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(71) Applicant(s)
Brightpulse Holding Ltd.

(72) Inventor(s)
Combette, Jean-Marc;Deloche, Catherine;Abadie, Claire;Mouz, Sebastien;Perino, Julien

(74) Agent / Attorney
FPA Patent Attorneys Pty Ltd, Level 43 101 Collins Street, Melbourne, VIC, 3000, AU

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(71) Applicant: **BRIGHTPULSE HOLDING LTD.** [CY/CY];
195 Arch Makarios III Ave, Neocleous House, Limassol,
3030 (CY).

(72) Inventors: **COMBETTE, Jean-Marc**; Rue de Terret 515,
F-74140 Saint-cergues (FR). **DELOCHE, Catherine**; Av-
enue William Favre 22, CH-1207 Geneva (CH). **ABADIE,
Claire**; 8 Rue des Francs Tireurs, F-74000 Annecy (FR).
MOUZ, Sebastien; 2757 Route des Vignes, F-74370
Villaz (FR). **PERINO, Julien**; 77bis Avenue de Gavot, F-
74500 Evian Les Bains (FR).

(74) Agent: **FIUSSELLO, Francesco**; STUDIO TORTA
S.p.A., Via Viotti, 9, I-10121 Torino (IT).

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Full      MASRNAYQLLVAAKEMKCCGSEALKCYTCKEEMISASCRITTRCKPEDIACMTILVTWEAE 80
Mature    -----LKYTCKEEMTSASCRITTRCKPEDIACMTILVTWEAE 38
          *****

Full      YFFNQSPVYIRSCSSSCVATRPDSIGAHLIFCCFRDLNSEL 103 SEQ ID NO:2
Mature    YFFNQSPVYIRSCSSSCVATRPDSIGAHLIFCCFRDLNSEL 81 SEQ ID NO:1
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FIG. 1

(57) Abstract: The present invention relates to a protein comprising SEQ ID NO:1 (mature form of SLURP-1) and to a composition comprising the same for use in inducing or accelerating cicatrisation, and/or in preventing infection in the eye of a subject.



PROTEIN SLURP-1 FOR USE IN THE TREATMENT OF OCULAR DISEASES

2013353957 17 Apr 2018

The present invention relates to a protein comprising SEQ ID NO:1 and a composition comprising the same for use in inducing or accelerating cicatrisation in the eye of a subject and/ or in preventing infection in the eye of a subject.

State of the art

Reference to any prior art in the specification is not an acknowledgment or suggestion that this prior art forms part of the common general knowledge in any jurisdiction or that this prior art could reasonably be expected to be understood, regarded as relevant, and/or combined with other pieces of prior art by a skilled person in the art.

Ocular, in particular corneal and conjunctival, lesions are one of the most diagnosed conditions in patients consulting their physician, and one of the major causes of sight loss. These lesions may be of various origins but are mainly due to allergy, infections (bacterial, viral and fungal), dry eye syndrome, surgery and other traumatisms. These lesions are harmful and very painful. Symptoms of these lesions may be dryness, burning and a sandy-gritty eye irritation. Symptoms may also be described as itchy, scratchy, stinging or tired eyes. Other symptoms are ocular pain, redness, a pulling sensation, and pressure behind the eye. The damage to the eye surface increases discomfort and sensitivity to bright light.

Ocular surface lesions need to be treated and healed up very rapidly to avoid worsening of the situation and complications such as ulceration, which may lead to loss of visual acuity and blindness in the most severe cases. For

the treatment of lesions due to dry eye syndrome, many lubricating solutions and hydrating hydrogels exist. However, these products only relieve the symptoms but do not accelerate the healing process of the lesions.

5 For deeper lesions, there are some vitamin A solutions and compositions enhancing mucin secretion, which may help in healing but have a limited efficiency.

Although there are some chemical compounds and biological molecules which have been found to be implicated
10 in wound healing in other tissues, the choice for the specific application to the eye is still very narrow and no effective molecules are currently available.

Human component B (hereinafter referred to as SLURP-1) is a member of the Ly-6/uPAR superfamily as demonstrated
15 via amino acid sequence comparison. This superfamily contains a C-terminal consensus sequence CCXXXXCN and different numbers of Ly-6/uPAR domain repeats. The whole sequence contains multiple cysteine residues (from eight to ten)^[1-3] leading to protein specific disulfide bonds.

20 The Ly-6/uPAR superfamily can be divided in two subfamilies depending on the presence of GPI anchoring signal sequences^[4].

SLURP-1 belongs to the first subfamily of the Ly-6/uPAR superfamily and has no GPI-anchoring signal
25 sequence. SLURP-2 is also a member of the same

subfamily^[4,5]. Patients with Mal de Meleda (MdM) are often associated with a mutation in SLURP-1 and the MdM gene is located in a cluster of Ly-6 genes on chromosome 8q24.3^[6-9].

5 The second subfamily comprises proteins with a GPI anchoring signal and includes several proteins: retinoic acid-induced gene E (RIG-E), E48 antigen (human Ly-6D), Ly6H, prostate stem cell antigen (PSCA), CD59 or protectin, Lynx1 and uPAR (urokinase receptor)^[10-15].

10 SLURP-1 is described in WO 94/14959. Briefly, the protein was first discovered and purified from a dialyzed urine concentrate after treatment with an adsorbing agent and a specific purification process. It is an 81 amino acid (SEQ ID NO:1) protein in its mature form (103 amino acid -
15 SEQ ID NO:2 - with the N-terminal signal peptide) which has a molecular weight around 8.9 kD. Its sequence is located on the long arm of human chromosome 8 as many of the Ly6/uPAR superfamily members, confirming a potential co-evolution after chromosomal duplication events.^[4]

20 SLURP-1 exists in two different forms depending on the presence of a sulfate group on the tyrosine at position 39: type 1 is sulfated while type 2 is not. SLURP-1 is detected in multiple organs and fluids like blood and urine. It is mainly produced by keratinocytes and epithelial cells,
25 hence having a main epithelial tissue distribution^[19-24].

Moreover, phylogenetic analysis demonstrated close relationship between SLURP-1 and Ly-6/uPAR superfamily members but also with snake neurotoxins^[4]. Snake neurotoxins show an inhibitory effect on acetylcholine receptors, hence inhibiting ion exchange between cells and extracellular media and subsequently preventing cell signalling^[16,17]. The important sequence homology between snake neurotoxins and SLURP-1 is emphasized by the three-dimensional structure similarity: SLURP-1 is likely to have a "three-finger" appearance, a specific feature of snake proteins^[18,19]. The resemblance of both proteins and the inhibitory activity of snake neurotoxins pointed out that SLURP-1 might be used to bind and interact with ion channels like nicotinic acetylcholine receptors.

