Immunomodulating and Revascularizing Activity of Kalanchoe pinnata Synergize with Fungicide Activity of Biogenic Peptide Cecropin P1

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Received 16 January 2017; Accepted 10 April 2017; Published 11 June 2017

Academic Editor: Sergey Morzunov

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Previously transgenic Kalanchoe pinnata plants producing an antimicrobial peptide cecropin P1 (CecP1) have been reported. Now we report biological testing Kalanchoe extracts containing CecP1 as a candidate drug for treatment of wounds infected with Candida albicans. The drug constitutes the whole juice from K. pinnata leaves (not ethanol extract) sterilized with nanofiltration. A microbicide activity of CecP1 against an animal fungal pathogen in vivo was demonstrated for the first time. However, a favorable therapeutic effect of the transgenic K. pinnata extract was attributed to a synergism between the fungicide activity of CecP1 and wound healing (antiscar), revascularizing, and immunomodulating effects of natural biologically active components of K. pinnata. A commercial fungicide preparation clotrimazole eliminated C. albicans cells within infected wounds in rats with efficiency comparable to CecP1-enriched K. pinnata extract. But in contrast to K. pinnata extract, clotrimazole did not exhibit neither wound healing activity nor remodeling of the scar matrix. Taken together, our results allow assumption that CecP1-enriched K. pinnata extracts should be considered as a candidate drug for treatment of dermatomycoses, wounds infected with fungi, and bedsores.

1. Introduction

Kalanchoe pinnata (Bryophyllum pinnatum) (Lamarck) Persoon (Crassulaceae) is a succulent perennial plant native to Madagascar. It is used for healing wounds in a traditional medicine to treat psychiatric disorders and as a tocolytic agent to prevent premature labour [1]. Extracts of K. pinnata and some other Kalanchoe species (K. crenata, K. brasiliensis, and K. daigremontiana) are reported to exhibit antimicrobial [2, 3] and virucide [4–6] activity. Further, they were found to kill leishmanias [7] and malaria plasmodium [8]. It induces the relaxation of smooth muscles [9]; exhibits antimutagenesis, antihistamine [10], and hepatoprotective activity [11]; causes immunomodulatory and anti-inflammatory effects [12, 13]; and inhibits thyroid peroxidase [14]. Baginskaya and Leskova [15] reported a positive effect of K. pinnata extract towards experimental gastric ulcer in mice. Eventually, it demonstrates an oncolytic activity on certain models [16]. To our best knowledge, primary experimental data about testing antifungal activity of Kalanchoe extracts have not been published, but occurrence of this effect is mentioned in a review by Kutsik and Zuzuk [17].
Like other species of the genus Kalanchoe, K. pinnata contains a number of biologically active compounds which may contribute to pharmacological properties of its extracts:

(1) Flavonoid glycosides: two phenolic glycosides, syringic acid \(\beta\)-D-glucopyranosyl ester and 4′-O-\(\beta\)-D-glucopyranosyl-cis-p-coumaric acid; nine flavonoids including kaempferol, quercetin, myricetin, acacetin, and diosmetin glycosides; and flavonol glycosides: quercetin \((3\text{-}O\text{-}\alpha\text{-L-arabinopyranosyl}(1\rightarrow2)-\alpha\text{-L-}

	 \text{rhamno} \text{y} \text{n} \text{os} \text{y} \text{ran} \text{os} \text{id} \text{e})\); 7-O-\(\beta\)-D-glucopyranoside); and myricetin \((3\text{-O\text{-}\alpha\text{-L-arabinopyranosyl}(1\rightarrow2)-\alpha\text{-L-}

	 \text{rhamno} \text{y} \text{n} \text{os} \text{y} \text{ran} \text{os} \text{id} \text{e})\) [18].

(2) Four bufadienolides: bersaldegenin-1-acetate, bryo-phyllin A, bersaldegenin-3-acetate, and bersaldegenin-1,3,5-orthoacetate [18].

(3) Blood-agglutinating lectins with Mr 44–47 kDa containing ~1.5% carbohydrate [19, 20].

