RANA RİDİBUNDA DERİ SEKRESYONU : ANTİFUNGAL ETKİSİ, KÜLTÜRDE YETİŞTİRİLEN RATTUS NORVEGİCUS EMBRYO FİBROBLAST VE KİMYASAL OLUŞTURULMUŞ TÜMOR HÜCRELERİNİN DNA SENTEZİNE VE MORFOLOJİSİNE ETKİSİ VE SAFLAŞTIRILMASI

THE RANA RIDIBUNDA SKIN SECRETION: ITS ANTIFUNGAL EFFECT, ITS EFFECT ON MORPHOLOGY AND ON DNA SYNTHESIS OF THE NORMAL EMBRYONIC FIBROBLAST CELLS AND CHEMICALLY INDUCED TUMOR CELLS OF RATTUS NORVEGICUS IN CULTURE AND ITS PURIFICATION

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SUMMARY

The extract (SSE) obtained from the skin secretion of the water frog Rana ridibunda, was purified by 95% cut $(NH_4)_2SO_4$ precipitation, Sephadax G25 or G75 and DEAE hephadex A25 column chromatographies. All of the biological activities were peresent in the II. protein fraction of Sephadex G75 column chromatography. Antibacterial, antifungal, INA synthesis inhibitor activities were present in the I. protein fraction of DEAE hephadex A25 column chromatography whereas the II. protein fraction had glycoprotein attacture with fibrinolytic system inhibitor activity.

Hendes the antibacterial activity of the extract, the antifungal activity against 11 funging that (Candida albicans 7650, C. parapsilosis KUEN 1010 (Y) $\rm C_1$ –12–1, C. krusei KUEN 1001 (Y) $\rm C_1$ –6–3, C. albicans klaur A–2, C. albicans, C.tropicalis KUEN 1022 (Y) $\rm C_1$ –19–3, the latest activity against XUEN 1018 (Y) $\rm C_1$ –18–1, C. albicans MIV–211, Cryptococcus neoformans MUEN 1047 (Y) $\rm C_2$ –1–9, Rhodotorula glutinis KUEN 1064 (Y) $\rm R_1$ –1–1, Saccharomycea materials, dermatofites Microsporum nanum KUEN 1089 (F) $\rm M_5$ –6–1, Trichophyton mantagrophytes 703.) was also detected. This extract was ineffective against Torulopsis labrata. A short review of this topic is also included.

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The effect of SSE on ³H-thymidine incorporation into Rattus norvegicus embryonic fibroblast cells (REF) and tumor cells (RT) in tissue culture was followed by autoradiography and morphological alterations caused by SSE were followed by light microscopy.

In tissue culture, in the medium with 5% SSE after 17 hrs of incubation, morphological alterations in REF and RT were observed. After 39 hrs of incubation at 10% SSE, the DNA synthesis was inhibited in both REF and RT. In 5% SSE containing medium after 39 hrs of incubation, ³H-thymidine incorporation into RT was 9.02 % whereas in control group (without SSE) this ratio was 36.62 %. The ³H-thymidine incorporation into REF after 100 hrs of incubation in a medium with 5 % SSE was 6.05 % whereas this ratio was 40.10 % in the REF cells grown in a medium without SSE. The DNA synthesis inhibitor effect of 850 was abolished when the cells, both REF and RT, were transferred into a medium without SSE. The time of recovery was found to be directly proportional to the time of preincubation of the cells in the medium with SSE.

ÖZET

Su kurbağası Rana ridibunda'nın derisinden salgılanan ekstre (KDS), % 95 (NH₄) presipitasyonu, Sephadex G25 veya G75 ve DEAE, Sephadex A 25 kolon kromatografilm sonucu saflaştırılmıştır. Sephadex G75 den elde edilen 2 protein fraksiyonundan birinde tüm biyolojik aktiviteler saptanmıştır. DEAE Sephadex A25 kolon kromatografisinden edilen I. protein fraksiyonundan antibakteriyel (AA) antifungal (AFA) ve DNA sonten inhibisyon (DNA–SIA) aktiviteleri görülürken II. protein de glikoprotein yapısı ve fibrinolotik sistemi inhibe edici aktivite saptanmıştır.

Ekstrenin antibakteriyal etkisi yanında 11 maya suşuna (Candida albicans 7650, parapsilosis KUEN 1010 (Y) $\rm C_1$ –12–1, C. krusei KUEN 1001 (Y) $\rm C_1$ –6–3, C. albicans klaus A–2, C. tropicalis KUEN 1022 (Y) 1–19–3, C. Stellatoidea KUEN 1018 (Y) $\rm C_1$ –18–13, C albicans MIV–211, Cryptococcus neoformans KUEN 1047 (Y) $\rm C_2$ –1–9, Rhodotorula glutinis KUEN 1064 (Y) $\rm R_1$ –1–1, Saccharomycea cerevisiae, dermatofitlerden Microsporum namus KUEN 1089 (F) $\rm M_5$ –6–1, Trichophyton mentagrophytes 703) antifungal etkisi saptanınıştır.

Rattus norvegicus embriyo fibroblast hücrelerine ve Rattus norvegicus tümür hücrelerine doku kültüründe ³H-timidin inkorporasyonuna otoradyografi yöntemi lib bakılmış morfolojik değişimler ise ışık mikroskobu ile takip edilmiştir.

Doku kültüründe % 5 KDS li mediumda inkube edilen hücrelerde 17 saatten sonra bası morfolojik değişiklikler izlenmiştir. % 10 yoğunlukta KDS lı besi ortamında 39 saat inkube edilen hücrelerde DNA sentezi her iki hücre grubunda inhibe olmaktadır.

Doku kültüründe % 5 yoğunlukta ki KDS li besi ortamında 39 saat inkube olan III hücrelerinde DNA sentezi % 9.02 iken kontrol tümor hücrelerinde DNA sentezi % 30.03 deney REF hücrelerinde DNA sentezi % 6.05, kontrol REF hücrelerinde % 40.10 oranında bulunmuştur.

Kurbağa deri sekresyonunun DNA sentezi inhibitor etkisi hücrelerin yeni besi artamına konmasıyla kaybolmuştur. Preinkubasyon zamanına bağlı olarak hücrelerin DNA sentezi normale dönmüştür. Bu konunun kısa derlemeside de yapılmıştır.

