

CONTRACT NO: DAMD17-91-C-1001

TITLE: MOLECULAR STUDIES OF HTLV-I INFECTION IN NEWLY RECOGNIZED HIGH RISK POPULATION

PRINCIPAL INVESTIGATOR: Yehuda Danon, M.D.

CONTRACTING ORGANIZATION: Tel Be:

Tel-Aviv University Beilinson Medical Center Petah-Tikva, 49100, Israel

AD

REPORT DATE: July 10, 1993

TYPE OF REPORT: Final Report



93-21915

PREPARED FOR: U.S. Army Medical Research and Development Command, Fort Detrick Frederick, Maryland 21702-5012

93 9 21 02

DISTRIBUTION STATEMENT: Approved for public release; distribution unlimited

The findings in this report are not to be construed as an official Department of the Army position unless so designated by other authorized documents.

DISCLAIMER NOTICE



THIS DOCUMENT IS BEST QUALITY AVAILABLE. THE COPY FURNISHED TO DTIC CONTAINED A SIGNIFICANT NUMBER OF PAGES WHICH DO NOT REPRODUCE LEGIBLY.

REPORT D	OCUMENTATION P	AGE	m Approved OMB No. 0704-0188
asthering and maintaining the data predeg, and	completing and reviewing the collection of	information. Send comments rega	viewing instructions, searching existing data sources, iding this burden estimate or any other aspect of this information Operations and Reports, 1215 Jefferson ect (0704-0188), Washington, DC 20503
1. AGENCY USE ONLY (Leave blan		3. REPORT TYPE AN	
	10 July 1993	Final Repor	<u>t (6/10/91 - 6/9/93)</u> 5. FUNDING NUMBERS
4. TITLE AND SUBTITLE Molecular Studies Newly Recognized			Contract No. DAMD17-91-C-1001
6. AUTHOR(S)			
Yehuda Danon, M.D	•		
7. PERFORMING ORGANIZATION NA	ME(S) AND ADDRESS(ES)		8. PERFORMING ORGANIZATION
Tel-Aviv Universi	ty		REPORT NUMBER
Beilinson Medical	Center		
Petah-Tikva, 4910	0, Israel		
9. SPONSORING/MONITORING AGE			10. SPONSORING / MONITORING AGENCY REPORT NUMBER
U.S. Army Medical	Research & Develo	opment Command	
Fort Detrick	1 01800 5010		
Frederick, Maryla	nd 21/02-5012		
11. SUPPLEMENTARY NOTES		د	
12a, DISTRIBUTION / AVAILABILITY	STATEMENT		126. DISTRIBUTION CODE
Approved for publ	ic release; distr	ibution unlimi	ted
13. ABSTRACT (Maximum 200 word	s)		
to define newly To define the immigrants to I Leukemia (ATL).	rizes the two years or recognized high risk extent of HTLV-I in srael with an incre Various serologica ed, including ELISA a	population for H fection among gr ased frequency 1 and molecula	TLV-I infection. roups of Jewish of Adult T-cell r screening of
by amplificatio mononuclear cell	n of HTLV-I provin Is DNA. Overall rate	al DNA from p of infection i	rripheral blood is 12% for Jews
	urusan-North-Eastern		
	other parts of Iran,		
	ould not identify F t Diabetes Mellitus		
	phomas, Psoriasis a		
	for HTLV-I isolates		
sequence to Afric			· · · · · · · · · · · · · · · · · · ·
- 14. SUBJECT TERMS			15. NUMBER OF PAGES
	om Dolimonas '	Virus Xida	2. AVHILL VI FALL
HTLV-I, Epidemiol Biotechnology, RA		virus, Alas,	16. PRICE CODE
17. SECURITY CLASSIFICATION OF REPORT	18. SECURITY CLASSIFICATION OF THIS PAGE	19. SECURITY CLASSIFI OF ABSTRACT	
Unclassified	Unclassified	Unclassifie	
ISN 7540-01-280-5500			Standard Form 298 (Rev. 2-89) Prescribed by ANSI Std. 239-18 298-102

•

•



HTLY-I INDUCED T-CELL LYMPHOMA IN ISRAELI PATIENTS OF IRANIAN ORICIN

J. Rosenblatt, D. Meitës, Y. Sidi, Y.L. Danon Rogoff-Wellcome Medical Research Institute, Dept. of Medicine and Div. of Pediatric Immunology, Edith Wolfson Hospital and Beilinson Medical Center Tel-Aviv University Sackler School of Medicine, Petah-Tikva 49100 ISRAEL.

T cell lymphoma-leukemia (ATL) is one of the several clinical entities linked to human T cell lymphotropic virus (HTLV-I). Few geographic endemic concentrations of HTLV-I infection were already described: The Ryukyu Islands in Southern Japan, Central Africa and the Caribbean Islands. This is the first description of endemic HTLV focus in the Middle East. The prevalence and clinical presentations of ATL in Israel were studied. We have diagnosed four Israeli Jewish ATL patients and one HAM (HTLV-I - Associated Myelopathy) in a nationwide survey performed in 1986-1990. In three of the patients evidence for HTLV-I infection was obtained. All those patients immigrated to Israel from the same region in Central Iran. The nationwide survey and the clinical course of this new group of patients will be presented.

 \mathbf{z}^{i}

(((

((

Abstract Reproduction Form

Japan

Type abstract within blue rectangle. DO NOT FOLD THIS FORM.



MOLECULAR CHARACTERIZATION OF IRANIAN HTLV-I ISOLATES

Y. Kilim¹, J.D. Rosenblatt², D. Meytes³, D. Stephens², H. Lee⁴, Y. Danon⁴

Children's Medical Center of Israel, Petach, Tiyva, Israel¹, UCLA School of Medicine, Los Angeles, CA², Edith Wolfson Hospital, Holon, Israel³, Abbott Laboratories, N. Chicago, IL⁴, USA

We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Mashad in Northeastern Iran (Lancet 336:1533-1535, 1990). We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerate chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR. using primers to the tax/rex region. All 10 samples were found to be HTLY-I and not HTLV-II using discriminatory PCR. An immortalized Tcell line was derived from a Mashadi HTLV-I carrier. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Further characterization of Iranian HTLV-I isolates is underway. (Supported by the Rashi Foundation, Doron Foundation, ICRF, and a grant from the USAMRDC).

	To be completed by presenting author: Name Vehucia Danon, M.D.	
	Athiation(University.Company.etc.) Bellincu: Medical Center of Isrcel	
	Mailing Address Petah Tikva 49100, ISRAEL	
Na Ar	Country	•••••
• • •	The number of the main polopony for the bore abetract is. A D I would prefer a poster presentation	
	Signature	••••



EPIDEMIOLOGIC AND MOLECULAR CHARACTERIZATION OF HTLV-I INFECTION IN ISRAEL

(()

Yehuda L. Danon, Yael Kilim and J. Rosenblatt, Kipper Institute of Child Immunology, Children's Medical Ctr. of Israel, Sackler School of Medicine, Tel-Aviv Univ. Kaplan Str. #14-16, Petah-Tikva 49100 Israel.

Human T-cell leukemia viruses type I (HTLV-I) and type II (HTLV-II) are the two human retroviruses directly implicated in pathogenesis of human leukemia. HTLV-I has been linked to adult T-cell leukemia/ lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. Epidemiologic and molecular studies of both viruses have identified several themes underlying the leukemogenic process. Leukemia is a rare consequence of infection, and both viral infection and secondary transforming events appear to be involved. Due to the absence of a virally encoded oncogene, or preferred viral integration sites, novel mechanisms of leukemogenesis must be invoked. We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Mashad in Northeastern Iran (Lancet 336:1533-1535,1990). We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. An immortalized T-cell line was derived from a Mashadi HTLV-I carrier. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 differrent restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Further characterization of Iranian HTLV-I isolates is underway. A potential role for the HTLV-I/II transregulatory proteins Tax and Rex in leukemogenesis is discussed. Insights derived from the study of HTLV-I/II can be applied to the search for other oncogenic viruses.

(Supported by the Rashi Foundation, Doron Foundation, ICRF, and a grant from the USAMRDC).

11

V

 \mathbf{i}

Yehuda L. Danon, Yael Kilim and Joseph Rosenblatt, Kipper Institute of Child Immunology, Children's Medical Ctr. of Israel, Sackler School of Medicine, Tel-Aviv Univ. Kaplan Str. #14-16, Petah-Tikva 49100 Israel.

100000

177

Human T-cell leukemia viruses type I-(HTLV-I) and type II (HTLV-II) are the two human retroviruses directly implicated in pathogenesis of human leukemia. HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL). and HTLV-II to rare cases of chronic T-cell leukemia. Epidemiologic and molecular studies of both viruses have identified several themes underlying the leukemogenic process. Leukemia is a rare consequence of infection, and both viral infection and secondary transforming events appear to be involved. Due to the absence of a virally encoded oncogene, or preferred viral integration sites, novel mechanisms of leukemogenesis must be invoked. We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Mashad in Northeastern Iran (Lancet 336:1533-1535,1990). We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. An immortalized T-cell line was derived from a Mashadi HTLV-I carrier. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Further characterization of Iranian HTLV-I isolates is underway. A potential role for the HTLV-I/II transregulatory proteins Tax and Rex in leukemogenesis is discussed. Insights derived from the study of HTLV-I/II can be applied to the search for other oncogenic viruses,

(Supported by the Rashi Foundation, Doron Foundation, ICRF, and a grant from the USAMRDC).

••

Ŋ

Ì

garan the part for a party		ACT SUBI FORM
V	Please read instructions on the reverse side of this sheet, before filling this form.)
Key words 1st	EPIDEMIOLOGIC AND MOLECULAR CHARACTERIZATION OF HTLV-I INFECTION IN ISRAEL	
	Yehuda L. Danon, Yael Kilim and Joseph Rosenblatt	
HTLV-1	Kipper Inst. of Child Immunology, Children's Med. Ctr	
	of Israel, Sackler School of Med., Tel-Aviv Univ. Israe	"
2nd	HILV-I has been linked to adult T-cell leukemia/lymphon	
EPIDENIOLOGY	(ATLL), and HTLV-II to rare cases of chronic T-cell leu	1-
	kemia. We recently reported a high rate of HTLV-I sero positivity among immigrants to Israel from Mashad i	
3rd •	Northeastern Iran. We have now characterized the Irania	1
0	HTLV-I isolate using a combination of Southern blotting	
HILL	polymerase chain reaction (PCR) and sequencing. DNA fro	
۱ <u>ــــــــــــــــــــــــــــــــــــ</u>	10 HTLV-I seropositive Mashadi carriers was isolate from peripheral blood mononuclear cells and amplified b	
PARTICULARS OF ABSTRACTS	PCR using primers to the tax-rex region. All 10 sample	
	were found to be HTLV-I and not HTLV-II using discrimi	-
TITLE:	natory PCR. The cell line contained integrated HTLV-	
	provirus. Southern blotting and restriction mapping o the Iranian isolate demonstrated marked similarity to	
	published Japanese isolates using 10 different restrict	-
APPROPRIATE WORK SHOP	tion enzymes. Amplification and preliminary sequencing	в
1st Choice : TR.B	of a 900 bp segment of env from two Mashadi isolates	
2nd Choice : TR.A	disclosed approximately 95% nucleic acid sequence homo- logy with the published sequence of Japanese isolates.	
3rd Choice :	and >98% homology between two Mashadi isolates. These	· •
(use codes as given on page	data indicate substantial conservation of sequence be-	- (
11 of the document)	tween Iranian and Japanese HTLV-I isolates. Insights	
	derived from the study of HTLV- I/II can be applied to the search for other retroviruses and oncogenic viruses.	
FUIL MAILING ADDRESS OF	(Supported by the USAMRDC and Doron Foundation)	
PRESENTING AUTHOR	ANON YEHUDA L.	یانی رو (میرانی)
PROF. / BR. LMR. AMS. Dt		
NATIONALITY ISPA		·
AFFILIATION TE	Aviv University	
ADDRESS CHILDRENS	MEDICAL CTR OF ISRAEL	
Kaplan	Str. # 14-16 - BEILINSON	• .
ON PETAH TI	AVA POSTALCODE 49100 COUNTRY ISRACL	
TELEPHONE 972-3-	3393905 FACSIMILE 92-3-929756ELEX	

.

•

New Daini.

.

•

.(

(

Original Abstract Form

 \mathcal{W} Viruses and Virus-Like Agents in >lsease

Molecular characterization of HTLV-I infection in Israel

<u>Yehuda L. Danon</u>, Yael Kilim and Joseph Rosenblatt Kipper Inst. of Child Immunology, Children's Med. Ctr. of Israel, Sackler School of Med., Tel-Aviv Univ. Israel

HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Northeastern Iran. We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Insights derived from the study of HTLV-I/II can be applied to the search for other retroviruses and oncogenic viruses.

Supported in part by USAMRDC Grant # DAMD17-91-C-1001

I wish to submit an abstract for presentation as poster on

Monday, March 8, 1993 🛛 🗍 T

🗌 Tuesday, March 9, 1993

Key Words

(

((

Name of Presenting Author Company/Institute

Yehuda L. Danon, M.D. Director, The Children's Medical Center of Israel Beilinson Medical Center Petah-Tikva 49100 ISRAEL

Mailing Address

Department

City

Country

Postal Code

Telephone 972-3-939 Telefax 7515 Signature Deadline for submission of abstracts is December 15, 1992

Please send the original abstract form and 3 photocopies by airmail to: S. Karger AG, 1993 Congress, P.O. Box, CH-4009 Basel, Switzerland

Lie, del pere

ý

· 3)

MOLECULAR CHARACTERISTICS OF HTLV-I INFECTION IN NEWLY CHARACTE-RISED HIGH RISK GROUP OF CARRIERS IN THE MIDDLE EAST.

HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Mashad in Northeastern Iran. We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Insights derived from the study of HTLV-I/II can be applied to the search for other retroviruses and oncogenic viruses.

(Supported by the Rashi Foundation, Doron Foundation, ICRF, and a grant from the USAMRDC).

EPIDEMIOLOGIC AND MOLECULAR CHARACTERIZATION OF HTLV-I INFECTION IN ISRAEL

Hay Fred.

t

4

Yehuda L. Danon, Yael Kilim and Joseph Rosenblatt, Kipper Institute of Child Kumunology, Children's Medical Center of Israel, Sackler School of Medicine, Tel-Aviv University Israel

HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. We recently reported a high rate of HTLV-I sero-positivity among immigrants () Israel from Mashad in Northeastern Iran. We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. The integrated HTLV-I provirus. Southern cell line contained mapping of blotting and restriction the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Insights derived from the study of HTLV- I/II can be applied to the search for other retroviruses and oncogenic viruses. (Supported by the USAMRDC and Doron Foundation)

15

-

IXth International Conference on AIDS, Berlin, June 7 - 11, 1993

in affiliation with the IVth STD World Congress



ABSTRACT FORM

Only this official form is acceptable as the original submission (no facsimile transmission). This form should be accompanied by 5 photocopies.

Choice of Topics

IMPORTANT:

Please indicate your preference according to the list of topics on the reverse side $(A1 \sim D38)$.

1st choice	
2nd choice	
STD*	
I prefer:	r
poster presentation	
oral presentation	

Choice of Key Words

See List of Key Words in "Call for Abstracts" and type in corresponding numbers

Affiliation

Address

Tel.

Fax

Mail original abstract and 5 photocopies to:

IX th International Conference on AIDS IV th STD World Congress Institute for Clinical and Experimental Virology of the Free University of Berlin Hindenburgdamm 27 D-1000 Berlin 45

NOLECULAR CHARACTERIZATION OF HTLV-I INFECTION IN ISRAEL <u>Yehuda L. Danon</u>, Yael Kilim and Joseph Rosenblatt, Kipper Inst. of Child Immunology, Children's Med. Ctr. of Israel, Sackler School of Med., Tel-Aviv Univ. Israel

HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Northeastern Iran. We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Insights derived from the study of HTLV- I/II can be applied to the search for other retroviruses and oncogenic viruses.

	· .			
S .		Instruct	tions:	Ć

1. Type within blue lines: title, author's name (6 or less), affiliations, city, state, country. (Underline name to indicate presenter; if there are more than 6 authors, type et al. to indicate additional authors). Do not include more than 6 authors' names.

2. Any handwritten symbols must be drawn in black ink.

3. REMEMBER: Your camera-ready abstract will be printed in the book of abstracts exactly as typed. No editorial corrections will be made.

4. For sample abstract see reverse side.

ABSTRACTS MUST BE RECEIVED NO LATER THAN JANUARY 15, 1993

*STD-related presentations should be assigned to the respective topic of each track.

HTLV-I VIRUS IN INSULIN-DEPENDENT DIABETES MELLITUS Y. Kilim, M.Sc.¹, M. Karp, M.D.² and Y.L. Danon, M.D.¹

Gueres

Kipper Institute of Immunology,

²Institute of Pediatric and Adolescent Endocrinology,

... The Children's Medical Center of Israel, Petah-Tikva, Israel

Human T-cell Leukemia Virus-I has been linked to adult T cell leukemia/lymphoma (ATLL) and HTLV-II to some cases of chronic T cell leukemia. We have recently reported a high rate of HTLV-I seropositive among immigrants to Israel from northeastern Iran, and especially the town of Mashad.

To determine the frequency of antibodies to HTLV-I virus in Insulin-Dependent Diabetes Mellitus (IDDM) patients, sera from 56 newly onset IDDM patients were tested by an enzyme immunoassay. According to our method the reactivity of antibodies detected by enzyme immunoassay against HTLV-I encoded antigens was determined by an assay which employs recombinant HTLV-I antigens. No antibodies to HTLV-I were detected in all 56 patients studied. Proliferative response to various species of insulin was performed in 26 of those patients, 23 out of 26 showed a positive response. Sera from 56 newly onset IDDM patients were screened for ICA. ICA were detected in 32 (57.1%) of the 56 patients.

-

()

It seems that HTLV-I is playing no role in IDDM.

Supported in part by USAMRDC Grant # DAMD17-91-C-1001

n

VIIIth INTERNATIONAL CONFERENCE ON AIDS IN AFRICA & VIIIth AFRICAN CONFERENCE ON STDs



ABSTRACT FORM DEADLINE : 31 AUGUST 1993

EPIDEMIOLOGIC AND MOLECULAR CHARACTERIZATION OF NEW HTLV-I INFECTION FOCUS IN THE MIDDLE EAST

Yehuda L. Danon, Yael Kilim and Joseph Rosenblatt, Kipper Institute of Child Immunology, Children's Medical Center of Israel, Sackler School of Medicine, Tel-Aviv University Israel

HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. We recently reported a high rate of HTLV-I sero-positivity among immigrants to Israel from Mashad in Northeastern Iran after a national serologic survey of blood donors. We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 17 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 17 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian, African (Zair) and Japanese HTLV-I isolates. Insights derived from the study of HTLV- I/II can be applied to the search for other retroviruses and oncogenic viruses. (Supported by the USAMRDC and Doron Foundation)

LANGUE OF PRESENTATION Frencip English	Oral presentation Poster
	KEY WORDS
AUTHOR'S NAME: PROF. Y. DANON Adress: CHILDRENS MEDILAL CTR Telephone: POBOR SS9 Fax: PLANT. KV9 49202 ISRAEL Presenting author's signature:	OF ISRAC, Tel AvivUnversity. 972-3-9247515 The:
Airmail this original abstract form piles 5 photos to :- CONFERENCE SECRETARIAT : VIIIth International Conference on AIDS and STDs in Africa	FOR SECRETARIAT USE ONLY Abstract N°:
lbis, place Charles Nicolle - Box:1818 - Casablanca. Morocco Tel : (212) 2 29-53-25 / (212) 2 29-53-26 Fax : (212) 2 29-53-27 / (212) 2 29-53-28 Felex : 45 382 M	Date received : Registration N°:
REVIEWER USE ONLY : A R	Score :
	Poster

A Decade with HTLV-I/HTLV-II: Lessons in Viral Leukemogenesis

Joseph D. Rosenblatt¹, Yehuda, Danon², and Alexander C. Black¹

¹Division of Hematology-Oncology, Department of Medicine and Jonsson Comprehencive Cancer Center, UCLA School of Medicine, Los Angeles, CA, USA and the ²Kipper Institute of Child Immunology, Children's Medical Centre of Israel, Beilinson Medical Center, Petach Tiqva, Israel

INTRODUCTION

The past decade has seen myriad advances in detection and characterization of human retroviruses. It began with initial description of human T-cell leukemia virus type I (HTLV-I) by Poiesz and Gallo in the US and Yoshida in Japan, which pointed to the involvement of the human retrovirus, HTLV-I, in an unusual form of T-cell malignancy, adult T-cell leukemia/lymphoma (ATLL) (1,2). The identification of HTLV-I intensified the search for related viruses, and soon thereafter, human T-cell leukemia virus type II (HTLV-II) was described by Kalynaraman and Gallo in a cell line derived from a patient with a chronic T-cell leukemia with features of hairy-cell leukemia (3). The rapid identification of HTLV-II on the heels of HTLV-I led to speculation that a host of human oncogenic retroviruses would soon be identified. The subsequent discovery of human immunodefiency virus types 1 (HIV-1) and 2 (HIV-2) and their implication in acquired immunodeficiency syndrome (AIDS) accelerated the pace and intensity of the search for oncogenic viruses. It was soon recognized that leukemic cells in malignancies associated with HTLV-I and -II contained clonally integrated provirus; in effect, a signature for direct viral involvement in the oncogenic process. In contrast, neoplasms frequently seen in the setting of HIV-1 infection (e.g. Kaposi's sarcoma and/or highgrade B-cell lymphomas) did not appear arise as a direct consequence of viral transformation of HIV-1infected cells. At the end of the decade, only HTLV-I and -II remain clearly implicated as directly leukemogenic human retroviruses. Therefore, we believe that insights gleaned from investigation of these viruses can and should be applied to the search for other oncogenic retroviruses.

EPIDEMIOLOGICAL LESSONS

The epidemiological link between HTLV-I and ATLL resulted from two complementary events. The first, an appreciation by Uchiyama and Takatsuki that ATLL represented a unique clinical entity (4) allowed geographic localization of the disease to southern islands of Japan: Kyushu, Shikoku, and the Ryuku chain of islands. Development of serological assays for HTLV-I led to correlation of HTLV-I infection to the presence of malignancy, as well as a determination of modes of

LEUKEMIA © 1992 Macmillan Press Ltd transmission (for review see 5). Epidemiological studies have suggested that exposure shortly after birth is a major risk factor for subsequent development of ATLL (5,6). In addition, these studies have demonstrated that twenty or more latent years may elapse between acquisition of infection and development of malignancy (5,6). Furthermore, only a minority (< 5%) of HTLV-I carriers actually develop ATLL (7), and ATLL as a consequence of transfusion-acquired HTLV-I is virtually unknown.

Hence, several general observations emerged from scrutiny of HTLV-I epidemiology; (a) leukemia may be an infrequent consequence of exposure to a fairly wide-spread virus; (b) leukemogenesis may depend on the timing and/or length of exposure, so that individuals infected in childhood may be at higher risk than those infected later in life; and (c) the long latency period suggests a multiple step process may be involved in leukemogenesis; while viral infection may be a prerequisite, it alone may be insufficient to produce the leukemic phenotype. These general epidemiologic features of ATLL suggest that a systematic re-evaluation of appropriate candidate malignancies and viruses may reveal other oncogenic viruses. Fairly prevalent or even ubiquitous viruses could conceivably manifest oncogenic potential in a sporadic fashion, and factors such as timing and length of exposure may be critical.

Careful cataloguing and description of clinical syndromes is essential to derive epidemiologic clues that may lead to virus identification. The recognition that non-Hodgkin's lymphomas could be divided into Tand B-cell subtypes and subsequent differentiation of ATLL from *mycosis fungoides* is a case in point. While ATLL was undoubtedly a frequent reason for in-patient hospitalizations in Japan prior to 1977, it was thought to be a variant of peripheral cutaneous T-cell lymphoma, and its characteristic features such as hypercalcemia and enhanced expression of interleukin 2 (IL-2) receptor alpha (IL-2R α) chain (Tac antigen) on the cell surface were initially overlooked. Recognition of the subtler clinical aspects of the syndrome allowed for subsequent epidemiologic observations (4).

In contrast to HTLV-I, it is premature to reach conclusions regarding pathogenesis by HTLV-II. Although originally isolated from the Mo T-cell line, a transformed T-cell line derived from the spleen of a patient with hairy-cell leukemia, the nature of the malignancy *in vivo* in the patient was not adequately addressed (8). We know that HTLV-I and -II can transform T-cell lines *in vitro*, and that the Mo T-cell line may have simply represented an outgrowth of HTLV-transformed cells *in vitro*. A second patient with HTLV-II and hairy-cell leukemia was found by our laboratory to have a biclonal lymphoproliferative

LEUKEMIA, Vol 6, Suppl 1, 1992; pp 18-23

Correspondence to: Joseph D. Rosenblatt, Division of Hematology-Oncology, Department of Medicine and Jonsson Comprehensive Cancer Center, UCLA School of Medicine, Los Angeles, CA 90024-1678, USA.

disorder in which a B-cell hairy-cell leukemia and a co-existant malignant CD8+ T-cell clone were observed (9,10). Oligoclonal integration of HTLV-II provirus into the CD8+ T-cells provided strong evidence for origin of malignancy in a virally infected cell. However, as additional cases of HTLV-II-induced malignancy have not been reported, there is considerable doubt as to whether we have as yet characterized the prototypic disease associated with HTLV-II.

An additional surprise that has emerged from epidemiological studies of HTLV has been the fact that screening procedures for HTLV-I identify a considerable number of crossreactive HTLV-II carriers. This raises the possibility that in the process of assaying for newly identified viruses. we may inadvertantly be assaving for a variety of crossreactive members of the same viral family. Specifically, intravenous drug abusers (IVDA) found to be seropositive for HTLV-I have been reported in several studies to have a higher incidence of HTLV-II infection and > 50% of seropositive random blood donors screened by HTLV-I ELISA were actually found by DNA amplification techniques to harbour HTLV-II (11-13). In the future. screening for newly identified viruses should be performed using both DNA amplification and serological techniques to avoid the initial confusion in delineating the epidemiology of HTLV-I and -II.

The HTLV-I model of malignancy as a rare consequence of infection with a prevalent virus suggests that careful molecular re-examination of the role of already recognized viruses in suspect malignancies is in order. In the case of HTLV-I, over a million infected individuals in Japan give rise to only 400-500 cases of ATLL per year. Hodgkin's disease (HD) may provide another case in point. The bimodal age distribution. prevalence, anecdotal descriptions of geographical clustering, and 'outbreaks' of HD suggest that an infectious agent may underlie pathogenesis (14-21). The increasingly frequent reports of Epstein-Barr virus (EBV) genome detection in some cases of HD suggest that EBV can be important in pathogenesis of a subset of HD patients (14-21). As another example, recognition that four of the last five cases of ATLL-like T-cell lymphoma in Israel occurred in Iranian immigrants from the northeastern city of Mashad, allowed identification of a new focus of HTLV-I infection (22,23). Recognition of geographic, familial and/or ethnic clustering of particular malignant disorders may yield important clues to viral etiology. It is important to note, however, that such time/space clustering may frequently relate to non-infectious risk factors rather than a virus. Some investigators have speculated on the likelihood of viral involvement in childhood acute lymphoblastic leukemia (ALL); however, the evidence is only mildly suggestive at best. Some studies suggest an association with geographic areas of high socioeconomic status, while others do not (24-29). Additional studies suggest a mild degree of clustering, particularly in children less than six years of age, although this remains controversial (29-31). Thus, little evidence points to an infectious cause or an underlying common leukemia virus in ALL. If a virus were involved,

analogy to ATLL would suggest that it might initially cause an insignificant acute infection that establishes latency and eventually leads to leukemia through secondary events. Given the lack of overt clustering, seroepidemiological studies are unlikely to settle the issue, and frank demonstration of molecular involvement of an infectious agent will likely be necessary.

New Molecular Mechanisms of Pathogenesis

The explosive growth in the study of oncogenes over the past decade came about as a result of recognition that in animal malignancies brought about by retroviral infection, the transduction of a cellular proto-oncogene and its inappropriate expression under control of the viral promoter was frequently observed in retrovirally induced tumors. A second type of molecular lesion frequently observed was integration of a retrovirus adjacent to a cellular oncogene and loss of normal patterns of proto-oncogene expression. These observations led to speculation that similar mechanisms may be operative in human malignancy. However, HTLV-I afforded a unique surprise, in that the viral sequences did not contain any transduced cellular sequences, and that integration sites appeared to be random. While clonal integration was observed by Yoshida and colleagues, the sites of integration often occurred on different chromosomes, and no specific integration patterns could be observed (2,32). These observations led to a search for new mechanisms of oncogenesis.

The demonstration by Seiki and colleagues of the unique coding regions located in the 3' end of the genome provided a first clue as to potential mechanisms (33). The 3' ends of the genome of HTLV-I, -II and the leukemogenic bovine leukemia virus (BLV) all contain coding sequences for at least two overlapping genes (34-37). These genes, known as tax and rex, are now known to encode a transcriptional and post-transcriptional regulator of viral expression. The initial discovery of the tax gene was surprising, in that such trans-acting transcriptional regulators had only been previously identified in DNA viruses such as herpes simplex virus, varicella-zoster virus, and adenovirus. The HTLV-I tax gene encodes a 40-kilodalton (kDa) protein, and the HTLV-II tax gene encodes a 37-kDa protein (34-37). These nuclear phosphoproteins are highly homologous, and have similar effects on transcription. Tax expression is necessary for transcriptional activation of virus replication and efficient RNA production from the viral long terminal repeat (LTR); however, an additional property noted early was the ability of Tax to trans-activate other viral and cellular promoters, including those involved in regulation of T-cell growth. Examples of such promoters include those for IL-2, IL-2R α , and the lymphokine, granulocyte-macrophage colony-stimulating factor (GM-CSF) (38-41). In the case of HTLV-I and -II, Tax appears to interact with three 21-base pair (bp) repeats located in the U3 portion of the viral LTR. Within the sequence of the 21-bp repeats are core sequences known to bind cellular transcription factors (42,43). A variety of these proteins have now been identified and

partially characterized.

