PRGF (Plasma Rich in Growth Factors)

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PRGF (Plasma Rich in Growth Factors) is a system for obtaining platelet and plasma proteins: autologous proteins are obtained from the patient’s own blood shortly before its therapeutic use. Its application accelerates the repair/regeneration mechanisms of various tissues.

This article describes specific characteristics of PRGF, differentiating it from other systems and techniques available on the market. Included are descriptive preclinical studies on the content of the following growth factors: PDGF (Platelet-Derived Growth Factor), TGFβ-1 (Transforming Growth Factor), EGF (Epidermal Growth Factor), VEGF (Vascular Endothelial Growth Factor), IGF-1 (Insulin-like Growth Factor Type I) and HGF (Hepatocyte Growth Factor). The relationship between the number of platelets and the concentration of these factors is analyzed.

The complexity of interactions between proteins and their interactions with the different cell types prevents establishing a relationship between dose and clinical effect. The information derived from our research shows that an effective and sufficient concentration does not imply large doses. As examples, we show the effect that treatment with PRGF has on cellular proliferation when the donors have a different number of platelets and different concentrations of growth factors. Also, two clinical cases illustrate that the efficacy of these preparations does not strictly depend on the number of platelets.

Key words: Growth Factors, Platelets, PRGF, Tissue Repair.

Introduction

Many times the objective of medical research is not so much to prolong life as to improve its quality. Similarly, in the field of implantology, research has led to the development of new geometric designs for implants, different surface treatments, new diagnostic imaging techniques, computerized systems to simulate the placement of implants and systems for obtaining and using autologous growth factors. All these factors have contributed to the achievement of very satisfactory clinical results. Osseointegration time has been reduced, and primary stability improved. Moreover, implant treatment has extended to more complex clinical situations, and currently different surgical strategies and techniques can...
be applied to allow the restoration of situations where the placement of implants could not be considered until now. This has improved the quality of life for many patients.

The situation is similar in the field of traumatology: articular prostheses have allowed patients affected by joint disease not to have to spend the rest of their lives in wheelchairs. But we should always keep in mind that the success of a treatment will be multifactorial and involves such things as surgical technique, implant or prosthesis design, imaging diagnostics, and the use of autologous biological techniques, such as PRGF, designed to accelerate tissue repair and regeneration process [1,2,3,4]. This involves using the patient's own proteins for regenerative purposes.

The use of PRGF is not limited to oral surgery, but has quickly spread to other medical specialties [5,6]. The reason for this is that the new treatments are based on the use of proteins: these are the nucleus of human physiology and regulate the various mechanisms involved in the repair of distinct tissues. PRGF can be widely applied because its mechanism of action affects basic molecular and cellular processes, generally common to all tissues.

**Complexity of the System**

The human body has some 100 trillion cells that govern themselves through an exchange of chemical signals. The cells secrete these chemical signals that influence the behavior of other cells and, in turn, receive external signals through specific receptors in their cellular membrane. These are complex systems, difficult to understand completely due to the multiple interactions between the system components. These dynamic interactions are regulated by physical and chemical laws and serve to tell the cell what its immediate function is and if its situation is the correct one. All this information communicates to the cell where the cell is at any given moment, which cells surround it, and the next thing it should do. Currently, a great deal of research is directed towards improving understanding of the different processes and, in the case of regenerative medicine, to decipher the mechanisms of action involved in repair and regeneration.

This is not a matter of invention, but of interpreting and using the information derived from research to achieve quicker and more efficient tissue regeneration. Until now, a large part of research has centered on in-depth study of the individual signaling proteins. But the challenge is to know the system as a whole, i.e., the interactions of these proteins among themselves and with the distinct cell types.

Two very clear facts must be kept in mind.
1. First, repair is the result of the interaction of different cell types with multiple signalling proteins.
2. Second, the biological condition of a tissue in repair/regeneration phase varies depending on the length of time since the injury and the distinct topography of the tissue.

**Platelets and Tissue Repair**

Platelets, most often studied for their role in hemostasis, have a very important physiological function which has recently been discovered and validated: they are protein carriers with an important role in tissue repair and regeneration.