INTRODUCTION

One of the most typical characterictics of animals is their capacity adopt to physical, chemical and biological conditions in their invironments. Although microorganisms are abundant throughout the nature, the interrelationships between animals and microorganisms under natural conditions are little investigated. Of course, the animals of monomical and medical importances are excluded in this generalization. This subject—matter includes many interesting topics of physiological and evolutionary importance. Extracts or substances from animals manufactured mammals) with biological activities like antitumoral, antimicrobial, anticoagulant etc. are listed on Table I.

Frogs could survive in dirty places with microorganisms as well as a clean water or land. Since they have nude skin, their skin could be sailly scratched when they are in motion. Their skin is covered with a thin mucous layer. Infection on their skin is almost never detected. This fact is probably do to the presence of antimicrobial substances on their skin or in their skin secretions. We have reported earlier the isolation of an extract with antibacterial activity from the frog Rana ridibunda (16). In our preliminary studies, fibrinolytic system inhibitor activity was also detected in this extract. In this study, this extract isolated from the skin corotion of the water frog Rana ridibunda is investigated for its antifungal activity and for its effects on Rattus norvegicus embryonic fibroblast cells (REFC) and chemically induced tumor cells (RTC) grown the substances with the above listed biological activities which were found to be proteins is antablished.

MARMARA ÜNİVERSİTESİ ECZACILIK DERGİSİ Table - I

Animals Group, Class	Origin	Biological Activity	Active Substance	Refe:
PROTOZOA Flagellata Gongaulax catenella (Dinoflagellata)	Crude extract	Neurotoxin	Saxitoxin I (STX)	12 38
Gongaulax tamarensis	Crude extract	Neurotoxin	Saxitoxin I (STX)	2 33
METAZOA Coelenterata Physalia physalis (Jelly fish)	nematocysts' venom	Proteolytic enzyme activitiy	Toxin	48
Stomolophus meleagris (Jelly fish)	nematocysts' venom	5' nucleotidase, hyaluronidase, phosphatase, phosphodiesterase, leucine aminopeptidase and protease activities	Toxin	88
Chironex fleckeri (Australian Jelly fish) Chrysaora quinquecurrha (Jelly fish) Physalia physalis (Jelly fish)	whole body extract	rises blood pressure, hemolytic, dermonecrotic and lethal effects	The state of the s	BB
Chironex fleckeri (Jelly fish) Chry guinquecirrha (Jelly fish)	Tentacles, nematocysts suspensions	ionic calcium uptake inhibitor	Toxin	11
Actinia equina	Tentacles' extract	(lethal)	eguinatoxin (Highly basic thermolabile protein)	26
Actina equina	Tentacles' extract	lethal, causes bradycardia	eguinatoxin Protein	81
Sea anemones	whole body extract	distriby a school day	Serotonin, homarine	66
Anemonia sulcate (Sea anemone)	whole body extract	Increases Na ⁺ /K ⁺ ATP ase activity.	Polypeptide toxins ATXI (Mr:4702) ATXII (Mr:4935) ATXIII (Mr:2678)	1

Animals Group, Class	Origin	Biological Activity	Active Substance	Refe-
Palythoa vestitus (Boft coral)	whole body extract	One of the most potent coronary vasoconstrictors known highly toxic	Polypeptide High Mr polypeptide eg. the deadly botulinus toxin	12 97
COLEMATA I Echinodermata Asteroidea: Asterias rubens (Bea star)	whole body extract	Haemolytic activity	Saponin like substances	49
Asteroids (flea stars)	whole body extract		Saponin and glycosides mostly linked to sterols	12
Asterias (Boa star)	whole body extract	Antimicrobial activity	Glycones	83
Asterias (Sea star)	whole body extract	Antimicrobial activity	Glycones, a glycones as cholesterol or pregnenolon type structure	37
Marthanterian graetalin	whole body extract	enit aman CACA There is been a marked and a marked a mar	Saponin like structures Asterosaponins (glycosidic sulphates) Aglycones (Steroidal saponin)	49 93
Helethuridea : Helethurians Pez cucumbers)	whole body extract	man of the A	Saponin and glycosides mostly linked to sterols	12
Hidothuria sp.	whole body extract	Antimicrobial and antitumoral activity	Glycones (glycosides)	83
H MOLLUSCA Fastropoda i Han marine Han marine	whole body extract	RBC agglutination (A, B, A + B, H erythrocyte haemaglutinin)	Aglutinin A, B, A+B, H blood group substances	65
Apinia valifornica lika hares)	Hemolymph	Opsonin activity a mechanism of bacterial clearance by the sea hare involving aglutinin, an opsonin activity.	Aglutinin, lack bactericidine	43 90 60

Animals Group, Class	Origin	Biological Activity	Active Substance	Hofe
Aplysia california (Sea hares)	Serum	Bacterial clearance, immune defense mechanism and agglutinating marine bacterial and vertebrate RBC	Bacterial aglutinin (Protein)	01
errio line al DSA TUS del del menero del consensado del consensado en la c	agis in in sodianyos voicin	Water-soluble toxin: produces a transient hypotension bradycardia and	dw o Yosha kashin	
Aplysiidae (Sea hares)	0	apnoea, convulsion, respiratory distress. Ether-soluble toxin: hypertension in rats, vasoconinstrictor action, irritability vicious and severe flaccid paralysis.	Water-soluble toxin Ether soluble toxin	00
ya keralaa		Inhibitor of aglutination of B erythrocytes	B substances	90
Helix hortansis (Garden snail)	whole body	RBC aglutination (B and H erythrocyte haemaglutinin).	Aglutinin (B, H substances)	64
Helix hortansis (Garden snail)		RBC aglutination (B erythrocyte haema- glutinin)	Aglutinin (B blood group substances)	76 45
Patalla milanta	whole body extract	RBC Aglutination (A ₁ , A ₂ and B erythrocyte haemaglutinin)	Aglutinin A ₁ , A ₂ , B blood group substances	64
Cassidaria echinophora	Buccal gland	Stimulates immune defense mechanism	Sulphur rich compounds (Low molecular weight)	96
Thais haemostoma (Clench) Sea snail	Hypobranchial gland extract	1st active component; produced a direct stimulatory effect on the blood pressure and heart actions. 2nd active component; as neuromuscular blocking agent depolarizing type.	2 active components	40

