ASSESSMENT OF CELL MEDIATED IMMUNITY AGAINST SALMONELLA TYPHI ANTIGEN IN GUINEA PIGS FOLLOWING INTRAPERITONEAL INJECTION OF SENSITIZED SPLEEN CELLS

Khalil H. AL-Joboury Department of Pathology, College of Veterinary Medicine, University of Baghdad Baghdad,Iraq

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ABSTRACT

In an assessment of cell mediated immunity against <u>Salmonella typhi</u> antigen in Guinea pigs following their intraperitoneal injection with sensitized spleen cells. The results were revealed a high level of cell mediated immunity against <u>Salmonella typhi</u> antigen in the Guinea pigs intraperitonealy received the sensitized spleen cells $(5x10^8/ml)$. The level of cell mediated immunity was detected using delayed type hypersensitivity test, Macrophages migration inhibition test and Erythrocytes rosette test.

The normal (nonsensitized) spleen cell recipient group (control group) was showed no level of cell mediated immunity using these tests.

INTRODUCTION

Adaptive transfer of cell mediated immunity to specific antigen in human was first demonstrated by Lawrence (1), who showed that transfer of whole viable lymphocytes from a normal tuberculin skin positive donor to a normal skin test negative recipient resulted in a conversion of the recipient to skin test positive.

Lawrence (2) demonstrated that delayed cutaneous hypersensitivity responsiveness to Streptococcal M substances could also be transferred by Soluble extracts of leukocytes and termed the factor responsible for this phenomenon transfer factor, the important immunological effects are the antigen specific conversion of delayed type hypersesitivity and production of lymphokines (3,4).

The successful transfer of delayed skin sensitivity, also was done in mice using spleen cells sensitized against other antigenic materials, such as a particulate antigen (Candida) and soluble protein antigens (ferritin, cytochrome C, horse radish peroxidase and purified protein derivatives) (5).

Cell mediated immunity could be transferred from sensitized mice to nonsensitized, using immune spleen cells against <u>Rhodococcus equi</u> (6). Also, by using immune peritoneal macrophages against <u>Salmonella typhimurium</u> and <u>Salmonella enteritidis</u> (7).

Because <u>Salmonella typhi</u> is facultative intracellular pathogens, cellular immunity specifically immne T lymphocytes and macrophages were proposed as the most important defense mechanism, for this reason, this study was aimed to evaluate the role of sensitized spleen cells of the Guinea pigs in transferring cell mediated immunity against whole killed <u>Salmonella typhi</u> antigen (local typhoid vaccine), using delayed type hypersensitivity-skin test, Erythrocyte-rosette test (E-rosette test) and macrophages migration inhibition test.

MATERIALS AND METHODS

Guinea pigs:

Eighteen adult Guinea pigs, weighing 400-500gm each, supplied by AL-Kindi Company of Veterinary drugs and Vaccine production were reared together on

concentrated food. Blood samples were taken from all the animals in order to test level of antibiodies against Salmonella typhi.

The animals were divided into 4 groups.

First group; immunization group (5 animals):

The immunization procedures were done by subcutaneous injection of the animals with 0.25ml of killed whole cell vaccine-phenol preserved containing 109 bacterial cell/ml. Booster doses of 0.25 and 0.5ml of vaccine were given subcutaneously on the 3rd and 5th week postinoculation respectively.

Second group; control group (5 animals):

Similarly, was injected with sterile phosphate buffer saline. At the 14th and 21st days after the second booster dose a delayed type hypersensitivity-skin test and macrophages migration inhibition test were done on immunization group and control group according to Dham and Thompson and Weir methods (8,9), to check the level of cell mediated immunity and immunity transfer by for preparation of sensitized and non-sensitized spleen cells.

Sensitized spleen cells were taken from the immunized group of animals which showed high level of cell mediated immunity against Salmonella typhi sonicate soluble antigen (DTH-skin test was > 5mm and migration index (MI) < 0.80).