Activation of nAChRs in non-neuronal cells could modify gene expression of proteins implicated in processes such as cell cycle regulation, apoptosis, cell-cell and cell-substrate interaction. The main studied homomeric receptor is composed of $\alpha 7$ subunits; the activation of this specific receptor is transduced by different initial signals however leading to a common end point. Transduced signals simultaneously involve ionic events and protein kinase signaling. Chernyavsky et al concluded that this twin activation (ionic events and protein kinase signaling) could lead to gene expression and simultaneous changes in

cell morphology (and subsequent locomotion of keratinocytes)^[20]. The simultaneous and complementary signaling was clearly demonstrated by inhibiting either ionic or protein signaling which led to partial inhibition while inhibiting both mechanisms led to complete inhibition of their effects.

The interaction between SLURP-1 and nicotinic receptors was confirmed in different publications and in the presence of acetylcholine, an agonist activity (contrary to snake neurotoxins) was discovered on human keratinocytes while using SLURP-1^[21]. This interaction regulates cell functions through cholinergic pathways and probably leads to epithelial cell adhesion, motility and wound healing^[22,23], however discordant results were obtained by a team leading to conclude that SLURP-1 was slowing down healing process while SLURP-2 was accelerating it.^[26] These results were yet to be compared with a third experiment from the same group stating that the combination of SLURP-1 and SLURP-2 leads to an additive positive effect on the skin healing process when compared to either SLURP-1 inhibiting or SLURP-2 accelerating activities on their own.

Further research is needed to ascertain whether SLURP-1 or SLURP-2 are useful for wound healing in general.

Moreover, the need is strongly felt in the field of ophthalmics to develop medicaments and related compositions

to promote healing following ocular disease with high efficiency, no counter-effects and recovery of complete corneal transparency.

It is an object of the present invention to provide
5 biological molecules and/or compositions for use in the treatment of ocular disease, in particular in inducing or accelerating cicatrisation, reducing inflammation, and preventing infection in the eye of a subject.

The above said object is achieved by the present
10 invention, as it relates to a protein as defined in claim 1.

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which
15 this invention pertains. Although many methods and materials similar or equivalent to those described herein may be used in the practice or testing of the present invention, preferred methods and materials are described below. Unless mentioned otherwise, the techniques described
20 herein for use with the invention are standard methodologies well known to persons of ordinary skill in the art.

Brief description of the drawings

Figure 1 shows a sequence comparison between the amino
25 acid sequence of the full-length, SEQ ID NO:2, and mature

SLURP-1 protein, SEQ ID NO:1, (respectively amino acids 1-103 and amino acids 23-103)

Figure 2 shows images of fluorescein staining of corneal epithelial wounds at the indicated time points after total alcohol-induced des-epithelialization in control and SLURP-1 treated groups. The control groups include one group that received no treatment (control) and one group that was treated with subconjunctival injection of the vehicle (PBS) (SCJ vehicle). Bold percentages indicate the group means of the corneal wound surface, and percentages in brackets indicate the value of the representative cornea for the considered group.

Figure 3A is a graph showing the time course of corneal wound healing after total des-epithelialization in control and SLURP-1 treated groups. Calculated healing rate means \pm SEM.

Figure 3B shows a graph of the healing rate on day two after total des-epithelialization in control and SLURP-1 treated groups (subconjunctival administration with 50 ng of SLURP and instillation).

Figure 3C shows a graph of the healing rate on day three after total des-epithelialization in control and SLURP-1 treated groups (subconjunctival administration with 50 ng of SLURP and instillation).

Figure 4 shows images of fluorescein staining of

corneal epithelial wounds at the indicated time points after calibrated alcohol-induced des-epithelialization in PBS and SLURP-1 treated groups. Bold percentages indicate the group mean of the corneal wound surface, and 5 percentages in brackets indicate the value of the representative cornea for the considered group.

Figure 5A is a graph showing the time course of measured wound area from day 0 to day 2 after des-epithelialization in control and SLURP-1 treated groups.

10 Figure 5B is a graph of the calculated healing rate between T0h and T47h after calibrated des-epithelialization in control and SLURP-1 treated groups.

Figure 5C is a graph of the degree of wound healing at T47h after calibrated des-epithelialization. The degree of 15 wound healing was calculated by the ratio: area of the fluorescein wound at T47h / area of the fluorescein wound at baseline, for each cornea, for each group, and expressed in percentage.

Figure 6 shows photographs of wound healing 20 experiments carried out by treating the human corneal epithelial cell line hTCEpi with SLURP-1.

Detailed description of the invention

According to the present invention a protein comprising SEQ ID NO:1 is used in inducing or accelerating 25 cicatrisation in the eye of a subject and/or in preventing

infection in the eye of a subject.

Preferably the ocular disease is selected from the group consisting of diabetic keratinopathy, keratitis, conjunctivitis, keratoconjunctivitis, uveitis, corneal
5 trauma, corneal abrasion, corneal burns, corneal chronic ulcer, corneal dystrophies, persistent corneal epithelial defect (PED), corneal epithelial defect after laser surgeries, corneal epithelial defect post-PRK, corneal epithelial defect post-transcorneal transplant.

10 SEQ ID NO:1 corresponds to the 81 amino acid mature form of SLURP-1.

In an alternative embodiment, according to the present invention a composition comprising a protein comprising SEQ ID NO:1 is used in inducing or accelerating cicatrisation,
15 and/or in preventing infection in the eye of a subject.

The composition preferably comprises at least one biocompatible polymer.