Animals Group,	Origin	Biological Activity	Active Substance	Refe- rences
Csphalopoda : Octopus bimaculates	Serum	Aglutination	A blood group substance	70
Octopus vulgaris	Posterior salivary glands	Neurotransmitter	Octopamine, p- hydroxyphenyl, Ethanol amine	44
Osphalopoda sps. 11053	whole body extract	Pain reducer effective on romatoid arthritis	Enteramine, Histamine Octopamine Tyramine	44
Lamellibranchiata : Oystor	whole body extract	RBC Aglutination (A and H erythrocyte haemaglutinin)	Aglutinin (A and H blood group substances)	15 89
Crassostrea virginica inyster)	whole body extract	Aglutination	Aglutinin (Protein)	89 1 100
Pinetada martensi (parl oyster)	whole body extract	Aglutination	Aglutinin A and H blood group substances	102
Velesunia ambiquas (mussel)	Hemolymph	Aglutination	H and P ₁ blood group substances.	30, 65 41
Mytilus valifornionus (California sea mussel)	whole body extract	Neurotoxin	Saxitoxin I (STX)*	33
HLARTHROPODA (Trustaceae : Panulirus argus (aping labster)	Hemolymph	Antibacterial activity	Bactericidin	23
Homorus americanus (Labster)	Hemolymph	Antibacterial activity	precipitin	84
Homorus vulgaris (Labster)	Digestive juice	secreta the secretary	Surface–active substances (Fatty acyl dipeptide)	39

^{**}Marka butter clam saxidomus gigantus, both feed on G. catenella

Animals Group, Class	Origin	Biological Activity	Active Substance	Refer
Crangon crangon Pandalus jordani	Hemolymph	Stimulated succinate oxidation in intact rat liver mitochondria in metabolic state 4	Eyes talk factor	89 84
Procamborus bicarinatus (crayfish)	Hemolymph	Antibacterial activity	opsonic factors	67
Procamborus clarkii (crayfish)	Serum	Aglutinates marines bacteria, chicken and rabbit RBC	Aglutinin (Protein)	56
Astacus leptodactylus (cray fish)	Digestive juices Hemolymph	Surface active substances RBC Aglutination	Fatty acyl-tauirine Aglutinin	30 17
Carsinus mediterraneus (crab)	Hemolymph	Fibrinolytic system activator-plasmin-like	Protein	98
L. polyphemis (Horses shoe crab)	Hemolymph	Aglutination	Aglutinin Protein Mr: 150,000, heat and pH sensitive.	56
Insecta : Dytiscus	pigidal gland	Antibacterial activity		50
Selenopis geminata	venom	Antibacterial activity	of a	77
Periplanata	chitin	Antibacterial activity	Amino-phenol	9
Atta sexdens	venom	Antibacterial activity	Anteropolitica	51
Acheta domesticus	gut contents	a microsomal enzyme inhibitor. (Inhibition apparently results from solubilization of NADPH Cytochrom C reductase from microsomal membranes)	Proteolytic enzme	10
Coccionella	Hemolymph	Stain and indicator	Carmic acid	76
Vespa orientalis	Venom	Venom acts directly on mitochondrial function and ultra- structure of the mice kidney.	melatus R supersoqueso stin I (STX) is also is desko butter clera za	73

Animals Group, Class	Origin	Biological Activity	Active Substance	Refe- rences
Boos	Venom	Antiarthritic action increases corticosteron secretion	Protein phospholipase A and hyaluronidase, mellitin and apamine (mast cell degranulating peptide) (MCD)	2 80
Carabidae	Venom	Defense mechanism	Formic acid, iso valeril aldehyde, asetic acid, tigric acid.	78
Notonecta	secretion	Antibacterial activity	p.hydroxibenzal- dehyde	76
Tenebrio molitor	whole body extract	Antibacterial activity		76
Forficula	whole body extract	Antibacterial activity	o sarigeada L. Shin	Post Edwa
Julus terrestris	whole body extract	Antibacterial activity	Tolohydrochion, Athydrochion	79
Plealeachi	Metatorax gland	Antibacterial activity	$\mathrm{H_2O_2}$	Aghan A
Araneldea: Androcteurus Australis hektor	venom	Neurotoxin paralyzes mammalia and insects	ale: Recoonder the second seco	71
Latrodectus sps.	venom on	the release of	Alternation Alternation Alternation Alternation Alternation Action Action and	71
Latradectus mactans	venom	Neurologic symptom in muscle	Alisempaikudikase (Sivretiva forlank	71
Loxonceles reclus	venom	Coagulating mechanism activator	ntale Chamadomicana new Maranoballoso	98
IV, VERMES Polymera (analida) spinculoidea: Npunculid worms	whole body extract	Antibacterial activity	powerful bactericidin facto	

Animals Group, Class	Origin	Biological Activity	Active Substance	Refe:
Dendrobaena venata	whole body extract	Antitumoral activity on the sarcoma tumor in vivo		18
Ascaris lumbricoides	whole body extract	Antimicrobial and toxic activity		76
Amera (Scolecida): Plathelminthes Turbellaria Trematodes Cestodes	whole body extract	Antibacterial and cytotoxic activity	Acid mucopoly saccharid	76
Hirudo medicinalis	gland	Anticoagulant	Hirudin	76
FROGS AMPHIBIA	Skin secretion	Antimicrobial activity	Antimicrobial substances	62, 63
Bombinavarigeata L.	Skin secretion	Antimicrobial activity Bufo toxins can be differentiated from other anura secretions by the content of cardio active steroids	Protein, basic polypeptides, aminoacid and 5- hydroxytriptamine	46
B. varigeata	Skin gland secretion	Bactericidal activity	Two nonpeptides, low Mr peptides	5 37
B. varigeata L.	Skin gland secretion	Bactericidal activity	Two oligopeptides	14
B. varigeata varigeata	Skin gland secretion	Bactericidal activity Bombesin causes contraction on the uterine and intestinal smooth muscles of several animals, increases arterial blood pressure and stimulates gastric secretion in dogs and chickens.	Bombesin	53
. varigeata pachpus	Skin gland secretion	uterine and intestinal smooth muscles of several animals, incre ase, arterial blood pressure and	Active fraction does not show any typical u.v. absorption band as does material containing aromatic groups and does not react with ninhydrine.	13