In contrast to the HTLV promoters, Tax activation of the IL-2R α gene involves induced nuclear expression of a cellular DNA binding protein, which interacts with an NF- κ B-like enhancer (44). NF- κ B is a DNAbinding factor first shown to interact with the enhancer of the κ light chain immunoglobulin gene (45). Tax interaction with NF- κ B is thought to account for inducibility of some other cellular and viral promoters such as the HIV-1 promoter. The lack of an NF- κ B-like binding site in the HTLV-I/-II promoter and deletion of NF- κ B-like binding sites from the GM-CSF promoter with retention of response to Tax indicate that Tax may act via different pathways in different cellular and promoter contexts (41).

To date, only limited evidence directly implicates Tax in T-cell transformation. Introduction of Tax coding sequences under the control of a herpes saimiri vector has resulted in continuously proliferating T-cell lines in vitro, although the transformed cell lines appear to retain dependence on IL-2 for continued growth (46). Expression of HTLV-I Tax under the control of the HTLV-I LTR in transgenic mice does not lead to Tax expression in T-cells, and T-cell malignancy is not observed (47). Some mice developed mesenchymal tumors reminiscent of neurofibromatosis, as well as muscular atrophy. Recent HTLV-I Tax transgenics under control of the T-cell-specific Thy-1 promoter also did not result in T-cell malignancy (48). Nevertheless, the promiscuous interaction of Tax with a variety of viral and cellular promoters suggests that it may play a pivotal role not only in the HTLV-I life-cycle, but also in definition of the malignant phenotype. Tax can *trans*-activate the IL-2R α gene, which offered an explanation for the high degree of Tac (high affinity IL-2 receptor) antigen expression in HTLV-I-transformed T-cells. Similarly, ectopic GM-CSF production due to Tax may cause eosinophilia, which is frequently seen in ATLL. In addition, Tax may also be involved in trans-activation of the parathyroid hormone-related protein (PTHRP) promoter, perhaps accounting for the ectopic expression of PTHRP in ATLL cells, thereby leading to altered calcium metabolism (49). However, HTLV-I mRNA expression in ATLL is so low that it has required use of RNA polymerase chain reaction (PCR) to be detected. Therefore, whether effects seen with Tax in vitro have applicability to HTLV-I in vivo remains unclear.

The ability of the viral *trans*-activator, Tax, to interface with several cellular transcriptional factor pathways suggests a new model for viral leukemogenesis. The presence of such *trans*-acting genes may allow development of new assays for the presence of as yet undiscovered retroviruses based on the ability of *trans*-acting 'Tax-like' transcriptional regulatory proteins to act on cellular and viral genes. Models for cooperation between oncogenes could undoubtedly be applied to help dissect a potential role for Tax in cooperation with other oncogenes. New models for oncogenic cooperation have emerged at this conference, such as superinfection of E_{μ} -myc transgenic mice

with Moloney murine leukemia virus (MoMuLV) (50,51).

Several conclusions can be derived from study of the *trans*-regulatory *tax* gene: (a) new mechanisms of retroviral leukemogenesis other than transduction of cellular proto-oncogenes and/or retroviral insertion adjacent to cellular proto-oncogenes may be operative in human malignancy; (b) human retroviruses possess transcriptional activators that may affect expression of cellular genes, and aberrant expression of cellular genes may contribute either to leukemogenesis *per se*, or to the leukemic phenotype; and (c) the effect of viral transregulatory genes may be felt early in leukemogenesis, and may be insufficient to elicit the full-blown leukemic phenotype.

An additional transregulatory gene studied more recently is the rex gene of HTLV-I, -II, and BLV. The rex gene is required for productive HTLV-I/-II infection. The rex gene of HTLV-I encodes two proteins, one of 27 kDa and one of 21 kDa, from an overlapping reading frame to that encoding $p40^{tax}$ (52). These proteins appear to result from utilization of an alternative initiator methionine. In HTLV-II, two proteins are also encoded of the apparent sizes, 26 and 24-kDa (53). In HTLV-II, these appear to derive from different degrees of phosphorylation, with the larger molecular weight species being a hyperphosphorylated form of the 24-kDa protein (54). In both HTLV-I and -II, the proteins appear to act as post-transcriptional regulators, and elicit export of full-length gag/pol mRNA and probably partially spliced env transcripts from cell nucleus to cytoplasm. The rex gene appears necessary to allow expression of non-spliced and partially spliced viral mRNA, which in turn allows synthesis of Env and Gag proteins and production of mature virions.

In HTLV-I, Rex has been found to act through a cis-acting Rex-responsive element (RxRE) located in the 3' LTR. Our group has studied Rex effects inediated through the 5' LTR of HTLV-II (55). In both cases. Rex appears to act through sequences located in the R region, downstream from the transcription initiation site. Assays of binding to radiolabeled viral * RNAs have demonstrated that purified HTLV-II Rex can directly bind to transcripts initiated from the 5' LTR, and that binding occurs to a portion of a cis-acting element responsible for Rex action, known as the RxRE (56,57). Mapping in our laboratory has demonstrated that Rex can bind directly to transcripts as short as 115 bp derived solely from sequences within the R region (57,58). These transcripts contain the 5' LTR splice donor site, and mutation of the splice donor site appears to impair Rex binding and function. Furthermore, Rex binding is dependent on retention of a specific stem-loop mRNA structure located downstream from the splice donor site (from nucleotide 465-501 within the HTLV-II 5' LTR) (57). This stem-loop structure is conserved in both HTLV-I and -II. Rex binding may be facilitated by hyperphosphorylation, and it would appear to be the higher molecular weight (26 kDa) Rex species of HTLV-II that binds efficiently, indicating that cellular controls on Rex function may exist at the level of phosphorylation

(Chen and Green, unpublished observations). Nucleolar localization and our results using RxRE mutations of the splice donor site suggest a direct interaction of Rex with the cellular splicing apparatus to facilitate bypass of cellular splicing mechanisms.

An intriguing observation regarding Rex of HTLV-I was first made by Rimsky and Greene, demonstrating that HTLV-I Rex can functionally substitute for the Rev protein of HIV-1 (59). The Rev protein of HIV-1 performs an analogous function to that described for Rex in HTLV-I. Their assay demonstrated the capacity of Rex to induce production of the truncated singleexon form of the HIV-1 Tat protein that reflects translation from unspliced env vector mRNA (57). HTLV-II Rex in our laboratory is also able to rescue replication of Rev-deficient mutants of HIV-1 (58). Rescue of HIV-1 Rev-deficient mutants by HTLV-II Rex is relatively inefficient, and this can partially be accounted for by the relatively low affinity of HTLV-II Rex binding to the HIV-1 Rev-responsive element (RRE) (58). In addition, HIV-1 Rev is unable to complement an HTLV-II Rex-deficient clone, indicating a non-reciprocal pattern of complementation. Nevertheless, the ability of Rex to complement the genetically distant HIV-1 virus in trans suggests that. like Tax. Rex may act promiscuously on a variety of non-HTLV target sequences. This raises the possibility that Rex may also interact with cellular RNA to elicit aberrant splicing and/or transport. Disrupted processing and expression of cellular mRNAs could conceivably be implicated in the process of leukemogenesis as well. Therefore, post-transcriptional regulators may also be involved in the process of retroviral leukemogenesis. Direct evidence supporting this hypothesis has not been obtained.

Disparate Disease Entities Related to HTLV-I

Approximately five years following its discovery, it was found that the pathology elicited by HTLV-I in one setting may not predict other forms of pathology related to the virus. A specific illustration is afforded by the two major non-overlapping syndromes associated with HTLV-I. While HTLV-I may cause ATLL. another subset of infected individuals, approximately half as many, will develop a slow neurologic disease characterized by gradual development of spastic paraparesis of the lower extremities with minimal sensory loss. This illness, tropical spastic paraparesis (TSP) (also known as HTLV-I-associated myelopathy (HAM)), is distinguished from multiple sclerosis (MS) by virtue of its chronic progressive and non-episodic nature, as well as the general limitation of pathology to motor control in the lower extremities and sphincter dysfunction. The association between this illness and the virus was discovered by Gessain and co-workers while screening neurologic illnesses for retroviral involvement in Martinique (60). As opposed to the leukemia, where HTLV-I has been observed to infect and transform T-cells in vitro, no adequate model for pathogenesis of the myelopathy exists. Regardless of underlying mechanisms, involvement of HTLV-I in a

slow neurologic disease was not predictable on the basis of its involvement in T-cell leukemia. The latency period for development of HAM is also appreciably shorter, and recently at UCLA, we saw a patient develop myelopathy approximately fifteen months following infection by transfusion (61). In contrast, development of ATLL following transfusion-acquired HTLV-I is almost never seen. Furthermore, co-existent ATLL and HAM have rarely been described. We have observed at least one case of multiple members of an Iranian Jewish family developing HAM (D. Meytes et al., unpublished), and this has been reported by other investigators. This would suggest that either differences in host genetic make-up and susceptibility or differences in viral isolates may account for familial HAM. It is important to note that the association between HAM and HTLV-I was made serendipitously. Quite possibly, if the link to HAM had been described first, no search for HTLV-I association with malignancy would have been made. This would suggest that other viruses that may not be associated in investigators' minds with development of malignancy may be candidates for potential oncogenic roles. Good candidates would be viruses with trans-acting transcriptional proteins, such as members of the herpes family, adenovirus, and/or other retroviruses.

DISCUSSION

Over a decade following initial identification of the role of the retrovirus in human disease. HTLV-I remains as the only example for human retroviral leukemogenesis. Several lessons pertinent to the search for leukemogenic viruses can be derived from the HTLV model. First, leukemia may be the rare consequence of infection with a relatively common virus. Second, a long latency period may be involved following acquisition of infection, and the leukemogenic process may involve multiple steps beyond initial infection. Third, the mechanism by which the virus may predispose to development of leukemia may be different from that seen with other retroviruses, and may involve transacting viral genes, either regulators of DNA transcription or regulators of viral mRNA expression. Finally, the role of HTLV-I as an etiologic agent in both HAM and ATLL suggests that viruses not thought to have oncogenic properties may on occasion predispose to malignancy. The overall lesson from the HTLV experience is that when suitable or suggestive epidemiologic factors are found, such as ethnic, geographic and/or familial clustering in the absence of a known inheritance pattern, a search for underlying viruses should be initiated. The search would involve removal of malignant tissue, when possible, for culture in vitro and a search using available molecular and serologic probes. The success in identifying HTLV-II by virtue of limited nucleic acid homology and antigenic similarity to HTLV-I suggests that new retroviruses can be identified. The demonstrated crossreactivity between HTLV-I and -II suggests that any search should be accompanied by rapid-isolation of nucleic acid probes for viral sequences of interest, so that crossreactive

entitites can be discerned. A fresh look should be taken using newly available probes as a means of determining viral clonality, particularly for DNA viruses such as herpesviruses to assess whether a particular malignant tissue has arisen from a single virally infected cell. Furthermore, scrutiny of viruses already known to be widespread in the population may prove fruitful, as already appears to be the case for EBV and a subset of Hodgkin's disease. A re-duplication of such efforts will determine whether new retroviruses with oncogenic potential will be identified in man in the upcoming decade, or whether HTLV will remain an isolated if fascinating example of retroviral leukemogenesis in man.

Acknowledgements. The authors are grateful to W. Aft for assistance in preparation of the manuscript. JDR is supported by NIH grants CA01314 and CA53632, and by the US Army Medical Research and Development Command (US AMRDC); YLD is supported by the Rashi Foundation, the US AMRDC and the Doron Foundation; and ACB is supported by an NIH Physician-Scientist Award the the Leukemia Society of America.

REFERENCES

- 1. Poiesz BJ, Ruscetti FW, Gazdar AF, Bunn PA, Minna JD, Gallo RC. Detection and isolation of type C retrovirus particles from fresh and cultured lymphocytes of a patient with cutaneous T-cell lymphoma. Proc Natl Acad Sci USA 1980:77:7415-7419.
- Yoshida M, Seiki M, Yamaguchi K, Takatsuki K. Monoclonal integration of human T-cell leukemia provirus in all primary tumors of adult T-cell leukemia suggests causative role of human T-cell leukemia virus in the disease. Proc Natl Acad Sci USA 1984;81:2534-2537.
- Kalyanaraman VS, Sarngadharan MG, Robert-Guroff M, Miyoshi I, Blayney D, Golde D, Gallo RC. A new subtype of human T-cell leukemia virus (HTLV-II) associated with a T-cell variant of hairy cell leukemia. Science 1982;218:571-573.
- 4. Uchiyama T, Yodoi J, Sagawa K, Takatsuki K. Uchino H: Adult T-cell leukemia: Clinical and hematologic features of 16 cases. Blood 1977;50:481-492.
- Blattner WA. Epidemiology of HTLV-I and associated diseases. In: Blattner WA, ed. Human Retrovirology: HTLV. New York: Raven Press Ltd, 1990, pp. 251-263.
- Blattner WA, Nomura A, Clark JW, Ho GYF, Nakao Y, Gallo R, Robert-Guroff M. Modes of transmission and evidence for viral latency from studies of human T-cell lymphotropic virus type I in Japanese migrant populations in Hawaii. Proc Natl Acad Sci USA 1986:83:4895-4898.
- 7. Tajima K, Ito SI, and the Tsushima ATL group. Prospective studies of HTLV-I and associated disease in Japan in human retrovirology. In: Blattner WA, ed. HTLV, New York: Raven Press, 1990.
- Saxon A, Stevens RH, Quan SG, Golde DW. Immunologic characterization of hairy cell leukemias in continuous culture. J Immunol 1978;120:777-782.
- Rosenblatt JD, Golde DW, Wachsman W, Jacobs A, Schmidt G, Quan S, Gasson JC, Chen ISY. A second HTLV-II isolate associated with atypical hairy-cell leukemia. New Engl J Med 1986;315:372-375.
- Rosenblatt JD, Giorgi JV, Golde DW, Ben Ezra J, Wu A, Winberg CD, Glaspy J, Wachsman W. Chen ISY. Integrated HTLV-II genome in CD8⁺ T-cells from a patient with 'atypical' hairy-cell leukemia: evidence for

distinct T- and B-cell lymphoproliferative disorders. Blood 1988,71:363-369.

29

30.

31

3.

3

- Lee HH, Swanson P, Rosenblatt JD, Chen ISY, Sherwood WC, Smith DE, Tegtmeier GE, Fernando LP, Fang CT, Osame M, Kleinman SH. Antibody screening of U.S. blood donors reveals similar prevalence of HTLV-I and HTLV-II infection in association with different risk factors. Lancet 1991;337:1435-1439.
- Lee H, Swanson P, Shorty VS, Zack JA, Rosenblatt JD, Chen ISY. High rate of HTLV-II infection in seropositive IV drug abusers from New Orleans. Science 1989:244:471-475.
- Robert-Guroff M, Weiss SH. Giron JA. Jennings AM, Ginzburg HM, Margolis IB. Blattner WA, Gallo RC. Prevalence of antibodies to HTLV-I. -II, and -III in intravenous drug abusers from an AIDS endemic region. J Am Med Assoc 1986:255:3133-3137.
- 14. Brousset P, Chittal S, Schlaifer D, et al. Detection of Epstein-Barr virus messenger RNA in Reed-Sternberg cells of Hodgkin's disease by *in situ* hybridizarion with biotinylated probes on specially processed modified acetone methyl benzoate xylene (ModAMeX) sections. Blood **1991**;77:1781-1786.
- Pallesen G, Hamilton-Dutoit SJ, Rowe M, Young LS. Expression of Epstein-Barr virus latent gene products in tumour cells of Hodgkin's disease. Lancet 1991; 337:320-322.
- Uccini S. Monardo F. Stoppacciaro A. et al. High frequency of Epstein-Barr virus genome detection in Hodgkin's disease of HIV-positive patients. Int J Cancer 1990;46:581-585.
- Herbst H. Niedobitek G, Kneba M, et al. High incidence of Epstein-Barr virus genomes in Hodgkin's disease. Am J Pathol 1990;137:13-18.
- 18. Kubonishi I, Equchi T, Kanzaki T, et al. EBV and Hodgkin's cells. Br J Haematol 1990;75:286-287.
- 19. Weiss LM. Movahed LA, Warnke RA, Sklar J. Detection of Epstein-Barr viral genomes in Reed-Sternberg cells of Hodgkin's disease. New Engl J Med 1989;320:502-506.
- Staal SP, Ambinder R, Beschorner WE, et al. A survey of Epstein-Barr virus DNA in lymphoid tissue: Frequent detection in Hodgkin's disease. Am J Clin Pathol 1989;91:1-5.
- Weiss LM, Warnke RA, Sklar J. Clonal antigen receptor gene rearrangements and Epstein-Barr viral DNA in cissues of Hodgkin's disease. Hematol Oncol 1988;6:233-238.
- Sidi Y, Meytes D, Shohat B, Fenig E, Weisbort Y, Lee H, Pinkhas J, Rosenblatt J. Adult T-cell lymphoma in Israeli patients of Iranian origin. Cancer 1990; 65:590-593.
- 23. Meytes D. Schochat B, Lee H. Nadel G. Sidi Y. Cerney M, Swanson P, Shaklai M, Kilim Y, Elgat M, Chin E, Danon Y, Rosenblatt JD. A serological and molecular survey for HTLV-I infection in a newly identified high-risk group. Lancet, 1991. in press.
- 24. Linet MS. Devesa SS. Descriptive epidemiology of childhood leukaemia. Br J Cancer 1991:63:424-429.
- Alexander FE, Ricketts TJ, McKinney PA, Cartwright RA. Community lifestyle characteristics and risk of acute lymphoblastic leukaemia in children. Lancet 1990;336:1461-1465.
- Okpala IE, Abayomi NA, Gevao SM, et al. Changing patterns of acute lymphoblastic leukaemia in Nigeria. Tokai J Exp Clin Med 1989;14:301-307.
- 27. Editorial. Childhood leukaemia: An infectious disease? Lancet 1990:336:1477-1479.
- 28. Kinlen LJ. Infective cause of childhood leukaemia.

ROSENBLATT ET AL.

22

Lancet 1989:i:378-379.

- 29. Lilleyman JS. Leukaemia 'outbreaks'. 1988; Lancet ii:1021.
- Till MM, Hardisty RM, Pike MC, et al. Childhood leukaemia in greater London: A search for evidence of clustering. Br Med J 1967;iii:755-758.
- Smith PG. Spatial and temporal clustering. In: Schottenfeld P, Fraumeni JF. eds. Cancer epidemiology and prevention, Philadelphia: Saunders, 1982, pp. 391-408.
- 32. Seiki M, Eddy R, Shows TB, Yoshida M. Nonspecific integration of the HTLV provirus genome into adult T-cell leukaemia cells. Nature **1984**.309:640-642.
- 33. Seiki M, Hattori S, Hirayama Y, Yoshida M. Human adult T-cell leukemia virus: Complete nucleotide sequence of the provirus genome integrated in leukemia cell DNA. Proc Natl Acad Sci USA 1983;80:3618-3622.
- 34. Slamon DJ, Shimotohno K, Cline MJ, Golde DW, Chen ISY. Identification of the putative transforming protein of the human T-cell leukemia viruses HTLV-1 and HTLV-II. Science 1984;226:61-65.
- 35. Lee TH, Coligan JE, Sodroski JG, Haseltine WA, Salahuddin SZ, Wong-Staal F, Gallo RC, Essex M. Antigens encoded by the 3'-terminal region of human T-cell leukemia virus: Evidence for a functional gene. Science 1984:226:57-61
- 36. Sodroski J, Rosen C, Goh WC, Haseltine W. A transcriptional activator protein encoded by the x-lor region of the human T-cell leukemia virus. Science 1985;228:1430-1434.
- Cann AJ, Rosenblatt JD, Wachsman W, Shah NP, Chen ISY. Identification of the gene responsible for human T-cell leukemia virus transcriptional regulation. Nature 1985:318:571-574.
- Greene WC. The human interleukin 2 receptor. Normal and abnormal expression in T cells and in leukemias induced by the human T-lymphotropic retroviruses. Ann Intern Med 1986;105:560-572.
- 39. Siekevitz M. Feinberg MB, Holbrook N, Wong-Staal F, Greene WC. Activation of interleukin 2 and interleukin 2 receptor (Tac) promoter expression by the trans-activator (tat) gene product of human T-cell leukemia virus, type I. Proc Natl Acad Sci USA 1987;84:5389-5393.
- Cross SL. Feinberg MB. Wolf JB, Holbrook NJ, Wong-Staal F. Leonard WJ. Regulation of the human interleukin-2 receptor alpha promoter: Activation of a nonfunctional promoter by the transactivator gene of HTLV-I. 1987; Cell 49:47-56.
- Nimer SD, Gasson JC, Hu K, Smalberg I, Williams JL, Chen ISY, Rosenblatt JD. Activation of the GM-CSF promoter by HTLV-I and -II *tax* proteins. Oncogene 1989;4:671-676.
- 42. Nyborg JK, Dynan WS, Chen ISY, Wachsman W. Binding of host-cell factors to DNA sequences in the long terminal repeat of human T-cell leukemia virus type I: Implications for viral gene expression. Proc Natl Acad Sci USA 1988;85:1457-1461.
- Ohtani K, Nakamura M, Saito S, Noda T, Ito Y, Sugamura K, Hinuma Y. Identification of two distinct elements in the long terminal repeat of HTLV-I responsible for maximum gene expression. Eur Mol Biol Org (EMBO) J 1987;6:389-395.
- Leung K, Nabel GJ. HTLV-I trans-activator induces interleukin-2 receptor expression through an NF_KB-like factor. Nature 1988;333:776-778.
- 45. Ruben S, Poteat H, Tan TH, Kawakami K, Roeder R, Haseltine W, Rosen CA. Cellular transcription factors and regulation of IL-2 receptor gene expression by HTLV-I tax gene product. Science 1988;241:89-92.

- 46. Grassmann R. Dengler C, Muller-Fleckenstein I, Fleckenstein B. McGuire K. Dokhelar M-C, Sodroski JG, Haseltine WA. Transformation to continuous growth of primary human T lymphocytes by human T-cell leukemia virus type I X-region genes transduced by a Herpesvirus saimin vector. Proc Natl Acad Sci USA 1989; 86:3351-3355.
- 47. Nerenberg M, Hinrichs SH, Reynolds RK, Khoury G, Jay G. The *tat* gene of human T-lymphotropic virus type I induces mesenchymal tumors in transgenic mice. Science 1987:237:1324-1329.
- Nerenberg MI, Minor T, Price J, Ernst DE, Shinohara T, Schwarz H. Transgenic thymocytes are refractory to transformation by human T-cell leukemia virus type I tax gene. J Virol 1991;65:3349-3353.
- 49. Watanabe T, Yamaguchi K, Takatsuki K, Osame M, Yoshida M. Constitutive expression of parathyroid hormone-related protein gene in human T cell leukemia virus type I (HTLV-I) carriers and adult T cell leukemia patients that can be trans-activated by HTLV-I tax gene. J Exp Med 1990;172:759-765.
- 50. van Lohuizen M. Veibeek S. Scheijen B. *et al.* Identification of cooperating oncogenes in E_{μ} -myc transgenic mice by provirus tagging. Cell **1991**:65:737-752.
- Haupt Y, Alexander WS. Barri G. et al. Novel zinc finger gene implicated as myc collaborator by retrovirally accelerated lymphomagenesis in E_µ-myc transgenic mice. Cell 1991:65:753-763.
- 52. Kiyokawa T, Seiki M, Iwashita S, Imagawa K, Shimizu F, Yoshida M, p27^{xIII} and p21^{xIII}, proteins encoded by the pX sequence of human T-cell leukemia virus type I. Proc Natl Acad Sci USA 1985:82:8359-8363.
- Shima H. Takano M. Shimotohno K. Miwa M: Identification of p26^{xb} and p24^{xb} of human T-cell leukemia virus type II. FEBS Lett 1986:209:289-294.
- 54. Green PL, Xie Y, Chen ISY. The Rex proteins of HTLV-II differ by serine phosphorylation. J Virol 1991;65:546-550.
- Rosenblatt JD. Cann AJ. Slamon DJ, Smalberg IS, Shah NP, Fujii J, Wachsman W, Chen ISY. HTLV-II transactivation is regulated by two overlapping nonstructural genes. Science 1988;240:916-919.
- Black AC, Chen ISY, Arrigo SJ, Ruland CT, Chin E, Allogiamento T. Rosenblatt JD. Different cis-acting regions of the HTLV-II 5' LTR are involved in regulation of gene expression by Rex. Virology 1991;181:433-444.
- 57. Black AC, Ruland CT, Yip MT, Luo J, Kalsi A, Quan E, Tran B, Chen ISY, Rosenblatt JD. HTLV-II Rex binding requires a specific RNA secondary structure and intact splice donor. J Virol, in press.
- 58. Yip MT, Dynan WS. Green PL. Black AC, Arrigo SJ, Torbati A, Heaphy S, Ruland C, Rosenblatt JD, Chen ISY. HTLV Rex protein binds specifically to RNA sequences of the HTLV LTR but not the HIV-I RRE, J Virol 1991;65:2261-2272.
- 59. Rimsky L. Hauber J. Dukovich M. Malim M. Langlois A, Cullen B, Greene W. Functional replacement of the HIV-1 rev protein by the HTLV-I rex protein. Nature 1988;335:738-740.
- Gessain A, Vernant JC, Maurs L. Barin F, Gout O, Calender A, de The G. Antibodies to human T-lymphotropic virus type-1 in patients with tropical spastic paraparesis. Lancet 1985;ii:407-409.
- 61. Saxton EH. Lee H. Swanson P. Chen ISY, Ruland C, Chin E, Aboulafia D. Delamarter R, Rosenblatt JD. Detection of human T-cell leukemia/lymphoma virus type I in a transfusion percipient with chronic myelopathy. Neurology 1989;39:841-844.

Immunoglobulin Prophylaxis against HTLV-I in a Rabbit Model

I. Miyoshi¹, N. Takehara¹, T. Sawada¹, Y. Iwahara¹, R. Kataoka¹, D. Yang², and H. Hoshino²

¹Department of Medicine, Kochi Medical School, Kochi 783, Japan, and ²Department of Hygiene, Gunma University School of Medicine, Gunma 371, Japan

We have investigated the protective effect of human T-cell leukemia virus I (HTLV-I) immune globulin (HTLVIG) against HTLV-I in rabbits. HTLVIG containing 77 mg/ml of IgG was prepared from pooled plasma from seropositive healthy persons. In the first experiment, four groups (A, B, C, and D) of three rabbits were transfused with 5 ml blood from an HTLV-Iinfected rabbit. Groups A, B, and C were infused 24 h later with 10, 5, and 2 mi HTLVIG, respectively, while group D was infused with 10 mi HTLVIG 48 h later. Seroconversion for HTLV-I occurred in none of group A, one of group B, and all of groups C and D after 2-5 weeks. In the second experiment, four litters (E, F, G, and H) born to another virus-infected rabbit and consisting of 7, 5, 7, and 7 newborns, respectively, were used. Litters E and H were allowed to grow normally as controls, while litters F and G were given intraperitoneal inoculation of 3 ml/kg of HTLVIG weekly four times until weaning. Although three of litters E and H each seroconverted after 5-8 weeks, none of litters F, and one of litter G became antibody-positive after 10 weeks. Presence or absence of HTLV-I infection in all these animals was confirmed by transfusion assay or gene amplification. These results indicate that passive immunization protects rabbits against blood- and milk-borne transmission of HTLV-I.

INTRODUCTION

A rabbit model of human T-cell leukemia virus I (HTLV-I) infection has been established, in which the virus was shown to be transmissible not only by blood transfusion (1,2) but also from dam to offspring via milk (3,4). In the blood transfusion experiment, as little as 0.01 ml blood from a virus-infected rabbit was capable of transmitting HTLV-I (2). Furthermore, milk or semen lymphocytes from seropositive healthy persons transmitted HTLV-I when inoculated intravenously into rabbits (5). This animal model, therefore, provided a unique opportunity to study the protective effect of passive immunization against HTLV-I (2,6). In the present experiment, immunoglobulin prophylaxis against blood- and milk-borne transmission of HTLV-I was further explored.

MATERIALS AND METHODS

Rabbits

Japanese white rabbits, weighing about 3 kg, purchased from a commercial breeder were used.

Detection of Antibodies to HTLV-I

Blood samples were taken from rabbits at intervals of 1-2 weeks and sera were titrated for HTLV-I antibodies by indirect

Correspondence to: Isao Miyoshi. MD, Department of Medicine, Kochi Medical School, Kochi 783, Japan.

LEUKEMIA © 1992 Macmillan Press Lid

24

immunofluorescence against the MT-2 cell line as described previously (2). The presence or absence of immunoglobulin G (IgG) antibodies was verified by Western blot using a MT-2 lysate as antigen. Sera were also tested for IgG and immunoglobulin M (IgM) antibodies by enzyme-linked immunosorbent assay (ELISA) against disrupted HTLV-I virions according to the manufacturer's instructions (Eisai. Tokyo). Neutralizing antibodies were assayed against vesicular stomatitis virus (VSV) bearing envelope antigens of HTLV-I as previously described (7).

HTLV-I Immune Globulin (HTLVIG)

HTLVIG containing 77 mg/ml of IgG was prepared from pooled plasma from seropositive healthy persons by the method of polyethylene glycol fractionation (8). The preparation had an immunofluorescence anti-HTLV-I titer of 1:5120 and a VSV (HTLV-I) pseudotype neutralizing antibody titer of 1:6250.

Transfusion Assay

To ascertain the status of HTLV-I infection. 20 ml of blood obtained from experimental rabbits were transfused into normal rabbits. Seroconversion of the recipient rabbits indicated a virus carrier state of the donor rabbits.

Polymerase Chain Reaction (PCR)

DNA extracted from peripheral blood mononuclear cells was analyzed for the presence of HTLV-I sequences by the method of Kwok *et al.* (9). DNA, 1 μ g, was subjected to 40 cycles of denaturation followed by annealing and extension. Oligonucleotide primers at 7341-7360 and 7460-7411 corresponding to the pX region of HTLV-I were used. Amplification was performed using a thermostable DNA polymerase on an automated DNA Thermal Cycler (Perkin-Elmer/Cetus, Norwalk, CT). The amplified products were electrophoresed on 6% polyacrylamide gels, transferred to nylon membranes, and hybridized with a ³²P end-labeled probe at 7364-7383.

RESULTS

Passive Immunization against Blood-borne Transmission of HTLV-I

Four groups (A, B, C, and D) of three rabbits were first transfused with 5 ml of blood from an HTLV-Iinfected rabbit. Groups A, B, and C were infused 24 h later with 10, 5, and 2 ml HTLVIG, respectively, while group D was infused with 10 ml HTLVIG 48 h later.

