discarded. Then, it is necessary to remove the reducing agent prior to refolding, by buffer exchange, since it can interfere with the oxidative folding pathway of the protein. The solution is buffer exchanged to an identical buffer, but with no reducing agent, and with the pH adjusted to 5.5, to prevent the formation of cysteine bridges. Solubilized inclusion bodies may be stored at -80 °C.

4. Refolding of inclusion bodies. The inclusion bodies are refolded by rapid dilution into a specific buffer, with gentle mixing. A dilution into a final protein concentration of 50 to 100 μg/ml is recommended. The refolding buffer consists in Tris buffer 0.1 M, 1 M NaCl, 2 mM EDTA, 0.5 M L-arginine, 0.3 M guanidine-HCl, 5 mM oxidized glutathione and 2 mM reduced glutathione, pH 8.5, but a combination of different concentrations of each component may be experimented. Refolding is performed initially on ice, for 30 min., to avoid protein precipitation, followed by incubation for a specific period of time, between 1h and 10 days, but mainly for 3 days. Prior to preparation, the refolding buffer is degassed and nitrogen-purged to ensure a buffer environment that protects the protein from amine alteration. Alternatively, the refolding is performed by stepwise dilution, by one-step dialysis or stepwise dialysis, involving a gradient of decreasing concentrations of guanidine-HCl, to test the influence of the removal rate of the protein denaturant from the solution. For a concentration of 0.4 M guanidine-HCl, a maximum of 200 μg/ml protein concentration is recommended; for 0.1 M guanidine-HCl, 50 μg/ml (as observed experimentally). Another option, involves the addition of pulses of denatured inclusion bodies into the refolding buffer, by slowly increasing the protein concentration over time.

5. Purification of BMP dimer by heparin affinity chromatography. The refolded protein is then dialysed against Tris 0.1 M, 4 M urea, pH 8.5 and applied to a pre-equilibrated heparin affinity chromatography column. The column is washed with Tris-urea buffer, with 0.3 M NaCl to remove the monomer. The dimer is eluted with Tris-urea buffer, with 0.8 M NaCl. The urea is used to avoid the protein precipitation in the column. The protein is temporarily unfolded during this step, refolding again after the removal of the urea. This purification step is optional as it does not interfere with the protein activity (as observed experimentally – data not shown).

6. Buffer exchange to physiological buffer and storage. The purified protein is then dialysed against 50 mM MES buffer, pH 5.0, to a maximum concentration of 50 μg/ml, and stored at -80 °C until bioactivity tests, or at 4°C for short-term use.
Bioactivity of BMPs refolded by the new method

The bioactivity of recombinant human BMPs, described in chapters IV and V, was much lower than commercial mammalian derived BMP-2 (Wyeth, UK) due its lack of induction of high levels of ALP activity after 5 days of culture, and bone mineralization after 21 days of culture in C2C12 cells (data not shown), despite the fact that it elevated significantly the expression of mRNA of several osteogenic markers in both C2C12 cells and human adipose derived stem cells.

For this reason, new variants of BMP-2 were cloned and expressed with N-terminal histidine tagged BMP-2, C-terminal histidine tagged BMP-2, non tagged BMP-2 and non-tagged BMP-2 with no heparin binding domain, which is postulated to reduce the BMP in vitro activity. These proteins were successfully overexpressed using pET-25 vector E. coli (data not shown) but still not showing full bioactivity, i.e. ability to induce high levels of ALP in C2C12 cells. After successive experiments, fully bioactive protein was obtained in some refolding batches, but not in a reproducible way, despite the fact of the controlled experimental conditions. In these cases, BMP-2 and BMP-4 were able to trigger high levels of ALP enzymatic activity, in C2C12 cells, after 5 days of culture (Figure 1), with clear differentiation into osteoblast morphology, similar to that observed with the positive control of BMP-2 (Figure 2). The saturation levels of ALP activity were achieved at approximately 1000 ng/ml of BMP-4, 500 ng/ml of refolded BMP-2 and 250 ng/ml of positive BMP-2 control. Alizarin Red staining revealed that the refolded BMP-2 and BMP-4 could successfully induce calcium-phosphate bone-like mineralization after two weeks of cell culture (Figure 3). Mineralization was achieved using both 200 and 1000 ng/ml of BMP-2 and only with 1000 ng/ml of BMP-4.
Figure 1. ALP activity in C2C12 cells after 5 days of culture with refolded BMP-2 (blue), BMP-4 (green) and positive BMP-2 control (red), at 25 to 1000 ng/ml. (mean ± S.D., n=3, * p < 0.05 relative to negative control)