Animals Group, Class	Origin	Biological Activity	INCOLVE	Refe- rences
B. varigeata	Skin secretion	Antimicrobial and some biologic activity. It causes systemic hypertension, bradycardia, constriction of the renal mesenteric and cardiac activity.	Bombesin (Tetradecapeptide)	27
Bombina bombina	Skin secretion	Antimicrobial activity	Low Mr peptides	37
Salamandra sp	Skin secretion	Antimicrobial activity venoms are active to microorganisms and their use is to protect the skin of amphibia against infection.	Azosteroids	8 34 37
Salamandra maculasa	Skin gland secretions	1 the membrane	Alkaloids Samandarone Samandarine Samondaridine	35 36 68 37
Leptodactylus pentadactylus (South American frog)	THE RESERVE OF THE PARTY OF THE	Antimicrobial activity	Alkaloids Spinaceamine 6-Methyl- Spinaceamine	19 20 68 36
Bufo spp.	Skin gland secretion	Pharmacological activity, Antimicrobia activity	Bufotenine Alvarobufotoxin Cinobufotoxin Fowlerbufotoxin Gamabufotoxin Marinobufotoxin Regularo— bufotoxine	n 30 e 1 ne 5

Animals Group, Class	Origin	Biological Activity	Active Substance	Refe-
Hyla arborea (European free frog)	Skin secretion	Pharmacological and antimicrobial activity	Proteins (pharmacologically active proteins)	46 12 68
Urodela spp. Friturus cristatus	Eggs extract	Aglutination	Aglutinin Various substances with biological activity	45
Phylobates urotaenia	Skin secretion	Aglutination	Batrachatoxin	37
Columbian poison Arrow frog)	Skin secretion	Antimicrobial activity and biological activity more toxic to mice	Batrachotoxin III	37
'arieba torosa California newt etradon spp.)	Eggs extract	Antibacterial activity Tetratoxin protects the egg clusters against predators and microorganisms.	Toxins Tarichatoxin Tetratoxin	28 91 37
endrobates spp.	Skin secretion	Antimicrobial activitiy	Alkaloids Cis- perhydroquinoline	37
endrobates striomcus	Skin secretion	Potentiates and blocks the indirectly elicited muscle twitch in a concentration dependent manner	Tricyclic alkaloid Gephyrotoxin (GyTx)	74
endrobatide (eotropical poison)	Skin secretion	Antimicrobial activity	Alkaloids	21
Or vor genammene Orrestant - cyclin II		tropoti princles Lergori, virgital Acressio, princip	Dermorphin Dermenkephalin	na thology na thology A stynois
25 Constitution 26 Constitutio	Skin secretion	Two powerful opioid peptides	The peptides issued from a common biosynthetic precursor. Dermophin from the precursor involves several posttranslational steps.	56 75
opis spp.	d	Antimicrobial activitiy	Xenopsin (biological active octa peptide)	85

Animals Group, Class	Origin	Biological Activity	Active Substance	Refe- rences
Nenopis leavis	Skin secretion	Antimicrobial activity	Xenopsin, Caerulein TRH, PGLA fractions	31
Xenapus laevis (Bouth African frog)	Skin secretion	Antimicrobial activity	Xenopsin, Caerulein, TRH, PGLA a HPLC chromatograms showed the secretion to be a complex mixture with over 30 components at similar levels to the four peptides previously isolated from X. laevis.	31 32
Xenopus laevis	Skin secretion	Antimicrobial activity	Caerulein was isolated and sequenced using restriction endonuclease	72
Nenopus laevis	Skin secretion	Antimicrobial activity	Antimicrobial active peptides	103 101
popple, special .	Alexandra y Haristan i	ereceitus de cateoraan	Antimicrobial active peptides	iona rid
Nenopus laevis	Skin secretion	Antimicrobial activity, antiprotozoal	The sequence of a partial C.DNA of precursor reveals that both peptides derive from a larger protein. These peptides	103 67
	TO SERVICE OF THE SER	and uses	appear to represent a previously unrecognized class of vertebrate antimicrobial activities.	gaj uno
Xenopis laevis	Skin secretion	Antimicrobial activity, non hemolytic and antiprotozoal activitiy	Peptides Magainin 1 Magainin 2 Neurotensin (like octapeptide)	103

Animals Group, Class	Origin	Biological Activity	Active Substance	Refe
Xenopis laevis	Skin secretion	Antimicrobial activity	Antimicrobial peptides (Magainin)	58
Xenopis laevis	Skin secretion	Astiminalia	Peptides Magainin 1 and 2, Levitidae, Xenopsin, Caerulin, Angiotensin,	6.9
TE PRINCE OF THE	Skill secretion	Antimicrobial activity	Bombesin, Bradykinin, Dermorphin, Sauvagine, Spasmolysin, Tachykinins, Thyrotropin.	6
Rana spp.	Skin secretion	Antibacterial activity	Serotonin (as active protein) toxic substance (in some species)	54
Rana esculenta (water frog)	Skin secretion	Antimicrobial and Phar- macological activity	Pharmacological active peptides	46
Rana esculanta	Secretory gland neurointermedi ate pituitary	Antibacterial activity	Polypeptide	6
Rana ridibunda	Skin secretion	Antibacterial activity	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	16
Rana pipiens	Skin secretion	dossieniena	Neuropeptide The skin is a complex secretory tissue, but the presence of large quantities of TRH and other biologically active neuropeptides make this tissue an attractive model to study the regulation of neuropeptide	7
The state of the s	of glibridg for	n secretion kon house arcticularity arcticularity	secretion	Xepo