Seroconversion for HTLV-I occurred in none of group A, one of group B, and all of groups C and D after 2-5 weeks (Figure 1). All five rabbits which were protected from seroconversion remained seronegative during an observation of six months. Sera taken immediately after infusion of HTLVIG showed anti-HTLV-I titers of 1:320 for groups A and D, 1:80 for group B, and 1:20 for group C. The VSV (HTLV-I) pseudotype neutralizing titers of these sera were 1:1250



DAVID W. GOLDE, M.D.

1533

identified populations of calcitation gene-related peptide (CORP)immunitestatic actions williaman with Hum Rev 1987, 414: 143–18

- 13 Wallengren J, Ekman F. Sundler F. Occurrence and distribution of neuropeptides in the human iden. An innumuentic hemical and immunichemical study on normal skin and blotter fluid from inflamed skin. Acta Rev. Voleccol 516(d), 1987;67:185-92.
- 14 Daligaard CJ, Jembech J, Stains W, et al. Calcitonin gene-related pepide-like immunereactivity in nerve fibres in the human white Relation to fibres emissiong substance. Proc simultation and substance intestinal polyppinde-like immuniveractivity. *Histocksongus* 1989, 91: 35-38.
- Pastriwski W, Fistman JC. Some effects of calcitonin gene-related peptide in human skin and on histamine release. Br J Dynamid 1986, 114: 37–46.
- Struchers AD, Brissin AJJ, MacDanield DWR, et al. Human calculosin gene-related peptide: a potent endogenous vasodilator in man. *Clin. Sci.* 1986, 70: 306-93.
- 17 Howden CW, Legue C, Gavin K, Collie L, Rubin PC Hachaudynamic efforts of intravenous human calentonin gene-related periode in min

1 Jun S. 1995, 74: 413-18

- 18 Builter (B. Foreman J, Reavley C, O'Shaughnessey D, Dovid PM (alciumin gene-related populat in the treatment of severe Ravinaud's phenemen-in. *Hi J Thomatic* 1989, 121: (suppl 34): 43-44.
- "10 Bunker (3) Keackly C. O'Shaughnessy D. Ehmed PAL Intracendus calentoning true-related peptide in subset Raynaud's phonomenon. Bi J. Rhenmatrik 1940, 29: (suppl 2), 1.
- Shawker S, Duckerson C, Harleman B, Brown All Selective supravenum to calcinonin-gene-related peptide in the hands in Raimaud spheriomenum Lance 1989, ii 1354-56.
- Runker CH, Foreman J, Dunid PM. Digital outaneous vascular responses to histamine, compound 48/80 and neuropeptides in normal subjects and Raynaudis phenomenon. J Invisi Demontol (in press)
- 22 Dovid PAL Bunker GB, Bult HA, et al. Raynaud's plictromenum, colotioningene-related peptide, endishelin, and curanesus vasculature Lancer (990, 336: 1014).
- 23 Zamma MR, O'Brien RF, Russerfind RB, Weil JY, Scrum endochelin-t emicentrations and cold providence in primary Raynaud's phenomenon Januer 1990, 336: 1144-47.

Serological and molecular survey for HTLV-I infection in a high-risk Middle Eastern group

DINA MEYTES BATYA SCHOCHAT HELEN LEE GIORA NADEL YECHEZKE! SIDI MICHAEL CERNEY PRISCILLA SWANSON MATHTYAHU SHAKLAT YAEL KILIM MAYA ELGAT EVA CHIN YEHUDA DANON JOSEPH D. ROSENBLATT

To define the extent of human T-cell leukaemia virus (HTLV-I) infection among a group of Jewish immigrants to Israel with an increased frequency of adult T-cell leukaemia, various serological and molecular screening methods, including enzymelinked immunosorbent assay (ELISA) for anti-HTLV-I, ELISA for antibody to recombinant HTLV-I p40tax protein, and molecular detection of infection by polymerase chain reaction (PCR) amplification of HTLV-I proviral DNA from peripheral blood mononuclear cell DNA, were used. By HTLV-I ELISA the overall rate of infection was 12% (24 of 208) among immigrants from Khurusan, northeastern Iran; no HTLV-I carriers were detected among 111 unselected Jewish immigrants from other parts of Iran. There was unexplained clustering of HTLV-I infection within a cohort of 32 elderly women of similar geographic origin in a home for old people-14 were seropositive by ELISA and 19 of 29 were positive by PCR. The findings in this newly identified high-risk population suggest that in addition to ELISA, other screening techniques may be required to detect all carriers in high-risk populations.

Lancer 1990 336: 1533-35

Introduction

Human T-cell leukacmia virus type 1 (HTLV-1) infection has been described in southern Japan, the Carihbean basin, and the northern parts of South America, and in certain high-risk groups, such as intravenous drug abusers in the United States.^{1,7} Previous reports of HTLV-1 infection among Ethiopian Jews in Israel were not confirmed.^{4,9} During the past 4 years, sporadic cases of adult T-cell leukacmia linked to HTLV-1 have been reported in Israel¹¹¹¹ and 1 of the 5 latest cases were among immigrants to Israel who originated from the city of Mashad in northeastern Iran ¹¹ Because of these findings, we undertook a systematic survey of Iranian Jews in Israel, focusing on immigrants with links to Mashad.

Subjects and methods

Blood samples from Israeli blood donors of Iranian origin were obtained from the Israeli Magen David Adom Blood Services Center, Jel Avis. The criterion for classification as an Iranian control was that the country of birth of the blood donor or at least one of his or her parents was Iran. Blood seriples were collected on three occasions from residents of a Mashadi home for elderly women in the Tel Avis area and from three Mashadi community synagopus in the cities of Briel Brak and Tel Avis. Samples were classified as Mashadi if the donor or at least one of his or her parents originated from Mashad, Iran. 20 samples from patients on long-term haemodialysis, 8 from patients with T-cell malignant diswders other than adult T-cell Rukaemia, and 12 from Ethiopian Jewish immigrants were also included.

Servivgical screening was done for HTLV-I antibodies on serum or plasma samples by means of an enzyme-linked immunoscreent assay (ELISA: Abbutt Laboratories). Confirmatory western blotting and or radioinmunoprecipitation assay (RIPA) with aulphur-35-labelled nicthionine HTLV-I-infected HUT 102B

> ter en el composition de la composition La composition de la c

ADDRESSES Edith Wolfson Hospital and Tal Aviv University. Sackler School of Medicine. Holon (D. Meyres, MD. M. Eiget BSc): Beilinson Medical Center, Petach Tikva (B. Schochet, Ph.D. Y. Sidu, MD. M. Shakar MD. Y. Kilm, MSr. Y. Danon, MD); Ministry of Health, Tel Aviv. Isreel (G. Nadol. MO); Diagnostic Division, Abbott Laboratories, North Chicago, Illinois (H. Lee. Ph.D. M. Ceney, BSc. P. Swamon, MSr.); and Division, of Hematology-Oncology, Department of Medicine. UCL'A. School of Medicine. Los Angeles, California, USA (E. Chin, BSc. J. D. Rosenhlatt. MO) Consepondence to Dr. Meytes, Department of Haematology Edith Wolfson Hospita', Tel Aviv. Israel 58100

		LISA and m blot	Anti-p40tax ELISA	
_	No tested	No (%) positive	No tested	No (%) positive
Mashadi Jows	208	24 / 12 15 1	127	1219-21
Other Ininian Jews	111	0	20	0
Ethiopian Jews	12	0'	12	0
Haemodialysis patients Patients with T-cell	20	n	20	0
malignant disorders		0	ND	

Is sate were also done.¹³ Samples positive in the ELISA were tested by both confirmatory methods. Antibodies to HTLV-1 p40*tax* were measured by means of an ELISA with recombinant p40*tax* as antigen on the solid phase (polystyrene beads) (Abbott). HTLV-1 seropositive infected samples with known reactivity against p40*tax* on RIPA were used as positive controls, and 4 samples negative for HTLV-1 on ELISA and western blot as negative controls. Samples were scored as positive for p40*tax* if the optical density exceeded 4.5 times the mean negative control value. The polymerase chain reaction (PCR) was used to amplify HTLV-1 sequences of DNA from peripheral blood mononuclear cells with primers to a 159 bp segment contained within the *tax*, rer gene as previously described.¹²

Results

The extensive social and ethnic tics between Jews of Mashadi origin allowed selective sampling of this population within Israel. On repeated assays, 24 of 208 (11.5%) Mashadi serum samples were positive by ELISA for antibodies to HTLV-I and were confirmed by western blotting. In contrast, none of the 151 control samples was positive by ELISA (table).

In an effort to survey the oldest members of the Mashadi population, we tested 32 unrelated elderly Mashadi women living in an old people's home in Tel Aviv. 14 (44%) were seropositive, a rate three times higher than that in the general Mashadi population. At our first sample collection in 1988, 12 (52%) of 23 long-standing residents of the home were seropositive; 2 of 9 new residents surveyed a year later were seropositive. Of 26 Mashadi women older than 60 years who lived elsewhere in Israel only 3 (12%) were seropositive.

To determine whether serological assays had detected all infected subjects, we carried out PCR amplification of HTLV-I DNA from 29 of the residents of the old people's home. 19 were positive for HTLV-I sequences: 14 of these were seropositive, 4 were seronegative, and 1 was seronegative by standard ELISA but reactive in the anti-p40*tax* ELISA. A second PCR on repeat samples from 10 of these women showed concordant results in 6 of 6 PCR-positive and 2 of 3 PCR-negative cases; 1 woman originally positive on PCR was negative on the second sample, and 1 woman originally PCR-negative was faintly positive on her second sample and had antibodies to p40*tax* by ELISA. This unusual clustering of HTLV-I infection within the old people's home suggested acquisition of infection within the institution.

In addition to antibudies to virion components, we assayed for antibudies to the non-structural p40*tax* protein. Of 128 Mashadi samples tested 12 had absorbance levels 4.5 or more times those of the negative control and were judged positive (table). 103 samples were negative by both assays. 8 samples were scropositive for both anti-HTLV-I and anti-p40*tax*; 13 samples were positive for anti-HTLV-I and negative for anti-p40*tax*; and 4 were positive for anti-p40*tax* but repeatedly negative for anti-HTLV-I by ELISA and western blot. 3 of the latter group of samples were negative on PCR, and the other, from a resident of the old people's home, was positive. All 4 were negative by western blotting and by RIPA. Therefore, the reason for the high titres in three samples in the anti-p40tax ELISA is unclear.

Direct comparison of anti-p40tax ELISA results and those of the RIPA was done for 28 subjects. 8 samples were positive for anti-p40tax by both RIPA and ELISA; 1 with traces of antibody by RIPA was negative by ELISA; the 3 samples mentioned previously were negative by RIPA but repeatedly positive by ELISA; and 16 were negative by both tests. The usefulness of the anti-p40tax ELISA in detecting true additional HTLV-1-infected seronegative infection was corroborated by other means of detection in only 1 of the 4 questionable cases.

The finding of a higher rate of seropositivity among elderly Mashadi women than in the general Mashadi population suggested a potential for gradual development of seropositivity with age, as has been reported by others. Because of the possibility that serological screening may not detect all positive subjects' we carried out HTLV-I-specific PCR on DNA from 68 randomly selected Mashadi blood samples. 5 of 7 known seropositive samples were positive in the PCR assay, and 61 seronegative samples were all negative by PCR. Hence, PCR did not increase sensitivity of detection within this sample above that achieved with serological assays alone. All PCR-positive samples contained HTLV-I and not HTLV-II, as shown by discriminatory PCR (data not shown).

Discussion

We have identified a high risk of HTLV-I infection in Iranian Jews originating from the city of Mashad in Khurusan, northeastern Iran. This group seems to have an overall infection rate of about 12%, though the rate was much higher among residents of a Mashadi home for old people. The reason for this clustering is unclear, although the findings suggest acquisition of infection within the institution. The overall level of infection among other Iranian Jews seems to be substantially lower than that among Mashadis. The explanation may be geographic, and Mashad may be within a previously unrecognised endemic region. Since we did not test non-Jewish Mashadis, this possibility cannot be excluded.

The unique history of the Mashadi community may help to explain their high rate of HTLV-I infection. In 1839, the Jewish community in Mashad was subjected to a series of violent attacks and forced to convert to Islam, though the majority of the community continued to practise Judaism envertly.¹¹ To safeguard the community secret necessitated a very high rate of intermarriage among community members. Members of this community continued to marry cluster relatives over the next 150 years. Markers of consanguinity are high among Mashadi Jews—for example, over 20% have a deficiency of glucose-6-phosphate dehydrogenase (D. M., unpublished). The chance introduction of HTLV-I infection into the community could have resulted in rapid transmission of the virus by sexual means and by breastfeeding.

An estimated 5000-6000 Mashadis now live in Israel. This is the first Israeli etimic group identified as having a high rate of HTLV-I infection. Members of the Mashadi Jewish community also migrated to the USA and parts of

Europe. We predict that the rate of infection among these migrants would be similar to that of the Israeli cohort.

In this study, we used several methods to detect HTLV-1 infection. Our rate of anti-p-f0tax scropositivity in carriers was somewhat lower than that found by others arning healthy HTLV-1 carriers.^{114 In} Scrum samples from 4 subjects were positive only for anti-p40tax antibodies. Independent evidence of infection was obtained by PCR in only 1 of the 4. Thus, detection of anti-p40tax antibodies did not appreciably add to the estimate of the rate of infection. Our findings on the use of PCR suggested that in a high-risk population, such as the old people's home we studied or in families of HTLV-1 carriers, PCR would increase the number of infected individuals above that detected by scrological means. The usefulness of PCR as a screening assay in appropriate settings requires further study.

We thank Prof Erresto Lubin for legistical support. The study was supported by (to J. D. R.) the Leakerna Research Foundation, the Rishi Foundation, NHH grants CA01314 and CA52410-01, and a perunal contribution from Edward and Dolly Loss; (to D. M.) the Chief Scienteria Bureau, Atinistry of Health, Israel, and the Israel Cancer Research Fund, and (to Y. D. and J. D. R.) a grant from the US Army Atelical Research and Development Command. J. D. R. was a visiting Fulbright scholar to the Sackler Schull of Atelica.

REFERENCES

- Himuma Y, Komoda H, Chosa T, et al. Antibodies to adult 1-cell leukema-virus-associated antigen: A ITA' in sera from patients with A TL and controls in Japan'a nation-wide serie-pidemiologic study. *Int.* J Cancer 1982, 29: 631–55.
- Himma Y. Sersepidemiology of adult. Leell leukemia virus (HTLV-37 A ULV): origin of virus carners in Japan. AUDS Rev. 1986; 2: 517–22.

- Biggar RJ, McBye M, Sarin PS, et al. ELISA HTLV recoverate antibody nuccivity associated with malaria and annuane complexes in healthy Africans. *Lanuer* 1985; ii. 520-23.
- 4 Blattner WA, Sasinger CA, Cleft J, et al. Human T-cell leukaemia/ hypohema virus-associated hypohemeticular neoplasie in Jamasca. Lancet 1983, ii: 61-61.
- Buim PA Jr. Schechter GP, Jaffe E, et al. (Jinical course of retrovirusassociated adult T-cell hymphome in the United States. N Engl J Med. 1993; 309: 237-64.
- Rohert-Gunvill M, Weiss SH, Girun JA, et al. Prevalence of antibodies to HTLLV-1, -11, and -111 in intravenous drug abusers from an AIDS endemic regim. JANIA 1986; 255: 3133-37.
- Ehrlich GD, Glaser JB, LaVigne K, et al. Prevalence of human T-cell leukernia/hymphoma virus (HTLV) type II infection among high-risk individuals type specific identification of HTLVs by polymerase chain reaction. Blowd 1989; 74: 1658-64.
- 8 Ben-Ishai Z, Hass M, Triglia D, et al. Human T-cell lymphotrophic virus type-1 antibudics in Falashas and other ethnic groups in Israel. Nature 1985, 315: 665-66.
- 9 Karpes A, Maayon S, Rar R. Lack of antibodies to adult T-cell leukaemin virus and to AIDS virus in Isracli Falavias. Nature 1986; 319: 794.
- 10 Lever J, Langiuitz P, Tran H, et al. HTLV-1 associated T-cell leukimia hymphoma in Israel Iar J Aled Sci 1988; 24: 397-400.
- Sidi Y, Meyter D, Shochar B, et al. Adult T-cell lymphoms in Israeli patients of Iranian origin. Cancer 1990; 65: 590-93.
- 12 Lee H, Swanson P, Shorty VS, Zack JA, Rosenblatt JD, Chen ISY. Highrate of HTLV-II infection in scropositive IV drug absuers from New Orleans. Science 1989; 244: 471–75.
- 11 Encyclopaedra Judaica 1972, 11:1399-400
- Ehrlich GD, Glaser JR, Abbrit MA, et al. Detection of anti-HTLV-I Tax antibudies in HTLV-1 enzyme linked immunosorbent assaynegative individuals. Blood 1989, 74: 1064-72.
- 15 Yukata L, Chu MJ, Lichibana N, et al. The prevalence of antibody to p42 of HTLV-1 among ATLL patients in companions with healthy carriers in Japan. Int J Cancer 1989;43: 970-71
- 16 Kamihara S, Toriya K, Amagasala T, et al. Antibedies against p40tax gene product of human 1-hymphotrophic virus type 1 under various conditions of HTLV-1 infection. Jpn J Concer Res 1989; 80: 1066-71.

Quinine-induced disseminated intravascular coagulation

RUTH L. SPEARING CHRISTINE M. HICKTON PETER SIZELAND ANTHONY HANNALL ROSS R. BAILEY

Recurrent disseminated intravascular coagulation occurred in 3 women after ingestion of quinine tablets for cramp. All had circulating quininedependent antibodies to platelets and in 2 there was initial evidence of antibody consumption, with low titres that rose steeply over the next few days and remained high for many months.

Lancet 1990; 338: 1535-37.

Introduction

Recognised haematological problems associated with ingestion of quinine include thrombucytopenia, erythrocyte haemolysis, and neutropenia. Quinine was first implicated as a cause of purpura in the late 19th century,¹ and there have been several reports of associated thrombucytopenia.²³ However, we are aware of only two published cases of disseminated intravascular coagulation induced by quinine,⁴⁴ and report three further cases.

Patients and methods

Case histories

A 71-year-old woman was admitted 5 times over 3 years with various symptoms, which included acute shortness of breath,

wheeze, generalised abdominal pain, fever, lower back and chest pain, melaena, haematemesis or haemopsysis, and bruising and petechiae. Most episodes occurred shortly after going to bed. Investigations on each occasion (table) showed evidence of disseminated intravascular congulation (DIC). On the first 2 admissions she was treated with antibiotics, although blood cultures were always negative. On the third admission she was treated for asthma, and on the last 2 occasions no specific treatment was given. On each occasion, fever and other symptoms resolved within 24 h with a subsequent resolution of congulation abnormalities. At least 2 similar, but milder, episodes also occurred for which she did not attend hospital. On 3 occasions recent quinine ingestion was clearly remembered by the patient or documented in the admission notes. A check on the number of tablets left in the bottle after her initial prescription indicated that 10 tablets had been taken over 7 years. Retrospective quinine-dependent platelet antibody analysis on samples stored from the last 3 admissions and from intervening periods showed low or undetectable anabody concentrations during the first 2 days of each acute episode, which then rose sharply

ADDRESSES Department of Haematology (R.L. Spearing, FRACP, C.M. Hickton, FN2IMLT) and Department of Nephrology (P. Sizeland, MRCP, A. Hannah, MB, R.B. Bailey, FRCP), Christohurch Hospital, Private Bag, New Zealand, Correspondence to Dr.R. L. Spoaring, Department of Haematology, Christohurch Hospital, Private Bag, Christohurch, New Zealand

MOLECULAR CHARACTERIZATION OF HTLV-I INFECTION IN ISRAEL

Yehuda L. Danon^{**}, Yael Kilim^{*} and Joseph D. Rosenblatt[@] *Kipper Institute of Immunology, #Basil and Gerald Felsenstein Research Center, Children's Medical Center of Israel, Sackler School of Medicine, Tel-Aviv University Israel and @Division of Hematology Oncology, Department of Medicine, UCLA School of Medicine Los Angeles Calif 90024, USA.

Address for Correspondence: Y.L. Danon, M.D.

Director,

The Children's Medical Center of Israel P.O. Box 559 Petah-Tikva 49202 Israel

This work was supported in part by the US Army Medical Research and Development Command

-

£.,

Short Title: Characterization of Israeli HTLV-I isolate

INTRODUCTION

Many studies were performed on human T-cell leukemia virus type I, ethiologically associated with adult T-cell leukemia (ATL) 1,2 and HTLV-associated myelopathy (HAM). $^{3-7}$

HTLV-I genomes isolated from ATL and HAM patients from different geographical origin demonstrated a high degree of homology (>96%).⁵⁻⁸ Sequence variation in different isolates were found mostly in the LTR (1.3-5.2%) and the region between the envelope and tax/rex reading frames $(0.1 \ 6.9\%)$.⁶ It was found that genetic diversity between different isolates is in association with the geographical origin and not with the clinical presentation.^{6,7,8} In 1988, HTLV-I was first discovered in Israel and the Middle East.⁹ Two years later a community of jewish immigrants from the city of Mashad in northeastern Iran was identified with an infection rate of about 12%.¹⁰

Our aim was to determine the nucleotide sequence of LTR and env gene from HTLV-I genome of an HAM patient who originated from Mashad. This data was compared to sequences derived from HTLV-I isolated from Japanese and African patients in order to locate the origin of the Mashadi virus.

MATERIALS AND METHODS

<u>Cell lines:</u> Lymphocytes were collected from 20ml peripheral blood by a Ficoll-Hypaque density gradient and were resuspended in RPMI-1640 medium supplemented with 15% fetal bovine serum, interleukin-2 (IL-2) at a concentration of 50 μ /ml, 0.5% PHA and 1% penicillin/streptomycin. Cells were incubated at 37^oC in the presence of 5% CO2 and were maintained for several weeks. The cells were cryopreserved at various intervals for DNA extraction.

DNA extraction and PCR amplification: DNA was extracted from lymphocytes with phenol/chloroform. Two regions in the HTLV-I genome were amplified: LTR and env gene. LTR amplification was performed with the primers R11/14 which defined a 741 base sequence from nucleotide 61 to 802 (nucleotides are numbered according to the sequence of Seiki et al.¹¹ The env gene was amplified by two pairs of primers : R15/17A which defined a 1163 base sequence from nucleotide 5201 to 6364 and R19/18 primers which defined 791 base sequence from nucleotide 5942 to 6733.

The patient's (HE) DNA was PCR amplified for 30 cycles in a reaction containing 100 μ l of 2mM MgCl, 200 ng from each primer, 0.2 mM from each dNTPs and 2.5 u of taq polymerase (USB, Cleveland, OH). A list of primers used for PCR amplification is presented in table 1.

DNA sequencing: PCR products were recovered from PCR reaction with DS primer Remover (Advanced Genetic Technologies Corp, Gaithersburg, MD) and ethanol precipitation. Nucleotide sequence analysis was performed by the Taq Dye Deoxy Terminator Cycle Sequencing kit using the 373A DNA Sequencer (Applied Biosystems) at the Biological Services of the Weizmann Institute Rehovot Israel. Sequence analysis was performed at least twice for each primer (Table 1).

- 4 --

<u>Sequence comparison</u>; HE sequence was compared to the Japanese (ATK), African (EL) and Papua New Guinea (PNG-1) sequences by gcg program with the accession numbers: JO2029 (ATK), S74562 (EL LTR), M85207 (PNG-1) and M69044 (EL).

RESULTS

<u>Comparison of HTLV-I (HE) LTR to other isolates.</u> PCR amplification of HTLV-I LTR produced a PCR product of 741bp of which 698 bases from it were sequenced. 7 nucleotides which are not verified yet are designated as N. HTLV-I-(HE) LTR sequence was compared to three other isolates. The Japanese (ATK) sequence showed nucleotide homology of 97.8% with 15 nucleotide differences.¹¹ The African sequence showed nucleotide homology of 95% with 35 nucleotide differences and the Papua New Guinea isolate, which was compared only by 629 nucleotides, showed 91% homology with 56 nucleotides differences.^{5,6}

Comparison of the sequence changes in the LTR region to the Japanese sequence showed that the differences are clustered mostly in the U3 (66.7%), 6.7% were in the R and 26.7% were in the U5 regions. There are no changes at the three 20bp enhancer elements at the U3 region and there is 98.7% homology for the Rex Responsive element located between bases 313 and 627.

<u>Comparison of the env gene amplified from HTLV-I (HE) genome to the</u> <u>Japanese and African sequences</u>. The PCR product was 1532bp in length from which 1496 bases were sequenced. The sequence between nucleotides 5625-5685 is not verified yet. Comparison to the Japanese (ATK) sequence showed nucleotide homology of 97.7% with 9 nucleotide differences while the frican (EL) sequence showed nucleotide homology of 98.4% with 6 nucleotide differences.^{12,14} There is a problem with base T at position 5400. It seems that it creates a stop codon while in the other two sequences it does not exist. This finding needs further examination. HE env2 sequence was compared to the Japanese (ATK) sequence.¹¹ 98.7% homology at the nucleotide level with 12 nucleotide differences were found. Comparison to the African (EL) sequence showed nucleotide homology of 99.5% with 5 nucleotide differences.¹²

- 6 -

DISCUSSION

A new focus of HTLV-I infection was recently identified in the Middle East.¹⁰ In this study we determined the sequence of two regions in HTLV-I isolated in Israel from an HAM patient belonging to this community: the LTR region and the env gene. Several studies have indicated a high degree of homology among HTLV-I isolates (>96%) and demonstrated that differences between variants are in association with their geographical origin.^{5,6,7,8} Comparison of Mideastern sequence which originates in Iran to the Japanese, African and Papua New Guinea sequences in order to examine whether there is indeed a higher degree of homology between isolates from the same geographical area. A high level of homology, at about 98%, to the Japanese isolate, while the African and Papua New Guinea isolates were more divergent with sequence homology of 95% and 91%, respectively.

21

Comparison of HTLV-I env gene to the Japanese and African isolates showed homology at the nucleotide level of about 98-99%. From these results we can conclude that the env gene is more conserved than the LTR region.

_ 7 _

With the exeptions of some nucleotides which are not verified yet, comparison of the LTR region showed a higher degree of homology between Iranian sequence and the Japanese isolate. As for the env gene, there is a high degree of homology between the Iranian sequence and the African isolate. With these results we cannot conclude if there is an influence of the geographical area on the virus genome.

There are two theories about HTLV-I origin and the way it was spread to the world. Gallo suggests that it originated in Africa and spread to the new world and Japan, while Saskena et al. suggest that HTLV-I originated in the Indo-Malay region.^{5,13,14} Based on the comparison of the LTR sequences, our results support the theory of HTLV-I originating in Africa, since we found higher degree of homology between HE sequence and the African isolate, than with the sequence of Papua New Guinea isolate.

۳۱

REFERENCES

- POIESZ, B.J., P.W. RUSCETTI, A.F. GAZDER, P.A.BUNN, J.D.MINNA & R.C. GALLO. 1980. Detection and isolation of type C retrovirus particles from fresh and cultured lymphocytes of a patient with cutaneous T-cell lymphoma. Proc.Natl.Acad.Sci. USA 77: 7415-7419.
- UCHIYAMA, T., J. YODOI, K. SAGAWA, K. TAKATSUKI & H. UCHINO.
 1977. Adult T-cell leukemia: clinical and hematologic features of 16 cases. Blood 50: 481-492.
- TSUJIMOTO, A., T. TERUUCHI, J. IMAMURA, K. SHIMOTOHNO, I. MIYOSHI & M. MIWA. 1988. Nucleotide sequence analysis of a provirus derived from HTLV-I-associated myelopathy (HAM). Mol. Biol. Med. 5: 29-42.
- GESSIAN, A., F. BARIN, J.C. VERNANT, O. GOUT, L. MAURS, A. CALENDER & G. de THE. 1985. Antibodies to human T-lymphotropic virus type I in patients with tropical spastic paraparesis. Lancet 11: 407-409.
- SAKSENA, N.K., M.P. SHERMAN, R. YANAGIHARA, D.K. DUBE & B.J. POIESZ. 1992. LTR sequence and phylogenetic analysis of a newly discovered variant of HTLV-I isolated from the Hagahai of Papua New Guinea. Virol. 189: 1-9.
- RATNER, L., T. PHILPOTT & D.B. TROWBRIDGE. 1991. Nucleotide sequence analysis of isolates of human T-lymphotropic virus type I of diverse geographical origins. AIDS Res. and Human Retroviruses. 7: 923-941.

۳,

- KINOSHITA, T., T. TSUJIMOTO & K. SHIMOTOHNO. 1991. Sequence variations in LTR and env regions of HTLV-I do not discriminate between the virus from patients with HTLV-I associated myelopathy and adult T-cell leukemia. Int. J. Cancer 4: 491-495.
- DAENKE, S., S. NIGHTHIGALE, J.K. CRUICKSHANK & C.R.M. BANGHAM.
 1990. Sequence variants of human T-cell lymphotropic virus type I from patients with tropical spastic paraparesis and adult T-cell leukemia do not distinguish neurological from leukemic isolates. J. Virol. 64: 1278-1282.
- LEOR J., P. LANGEUITZ, H. TRAU, E.O.M. SCHNIDER, D. DOUER & I. BEN-BASSAT. 1988. HTLV-I associated T-cell leukemia/lymphoma in Israel. Isr. J. Med. Sci. 24: 397-400.
- MEYTES, D., B. SHOCHAT, H. LEE, G. NADEL, Y. SIDI, N. CERENY,
 P. SWANSON, M. SHAKLAI, Y. KILIM, M. ELGAT, E. CHIN, Y. DANON
 & J.D. ROSENBLATT. 1990. Serological and molecular survey for
 HTLV-I infection in a high-risk Middle Eastern group. Lancet
 336: 1533-1535.
- SEIKI, M., S. HATTORI, Y. HIRAYAMA & M. YOSHIDA. 1983. Human adult T-cell leukemia virus: complete nucleotide sequence of the provirus genome integrated in leukemia cell DNA. Proc. Natl. Acad. Sci. USA 80: 3618-3622.
- PAINE, E., J. GARCIA, T.C. PHILPOTT, G. SHAW & L. RATNER. 1991.
 Limited sequence variation in human T-lymphotropic virus type

 isolates from North American and African patients. Virol.
 182: 111-123.