Figure 2. Morphology of C2C12 cells after 4 days of culture with refolded BMP-2 and BMP-4, positive BMP-2 control (mammalian-derived, Wyeth, UK), at 1000 ng/ml, and negative control (no growth factor). Osteoblast-like morphology is observed in all conditions except in negative control.
Figure 3. Alizarin Red staining of C2C12 cells treated with refolded BMP-2 and BMP-4, at 1000 ng/ml, in comparison with the positive control (mammalian-derived, Wyeth, UK), at 1000 ng/ml, and negative control (no growth factor), after 15 days of culture. Mineralized extracellular matrix is stained in cells treated with refolded BMP-2, BMP-4 and positive BMP-2 control.

Stability of bioactive BMP-2

The refolded BMP-2 showing activity was also studied for its stability in different conditions, such as pH and ionic strength, when incubated overnight. The protein retains the activity at pH between 4.5 and 9.0 (Figure 4), and at ionic strength above 10 mM phosphate buffer (Figure 5). This shows that physiological buffers, or slightly acidic conditions, are preferred for storage of the recombinant protein, which is of interest for tissue engineering and biomedical applications.
Figure 4. ALP activity in C2C12 cells after 5 days of culture of refolded BMP-2 stored at different pH conditions, added at 150 ng/ml, and a negative control with no growth factor (C). Loss of ALP activity is observed at pH 3.0 and pH 11.0. Activity is retained between pH 4.5 and pH 9.0.

Figure 5. ALP activity in C2C12 cells after 5 days of culture of refolded BMP-2 stored at different ionic strength buffer conditions (phosphate buffer, pH 7.4), added at 150 ng/ml, and a negative control with no growth factor (C). Loss of ALP activity is observed at buffer conditions below 10 mM phosphate. Activity is retained in 10 mM to 1 M phosphate buffer.
Correct refolding versus misfolding

The protein is initially in the form of unfolded and reduced inclusion bodies, in the presence of a high concentration of chaotrophic salt (guanidine-HCl), before the refolding (Figure 6A). Then, as the concentration of the denaturant decreases during the refolding steps (e.g. dialysis or dilution), and oxidizing conditions are present, the protein gradually folds into its native structure, and forms dimers by cysteine bridge arrangements (Vallejo and Rinas 2004a). It was observed, by Western-blot, that the renaturation typically resulted in about 20 to 30% of dimer formation (Figure 6B), as analysed by Gel Pro Analyzer image software (Media Cybernetics). However, as explained before, in several refolding batches, this did not correspond to bioactivity, when the protein was tested in the cells, even when the dimeric form was isolated by heparin affinity chromatography (Figure 6C) or by size exclusion chromatography (data not shown).

As shown in Figure 7, the correct protein folding pathway (1) often competes with misfolding (2) and aggregation (3), depending in the several variables of the reaction mix (listed in next page). Disulfide-bond knot proteins, such as BMPs, can easily form misfolded products, due to the complicate nature of their intra- and intermolecular disulfide bonds (Arolas et al. 2006). The misfolded dimers would have the correct molecular weight but no bioactivity. If the percentage of correct refolding is low, below a certain threshold, no activity will be detected in the cell cultures; whether if the percentage of refolding is slightly increased, then bioactivity would be observed.
Figure 7. Simplified model of correct folding versus misfolding and aggregation. The correct protein folding pathway (1) often competes with misfolding (2) and aggregation (3), depending in several variables. Disulfide knot proteins, such as BMPs, can easily form misfolded products, due to the complicate nature of intra- and intermolecular disulfide bridges. These misfolded dimers would have the correct molecular weight but no bioactivity.

Because the misfolded protein would appear with the same molecular weight, but not showing activity, it is therefore important to state that bioactivity assays are required to screen for the correct refolding of the protein and its full biological activity.