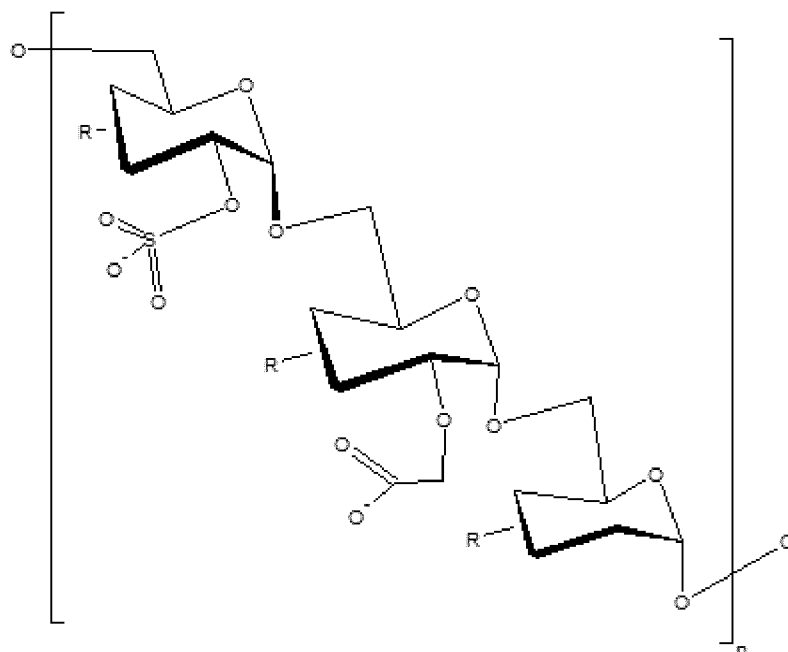
More preferably, the biocompatible polymer is selected from the group consisting of hyaluronic acid, sugar
20 polymers, lecithin gels, polyalanine derivates, pluronics, poly(ethylene)glycol, poloxamers, chitosan, xyloglucan, collagen, fibrin, polyorthoesters and mixtures thereof.

Even more preferably the biocompatible polymer is hyaluronic acid.

25 Alternatively, the biocompatible polymer is preferably

a sugar polymer, even more preferably dextran (a glucose polymer).

Preferably, the dextran is a carboxymethyl dextran sulphate polymer, more preferably a ReGeneraTing Agent
 5 (RGTA®), even more preferably OTR4120 (formula I) which is marketed under the name of Cacicol 20®.



Formula I represents an analogue subunit of heparan
 10 sulfate with a glucose subunit based backbone.

The composition can be solid, semi-solid, gel-like or liquid; moreover, it can be a solution, a suspension, an emulsion or a thermosetting gel.

In an alternative embodiment the composition comprises

at least one nanoparticle carrier.

The nanoparticle carrier is preferably selected from the group consisting of poly- ϵ -caprolactone, polycyanocrylate and chitosan.

5 In an alternative embodiment, the composition is in the form of an injectable viscous polymer composition. In particular the polymers of the composition are poly-orthoesters.

The composition is preferably administered by topical
10 treatment or subconjunctival injection.

More preferably, the amount of protein comprising SEQ ID NO:1 is from 5 ng to 50 μ g per administration unit, more preferably from 10 ng to 10 μ g per administration unit.

In a preferred embodiment the composition is
15 administered by subconjunctival injection and the amount of protein comprising SEQ ID NO:1 is from 20 ng to 90 ng per administration unit.

Examples

Example 1: corneal wound healing after topical or sub-
20 conjunctival administration.

Figures 2 and 3 report the data of a first experiment in which the cicatrizing/ocular wound healing activity of SLURP-1 was compared using two routes of administration in a des-epithelialization model in rats. The first route of
25 administration is a common route for ocular products used

for moisture maintenance of the cornea and specifically for its ease of use: instillation (topical application). This route was used with a six times/day administration frequency at days 0 and 3 after a surgical complete des-
5 epithelialization. Following the same surgical treatment, a second set of animals was treated by a different route of administration: the sub-conjunctival route. The latter administration was practiced only once daily at days 0 and 3.

10 Both routes of administration have a common feature, namely the SLURP-1 applied quantities: the group receiving instillations was treated with a solution corresponding to the highest dose used in the group treated sub-
15 conjunctively. It could be easily conceivable that products administered via the sub-conjunctival route would be available for a longer time during wound treatment as they might be slowly released by the conjunctiva tank they are stored in; conversely treatment administered topically would be quickly eliminated from the eye surface. Following
20 this hypothesis, the dose regimen used in the sub-conjunctival treatment was in a range from the dose used topically (5 μ g) and gradually decreasing to 50 ng.

As shown in Figure 3A, both routes of administration were capable of accelerating the wound healing process.

25 On day 2 after total des-epithelialization, the

healing rate after a sub-conjunctival injection of 50 ng SLURP-1 was significantly higher as compared to a sub-conjunctival injection of PBS ($p < 0.05$) (Figure 3B).

The unexpected discovery was concerning the sub-
5 conjunctival route of administration in which the cicatrizing effect was more pronounced at the lowest dose (50 ng) when compared to the dose used in topical administration which in terms of quantity is much higher.

The total administered quantity was calculated in both
10 routes of administration regimen giving similar efficacy results (that is 1 $\mu\text{g}/\text{mL}$ for sub-conjunctival and 100 $\mu\text{g}/\text{mL}$ for topical route). In the topical treatment, 6 administrations of 50 μL drops were performed per day on two timely separate days with a 100 $\mu\text{g}/\text{mL}$ solution. Then
15 the final quantity used in this treatment group was $6 * 2 * 5 \mu\text{g} = 60 \mu\text{g}$. In the sub-conjunctival treatment, a 1 $\mu\text{g}/\text{mL}$ solution was injected twice (50 μL volume) thus the overall quantity given by this route was $2 * 0.05 \mu\text{g} = 0.1 \mu\text{g}$. When
20 comparing both administered quantities leading to similar clinical efficacy, a ratio of 600 between sub-conjunctival and topical quantities is observed.

This is clearly in favour of a severely increased efficacy of the product by this specific less common route of administration for this product in order to treat ocular
25 wound healing and other stated clinical conditions.

More in particular, the experiment involved mechanical resection of the whole corneal epithelium performed under a surgical microscope using a 70% ethanol-soaked microsponge and a 15° surgical knife. The cornea was then rinsed with
5 0.9% NaCl and the effect of the treatment on corneal re-epithelialization was evaluated using topical 0.5% fluorescein, at 5 time-points after des-epithelialization (D1, D2, D3, D6 and D7 corresponding respectively to T24h, T48h, T72h, T144h and T168h).

10 Six groups of 5 animals received the following treatment:

Group 1: 50 µL vehicle subconjunctival injection at D0 and D3

15 Group 2: 1 µg/mL (50 µL) SLURP-1 subconjunctival injection (50 ng per administration), at D0 and D3

Group 3: 10 µg/mL (50 µL) SLURP-1 subconjunctival injection (500 ng per administration) at D0 and D3

Group 4: 100 µg/mL (50 µL) SLURP-1 subconjunctival injection (5 µg per administration) at D0 and D3

20 Group 5: 100 µg/mL (50 µL) SLURP-1, 6 topical instillations (5 µg per administration) at D0 and D3

Group 6: No treatment

At each time-point (D1, D2, D3, D6 and D7), full-face photographs were taken using cobalt blue biomicroscope
25 light and the green fluorescent labeling of the cornea was

used to determine the shape and area of the remaining ulcer (Figure 2). The time course of the healing rate is presented on graph for each group in order to compare "control" groups (vehicle and/or control without any treatment) versus SLURP-1 treated groups (Figure 3A). The wound healing follow up of each cornea was performed through computer-assisted measures of the wound area (A_t): ratio of the fluorescein-stained area to the total corneal area. For each time point, the healing rate: $(A_{t0}-A_t)/A_{t0}$ was calculated. Results are presented as mean \pm SEM. An ANOVA test followed by the Dunnett's multiple comparison test or the non-parametric Mann Whitney comparison test were performed using *GraphPad Prism* (GraphPad Software, San Diego, U.S.A.).

