PAPER DETAILS

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ANURAN SPECIES FROM TURKEY

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PAGES: 43-52

ORIGINAL PDF URL: https://dergipark.org.tr/tr/download/article-file/912470

Dumlupınar Üniversitesi Fen Bilimleri Enstitüsü Dergisi E-ISSN: 2651-2769



Journal of Science and Technology of Dumlupinar University E-ISSN: 2651-2769

Number 43, December 2019

Sayı 43, Aralık 2019 ARAŞTIRMA MAKALESİ (Research Article)

ASSESSMENT OF ANTIMICROBIAL ACTIVITY ON THE SKIN SECRETIONS OF NINE ANURAN SPECIES FROM TURKEY

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Geliş Tarihi:25.09.2018

Kabul Tarihi: 03.07.2019

ABSTRACT

A dramatic decline of amphibians was reported for the first time about 40 years ago and then, this decline has continued at an alarming rate. Nearly one-third (32%), or 6593 amphibian species are threatened according to the most recent 2008 global assessment. Since 1980, the population size of at least 43% of amphibian species is declining and 122 amphibian species have become extinct. The causes of this declines are likely to be complex. Amphibians, like higher vertebrates, have immune systems including both adaptive and innate immunity. Innate immunity has different mechanisms that can provide an instant response nonspecifically to many pathogens. Antimicrobial peptides are the first step in these mechanisms. As each frog species has its own unique peptides, significant variation obtains among species in the number, structure and antimicrobial activity of these peptides. In our knowledge, there is no study previously performed in our country on the amphibian immune system. This study aims to evaluate the skin secretions of the nine anuran species and antimicrobial activity.

Keywords: Amphibian populations, decline causes, amphibian immune system, antimicrobial peptids

TÜRKİYE'DEN DOKUZ ANURA TÜRÜNÜN DERİ SEKRESYONLARININ ANTİMİKROBİYAL AKTİVİTESİNİN DEĞERLENDİRİLMESİ

ÖZ

İlk kez yaklaşık kırk yıl önce amfibilerde dramatik bir azalma rapor edilmiş ve bu düşüş endişe verici bir oranda devam etmiştir. 2008 yılı son küresel değerlendirmesine göre amfibilerin yaklaşık üçte biri (% 32) veya 6593 amfibi türü tehdit altındadır. Bu azalışın birçok karmaşık sebebi olabilir. Amfibiler omurgalılar da olduğu gibi hem doğuştan hem de kazanılmış immün sisteme sahip canlılardır. Doğal bağışıklık bireyin birçok farklı patojene karşı anlık tepki göstermesini sağlayan farklı mekanizmaları içerir. Bu mekanizmaları içerisinde antimikrobiyal peptidler ilk sırada yer alır. Her bir kurbağa türünün kendine özgü peptidleri olduğu için, bu peptidlerin sayısı, yapısı ve antimikrobiyal aktivitesindeki türler arasında önemli farklılıklar gösterir. Türkiye'de amfibi bağışıklık sistemi ile ilgili yapılmış

herhangi bir çalışma bulunmamaktadır. Bu çalışmada 9 amfibi türünden edilen deri sekresyonlarının antimikrobiyal aktivitelerinin belirlenmesi amaçlanmıştır.

Anahtar Kelimeler: Amfibi populasyonu, Azalma sebepleri, Amfibi immün sistemi, Antimikrobiyal peptidler

1. INTRODUCTION

Global declines of amphibian populations are a source of great concern. Many possible causes for amphibian population declines have been proposed [1]. According to researchers, deaths of amphibians are generally originate from three different factors. The first one of these factors is the abiotic factors and amphibians are interaction with this factor. People cause the change of ecosystems due to effect of abiotic factors consciously or unconsciously. The other factor is increase of number and virulance of pathogen microorganisms and organisms are adversely affected these microorganisms. Another factor is weakened immune system of organisms which live in ecosystems and organisms are under threat.