Similarly, normal (non-sensitized) spleen cells were taken from the control group (2nd

The spleen cells from each animals in both groups were surgically taken-out in RPMI-1640, containing 10% fetal calf serum, and after trimming off all the adherent tissue, cut into small pieces, minced and teased on a sterile stainless sieve to obtain single cell suspension.

The leukocytes from the spleen cell suspension were made free from erythrocytes by treatment with 0.83% ammonium chloride for 10 minutes, after that the spleen cells suspension was washed 3 times by phosphate buffer saline and checked for viability using 0.2% trypan blue dye.

The number of spleen cells per ml of phosphate buffer saline was adjusted to 5X108 cell/ml and used as viable cells for immunity transfer by intraperitoneal injection of the third group of animals (sensitized cell recipient group, containing 4 animals). Similarly, normal splcen cell suspension was prepared and intraperitoneally injected into the fourth group of animals (normal spleen cell recipient group, containing 4 animals).

After the injection of sensitized and normal spleen cells, an assessment of the cell mediated immunity was done in these two groups, using DTH-skin test which was done after 24hrs and MIF Test was done during 5 to 14th days postinoculation of spleen cells. Also, Erythrocytes-rosette test was done during this period according to Braganza et al. Technique (10)

RESULTS

This study was revealed the following findings:

Sensitized spleen cell recipient group; This group of animals was showed high level of cell mediated immunity in compare to the normal cell recipient group (control group), using the following tests:

DTH-skin test:

Marked skin reactive areas were seen in sensitized spleen cell recipient group, the mean diameters of reactive areas were (>5mm), 8.75 ± 1.708 mm, 6.5 ± 1.291 mm and 5.0 ± 816 mm after 24hrs of inoculation of concentrated Sonicate antigen (400, 40, 4 μ g/ml) respectively, and then gradually decreased thereafter (table -1). The normal spleen cell recipient group (negative control) showed skin reactive areas, with mean diameters < 5mm (3.25 ± 1.258 , 2.25 ± 1.258 and 1.75 ± 0.957) after 24hrs of using concentrated Sonicate antigens (400, 40, 4 μ g/ml) respectively.

2. Macrophage migration inhibition test:

Macrophage migration inhibition test (MIF) activity was determined in all animals, intraperitoneally received the sensitized spleen cells. The mean indices of macrophages migration were <0.80 (0.252 \pm 0.260, 0.427 \pm 0.212 and 0.669 \pm 0.348) at the different concentrations of Sonicate antigen (400, 40, 4 μ g/ml) respectively (Table-2), whereas, the mean indices of macrophages migration were >0.80 (0.847 \pm 0.285, 1.530 \pm 0.736 and 2.249 \pm 1.732) in the normal spleen cell recipient group against different concentrations of Sonicate antigen (400, 40, 4 μ g/ml) respectively.

3. Erythrocytes-rosette Test:

E-rosette formation active of and total T-lymphocytes were increased in all animals received sensitized spleen cells against Salmonella typhi antigen. The mean numbers of E-rosette formation active and total T-lymphocytes were 0.625 ± 0.250 and 0.750 ± 0.288 respectively before treatment.It increased to a mean of 12.250 ± 2.872 and 13.500 ± 3.109 for active and total T-lymphocytes respectively in sensitized spleen cell recipient whereas, the mean numbers of E-rosette formation active and total T-lymphocytes were 0.625 ± 0.250 and 0.750 ± 0.288 respectively before treatment. It increased to 1.250 ± 0.500 and 1.500 ± 0.577 respectively in the normal spleen cell recipient group (Table-3).