The model induced a limbic insufficiency leading to corneal neovascularization and slowing the healing process. Corneas that presented a strong central ulcer were excluded from the study.

As shown in Figure 3B, on day 2 after total des-epithelialization, the healing rate after a sub-conjunctival injection of 50 ng of SLURP-1 is significantly higher as compared to a sub-conjunctival injection of PBS ($p < 0.05$ Dunnett) and controls ($p < 0.01$ Dunnett). The healing rate after instillation of SLURP-1 is also significantly higher than controls ($p < 0.05$, Dunnett), but is not

significantly different than after PBS injection.

As shown in Figure 3C, on day 3 after total des-epithelialization, the healing rates of the SCJ 50 ng SLURP-1 treated group and instillation group are
5 significantly higher than the control group ($p < 0.05$, Dunnett) but are not significantly different than the healing rate with PBS.

This observation suggests that the SCJ procedure (or PBS) itself induces a response promoting corneal healing.

10 As regards neovascularization, neovessels were visible in each group just after the beginning of the re-epithelialization process. The qualitative follow up of the corneal neovascularization did not demonstrate any difference between the SLURP-1-treated groups and controls.
15 This is a very important observation showing that the healing properties of SLURP-1 are not associated with proangiogenic effects and do not impair restoration of corneal transparency.

This means that SLURP-1 showed a significant corneal
20 effect after sub-conjunctival administration and instillation, in the absence of side effects, and particularly, in the absence of corneal neovascularization that is associated with limbal deficiency, demonstrating that the healing effect is not linked to the corneal trans-
25 differentiation process.

Example 2: corneal wound healing after topical or sub-conjunctival administrations in a calibrated corneal des-epithelialization model

Following the preliminary results obtained in the first corneal des-epithelialization model that implied not only a healing process but also a limbic insufficiency, a calibrated model of alcohol-induced des-epithelialization was chosen as a second study model in order to focus on the corneal healing properties of SLURP-1. The objective of this study was to confirm the preliminary results showing an enhancement of the re-epithelialization by the tested protein as compared to controls, and to evaluate a possible dose-response induced by the protein.

The model involved a calibrated wound created using a trephine causing a 4mm diameter circular incision. Healing of such a minute wound is very fast even in the absence of any treatment (sometime around 2 days). Hence administrations were performed only once, on the day of incision, and efficacy was evaluated during the first 48h by an examination twice a day.

The injected volume was adjusted from the first experiment; doses were equivalent with twice the administration volume and half concentrated solutions. This different dosing was chosen to increase the retention time in the tank created by sub-conjunctival injection

leading to a prolonged presence of SLURP-1 on the corneal surface to compensate the unique administration performed in this standardized model.

In this second experimental model, SLURP-1 effects were confirmed; namely the sub-conjunctival route with low injected quantities was 6 to 60 times more efficient in accelerating ocular wound healing when compared to the topical route.

More in particular, corneal epithelial wounds were obtained as described by Hattori *et al* [25]. Briefly, after systemic and topical anesthesia, a trephine was used to make a 4-mm-diameter circular incision that was centered on the cornea. Then, the 4-mm diameter circular filter paper, which had been soaked with 70% ethanol, was placed on the incised area for 5 seconds. After a gentle wash with 5 mL saline, the detached corneal epithelium was removed.

Treatment was administered only in the right eye via 1 single subconjunctival injection or 6 instillations at D0 (time of corneal des-epithelialization). Five groups of 4-6 animals received the following treatment:

Group 1: 0.5 µg/mL (100 µL) SLURP-1 sub-conjunctival injection (50 ng per administration), at D0

Group 2: 5 µg/mL (100 µL) SLURP-1 sub-conjunctival injection (500 ng per administration), at D0

Group 3: 50 µg/mL (100 µL) SLURP-1 sub-conjunctival

injection (5 µg per administration), at D0

Group 4: 100 µg/mL (50 µL) SLURP-1 (5 µg per administration), 6 instillations at D0

Group 5: Vehicle subconjunctival injection (100 µL) at
5 D0

The right corneas were examined with a biomicroscope. Each right cornea received one drop of 0.5% fluorescein (Novartis pharma S.A.S) and was examined using a cobalt blue light. Digital photographs were taken through the
10 binocular. During the healing follow-up in the calibrated corneal des-epithelialization models, corneas were examined at day 0, day 1 AM (T23h), day 1 PM (T28h), day 2 AM (T47h) and day 2 PM (T56h) (Figure 4).

Digital photographs were analyzed with imaging
15 software (*Adobe Photoshop*). For each eye and each time-point, the area of the remaining ulcer was compared to the total corneal area. Time course of the healing rate is presented on graph for each group in order to compare control (vehicle and/or control without any treatment)
20 versus SLURP-1 treated groups. Results are presented as mean ± SEM. An ANOVA test followed by the Dunnett's multiple comparison test or the non-parametric Mann Whitney comparison test were performed using *GraphPad Prism* (GraphPad Software, San Diego, U.S.A.).

The time courses of the corneal wound closure in the control and SLURP-1 treated groups are shown in Figure 5A. When comparing the healing rate calculated between T0h and T47h, SCJ injection of 500 ng and 5 µg of SLURP-1 significantly increased the healing rate as compared to the instillation or PBS sub-conjunctival administration groups (Figure 5B). There was a dose response effect since no significant effect was observed in the 50 ng-SCJ injection group. Moreover, at T47h, the number of totally healed corneas was higher in the groups treated with sub-conjunctival injection of SLURP-1 at either 500 ng or 5 µg (Figure 5C), with a noticed difference between the 500 ng (4 out of 5 corneas) and 5 µg (5 out of 5 corneas) as compared to the control group (3 out of 6 corneas).

Example 3: Wound closure assay with SLURP-1 on human corneal epithelial cell line hTCEpi.