Amphibians are a transition species from the aquatic ancestors to the reptiles. Their skin is a morphologically, physiologically, and biochemically complex organ. Functions of amphibian skin are necessary for survival, including respiration, antipredator, water regulation, and antimicrobial defense, temperature control, excretion, and reproduction [2]. Their skin contains mucous and granular glands which responsible for the secretion of toxins and defences against microorganisms and predators [3,4]. The glands are usually activated by injury or stress and secrete a complex chemical mixture which vary from species to species [5, 6, 7, 8]. The skin glands of amphibians have various bioactive molecules such as peptides, steroids, proteins, alkaloids and biogenic amines that possess antibacterial and other important biological activities. According to their interactions with the environment, the compositions of the skin secretions differ among anuran groups [5, 7, 8]. These substances play different roles either in the defence against microorganisms and predators or in the regulation of physiological functions of the skin [9]. Besides, they are related to a variety of biological effects such as cytotoxic, fungicidal, bactericidal, lytic, anesthetic, neuromimetic, and phenomenal [4, 10, 11, 12, 13]. Recently, skin secretions were also noted as a rich source of multiple antimicrobial peptides effective against multidrug resistant strains[14]. Antimicrobial peptides were divided into two categories: those obtained from protein hydrolysis which called cryptides [15] and those obtained by ribosomal synthesis. Peptids which were obtained by ribosomal synthesis a significant part of innate immunity. Antimicrobial peptides constitute the first line of defense of the body [16, 17, 18]. Some antimicrobial peptides constitude by the proteolytic degradation of some proteins present within the organism. Studies on this subject is limited to a few species of frog in Turkey.

The purpose of present study is to test the antimicrobial activity of different anuran skin secretions against Gram (+), Gram (-) bacteria and *Candida albicans* cultures.

2. MATERIALS AND METHODS

2.1. Specimen Biodata and Secretion Harvesting

All the animal experiments were carried out in accordance with the approved guidelines with the regulations of the Turkish Department of Nature Conservation (Permit number, DKMP-

Aykıl vd., DPÜ Fen Bilimleri Enstitüsü Dergisi, Sayı 43, 43-52, Aralık 2019. Aykıl et all., Journal of Science and Technology of DPU, Number 43, 43-52, December 2019.

0.15.01.510.39719) and were approved by the Animal Welfare and Research Ethics Committee of Afyon Kocatepe University (permit number, AKU.0.A2.00.00-222). Adult specimens of frogs of both sexes (*Bombina bombina, Rana dalmatina, Rana macrocnemis, Bufo bufo, Bufotes variabilis, Bufo verrucosissimus, Pelodytes caucasicus, Pelophylax ridibundus, Pelophylax caralitanus*) obtain from different regions in Turkey (Fig1).



Figure 1 Turkey on the map of study areas, 1. Hyla orientalis (İğne ada (Kırklareli)) 2. Bombina bombina (Akgöl (Adapazarı)) 3. Bufo bufo (Kabaca köyü (Zonguldak)) 4. Pelodytes caucasicus (Trabzon Uzungöl)) 5. Rana dalmatina (Uzungöl (Trabzon)) 6. Bufo verrucosissimus (Uzungöl (Trabzon)) 7. Rana macrocnemis (Uzungöl (Trabzon)) 8. Bufotes variabilis (Sahara (Artvin)) 9. Pelophylax caralitanus (Eğirdir Gölü (Isparta)).

2.2. Preparation of Skin Secretion

The frogs were washed first with tap water for the first step in the experimentand then with distilled water. Skin secretion was taken from the dorsal skin with gentle transdermal electrical stimulation [19]. Secretions were washed from the skin using deionized water and collected solutions were held in 80°C water bath for 30 min and centrifuged at 5500 rev/min for 30 min. The precipitate which constitute in centrifugation was diluted with distilled water,0.1 N HCl, 0.1 N NH₄OH and 1 M phosphate buffers (pH: 4 and pH: 7)was used in the experiments [20].