Platelets are small discoid cellular elements, heterogeneous in size and density; they are cytoplasmic fragments of the megakaryocyte, a giant cell in the medullary bone; they circulate through the blood stream for around 8-10 days.

In addition to being involved in hemostasis for their pro-coagulant function, they contain various growth factors involved in the repair of different tissues. They act as a carrier for these growth factors and release them in the areas where there is tissue damage.

GFs (Growth Factors) are stored inside, in special secreting granules, the α granules. Among others, some which are stored are: PDGF, TGFβ-1, EGF and VEGF. These substances were synthesized by the megakaryocyte, since the platelet does not contain a nucleus or the necessary elements for protein synthesis.

On the other hand, during its journey through the circulatory system, the platelet captures plasma proteins that it also stores inside its granules. It therefore, contains a very complex mixture of proteins [7], some from its precursor cell, the megakaryocyte, and others captured by endocytosis from the blood stream.

Platelet concentrates have become popular in the clinical environment as a tool that, along with surgery, allows quicker and more efficient tissue regeneration. The first publications on the use of autologous platelet proteins appeared in the late 90s in the area of oral and maxillofacial surgery [8,9,10,11,12]. In the years following, different systems for obtaining
and preparing platelet concentrates for therapeutic purposes have become available on the market.

Preparation protocols vary from system to system, along with the concentrations of the different integrating proteins. PRGF (Plasma Rich in Growth Factors) is a system for preparing platelets and plasma proteins; it has particular characteristics that differentiate it from other systems available on the market. The objective of this article is to describe PRGF and its principal characteristics.

**Characteristics of PRGF**

PRGF is a mixture of autologous proteins, prepared from a determined volume of platelet rich plasma (PRP). PRP is a term coined by hematologists to describe a plasma rich in platelets. According to hematologists' criteria, PRP is plasma that contains more than 300-350,000 platelets/uL. PRGF is the only technique described that prepares platelet-enriched plasma that does not contain leukocytes. PRGF always and exclusively uses the patient's own autologous proteins, prepared at the same time they are used. PRGF contains platelet and plasma growth factors involved in the repair process; it also contains sticky plasma proteins such as fibrin, fibronectin and vitronectin, among others.

**Preparation:**

PRGF is prepared from a small volume of blood that is adapted to each specific clinical case. It can vary from 5 - 80 cm³.

To prepare the PRGF, blood is taken from a peripheral vein using sodium citrate as an anticoagulant. Other anticoagulants induce changes in the platelet morphology, when the blood is collected into EDTA, the platelets swell and become spherical instead of discoid. Another commonly used anticoagulant, ACD, has a lower pH (6.5) and interferes with platelet aggregation. The traditional preparation of PRP consists of a slow centrifugation, which allows the platelets to remain suspended in the plasma while the leukocytes and erythrocytes are displaced to the bottom of the tube. A rapid centrifugation can cause mechanical forces and can raise the temperature which can induce changes in the ultrastructure and form of the platelet which, in turn, can initiate a partial activation, causing it to lose part of its content.

The current systems for preparing platelet concentrations use various centrifuges, at different speeds. The final objective is to obtain a platelet pellet, but this precipitation is not selective and the precipitate itself includes all the leukocytes corresponding to the initial volume of blood.

The PRGF is prepared with a single stage centrifugation. Using the parameters of time and speed established in the protocol, the platelets are concentrated in the cm³ of plasma situated immediately above the red cells. Also, the leukocytes end up in a single layer (buffy coat) immediately on top of the erythrocytes; this allows collection of the PRGF

**Confocal microscopy of the PRGF clot**

**Fig. 1a:** shows the layout of the fibrin fibers marked with green fluorescence.
**Fig. 1b:** shows the layout of the platelets marked in red.
**Fig. 2:** Global image of PRGF clot in which the fibrin and platelets regroup.
**Therapeutic dose and effect:**
There is a belief that in order to obtain a therapeutic PRP, the platelets must be concentrated to a maximum. Although some authors speculate that the minimum concentration of platelets to achieve a clinical effect should be one million platelets per microliter, there are no conclusive experimental results that support this hypothesis [14].

Although this criterion is assumed many times by clinicians with no questions as to its reality, we believe that it is at least a subject open to debate.