Over the last decade, ethanol extracts became the most popular form of Kalanchoe medicinal application [21–23]. It means that lipophilic constituents of the fresh juice (bufadienolides, polyphenols, and flavonoids) are suggested to confer essential part of their total activity. However, an evident biological effect of Kalanchoe lectins has been formerly documented. Lapchik et al. demonstrated a high mitogenic and blast-transforming activity of lectins from different Kalanchoe species, particularly from K. blossfeldiana [24]. A study of lectins from 52 Kalanchoe species cultivated in Kiev State University’s botanical garden in 1982 demonstrated a drastic difference in their virucide activity. For instance, K. daigremontianum broadly used in pharmacy did not exhibit virucide activity. In contrast, K. velutina, K. blossfeldiana, K. pinnata, and K. crenata demonstrated a high neutralizing activity in vitro towards nonenveloped RNA-containing viruses (vaccine poliovirus Sabin type II, Coxackie B1, and Coxackie B6). 50% reduction of the virus titer was achieved after incubation of these viruses with fresh juice of K. velutina diluted with cultural medium in ratio 1:32768, and K. pinnata, K. blossfeldiana, and K. crenata diluted 1:16384–1:8192 times. An enveloped virus A (Hong Kong) 1/68/H3N2 could be 50% inactivated by a juice of K. velutina in dilution 1:400, by juice of K. crenata—in dilution 1:200, and K. pinnata, K. blossfeldiana, and K. beharensis—in dilution 1:100 [20]. The rate of the virus neutralization did not depend on the temperature. Electronic microscopy demonstrated the contact of the viral particles with Kalanchoe juice induced their deformation and aggregation. Taken together, these observations may be explained by agglutination of the viral particles by Kalanchoe lectins or blocking their cell-specific receptors.

Hence, lectins may contribute to both antipathogenic and immunomodulatory activities of Kalanchoe extracts by binding carbohydrate moieties of extracellular receptors. Immunomodulatory functions may be mediated by nonproteinaceous components of these extracts as well. Costa et al. [25] reported inhibition of human lymphocyte proliferation with patuletin acetylhamnosides from K. brasiliensis. Da-Silva et al. [26] communicated about the induction of Th1-mediators (IL-2 and IFN-γ) and suppression of Th2-mediator IL-4 in mice after the administration of K. brasiliensis ethanol extract.

Recently, transgenic plants of K. pinnata producing antimicrobial peptide (AMP) cecropin P1 (CecP1) from Ascaris suis was described [27]. An antimicrobial, antifungal, and antiviral activity was attributed to this AMP [28]. However, data about biological trials of CecP1 towards human or animal microbial pathogens are scarce due to poor availability of the synthetic peptide and difficulty of its biogenic synthesis in bacteria or yeast which suffer from its toxicity [29].

This work pursues testing antifungal and wound healing activity of K. pinnata plants producing CecP1 towards highly prevalent human fungal pathogen Candida albicans on a model of infected wounds in rats. Taking into account the probability of existence of an intrinsic antifungal activity in K. pinnata juice (although not formerly reported), a juice of nontransgenic K. pinnata was tested in parallel as a negative control. Since CecP1 is not soluble in ethanol, a water extract of the plants was tested. In this respect, occurrence of lectins exhibiting lymphoproliferative and immunomodulatory properties along with bufadienolides and flavonoids in this type of preparations may be noteworthy. A commonly used commercial medical fungicide clotrimazole was used as a positive control (standard).

2. Methods

2.1. Producing K. pinnata Extracts. K. pinnata extracts were prepared form leaves of K. pinnata transgenic plants bearing a binary vector pBM-cecP1 for Agrobacterium tumefaciens vector with T-element randomly integrated to a plant genome. The vector was free from drug-resistance markers. The transgenic plants were selected after Agrobacterium-mediated transformation by a direct immunological testing exhibited a steady yield of CecP1 for at least two years as checked by three independent methods (immunoblotting, plate test for microbicide activity, and HPLC combined with mass-spectroscopy detection). 5.07 L of extract was produced from 3 kg of the recombinant plant leaves using deionized water as a solvent. After heating the fresh extract at +80°C and its sterilizing filtration through a nylon membrane with 0.22 μm pores, the extracts were adjusted to a total protein concentration ~1 mg/ml. Usually, they contained 0.69–0.82 μg/ml CecP1. The extracts of the wild-type (nonrecombinant) K. pinnata was produced by the same method and adjusted to the same concentration of the total protein.