2.3. Test Microorganisms and Growth Conditions

Test microorganisms (*Escherichia coli, Staphylococcus aureus, Pseudomonas aeruginosa, Listeria, Micrococcus luteus, Bacillus subtilis, Klebsiella pneumoniae, Salmonella thyphi, Proteus vulgaris, Bacillus cereus, Candida albicans*) were used from the culture collection of Afyon Kocatepe University, Faculty of Science and Art, Biology Department, Afyonkarahisar/Turkey. Cultures of these bacteria were grown in nutrient broth (NB) at 37°C for 24 h [21].

2.4. Determination of Antimicrobial Activity

Agar-Disc Diffusion Method were carried out *in vitro* antimicrobial activity studies. For antimicrobial activity Nutrient agar (NA) was preferred as the most suitable medium. 20 µl extract was inoculated into a sterile 6 mm diameter disc [22]. The turbidity of bacterial suspension was adjusted according to Mcfarland Standard Tube (0.5) with physiologic serum and suspension of the tested microorganism was spread on the solid media plates. Filter paper discs placed on the inoculated plates. These plates,

were incubated at 37°C for 24 h for bacteria and, at 30°C for 24 h, for yeasts [23]. The diameters of the inhibition zones were measured in millimetres. All tests were performed in duplicate. In addition, the solvent was used as negative control. 0.1 N HCl, 0.1 N NH₄OH, 1 M phosphate buffers (pH: 4), 1 M phosphate buffers (pH:7) were used as positive controls. Experiments were repeated two times and results were evaluated as average values.

3. RESULTS

In this study, antimicrobial effects of prepared extracts on the tested microorganisms were determined by using different solvents. According to our findings, all the extracts were obtained from diffrent amphibians, exhibit antimicrobial activity effects of different extracts were obtained from the skin secretions against bacteria and yeast cultures results are given in Table 1. Adult specimens of frogs of both sexes obtain from different regions in Turkey (Table 2).

In our results show that, The highest antibacterial effect showed by 0.1 N NH₄OH of *Bufotes* variabilis extract against *Micrococcus luteus* (ATCC-9341). Extracts of *Bufo bufo* skin secretion exhibited no effects against *Pseudomonas aeruginosa* and *Micrococcus luteus*. However the highest antibacterial effect showed against *E.coli* (10,5-14mm inhibition zone). The more antibacterial effect showed by 0.1 N HCL and 1M phosphate buffers (pH 7) extracts of *Bufo variabilis* than 0.1 N NH₄OH extract and 1M phosphate buffers (pH 4). According to our findings, the highest antimicrobial activity was observed by 0.1 N HCI extract of *Rana dalmatina* and 1M phosphate buffers (pH 7) extracts of *Rana macrocnemis* against *Klepsiella pneumonia*.

 Table 1 Antimicrobial activity effects of different extracts were obtained from the skin secretions against bacteria and yeast cultures.

Frog Samples	Test strains	S. typhimurium	K. pneumoniae	E. coli	P. vulgaris	P. aeruginosa	B. cereus	B. subtilis	S. aureus	M. luteus	L. monoaytopenus	C. albicans
Destriction	Sulbactam/ampicillin	9	25	6	9	20	12	13	10	17	10	6
Positive control	AM/Amikacin	0	6	6	0	6	20	22	10	6	6	0
		6	6	10	6	6	6	23	13	11	6	6
	0.1 N NH.OH	6	6	10	6	6	6	7	6	6	6	6
	1 M phosphate buffers	6	6	10	6	6	6	7	6	6	6	6
Negative control	(nH·4)	0	0	10	0	0	0	/	0	0	0	0
	1 M phosphate buffers	6	6	8	6	6	18	7	14	20	6	6
	(pH:7)											
	0.1 N HCl	13	13	12	12	10.5	11	11	11.5	12.5	9.5	12
	0.1 N NH ₄ OH	12.5	12.5	10.5	11	11	10	10	11	12	10	6
Bombina	1 M phosphate buffers	17	13.5	11	14.5	13	11.5	11	13	14	9.5	6
Dombina	(pH: 4) 1 M phosphate buffers (pH:7)	12.5	14	10	12.5	11.5	9.5	10	13.5	10.5	9.5	6