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Table-1: DTH-skin test in different recipient groups of G. pigs following I/D injection of $\frac{1}{2}$ injection of

		PBS		0	0		0	0	0		0
	×	lm	4	3-4	3.25	+1	0.5	0-1	0.25	+1	0.50
	72 hrs	Ag concent 4g/ml	40	3-6	4.5	+	1.291	0-2	0.5	+1	1.00
		Agcor	400	4-7	5.5	+1	1.291	0-2	1.0	+1	0.816
		PBS		0	0		0	0	0		0
	rs.	lm/s	4	3-5	4.0	+1	0.816	0-2	0.75	+	0.957
	48 hrs.	Ag concent µg/ml	40	4-7	5.5	+1	1.291	0-3	1.25	+1	1.258
hours		Ag co	400	6-9	7.0	+1	1.414	1-3	2.0	+1	0816
Time/hours		PBS		0	0		0	0	0		0
	i.	lm/s	4	4-6	5.0	+1	0.816	1-3	1.75	+	0.957
	24 hrs.	Ag concent µg/ml	40	2-8	6.5	+1	1.291	1-4	2.25	+1	1.258
		Ag co	400	7-11	8.75	+1	1.708	2-5	3.25	+	1.258
	Diameters	mm		Range	Mean	+	S.D		Range	Mean	S.D
	(Group		Sensitized	spleen cell	recip.		Normal	spleen cell	recip.	

PBS: Phosphate Buffer Saline

ation inhibition indices (MI) following various treatments in G. pigs in response to S. typhi Ag.

Animal		Ag concent µg/ml		PHA (10 μg/ml)
.00.	400	40	4	
1	0.640	0.640	1.000	0
2	0.140	0.562	0.765	0
3	0.081	0326	0.734	0
4	0.148	0.179	0.179	0.213
D	0.252±0.260	0.427±0.212	0.669±0.348	0.053±0.106
-	0.694	1.000	1.361	0
2	1.265	1.562	1.562	0
3	0.790	1.000	1.234	0.197
4	0640	2.560	4.840	0
O	0.874±0.285	1.530±0.736	2.249±1.732	0.049±0.098

ohemagglutinine

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Table-3: Mean percentage values of active and total T-lymphocytes in the recipient groups (E-rosette Test).

Pretreat	Pretreatment neriod		Post	Post treatment	
		Sensitized spl	Sensitized spleen cell recip.	Normal	Normal spleen cell recip.
AT	TT^2	AT	TT	AT	TT
0.625±0.250	0.750±0.288	12.250±2.872	13.500±3.109	1.250±0.500	1.500 ± 0.577

'AT= Active T lymphocytes. %
'TT= Total Tlymphocytes. %

DISCUSSION

The major host defense mechanism for Salmonella infection is controversial, but it is clear that both humoral and cell mediated immunity responses are important (11). Because Salmonella are facultative intracellular pathogens, cellular immunity specifically immune macrophages and T lymphocytes is proposed as the most important defense mechanism (12).

A similar findings were correlated with the present study in which sensitized spleen cells had ability to transfer cell mediated immunity against the Salmonella typhi antigen, detected by DTH-skin test and MIF test. Histologically, spleen tissue was composed of T cells (85%), responsible for lymphokines production, B cells (15%) responsible for antibody production. Other cell type was also present in spleen tissue such as dendritic cells (macrophages) responsible for the antigen processing and some lymphokines production (13) and therefor following the injection of the animals with antigen (Salmonella typhi antigen) in the present study, the spleen cell became sensitized and had the ability to transfer the cell mediated immunity to the recipient group which gave DTHskin reactive areas >5mm diameter and migration indices (MI)<0.80. similarly, the sensitized peritoneal macrophages and spleen cells gave protection against Salmonella typhimurium and Salmonella enteritidis (7) and against other type of microorganisms such as Rhodococcus equi (6). Also, similar findings were detected following sensitization of spleen cells with particulate antigen (Candida) and Soluble protein antigens (Ferritin, cytochrom C, horse radish peroxidase and purified protein derivatives) (5) and also with tuberculin (14) using sensitized lymphocytes of tuberculin hypersensitive Guinea pigs.

تقييم للمناعة الخلوية ضد مستضد عصيات التايفوئيد في خنازير غينيا بعد حقنها بخلايا الطحال المحسسة داخل تجويف البطن.