Animals Group, Class	Origin	Biological Activity	Active Substance	Refe- rences
INAICE Clapide by boards	inglidaswova a diej	Neurotoxin and cytotoxin	(Protein) Neurotoxin I Neurotoxin II Neurotoxin III Neurotoxin IV Toxin A Toxin α Toxin B Toxin Y Cobratoxin Toxicin F2 Toxicin F1 Toxicin F3 Toxicin E5 Toxicin E6 Toxicin E7 Principal neurotoxin Minor neurotoxin Notexin	92 86 87 22
Hydrophiidae (Bea snakes)	Venom	Neurotoxin and cytotoxin	(Protein) Toxin a Toxin b Erabutoxin a Erabutoxin b	rocedi onceni umma
Viperidae (viperids, vipers)	Venom	Neurotoxin hemolytic and Coagulant activities	Viperotoxin Toxin	Teo.
Crotalidae (Crotalida, pit vipera)	Venom	Neurotoxin hemolytic and coagulant activities	Crototoxin Crotamin	83
FISH Anguilla	Serum	Hemolytic and Ichthyotoxic activitiy	Ichthyotoxin, hemolysin cytotoxin	4
Hypticus saponoccus (Atlantic soapfish)	Cutaneous gland toxin Foamy secretion	Highly toxic to fish and mice	Protein, polypeptide	52
Pardachirus marmoratus (flat-fish)	Skin secretion	Hemolytic and Ichthyotoxic activitiy	Protein	69
Gobiodon spp.	Skin secretion	Hemolytic and Ichthyotoxic activitiy	Toxin	38

Column chromatography; (1): Activity present, (4), addivity also at

MATERIAL AND METHODS

Isolation of the crude skin secretion extract and purification of the biologically active proteins from the crude extract: The water frogs, Rana ridibunda were obtained from Alibeyköy and Çorlu in Istanbul. Under ether anestesia the dorsal skin of the frog was massaged with fingertips back and forth to situmulate the foamy skin secretion. The secretion was collected with a spatula into a test tube, left in water bath at 100°C for 30 min, centrifuged at 5500 rpm for 30 min (T30 Janetzled 30 type centrifuge). The supernatant was used as the crude extract (CE). CE was precipitated with 95% (NH₄)₂SO₄ cut. Centrifuged at 10,000 rpm for 10 min (Beckman 21 rotor). Precipitate was dissolved in and dialyzed against 0,9 % NaCl. The skin secretion was further purified by either sephadex G25 or Sephadex G75 column chromotography (column height : 25 cm, column with 10 cm). 1 mg/ml protein containing crude extracet was applied to the column. 1 ml fractions were collected. The elution was done with physiological serum. The absorption at 280 nm was recorded for each fraction. The protein fractions with the biological activities obtained from Sephadex G75 column chromatography was further purified by DEAE Sephadex A25 column chromatography using the same procedure as above except the elution was done with stepwise NaCl concentrations (0.02 M - 1.00 M). The purification procedure is summarized in Fig. 1.

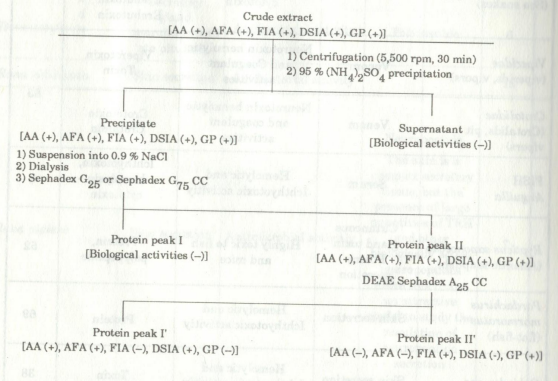


Fig. 1: Purification steps of the skin secretion and biological activities present (AA: Antibacterial activity; AFA: Antifungal activity; FIA: Fibrinolytic system inhibitor activity; GP: Glycoprotein; DSIA: DNA synthesis inhibitor activity; CC: Column chromatography; (+): Activity present, (-); activity absent).

Determination of antifungal effect: Fungi and mold were blained from Istanbul University, Istanbul Medical School, Department Microbiology. Sabouraud dextrose agar medium sterilized at 115°C in the autoclave for 15 min was used. After sterilization, 30 U/ml penicillin, medium streptomycine and 0.5 mg/ml actidion and 0.25 ml/ml skin medium were added into the medium. Two ml aliquats were put into the tubes. The controls contained physiological serum instead of skin medium. Fungi and dermatophyte strains were inoculated. The tubes mediated with dermatophytes were incubated at 26°C for 10 days. The mediated and incubated tubes by following the cell growth.

Isolation of Rattus norvegicus tumor cells (RTC): Tumor was developed by chemical carcinogenesis. The neck skin was treated with croton oil (H.J. Muller, Hamburg) or 7,12—dimethyl meanthracene (Fluka AG. Buscks SG). 8 months later, the tumor developed was transferred into tissue culture medium under aseptic multions. Passages were done at certain intervals into the medium 10% bovine serum where an antibiotic was also included. Tumor cells were kept after freezing in liquid nitrojen in deep freeze at—an Cuntil use.

Inolation of Rattus norvegicus embryonic fibroblast cells (HEFC): The embrionic cells were obtained from 10–15 days pregnant. The embrionic tissue was separated into its cells by trypsinization. The culture procedure of the embriyonic cells, primary cell that was obtained. The cultures obtained from the secondary passages were used in the experiments.

Investigation of the effect of frog skin secretion on monology of Rattus norvegicus tumor (RTC) and the normal monology of Rattus norvegicus tumor (RTC) and the normal monology of Rattus norvegicus tumor (RTC) and the normal monology of Rattus norvegicus tumor (RTC). Four sterile cover—slip were put monology of the normal monology of REFC): Four sterile cover—slip were put on the monology of the normal monology of Rattus norvegicus tumor (RTC) and the normal monology of RTC norvegicus tumor (RTC) and the normal monology of RTC norvegicus tumor (RTC) and the normal monology of RTC norvegicus tumor (RTC) and the normal monology of RTC norvegicus tumor (RTC) and the normal monology of RTC norvegicus tumor (RTC) and the normal monology of RTC norvegicus tumor (RTC) and the normal monol

were incubated in petri dishes for 1 hr in an incubator containing the air and 5% CO₂. Then the medium was withdrawn and 10 ml growth medium (GM) was added to each petri dish containing 5% or 10% CB mprotein peak I' (8%, 4%, 2%, 1%) over cover slips including cells were incubated for various periodic times (1-72 hrs) at 37°C later the slips were removed from the medium having CE or protein peak medium. The cover slips were put on a self constructed special microscope slide chamber, having the cell growing surface inverted for downward on the holes (a special microscope slide chamber. 10 mm diameter x 1,5 mm depth) surface touching medium containing above mentioned percent ages of CE or protein peak I'. They were investigated under the phase—contrast microscope vitally and their microphotographs were taken at 40x10 enlargements.