	0.1 N HCl	9.5	14.5	11	13	11	10	9.5	10.5	12	10.5	12
	0.1 N NH ₄ OH	6	6	11	6	10	6	6	6	6	6	9
Rana dalmatina	1 M phosphate buffers (pH: 4)	6	6	11	6	6	6	6	6	6	9.5	6
	1 M phosphate buffers (pH:7)	7.5	14	10.5	10	9.5	9	10.5	10	6	11.5	8.5
	0.1 N HCl	14	13.5	12	13.5	13.5	10.5	10.5	14.5	11	11	10.5
	0.1 N NH4OH	12	13.5	13	12.5	6	14	11.5	13.5	12	7.5	6
Rana	1 M phosphate buffers	9	10	9.5	6	6	11	6	9.5	6	6	6
macrocnemis	(pH: 4)					-		-		-		
	1 M phosphate buffers (pH:7)	12.5	14.5	11	12	13.5	8	9.5	9.5	6	8.5	6
	0.1 N HCl	11.5	16	14	10.5	6	9	9	9	6	8.5	6
	0.1 N NH ₄ OH	9	12	11	9	6	6.5	10	7.5	6	8	6.5
Bufo bufo	1 M phosphate buffers (pH: 4)	6	6	11.5	6	6	8	6.5	6	6	6	8.5
	1 M phosphate buffers (pH:7)	9	6	10.5	9.5	-	8.5	8.5	8	6	9	9
	0.1 N HCl	11	11	10.5	11	12.5	11	11	9.5	15.5	9	12.5
	0.1 N NH ₄ OH	6	6	9	6	8.5	6	6	6	6	6	6
Bufotes variabilis	1 M phosphate buffers (pH: 4)	6	6	10	6	6	6	6	6	6	7.5	9
	1 M phosphate buffers (pH:7)	10	6	8.5	10	10	9	9	11.5	6	8	6
	0.1 N HCl	8.5	11.5	11	12	12	9.5	9.5	10.5	10.5	12	11
	0.1 N NH₄OH	6	8	13.5	6	6	6	6	6	6	6	6
Bufo	1 M phosphate buffers (pH: 4)	6	6	11.5	6	6	6	6	6	6	6	6
verrucosissimus	1 M phosphate buffers (pH:7)	8.5	8	8	11	10	9.5	7.5	9.5	6	10	6
	0.1 N HCl	9	11	10	11.5	10.5	9.5	10	10	11	10.5	10
	0.1 N NH₄OH	6	6	11.5	6	6	6	6	6	6	6	7
Pelodytes	1 M phosphate buffers	6	6	12	6	6	9	6	6	6	11	6
caucasicus	(pH: 4)											
	1 M phosphate buffers (pH:7)	9.5	9.5	11.5	10	8.5	8.5	10	9.5	10	9.5	6.5
	0.1 N HCl	11	6	6	10	6	8.5	8.5	9.5	7	12	6
	0.1 N NH ₄ OH	10	9	6	10	6	12	10	11	13.5	11	6
Pelophylax ridibundus	1 M phosphate buffers (pH: 4)	12	9	8	11.5	6	15	13.5	11.5	10.5	11	6.5
	1 M phosphate buffers (pH:7)	11	8	6	9.5	6	11	11	10.5	8.5	8	6
	0.1 N HCl	9.5	10	11	11.5	12	9	9	9	12.5	10	11
	0.1 N NH ₄ OH	6	9	12.5	8	6	6	10	6	8.5	6	6
Pelophylax caralitanus	1 M phosphate buffers (pH: 4)	6	6.5	12.5	6	6	6	8.5	6	6	7.5	6
	1 M phosphate buffers (pH:7)	10.5	6	10.5	11	10.5	10.5	9.5	11	10.5	9	6