When PRGF or another platelet concentrate is used, a multiple combination of proteins is applied [7] with varying interactions and effects on different cellular mechanisms. This seems to be an over-simplification for such a complex situation to believe that the more platelets the better and the greater the clinical result, or that these preparations are effective only above one million platelets per microliter. Based on our experiments, however, we believe that an effective and sufficient concentration does not involve large doses.

In an injury site, there are different types of cells in different situations - some will be irreversibly damaged and will suffer a process of apoptosis; still other cells will be less damaged and will react to stimuli from the nearest microenvironment to maintain tissue homeostasis, i.e., the conditions prior to the injury.

Cells from the nearby tissues will begin a displacement in response to specific signals (chemotaxis) and inflammatory cells infiltrating from the blood stream and precursor cells will arrive.

This is only a glimpse of the complexity in which the damaged tissue finds itself in the moments following injury.

**Preclinical studies:**
In preclinical studies carried out by our research department, concentrations of growth factors in PRGF have been studied along with their relationship with the number of platelets. Plasma from 55 healthy individuals was analyzed: 22 females and 33 males between the ages of 18 and 60 years.

![Fig. 4a](image1.png)  
**Number of platelets in peripheral blood and in the three fractions of plasma**

![Fig. 4b](image2.png)  
**Concentrations of PDGF-A, TGF beta 1 and IGF-I in the three fractions of plasma**

![Fig. 4c](image3.png)  
**Concentrations of EGF, VEGF and HGF in the three fractions of plasma**

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**Fig. 4 (a, b, c): Quantitative description (mean values ± standard error) of the different fractions of plasma.**
Biological variability in the number of platelets and in the concentration of growth factors:
Peripheral blood contains 150-400 million platelets per ml, therefore, the range is very broad. PRGF in any given individual concentrates platelets around 3 times the number in the peripheral blood. The platelet population is not homogenous and there is a size distribution of a mean diameter of around 2 μm; the volume and area also present a distribution.

Just as in all biological parameters, there is also a biological variability in the concentration of GFs that are inside the platelets of each individual and there is also variability in the concentration of the plasma growth factors.

Platelet growth factors:
Platelet-derived growth factor (PDGF-AB), transforming growth factor (TGFβ-1), epidermal growth factor (EGF) and vascular endothelial growth factor (VEGF) are platelet secreted growth factors, are synthesized by the megakaryocyte and stored in the α granule of the platelets.

Figures 5a, 5b, 5c and 5d show the number of platelets beside the concentration of these growth factors.

In the case of VEGF (Fig. 5d) the relationship between the number of platelets and the concentration is weak, meaning that there are individuals who have a high concentration of VEGF and others who have a very low concentration. The concentration of VEGF is a characteristic of each individual.

However, the concentrations of PDGF-AB, TGFβ-1 and EGF (Figs. 5a-c) are very much interrelated and also related to the number of platelets. This relationship is statistically significant to 99% certainty. A mathematical model exists that explains an 82% of the variability. In practical terms, this relationship means that for any individual a platelet concentrate will always contain a larger amount of TGFβ-1 than PDGF-AB and, in turn, this will be more concentrated than the EGF.

In short, the relationship between the concentrations of these factors is always similar for different persons; there will always be 2.5-3 times more TGFβ-1 than PDGF.
This means that whatever number of platelets the preparation contains, there will always be 2.5-3 times more TGFβ-1 than PDGF-AB. This fact is very important and we will consider it in the following discussion on platelet dose and therapeutic effect.

**Plasma growth factors:**

As seen in figures 5e and 5f, the concentrations of IGF-I and HGF have no relation to the number of platelets. The concentration of these factors inside the platelets is very small compared to the plasma concentration.

This leads us to the conclusion that the relationship between the number of platelets and concentration for these GFs is very weak, since the contribution of platelets to the total concentration is very small.

**What are the repercussions of this variability in the clinical result?**

An effective and sufficient concentration does not imply large doses.

Based on the belief that if a little is good then more is better, others seek to concentrate platelets up to 8 - 10 times (basal reference is always peripheral blood).

Technically, this is very easy to achieve; all we need to do is to precipitate the platelets and resuspend the pellet in a small volume of platelet poor plasma.