Different variables are known to affect refolding

Several factors are known to affect protein refolding (Rudolph and Lilie 1996, Singh and Panda 2005, Tsumoto et al. 2003, Vallejo and Rinas 2004a). The protein concentration, the removal rate of denaturant and the temperature are three key factors. pH is another critical factor, as it is the redox environment and the buffer composition (e.g. the addition of additives that prevent protein aggregation or enhance folding).

Thus, the following variables are known to affect protein refolding and were investigated:

- **Removal rate of denaturant.** Because of the competing processes of misfolding and aggregation, a key to refolding is in the intermediate concentration of the denaturant, where denaturant concentration is low enough to force protein molecules to collapse.
into folded configuration, yet can allow them to stay in solution and be flexible to reorganize their structures. Such balance can be manipulated by two factors. One is the rate by which denaturant concentration is reduced, which depends on the method used (rapid dilution, one-step dialysis, stepwise dialysis or stepwise dilution). The other is to add additives that reduce protein aggregation or enhance folding. Another option that was evaluated was the slowly addition of pulses of denatured protein into the refolding buffer. Since completely folded proteins are usually not prone to precipitation, higher yields are theoretically possible, at higher protein concentrations.

- **Protein concentration.** In general, for the refolding of BMPs, concentrations between 30 μg/ml and 200 μg/ml. Higher concentrations, up to 500 μg/ml, were tested, but precipitation of the protein was frequently observed (data not shown). Since refolding is a first order reaction, and the competing misfolding and aggregation reactions are second-order processes, refolding is favoured by working at low protein concentration. One drawback of working with low concentrations, is that it may expose the protein to degradation or surface binding, resulting in lower recovery, and requires time-consuming concentrating steps, after refolding.

- **Temperature.** While increased temperature accelerates the rate of refolding, it also increases greatly the likelihood of protein precipitation and misfolding. Therefore, as a rule, refolding was mainly performed at 4 °C. Whereas, other temperatures were evaluated, e.g. 10 °C or room temperature, we have found that the incubation of protein on ice on the first stages of refolding (first 1-3 hours) was enough to avoid protein precipitation.

- **pH.** The formation of cysteine bridges in the BMPs is reported to occur mostly at a very narrow pH range, between 8.0 and 9.0, optimally at pH 8.5 (Vallejo and Rinas 2004b). In the context of this thesis, refolding experiments were conducted mainly at this pH, but a range of pH 5.0 to 9.0 was also tested. After refolding, buffer should be exchanged to acidic conditions, such as 50 mM MES, pH 5.0, since the solubility of the protein is best.

- **Buffer composition (e.g. ionic strength and presence of additives).** The addition of additives that either reduce aggregation, enhance folding or protein interaction, is also recommended. Particularly, L-arginine is generally recommended (Tsumoto et al. 2004) as it suppresses protein aggregation during the early stages of refolding, therefore aiding in refolding. Routinely, concentrations of 0.5 to 1 M were used. Other additives include MgCl₂, glycine, ammonium buffers, sucrose, glycerol, PEG, CHES and EDTA.
General salt conditions include 0.1M Tris and 1 M NaCl, but other conditions were also studied.

- **Redox conditions.** The formation of intra and intermolecular cysteine bridges, required for correct folding and the formation of dimers, is achieved by adding a mix of oxidized and reduced thiol reagents such as glutathione, in both oxidized and reduced forms. In general, because thiol-disulfide exchange reactions are reversible, these reagents aid in increasing both the rate and the yield of correct disulfide bonds, by rapid reshuffling of improper disulfide bonds, that lead to wrong folding (Rudolph and Lilie, 1996). In the context of this thesis, we have experimented with different ratios of glutathione forms.

- **Preparation of inclusion bodies prior to refolding.** Other factors that can affect the efficiency of refolding are the purification degree of inclusion bodies prior to folding, the expression conditions during fermentation (lower rates of expression can correspond to increased yields of soluble and native protein), and the conditions used during inclusion body solubilization (milder solubilization, with lower concentrations of denaturant, may result in increased efficiency of refolding).

Future work will involve the optimization of the refolding conditions necessary to achieve the reproducibility of fully bioactive protein, under controlled conditions.
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