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No	Species Region Coordina Altit		Altitude		Average					
					1	2	3	4	5	
1	Hyla orientalis	İğne	41 11 13	39	36.88	37.6	-	-	-	37.24
		ada(Kırklareli)	40 59 19							57.24
2	Bombina	Ak lake	40 52 36	39	37.3	37.6	36.34	-	-	37 08
	bombina	(Adapazarı)	30 26 02							57.08
3	Bufo bufo	Kabaca köyü (Zonguldak)	41 17 53	482	117.04	-	-	-	-	117.04
1	Pelodytes	Uzun lake	40 37 10	1164	52.28	17	123	13.26	11.6	15 888
-	caucasicus	(Trabzon)	40 17 27		52.20		72.5	т <i>3.2</i> 0	0	+J.000
5	Rana	Uzun lake l	40 35 27	1272	32	-	-	-		32
	dalmatina	(Trabzon)	40 20 45							
(Bufo	Uzun lake	40 35 27	1272	046	60.24	60.20	60 60	66	72 500
0	verrucosissimus	(Trabzon)	40 20 45		94.0	09.54	09.29	08.08	00	15.362
-	Rana	Uzun lake	40 34 38	1701	72.1					72 1
/	macrocnemis	(Trabzon)	40 23 47		/3.1	-	-	-	-	/3.1
8	Bufotes variabilis		41 13 47	1876	876 71.88	-	-	-	-	71.00
		Sanara (Artvin)	42 27 05							/1.88
9	Pelophylax	Eğirdir lake	38 12 44	923	132	89.82	60.6	84.48	90.5	01.40
	caralitanus	(Isparta)	30 45 05							91.48

Table 2 Adult specimens of frogs of both sexes obtain from different regions in Turkey.

4. DISCUSSION

The skin of amphibians is an extraordinarily rich source of antimicrobial peptides (AMPs). The first AMP was detected in the skin of the European frog *Bombina variegata* 42 years ago [24]. Since that time, several hundreds of different antimicrobial peptides have been obtained from the skin of amphibians belonging to the families *Bombinatoridae*, *Hylidae*, *Hyperoliidae*, *Leiopelmatidae*, *Leptodactylidae*, *Myobatrachidae*, *Pipidae*, and *Ranidae* [28]. AMPs belong to a large group of linear amphipathic helical peptide which are cationic, containing a flexible number of positively charged residues and hydrophobic regions. These properties supply them with an ability to bind to membrane lipids and/or negatively charged molecules and disrupt the membrane structure (Rollins-Smith et al. 2005).

In general, the antimicrobial activity of frog skin peptides is determined with some bacterial and fungal strains. These include the Gram-negative *Escherichia coli* and *Pseudomonas aeruginosa*, the Gram-positive *Staphylococcus aureus*, and the yeast *Candida albicans* [24].

The antimicrobial activity of bombinin-like peptides or bombinins H obtain from *Bombina sp.* can be distinguished on the basis of their cytolytic properties. Bombinins were found to be active against Gram-positive [G(+)] (*Bacillus megaterium, Staphylococcus aureus*) and Gram-negative [G(-)] (*Escherichia coli, Yersinia pseudotuberculosis, Pseudomonas aeruginosa*) bacteria as well as against *Candida albicans* [25]. According to our findings, skin secretion from *Bombina bombina* was

observed antimicrobial activities against both G(-) and G(+) bacteria. The highest antibacterial effect showed by 1M phosphate buffer (pH4) of *Bombina bombina* skin extract against *Proteus vulgaris*, *Salmonella typhimurium* (NRRLB-4420) and *Bacillus cereus* (ATCC-11778). Only 0.1N HCI extracts of skin secretions showed antiyeast effects. The results of our study are similar to the other literatures.