- 9 -

- 13. GALLO, R.C. 1991. Human retroviruses: a decade of discovery and link with human disease. J. Infect. Dis. 164: 235-243.
- ROSENBLATT, J.D., Y. DANON & A.C. BLACK. 1992. A decade with HTLV-I/HTLV-II: Lessons in viral leukemogenesis. Leukemia 6 (1): 18-23.

٠,

Prim er	From 1	nt. to nt.	Used for	- Seq.	Remarks
R11	61	77	amp. & Seq.	5' TAGAGCCTCCCAGTGAA	
R12	494	470	seq.	5' CCTAGACGGCGGACGCAG	Comp.
R14	802	78 6	amp. & seq	5' CTCGTATCCCGGACGAG	Comp.
R15	5201	5218	amp. & seq.	5' CATGGGTAAGTTTCTCGC	
R16	5660	5645	seq.	5' ATGGAGATTAATATTG	Comp.
R17	5641	5658	seq.	5' GCCTCAATATTAATCTCC	Comp.
R19	5942	595 9	amp. & seq.	5' TCCATCCTCTTCTTCTAC	
R17A	6364	6347	amp.	5' TCCCAGAACAGGAGATCA	Comp.
R18	6733	6716	amp. & seq.	5' GGGAGAGGTAATTATTG	

Table 1: LIST OF PRIMERS FOR PCR AMPLIFICATION AND SEQUENCING

Δ,

Editorial

T-LYMPHOCYTES IN CHILDHOOD LEUKEMIA

Yehuda L. Danon, MD D Kipper Institute of Immunology, Chilren's Medical Center of Israel, Tel-Aviv University, Sackler School of Medicine, Kaplan Street 14, Petah-Tikva 49202, Israel

[□] The recent decade is associated with major breakthroughs in lymphocyte immunotyping. In view of the excellent improvement in chemotherapy results in acute lymphoblastic leukemia (ALL) and the chemotherapy associated mortality, Dr. Lovat and associates studied serially the T-lymphocytes in childhood leukemia¹.

The immunophenotyping classification of ALL is part of the diagnostic profile that includes morphology cytochemistry and genetic karyotypic assay.

Immunophenotypic classification divides ALL to common ALL with blasts expressing HLA-DR, CD-19 and CD-10; null ALL with HLA/DR and/or CD-19 antigens presenting on cell surfaces; and pre B-ALL with immunophenotyping profile including HLA-DR, CD-19 and CD-10 with cytoplasmic μ -chain²⁻³.

The serial study of T-lymphocyte subpopulation quantity and function published in this journal¹ reveals a significant and marked decrease in circulating T helper cells CD4—number and significant but less profound fall in CD8 Cytotoxic T cells compared to normal controls.

Polyclonal and specific HSV-1 proliferation responses showed a slightly but significantly decreased response in cALL patients; however, marked impairment of T-cell response to specific recall antigens or polyclonal stimulation is not demonstrated. Although there were persistent reports of heterogeneity of the helper T-cell compartment, only relatively recent were distinct types of helper T cells resolved defined primarily by the different patterns of lymphokine synthesis that became also a convenient marker to describe T help cells subclass differences, but an extensive study of IL-2 and IL-4 production could not show any difference from control patients.

A recent study, aimed at investigating whether CD4 + T cells are predetermined to produce given patterns of lymphokynes, showed that IL-2producing clones can be derived from the same cells as IL-4-producing clones,

Pediatric Hematology and Oncology, 10:ix-z, 1993 Copyright © 1993 Taylor & Francis 0888-0018/93 \$10.00 + .00

he

۲,
X YEHUDA L DANON

supporting the view that the IL-2-producing Th1 or the IL-4-producing Th2 phenotype of a T cell clone is acquired during T cell differentiation³.

Differential production of cytokines by helper T cells during the immune reaction has important regulatory effects on the nature of the response and induction of Th1, and because Th2 responses play a key role in the natural response, it seems that availability of additional monoclonal antibodies, improved immunophenotyping methods and better understanding of cytokine mechanisms will improve our monitoring capabilities in the future. It was shown that CD8 + T cells may switch the response induced by antigens and antigen-presenting cells from humoral to a cell mediated role.

Lymphocyte phenotyping was assessed by Lovat et al.¹ using indirect immunofluorescence on blood smears. The availability of a variety of monoclonal antibodies dramatically increased our possibilities to study cellular subpopulations. Microscopic quantitation methods of immunofluorescence are ill suited to accurately count large numbers of cells; direct or indirect immunofluorescence is more accurately analyzed by flow cytrometry with the possibility of sorting and functually studying target populations. Even with flow cytrometry, the differences in instrumentation, antibodies and techniques may introduce significant sources of variation.

Infection is a major cause of morbidity and mortality in leukemic patients. Infections are bacterial, protozoal, viral and fungal, although the underlying mechanism for the increased susceptibility to infection is not clearly defined. The present work widens our knowledge and scope of the prediction of susceptible populations. It seems that a wide, long-range prospective study is needed defining predictors of infections, possible modifications needed in chemotherapy and other ways to treat immunosuppression.

REFERENCES

- 1. Lovat EP, Robinson JH, Windebank KP, Kernaban J, Watson JG. Serial study of T lymphocytes in childhood leukemia during remission. *Pediatr Hematol Oncol.* 1993.
- Ludwig WD, Barram CR, Harbott J, et al. Phenotypic and genotypic heterogeneity in infant acute leukemia. Leukemia 1989; 3:431-439.
- 3. Rocken M, Saurat J, Hand Hauser C. A common precursor for CD4+ T cells producing IL-2 or IL-4. J Immunol. 1992;148:1031-1036.

Address correspondence to Y. Danon.

Annotation

EPIPOE

Richard & Philadelphi

Hente phagocyti a myelody teniposide predisposi which has this diseas Presumabl saved him arisen fron Eviden teniposide Jude Child acute lymp side or ten myeloid leu secondary h an alternate United Kin most convin study of ove and 1983. th ters were m dophyllotoxi with a second develop secor (five cases ha tients). This bined modalit

> Pediatric Hema Copyright © 1 0888-0018/93



FOREWORD

Opinions, interpretations, conclusions and recommendations are those of the author and are not necessarily endorsed by the U.S. Army.

 \mathbb{N}) Where copyrighted material is quoted, permission has been obtained to use such material.

()) Where material from documents designated for limited distribution is quoted, permission has been obtained to use the material.

 $(\sqrt{)}$ Citations of commercial organizations and trade names in this report do not constitute an official Department of the Army endorsement or approval of the products or services of these organizations.

() In conducting research using animals, the investigator(s) adhered to the "Guide for the Care and Use of Laboratory Animals," prepared by the Committee on Care and Use of Laboratory Animals of the Institute of Laboratory Animal Resources, National Research Council (NIH Publication No. 86-23, Revised 1985).

(||) For the protection of human subjects, the investigator(s) have adhered to policies of applicable Federal Law 45 CFR 46.

() In conducting research utilizing recombinant DNA technology, the investigator(s) adhered to current guidelines promulgated by the National Institutes of Health.





DTIC OCALVES INCEDOTED L

MOLECULAR STUDIES OF HTLV-I INFECTION IN NEWLY RECOGNIZED HIGH RISK POPULATION - MIDTERM REPORT

-

TABLE OF CONTENTS

<u>Paqe</u>

1.	Front Cover	1
2.	Report-Documentation Page	2
3.	Foreword	3
4.	Table of Contents	4
5.	Introduction	5
6.	Final Report	10
7.	Methods	10
8.	Results	14
9.	Discussion	23
10.	References	28
11.	Conclusions	35
12.	Bibliography of publications	37
13.	List of Personnel	38
14.	Appendix	

*

τ,

INTRODUCTION:

Human retroviruses have been recognized and well characterized during the last decade after the detection of human T-cell leukemia virus type I (HTLV-I) by Poiesz and Gallo (1) and Yoshida (2) endemic areas were described in southern Japan and parts of the Caribbean and South America (3-7), as well as high risk groups of IVDA (Intravenous Drug Abusers) (8). Following the discovery of human immunodeficiency virus types 1 (HIV-1) and 2 (HIV-2) and their implication in AIDS accelerated the intensity of research for possible oncogenic viruses. HTLV-I is the human retrovirus known as the cause or ethiologic agent of Adult T-cell Leukemia (ATLC) and of the chronic progressive demyelinating process HTLV-I-associated Myelopathy HAM, known also as Tropical Spastic Paraparesis - TSP.

It was found that leukemic cells in malignancies associated with HTLV-I contained clonically integrated provirus, leading to the direct viral involvement in the oncogenic process, while in contrast, neoplasms frequently seen in the setting of HIV infection - (Kaposis Sarcoma and high grade B-cell Lymphoma) did not appear to arise as a direct consequence of viral transformation of HIV-1 infected cells. HTLV-I studies were not successful to detect a viral message in leukemic cells of ATL, leading to the hypothesis that the HTLV-I gene expression is not required for the maintenance of T-cell leukemia or lymphoma.

The epidemiological link between HTLV-I and ATLL resulted from two complementary events. The first, an appreciation of Uchiyame and Takatsuki, that ATLL represents a unique clinical entity (9), allowed the geographic localization of the disease to the southern islands of Japan. Epidemiological studies showed that exposure shortly after birth is a major risk factor for subsequent development of HTLV-I associated malignancy (10), while a very small proportion (<5%) of contacts, after a period of twenty or more latent years, develop malignancies, suggesting a multiple step process that may be involved in leukemogenesis, while a viral infection may be a prorequisite, it alone may be insufficient to produce the leukemic phenotype.

These general epidemiologic features of ATLL suggest that systematic reevaluation of appropriate candidate malignancies and viruses may reveal other oncogenic viruses. Careful cataloguing and description of clinical syndromes is essential to derive epidemiological clues that may lead to virus identification. As example: the differentiation of ATLL from mycosis-fungoides is a case in point. While some of the special features of ATLL, as the enhanced expression of interleukin-2 (IL-2) receptor alpha (IL-2Rg) chain (Tac antigen) on the cell surface, was initially ignored while recognition of the subtler clinical aspects of the syndrome allowed for subsequent epidemiologic observations (9). The HTLV-I model for malignancy as a consequence of viral infection suggests that careful molecular reexamination of the role of already recognized viruses in suspect malignancies is in order. In the case of HTLV-I over one million of infected individuals in Japan give rise to about 500 cases of ATLL per year. Hodgkin's disease may provide another case in point. The bimodal age distribution, the age prevalence, anectodal description of geographic clustering and outbreaks of HD suggest possible viral ethiology (11-13).

The increasingly frequent reports of Epstein Bar Virus (EBV) genome detection in some cases of HD suggest that EBV can be important in the pathogenesis of subsets of HD patients (14), as well as in some ENT malignancies (15).

The example of the last five cases of ATLL-like T-cell Lymphoma in Israel, occurring in Iranian immigrants from the area of north eastern Iran (16), raised the possibility of a new population at risk for HTLV-I infection. This is an example of geographic, familial and ethnic clustering of HTLV-I infection that may yield important clues to viral etiology, in view of conflicting reports about clustering in children less than six years of age (17-19) or reports about geographic clustering associated with high socioeconomic status (20-21), while others question those reports (22-23).

In view of the clustering of ATLL in Jewish newcomers from north eastern Iran (16,24) we have designed this study being a systematic survey of HTLV-I serology in newcoming communities to Israel, with an emphasis on middle eastern communities. The methodology of the research was based on a general survey of Israeli blood donors. Annually about 250.000 units of blood are donated in Israel, big part of them through the Israel Defence Forces. All data of blood donors is computerized and part of the National Medical Database. This national database is based on the fact that virtually all non-Arab citizens of the country are drafted (25). Conditions in Israel are favorable for epidemiologic research. The country is small, densely populated, with a special ethnic mixture of the Jewish Community and quite society stable population.

Reliable population registers are available and any individual may be traced by his personal, seven digit, identification number, used ago for administrative and research purposes. Data is available from records of hospital admissions and discharges, as well as from the National Compensatory Health Draft Registry, enabling total population studies on Jewish young adults. Numerous epidemiologic studies, done by our group, have used this database (26-34). In addition to conducting an epidemiologic survey, using serologic methods, two additional aimes of this study are to disparate additional disease entities related to HLTV-I and to conduct a molecular survey of HTLV-I.

Approximately five years following the discovery of HTLV-I it was found that the pathology caused by HTLV-I in one setting may not predict other forms of pathology related to the virus. It seems that the role of HTLV-I in ATL is restricted to generating a polyclonal T-lymphocyte proliferation after infection (35).

A specific illustration is afforded by the two major non-overlapping syndromes associated with HTLV-I. While HTLV-I may cause ATLL, another subset of infected individuals that some of them will develop a slow neurologic disease characterized by a gradual development of spastic paresis of the lower extremities with minimal sensory loss. This illness of tropical spastic paresis or HTLV-I-associated myelopathy (HAM), differentially diagnosed from multiple sclerosis, is connected to the virus. The association between the illness - HAM and HTLV-I - was discovered by Gessian et al (36), while screening for neurologic diseases in Martinique. Regardless of underlying mechanism involvement of HTLV-I, in a slow neurologic disease, is guite surpri-

- 7 -

sing. So far, up to the present research, no HAM cases were described in Israel in view of the description of ATLL, an additional aim of our work, as part of the epidemiologic work to screen for possible HTLV-Iassociated neurologic disease in Israel.

Serologic survey is backed by western blotting. The demonstration of Seiki and colleagues of the unique coding regions located in the 3' end of the genome provided a first clue as to potential oncologic mechanisms of HTLV-I (37). The 3' ends of the genome of HTLV-I, -II and the leukemogenic bovine leukemia virus (BLV) all contain coding sequences for at least two overlapping genes (38-41). These genes, known as tax and rex, are now known to encode a transcriptional and post-transcriptional regulator of viral expression. The initial discovery of the tax gene was surprising, in that such transacting transcriptional regulators had only been previously identified in DNA viruses such as herpes simplex virus, varicella-zoster virus, and adenovirus.

The HTLV-I tax gene encodes a 40-kilodalton (kDa) protein, and the HTLV-II tax gene encodes a 37-kDa protein (36-41). These nuclear phosphoproteins are highly homologous, and have similar effects on transcription. Tax expression is necessary for transcriptional activation of virus replication and efficient RNA production from the viral long terminal repeat (LTR); however, an additional property noted early was the ability of Tax to trans-activate other viral and cellular promoters, including those involved in regulation of T-cell growth. Examples of such promoters include those for IL-2, IL-2Ra, and the lymphokine, granulocyte-macrophage colony-stimulating factor (GM-CSF) (42-45). In the case of HTLV-I and -II, Tax appears to interact with three 21-base pair (bp) repeats located in the U3 portion of the viral LTR. Within the sequence of the 21-bp repeats are core sequences known to bind cellular transcription factors (46-47). A variety of these proteins have now been identified and partially characterized.

ά,

- 8 -

The aim of this research is also to back the serologic and epidemiological work with molecular screening in addition to western blotting, measuring of antibodies to HTLV-I p40tgx, with recombinant p40tax as an antigen, since ATL and HTLV-I show the same geographic distribution, and ATL patients are always infected with HTLV-I (48-49,50) it is expected that every new geographic endemic focus will behave similarly, meaning identification of HTLV-I carreirs ATL and HAM population, and viral isolates as part of this study. Moreover, it is important to study the cell surface antigens (CSA) of the isolated infected cells, and viral structure (51).

The genetic structure of HTLV-I is similar to that of the other known retroviruses with three well identified regions: gag region, a pol region and an env region (52-54) with an additional unique pX region (54) that is crucial to the activated host genes in view of those important relations of viral structure and pathogenesis of HTLV-I in particular, and retroviral disease in general, part of our aim was to study viral isolate structure.

...

- 9 -

FINAL REPORT:

Blood samples were drawn from Israeli blood donors, through the central blood services of Magen David Adom. Additional sources of samples was the collection of blood through the Association of Iranian immigrants to Israel and through the community services of Iranian and Mashadi Jews in Israel: clinics, elderly homes, schools and synagogues in Bney Barak, Holon, Bat-Yam, Herzlia and Tel-Aviv, all towns and communities in the central part of Israel. Additional parts of this survey were samples from patients on long-term hemodia-lysis, peritoneal dialysis, Adult T-cell Leukemia (ATLL), T-cell malignant disorders, other than Adult T-cell Leukemia, Mycosis fungoides, Sezary's Syndrome Complex, Parapsoriasis, non-Burkitt's Lymphoma patients, young adults with Insulin Dependent Diabetes Mellitus (IDDM) and the collection of blood samples from immigrants from Ethiopia.

During the report period, up to the end of June 1993, we have studied altogether 11230 blood samples including 10122 blood donors (all of them negative for HTLV-I), 480 Iranian Jews of them 212 Mashadi Jews, 181 Ethiopian immigrants, 36 T-cell malignancies, 41 hemodialysis patients, 40 ICPD patients, 32 Mycosis Fungoides/Parapsoriasis patients, 90 non-Burkitt's Lymphoma patients and 208 IDDM patients. Unfortunately, at present we could not collect any data from native Iranians in Mashad or the region, as well as Iranian refugees from Mashad in Europe.

METHODS:

Serologic screening for HTLV-I antibodies in serum or plasma samples was done by means of an enzyme-linked immunosorbent assay-ELISA (Abbott Laboratories). Confirmatory was done by western blotting and/or radioimmunoprecipitation assay - RIPA with Sulphur S³⁵-labeled methionine. HTLV-I-infected HUT 102B lysates were also done (55). Samples, positive in the ELISA, were tested by both confirmatory methods. Antibodies to HTLV-I p40tax were measured by means of an ELISA with recombinant p40 tax as an antigen on solid phase of polystrene beads (Abbott Laboratories Diagnostic Division North Chicago Illinois, USA). HTLV-I seropositive-infected samples, with a known reactivity against p40tax on RIPA, were used as positive controls (Dr. Rosenblatt's Laboratory Division of Hematology-Oncology Department of Medicine UCLA School of Medicine, Los Angeles California USA) and samples negative for HTLV-I on ELISA and western blot as negative controls. Samples were scored as a positive for p40tax, if the optimal density exceeded 4-5 times the mean negative control value. The polymerase chain reaction - PCR - was used to amplify HTLV-I sequence of DNA from peripheral blood mononuclear cells with primers to a 159bp segment contained with the tax/rex gene as described before (56).

<u>Cell-lines:</u> Lymphocytes were collected from 20ml of heparinized peripheral blood by a Ficoll-Hypaque density centrifugation and resuspended in RPMI-1640 medium supplemented with 15% fetal bovine serum, interleukin-2 (IL-2) at a concentration of 50u/ml, 0.5% PHA and 1% penicillin/streptomycin. Cells were incubated at 37° C in the presence of 5% CO² and were maintained for several weeks. The cells were cryopreserved at various intervals for DNA extraction.

DNA extraction and PCR amplification: DNA was extracted from lymphocytes with phenol/chloroform. Two regions in the HTLV-I genome were amplified: LTR and env gene. LTR amplification was performed with the primers R11/14 which flank a 741 base sequence from nucleotide 61 to 802 (nucleotides are numbered according to the sequence of Seiki et al (37). The env gene was amplified by two pairs of primers: R15/17A which flanks a 1163 base sequence from nucleotide 5201 to 6364 and R19/18 primers which flank 791 base sequence from nucleotide 5942 to 6733. The patient's (HE) DNA was PCR amplified for 30 cycles in a reaction containing 100 μ l of 2nM MgCl, 200 ng from each primer, 0.2 mM from each dNTPs and 2.5 u of tag polymerase (ESB, Cleveland, OH). A list of primers used for PCR amplification is presented in table 1.

DNA sequencing: PCR products were recovered from the PCR reaction with DS primer remover (Advanced Genetic Technologies Corp. Gaithersburg, MD) and ethanol precipitation. Nucleotide sequence analysis was performed by the bTaq Dye Deoxy Terminator Cycle Sequencing kit (Applied Biosystems) using the 373A DNA sequencer (Applied Biosystems) at the Biological Services of the Weizmann Institute Rehovot Israel. Sequence analysis was performed at least twice for each primer (Table 1).

<u>Sequence comparison:</u> HE sequence was compared to the Japanese (ATK), African (Zair, EL) and Papua New Guinea (PNG-1) sequences by geg program with the accession numbers: JO2029 (ATK), S74562 (Zairian LTR), M85207 (PNG-1) and M69044 (EL).

Primer	From nt.	To nt.	Used for	· Seq.	Remarks
RIL		77	amp. & seq.	5' TAGAGCCTCCCAGTGAA	
R12	494	470	seq.	5' CCTAGACGGCGGACGCAG	comp.
R14	802	786	amp. & seq.	5' CTCGTATCCCGGACGAG	comp.
R15	5201	5218	amp. & seq.	5' CATGGGTAAGTTTCTCGC	
R16	5660	5645	seq.	5' ATGGAGATTAATATTG	comp.
R17	. 5641	5458	seq.	5' GCCTCAATATTAATCTCC	comp.
R19	5942	5959	amp. & seq.	5' TECATECTETTETTETAC	
R17A	_6364	6347	amp.	5' TECCAGAACAGGAGATCA	comb -
R18	6733	6716	amp. & seq.	5' GGGAGAGGTAATTATTG	

able 1: list of primers for PCR amplification and sequencing.

•

- 13 -

-

Out of 11230 blood samples tested for HTLV-I we have found 26 positive for HTLV-I in ELISA and western blot and 12 positive in anti p40tax ELISA. The positive results are detailed in the following tables.

TABLE 2: Prevalence of HTLV-I infection

Patients #		& western blot # Positive (%)
Iranian Jews	480	5.41% (26)*
Mashadi Jews	212	12.3% (26)
Ethiopian Jews	181	0 %
T-cell malignancy	136	0%
Haemodialysis	41	0%
CIPD	40	0%
Mycosis Fungoisis	&	
Parapsoriasis	32	0%
Non-Burkitt's		
Lymphoma	90	0%
IDDM	208	0%

*All positive patients were of Mashadi origin

Anti p40tax ELISA # tested # Positive (%)		
190	19 (10%)	
92	0%	
65	0%	
20	0	
40	0	
30	0%	
22	0%	
	# tested 190 92 65 20 40 30	

TABLE 3: Anti p40tax ELISA results

The extensive social and ethnic ties between Jews of Mashadi origin allowed selective sampling of this population within Israel. On repeated assys, 26 of 212 (12.3%) Mashadi serum samples were positive by ELISA for antibodies to HTLV-I and were confirmed by western blotting. In contrast, none of the control samples was positive by ELISA (Tables). In an effort to survey the oldest members of the Mashadi population, we tested 32 unrelated elderly Mashadi women living in an old people's home in Tel-Aviv, 14 (44%) were seropositive, a rate three times higher than that in the general Mashadi population. At our first sample collection, 12 (52%) of 23 long-standing residents of the home were seropositive, 2 of 9 new residents surveyed a year later were seropositive. Of 26 Mashadi women, older than 60 years, who lived elsewhere in Israel only 3 (12%) were seropositive.

To determine whether serologocal assays had detected all infected subjects, we carried out PRC amplification of HTLV-I DNA from 29 of the residents of the old people's home. '9 were positive for HTLV-I sequences: 14 of these were seropositive, 4 were seronegative, and 1 was seronegative by standard ELISA but reactive in the anti-p40tax ELISA. A second PCR on repeat samples from 10 of these women showed concordant results in 6 of 6 PCR-positive and 2 of 3 PCR-negative cases; 1 woman originally positive on PCR was negative on the second sample, and 1 woman originally PCR-negative was faintly positive on her second sample and had antibodies to p40tax by ELISA. This unusual clustering of HTLV-I infection within the old people's home suggested acquisition of infection within the institution.

In addition to antibodies to virion components, we assayed for antibodies to the non-structural p40tax protein. Of 190 Mashadi samples tested 19 had absorbance levels 4-5 or more times those of the negative control and were judged positive (Tables). 92 samples were negative by both assays, 8 samples were seropositive for both anti-HTLV-I and anti-p40tax; 13 samples were positive for anti-HTLV-I and negative for anti-p40tax; and 4 were positive for anti-p40tax but repeatedly negative for anti-HTLV-I by ELISA and western blot. 3 of the latter group of samples were negative on PCR and the other, from a resident of the old people's home, was positive. All 4 were negative by western blotting and by RIPA. Therefore, the reason for the high titres in three samples in the anti-p40tax ELISA is unclear.

Ψ,

- 15 -

Direct comparison of anti-p40tax ELISA results and those of the RIPA was done for 28 subjects. 8 samples were positive for anti-p40tax by both RIPA and ELISA; 1 with traces of antibody by RIPA was negative by ELISA; the 3 samples mentioned previously were negative by RIPA but repeatedly positive by ELISA and 16 were negative by both tests. The usefulness of the anti-p40tax ELISA in detecting true additional HTLV-I-infected seronegative individuals remains unclear, since seronegative infection was corroborated by other means of detection in only 1 of the 4 questionable cases.

The finding of a higher rate of seropositivity among elderly Mashadi women, than in the general Mashadi population, suggested a potential for gradual development of seropositivity with age, as has been reported by others. Because of the possibility that serological screening may not detect all positive subjects we carried out HTLV-Ispecific PCR on DN from 68 randomly selected Mashadi blood samples. 5 of 7 known seropositive samples were positive in the PCR assay, and 61 sero-negative samples were all negative by PCR. Hence, PCR did not increase sensitivity of detection within this sample above that was achieved with serological assays alone. All PCR-positive samples contained HTLV-I and not HTLV-II, as shown by discriminatory PCR.

We have characterized the Iranian HTLV-I isolate using a combination of Southern blotting, PCR and sequencing. We have developed a primary T-cell line from a Mashadi HTLV-I carrier containing integrated HTLV-I provirus.

Comparison of HTLV-I(HE) LTR to other isolates: PCR amplification of HTLV-I LTR produced a PCR product of 741bp of which 698 bases were sequenced. 7 nucleotides, which are not verified yet, are designated as N. HTLV-I (HE) LTR sequence was compared to three other isolates. The Japanese (ATK) sequence showed nucleotide homology of 97.8% with 15 nucleotide differences (37). The Zairian sequence showed nucleotide homology of 95% with 35 nucleotide differences and the Papua New Guinea isolate showed 91% homology with 56 nucleotides differences, in 629 nucleotides examined (56,57,58). Comparison of the sequence changes in the LTR region to the Japanese sequence showed that the differences are clustered mostly in the U3 (66.7%), 6.7% were in the R and 26.7% were in the U5 regions. There are no changes at the three 20bp enhancer elements at the U3 region and there is 98.7% homology for the Rex responsive element located between bases 313 and 627 (Fig. 1,2).

Comparison of the env gene amplified from HTLV-I (HE) genome to the Japanese and African sequences. The PCR product was 1532 bp in length of which 1496 bases were sequenced. The sequence between nucleotides 5625-5685 is not verified yet (Fig. 3).

Comparison to the Japanese 'ATK sequence showed nucleotide homology of 97.7% with 9 nucleotide differences, while the African (EL) sequence showed nucleotide homology of 98.4% with 6 nucleotide differences. There is a problem with base T position 5400. It seems that it creates a stop codon while in the other two sequences it does not exist. This finding needs further examination, but may be due to a defective HE provirus or PCR cloning artifact. HE env2 sequence was compared to the Japanese (ATK) sequence (37). 98.7% homology at the nucleotide level with 12 nucleotide differences were found. Comparison to the African (EL) sequence showed nucleotide homology of 99.5% with 5 nucleotide differences (58,59). In contrast, comparison of nucleotide sequences 5424-5839 within env between the Iranian isolate and MEL5, a Melanesian isolate from the Solomon Islands shows 11 nucleotide changes, as compared to none between the Iranian and Japanese (ATK) isolates (V. Nerurkar, pers. communication).

۳,

U3 78 HE aaacatttccgcgaaacagaagtctgaaaaggtcagggcccagactaagg ATK Α G Zair Α PNG-1 HE 128 ctctgacgtctccccccggagggacagctcagcaccggctcaggctaggc ATK G G Zair enhancer A Α TA T PNG-1 Α HE 177 cctgacgtgtccccctaaagacaaatcataagctcagacctccgggaagc ATK. G Α G G G Zair enhancer CT -GG GT G G PNG-1 cace ggaaccacceatttectecceatgtttgtegageegeecteagge HE 227 ATK AA Α Т T T G А A Zair Τ PNG-1 AA G Т CG HE 277 gttgacgacaacccctcacctcaaaaaacttttcatggcacgcatatggc ATK A enhancer Т Α C. Zair TATA box U3 R PNG-1 poly(A) signal 327 tgaataaactaacaggagtetataaaagegtggagacagtteaggagggg HE ATK С G C Α Zair C AC PNG-1 Α 377 HE getegeateteteetteacgegecegecetacetgaggeegecatee ATK Т Zair Т С PNG-1 C +GC Т HE 427 acgccggttgagtcgcgttctgccgcctcccgcctgtggtgcctcctgaa ATK Zair PNG-1 477 HE ctgcgtccgccgtctaggtaagtttagagctcangtcgagaccgggcctt ATK. Α Zair PNG-1 CG HE 527 tgtccggcgctcccttggagcctacctagactcagccggctctccacgct ATK Zair Α PNG-1 Α С Т T G

		R U5
HE ATK Zair PNG-1	577	
HE ATK Zair PNG-1	627	tgegeegetacagategaaagtteeaceettteeennneattenegaet T T T C G T
HE ATK Zair PNG-1	677	gactgccggcttggcccacggccaagtaccggcgactccgttggctcgga T TTAC AC T AC
HE ATK Zair PNG-1	72 7	gccagcgacagcccattctatagcactctccaggagagaaanttagtaca C – T C C A A
HE	776	c

Fig. 1: Nucleotide sequence of HTLV-I(HE) LTR. Nucleotide differences from the Japanese (ATK), Zairian (Zair) and Papua New Guinea (PNG-1) sequences are shown underline. Nucleotides are numbered according to ATK sequence

τ,

SU cactttgattttattcttccagttctgccccntcatcctcggtgattaca HE 5219 ATK C Α EL gccccagctgctgtactctcacaattggagtctcctcataccactctaaa HE 5269 ATK EL HE ccctgcaatcctgcccagccagtttgttcgtggaccctcgacctgctggc 5319 ATK EL cctttcagcagatnaggccctacagccccctngccctaatctagtaagt HE 5369 С ATK G EL tactctagctaccatgccacctattccctatatctattccctcattggat HE 5418 C. ATK С Т EL taaaaagccaaaccgaaatggcggaggctattattcagcctcttattcag HE 5468 ATK G EL ${\tt acccttgttccttaaagtgcccatacctggggtgncaatcatggacctgc}$ HE 5518 ATK EL $\verb|ccntacanaggagncgnctncaggccntactggnagtttcannaagatgt||$ HE 5568 С С ATK Т С EL т HE 5618 naatttt

Fig. 2: Comparison of HE env1 sequence to the Japanese (ATK) and African (EL) sequences.