A wound closure assay used to monitor cell migration is the Oris Cell Migration Assay-Collagen I Coated from Platypus Technologies. Cell seeding density was determined visually using an inverted microscope. 100 microliter of optimal cell seeding density was pipetted into test wells and incubated in a humidified chamber (37°C, 5% CO₂) for 1-4 hours to allow cell attachment. Cytochalasin D was used as positive control.

Experiments were carried out in triplicate with increasing amounts of SLURP-1. The amounts tested were 0 µg/ml (negative control), 1 µg/ml, 5 µg/ml, 10 µg/ml, 15 µg/ml, 25 µg/ml and 50 µg/ml. After 24h the wounded areas on the plate were healed at different degrees. Cells treated with 10 µg/ml of SLURP-1 showed optimal wound healing as may be seen in Figure 6.

From an analysis of the above data the advantages the present invention allows to achieve are apparent.

In particular, an effective association between a specific concentration of SLURP-1 and a particular route of administration gives optimal results in terms of accelerated healing rate and faster reduced wound area, in particular protecting ocular surface from infection.

Moreover, the healing properties of SLURP-1 are not associated with proangiogenic effects and do not impair restoration of corneal transparency.

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CLAIMS

1. A method of inducing or accelerating cicatrisation in the eye of a subject which comprises administering to the subject an effective amount of a protein comprising SEQ ID NO:1.

2. A method of preventing infection in the eye of a subject which comprises administering to the subject an effective amount of a protein comprising SEQ ID NO:1.

3. A method of inducing or accelerating cicatrisation in the eye of a subject or preventing infection in the eye of a subject which comprises administering to the subject a composition comprising a protein comprising SEQ ID NO:1 and at least one biocompatible polymer.

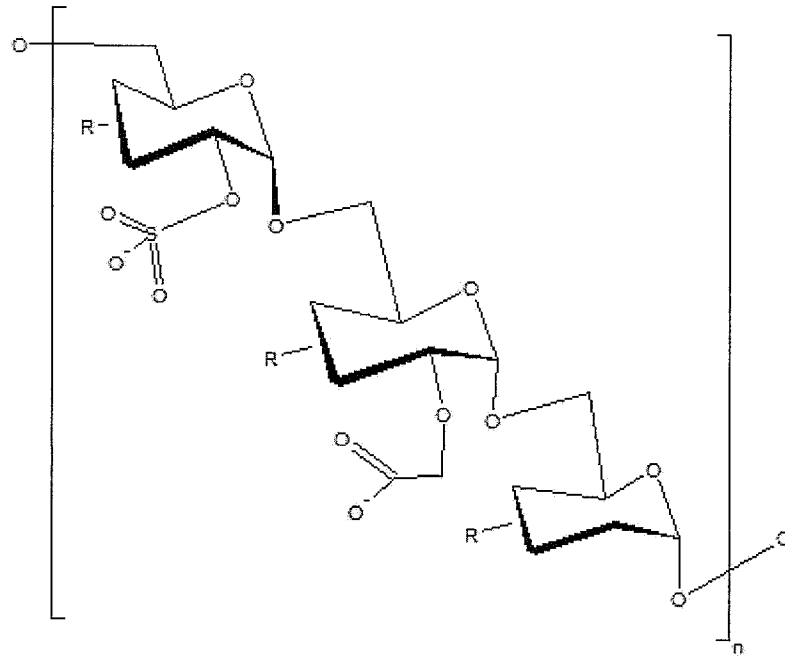
4. The method according to claim 3, wherein the biocompatible polymer is selected from the group consisting of hyaluronic acid, sugar polymers, lecithin gels, polyalanine derivates, pluronics, poly(ethylene)glycol, poloxamers, chitosan, xyloglucan, collagen, fibrin, polyorthoesters and mixtures thereof.

5. The method according to claim 4, wherein the biocompatible polymer is hyaluronic acid and/or a sugar polymer.

6. The method according to claim 5, wherein the sugar polymer is a dextran.

7. The method according to claim 6, wherein the dextran is a carboxymethyl dextran sulphate polymer.

8. The method according to claim 7, wherein the carboxymethyl dextran sulphate polymer is



9. A method of inducing or accelerating cicatrisation in the eye of a subject or preventing infection in the eye of a subject which comprises administering to the subject a composition comprising a protein comprising SEQ ID NO:1 and at least one nanoparticle carrier.

10. The method according to claim 9, wherein the nanoparticle carrier is selected from the group consisting of poly- ϵ -caprolactone, polycyanocrylate and chitosan.

11. The method according to any one of claims 3 to 10, wherein the composition is administered by topical treatment or subconjunctival injection.

12. The method according to claim 11, wherein the amount of protein comprising SEQ ID NO:1 is from 5 ng to 50 μ g per administration.

13. The method according to claim 12, wherein the amount of protein comprising SEQ ID NO:1 is from 10 ng to 10 μ g per administration.

14. The method according to claim 13, wherein the composition is administered by subconjunctival injection and the amount of protein comprising SEQ ID NO:1 is from 20 ng

to 90 ng per administration.

15. The method according to claim 14, wherein the amount of protein comprising SEQ ID NO:1 is from 40 ng to 60 ng per administration.

5 16. Use of a protein comprising SEQ ID NO:1 for the preparation of a medicament for inducing or accelerating cicatrisation in the eye of a subject.

10 17. Use of a protein comprising SEQ ID NO:1 for the preparation of a medicament for preventing infection in the eye of a subject.

15 18. Use of a composition comprising a protein comprising SEQ ID NO:1 and at least one biocompatible polymer for the preparation of a medicament for inducing or accelerating cicatrisation in the eye of a subject or for preventing infection in the eye of a subject.

20 19. Use of a composition comprising a protein comprising SEQ ID NO:1 and at least one nanoparticle carrier for the preparation of a medicament for inducing or accelerating cicatrisation in the eye of a subject or for preventing infection in the eye of a subject.