2.2. Animals. In vivo trial was designed according to the European Convention about defense of the vertebrates used for experiments or for another scientific aims (Strasbourg, Mar. 18, 1986) of ETS N123. In total, 120 adult male Wistar rats weighing 180 ± 20.0 g 3–4 months old were allocated for the experiment. After quarantine, they were kept in individual cages. All animals were contained in equal terms on a standard diet and photoperiod (twelve hours of darkness and twelve hours of light). They had a ready access to water and food.
The animals were randomly divided into experimental, positive control, reference, and negative control groups (30 animals per each). All 120 animals after wounding were infected with C. albicans. The experimental group was treated with CecP1-enriched recombinant K. pinnata extract (K. pinnata + CecP1). The positive control group was treated with clotrimazole. The negative control group was subjected to mock treatment with a saline, and the reference group was treated with wild-type K. pinnata extract. Each group was divided into three echelons withdrawn from the experiment at 3th, 10th, and 14th day after the beginning of the curing.

2.3. Fungal Pathogen Strain. The human fungal pathogen C. albicans (NCTC 2625) formerly isolated from a human clinical specimens was purchased from a type strain collection of Tarasevich Research Institute for Standardization and Control of Medical Biological Preparations. C. albicans culture was obtained by cultivation on a slant meat-peptone nutrient agar supplemented with 1% glucose for at 37 °C for 18–20 h. The cells were washed from the slant agar with a sterile saline, thoroughly suspended, adjusted to a concentration ~10⁹ CFU per ml by using an optical turbidity standard CCA 42-28-29-85 and used for infecting the wounds in rats.

2.4. Surgical Manipulations, Treatment, and Planimetry Assay of the Wounds. A purulent infection of wounds was modeled in rats using a method described previously [30]. The animals were anesthetized with diethyl ether. A square model in rats using a method described previously [30]. Assay of the Wounds.

2.4. Surgical Manipulations, Treatment, and Planimetry

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The animals were examined daily, and stages of wound healing (inflammation, granulation, and maturation (marginal epithelization)) were fixed. The planimetric analysis of the wound recovery percentage was carried out at 3th, 10th, and 14th day after the beginning of curing. After this, one echelon (10 animals from each group) was withdrawn from the experiment. The animals were sacrificed with overdose of the ether anesthesia. Residual wound square was measured individually in each animal as described formerly [32], using an imprint on a transparent polyethylene sheet which was scanned at resolution 200 pcs/inch using a common office scanner. The images were acquired in a format Adobe Photoshop CS Extended. The object was selected using the abovementioned standard software, and its square was automatically calculated by selecting a menu command “Analysis.” An average mean and a standard deviation (M ± Std. Dev.) were calculated for each group/echelon (10 animals). The recovery percentage was evaluated with following formula:

Recovery percentage = (initial wound square – wound square on the day \( X \)) \times 100/(initial wound surface), where \( X \) = the day of wound square measurement (0 = beginning of the curing).

2.5. Preparing Histological Slides, Their Staining, and Analysis. After the completion of treatment, animals were sacrificed, and tissue samples were prepared as described previously [32]. The resulting biological material was fixed in 10% neutral formalin. After fixation, the tissues dissected for 1 × 1 cm fragments, washed, dehydrated, and impregnated with paraffin by standard methods. 5–7 μm thick microtome sections were stained with hematoxylin and eosin to assess the density of collagen fibers. A light microscope Leica CME with magnifications ∠40, ∠100, ∠200, and ∠400 and camera DCM-510 were used. Ten sections from each wound were investigated and evaluated. Image tools 3 software (UTHSCSA ImageTool) was used for accumulating and analyzing the images.

Microscopy and photographing the slides were performed using an optical system by Leica comprising an eyepiece camera and software for documenting images FUTURE WINJOE (supplied by the manufacturer of the hardware). Cell content nearby the wound edge and underlying tissue were examined at each micrograph. The cells were categorized into neutrophils, macrophages, mast cells, fibroblasts, and capillary endotheliocytes according to their morphological features (shape of the cell, shape of the nucleus, and cytoplasm/nucleus ratio) as described previously [33]. Regularity of the collagen fiber alignment was evaluated. The percentage of those representatives of a cell population was calculated after counting 100 cells in several nonoverlapping visual fields (at least 10).