۳,

HE 5686 cccttctagtcgangntccagganatgaccccatctggttccttaatacc ATK EL HE 5736 gancecagneaactgneteceacegnecetectaetececeactetaa ATK EL HE 5786 nctagancacatcctcgagccctctataccatggaaatcaaaactcctga ATK EL 5836 HE cccttgnccagttaaccctacaaagcactaattatacttgnattgnctgtATK EL HE 5886 atcgatcgtgncagnetatccacttggnacgtcctatannntcccaacgtATK С EL HE 5936 ctctgntccatcencttcttctacccccncctttacccatcgttagege ATK EL HE 5986 ttccagccccccacctgacgttaccatttaactggacccactgctttgac ATK EL HE 6036 ccccagattcaagctatagtctcctcccctgtcataactccctcatcct ATK EL HE 6086 gcccccttttccttgtcacctgttcccaccctgggatcccgctcccgcc ATK SU EL TM Α HE 6136 gageggtaceggtggeggnetggettgteteegeeetggwwatgggagee ATK EL HE 6186 ggagtggctggcgggattaccggctccatgtccctcgcctcaggaaagag ATK EL HE 6236 nctcctacatgaggtggacaaagatatttcccaattaactcaagcaatag ATK G. EL HE 6286 tcaaaaaccacaaaaatctnctcaaaattgcgcagtatgctgcccagaac ATK EL HE 6336 agacganggcnngatctcctgttctgggagcaaggaggattatgnaaagcATK С EL С

HE 6386 attacaagaacagtgctgttttctgaatattactaattccnatgtctcna ATK С С С EL HE 6435 atactacaagaaagaccccccttgagaatcgagtcctgactgggg ATK EL HE 6485 ccttaactgggaccttggcctctcacagtgggctcgagaggccttacaaa ATK EL HE 6535 ctggaatcacccttgtcgcgctactccttcttgtcatccttgcaggacca ATK Т EL Т 6585 HE tgcatcctccgtcagctacggcacctcccctcgcgcgtcagataccccca ATK A EL A HE 6635 ttactctcttataaaccctgagtcatnccctgtaaaccaagcacacaatt ATK Α G EL HE 6684 attgcaaccacatcgcctccagcctcccctgc

Fig. 3: Comparison of HE env2 sequence to the Japanese (ATK) and African (EL) sequences.

 \mathbf{T}^{i}

DISCUSSION:

We have identified a high risk of HTLY-I infection in Iranian Jews originating from the city of Mashad in Khurusan, northeastern Iran. This group seems to have an overall infection rate of about 12%, though the rate was much higher among residents of a Mashadi home for old people. The reason for this clustering is unclear, although the findings suggest acquisition of infection within the institution. The overall level of infection among other Iranian Jews seems to be substantially lower than that among Mashadis. The explanation may be geographic, and Mashad may be within a previously unrecognised endemic region. Since we did not test non-Jewish Mashadis, this possibility cannot be excluded. In the present situation we will not be able to test this population unless USAMRD, or any other federal agency, may help us with a proper connection with the Mashad University or any other scientific group in Iran.

The unique history of the Mashadi community may help to explain their high rate of HTLV-I infection. In 1839, the Jewish community in Mashad was subjected to a series of violent attacks and forced to convert to the Islam, though the majority of the community continued to practise Judaism covertly (62). To safeguard the community secret necessitated a very high rate of intermarriage among community members. Members of this community continued to marry close relatives for over the next 150 years. Markers of consanguinity are high among Mashadi Jews - for example, over 20% have a deficiency of glucose-6-phosphate dehydrogenase (63). The chance introduction of HTLV-I infection into the community could have resulted in rapid transmission of the virus by sexual means and by breastfeeding.

It was recently shown by Krivine (64) that there is an retrovirus replication in the first weeks of life. In a prospective longitudinal study of 50 infants, born to HIV-seropositive women, blood samples were obtained at birth at 4-9 weeks, and 5-9 month of age and were tested for HIV RNA by the polymerase chain reaction (PCR), viral culture and p24 antigen measurement. 16 were diagnosed as HIV infected by the age of 4-9 weeks by PCR and culture, while only 10% of the newly born - 5 were detected as HIV-positive at birth. No changes

in HIV infection were detected on both ages of testing for HIV. We started a similar study in HTLV-I-seropostive women, assuming that perinatal HTLV-I infection could also be demonstrated by PCR or culture, after the first two months of life. Such a work may also give some clues about the ways of transmission of HTLV-I and possibility of HTLV-I-like HIV during the first weeks of life, unfortunately on writing this final report we still do not have HTLV-I neonatal data from Israel.

An estimated 5000-6000 Mashadis now live in Israel. This is the first Israeli ethnic group identified as having a high rate of HTLV-I infection. Members of the Mashadi Jewish community also migrated to the USA and parts of Europe. We estimate that similar prevalence of HTLV-I infection could be found in other Jewish Mashadi communities. We intend to continue this work during the next year, and formed already initial conncetion with the Mashadi community in Milano, Northern Italy. Unfortunately we cannot do any work at present with native Iranian residing in Mashad.

In addition, it is estimated that Mashadi and Iranian Jews have moved to the East, to the regions of Uzbekistan, Afganistan, Kazahstan, Armenia and Pakistan (supplement map). Initial study of samples of those newly coming immigrants from Eastern Soviet republics (parts of former USSR) were negative but the data is still to small. We estimate that at least 20.000 immigrants have arrived to israel from those areas. There was also a settlement of Iranian Jews in China, in the area of Kaifen. But at present we have no possibility to explore this community, that partly assimilated in the general Chinese population.

In this study we use several methods to detect HTLV-I infection. Our rate of anti-p40tax seropositivity in carriers was somewhat lower than that found by others among healthy HTLV-I carriers (63-66), independent evidence for infection using PCR was obtained only in 1 out of 4 samples, thus detection of anti-p40tax antibodies did not appreciably add to the estimate of the rate of infection, and there is no point of its further use in our study. Our findings on the use of PCR suggest that in a high risk population such as the Mashadi population in the old people's home we have studied, or in families with HTLV-I

- 24 -

carriers, PCR would increase the number of detected infected individuals above those that are detected by serological survey.

The results in general population survey, in quite big sample, suggests that there is no point to continue such screening in high risk populations only. The usefulness of PCR as a screening assay in approriate setting requires further study.

The limited number of Ethiopian Jews studied does not confirm previously published results (67) about HTLV-I infection in the Falashi community and further work is needed in this group too. We have not summarized, yet, the data of Mycosis Fungoides and Parapsoriasis patients. Further work is also needed with the study of Mashadi isolates and their sequence as compared to Japanese and other isolates. The successful identification of HTLV-II by vitro of limited nucleic acid homology and antigenic similarity to HTLV-I suggests that variants and new retroviruses can be identified. We have published our initial result in a paper attached to this report (68). We have determined the sequence of two regions in HTLV-I isolated in Israel from an HAM patient belonging to this community: the LTR region and the env gene.

Several studies have indicated a high degree of homology among HTLV-I isolates (>96%) and demonstrated that differences between variants are primarily associated with their geographical origin (69-70). We compared HTLV-I in an Iranian immigrant with HAM to the Japanese, African and Papua New Guinea isolates. A high level of homology of about 98% was observed to the Japanese isolate, while the African and Papua New Guinea isolates were more divergent with sequence homology of 95% and 91% in the LTR respectively.

We have summarized, recently, our sequence data appearing in the appendix of this report (71-72).

Comparison of HTLV-I env gene to the Japanese and African isolates showed homology at the nucleotide level of about 98-99%. From these results we can conclude that the env gene is more conserved than the LTR region. With the exeptions of some nucleotides which are not verified yet, comparison of the LTR region showed the highest homology between the Iranian and Japanese isolate. As for the env gene, the Iranian isolate was most similar to the Japanese isolates. Therefore, these results suggest that the Iranian isolate is phylogenetically most similar to the prototypic Japanese/African isolates, and divergent from the Melanesian variants of HTLV-I.

There are two theories about HTLV-I origin and the way it was spread to the world. Gallo suggests that it originated in Africa and spread to the new world and Japan, while Saskena et al. suggest that HTLV-I originated in the Indo-Malay region (56,67-68). Based on the comparison of the LTR sequences, our results support the theory of HTLV-I may have originated in Africa, since we found a higher degree of homology between the HE sequence and the Zairian isolate, than with the sequence of Papua New Guinea isolate. It is likely that HTLV-I may have reached Iran from Africa by overland trade routes traversing through Mashad. These routes may have extended to the Far East (Japan) rather than the more commonly held notion of seaborne extension to Southern Japan, thus linking the African/Iranian and Japanese isolates. Comparative sequence analysis once again confirms the striking homology between distant HTLV-I isolates compared to HIV-I.

Over a decade following initial identification of the role of the retrovirus in human disease, HTLV-I remains as the only example for human retroviral leukemogenesis. Several viruses can be derived from the HTLV model. First, leukemia may be the rare consequence of infection with a relatively common virus. Second, a long latency period may be involved following acquisition of infection, and the leukemogenic process may involve multiple steps beyond initial infection. Third, the mechanism by which the virus may predispose to development of leukemia may be different from that seen with other retroviruses, and may involve transacting viral genes, either regulators of DNA transcription or regulators of viral mRNA expression. Finally, the role of HTLV-I as an etiologic agent in both HAM and ATLL suggests that viruses, not thought to have oncogenic properties, may on occasion predispose to malignancy. The overall lesson from the HTLV experience is that when suitable or suggestive epidemiologic factors are found, such as ethnic, geographic and/or familial clustering in the absence of a known inheritance pattern, a search for underlying viruses should be initiated. The search would involve removal of malignant tissue, when possible, for culture in vitro and a search using available molecular and serologic probes.

٣'

÷.,

REFERENCES:

- Poiesz BJ, Ruscetti FW, Gazdar AF, Bunn PA, Minna JD, Gallo RC. Detection and isolation of type C retrovirus particles from fresh and cultured lymphocytes of a patient with cutaneous T-cell lymphoma. Proc Natl Acad Sci USA 1980;77:7415-7419.
- Yoshida M, Seiki M, Yamaguchi K, Takatsuki K. Monoclonal integration of human T-cell leukemia provirus in all primary tumors of adult T-cell leukemia suggests causative role of human T-cell leukemia virus in the disease. Proc Natl Acad Sci USA 1984;81:2534-2537.
- 3. Hinuma Y, Komoda H, Chosa T, et al. Antibodies to adult T-cell leukemia-virus-associated antigen (ATLA) in sera from patients with ATL and controls in Japan: a nationwide seroepidemiologic study. Int J Cancer 1982;29:631-635.
- Hinuma Y. Seroepidemiology of adult T-cell leukemia virus (HTLV-II ATLV) origin of virus carriers in Japan. AIDS Res 1986;2:517-522.
- Riggar RJ, Melbye M, Sarin PS, et al. ELISA HTLV retrovirus antibody reactivity associated with malaria and immune complexes in healthy Africans. Lancet 1985;ii:520-523.
- Blattner WA, Saxinger CA, Clark J. et al. Human T-cell leukemia/lymphoma virus-associated lymphoreticular neoplasia in Jamaica. Lancet 1983;ii:61-61.
- Bunn PA Jr., Schechter GP, Jaffe E. et al. Clinical course of retrovirus-associated adult T-cell lymphoma in the United States. N Engl J Med 1983;309:257-264.
- Robert-Guroff M, Weiss SH, Giron JA. et al. Prevalence of antibodies to HTLV-I, -II, and-III in intravenous drug abusers from an AIDS endemic region. JAMA 1986;255:3133-3137.
- Uchiyama T, Yodoi J, Sagawa K, Takatsuki K, Uchino H. Adult T-cell leukemia: Clinical and hematologic features of 16 cases. Blood 1977;50:481-492.
- 10. Blattner WA, Nomura A, Clark JW, Ho GYF, Nakao Y, Gallo R, Robert-Guroff M. Modes of transmission and evidence for viral latency from studies of human T-cell lymphotropic virus type I in Japanese migrant populations in Hawaii. Proc Natl Acad Sci USA 1986;83:4895-4898.

<u>REFERENCES</u> (Cont'd):

- 11. Brousset P, Chittal S, Schlaifer D. et al. Detection of Epstein-Barr virus messenger RNA in Reed-Sternberg cells of Hodgkin's disease by in situ hybridization with bioinylated probes on specially processed modified acetone methyl benzoate xylene (ModAMeX) sections. Blood 1991;77:1781-1786.
- 12. Pallesen G, Hamilton-Dutoit SJ, Rowe M, Young LS. Expression of Epstein-Barr virus latent gene products in tumour cells of Hodgkin's disease. Lancet 1991;337:320-322.
- 13. Herbst H, Niedobitek G, Kneba M. et al. High incidence of Epstein-Barr virus genomes in Hodgkin's disease. Am J Pathol 1990;137:13-18.
- 14. Kubonishi I, Equchi T, Kanzaki T. et al. EBV and Hodgkin's cells. Br J Haematol 1990;75:286-287.
- 15. Feinmesser R, Miyazaki I, Cheung R, Freeman J, Noyek AM, Dosch H-M. Diagnosis of nasopharyngeal carcinoma by DNA amplification of tissue obtained by fine-needle aspiration. New Engl J Med 1992;326:17-21.
- 16. Sidi Y, Meytes D, Shohat B, Fenig E, Weisbrot Y, Lee H, Pinkhas J, Rosenblatt J. Adult T-cell lymphoma in Israeli patients of Iranian origin. Cancer 1990;65:590-593.
- 17. Lilleyman JS. Leukaemia "outbreaks". Lancet 1988;ii:1021.
- 18. Till MM, Hardisty R, Pike MC. et al. Childhood leukaemia in greater London: A search for evidence of clustering. Br Med J 1967;iii:755-758.
- 20. Linet MS, Devesa SS. Descriptive epidemiology of childhood leukaemia. Br J Cancer 1991;63:424-429.
- 21. Alexander FR, Ricketts TJ, McKinney PA, Cartwright RA. Community lifestyle characteristics and risk of acute lymphoblastic leukaemia in children. Lancet 1990;336:1461-1465.

• ج

22. Editorial. Childhood leukaemia: An infectious disease ? Lancet 1990;336:1477-1479. REFERENCES (Cont'd):

- 23. Okpala IE, Abayomi NA, Gevao SM. et al. Changing patterns of acute lymphoblastic leukaemia in Nigeria. Tokai J Exp Clin Med 1989;14:301-307.
- 24. Laor J, Langeuitz P, Tran H. et al. HTLV-I-associated T-cell leukemia lymphoma in Israel. Isr J Med Sci 1988;24:397-400.
- 25. Kark JD, Kedem R, Revach M. Medical examination of Israeli 17-years-old before military service as a national resource for health information. Isr J Med Sci 1986;22:318-325.
- 26. Seidman DS, Laor A, Gale R, Stevenson DK, Danon YL. Is low birthweight a risk factor for asthma during adolescence ? Arch Dis Childhood 1991;66:584-587.
- 27. Seidman DS, Paz I, Laor A, Gale R, Stevenson DK, Danon YL. Apgar scores and cognitive performance at 17-years of age. Obst Gynecol 1991;77:875-878.
- 28. Seidman DS, Laor A, Gale R, Stevenson DK, Mashiach S, Danon YL. Birthweight, current body weight and blood pressure in late adolescence. Br Med J 1991;302:1235-1237.
- 29. Seidman DS, Laor A, Gale R, Stevenson DK, Mashiach S, Danon YL. Long-term effects of vacuum and forceps deliveries. Lancet 1991;337:1583-1585.
- 30. Laor A, Seidman DS, Danon YL. Changes in body height among selected ethnic groups. J Epidem & Commun Health 1991;45:169-171.
- 31. Seidman DS, Laor A, Gale R, Stevenson DK, Danon YL. A longitudinal study of birth weight and being overweight in late adolescence. Am J Dis Child 1991:145:782-785.
- 32. Seidman DS, Laor A, Gale R, Stevenson DK, Danon YL. Pre-eclampsia and offspring's blood pressure, cognitive ability and physical development at 17-years of age. Br J Obstet Gynaecol 1991;98:1009-1014.
- 33. Seidman DS, Laor A, Gale R, Stevenson DK, Mashiach S, Danon YL. Long-term effects of obstetical intervention: a population based study. Obstet Gynecol 1992;47:33-34.
- 34. Seidman DS, Laor A, Gale R, Stevenson DK, Danon YL. Birthweight and intellectual performance in late adolescence. Obstet Gynecol 1992;79:543-546.
- 35. Yoshida M, Seiki M. Recent advances in the molecular biology of HTLV-I: Trans-activation of viral and cellular genes. Ann. Rev. Immunol., 1987;5:541-559.

<u>REFERENCES</u> (Cont'd):

- 36. Gessain A, Vernant JC, Maurs L, Barin F, Gout O, Calender A, de The G. Antibodies to human T-lymphotropic virus type-1 in patients with tropical spastic paraparesis. Lancet 1985;ii:407-409.
- 37. Seiki M, Hattori S, Hirayama Y, Yoshida M. Human adult T-cell leukemia virus: Complete nucleotide sequence of the provirus genome integrated in leukemia cell DNA. Proc Natl Acad Sci USA 1983;80:3618-3622.
- 38. Slamon DJ, Shimotohno K, Cline MJ, Golde DW, Chen ISY. Identification of the putative transforming protein of the human T-cell leukemia virus HTLV-I and HTLV-II. Science 1984;226:61-65.
- 39. Lee TH, Coligan JE, Sodroski JG, Haseltine WA, Salahuddin SZ, Wong-Staal F, Gallo RC, Essex M. Antigens encoded by the 3'-terminal region of human T-cell leukemia virus: Evidence for a functional gene. Science 1984;226:57-61.
- 40. Sodroski J, Rosen C, Goh WC, Haseltine W. A transcriptional activator protein encoded by the X-lor region of the human T-cell leukemia virus. Science 1985;228:1430-1434.
- 41. Cann AJ, Rosenblatt JD, Wachsman W, Shah NP, Chen ISY. Identification of the gene responsible for human T-cell leukemia virus transcriptional regulation. Nature 1985;318:571-574.
- 42. Greene WC. The human interleukin-2 receptor. Normal and abnormal expression in T cells and in leukemias induced by the human T-lymphotropic retroviruses. Ann Intern Med 1986;105:560-572.
- 43. Siekevitz M, Feinberg MB, Holbrook N, Wong-Staal F, Greene WC. Activation of interleukin 2 and interleukin 2 receptor (Tac) promoter expression by the trans-activator (tat) gene product of human T-cell leukemia virus, type I. Proc Natl Acad Sci USA 1987;84:5389-5393.
- 44. Cross SL, Feinberg MB, Wolf JB, Holbrook NJ, Wong-Staal F, Leonard WJ. Regulation of the human interleukin-2 receptor alpha promoter: Activation of a nonfunctional promoter by the transactivator gene of HTLV-I. Cell 1987:49:47-56.
- 45. Nimer SD, Gasson JC, Hu K, Smalberg I, Williams JL, Chen ISY, Rosenblatt RD. Activation of the GM-CSF promoter by HTLV-I and -II tax protein. Oncogene 1989;4:671-676.

REFERENCES (Cont'd):

- 46. Nyborg JK, Dynan WS, Chen ISY, Wachsman W. Binding of host-cell factors to DNA sequences in the long terminal repeat of human T-cell leukemia virus type I: Implications for viral gene expression. Proc Natl Acad Sci USA 1988:85:1457-1461.
- 47. Ohtani K, Nakamura M, Saito S, Noda T, Ito Y, Sugamura K, Hinuma Y. Identification of two distinct elements in the long terminal repeat of HTLV-I responsible for maximum gene expression. Eur Mol Biol Org (EMBO) J 1987;6:389-395.
- 48. Hunsmann G, Schneider J, Schmitt J, Yamamoto N. Detection of serum antibodies to adult T-cell leukemia virus in non-human primates and in people from Africa. Int J Cancer 1983;32:329-332.
- 49. Hinuma Y, Nagata K, Misoka M, Nakai M, Matsumoto T, Kinoshita K, Shirakawa S, Miyoshi I. Adult T-cell leukemia: Antigen in an ATL cell line and detection of antibodies to the antigen in human sera. Proc Natl Acad Sci USA 1981;73:6476-6480.
- 50. Blattner WA, Kalyanaraman VS, Robert-Guroff M, Lister A, Galton DAG, Sarin PSD, Crawford MH, Catovski D, Greaves M, Gallo RC. The human type C retrovirus, HTLV, in blacks from the Caribbean region, and relationship to adult T cell leukemia/lymphoma. Int J Cancer 1982;30:257-264.
- 51. Hattori T, Uchiyama T, Topibana K, Takatsuki K, Uchino H. Surface phenotype of Japanese adult T-cell leukemia cells characterized by monoclonal antibodies. Blood 1981;58:645-647.
- 52. Lee H, Swanson P, Shorty VS, Zack JA, Rosenblatt JD, Chen ISY. High rate of HTLV-II infection in seropositive IV drug abusers from New Orleans. Science 1989;244:471-475.
- 53. Seiki M, Hattori S, Yoshida M. Human adult T-cell leukemia virus: molecular cloning of the provirus DNA and the unique terminal structure. Proc Natl Acad Sci USA 1982;79:6899-6902.
- 54. Seiki M, Hattori S, Hirayama Y, Yoshida M. Human adult T-cell leukemia virus: complete nucleotide sequence of the provirus genome integrated in leukemia cell DNA. Proc Natl Acad Sci USA 1983;80:3618-3622.
- 55. Cann AJ, Chen ISY. Human T-cell leukemia virus types I and II. In: Fiëlds BN, Knipe DM, Chanock RM. et al eds. Fileds virology. 2nd ed. New York, Raven Press, 1990:1501-1527.

<u>REFERENCES</u> (Cont'd):

- 56. Lee HH, Swanson P, Rosenblatt JD, Chen ISY, Sherwood WC, Smith DE, Tegtmeier GE, fernando LP, Fang CT, Øsame M, Kleinman SH. Antibody screening of U.S. blood donors reveals similar prevalence of HTLV-I and HTLV-II infection in association with different risk factors. Lancet 1991;337:1435-1439.
- 57. Saksena NK, Sherman MP, Yanagihara R, Dube DK, Poiesz BJ. LTR sequence and phylogenetic analysis of a newly discovered variant of HTLV-I isolated from the Hagahai of Papua New Guinea. J Virol 1992;189:1-9.
- 58. Ratner L, Philpott T, Trowbridge DB. Nucleotide sequence analysis of isolates of human T-lymphotropic virus type I of diverse geographical origins. AIDS Res and Human Retroiruses 1991;7:923-941.
- 59. Kinoshita T, Tsujimoto T, Shimotohno K. Sequence variations in LTR and env regions of HTLV-I do not discriminate between the virus from patients with HTLV-I associated myelopathy and adult T-cell leukemia. Int J Cancer 1991;4:491-495.
- 60. Daenke S, Nightigale S, Cruickshank JK, Bangham CRM. Sequence variants of human T-cell lymhotropic virus type I from patients with tropical spastic paraparesis and adult T-cell leukemia do not distinguish neurological from leukemic isolates. J Virol 1990;64:1278-1282.
- 61. Paine E, Garcia J, Philpott TC, Shaw G, Ratner L. Limited sequence variation in human T-lymphotropic virus type I isolates from North American and African patients. Virol 1991;182:111-123.
- 62. Encyclopaedia Judaica 1972;11:1399-1400.
- 63. Ehrlich GD, Glaser JB, LeVigne K. et al. Prevalence of human T-cell leukemia lymphoma virus (HTLV) type II infection among high-risk individuals type specific identification of HTLV's by polymerase chain reaction. Blood 1989;74:1658-1661.
- 64. Krivine A, Firtion G, Cao L, Francoual C. et al. HIV replication during the first weeks of life. Lancet 1992;339:1187-1189.
- 65. Ehrlich GD, Glaser JR, Abbott MA. et al. Detection of anti-HTLV-I Tax antibodies in HTLV-I enzyme-linked immunosorbent assay-negative individuals. Blood 1989;74:1066-1072.
- 66. Kamihara S, Toriya K, Amagasaki T. et al. Antibodies against p-10tax gene product of human T-lymphotrophic virus type I under various conditions of HTLV-I infection. Jpn J Cancer Res 1989;80:1066-1071.

<u>REFERENCES</u> (Cont'd):

- 67. Ben-Ishai Z, Haas M, Triglia D. et al. Human T-cell lymphotrophic virus type-I antibodies in Falashas and other ethnic groups in Israel. Nature 1985;315:665-666.
- 68. Meytes D, Shochat B, Lee H, Nadel G, Sidi Y, Cerney M, Swanson P, Shaklai M, Kilim Y, Elgat M, Chin E, Danon YL, Rosenblatt JP. Serological and molecular survey for HTLV-I infection in a newly identified high-risk group in Israel. Lancet 1991; 336:1533-1535.
- 69. Gallo RC. Human retroviruses: a decade of discovery and link with human disease. J Infect Dis 1991;164:235-243.
- 70. Rosenblatt JP, Danon Y, Black AC. A decade with HTLV-I/HTLV-II: lessons in viral leukemogenesis. Leukemia 1992;6:18-23.
- 71. Y.L. Danon, Y. Kilim, J.D. Rosenblatt, Molecular characterization of HTLV-I infection in Israel. <u>Ann. N.Y. Acad. Sci.</u> (In Press - 1993)
- 72. Y. Kilim, J.D. Rosenblatt, Y.L. Danon, Molecular characterization of HTLV-I infection in Israel. <u>Isr J Med Sci</u> (Submitted - 1993)

 \mathbf{t}^{*}

CONCLUSIONS

During the two years of the project, summarized in this final report, we have studied more than 11.000 blood samples for HTLV-I serology and molecular detection and characterization of the virus.

We have been successful in identifying a new high risk population for HTLV-I in the Middle East, a population of immigrant Jews from Iran, from the region of Khurusan (North East Iran - NEI) and mainly from the town of Mashad. It seems that HTLV-I infection is prevalent among this special community, like some other areas of the world, like southern Japan and Pacific Islands, several Caribbean countries and central Africa. The prevalence of HTLV-I infection in native Iranians and large population of Iranians and Mashadies residing in North America and Europe has not been studied yet.

Because persons infected with HTLV-I may be screened medically for HTLV-I associated diseases and because changes in breast feeding, sexual behavior, and blood donation can interrupt the transmission of the virus, studies of this kind have a major importance in preventive medicine.

This special population, with first description and virus isolates of Adult T cell Leukemia (ATTL) and HTLV-I Associated Myelopathy (HAM) gives a special and new insight on HTLV-I epidemiology, with special interest focusing on a concentration of HTLV-I positive patients, in a very senior population residing in a geriatric center.

Various serological and molecular screening methods, including enzyme-linked immunosorbent assay (ELISA) for anti-HTLV-I, ELISA for an antibody for recombinant HTLV-I p40tax protein, and molecular detection of infection by a polymerase chain reaction (PCR) amplification of HTLV-I proviral DNA, from peripheral mononuclear cells DNA, were used.

By HTLV-I ELISA the overall rate of infection was 12.2% among immigrants from Khurusan (northeastern Iran), non HTLV-I carriers were detected in a general survey of the population and other high risk groups, including other Iranian and Ethiopian Jews, as well as some clinical conditions such as ATLL, other T-cell malignancies and haemodialysis patients. We have found an unexplained clustering of HTLV-I infection in a cohort of 32 elderly women of similar geographic origin (Mashad) in a home of senior citizens, 14 were seropositive in ELISA and 19 of 28 were positive by PCR. These findings and this newly identified high risk population suggests that, in addition to ELISA, other screening techniques may be required to detect all carriers in high risk populations.

We have done detailed sequencing work on an HAM Iranian isolate of HTLV-I virus.

In this study we determined the sequence of two regions in HTLV-I isolated in Israel from an HAM patient belonging to this community: the LTR region and the env gene. Several studies have indicated a high degree of homology among HTLV-I isolates (>96%) and demonstrated that differences between variants are primarily associated with their geographical origin. We compared HTLV-I in an Iranian immigrant with

HAM to the Japanese, African and Papua New Guinea isolates. A high level of homology of about 98% was observed to the Japanese isolate, while the African and Papua New Guinea isolates were more divergent with sequence homology of 95% and 91% in the LTR respectively.

Comparison of HTLV-I env gene to the Japanese and African isolates showed homology at the nucleotide level of about 98-99%. From these results we can conclude that the env gene is more conserved than the LTR region.

With the exeptions of some nucleotides which are not verified yet, comparison of hte LTR region showed the highest homology between the Iranian and Japanese isolate. As for the env gene, the Iranian isolate was most similar of the Japanese isolates. Therefore, these results suggest that the Iranian isolate is phylogenetically most similar to the prototypic Japanese/African isolates, and divergent from the Melanesian variants of HTLV-I.

There are two theories about HTLV-I origin and the way it was spread to the world. Gallo suggests that it originated in Africa and spread to the new world and Japan, while Saskena et al. suggest that HTLV-I originated in the Indo-Malay region. Based on the comparison of the LTR sequences, our results support the theory of HTLV-I may have originated in Africa, since we found a higher degree of homology between the HE sequence and the Zairian isolate, than with the sequence of Papua new Guinea isolate. It is likely that HTLV-I may have reached Iran from Africa by overland trade routes traversing through Mashad. These routes may have extended to the Far East (Japan) rather than the more commonly held notion of seaborne extension to Southern Japan, thus linking the African/Iranian and Japanese isolates. Comparative sequence analysis once again confirms the striking homology between distant HTLV-I isolates compared to HIV-I.

HTLV-I has been recently associated with some new additional disease, like HTLV-I associated arthropathy, HTLV-I polymyositis, HTLV-I uneitis, and pediatric HTLV-I infectious dermatitis. Unfortunately, we have not been able at this stage to detect any of those, and our studies in IDDM newly diagnosed patients, non-Burkitt's Lymphoma patients, and Psoriasis/Parapsoriasis patients were all negative.