CLUSTAL 2.1 multiple sequence alignment

Full YASRWAVQLLLVAAWEMGCGEALKCYTCKEPMTSASCRITIRCKPEDTACMTILVTVEAE 60
 Mature -----LRCYTCKEPMTSASCRITIRCKPEDTACMTILVTVEAE 38

Full YFFNQSPVWIRSCSSSCVAIDPDSIGAAHLIFCCFRDLNSEL 103 SEQ ID NO:2
 Mature YFFNQSPVWIRSCSSSCVAIDPDSIGAAHLIFCCFRDLNSEL 81 SEQ ID NO:1

FIG. 1

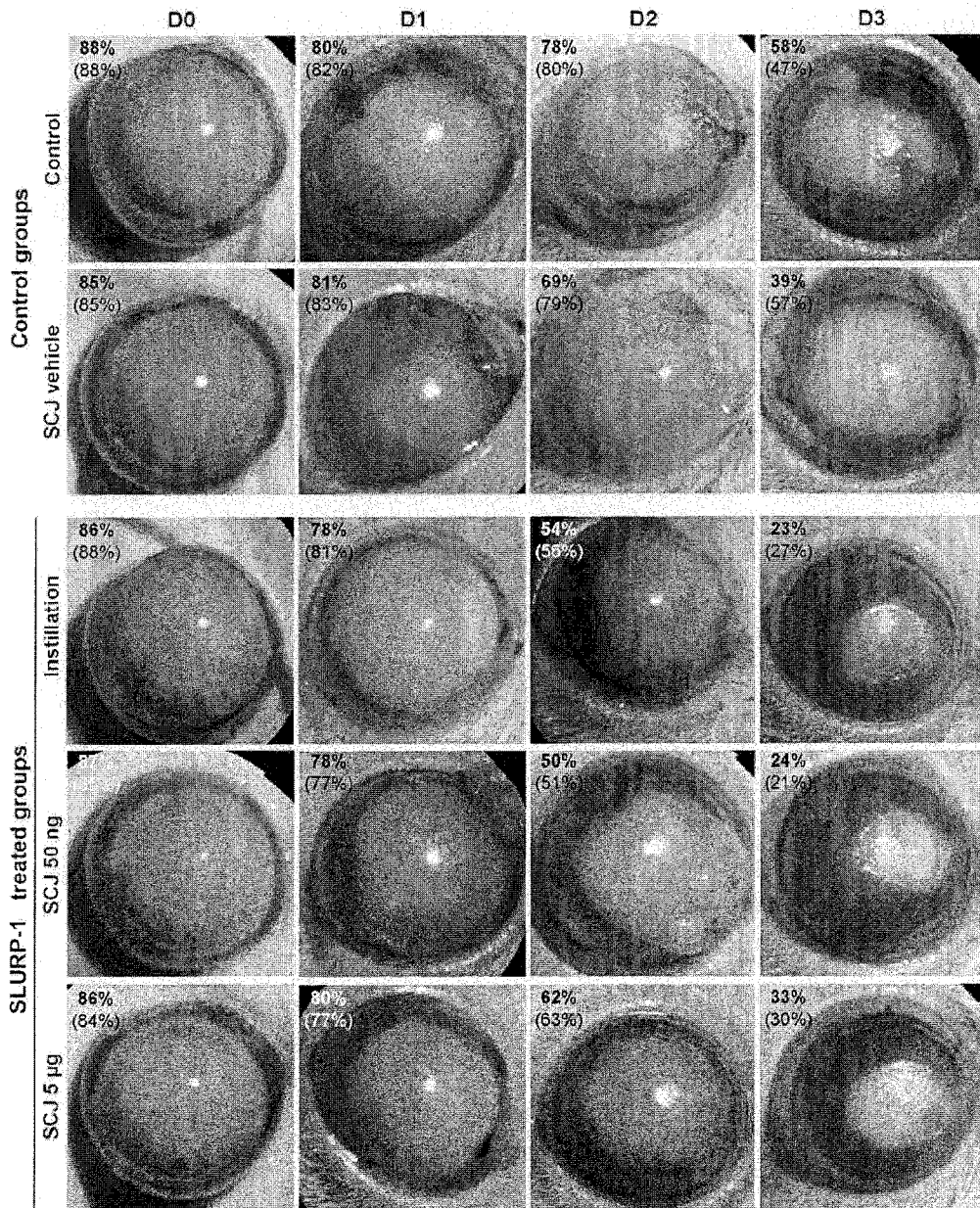


FIG. 2

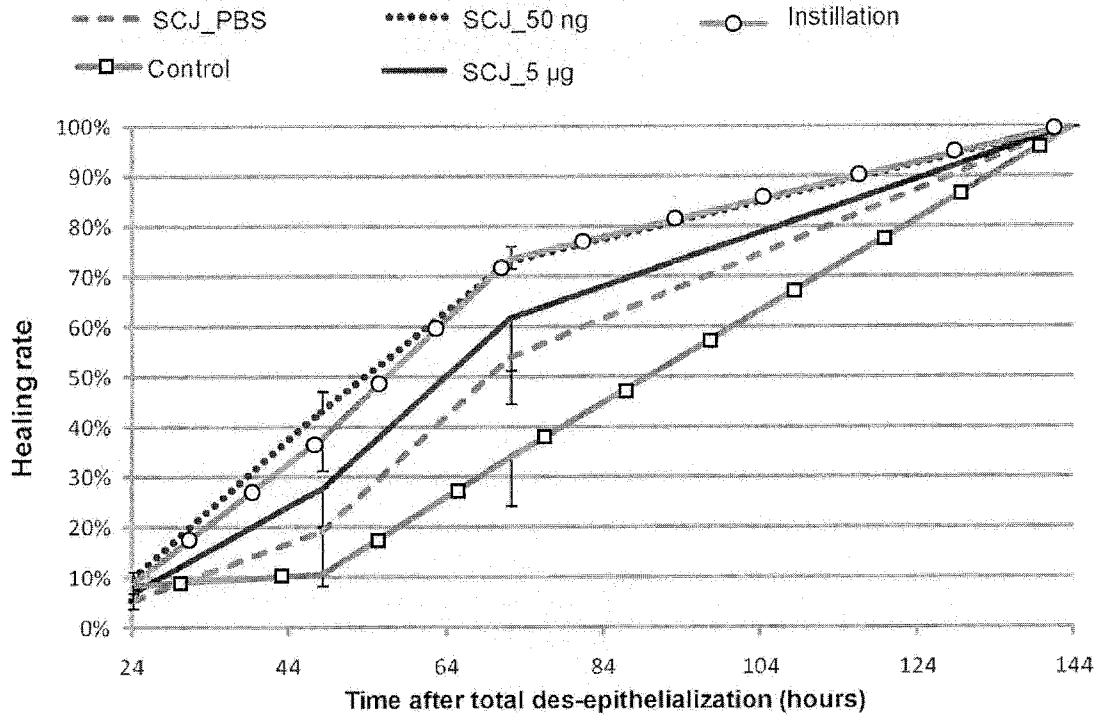


FIG. 3a

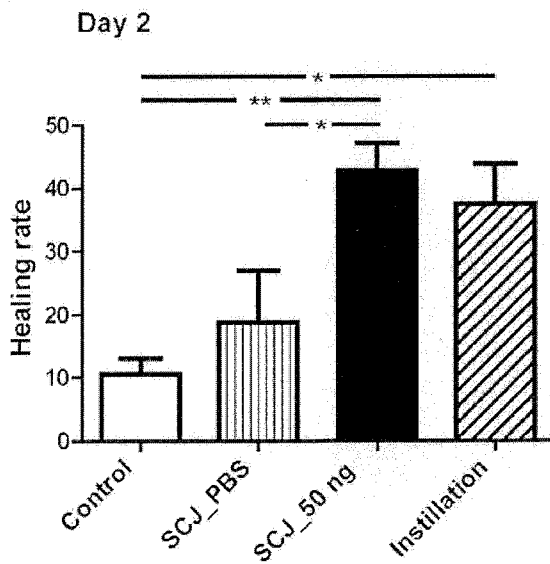


FIG. 3b

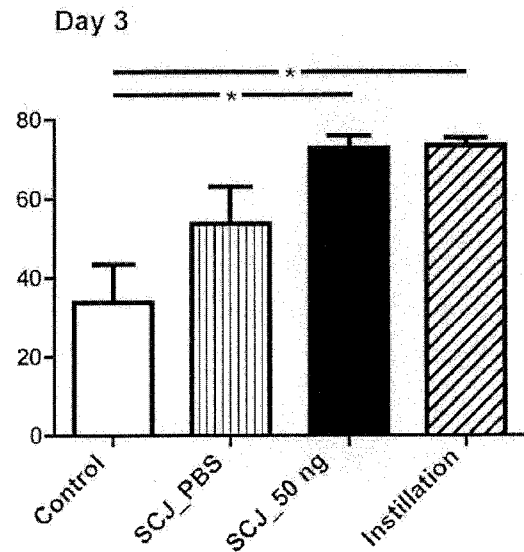


FIG. 3c

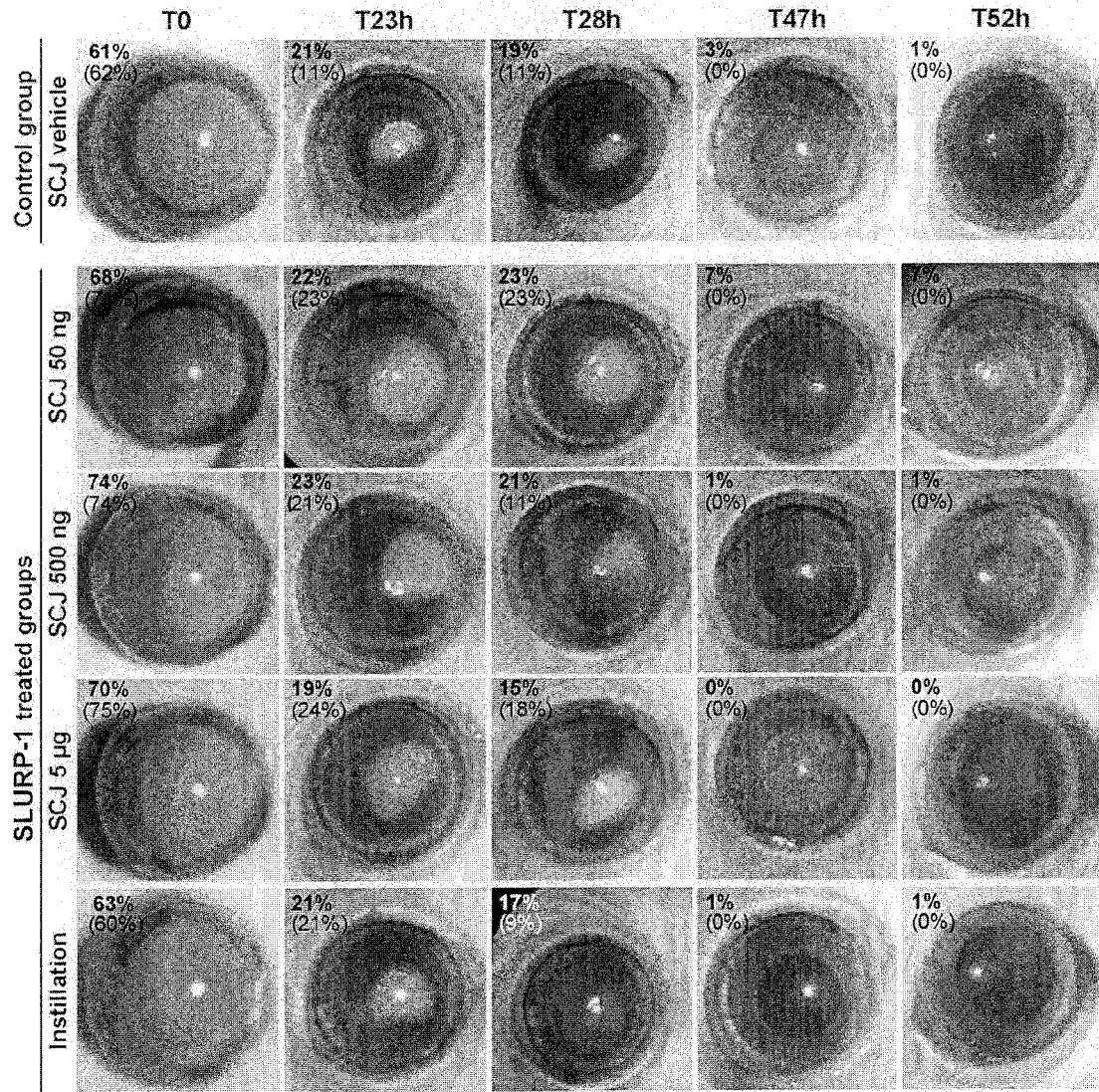


FIG. 4