2.6. Microbiological Analysis of the Wound Bed Microbial Contamination. App. 0.5g tissue (fibrous mass, infiltrate, and underlying derma) was sampled from each sacrificed animal under aseptic conditions, weighed at analytical grade balances (accuracy 0.1 mg), placed into a sterile porcelain mortar, mixed with a sterile saline in a weight ratio 1:10, and homogenized with a sterile pestle for 3 min at room temperature. The homogenate was diluted 1000 times with a sterile saline (with three consequent steps 1:10 using 1 mL samples) and 100 μl aliquot of each dilution was inoculated to Petri dishes with meat-peptone nutrient agar supplemented with 0.1% glucose. The inoculated dishes were incubated at 37 ± 1°C for 72 h and then 1 day more at a room.
temperature. The colonies were counted and number of CFU recalculated per 1 g tissue. The count was suggested to be valid if number of colonies was between 30 and 300 per dish.

2.7. Statistical Analysis. Statistical analysis was provided by Microsoft Excel 2007 and program “Statistics” 8.0 StatSoft. The average values and standard deviations of quantitative indexes obtained by planimetry and microbiological methods were calculated for each group/echelon (10 animals). The statistical significance of differences between the groups/echelons was estimated by the Mann–Whitney U test (p < 0.05).

3. Results

3.1. Study of the Wound Healing Activity by Planimetry. The wound healing process was divided into three partially overlapping stages: inflammation, granulation, and maturation. The stage of inflammation was considered to be finished once perifocal edema around the wound disappeared. Start of the granulation stage was registered once the wound bed began filling with a newly tissue of a pale pink color. Start of the maturation stage was registered once a marginal epithelialization became visible. Each animal was analyzed daily in order to reveal perifocal edema, granulation, and marginal epithelialization (Figure 1). Residual square of the wound was measured at 10th and 14th days after the beginning of curing (Figure 2). Significance of the difference between the groups is shown on Figure 3.

3.2. Histological Analysis of the Wound Healing Activity. Three days after the beginning of curing, the wound beds in all four groups were filled with purulent-necrotized cell mass (Figure 3). Leucocyte infiltration was visible deeply in the derma. An edema and plethora of dilated capillaries were found in the underlying tissues. The collagen fibers were disorganized and pushed away from each other due to interstitial edema.

Ten days after the beginning of curing, the negative control group exhibited rising of the inflammation that was manifested as an appearance of a polynuclear inflammatory infiltrate, composed of neutrophils, macrophages, and must cells (Figure 4).

This period was characterized with melting of the necrotic tissue, their removal, and partial emptying of the wound cavity. The interstitial edema was conserved beneath the wound cavity indicating an increased permeability of the capillaries. Single microabscesses were found in deep layers of the derma. In the same period, noninfected animals did not exhibit traits of the inflammation. The surface of the wound was covered by separate islands of new-formed thin epithelium growing from the margins of the wound. Layers of derma beneath the wound were impregnated with fibrin and contained single fibroblasts. A wound bed in the positive control group (clotrimazole) contained granulation focuses constituted from inflammatory and proliferative cells and enriched with newly collagen fibers. A thin epithelial layer without abnormalities appeared along the margins of the wound. The reference group (extract of the wild-type K. pinnata) contained much more collagen fibers and fibroblasts beneath the edge of the wound then the negative and positive control groups. Marginal epithelialization was more pronounced in the reference group. However, the inflammatory cell component was more abundant than in the positive control group. The animals of the experimental group (CecP1) were characterized with the fastest progression in the collagen matrix formation, its optimal structuration, the fastest marginal epithelialization, and the minimal number of the polynuclear inflammatory infiltrate in the wound bed.

Fourteen days after the beginning of curing, the wound cavity of the negative control group was still filled with polynuclear inflammatory infiltrate (Figure 5).

Only single spindle-shaped fibroblasts were found. The new-formed granulation tissue was located in the bottom of the wound bed. The inflammatory infiltration and the interstitial edema were conserved in the depth of the derma.
The singular thin collagen fibers were located irregularly. Most fibroblasts were clustered around them. A scar formation appeared as a solid layer of collagen fibers without vessels and skin appendages. The epidermis was much thicker than normal. The positive control group exhibited a similar picture although the collagen fibers were more...
abundant and less regular than in the negative control. The reference group (wild-type *K. pinnata* extract) did not contain solid collagen layer neither focuses; isolated inflammatory infiltrate focuses were conserved. However, overall structure of the collagen was more regular that in positive and negative control group. Revascularization of the scar was beginning. The experimental group combined positive traits of the positive control and the reference group. In this period, solid collagen layer completely covered the former wound cavity in this group. This layer was of the most regular, partially revascularized, and was completely free from the residual inflammation focuses.