BIBLIOGRAPHY OF PUBLICATIONS AND MEETING ABSTRACTS

- Rosenblatt JP, Danon Y, Black AC. A decade with HTLV-I/HTLV-II: lessons in viral leukemogenesis. Leukemia 1992;6:18-23.
- Meytes D, Shochat B, Lee H, Nadel G, Sidi Y, Cerney M, Swanson P, Shaklai M, Kilim Y, Elgat M, Chin E, Danon YL, Rosenblatt JP. Serological and molecular survey for HTLV-I infection in a newly identified high-risk group in Israel. Lancet 1991; 336:1533-1535.
- Y.L. Danon, T-lymphocytes in childhood leukemia <u>Pediatr. Hematol. Oncol.</u>, 10:9-10, 1993
- 4. Y.L. Danon, Y. Kilim, J.D. Rosenblatt, Molecular characterization of HTLV-I infection in Israel. <u>Ann. N.Y. Acad. Sci.</u> (In Press - 1993)
- 5. Y. Kilim, J.D. Rosenblatt, Y.L. Danon, Molecular characterization of HTLV-I infection in Israel. <u>Isr J Med Sci</u> (Submitted - 1993)
- 6. J. Rosenblatt, D. Meites, Y. Sidi, Y.L. Danon, HTLV-I induced T-Cell lymphoma in Israeli patients of Iranian origin. <u>The 3rd Int. Symp. Immune Def.</u>, 20-21 April, 1990, Warsaw, Poland.
- 7. Y. Kilim, J.D. Rosenblatt, D. Meytes, D. Stephens, H. Lee, Y. Danon, Molecular characterization of Iranian HTLV-I isolates. <u>5th Int. Conf. on Human Retrovirol.: HTLV</u>, May 11-13, 1992, Japan.
- Y.L. Danon, Y. Kilim, J. Rosenblatt Epidemiologic and Molecular Characterization of HTLV-I Infection in Israel. <u>UJA Medical Symposia</u>, Jerusalem - August, 1992.
- 9. Y.L. Danon, Y. Kilim, J. Rosenblatt, Epidemiologic and molecular characterization of HTLV-I infection in Israel. <u>8th Int. Congress Immunol.</u>, Budapest, August, 1992.
- 10. Y.L. Danon, Y. Kilim, J. Rosenblatt, Epidemiologic and molecular characterization of HTLV-I infection in Israel. <u>2nd Int. Congress on Aids in Asia and the Pacific</u>,
- 11. Y.L. Danon, Y. Kilim, J. Rosenblatt, Molecular characterization of HTLV-I infection in Israel. <u>Viruses & Virus-like Agents in Disease</u>, March 7-9, 1993, Basel Switzerland
BIBLIOGRAPHY OF PUBLICATIONS AND MEETING ABSTRACTS (Cont'd)

- 12. Y.L. Danon, Y. Kilim, J. Rosenblatt, Molecular characteristics of HTLV-I infection in newly characterised high risk group of carriers in the middle east. <u>27th Ann. Sci. Meeting</u>, April 14-17, 1993 Heidelberg Cermany
- 13. Y.L. Danon, Y. Kilim, J. Rosenblatt, Epidemiologic and Molecular characterization of HTLV-1 infection in Israel <u>The New York Acad. of Sci.</u>, June 2-5, 1993.
- 14. Y.L. Danon, Y. Kilim, J. Rosenblatt, Molecular characterization of HTLV-I infection in Israel. <u>IXth Int. Conf. on AIDS</u>, Berlin June 7-11, 1993.
- 15. Y. Kilim, M. Karp, Y.L. Danon, HTLV-I virus in insulin-dependent diabetes mellitus. <u>19th Ann. Meeting of the Int. Study Group of Diabetes in Children</u> and Adolescents, September 2-6, 1993, Greece
- 16. Y.L. Danon, Y. Kilim, J. Rosenblatt, Epidemiologic and molecular characterization of new HTLV-I infection focus in the Middle East. <u>VIIIth Int. Conf. on AIDS in Africa</u>, 12-16 Dec. 1993 - Morocco

۳١.

LIST OF PERSONNEL RECEIVING PAY ON THIS CONTRACT:

- 1. Benjamin Kornbrot, M.D.
- 2. Elisabeth Kaminsky, M.Sc.
- 3. Yael Kilim, M.Sc.
- 4. Merav Crook

13

HTLV-I INDUCED T-CELL LYMPHOMA IN ISRAELI PATIENTS OF IRANIAN ORICIN

J. Rosenblatt, D. Meites, Y. Sidi, Y.L. Danon Rogoff-Wellcome Medical Research Institute, Dept. of Medicine and Div. of Pediatric Immunology, Edith Wolfson Hospital and Beilinson

Medical Center Tel-Aviv University Sackler School of Medicine, Petah-Tikva 49100 ISRAEL.

T cell lymphoma-leukemia (ATL) is one of the several clinical entities linked to human T cell lymphotropic virus (HTLV-I). Few geographic endemic concentrations of HTLV-I infection were already described: The Ryukyu Islands in Southern Japan, Central Africa and the Caribbean Islands. This is the first description of endemic HTLV focus in the Middle East. The prevalence and clinical presentations of ATL in Israel were studied. We have diagnosed four Israeli Jewish ATL patients and one HAM (HTLV-I - Associated Myelopathy) in a nationwide survey performed in 1986-1990. In three of the patients evidence for HTLV-I infection was obtained. All those patients immigrated to Israel from the same region in Central Iran. The nationwide survey and the clinical course of this new group of patients will be presented.

(

(((

((

Abstract Reproduction Form

Japan

Type abstract within blue rectangle. DO NOT FOLD THIS FORM.



MOLECULAR CHARACTERIZATION OF IRANIAN HTLV-I ISOLATES

Y. Kilim¹, J.D. Rosenblatt², D. Meytes³, D. Stephens², H. Lee⁴, Y. Danon⁴

Children's Medical Center of Israel, Petach, Tiqva, Israel¹, UCLA School of Medicine, Los Angeles, CA², Edith Wolfson Hospital, Holon, Israel³, Abbott Laboratories, N. Chicago, IL⁴, USA

We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Mashad in Northeastern Iran (Lancet 336:1533-1535, 1990). We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-J seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax/rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. An immortalized Tcell line was derived from a Mashadi HTLV-I carrier. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Further characterization of Iranian HTLV-I isolates is underway. (Supported by the Rashi Foundation, Doron Foundation. ICRF, and a grant from the USAMRDC).

To be completed by presenting author: Yehuda Danon, M.D.	
Elen Mart	••••••••••••••••••••••••••••••••••••••
Athlation(University.Company.etc.) Beilleschi Medical Center of Israel	· · · · · · · · · · · · · · · · · · ·
Mailing Address Petah Tikva 49100, ISRAEL	
Country	
Phone 972 3 924-7515 Fax: Same T	eier:
The number of the main polopony for we above abstract B. A P I would	Id prefer a poster presentation
	_
Sprature	***************************************

EPIDEMIOLOGIC AND MOLECULAR CHARACTERIZATION OF HTLV-I INFECTION IN ISRAEL

UTA

Yehuda L. Danon, Yael Kilim and J. Rosenblatt, Kipper Institute of Child Immunology, Children's Medical Ctr. of Israel, Sackler School of Medicine, Tel-Aviv Univ. Kaplan Str. #14-16, Petah-Tikva 49100 Israel.

Human T-cell leukemia viruses type I (HTLV-I) and type II (HTLV-II) are the two human retroviruses directly implicated in pathogenesis of human leukemia. HTLV-I has been linked to adult T-cell leukemia/ lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. Epidemiologic and molecular studies of both viruses have identified several themes underlying the leukemogenic process. Leukemia is a rare consequence of infection, and both viral infection and secondary transforming events appear to be involved. Due to the absence of a virally encoded oncogene, or preferred viral integration sites, novel mechanisms of leukemogenesis must be invoked. We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Mashad in Northeastern Iran (Lancet 336:1533-1535,1990). We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. An immortalized T-cell line was derived from a Mashadi HTLV-I carrier. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 differrent restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Further characterization of Iranian HTLV-I isolates is underway. A potential role for the HTLV-I/II transregulatory proteins Tax and Rex in leukemogenesis is discussed. Insights derived from the study of HTLV-I/II can be applied to the search for other oncogenic viruses.

(Supported by the Rashi Foundation, Doron Foundation, ICRF, and a grant from the USAMRDC).

11

V

Yehuda L. Danon, Yael Kilim and Joseph Rosenblatt, Kipper Institute of Child Immunology, Children's Medical Ctr. of Israel, Sackler School of Medicine, Tel-Aviv Univ. Kaplan Str. #14-16, Petah-Tikva 49100 Israel.

Budars

177

Human T-cell leukemia viruses type I-(HTLV-I) and type II (HTLV-II) are the two human retroviruses directly implicated in pathogenesis of human leukemia. HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. Epidemiologic and molecular studies of both viruses have identified several themes underlying the leukemogenic process. Leukemia is a rare consequence of infection, and both viral infection and secondary transforming events appear to be involved. Due to the absence of a virally encoded oncogene, or preferred viral integration sites, novel mechanisms of leukemogenesis must be invoked. We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Mashad in Northeastern Iran (Lancet 336:1533-1535,1990). We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. An immortalized T-cell line was derived from a Mashadi HTLV-I carrier. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Further characterization of Iranian HTLV-I isolates is underway. A potential role for the HTLV-I/II transregulatory proteins Tax and Rex in leukemogenesis is discussed. Insights derived from the study of HTLV-I/II can be applied to the search for other oncogenic viruses.

(Supported by the Rashi Foundation, Doron Foundation, ICRF, and a grant from the USAMRDC).

<u>س</u>،

X

)



Original Abstract Form

Viruses and Virus-Like Agents In Disease

Nolecular characterization of HTLV-I infection in Israel

Yehuda L. Danon, Yael Kilim and Joseph Rosenblatt Kipper Inst. of Child Immunology, Children's Med. Ctr. of Israel, Sackler School of Med., Tel-Aviv Univ. Israel

1/

 \mathcal{W}

HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Northeastern Iran. We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Insights derived from the study of HTLV-I/II can be applied to the search for other retroviruses and oncogenic viruses.

Supported in part by USAMRDC Grant # DAMD17-91-C-1001

I wish to submit an abstract for presentation as poster on

Monday, March 8, 1993 🔲 Tuesday, March 9, 1993

Key Words

Ć

((

Name of Presenting Author	Yehu
	Dire
Company/Institute	Medi

Yehuda L. Danon, M.D. Director, The Children's Medical Center of Israel Beilinson Medical Center Petah-Tikva 49100 ISRAEL

Mailing Address City

Department

Country

Postal Code

Telephone **Telefax** 12-3-930 Signature Deadline for submission of abstracts is December 15, 1992

Please send the original abstract form and 3 photocopies by airmail to: S. Karger AG, 1993 Congress, P.O. Box, CH-4009 Basel, Switzerland

MOLECULAR CHARACTERISTICS OF HTLV-I INFECTION IN NEWLY CHARACTE-RISED HIGH RISK GROUP OF CARRIERS IN THE MIDDLE EAST.

Lleidelpera.

Ŋ

-**)**

HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Mashad in Northeastern Iran. We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Insights derived from the study of HTLV-I/II can be applied to the search for other retroviruses and oncogenic viruses.

(Supported by the Rashi Foundation, Doron Foundation, ICRF, and a grant from the USAMRDC).

My Fred.

ł

ĺ

EPIDEMIOLOGIC AND MOLECULAR CHARACTERIZATION OF HTLV-I INFECTION IN ISRAEL

12

ς.,

Yehuda L. Danon, Yael Kilim and Joseph Rosenblatt, Kipper Institute of Child Immunology, Children's Medical Center of Israel, Sackler School of Medicine, Tel-Aviv University Israel

HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. We recently reported a high rate of HTLV-I sero-positivity among immigrants to Israel from Mashad in Northeastern Iran. We have now characterized the Iranian HTLV-I isolate using a combination of Southern Jotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. The integrated HTLV-I provirus. Southern cell line contained blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Insights derived from the study of HTLV- I/II can be applied to the search for other retroviruses and oncogenic viruses.

(Supported by the USAMRDC and Doron Foundation)

IXth International Conference on AIDS, Berlin, June 7 – 11, 1993

in affiliation with the IVth STD World Congress

ABSTRACT FORM

Only this official form is acceptable as the original submission (no facsimile transmission). This form should be accompanied by 5 photocopies.

Choice of Topics

IMPORTANT:

Please indicate your preference according to the list of topics on the reverse side (A1 - D38).

1st choice	
2nd choice	
STD*	
I prefer: poster presentation	
oral presentation	

· Choice of Key Words

See List of Key Words in "Call for Abstracts" and type in corresponding numbers.

Name

Affiliation

Address

Tel.

Fax

Mail original abstract and 5 photocopies to:

IXth International Conference on AIDS IVth STD World Congress Institute for Clinical and Experimental Virology of the Free University of Berlin Hindenburgdamm 27 D-1000 Berlin 45

MOLECULAR CHARACTERIZATION OF HTLV-I INFECTION IN ISRAEL Yehuda L. Danon, Yael Kilim and Joseph Rosenblatt, Kipper Inst. of Child Immunology, Children's Med. Ctr. of Israel, Sackler School of Med., Tel-Aviv Univ. Israel

HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. We recently reported a high rate of HTLV-I seropositivity among immigrants to Israel from Northeastern Iran. We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 10 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 10 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian and Japanese HTLV-I isolates. Insights derived from the study of HTLV- I/II can be applied to the search for other retroviruses and oncogenic viruses.

 	-		

Instructions:

1. Type within blue lines: title, author's name (6 or less), affiliations, city, state, country. (Underline name to indicate presenter; if there are more than 6 authors, type et al. to indicate additional authors). Do not include more than 6 authors' names.

2. Any handwritten symbols must be drawn in black ink.

3. REMEMBER: Your camera-ready abstract will be printed in the book of abstracts exactly as typed. No editorial corrections will be made.

4. For sample abstract see reverse side.

ABSTRACTS MUST BE RECEIVED NO LATER THAN JANUARY 15, 1993

* STD-related presentations should be assigned to the respective topic of each track.

HTLV-I VIRUS IN INSULIN-DEPENDENT DIABETES MELLITUS Y. Kilim. M.Sc.¹, M. Karp, M.D.² and Y.L. Danon, M.D.¹

Kipper Institute of Immunology,

Greece

²Institute of Pediatric and Adolescent Endocrinology,

... The Children's Medical Center of Israel, Petah-Tikva, Israel

Human T-cell Leukemia Virus-I has been linked to adult T cell leukemia/lymphoma (ATLL) and HTLV-II to some cases of chronic T cell leukemia. We have recently reported a high rate of HTLV-I seropositive among immigrants to Israel from northeastern Iran, and especially the town of Mashad.

To determine the frequency of antibodies to HTLV-I virus in Insulin-Dependent Diabetes Mellitus (IDDM) patients, sera from 56 newly onset IDDM patients were tested by an enzyme immunoassay. According to our method the reactivity of antibodies detected by enzyme immunoassay against HTLV-I encoded antigens was determined by an assay which employs recombinant HTLV-I antigens. No antibodies to HTLV-I were detected in all 56 patients studied. Proliferative response to various species of insulin was performed in 26 of those patients, 23 out of 26 showed a positive response. Sera from 56 newly onset IDDM patients were screened for ICA. ICA were detected in 32 (57.1%) of the 56 patients.

()

It seems that HTLV-I is playing no role in IDDM.

Supported in part by USAMRDC Grant # DAMD17-91-C-1001

ñ

VIIIth INTERNATIONAL CONFERENCE ON AIDS IN AFRICA & VIIIth AFRICAN CONFERENCE ON STDs



ABSTRACT FORM DEADLINE : 31 AUGUST 1993

EPIDEMIOLOGIC AND MOLECULAR CHARACTERIZATION OF NEW HTLV-I INFECTION FOCUS IN THE MIDDLE EAST

Yehuda L. Danon, Yael Kilim and Joseph Rosenblatt, Kipper Institute of Child Immunology, Children's Medical Center of Israel, Sackler School of Medicine, Tel-Aviv University Israel

HTLV-I has been linked to adult T-cell leukemia/lymphoma (ATLL), and HTLV-II to rare cases of chronic T-cell leukemia. We recently reported a high rate of HTLV-I sero-positivity among immigrants to Israel from Mashad in Northeastern Iran after a national serologic survey of blood donors. We have now characterized the Iranian HTLV-I isolate using a combination of Southern blotting, polymerase chain reaction (PCR) and sequencing. DNA from 17 HTLV-I seropositive Mashadi carriers was isolated from peripheral blood mononuclear cells and amplified by PCR using primers to the tax-rex region. All 17 samples were found to be HTLV-I and not HTLV-II using discriminatory PCR. The cell line contained integrated HTLV-I provirus. Southern blotting and restriction mapping of the Iranian isolate demonstrated marked similarity to published Japanese isolates using 10 different restriction enzymes. Amplification and preliminary sequencing of a 900 bp segment of env from two Mashadi isolates disclosed approximately 95% nucleic acid sequence homology with the published sequence of Japanese isolates, and >98% homology between two Mashadi isolates. These data indicate substantial conservation of sequence between Iranian, African (Zair) and Japanese HTLV-I isolates. Insights derived from the study of HTLV-I/II can be applied to the search for other retroviruses and oncogenic viruses. (Supported by the USAMRDC and Doron Foundation)

LANGUE OF PRESENTATION French English	PREFERENCE Oral presentation Poster
$(A) \begin{array}{c} TOPIC \\ B \\ C \\ D \\ E \end{array}$	KEY WORDS
B L L L L L	איזא ו-עדא (אאא ו-עדא) (דעדא)
AUTHOR'S NAME: POF. Y. DANON Adress: CHILDRENS MEDICAL CTR Telephone: P.O. Box SS9 Fax: PLONT. KV9 Y9202 ISRAEL Presenting author's signature:	OF ISRACL, Tel Av. VUNVERity. 972-3-9247515 Tix:
Airmail this original abstract form plus 5 photos to :- CONFERENCE SECRETARIAT : VIIIth International Conference on AIDS and STDs in Africa	FOR SECRETARIAT USE ONLY Abstract N°:
Ibis, place Charles Nicolle - Box: 1818 - Casablanca - Morocco Tel : (212) 2 29-53-25 / (212) 2 29-53-26 Fax : (212) 2 29-53-27 / (212) 2 29-53-28 Fex : 45 382 M	
REVIEWER USE ONLY : A	Score · Oral 5'
COMMENTS :	Poster
Type title, author's names affiliation	n, city, province/state, country and abstract. of presentating authors.

A Decade with HTLV-I/HTLV-II: Lessons in Viral Leukemogenesis

Joseph D. Rosenblatt¹, Yehuda, Danon², and Alexander C. Black¹

¹Division of Hematology-Oncology, Department of Medicine and Jonsson Comprehensive Cancer Center, UCLA School of Medicine, Los Angeles, CA, USA and the ²Kipper Institute of Child Immunology, Children's Medical Centre of Israel, Beilinson Medical Center, Petach Tiqva, Israel

INTRODUCTION

The past decade has seen myriad advances in detection and characterization of human retroviruses. It began with initial description of human T-cell leukemia virus type I (HTLV-I) by Poiesz and Gallo in the US and Yoshida in Japan, which pointed to the involvement of the human retrovirus, HTLV-I, in an unusual form of T-cell malignancy, adult T-cell leukemia/lymphoma (ATLL) (1,2). The identification of HTLV-I intensified the search for related viruses, and soon thereafter, human T-cell leukemia virus type II (HTLV-II) was described by Kalynaraman and Gallo in a cell line derived from a patient with a chronic T-cell leukemia with features of hairy-cell leukemia (3). The rapid identification of HTLV-II on the heels of HTLV-I led to speculation that a host of human oncogenic retroviruses would soon be identified. The subsequent discovery of human immunodefiency virus types 1 (HIV-1) and 2 (HIV-2) and their implication in acquired immunodeficiency syndrome (AIDS) accelerated the pace and intensity of the search for oncogenic viruses. It was soon recognized that leukemic cells in malignancies associated with HTLV-I and -II contained clonally integrated provirus; in effect, a signature for direct viral involvement in the oncogenic process. In contrast, neoplasms frequently seen in the setting of HIV-1 infection (e.g. Kaposi's sarcoma and/or highgrade B-cell lymphomas) did not appear arise as a direct consequence of viral transformation of HIV-1infected cells. At the end of the decade, only HTLV-I and -II remain clearly implicated as directly leukemogenic human retroviruses. Therefore, we believe that insights gleaned from investigation of these viruses can and should be applied to the search for other oncogenic retroviruses.

EPIDEMIOLOGICAL LESSONS

The epidemiological link between HTLV-I and ATLL resulted from two complementary events. The first, an appreciation by Uchiyama and Takatsuki that ATLL represented a unique clinical entity (4) allowed geographic localization of the disease to southern islands of Japan: Kyushu, Shikoku, and the Ryuku chain of islands. Development of serological assays for HTLV-I led to correlation of HTLV-I infection to the presence of malignancy, as well as a determination of modes of

LEUKEMIA © 1992 Macmillan Press Lid transmission (for review see 5). Epidemiological studies have suggested that exposure shortly after birth is a major risk factor for subsequent development of ATLL (5,6). In addition, these studies have demonstrated that twenty or more latent years may elapse between acquisition of infection and development of malignancy (5,6). Furthermore, only a minority (< 5%) of HTLV-I carriers actually develop ATLL (7), and ATLL as a consequence of transfusion-acquired HTLV-I is virtually unknown.

Hence, several general observations emerged from scrutiny of HTLV-I epidemiology: (a) leukemia may be an infrequent consequence of exposure to a fairly wide-spread virus; (b) leukemogenesis may depend on the timing and/or length of exposure, so that individuals infected in childhood may be at higher risk than those infected later in life; and (c) the long latency period suggests a multiple step process may be involved in leukemogenesis; while viral infection may be a prerequisite, it alone may be insufficient to produce the leukemic phenotype. These general epidemiologic features of ATLL suggest that a systematic re-evaluation of appropriate candidate malignancies and viruses may reveal other oncogenic viruses. Fairly prevalent or even ubiquitous viruses could conceivably manifest oncogenic potential in a sporadic fashion, and factors such as timing and length of exposure may be critical.

Careful cataloguing and description of clinical syndromes is essential to derive epidemiologic clues that may lead to virus identification. The recognition that non-Hedgkin's lymphomas could be divided into Tand B-cell subtypes and subsequent differentiation of ATLL from *mycosis fungoides* is a case in point. While ATLL was undoubtedly a frequent reason for in-patient hospitalizations in Japan prior to 1977, it was thought to be a variant of peripheral cutaneous T-cell lymphoma, and its characteristic features such as hypercalcemia and enhanced expression of interleukin 2 (IL-2) receptor alpha (IL-2R α) chain (Tac antigen) on the cell surface were initially overlooked. Recognition of the subtler clinical aspects of the syndrome allowed for subsequent epidemiologic observations (4).

In contrast to HTLV-I, it is premature to reach conclusions regarding pathogenesis by HTLV-II. Although originally isolated from the Mo T-cell line, a transformed T-cell line derived from the spleen of a patient with hairy-cell leukemia, the nature of the malignancy *in vivo* in the patient was not adequately addressed (8). We know that HTLV-I and -II can transform T-cell lines *in vitro*, and that the Mo T-cell line may have simply represented an outgrowth of HTLV-transformed cells *in vitro*. A second patient with HTLV-II and **hairy**-cell leukemia was found by our laboratory to have a biclonal lymphoproliferative

LEUKEMIA, Vol 6, Suppl 1, 1992; pp 18-23

Correspondence to: Joseph D. Rosenblatt, Division of Hematology-Oncology, Department of Medicine and Jonsson Comprehensive Cancer Center, UCLA School of Medicine, Los Angeles, CA 90024-1678, USA.

disorder in which a B-cell hairy-cell leukemia and a co-existant malignant CD8+ T-cell clone were observed (9,10). Oligoclonal integration of HTLV-II provirus into the CD8+ T-cells provided strong evidence for origin of malignancy in a virally infected cell. However, as additional cases of HTLV-II-induced malignancy have not been reported, there is considerable doubt as to whether we have as yet characterized the prototypic disease associated with HTLV-II.

An additional surprise that has emerged from epidemiological studies of HTLV has been the fact that screening procedures for HTLV-I identify a considerable number of crossreactive HTLV-II carriers. This raises the possibility that in the process of assaying for newly identified viruses, we may inadvertantly be assaying for a variety of crossreactive members of the same viral family. Specifically, intravenous drug abusers (IVDA) found to be seropositive for HTLV-I have been reported in several studies to have a higher incidence of HTLV-II infection and > 50% of seropositive random blood donors screened by HTLV-I ELISA were actually found by DNA amplification techniques to harbour HTLV-II (11-13). In the future, screening for newly identified viruses should be performed using both DNA amplification and serological techniques to avoid the initial confusion in delineating the epidemiology of HTLV-I and -II.

The HTLV-I model of malignancy as a rare consequence of infection with a prevalent virus suggests that careful molecular re-examination of the role of already recognized viruses in suspect malignancies is in order. In the case of HTLV-I, over a million infected individuals in Japan give rise to only 400-500 cases of ATLL per year. Hodgkin's disease (HD) may provide another case in point. The bimodal age distribution, prevalence, anecdotal descriptions of geographical clustering, and 'outbreaks' of HD suggest that an infectious agent may underlie pathogenesis (14-21). The increasingly frequent reports of Epstein-Barr virus (EBV) genome detection in some cases of HD suggest that EBV can be important in pathogenesis of a subset of HD patients (14-21). As another example, recognition that four of the last five cases of ATLL-like T-cell lymphoma in Israel occurred in Iranian immigrants from the northeastern city of Mashad, allowed identification of a new focus of HTLV-I infection (22,23). Recognition of geographic. familial and/or ethnic clustering of particular malignant disorders may yield important clues to viral etiology. It is important to note, however, that such time/space clustering may frequently relate to non-infectious risk factors rather than a virus. Some investigators have speculated on the likelihood of viral involvement in childhood acute lymphoblastic leukemia (ALL); however, the evidence is only mildly suggestive at best. Some studies suggest an association with geographic areas of high socioeconomic status, while others do not (24-29). Additional studies suggest a mild degree of clustering, particularly in children less than six years of age, although this remains controversial (29-31). Thus, little evidence points to an infectious cause or an underlying common leukemia virus in ALL. If a virus were involved,

VIRAL LEUKEMOGENESIS

analogy to ATLL would suggest that it might initially cause an insignificant acute infection that establishes latency and eventually leads to leukemia through secondary events. Given the lack of overt clustering, seroepidemiological studies are unlikely to settle the issue, and frank demonstration of molecular involvement of an infectious agent will likely be necessary.

New Molecular Mechanisms of Pathogenesis

The explosive growth in the study of oncogenes over the past decade came about as a result of recognition that in a nimal malignancies brought about by retroviral infection, the transduction of a cellular proto-oncogene and its inappropriate expression under control of the viral promoter was frequently observed in retrovirally induced tumors. A second type of molecular lesion frequently observed was integration of a retrovirus adjacent to a cellular oncogene and loss of normal patterns of prote-oncogene expression. These observations led to speculation that similar mechanisms may be operative in human malignancy. However, HTLV-I afforded a unique surprise, in that the viral sequences did not contain any transduced cellular sequences, and that integration sites appeared to be random. While clonal integration was observed by Yoshida and colleagues, the sites of integration often occurred on different chromosomes, and no specific integration patterns could be observed (2.32). These observations led to a search for new mechanisms of oncogenesis.

The demonstration by Seiki and colleagues of the unique coding regions located in the 3' end of the genome provided a first clue as to potential mechanisms (33). The 3' ends of the genome of HTLV-I, -II and the leukemogenic bovine leukemia virus (BLV) all contain coding sequences for at least two overlapping genes (34-37). These genes, known as tax and rex, are now known to encode a transcriptional and post-transcriptional regulator of viral expression. The initial discovery of the tax gene was surprising, in that such trans-acting transcriptional regulators had only been previously identified in DNA viruses such as herpes simplex virus, varicella-zoster virus, and adenovirus. The HTLV-I tax gene encodes a 40-kilodalton (kDa) protein, and the HTLV-II tax gene encodes a 37-kDa protein (34-37). These nuclear phosphoproteins are highly homologous, and have similar effects on transcription. Tax expression is necessary for transcriptional activation of virus replication and efficient RNA production from the viral long terminal repeat (LTR); however, an additional property noted early was the ability of Tax to trans-activate other viral and cellular promoters, including those involved in regulation of T-cell growth. Examples of such promoters include those for IL-2, IL-2R α , and the lymphokine, granulocyte-macrophage colony-stimulating factor (GM-CSF) (38-41). In the case of HTLV-I and -II, Tax appears to interact with three 21-base pair (bp) repeats located in the U3 portion of the viral LTR. Within the sequence of the 21-bp repeats are core sequences known to bind cellular transcription factors (42,43). A variety of these proteins have now been identified and

partially characterized.

In contrast to the HTLV promoters, Tax activation of the IL-2R α gene involves induced nuclear expression of a cellular DNA binding protein, which interacts with an NF- κ B-like enhancer (44). NF- κ B is a DNAbinding factor first shown to interact with the enhancer of the κ light chain immunoglobulin gene (45). Tax interaction with NF- κ B is thought to account for inducibility of some other cellular and viral promoters such as the HIV-1 promoter. The lack of an NF- κ B-like binding site in the HTLV-I/-II promoter and deletion of NF- κ B-like binding sites from the GM-CSF promoter with retention of response to Tax indicate that Tax may act via different pathways in different cellular and promoter contexts (41).

To date, only limited evidence directly implicates Tax in T-cell transformation. Introduction of Tax coding sequences under the control of a herpes saimiri vector has resulted in continuously proliferating T-cell lines in vitro, although the transformed cell lines appear to retain dependence on IL-2 for continued growth (46). Exp. ession of HTLV-I Tax under the control of the HTLV-I LTR in transgenic mice does not lead to Tax expression in T-cells, and T-cell malignancy is not observed (47). Some mice developed mesenchymal tumors reminiscent of neurofibromatosis, as well as muscular atrophy. Recent HTLV-I Tax transgenics under control of the T-cell-specific Thy-1 promoter also did not result in T-cell malignancy (48). Nevertheless, the promiscuous interaction of Tax with a variety of viral and cellular promoters suggests that it may play a pivotal role not only in the HTLV-I life-cycle, but also in definition of the malignant phenotype. Tax can trans-activate the IL-2R α gene. which offered an explanation for the high degree of Tac (high affinity IL-2 receptor) antigen expression in HTLV-I-transformed T-cells. Similarly, ectopic GM-CSF production due to Tax may cause eosinophilia, which is frequently seen in ATLL. In addition, Tax may also be involved in trans-activation of the parathyroid hormone-related protein (PTHRP) promoter, perhaps accounting for the ectopic expression of PTHRP in ATLL cells, thereby leading to altered calcium metabolism (49). However, HTLV-I mRNA expression in ATLL is so low that it has required use of RNA polymerase chain reaction (PCR) to be detected. Therefore, whether effects seen with Tax in vitro have applicability to HTLV-I in vivo remains unclear.