Dülger et al. [26] investigated antimicrobial activity of skin secretions from *Bufo viridis*. In this study, antimicrobial activity was determined with *Escherichia coli*, *Staphylococcus aureus*, *Klebsiella pneumoniae*, *Pseudomonas aeruginosa*, *Proteus vulgaris*, *Bacillus cereus*, *Mycobacterium smegmatis*, *Listeria monocytogenes*, *Micrococcus luteus*, *Candida albicans*, *Rhodotorula rubra*, and *Kluyveromyces fragilis*. Park et al. [27] report a novel antimicrobial peptide named buforin I and II purified from the stomach of *Bufo bufo gargarizans*, an Asian toad, which has been used as a woundhealing agent in traditional Korean medicine. Both buforin I and buforin II displayed strong antimicrobial activities against a broad spectrum of bacteria including, *Bacillus subtilis*, *Staphylococcus aureus*, *Streptococcus mutans*, *Streptococcus pneumoniae*, *Escherichia coli*, *Serratia* sp., *Pseudomonas putida*, and *Salmonella typhimurium*. Furthermore, *Candida albicans*, *Saccharomyces cerevisiae* and *Cryptococcus neoformance* were also killed.

In our results show that, the highest antibacterial effect was observed by $0.1 \text{ N NH}_4\text{OH}$ of *Bufotes variabilis* extract against *Micrococcus luteus* (ATCC-9341). Extracts of *Bufo bufo* skin secretion exhibited no effects against *Pseudomonas aeruginosa* and *Micrococcus luteus*. However the highest antibacterial effect showed against *E.coli* (10,5-14mm inhibition zone). The more antibacterial effect showed by 0.1 N HCL and 1M phosphate buffers (pH7) extracts of *Bufotes variabilis* than 0.1 N NH₄OH extract and 1M phosphate buffers (pH4).

Frogs which belong to family Ranidae represent a important source of antimicrobial peptides [28]. Çevikbas [29] reported that skin secretion of *Rana ridibunda* determines antibacterial activity at different levels. However, Afsar et al. [20] showed that, skin secretions of *Rana macrocnemis* against the yeast cultures show more antimicrobial activity than that of the bacterial cultures. According to our results, the highest antimicrobial activity was observed by 0.1 N HCI extract of *Rana dalmatina* and 1M phosphate buffers (pH7) extracts of *Rana macrocnemis* against *Klepsiella pneumoniae*. The highest antibacterial effect showed by 1M phosphate buffers (pH 7) *Rana dalmatina* extract against *Listeria monoaytopenus* (ATCC-7644); 0.1 N NH₄OH and 1M phosphate buffer (pH 7) of *Rana macrocnemis* skin extract against *Pseudomonas aeruginosa*. The highest antimicrobial activity was observed by 0.1 N HCI extract against *Bacillus subtilis* (NRS-744). Sensitivity of the microorganisms to the different agents changes from strain to strain [30]. According to our results, all the extracts of skin secretion, were obtained from diffrent anurans, exhibit antimicrobial activity. The present study has demonstrated that antimicrobial activity of skin secretions of varries at both the generic and ecological.

In conclusion, amphibians are the first group of organism forming a connecting link between land and water. Moreover, they have to fight to adopt a different conditions of environment and laden with pathogenic microbes. Therefore, theywere endowed with an excellent chemical defense system composed of pharmacological and antimicrobial peptides [31]. New peptides have been found that could inspire the design of substance prevent or treat infections. Peptide-based antibiotics are considered a potential solution to the growing problem of resistance to traditional antibiotics. In the development of AMPs as therapeutics, peptides obtained from amphibian skin and their synthetic analogues will play a very important role.

ACKNOWLEDGMENTS

This research was supported by Project no. Tubitak 113Z139. Ethical endorsement was ratified by the Ethical Committee of Afyon Kocatepe University and the Turkish Department of Nature Conservation (Permit number, DKMP-51039719).

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