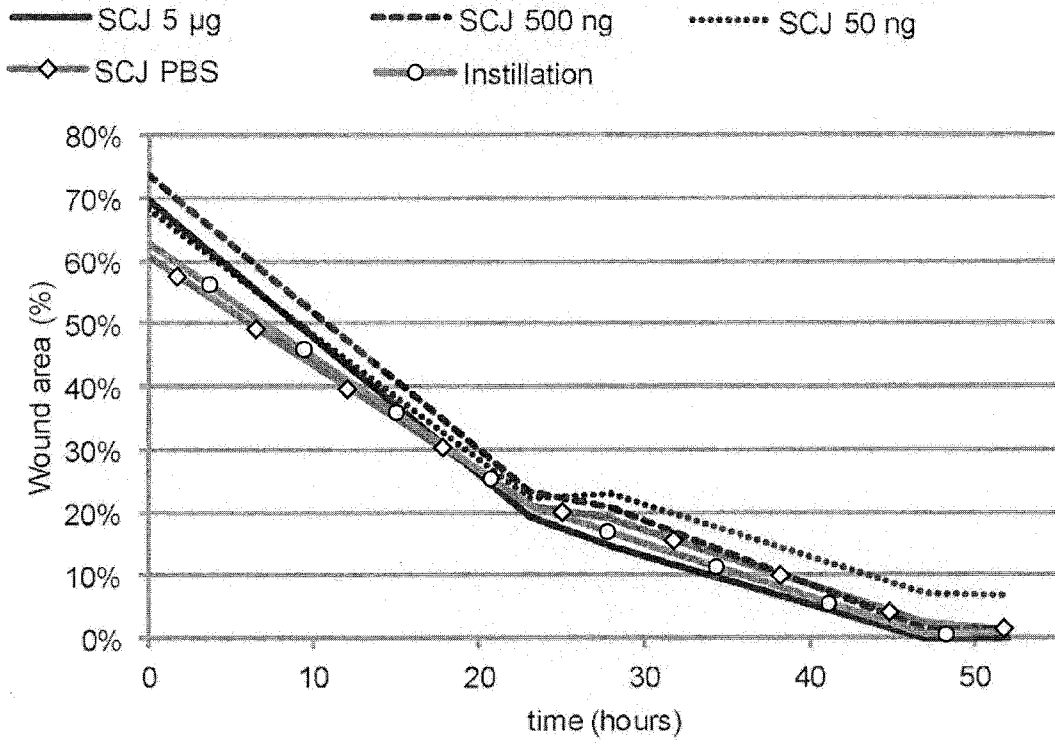


FIG. 5a

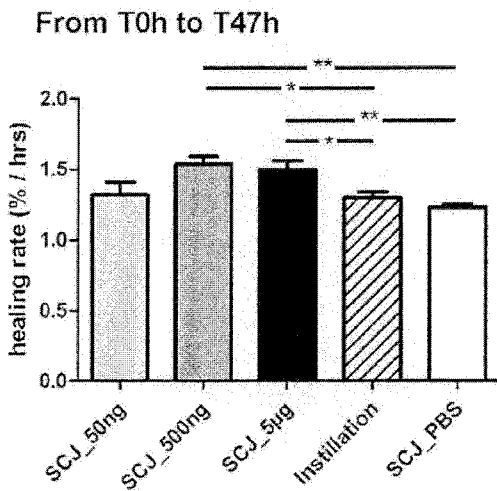


FIG. 5b

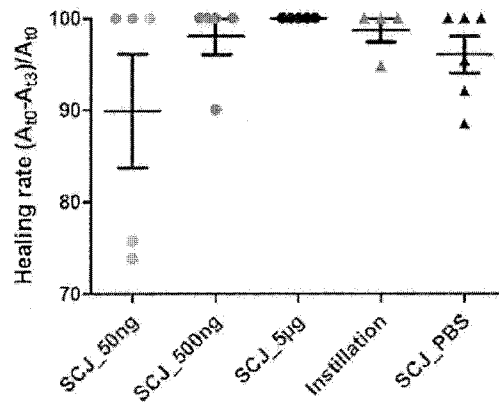
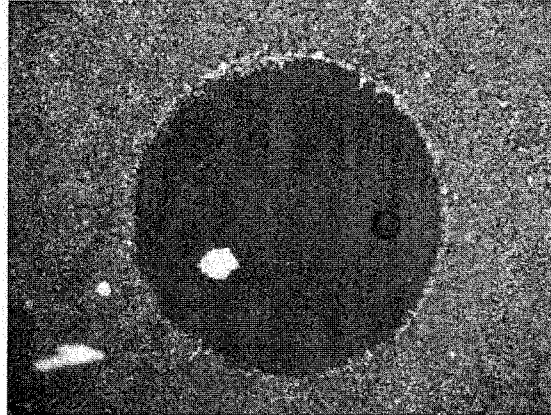
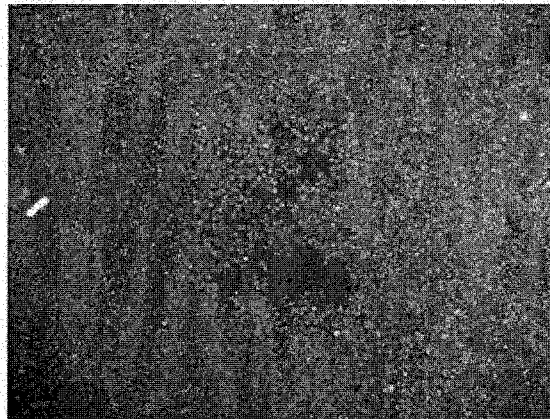


FIG. 5c

100% open wound, 0% wound closure



10 μ g/ml SLURP-1



Control (No treatment)

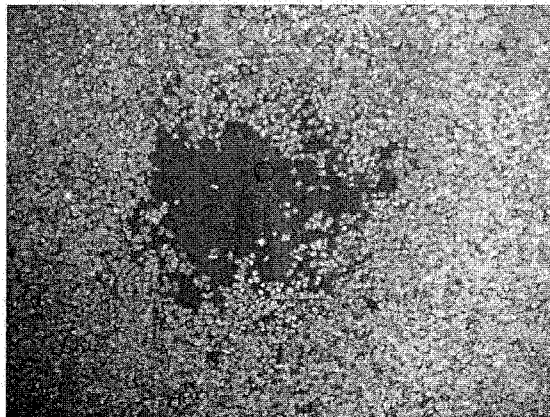


FIG. 6

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35          40          45
Arg Ser Cys Ser Ser Ser Cys Val Ala Thr Asp Pro Asp Ser Ile Gly
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65          70          75          80
Leu

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Thr Ser Ala Ser Cys Arg Thr Ile Thr Arg Cys Lys Pro Glu Asp Thr
35          40          45
Ala Cys Met Thr Thr Leu Val Thr Val Glu Ala Glu Tyr Pro Phe Asn
50          55          60
Gln Ser Pro Val Val Thr Arg Ser Cys Ser Ser Ser Cys Val Ala Thr
65          70          75          80
Asp Pro Asp Ser Ile Gly Ala Ala His Leu Ile Phe Cys Cys Phe Arg
85          90          95
Asp Leu Cys Asn Ser Glu Leu
100

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