The ability of the viral *trans*-activator. Tax, to interface with several cellular transcriptional factor pathways suggests a new model for viral leukemogenesis. The presence of such *trans*-acting genes may allow development of new assays for the presence of as yet undiscovered retroviruses based on the ability of *trans*-acting 'Tax-like' transcriptional regulatory proteins to act on cellular and viral genes. Models for cooperation between oncogenes could undoubtedly be applied to help dissect a potential role for Tax in cooperation have emerged at this conference, such as superinfection of E_{μ} -myc transgenic mice

with Moloney murine leukemia virus (MoMuLV) (50,51).

Several conclusions can be derived from study of the *trans*-regulatory *tax* gene: (a) new mechanisms of retroviral leukemogenesis other than transduction of cellular proto-oncogenes and/or retroviral insertion adjacent to cellular proto-oncogenes may be operative in human malignancy; (b) human retroviruses possess transcriptional activators that may affect expression of cellular genes, and aberrant expression of cellular genes may contribute either to leukemogenesis *per se*, or to the leukemic phenotype; and (c) the effect of viral transregulatory genes may be felt early in leukemogenesis, and may be insufficient to elicit the full-blown leukemic phenotype.

An additional transregulatory gene studied more recently is the rex gene of HTLV-I, -II, and BLV. The rex gene is required for productive HTLV-I/-II infection. The rex gene of HTLV-I encodes two proteins, one of 27 kDa and one of 21 kDa, from an overlapping reading frame to that encoding p40^{tax} (52). These proteins appear to result from utilization of an alternative initiator methionine. In HTLV-II, two proteins are also encoded of the apparent sizes. 26 and 24-kDa (53). In HTLV-II, these appear to derive from different degrees of phosphorylation, with the larger molecular weight species being a hyperphosphorylated form of the 24-kDa protein (54). In both HTLV-I and -II, the proteins appear to act as post-transcriptional regulators, and elicit export of full-length gag/pol mRNA and probably partially spliced env transcripts from cell nucleus to cytoplasm. The rex gene appears necessary to allow expression of non-spliced and partially spliced viral mRNA, which in turn allows synthesis of Env and Gag proteins and production of mature virions.

In HTLV-I. Rex has been found to act through a cis-acting Rex-responsive element (RxRE) located in the 3' LTR. Our group has studied Rex effects mediated through the 5' LTR of HTLV-II (55). In both cases, Rex appears to act through sequences located in the R region, downstream from the transcription initiation site. Assays of binding to radiolabeled viral ' RNAs have demonstrated that purified HTLV-II Rex can directly bind to transcripts initiated from the 5' LTR, and that binding occurs to a portion of a cis-acting element responsible for Rex action, known as the RxRE (56.57). Mapping in our laboratory has demonstrated that Rex can bind directly to transcripts as short as 115 bp derived solely from sequences within the R region (57,58). These transcripts contain the 5' LTR splice donor site, and mutation of the splice donor site appears to impair Rex binding and function. Furthermore, Rex binding is dependent on retention of a specific stem-loop mRNA structure located downstream from the splice donor site (from nucleotide 465-501 within the HTLV-II 5' LTR) (57). This stem-loop structure is conserved in both HTLV-1 and -II. Rex binding may be facilitated by hyperphosphorylation, and it would appear to be the higher molecular weight (26 kDa) Rex species of HTLV-II that binds efficiently, indicating that cellular controls on Rex function may exist at the level of phosphorylation

(Chen and Green, unpublished observations). Nucleolar localization and our results using RxRE mutations of the splice donor site suggest a direct interaction of Rex with the cellular splicing apparatus to facilitate bypass of cellular splicing mechanisms.

An intriguing observation regarding Rex of HTLV-I was first made by Rimsky and Greene, demonstrating that HTLV-I Rex can functionally substitute for the Rev protein of HIV-1 (59). The Rev protein of HIV-1 performs an analogous function to that described for Rex in HTLV-I. Their assay demonstrated the capacity of Rex to induce production of the truncated singleexon form of the HIV-1 Tat protein that reflects translation from unspliced env vector mRNA (57). HTLV-II Rex in our laboratory is also able to rescue replication of Rev-deficient mutants of HIV-1 (58). Rescue of HIV-1 Rev-deficient mutants by HTLV-II Rex is relatively inefficient, and this can partially be accounted for by the relatively low affinity of HTLV-II Rex binding to the HIV-1 Rev-responsive element (RRE) (58). In addition. HIV-1 Rev is unable to complement an HTLV-II Rex-deficient clone, indicating a non-reciprocal pattern of complementation. Nevertheless, the ability of Rex to complement the genetically distant HIV-1 virus in trans suggests that. like Tax. Rex may act promiscuously on a variety of non-HTLV target sequences. This raises the possibility that Rex may also interact with cellular RNA to elicit aberrant splicing and/or transport. Disrupted processing and expression of cellular mRNAs could conceivably be implicated in the process of leukemogenesis as well. Therefore, post-transcriptional regulators may also be involved in the process of retroviral leukemogenesis. Direct evidence supporting this hypothesis has not been obtained.

Disparate Disease Entities Related to HTLV-I

Approximately five years following its discovery, it was found that the pathology elicited by HTLV-I in one setting may not predict other forms of pathology related to the virus. A specific illustration is afforded by the two major non-overlapping syndromes associated with HTLV-I. While HTLV-I may cause ATLL. another subset of infected individuals, approximately half as many, will develop a slow neurologic disease characterized by gradual development of spastic paraparesis of the lower extremities with minimal sensory loss. This illness, tropical spastic paraparesis (TSP) (also known as HTLV-I-associated myelopathy (HAM)), is distinguished from multiple sclerosis (MS) by virtue of its chronic progressive and non-episodic nature, as well as the general limitation of pathology to motor control in the lower extremities and sphincter dysfunction. The association between this illness and the virus was discovered by Gessain and co-workers while screening neurologic illnesses for retroviral involvement in Martinique (60). As opposed to the leukemia, where HTLV-I has been observed to infect and transform T-cells in vitro, no adequate model for pathogenesis of the myelopathy exists. Regardless of underlying mechanisms, involvement of HTLV-1 in a

slow neurologic disease was not predictable on the basis of its involvement in T-cell leukemia. The latency period for development of HAM is also appreciably shorter, and recently at UCLA, we saw a patient develop myelopathy approximately fifteen months following infection by transfusion (61). In contrast, development of ATLL following transfusion-acquired HTLV-I is almost never seen. Furthermore, co-existent ATLL and HAM have rarely been described. We have observed at least one case of multiple members of an Iranian Jewish family developing HAM (D. Meytes et al., unpublished), and this has been reported by other investigators. This would suggest that either differences in host genetic make-up and susceptibility or differences in viral isolates may account for familial HAM. It is important to note that the association between HAM and HTLV-I was made serendipitously. Quite possibly, if the link to HAM had been described first. no search for HTLV-I association with malignancy would have been made. This would suggest that other viruses that may not be associated in investigators' minds with development of malignancy may be candidates for potential oncogenic roles. Good candidates would be viruses with trans-acting transcriptional proteins, such as members of the herpes family, adenovirus, and/or other retroviruses.

DISCUSSION

Over a decade following initial identification of the role of the retrovirus in human disease. HTLV-I remains as the only example for human retroviral leukemogenesis. Several lessons pertinent to the search for leukemogenic viruses can be derived from the HTLV model. First, leukemia may be the rare consequence of infection with a relatively common virus. Second, a long latency period may be involved following acquisition of infection, and the leukemogenic process may involve multiple steps beyond initial infection. Third, the mechanism by which the virus may predispose to development of leukemia may be different from that seen with other retroviruses, and may involve transacting viral genes, either regulators of DNA transcription or regulators of viral mRNA expression. Finally, the role of HTLV-I as an etiologic agent in both HAM and ATLL suggests that viruses not thought to have oncogenic properties may on occasion predispose to malignancy. The overall lesson from the HTLV experience is that when suitable or suggestive epidemiologic factors are found, such as ethnic, geographic and/or familial clustering in the absence of a known inheritance pattern, a search for underlying viruses should be initiated. The search would involve removal of malignant tissue, when possible, for culture in vitro and a search using available molecular and serologic probes. The success in identifying HTLV-II by virtue of limited nucleic acid homology and antigenic similarity to HTLV-I suggests that new retroviruses can be identified. The demonstrated crossreactivity between HTLV-I and -II suggests that any search should be accompanied by rapid isolation of nucleic acid probes for viral sequences of interest, so that crossreactive

2

entitites can be discerned. A fresh look should be taken using newly available probes as a means of determining viral clonality, particularly for DNA viruses such as herpesviruses to assess whether a particular malignant tissue has arisen from a single virally infected cell. Furthermore, scrutiny of viruses already known to be widespread in the population may prove fruitful, as already appears to be the case for EBV and a subset of Hodgkin's disease. A re-duplication of such efforts will determine whether new retroviruses with oncogenic potential will be identified in man in the upcoming decade, or whether HTLV will remain an isolated if fascinating example of retroviral leukemogenesis in man.

Acknowledgements. The authors are grateful to W. Aft for assistance in preparation of the manuscript. JDR is supported by NIH grants CA01314 and CA53632, and by the US Army Medical Research and Development Command (US AMRDC); YLD is supported by the Rashi Foundation, the US AMRDC and the Doron Foundation; and ACB is supported by an NIH Physician-Scientist Award the the Leukemia Society of America.

REFERENCES

- Poiesz BJ, Ruscetti FW, Gazdar AF, Bunn PA, Minna JD, Gallo RC. Detection and isolation of type C retrovirus particles from fresh and cultured lymphocytes of a patient with cutaneous T-cell lymphoma. Proc Natl Acad Sci USA 1980;77:7415-7419.
- Yoshida M, Seiki M, Yamaguchi K, Takatsuki K. Monoclonal integration of human T-cell leukemia provirus in all primary tumors of adult T-cell leukemia suggests causative role of human T-cell leukemia virus in the disease. Proc Natl Acad Sci USA 1984;81:2534-2537.
- Kalyanaraman VS, Samgadharan MG, Robert-Guroff M, Miyoshi I, Blayney D. Golde D, Gallo RC. A new subtype of human T-cell leukemia virus (HTLV-II) associated with a T-cell variant of hairy cell leukemia. Science 1982;218:571-573.
- 4. Uchiyama T, Yodoi J, Sagawa K, Takatsuki K, Uchino H: Adult T-cell leukemia: Clinical and hematologic features of 16 cases. Blood 1977;50:481-492.
- Blattner WA. Epidemiology of HTLV-I and associated diseases. In: Blattner WA, ed. Human Retrovirology: HTLV. New York: Raven Press Ltd, 1990, pp. 251-263.
- 6. Blattner WA, Nomura A, Clark JW, Ho GYF, Nakao Y, Gallo R, Robert-Guroff M. Modes of transmission and evidence for viral latency from studies of human T-cell lymphotropic virus type I in Japanese migrant populations in Hawaii. Proc Natl Acad Sci USA 1986;83:4895-4898.
- 7. Tajima K, Ito SI, and the Tsushima ATL group. Prospective studies of HTLV-I and associated disease in Japan in human retrovirology. In: Blattner WA, ed. HTLV, New York: Raven Press, 1990.
- Saxon A, Stevens RH, Quan SG, Golde DW. Immunologic characterization of hairy cell leukemias in continuous culture. J Immunol 1978;120:777-782.
- Rosenblatt JD, Golde DW, Wachsman W, Jacobs A, Schmidt G, Quan S, Gasson JC, Chen ISY. A second HTLV-II isolate associated with atypical hairy-cell leukemia. New Engl J Med 1986;315:372-375.
- Rosenblatt JD, Giorgi JV, Golde DW, Ben Ezra J, Wu A, Winberg CD, Glaspy J, Wachsman W, Chen ISY. Integr.ted HTLV-II genome in CD8⁺ T-cells from a patient with 'atypical' hairy-cell leukemia: evidence for

distinct T- and B-cell lymphoproliferative disorders. Blood 1988,71:363-369.

29

30.

31

31

3

- Lee HH, Swanson P, Rosenblatt JD, Chen ISY, Sherwood WC, Smith DE, Tegtmeier GE, Fernando LP, Fang CT, Osame M, Kleinman SH. Antibody screening of U.S. blood donors reveals similar prevalence of HTLV-I and HTLV-II infection in association with different risk factors. Lancet 1991;337:1435-1439.
- Lee H. Swanson P. Shorty VS. Zack JA, Rosenblatt JD, Chen ISY. High rate of HTLV-II infection in seropositive IV drug abusers from New Orleans. Science 1989;244:471-475.
- Robert-Guroff M, Weiss SH, Giron JA, Jennings AM, Ginzburg HM, Margolis IB, Blattner WA, Gallo RC. Prevalence of antibodies to HTLV-I, -II, and -III in intravenous drug abusers from an AIDS endemic region. J Am Med Assoc 1986;255:3133-3137.
- 14. Brousset P, Chittal S, Schlaifer D, et al. Detection of Epstein-Barr virus messenger RNA in Reed-Sternberg cells of Hodgkin's disease by in situ hybridization with biotinylated probes on specially processed modified acetone methyl benzoate xylene (ModAMeX) sections. Blood 1991:77:1781-1786.
- Pallesen G, Hamilton-Dutoit SJ, Rowe M, Young LS. Expression of Epstein-Barr virus latent gene products in tumour cells of Hodgkin's disease. Lancet 1991: 337.320-322.
- Uccini S, Monardo F, Stoppacciaro A, et al. High frequency of Epstein-Barr virus genome detection in Hodgkin's disease of HIV-positive patients. Int J Cancer 1990;46:581-585.
- Herbst H, Niedobitek G, Kneba M, et al. High incidence of Epstein-Barr virus genomes in Hodgkin's disease. Am J Pathol 1990;137:13-18.
- 18. Kubonishi I, Equchi T, Kanzaki T, et al. EBV and Hodgkin's cells. Br J Haematol 1990;75:286-287.
- Weiss LM. Movahed LA, Warnke RA, Sklar J. Detection of Epstein-Barr viral genomes in Reed-Sternberg cells of Hodgkin's disease. New Engl J Med 1989;320:502-506.
- Staal SP, Ambinder R, Beschorner WE, et al. A survey of Epstein-Barr virus DNA in lymphoid tissue: Frequent detection in Hodgkin's disease. Am J Clin Pathol 1989;91:1-5.
- Weiss LM, Warnke RA, Sklar J. Clonal antigen receptor gene rearrangements and Epstein-Barr viral DNA in tissues of Hodgkin's disease. Hematol Oncol 1988;6:233-238.
- Sidi Y, Meytes D, Shohat B, Fenig E, Weisbort Y, Lee H, Pinkhas J, Rosenblatt J. Adult T-cell lymphoma in Israeli patients of Iranian origin. Cancer 1990; 65:590-593.
- 23. Meytes D, Schochat B, Lee H, Nadel G, Sidi Y, Cerney M, Swanson P, Shaklai M, Kilim Y, Elgat M, Chin E, Danon Y, Rosenblatt JD. A serological and molecular survey for HTLV-I infection in a newly identified high-risk group. Lancet, 1991, in press.
- 24. Linet MS. Devesa SS. Descriptive epidemiology of childhood leukaemia. Br J Cancer 1991:63:424-429.
- Alexander FE, Ricketts TJ, McKinney PA, Cartwright RA. Community lifestyle characteristics and risk of acute lymphoblastic leukaemia in children. Lancet 1990;336:1461-1465.
- Okpala IE, Abayomi NA, Gevao SM, et al. Changing patterns of acute hymphoblastic leukaemia in Nigeria. Tokai J Exp Clin Med 1989;14:301-307.
- Editorial. Childhood leukaemia: An infectious disease? Lancet 1990;336:1477-1479.
- 28. Kinlen LJ. Infective cause of childhood leukaemia.

ROSENBLATT ET AL.

22

Lancet 1989;i:378-379.

- 29. Lilleyman JS. Leukaemia 'outbreaks'. 1988; Lancet ii:1021.
- Till MM, Hardisty RM. Pike MC, et al. Childhood leukaemia in greater London: A search for evidence of clustering. Br Med J 1967;iii:755-758.
- Smith PG. Spatial and temporal clustering. In: Schottenfeld P. Fraumeni JF. eds. Cancer epidemiology and prevention, Philadelphia: Saunders, 1982, pp. 391-408.
- 32. Seiki M, Eddy R, Shows TB, Yoshida M. Nonspecific integration of the HTLV provirus genome into adult T-cell leukaemia cells. Nature 1984:309:640-642.
- 33. Seiki M, Hattori S, Hirayama Y, Yoshida M. Human adult T-cell leukemia virus: Complete nucleotide sequence of the provirus genome integrated in leukemia cell DNA. Proc Natl Acad Sci USA 1983;80:3618-3622.
- 34. Slamon DJ, Shimotohno K, Cline MJ, Golde DW, Chen ISY. Identification of the putative transforming protein of the human T-cell leukemia viruses HTLV-I and HTLV-II. Science 1984;226:61-65.
- 35. Lee TH, Coligan JE, Sodroski JG, Haseltine WA, Salahuddin SZ, Wong-Staal F, Gallo RC, Essex M. Antigens encoded by the 3'-terminal region of human T-cell leukemia virus: Evidence for a functional gene. Science 1984:226:57-61
- 36. Sodroski J, Rosen C, Goh WC, Haseltine W. A transcriptional activator protein encoded by the x-lor region of the human T-cell leukemia virus. Science 1985;228:1430-1434.
- Cann AJ, Rosenblatt JD, Wachsman W, Shah NP, Chen ISY. Identification of the gene responsible for human T-cell leukemia virus transcriptional regulation. Nature 1985:318:571-574.
- Greene WC. The human interleukin 2 receptor. Normal and abnormal expression in T cells and in leukemias induced by the human T-lymphotropic retroviruses. Ann Intern Med 1986:105:560-572.
- 39. Siekevitz M. Feinberg MB, Holbrook N, Wong-Staal F, Greene WC. Activation of interleukin 2 and interleukin 2 receptor (Tac) promoter expression by the trans-activator (tat) gene product of human T-cell leukemia virus, type I. Proc Natl Acad Sci USA 1987:84:5389-5393.
- Cross SL. Feinberg MB. Wolf JB, Holbrook NJ, Wong-Staal F, Leonard WJ. Regulation of the human interleukin-2 receptor alpha promoter: Activation of a nonfunctional promoter by the transactivator gene of HTLV-I. 1987; Cell 49:47-56.
- 41. Nimer SD, Gasson JC, Hu K, Smalberg I, Williams JL, Chen ISY, Rosenblatt JD. Activation of the GM-CSF promoter by HTLV-I and -II *tax* proteins. Oncogene 1989;4:671-676.
- 42. Nyborg JK. Dynan WS, Chen ISY, Wachsman W. Binding of host-cell factors to DNA sequences in the long terminal repeat of human T-cell leukemia virus type I: Implications for viral gene expression. Proc Natl Acad Sci USA 1988:85:1457-1461.
- Ohtani K, Nakamura M, Saito S, Noda T, Ito Y, Sugamura K, Hinuma Y. Identification of two distinct elements in the long terminal repeat of HTLV-I responsible for maximum gene expression. Eur Mol Biol Org (EMBO) J 1987:6:389-395.
- Leung K, Nabel GJ. HTLV-I trans-activator induces interleukin-2 receptor expression through an NFxB-like factor. Nature 1988;333:776-778.
- 45. Ruben S, Poteat H, Tan TH, Kawakami K, Roeder R, Haseltine W, Rosen CA. Cellular transcription factors and regulation of IL-2 receptor gene expression by HTLV-I tax gene product. Science 1988;241:89-92.

VIRAL LEUKEMOGENESIS

- 46. Grassmann R. Dengler C, Muller-Fleckenstein I, Fleckenstein B. McGuire K. Dokhelar M-C, Sodroski JG, Haseltine WA. Transformation to continuous growth of primary human T lymphocytes by human T-cell leukemia virus type I X-region genes transduced by a Herpesvirus saimiri vector. Proc Natl Acad Sci USA 1989; 86:3351-3355.
- Verenberg M, Hinrichs SH, Reynolds RK, Khoury G, Jay G. The *tat* gene of human T-lymphotropic virus type 1 induces mesenchymal tumors in transgenic mice. Science 1987;237:1324-1329.
- Nerenberg MI, Minor T. Price J. Ernst DE, Shinohara T, Schwarz H. Transgenic thymocytes are refractory to transformation by human T-cell leukemia virus type I tax gene. J Virol 1991;65:3349-3353.
- 49. Watanabe T, Yamaguchi K, Takatsuki K, Osame M, Yoshida M. Constitutive expression of parathyroid hormone-related protein gene in human T cell leukemia virus type I (HTLV-I) carriers and adult T cell leukemia patients that can be trans-activated by HTLV-I tax gene. J Exp Med 1990;172:759-765.
- 50. van Lohuizen M, Veibeek S, Scheijen B, et al. Identification of cooperating oncogenes in E_{μ} -myc transgenic mice by provirus tagging. Cell **1991**;65:737-752.
- 51. Haupt Y, Alexander WS, Barri G, et al. Novel zinc finger gene implicated as myc collaborator by retrovirally accelerated lymphomagenesis in E_{μ} -myc transgenic mice. Cell 1991;65:753-763.
- 52. Kiyokawa T. Seiki M. Iwashita S. Imagawa K, Shimizu F, Yoshida M. p27^{all1} and p21^{all1}. proteins encoded by the pX sequence of human T-cell leukemia virus type I. Proc Natl Acad Sci USA 1985;82:8359-8363.
- Shima H, Takano M, Shimotohno K, Miwa M: Identification of p26^{xb} and p24^{xb} of human T-cell leukemia virus type II. FEBS Lett 1986:209:289-294.
- Green PL, Xie Y, Chen ISY. The Rex proteins of HTLV-II differ by serine phosphorylation. J Virol 1991;65:546-550.
- Rosenblatt JD, Cann AJ, Slamon DJ, Smalberg IS, Shah NP, Fujii J, Wachsman W. Chen ISY. HTLV-II transactivation is regulated by two overlapping nonstructural genes. Science 1988;240:916–919.
- Black AC, Chen ISY, Arrigo SJ, Ruland CT, Chin E, Allogiamento T, Rosenblatt JD. Different cis-acting regions of the HTLV-II 5' LTR are involved in regulation of gene expression by Rex. Virology 1991:181:433-444.
- 57. Black AC, Ruland CT, Yip MT, Luo J, Kalsi A, Quan E, Tran B, Chen ISY, Rosenblatt JD. HTLV-II Rex binding requires a specific RNA secondary structure and intact splice donor. J Virol, in press.
- 58. Yip MT, Dynan WS, Green PL, Black AC, Arrigo SJ, Torbati A, Heaphy S, Ruland C, Rosenblatt JD, Chen ISY. HTLV Rex protein binds specifically to RNA sequences of the HTLV LTR but not the HIV-I RRE, J Virol 1991;65:2261-2272.
- Rimsky L, Hauber J, Dukovich M. Malim M, Langlois A, Cullen B, Greene W. Functional replacement of the HIV-1 rev protein by the HTLV-I rex protein. Nature 1988;335:738-740.
- Gessain A, Vernant JC, Maurs L. Barin F, Gout O, Calender A, de The G. Antibodies to human T-lymphotropic virus type-1 in patients with tropical spastic paraparesis. Lancet 1985;ii:407-409.
- 61. Saxton EH, Lee H, Swanson P, Chen ISY, Ruland C, Chin E, Aboulafia D, Delamarter R, Rosenblatt JD. Detection of human T-cell leukemia/lymphoma virus type I in a transfusion eccipient with chronic myelopathy. Neurology 1989;39:841-844.

Immunoglobulin Prophylaxis against HTLV-I in a Rabbit Model

I. Miyoshi¹, N. Takehara¹, T. Sawada¹, Y. Iwahara¹, R. Kataoka¹, D. Yang², and H. Hoshino²

¹Department of Medicine, Kochi Medical School, Kochi 783, Japan, and ²Department of Hygiene, Gunma University School of Medicine, Gunma 371, Japan

We have investigated the protective effect of human T-cell leukemia virus I (HTLV-I) Immune globulin (HTLVIG) against HTLV-I in rabbits. HTLVIG containing 77 mg/mi of IgG was prepared from pooled plasma from seropositive healthy persons. In the first experiment, four groups (A, B, C, and D) of three rabbits were transfused with 5 ml blood from an HTLV-Iinfected rabbit. Groups A, B, and C were infused 24 h later with 10. 5, and 2 mi HTLVIG, respectively, while group D was infused with 10 ml HTLVIG 48 h later. Seroconversion for HTLV-I occurred in none of group A, one of group B, and all of groups C and D after 2-5 weeks. In the second experiment, four litters (E, F, G, and H) born to another virus-infected rabbit and consisting of 7, 5, 7, and 7 newborns, respectively, were used. Litters E and H were allowed to grow normally as controls, while litters F and G were given intraperitoneal inoculation of 3 ml/kg of HTLVIG weekly four times until weaning. Although three of litters E and H each seroconverted after 5-8 weeks, none of litters F, and one of litter G became antibody-positive after 10 weeks. Presence or absence of HTLV-I infection in all these animals was confirmed by transfusion assay or gene amplification. These results indicate that passive immunization protects rabbits against blood- and milk-borne transmission of HTLV-I.

INTRODUCTION

A rabbit model of human T-cell leukemia virus I (HTLV-I) infection has been established, in which the virus was shown to be transmissible not only by blood transfusion (1,2) but also from dam to offspring via milk (3,4). In the blood transfusion experiment, as little as 0.01 ml blood from a virus-infected rabbit was capable of transmitting HTLV-I (2). Furthermore, milk or semen lymphocytes from seropositive healthy persons transmitted HTLV-I when inoculated intravenously into rabbits (5). This animal model, therefore, provided a unique opportunity to study the prc⁻ective effect of passive immunization against HTLV-I (2,6). In the present experiment, immunoglobulin prophylaxis against blood- and milk-borne transmission of HTLV-I was further explored.

MATERIALS AND METHODS

Rabbits

Japanese white rabbits, weighing about 3 kg, purchased from a commercial breeder were used.

Detection of Antibodies to HTLV-I

Blood samples were taken from rabbits at intervals of 1-2 weeks and sera were titrated for HTLV-I antibodies by indirect

Correspondence to: Isao Miyoshi, MD, Department of Medicine, Kochi Medical School, Kochi 783, Japan.

LEUKEMIA © 1992 Macmillan Press Ltd immunofluorescence against the MT-2 cell line as described previously (2). The presence or absence of immunoglobulin G (IgG) antibodies was verified by Western blot using a MT-2 lysate as antigen. Sera were also tested for IgG and immunoglobulin M (IgM) antibodies by enzyme-linked immunosobent assay (ELISA) against disrupted HTLV-I virions according to the manufacturer's instructions (Eisai, Tokyo). Neutralizing antibodies were assayed against vesicular stomatitis virus (VSV) bearing envelope antigens of HTLV-I as previously described (7).

HTLV-I Immune Globulin (HTLVIG)

HTLVIG containing 77 mg/ml of IgG was prepared from pooled plasma from seropositive healthy persons by the method of polyethylene glycol fractionation (8). The preparation had an immunofluorescence anti-HTLV-1 titer of 1:5120 and a VSV (HTLV-I) pseudotype neutralizing antibody titer of 1:6250.

Transfusion Assay

To ascertain the status of HTLV-I infection. 20 ml of blood obtained from experimental rabbits were transfused into normal rabbits. Seroconversion of the recipient rabbits indicated a virus carrier state of the donor rabbits.

Polymerase Chain Reaction (PCR)

DNA extracted from peripheral blood mononuclear cells was analyzed for the presence of HTLV-I sequences by the method of Kwok *et al.* (9). DNA, $1 \mu g$, was subjected to 40 cycles of denaturation followed by annealing and extension. Oligonucleotide primers at 7341-7360 and 7460-7411 corresponding to the pX region of HTLV-I were used. Amplification was performed using a thermostable DNA polymerase on an automated DNA Thermal Cycler (Perkin-Elmer/Cetus. Norwalk, CT). The amplified products were electrophoresed on 6% polyacrylamide gels, transferred to nylon membranes. and hybridized with a ³²P end-labeled probe at 7364-7383.

RESULTS

Passive Immunization against Blood-borne Transmission of HTLV-I

Four groups (A, B, C, and D) of three rabbits were first transfused with 5 ml of blood from an HTLV-Iinfected rabbit. Groups A, B, and C were infused 24 h later with 10, 5, and 2 ml HTLVIG, respectively, while group D was infused with 10 ml HTLVIG 48 h later.

Seroconversion for HTLV-I occurred in none of group A, one of group B, and all of groups C and D after 2-5 weeks (Figure 1). All five rabbits which were protected from seroconversion remained seronegative during an observation of six months. Sera taken immediately after infusion of HTLVIG showed anti-HTLV-I titers of 1:320 for groups A and D, 1:80 for group B, and 1:20 for group C. The VSV (HTLV-I) pseudotype neutralizing titers of these sera were 1:1250



DAVID W. GOLDE, M.D.

1533

identified populations of calcitonin gene-related periode (CGRP+ immunifective asons in human skin. Brain Rev 1987, 414: 14-48

13 R allengren J, Eleman R, Sundler J: Occurrence and distribution of neuropeptides in the human plin. An impunoversity hermical and immunochemical study on normal skin and blister fluid from willamed gkin. Acad 2000. Unserved: Studyh. 1987; 67: 185-92.

VOL 336

- 14 Dalsgaard CJ, Jernbeck J, Stains W, et al. Calcitonin gene-related peptide-like immunoreactivity in nerve fibres in the human with Relation in fibres containing substance. P., sumatostanti- and vasuactive intestinal polyappide-like immunoreactivity. Hitted-source 1080, 91: 35-88.
- Pastrowski W, Foreman JC. Some effects of calcitonin gene-related popule in human skin and on histamine release. II: J Dynamid 1986, 114: 37-46.
- Struchers AD, Brown AJJ, MacDunald DWR, et al. Human calcumin gene-related peptide: a potent endogenous vasedilator in man. *Clin. Sci.* 1986, 70: 388-93.
- 17. Huwden CW, Legue C, Gavin K, Gollie L, Ruhin PC. Haemadynamic effects of intravenous human calentonin gene-related peptide in man.