Investigation of the effect of skin secretion on the thymidine incorporation into DNA of RTC or REFU (autoradiographic studies)

The cells were incubated in 5% or 10 % skin secretion containing GM in petri dishes at 37 °C for 39 hours. The cells were separated from the medium and washed with Basalt salt solution (BSS) and were transferred into 10 ml of culture medium containing 3 mCi/ml of ³H-thymidine (Amersham Co. sp. act. 5000 mCi/mmole) incubated at 37° C for 30 min. The coverslips were fixed with acetic acid: ethanol (13) for 15 min, washed for 5 min with 70 % ethanol, dried with air and the coverslip containing these cells were mounted on gelatinized microscope slides. Then covered with Kodak AR-10 stripping film in a dark room They were placed in black light-tight boxes together with some silicage! and a small amount of solid CO_2 to remove O_2 . The boxes sealed with black tape and stored at +4 °C for one week for exposure. All procedures were carried out in safe light illumination. Also all the developing procedures were carried out in safe light illumination. The slides were allowed to reach room temperature then developed at 18-19 °C by placing in D₁₉ for 21/2 minutes and washed in 5% acetic acid for 1/2 min The slides were then transfered to 1/4 fixel for 10 minutes. Following washing by standing in water for 10 minutes, they were drained and dried. The slides were stained with a diluted solution of Giemsa blood stain containing citric acid, NaHPO4, methyl alcohol and Giemsa stock The slides were stained for 7 minutes at room temperature. Then dipped

the in distilled water and dried at room temperature. The slides were then examined by oil immersion for the presence of silver black grain.

Anothing to the intensity of the incorporation of radioactivity, the labelling of the cells were grouped as very little labelled, little labelled or little labelled and embrionic fibroblast cells and tumor cells were mated according to the intensity of ³H--thymidine labelling.

Recovery studies: The cells were incubated in a medium making CE (5% or 10%) at 37 °C for 17 hrs. Then washed 3 times with and transferred into a growth medium without skin secretion (CE) medium at 37 °C for 22 hrs. As the positive control, cells were medium with 5% CE for 39 hrs. As the negative control, were incubated in growth medium without skin secretion (CE) for the cells used were either RTC or REFC mentioned above.

The same procedure was repeated to show the effect different methation times on recovery where 17, 39, 48 and 72 hours of incubation were chosen. The negative and positive controls with respective methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells showed methation times were also included and followed until the cells are considered to the contract

RESULTS

Purification of skin secretion: In Fig. 1 together with methods the biological activities and presence of glycoprotein marized. (NH₄)₂SO₄ precipitated (95 % cut) protein fraction of the method (CE), isolated by centrifugation of the boiled skin secretion method, was applied either to Sephadex A25 or Sephadex G75 and Sephadex A25 column chromatographies. The biological manual manual manual (AA) to Gram negative and Gram positive manual manual (AFA), DNA synthesis inhibitor (DSIA) and molytic system inhibitor activities (FIA) determined with euglobuling and fibrin plate methods were all detected in the protein peak which comes out after the void volume which implies molecular under 25,000 Da's. In protein peak I' all of the above activities FIA were detected. FIA was detected in protein peak II'.

The elution profile of Sephadex G75 column chromatography is seen The biological activities were all positive in fractions 21, 22 and

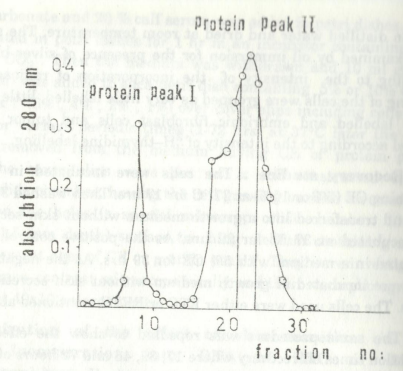


Fig. 2: Sephadex G₇₅ column chromatography of (NH₄)₂SO₄ precipitated frog skin secretion (FSS).

23. These fractions were poled together, concentrated with Amicon ultrafiltration using PM 10 membranes and passed through DEAN Sephadex A25 column chromatography, (Fig. 3) The protein peak I

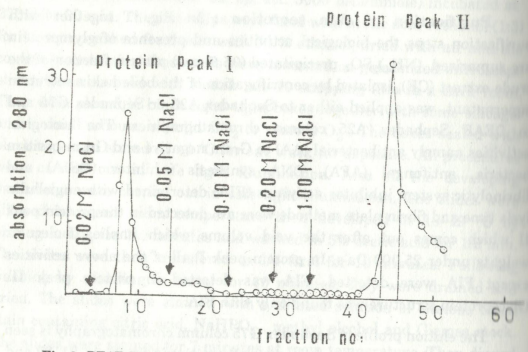


Fig. 3: DEAE Sephadex A25 column chromatography of pooled protein peak I

With 0.02 M NaCl elution had both the AA and AFA together the IA. The protein peak II' appearing upon 1 M NaCl elution had Throughout the purification steps the GP presence was also with periodic acid shiff (PAS) reagent and the glycoprotein was found to be related with fibrinolytic system inhibitor. The summary of the purification steps and spesific antibacterial at each step are summarized at Table II to show the efficiency faction at each step. After DEAE Sephadex A25 column at each step, both proteins I' and I" showed a single protein cellulose acetate electrophoresis at pH 8.6. It looks as if all biological activities namely antibacterial, antifungal, antitumoral are due to a protein with a molecular weight between 10,000 to 15,000 Da.

The efficiency of purification followed with antibacterial activities upon a puldermidis 69 micrococcus using hole method (16).