This can be done easily, so it is not a technical problem that leads us to favor a moderate platelet concentration more akin to human physiology.

To measure the efficacy of something, an effect must be measured. As to the effect, which one do we measure? The GFs that concern us influence proliferation, inflammation, chemotaxis, differentiation, etc. They also deal with a complex mixture whose components show multiple interactions conditioned by relative concentrations and biological surroundings.

An example will better illustrate the situation: PDGF is a factor that induces proliferation in mesenchymal type cells. TGFβ-1, on the other hand, is an inhibitor of the proliferation of these same types of cells.

According to findings from our research, regardless of the number of platelets we use, we always applied 2-3 times more TGFβ-1 than PDGF-AB.

This means that the quantitative relationship between two platelet factors that are recognized as proliferation modulators, one positive and the other negative, is going to be the same. This fact is independent of the number of platelets the preparation contains.

We have proven this with experiments of *in vitro* cell cultures, using plasma with different numbers of platelets, but with the same PDGF-AB/TGFβ-1 relationship. (Fig. 6)

**Therapeutic dose and effect: Clinical Cases**

Will the clinical effect of these preparations depend on the number of platelets? Can the number of platelets predict clinical outcome? Will those patients with a greater number of platelets have a better outcome after application of PRGF?

To answer these questions, we chose the clinical case of an ulcer, since in ulcers the healing progress is seen from day-to-day and the therapeutic effect of these PRGF applications can be evaluated.
**Clinical case 1**

Clinical case 1: A 73-year-old female patient with osteoarthritis in the left knee, fatty hypertrophy and residual edema secondary to thrombophlebitis. Clinical history reveals renal failure, hypothyroidism, and chronic obstructive pulmonary disease (COPD).

A total arthroplasty of the knee was performed that developed a deep skin necrosis in which the patellar tendon was left exposed. (Fig. 7a). The patient was treated with antibiotics and a bacterial culture showed that the necrosis was not caused by an infection.

The surgeon recommended debridement and secondary closure. Over a period of 1 week, the wound was perfused with saline with no result and with a progressive increase in the size of the ulcer, which reached an approximate diameter of 6 cm (Fig. 7b). It was then decided to begin treatment with PRGF.

After debridement, newly-activated PRGF was placed so that the clot would form *in situ* inside the necrotized area (Fig. 7c).

This treatment was repeated weekly for 6 weeks. The ulcer healed from week to week (Figs. 7d, e, f and g) and was totally cured by the seventh week (Fig. 7h).

The number of the patient's platelets in the PRGF was very low (Fig 8a) in relation to the population mean values.

This anomaly in the number of platelets is related to her clinical history. The platelets in the PRGF, although scarce, were of a larger size.

The result, 3 years post-intervention and PRGF treatment of the ulcer is stable and the area is completely regenerated.

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Figs. 7a and b: The images show an ulcer approximately 6 cm in diameter and inflammatory reaction.
Fig. 7c: Image of clotted PRGF covering the entire area of the ulcer.

Fig. 7d: Progress of the ulcer in response to weekly treatment with PRGF (Image at 2 weeks).

Figs. 7e and f: Images at the second and fourth week of treatment.

Fig. 7g: Image of the ulcer at 5 weeks at the beginning of treatment with PRGF.

Fig. 7h: The image shows total cure of the ulcer after 7 weeks of treatment.
Fig. 8a: Number of platelets (millions/ml of PRGF) in a population of 65 individuals (mean values ± standard error) and number of platelets in PRGF of treated patient.

Fig. 8b: Concentration of PDGF-AB, TGFβ-1, IGF-I (pg/ml of PRGF) in a population of 65 individuals (mean values ± standard error) and concentrations of the aforementioned growth factors in the PRGF of the treated patient.

Fig. 8c: Concentration of EGF, VEGF, HGF (pg/ml of PRGF) in a population of 65 individuals (mean values ± standard error) and concentrations of these growth factors in the PRGF of the treated patient.