3.3. Analysis of *C. albicans* Cell Survival In Vivo. Fungicide activity of the tested medicines was evaluated as a decrease of alive *C. albicans* cell number per g of the granulate tissue determined by microbiological method as described in Method (Figure 6). An evident fungicidal activity of *K. pinnata* extract in vivo was found, and it was substantially increased by CecP1. The load of yeast cells in the granulate tissue at the 3rd day upon a treatment with the experimental preparation was even less than in the positive control (clotrimazole). In contrast, at the 10th day, the fungicidal effect of clotrimazole was stronger than in the experimental *K. pinnata* extract with CecP1. In this period, distribution of the pathogen in the granulate tissue of the experimental group was much less homogeneous than the positive control group. At 14th day after the beginning of curing, *C. albicans* was eliminated over the most surface of the wound bed in the positive control and reference and experimental groups; however, single focuses of infection were conserved. In contrast, the infection evolved in the negative control group.

4. Conclusion

An evident fungicidal activity of *K. pinnata* against the most common human fungal pathogen *C. albicans* has been demonstrated for the first time. Moreover, the data of histological analysis make an evidence that *K. pinnata* exhibits a positive effect on remodeling of the collagen matrix and reepithelialization of the wound cavity along with its repairing. As specified in Introduction, *K. pinnata* is known with the production of bufadienolides and polyphenols which exhibited a toxic activity of animal cells in vitro. However, our results give show that these compounds do not compromise wound healing activity when administrated within the water extracts of *K. pinnata* leaves normalized by the total protein load 1 mg/ml. CecP1 within the recombinant *K. pinnata* extract

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**Figure 4:** Histological analysis of derma within the wound bed, day 10 after the beginning of curing (magnif. ×70). (a) Positive control (clotrimazole); (b) negative control (no specific treatment); (c) reference group (wild-type *K. pinnata* extract); (d) experimental group (CecP1). Selected area: arrow—new-formed thin epithelium, square—area of melting of the necrotic tissue.
**Figure 5**: Histological analysis of derma in the wound bed, day 14 after the beginning of curing (magnif. ×280). (a) Positive control (clotrimazole); (b) negative control (no specific treatment); (c) reference group (wild-type *K. pinnata* extract); (d) experimental group (CecP1). Selected area: arrow—fibroblasts, circle—collagen fibers, and double arrow—thickened epidermis.

**Figure 6**: Dynamics of the wound bed contamination with *C. albicans*. 

|                | Negative control | Reference group | Experimental group | Positive control |
|----------------|------------------|-----------------|--------------------|------------------|
| 3rd day        | 0.295 ± 0.09     | 0.139 ± 0.04    | 0.035 ± 0.01       | 0.011 ± 0.03     |
| 10th day       | 0.329 ± 0.14     | 0.138 ± 0.09    | 0.032 ± 0.05       | 0.0115 ± 0.03    |
| 14th day       | 0.1865 ± 0.08    | 0.1065 ± 0.05   | 0.0325 ± 0.03      | 0.0115 ± 0.03    |
substantially ameliorates its natural antifungal properties making it comparable to the commonly used commercial antifungal preparation. In a certain extent, this result is unexpected, since many formerly studied AMPs with α-helical secondary structure, on the one hand, exhibited unacceptable side-toxicity and, on the other hand, were unstable in vivo [34]. Contribution of immunomodulating activity of K. pinnata natural compounds to C. albicans elimination in vivo is confirmed by the fact that although CecP1 exhibits 80–300-fold lower molar activity against C. albicans in vitro than clotrimazole, CecP1-enriched extract of K. pinnata provides even more fast and complete elimination of the fungal pathogen from the infected wound than the commercial fungicide. Taken together, our data demonstrate a high perspective of pharmaceutical application of K. pinnata extracts containing CecP1 for curing C. albicans infection and possibly other fungal infections. This approach is technologically and economically affordable. However, an extensive toxicological trials and testing of rising resistance mechanisms in the fungal pathogens against the novel potential drug are required before it can be recommended for a practical implementation.

Conflicts of Interest
The authors declare no conflicts of interest.

Acknowledgments
This work is supported by a grant from the Ministry of Education and Science of the Russian Federation (Agreement no. RFMEFI60714X0016).

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