(Jm S. 1985. 74-413-18

- 18 Rusker (2): Foreman J, Reasley C, O'Shaughnessey D, Dovid PM, Galeitonin gene-related populae in the treatment of severe Raynaud's phenomenon. *Hi J Dynatid* 1989, 121: (suppl 34): 43-44.
- [19] Bunker (2), Reasky C. O'Shaughnessy D. Dowd PAL Intrasenous calculum gene-related peptide in severe Raynaud's phenomenon. B: J. Richmand (1993), 29: (suppl 2), 1.
- Shawket S. Dickerson C. Hazleman B. Brown All Selective supravensitivity to calcitonin-gene-related peptide in the hands in Raynaud's physiceneous Lancet 1989, ii: 1354-56
- Bunker CH, Foreman J, Duni d PAL Digital cutaneous vascular responses in histamine, compound 48 701 and neuroprepaides in normal subjects and Raynoud's phenomenon. J Invisi Dermatel (in press)
 Dand PAL Bunker CB, Bull HA, et al. Raynaud's phenomenon,
- 22 Dawd PAL, Bunker GB, Bull \$1A, et al. Raynaud's phenomenon, calculum gene-related peptide, endethelin, and curaneous vasculature *Lances* 1990, 336: 1014.
- 23 Zamnra MR, U'Brien RF, Rutterford RB, Weil JY, Scrum endothelan-1 ecricentrations and cold provincation in primary Raynaud's phenometrics funct 1990, 336: 1144-47.

Serological and molecular survey for HTLV-I infection in a high-risk Middle Eastern group

DINA MEYTES BATYA SCHOCHAT HELEN LEE GIORA NADEL YECHEZKEL SIDI MICHAEL CERNEY PRISCILLA SWANSON MATITTYAHU SHAKLAT YAEL KILIM MAYA ELGAT EVA CHIN YEHUDA DANON JOSEPH D. ROSENRLATT

To define the extent of human T-cell leukaemia virus (HTLV-I) infection among a group of Jewish immigrants to Israel with an increased frequency of adult T-cell leukaemia, various serological and molecular screening methods, including enzymelinked immunosorbent assay (ELISA) for anti-HTLV-I, ELISA for antibody to recombinant HTLV-I p40tax protein, and molecular detection of infection by polymerase chain reaction (PCR) amplification of HTLV-I proviral DNA from peripheral blood mononuclear cell DNA, were used. By HTLV-I ELISA the overall rate of infection was 12% (24 of 208) among immigrants from Khurusan, northeastern Iran; no HTLV-I carriers were detected among 111 unselected Jewish immigrants from other parts of Iran. There was unexplained clustering of HTLV-I infection within a cohort of 32 elderly women of similar geographic origin in a home for old people-14 were seropositive by ELISA and 19 of 29 were positive by PCB. The findings in this newly identified high-risk population suggest that in addition to ELISA, other screening techniques may be required to detect all carriers in high-risk populations.

Lancet 1990 336: 1533-35.

Introduction

Human T-cell leukacmia virus type I (HTLV-I) infection has been described in southern Japan, the Caribbean basin, and the northern parts of South America, and in certain high-risk groups, such as intravenous drug abusers in the United States.^{1,1} Previous reports of HTLV-I infection among Ethiopian Jews in Israel were not confirmed.¹¹ During the past 4 years, sporadic cases of adult T-cell leukaemia linked to HTLV-I have been reported in Israel¹⁹³¹ and 1 of the 5 latest cases were among immigrants to Israel who originated from the city of Mashad in northeastern Iran ¹¹ Because of these findings, we undertook a systematic survey of Iranian Jews in Israel, focusing on immigrants with links to Mashad.

Subjects and methods

Blood samples from Israeli blood donors of Iranian origin were obtained from the Israeli Magen David Adom Blood Services Center, Tel Aux. The criterion for classification as an Iranian control was that the country of birth of the blood donor or at least one of his or her parents was Iran. Blood samples were collected on three occasions from residents of a Mashadi home for elderly women in the Tel Aviv area and from three Mashadi community synagopues in the cities of Briel Brak and Tel Aviv. Samples were classified as Mashadi it the donor or at least one of his or her parents originated from Mashad, Iran. 20 samples from patients on long-term haemodialysis, 8 from patients with T-cell malignant diswders other than adult E-cell leukaemia, and 12 from Ethiopian Jewish immigrants were also included.

Scroligical screening was done for HTLV-1 antihodies on serum or playna samples by means of an enzyme-linked immunosystem astay (ELISA; Abbisti Laboratories). Confirmatory western blotting and or radioinmunoprecipitation assay (RIPA) with sulphur-35-labelled methionine HTLV-1-infected HUT 1028

ADDRESSES Edith Wolfson Nospital and Tel Aviv University, Sacklei School of Medicine, Holon (D. Meytes MD. M. Eight BSC); Beilinson Medicat Center, Petach Tikve (B. Schochst, PhD, Y. Sid, MD. M. Shaklei MD, Y. Kilm, MS, Y. Danon, MD); Ministry of Health, Tel Aviv, Iarsel (G. Nadel MD); Diagnostic Division, Abbott Laboratories, North Chicago, Illinois (H. Lee, PhD M. Center, BSc, P. Swamon, MSc); and Division, of Hematology Oncology, Department of Medicine, UCLX School of Medicine, Los Angeles, California, USA (E. Chin, BSc, J. D. Rosenhist, MD) Correspondence to Dr. D. Meytus, Department of Haematology, Edith Wolfson Hospis' Tel Aviv Israel 58100

PREVALENCE OF HTLV-1 INFECTION

		LISA and m blot	Anti-p40182 ELISA		
~	No tested	No (%) positive	No tested	No (%) positive	
Mashadi Jewa	208	24 (12%)	127	1219-11	
Other Innian Jewa	1 111	0	20	0	
Ethiopian Jews	12	0'	12	C	
Haemodialysis patients Patients with T-cell	20	n	20	0	
malignant disorders		0	ND		

lysate were also done.¹¹ Samples positive in the ELISA were tested by both confirmatory methods. Antibodies to HTLV-1 p40*tax* were measured by means of an ELISA with recombinant p40*tux* as antigen on the solid phase (polystyrene beads) (Abbott). HTLV-1 seropositive infected samples with known reactivity against p40*tax* on RIPA were used as positive controls, and 4 samples negative for HTLV-1 on ELISA and western blot as negative controls. Samples were scored as positive for p40*tax* if the optical density exceeded 4.5 times the mean negative control value. The polymerase chain reaction (PCR) was used to amplify HTLV-1 sequences of DNA from peripheral blood mononuclear cells with primers to a 159 bp segment contained within the *tax,rex* gene as previously described.¹⁰

Results

The extensive social and ethnic ties between Jews of Mashadi origin allowed selective sampling of this population within Israel. On repeated assays, 24 of 208 (11.5%) Mashadi serum samples were positive by ELISA for antibodies to HTLV-1 and were confirmed by western blotting. In contrast, none of the 151 control samples was positive by ELISA (table).

In an effort to survey the oldest members of the Mashadi population, we tested 32 unrelated elderly Mashadi women living in an old people's home in Tel Aviv 14 (44%) were seropositive, a rate three times higher than that in the general Mashadi population. At our first sample collection in 1988, 12 (52%) of 23 long-standing residents of the home were seropositive; 2 of 9 new residents surveyed a year later were seropositive. Of 26 Mashadi women older than 60 years who lived elsewhere in Istael only 3 (12%) were seropositive.

To determine whether scrological assays had detected all infected subjects, we carried out PCR amplification of HTLV-I DNA from 29 of the residents of the old people's home. 19 were positive for HTLV-I sequences: 14 of these were scropositive, 4 were scronegative, and 1 was scronegative by standard ELISA but reactive in the anti-p40tax ELISA. A second PCR on repeat samples from 10 of these women showed concordant results in 6 of 6 PCR-positive and 2 of 3 PCR-negative cases; 1 woman originally positive on PCR was negative on the second sample, and 1 woman originally PCR-negative was faintly positive on her second sample and had antibodies to p40tax by ELISA. This unusual clustering of HTLV-I infection within the old people's home suggested acquisition of infection within the institution.

In addition to antibodies to virion components, we assayed for antibodies to the non-structural p40rax protein. Of 128 Mashadi samples tested 12 had absorbance levels 4.5 or more times those of the negative control and were judged positive (table). 103 samples were negative by both assays. 8 samples were scropositive for both anti-HTLV-1 and anti-p40rax; 13 samples were positive for anti-HTLV-1 and negative for anti-p40rax; and 4 were positive for anti-p10rax but repeatedly negative for anti-HTLV-I by ELISA and western blot. 3 of the latter group of samples were negative on PCR, and the other, from a resident of the old people's home, was positive. All 4 were negative by western blotting and by RIPA. Therefore, the reason for the high titres in three samples in the anti-p40tar ELISA is unclear.

Direct comparison of anti-p40tax ELISA results and those of the RIPA was done for 28 subjects. 8 samples were positive for anti-p40tax by both RIPA and ELISA; 1 with traces of antihody by RIPA was negative by ELISA; the 3 samples mentioned previously were negative by RIPA but repeatedly positive by ELISA; and 16 were negative by both tests. The usefulness of the anti-p40tax ELISA in detecting true additional HTLV-I-infected seronegative infection was corroborated by other means of detection in only 1 of the 4 questionable cases.

The finding of a higher rate of seropositivity among elderly Mashadi women than in the general Mashadi population suggested a potential for gradual development of seropositivity with age, as has been reported by others. Because of the possibility that serological screening may not detect all positive subjects' we carried out HTLV-I-specific PCR on DNA from 68 randomly selected Mashadi blood samples. 5 of 7 known seropositive samples were positive in the PCR assay, and 61 seronegative samples were all negative by PCR. Hence, PCR did not increase sensitivity of detection within this sample above that achieved with serological assays alone. All PCR-positive samples contained HTLV-I and not HTLV-II, as shown by divertiminatory PCR (data not shown).

Discussion

We have identified a high risk of HTLV-1 infection in Iranian Jews originating from the city of Mashad in Khurusan, northeastern Iran. This group seems to have an overall infection rate of about 12%, though the rate was much higher among residents of a Mashadi home for old people. The reason for this clustering is unclear, although the findings suggest acquisition of infection within the institution. The overall level of infection among other Iranian Jews seems to be substantially lower than that among Mashadis. The explanation may be geographic, and Mashad may be within a previously unrecognised endemic region. Since we did not test non-Jewish Mashadis, this possibility cannot be excluded.

The unique history of the Mashadi community may help to explain their high rate of HTI.V-I infection. In 1839, the Jewish community in Mashad was subjected to a series of violent attacks and forced to convert to Islam, though the majority of the community continued to practise Judaism coverily.¹¹ To safeguard the community secret necessifated a very high rate of intermarriage among community members. Members of this community continued to marry close relatives over the next 150 years. Markers of comsanguinity are high among Mashadi Jews—for example, over 20% have a deficiency of glucose-6-phosphate dehydrogenase (D. M., unpublished). The chance introduction of HTELV-I infection into the community could have resulted in rapid transmission of the virus by sexual means and by breastfeeding.

An estimated 5000-6000 Mashadis now live in Israel. This is the first Israeli culture group identified as having a high rate of HTLV-1 infection. Members of the Mashadi Jewish community also migrated to the USA and parts of **VOL 336**

In this study, we used several methods to detect HTLV-1 infection. Our rate of anti-p40tax scropositivity in carriers. was somewhat lower than that found by others among healthy HTLV-I carriers."14 In Scrum samples from 4 subjects were positive only for anti-p40tax antibodies. Independent evidence of infection was obtained by PCR in only 1 of the 4. Thus, detection of anti-p40/ax antibodies did not appreciably add to the estimate of the rate of infection. Our findings on the use of PCR suggested that in a high-risk population, such as the old people's home we studied or in families of HTLV-I carriers, PCR would increase the number of infected individuals above that detected by scrological means. The usefulness of PCR as a screening assay in appropriate settings requires further study.

We thank Prof Ernetto Lubin for Ingistical support. The study was supported by (to J. D. R.) the Leukemia Research Foundation, the Rashi Foundation, NIH grants CA01314 and CA52410-01, and a personal contribution from Edward and Dolly Ives; (to D. M.) the Chief Scientist's Bureau, Ministry of Health, Israel, and the Israel Cancer Research Fund, and (to Y, D and J, D, R.) a grant from the US Army Medical Research and Development Command. J. D. R. was a visiting Fulbright schelar to the Sackler School of Medicine

REFERENCES

- J Hunuma Y, Komoda H, Chosa T, et al. Antibodies to adult T-cell leukemia-virus-associated antigen. A LLA3 in sera from patients with ATE and controls in Japan'a nation-wide scroepidemiol-gie study. Int 7 Curker 1992; 29: 631-35.
- 2. Himuma Y. Servepidemiology of adult T-cell leukemia virus (HTLV-I) A FLV+ origin of virus carriers in Japan AIDS Rev 1986; 2: 517-22.

- J. Riggar RJ. Methye M, Sarin PS, et al. ELISA HTLV retrovinus antihody reactivity associated with malaris and immune complexes in healthy Africans, Luna et 1985; ii: 520-23.
- 4. Blattner WA, Saxinger CA, Clark J, et al. Human T-cell leukacmia/ hmohema virus-associated temphareticular neoplasia in Tamaica. Luner 1983; ii: 61-61
- 5 Bunn PA Jr. Schechter GP, Jaffe E, et al. Clinical course of retrovin associated adult T-cell lymphome in the United States. N Engl J Aled 1983: 309: 257-64.
- 6. Rohert-Gundf M, Weiss SH, Giron JA, et al. Prevalence of antibodies to HTTLV-1, -11, and -111 in intravenous drug abusers from an AIDS endemic regim. JAAIA 1986; 255: 3133-37.
- 7. Ehrlich GD, Glaser JB, LaVigne K, et al. Prevalence of human T-cell leukemia/lymphome virus (HTTLV) type II infection among high-risk individuals: type specific identification of HTLVs by polymerase chain reaction. Blood (989; 74; 1658-64.
- 8 Ben-Ishai Z, Haas M, Triglia D, et el Human T-cell lymphotrophic virus type-I antibudies in Falashas and other ethnic groups in Israel. Nature 1985; 315: 665-66.
- 9 Karpas A, Masyon S, Raz R. Lack of antibodies to adult T-cell leuka virus and to AIDS virus in Isracli Falashas. Nature 1986; 319: 794.
- 10 Leve J, Langeuitz P, Tran H, et al. HTLV-1 associated T-cell leukomis hymphomia in Israel. In J Aled Sci 1988; 24: 397-400. 11. Sidi Y, Meytes D, Shochat B, et al. Adult T-cell lymphomia in Israeli
- patients of Iranian origin. Canar 1990; 65: 590-93.
- 12. Let H. Swanson P. Shorty VS, Zack JA, Rosenblan JD, Chen ISY. High rate of H1LV-II infection in scropositive IV drug absuers from New Orienny. Science 1989; 244: 471-75.
- 13 Encyclopaedia Judanca 1972; 11: 1399-400.
- 11 Ehrlich GD, Glaser JB, Abbon MA, et al. Detection of anti-HTLV-I Tax antibodies in HTLV-1 enzyme linked immunosorbenit assaynegative individuals Blood 1989; 74: 1064-72
- 15. Yokata'E, Cho MJ, Fachibana N, et al. The prevalence of antibody to p42 of IEEEV-I among ATEL patients in companisons with healthy carriers in Japan. Int J Cuncer 1989; 43: 970-71
- 16 Kamihara S, Toriya K, Amagasaki T, et al. Antibodiet against p40tax gene product of human T-lymphotrophic virus type I under various conditions of HTLV-1 infection Ton 7 Concer Res 1989; 80: 1066-71.

Quinine-induced disseminated intravascular coagulation

RUTH L. SPEARING CHRISTINE M. HICKTON PETER SIZELAND ANTHONY HANNAH ROSS R. BAILEY

Recurrent disseminated intravascular coagulation occurred in 3 women after ingestion of guinine tablets for cramp. All had circulating quininedependent antibodies to platelets and in 2 there was initial evidence of antibody consumption, with low titres that rose steeply over the next few days and remained high for many months.

Lancet 1990; 336: 1535-37.

Introduction

Recognised haematological problems associated with ingestion of quinine include thrombocytopenia, erythrocyte haemolysis, and neutropenia. Quinine was first implicated as a cause of purpura in the late 19th century, 1 and there have been several reports of associated thrombocytopenia.23 However, we are aware of only two published cases of disseminated intravascular coagulation induced by quinine,** and report three further cases.

Patients and methods

Case histories

A 74-year-old woman was admitted 5 times over 3 years with various symptoms, which included acute shortnes: of breath,

wheeze, generalised abdominal pain, fever, lower back and chest pain, melaena, haematemesis or haemophysis, and bruising and petechiae. Most episodes occurred shortly after going to bed. Investigations on each occasion (table) showed evidence of disseminated intravascular coogulation (DIC). On the first 2 admissions she was treated with antibiotics, although blood cultures were always negative. On the third admission she was treated for asthma, and on the last 2 occasions no specific treatment was given. On each occasion, fever and other symptoms resolved within 24 h with a subsequent resolution of congulation abnormalities. At least 2 similar, but milder, episodes also occurred for which she did not attend hospital. On 3 occasions recent quinine ingestion was clearly remembered by the patient or documented in the admission notes. A check on the number of tablets left in the bottle after her initial prescription indicated that 10 tablets had been taken over 7 years. Retrospective quinine-dependent platelet antibody analysis on samples stored from the last 3 admissions and from intervening periods showed low or undetectable antibody concentrations. during the first 2 days of each acute episode, which then rose sharply

ADDRESSES Department of Haematology (R.L. Spearing, FRACP. C.M. Hickton, FN2/MLT) and Department of Nephrology (P. Sizeland, MRCP, A. Hannah, MB, R. 8. Bailey, FRCP), Christochurch Hospital, Private Bag, New Zealand, Correspondence to Dr.R. L. Spearing, Department of Haematology, Christchurch Hospital, Private Bag, Christchurch, New Zealand

MOLECULAR CHARACTERIZATION OF HTLV-I INFECTION IN ISRAEL

Yehuda L. Danon^{**}, Yael Kilim^{*} and Joseph D. Rosenblatt[@] *Kipper Institute of Immunology, #Basil and Gerald Felsenstein Research Center, Children's Medical Center of Israel, Sackler School of Medicine, Tel-Aviv University Israel and @Division of Hematology Oncology, Department of Medicine, UCLA School of Medicine Los Angeles Calif 90024, USA.

Address for Correspondence: Y.L. Danon, M.D.

Director,

The Children's Medical Center of Israel P.O. Box 559 Petah-Tikva 49202 Israel

This work was supported in part by the US Army Medical Research and Development Command

Short Title: Characterization of Israeli HTLV-I isolate

_

 \mathbb{T}^{1}

INTRODUCTION

Many studies were performed on human T-cell leukemia virus type I, ethiologically associated with adult T-cell leukemia (ATL) 1,2 and HTLV-associated myelopathy (HAM). $^{3-7}$

HTLV-I genomes isolated from ATL and HAM patients from different geographical origin demonstrated a high degree of homology (>96%).⁵⁻⁸ Sequence variation in different isolates were found mostly in the LTR (1.3-5.2%) and the region between the envelope and tax/rex reading frames (0.1-6.9%).⁶ It was found that genetic diversity between different isolates is in association with the geographical origin and not with the clinical presentation.^{6,7,8} In 1988, HTLV-I was first discovered in Israel and the Middle East.⁹ Two years later a community of jewish immigrants from the city of Mashad in northeastern Iran was identified with an infection rate of about 12%.¹⁰

Our aim was to determine the nucleotide sequence of LTR and env gene from HTLV-I genome of an HAM patient who originated from Mashad. This data was compared to sequences derived from HTLV-I isolated from Japanese and African patients in order to locate the origin of the Mashadi virus.

MATERIALS AND METHODS

<u>Ceil lines:</u> Lymphocytes were collected from 20ml peripheral blood by a Ficoll-Hypaque density gradient and were resuspended in RPMI-1640

- 3 -

medium supplemented with 15% fetal bovine serum, interleukin-2 (IL-2) at a concentration of 50u/ml, 0.5% PHA and 1% penicillin/streptomycin. Cells were incubated at 37° C in the presence of 5% CO2 and were maintained for several weeks. The cells were cryopreserved at various intervals for DNA extraction.

DNA extraction and PCR amplification: DNA was extracted from lymphocytes with phenol/chloroform. Two regions in the HTLV-I genome were amplified: LTR and env gene. LTR amplification was performed with the primers R11/14 which defined a 741 base sequence from nucleotide 61 to 802 (nucleotides are numbered according to the sequence of Seiki et al.¹¹ The env gene was amplified by two pairs of primers : R15/17A which defined a 1163 base sequence from nucleotide 5201 to 6364 and R19/18 primers which defined 791 base sequence from nucleotide 5942 to 6733.

The patient's (HE) DNA was PCR amplified for 30 cycles in a reaction containing 100 μ l of 2mM MgCl, 200 ng from each primer, 0.2 mM from each dNTPs and 2.5 u of taq polymerase (USB, Cleveland, OH). A list of primers used for PCR amplification is presented in table 1.

DNA sequencing: PCR products were recovered from PCR reaction with DS primer Remover (ALvanced Genetic Technologies Corp., Gaithersburg, MD) and ethanol precipitation. Nucleotide sequence analysis was performed by the Taq Dye Deoxy Terminator Cycle Sequencing kit using the 373A DNA Sequencer (Applied Biosystems) at the Biological Services of the Weizmann Institute Rehovot Israel. Sequence analysis was performed at least twice for each primer (Table 1).

- 4 --

Sequence comparison: HE sequence was compared to the Japanese (ATK), African (EL) and Papua New Guinea (PNG-1) sequences by gcg program with the accession numbers: JO2029 (ATK), S74562 (EL LTR), M85207 (PNG-1) and M69044 (EL).

RESULTS

Comparison of HTLV-I (HE) LTR to other isolates. PCR amplification of HTLV-I LTR produced a PCR product of 741bp of which 698 bases from it were sequenced. 7 nucleotides which are not verified yet are designated as N. HTLV-I-(HE) LTR sequence was compared to three other isolates. The Japanese (ATK) sequence showed nucleotide homology of 97.8% with 15 nucleotide differences.¹¹ The African sequence showed nucleotide homology of 95% with 35 nucleotide differences and the Papua New Guinea isolate, which was compared only by 629 nucleotides, showed 91% homology with 56 nucleotides differences.^{5,6}

Comparison of the sequence changes in the LTR region to the Japanese sequence showed that the differences are clustered mostly in the U3 (66.7%), 6.7% were in the R and 26.7% were in the U5 regions. There are no changes at the three 20bp enhancer elements at the U3 region and there is 98.7% homology for the Rex Responsive element located between bases 313 and 627.

<u>Comparison of the env gene amplified from HTLV-I (HE) genome to the</u> <u>Japanese and African sequences</u>. The PCR product was 1532bp in length from which 1496 bases were sequenced. The sequence between nucleotides 5625-5685 is not verified yet. Comparison to the Japanese (ATK) sequence showed nucleotide homology of 97.7% with 9 nucleotide differences while the African (EL) sequence showed nucleotide homology of 98.4% with 6 nucleotide differences.^{12,14} There is a problem with base T at position 5400. It seems that it creates a stop codon while in the other two sequences it does not exist. This finding needs further examination. HE env2 sequence was compared to the Japanese (ATK) sequence.¹¹ 98.7% homology at the nucleotide level with 12 nucleotide differences were found. Comparison to the African (EL) sequence showed nucleotide homology of 99.5% with 5 nucleotide differences.¹²

DISCUSSION

A new focus of HTLV-I infection was recently identified in the Middle East.¹⁰ In this study we determined the sequence of two regions in HTLV-I isolated in Israel from an HAM patient belonging to this community: the LTR region and the env gene. Several studies have indicated a high degree of homology among HTLV-I isolates (>96%) and demonstrated that differences between variants are in association with their geographical origin.^{5,6,7,8} Comparison of Mideastern sequence which originates in Iran to the Japanese, African and Papua New Guinea sequences in order to examine whether there is indeed a higher degree of homology between isolates from the same geographical area. A high level of homology, at about 98%, to the Japanese isolate, while the African and Papua New Guinea isolates were more divergent with sequence homology of 95% and 91%, respectively.

Δ,

- 6 -

Comparison of HTLV-I env gene to the Japanese and African isolates showed homology at the nucleotide level of about 98-99%. From these results we can conclude that the env gene is more conserved than the LTR region.

With the exeptions of some nucleotides which are not verified yet, comparison of the LTR region showed a higher degree of homology between Iranian sequence and the Japanese isolate. As for the env gene, there is a high degree of homology between the Iranian sequence and the African isolate. With these results we cannot conclude if there is an influence of the geographical area on the virus genome.

There are two theories about HTLV-I origin and the way it was spread to the world. Gallo suggests that it originated in Africa and spread to the new world and Japan, while Saskena et al. suggest that HTLV-I originated in the Indo-Malay region.^{5,13,14} Based on the comparison of the LTR sequences, our results support the theory of HTLV-I originating in Africa, since we found higher degree of homology between HE sequence and the African isolate, than with the sequence of Papua New Guinea isolate.

τ,

REFERENCES

- POIESZ, B.J., F.W. RUSCETTI, A.F. GAZDER, P.A.BUNN, J.D.MINNA & R.C. GALLO. 1980. Detection and isolation of type C retrovirus particles from fresh and cultured lymphocytes of a patient with cutaneous T-cell lymphoma. Proc.Natl.Acad.Sci. USA 77: 7415-7419.
- UCHIYAMA, T., J. YODOI, K. SAGAWA, K. TAKATSUKI & H. UCHINO. 1977. Adult T-cell leukemia: clinical and hematologic features of 16 cases. Blood 50: 481-492.
- TSUJIMOTO, A., T. TERUUCHI, J. IMAMURA, K. SHIMOTOHNO, I. MIYOSHI & M. MIWA. 1988. Nucleotide sequence analysis of a provirus derived from HTLV-I-associated myelopathy (HAM). Mol. Biol. Med. 5: 29-42.
- GESSIAN, A., F. BARIN, J.C. VERNANT, O. GOUT, L. MAURS, A. CALENDER & G. de THE. 1985. Antibodies to human T-lymphotropic virus type I in patients with tropical spastic paraparesis. Lancet 11: 407-409.
- 5. SAKSENA, N.K., M.P. SHERMAN, R. YANAGIHARA, D.K. DUBE & B.J. POIESZ. 1992. LTR sequence and phylogenetic analysis of a newly discovered variant of HTLV-I isolated from the Hagahai of Papua New Guinea. Virol. 189: 1-9.
- RATNER, L., T. PHILPOTT & D.B. TROWBRIDGE. 1991. Nucleotide sequence analysis of isolates of human T-lymphotropic virus type I of diverse geographical origins. AIDS Res. and Human Retroviruses. 7: 923-941.

~

- KINOSHITA, T., T. TSUJIMOTO & K. SHIMOTOHNO. 1991. Sequence variations in LTR and env regions of HTLV-I do not discriminate between the virus from patients with HTLV-I associated myelopathy and adult T-cell leukemia. Int. J. Cancer 4: 491-495.
- DAENKE, S., S. NIGHTHIGALE, J.K. CRUICKSHANK & C.R.M. BANGHAM.
 1990. Sequence variants of human T-cell lymphotropic virus type I from patients with tropical spastic paraparesis and adult T-cell leukemia do not distinguish neurological from leukemic isolates. J. Virol. 64: 1278-1282.
- LEOR J., P. LANGEUITZ, H. TRAU, E.O.M. SCHNIDER, D. DOUER & I. BEN-BASSAT. 1988. HTLV-I associated T-cell leukemia/lymphoma in Israel. Isr. J. Med. Sci. 24: 397-400.
- MEYTES, D., B. SHOCHAT, H. LEE, G. NADEL, Y. SIDI, N. CERENY,
 P. SWANSON, M. SHAKLAI, Y. KILIM, M. ELGAT, E. CHIN, Y. DANON
 & J.D. ROSENBLATT. 1990. Serological and molecular survey for
 HTLV-I infection in a high-risk Middle Eastern group. Lancet
 336: 1533-1535.
- SEIKI, M., S. HATTORI, Y. HIRAYAMA & M. YOSHIDA. 1983. Human adult T-cell leukemia virus: complete nucleotide sequence of the provirus genome integrated in leukemia cell DNA. Proc. Natl. Acad. Sci. USA 80: 3618-3622.
- PAINE, E., J. GARCIA, T.C. PHILPOTT, G. SHAW & L. RATNER. 1991. Limited sequence variation in human T-lymphotropic virus type
 I isolates from North American and African patients. Virol. 182: 111-123.

ت،

- 13. GALLO, R.C. 1991. Human retroviruses: a decade of discovery and link with human disease. J. Infect. Dis. 164: 235-243.
- ROSENBLATT, J.D., Y. DANON & A.C. BLACK. 1992. A decade with HTLV-I/HTLV-II: Lessons in viral leukemogenesis. Leukemia 6 (1): 18-23.

ۍ:

,

Primer	From	nt. to nt.	Used for	Seq.	Remarks
R11	61	77	amp. & Seq.	5' TAGAGCCTCCCAGTGAA	
R12	494	470	seq.	5' CCTAGACGGCGGACGCAG	Comp.
R14	802	786	amp. & seq	5' CTCGTATCCCGGACGAG	Comp.
R15	5201	5218	amp. & seq.	5' CATGGGTAAGTTTCTCGC	
R16	5660	5645	seq.	5' ATGGAGATTAATATTG	Comp.
R17	5641	5658	seq.	5' GCCTCAATATTAATCTCC	Comp.
R19	5942	595 9	amp. & seq.	5' TCCATCCTCTTCTTCTAC	
R17A	6364	6347	amp.	5' TCCCAGAACAGGAGATCA	Comp.
R18	6733	6716 _.	amp. & seq.	5' GGGAGAGGTAATTATTG	

4,

Table 1: LIST OF PRIMERS FOR PCR AMPLIFICATION AND SEQUENCING