Clinical case 2

A 26-year-old male patient suffered traumatic amputation of the distal end of the index fingertip of the right hand. The patient was attended to in a trauma center where two grafts were performed to close the wound (Figs. 9a, b, c and d). When treated at the hospital, amputation was elected due to repeated necrosis of the grafts and patient was referred to us by a family member. PRGF treatment was initiated and applied weekly. Wound healing progressed positively from week to week (Figs. 9e - i) until it was totally cured at 5 weeks after commencement of treatment.
Figs. 9b and c: Initial situation.

Fig. 9d: 1st application of PRGF.

Fig. 9e: Exposed bone was remodelled with a gouge. The necrotic graft was painlessly eliminated without anesthesia.

Fig. 9f: Image of fingertip after 2nd application.

Fig. 9g: The image shows the cure after 3 weeks of treatment.
Figs. 9h and 9i: Image of fingertip after 5th week of treatment. Observe remodelling of bone made with a gouge and repair of injury.

Figs. 9j and 9k: Image one year after termination of treatment. The patient tells us that he uses the finger with total normality in daily life.

**Fig. 10a:**
Number of platelets (millions/ml) of PRGF in a population of 65 individuals (mean values ± standard error) and number of platelets in PRGF of patient with partial amputation of fingertip.
Final considerations:

Two clinical cases treated with PRGF have been described. In both cases, other conventional treatments were discontinued because they had been ineffective. Once the possibility of an infection that could impede the repair process had been discounted, the inability to repair could be due to the absence of cellular signals. The objective of PRGF is to provide those cellular signals in a matrix formed by sticky proteins such as fibrin, fibronectin and vitronectin.

The combined proteins are going to trigger angiogenesis and cellular proliferation. Furthermore, different types of cells will proliferate: cells reconstructing the localized injured tissue, as well as endothelial cells forming new vessels which are going to transport precursor cells and the nutrients necessary for tissue regeneration.

A very high number of platelets does not appear to be a requirement to start the repair mechanisms, as illustrated by the clinical cases. The use of PRGF offers considerable therapeutic benefits for treatment in a large number of clinical situations. Given the customary decrease of sequelae and the significant advantages for patients such as pain reduction, recovery of function and the fact that delayed healing or prolonged suffering caused by other more palliative treatments, even as irreversible as amputation in some cases, we clearly recognize a remarkably promising future for PRGF in regenerative medicine.

Legal treatment:

The Spanish Health Ministry (Ministerio de Sanidad Español), after consultation with the author and a detailed study of the PRGF technique, concluded that it cannot be considered a derived hemo therapy. Therefore, the requirements for its use are similar to those for other surgical procedures, regarding the quality of equipment, control of environmental conditions and TRAINING OF PERSONNEL that prepares and uses it and who should know its properties and limitations.

It is used following the methodology of an autograft, i.e., using only the patient's own blood which prevents immunological responses of rejection and complications intrinsic to the use of heterologous or homologous materials and substances. Its use is immediate so it does not require storage or transport.

As with all new technological advances, these new tools take time to become fully integrated into society and there are always sectors among professionals who show a great resistance to their incorporation. It is evident that, associated with these new therapeutic tools, there is the need to acquire new knowledge directed toward understanding the application and indications of these new treatments.
Biographical information

• Bachelor Degree in Medicine and Surgery from the Universidad de Salamanca in 1979.
• Doctor in medicine and surgery.
• Specialization in Stomatology from the Universidad del Pais Vasco, continuing studies on numerous visits to the United States (Philadelphia, New York, Miami, San Francisco, Chicago) and Europe (Italy, Germany, France, and Spain).
• Has led conferences in various Spanish universities, as a post-graduate assistant professor in Prosthetics and Occlusion at the Universidad de Valencia and post-graduate studies in Implantology at the Universidad de Oviedo.
• Has delivered more than two hundred courses and conferences at national and international conferences (in Europe, U.S., Mexico, South America, Asia) on Implants, Prosthetics, Dental Esthetics and Tissue Regeneration.
• Program director for “Continuing Education in Oral Implantology and Rehabilitation” in Spain and several countries around the world.
• Visiting Professor for dental school of the Universidad de: Guatemala, Intercontinental de Mexico, Javeriana de Colombia, República de Argentina, Uruguay, Portugal (School of Oporto and Lisboa).
• Scientific director at BTI - Biotechnology Institute.
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