خليل حسن الجبوري فرع الأمراض، كلية الطب البيطري،جامعة بغداد،بغداد،العراق

لخلاصة

في تجربة صممت لتقييم مستوى المناعة الخلوية صد مستضد عصيات التايفونيد في خناز ير غينيا بعد حقنها بخلايا طحال محسسة داخل تجويف البطن و بجرعة ٥ ٪١٠ أخلية/مللتر، حيث دلت النتائج إلى وجود مستوى عالى للمناعة الخلوية ضد مستضد عصيات التايفونيد في خناز ير غينيا بعد حقنها بخلايا الطحال المحسسة. وقد قيست المناعة الخلوية باستعمال الاختبارات التالية:

و فحص تفاعل الجلد الناجم عن فرط الحساسية المتأخر، فحص منع هجرة البلاعم الكبيرة و فحص تكوين الوردة بو اسطة الخلايا التائية الفعالة و الكلية, أما مجموعة السيطرة و المستلمة خلايا طحال طبيعية فلا يوجد عليها مستوى واضح للمناعة الخلوية ضد مستضد التائفونيد

REFERENCES

- Lawrence, H. S. (1949). The cellular transfer of cutaneous hypersensitivity to tuberculin in man. Proc. Soc. EXP. Biol. Med. 71: 516-522.
- Lawrence, H. S. (1955). The transfer in humans of delayed skin sensitivity to Streptococcal M substance and to tuberculin with disrupted leukocytes J. Clin. Invest. 34: 219-230.
- Kirkpatrik, C. H.; Rozzo, S. J. and Mascali, J. J (1985). Murine transfer factor. III-specific interactions between transfer factor and antigen. J. Immunol. 135 (6): 4027-4033.
- Wilson, G. B. and Fudenberg, H. H. (1981). Leukocytes migration inhibition as a method for assaying transfer factor activities. Lymphokines 4: 107-172
- Peterson, E. A.; Greenberg, L. E.; Manzara, T. and Kirkpatrick, C. H. (1981). Murine transfer factor 1-Description of the model and evidence for specificity. J. immunol. 126 (6): 2480-2484.
- Madarame, H.; Takai, S.; Matsumoto, C.; Minamiyama, K.; Sasaki, Y.; Tsubaki, S.; Hasegawa, Y. and Nakane, A. (1997). Virulent and avirulent Rhodococcus equi infection in T cell deficient athymic nude mice: pathologic, bacteriologic and immunologic responses. FEMS Immunol and Med. Microbiol. 17: 251-262.
- Rowley, D.; Turner, K. J. and Jenkin, C. R. (1964). The basis for immunity to mouse typhoid. 3-cell-bound antibody Aust. J. EXP. Biol. Med. Sci 42: 237-248.
- 3. Dham, S. K. and Thompson, R. A. (1982). Humoral and Cell mediated immune responses in chronic typhoid carriers. Clin. EXP. Immunol. 50: 34-40.
- Weir, D. M. (1978). Handbook of experimental immunology, 3rd. ed. Blakwell Scientific Publications, Oxford. Vol. II-cellular Immunity, pp: 27-21.
- Braganza, C. M.; Stathopoulus, G.; Davis, A. J. S.; Elliot, E. V. and Kerbel, R. S. (1975). Lymphocytes: Erythrocytes (L. E) Rosette as indicators of the heterogeneity of lymphocytes in a variety of mammalian species. Cell, 4: 103-106.
- Eisenstein, T. K. and Sultzer, B. M. (1983) Immunity to Salmonella infection. Adv. Exp. Med. Biol. 162: 261-296.
- Spier, S. J. (1993). Salmonellosis. Veterinary Clinics of North America: Equine practice. 9 (2): 385-397.
- Chapel, H. and Haeney, M.(1995). Essentials of clinical immunology, 3rd. ed. Blackwell Science.
- Smogorzewska, F. (1990). Transfer of delayed hypersensitivity in Guinea pigs. Probl-Med-Wieku-Rozwoi, 16: 167-188.