	Spesific	Purification fold	Total before			Total after		
	activity inhibition gone (em/mg)		Volume (ml)	Protein (mg)	Activity unit (inhibition zone cm)	Volume (ml)	Protein (mg)	Activity unit (inhibition zone cm)
	1.9	ppss-dad	29	1177	2236	166 _ 16 CK	1978 <u>(</u> 30) 1986 700V	provided trans
hy	4.7	2.5	20	1030	2000	22	141	661
	8.4	4.6	20	126	600	48	58	484
	12.2	6.4	45	54	453	40	37	451

The effect of skin secretion on fungi: The antifungal effect is an Table III. The skin secretion was effective on all of the fungi except to Torulopsis glabrata. All of the strains grew on control alarmed medium.

Table - III: The effect of frog skin secretion on the fungi strains obtained from Istanbul University, Istanbul Medical Faculty, department of Microbiology (AFA).

Fungus	Code no.	Frog skin secretion.	Control
Yeasts Candida albicans C. parapsilosis KUEN C. krusei C. albicans klaur C. albicans C. tropicalis C. stellatoidea C. albicans Cryptococcus neoformans Torulopsis glabrata	7650 1010 (Y) C1–12–1 KUEN1001 (Y) C1–6–3 A–2 KUEN1022 (Y) C1–19–3 KUEN1018 (Y) C1–18–1 MIV–211 KUEN1047 (Y) C2–1–9	on et ead phy step, b Muloge acc	+
Rhodotorula glutinis Saccharomyces cereviseae	KUEN1064 (Y) R1-1-1	olm 63 sibhee SO _{4 M} oleimin	+
Mould Microsporum nanum Frichopyton mentagrophytes	KUEN1089 (F) M5–6–1 703	is diversification diversification	didni Am

^{+:} growth was seen

The effect of frog skin secretion on the morphology of Rattus norvegicus tumor (RTC) and normal embryonic fibroblast cells (REFC):

In the cells left in the medium with 5 % crude skin secretion, no differences from controls were observed in 1 and 3 hrs of incubation. After 17 hrs of incubation, some differences like darkining of the nucleolus were observed. After 26 hrs, further darkining of the nucleolus and decrease in cytoplasm volume of some of the cells were observed but still the cells were alive. After 39 hrs 1 decrease in the cell number was seen. The individual cells became more circular and sharper, appearence of nuclei were noticible. 48 hrs later most of the cells were circular and their nuclei appeared darker. 72 hrs later most of the cells were circular.

^{-:} no growth

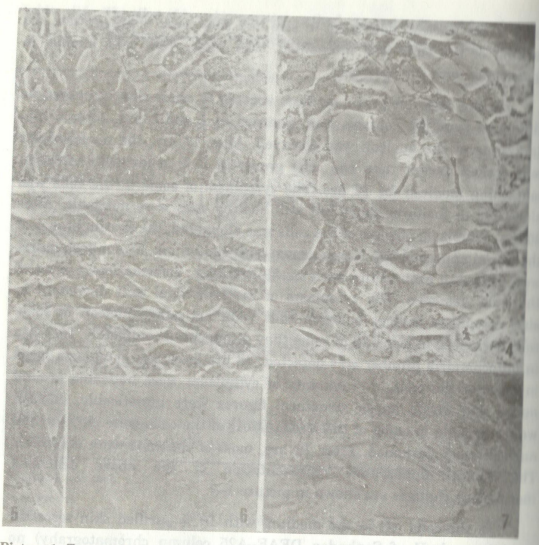
the nucleoli decreased in size and they were disintegrated, the tells was intensified. Still there were cells similar in which with the controls.

In the cells left in the medium with 10 % crude skin secretion, no from the controls was observed in 1 and 3 hrs. After 6 hrs, described of the nucleoli was observed. After 17 hrs, the nuclei of the the looked darker than the cells grown in 5 % crude skin secretion medium for 17 hrs. The cytoplasm of some of the cells was alanguated in fusiform shape but still healthy cells were noticable. After was elongated, the circumference of was drawn with Indian ink. The mitochondria wathered around the nucleus. 39 hrs later, deformation in most of the sells was seen. The cytoplasm of the individual cells decreased in and sytoplasm bridges between cells were noticable, nucleoli in some sells decreased in size and in some they disappeared completely. Increase member was detected. 48 hrs later, the cytoplasm lysed ment of the cells, or remnants of cytoplasm were noticable around and inside nuclei structural figures were unnoticable, nucleoli man noticable in some of the nuclei. Most of the cells were circular and were darkened. After 72 hrs, most of the cells were deformed. typical appearance representation of the above described marphalogy changes are shown in picture 1-7.

In the cells left in the medium with 10 % purified skin secretion peak II of Sephadex DEAE-A25 column chromatograhy) no from the controls was observed in 1 and 3 hrs. After 6 hrs, the the cells looked darker. After 17 hrs, decrease in cell number abserved. Some of the cells stood up on the coverslip and decrease in the little number of cells sticking onto the surface with a structure were noticable.

In the cells left in the medium with 8 % purified skin secretion, no the cells left in the controls were noticable in 1,3 and 6 hrs.

Thus, some of the cells had decreased cytoplasm 24 hrs later the median some of the cells were darkened, decrease in cytoplasm in the cells was noticable.



Picture 1: Tumor cells kept for 17 hrs in CE containing medium

Picture 2: Tumor cells kept for 39 hrs in CE containing medium

Picture 3: The control cells (non treated tumor cell with CE)

Picture 4: Tumor cells kept for 17 hrs in CE containing medium and transferred to new growth medium (after recovery)

Picture 5: The control cells (non treated embrionic cells with CE.

Picture 6, 7: The normal embryonic cells kept for 39 hrs in CE containing medium.

Most of the cells left in the medium with either of 4 %, 2 % or 1% CE no differences from the normal controls were noticable upto 24 hrs.

When the cells were incubated for 24 hrs in 10 % skin secretion, they recovered after 4 days, when incubated for 39 hrs, they recovered

after 6 days, when incubated for 48 hrs, they recovered after 9 days, when incubated for 72 hrs, they recovered after 12 days. The recovery time was elongated proportionally to the incubation time in a medium with skin secretion. (Fig 4)

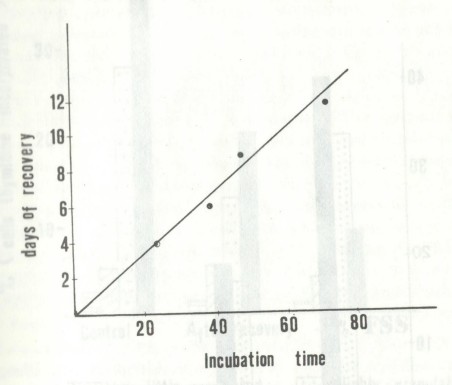


Fig. 4: The effect of incubation time in a medium with 10 % skin secretion on recovery days after transfer into new growth medium without skin secretion.

The effect of skin secretion on ³H-thymidine incorporation in DNA of the RTC and of REFC: Thymidine incorporation was detected in 2.75 % of the RTC grown in 5 % skin secretion containing medium. This percentage was 1.35 % for embryonic fibroblasts. In the tumor cells at 39 hrs of incubation in the medium without skin secretion this percentage was 27.95 %. It was 34.15 % for the embryonic fibroblast cells.

The cells grown in the medium with 10 % skin secretion, the DNA synthesis was 0 % for both kinds of cells at 39 hrs. After incubation of cells for 17 hrs with 5 % skin secretion they were transferred into normal medium ³H-thymidine incorporation in tumor cells after 22 hrs of

recovery the percentage was 13.55 %. This ratio was 17.75 % for embryonic fibroblast cells, the cells kept in the medium having 5 % skin secretion for 39 hrs (as control) ³H-thymidine incorporation for RTC were 2.75 % and it was 1.35 % for embryonic fibroblast cells (Fig 5, Fig 6)

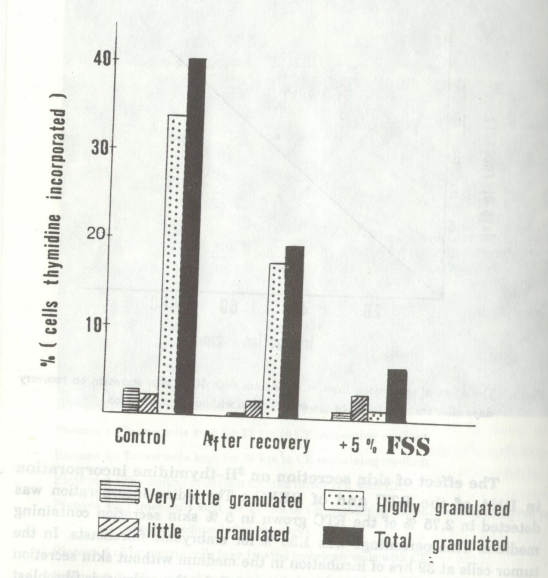


Fig 5: The effect of skin secretion on DNA synthesis of Rattus norvegicus embryonic fibroblast cells. Thymidine incorporation is represented by the intensity of granulation.

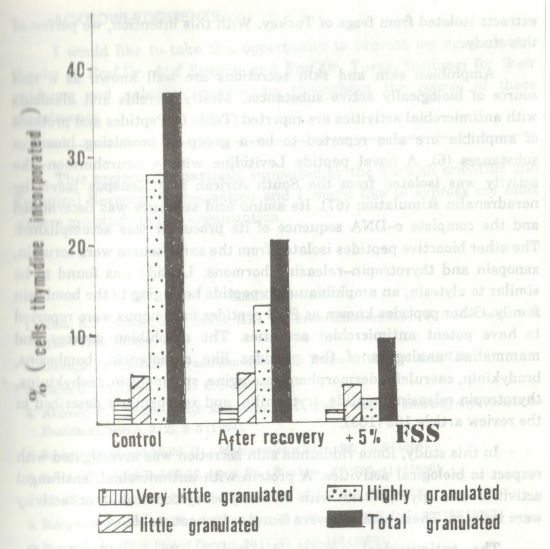


Fig 6: The effect of skin secretion on DNA synthesis of Rattus norvegicus of tumor cells.

Thymidine incorporation is represented by the intensity of granulation.

DISCUSSION

Various substances from different animal species with biological activities are reported in the literature. It is possible to detect these substances in protozoa coelenterata, coelamata and in the lower vertebrata like fish snake and amphibia (Table I). As seen at this table, most of the subtances with biological activities are of protein origin. Although there are quite a number of articles reported in the literature besides our work on amphibia, we believed that it would be very interesting to isolate substances with biological activities from the

extracts isolated from frogs of Turkey. With this intention, we perfored this study.

Amphibian skin and skin secretions are well known as a rich source of biologically active substances. Mostly steroids and alkaloids with antimicrobial activities are reported (Table I). Peptides and proteins of amphibia are also reported to be a group of promising bioactive substances (6). A novel peptide Levitidine with a neurohormon-like activity was isolated from the South African frog Xenopus laevis by noradrenalin stimulation (67). Its amino acid sequence was determined and the complete c-DNA sequence of its precursor was accomplished. The other bioactive peptides isolated from the same source were cerulein, xenopsin and thyrotropin-releasing hormone. Levtide was found to be similar to alytesin, an amphibian skin peptide belonging to the bombesin family. Other peptides known as PGS peptides in Xenopus were reported to have potent antimicrobial activities. The amphibian sources and mammalian analogues of the peptides like angiotensin, bombesins, bradykinin, caerulein, dermorphen, sauvagine, spasmolysin, tachykinins, thyrotropin releasing peptide, trytophilin, and xenopsin are described in the review article 103 (103).

In this study, Rana ridibunda skin secretion was investigated with respect to biological activities. A protein with antimicrobial, antifungal activities and glycoprotein with fibrinolytic system inhibitor activity were isolated. These proteins were found to be heat stable.

The antimicrobial activity detected in Rana ridibunda skin secretion belonging to a protein is inaccordance with the literature findings of other amphibia species (16).

This protein with M of 10,000 – 25,000 and with antibacterial and antifungal activities was found to be nontoxic to normal embryonic fibroblast cells and to tumor cells in vitro at the doses of antimicrobial effects. At very high doses, the thymidine incorporation into tumor and normal fibroblast cells of Rattus norvegicus were inhibited but when transferred to a medium without the Rana ridibunda skin secretion protein, the recovery of the DNA synthesis was established and they had a strong potency for becoming medically important substances. The most important advantages of these proteins are the antibacterial activity to a large spectrum of microorganisms and the fibrinolytic system inhibitor activity appearing in another molecule.

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