AD_____

Award Number: DAMD17-99-1-9406

TITLE: hRAD51 Involvement in Genomic Instability and Development of Breast Cancer

PRINCIPAL INVESTIGATOR: Richard A. Fishel, Ph.D. Christoph Schmutte, Ph.D.

CONTRACTING ORGANIZATION: Thomas Jefferson University Philadelphia, Pennsylvania 19107

REPORT DATE: September 2002

TYPE OF REPORT: Annual

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

DISTRIBUTION STATEMENT: Approved for Public Release; Distribution Unlimited

The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision unless so designated by other documentation.

20030602012

REPORT	Form Approved OMB No. 074-0188					
Public reporting burden for this collection of infor	mation is estimated to average 1 hour part reserves					
reducing this burden to Washington Headquarte Management and Budget, Paperwork Reduction	rs Services Directorate for Information Operation	parding this burden estimate or any oll s and Reports, 1215 Jefferson Davis I	er aspect of this collection of informatight of the source	ation, including suggestion 22202-4302, and to the Office		
1. AGENCY USE ONLY (Leave blar		3. REPORT TYPE AND				
4. TITLE AND SUBTITLE	September 2002	Annual (23 Aug	01 - 22 Aug 02)			
	in Conomin Turk		5. FUNDING NUMBERS			
Development of Deve	in Genomic Instab:	ility and	DAMD17-99-1-9	406		
Development of Bre	ast Cancer					
6. AUTHOR(S): Richard A. Fishel,						
Christoph Schmutte						
enriscoph sennacce	, PIL.D.					
7. PERFORMING ORGANIZATION	NAME(S) AND ADDRESS(ES)					
		8. PERFORMING ORGANIZATION REPORT NUMBER				
Thomas Jefferson U						
Philadelphia, Penn	sylvania 19107					
E-MAIL:						
RFishel@lac.jci.tju.edu; cschmu	utte@lac.jci.tju.edu					
9. SPONSORING / MONITORING	AGENCY NAME(S) AND ADDRESS	ES)	10. SPONSORING / MON	ITORING		
U.S. Army Medical Research an	d Material Command		AGENCY REPORT NU			
Fort Detrick, Maryland 21702-5	5012					
••••						
		-				
44 01100 01100	anti	20	030602	010		
11. SUPPLEMENTARY NOTES		21		017		
				· · · ·		
		<u> </u>				
12a. DISTRIBUTION / AVAILABILIT Approved for Public Re	TY STATEMENT	<u></u>		- · · -		
12a. DISTRIBUTION / AVAILABILI Approved for Public Re	TY STATEMENT elease; Distribution Ur	<u></u>				
12a. DISTRIBUTION / AVAILABILI Approved for Public Re	TY STATEMENT elease; Distribution Ur	<u></u>		- · · •		
Approved for Public Re	elease; Distribution Ur	<u></u>				
 12a. DISTRIBUTION / AVAILABILIT Approved for Public Re 13. ABSTRACT (Maximum 200 WARD) 	elease; Distribution Ur	<u></u>				
Approved for Public Re	elease; Distribution Ur	<u></u>				
Approved for Public Re 13. ABSTRACT (Maximum 200 W	elease; Distribution Ur	alimited	12b. DIS	STRIBUTION CODE		
Approved for Public Re 13. ABSTRACT (Maximum 200 W) During the third year interactions between h	of the funded period,	we focused on the	2 characterizatio	STRIBUTION CODE		
Approved for Public Re 13. ABSTRACT (Maximum 200 W During the third year interactions between h suggesting a stable co	of the funded period, NRAD51 and the five hRA	we focused on the D51 paralogs. We	e characterizatic detected strong	on of interaction		
Approved for Public Re 13. ABSTRACT (Maximum 200 W) During the third year interactions between h suggesting a stable co signals between hRAD51	of the funded period, NRAD51 and the five hRA paralogs increased in	we focused on the D51 paralogs. We eractions. Some of	e characterizatic detected strong of these weaker i	on of interaction		
Approved for Public Re 13. ABSTRACT (Maximum 200 W) During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which	of the funded period, NRAD51 and the five hRA pmplex, and weaker inte paralogs increased in may indicate a regulat	we focused on the D51 paralogs. We eractions. Some of the presence of	e characterizatic detected strong of these weaker i ATP and decrease	on of interaction ed in the		
Approved for Public Re 13. ABSTRACT (Maximum 200 W) During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor	of the funded period, NRAD51 and the five hRA Demplex, and weaker inte paralogs increased in may indicate a regulat	we focused on the D51 paralogs. We eractions. Some of the presence of ory role for ader	2 characterizatic detected strong of these weaker i ATP and decrease hosine nucleotide	on of interaction nteraction ed in the es. We also		
Approved for Public Re 13. ABSTRACT (Maximum 200 W) During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor	of the funded period, NRAD51 and the five hRA mplex, and weaker inter paralogs increased in may indicate a regulat of to purify the paraloc cance of these interac	we focused on the D51 paralogs. We eractions. Some of the presence of ory role for ader	2 characterizatic detected strong of these weaker i ATP and decrease hosine nucleotide	on of interaction ed in the es. We also		
Approved for Public Re 13. ABSTRACT (Maximum 200 W) During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor the functional signifi currently in progress.	of the funded period, NRAD51 and the five hRA mplex, and weaker inter paralogs increased in may indicate a regulat of to purify the paraloc cance of these interac	we focused on the D51 paralogs. We eractions. Some of the presence of ory role for ader	2 characterizatic detected strong of these weaker i ATP and decrease hosine nucleotide	on of interaction nteraction ed in the es. We also		
Approved for Public Re 13. ABSTRACT (Maximum 200 W) During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor the functional signifi	of the funded period, NRAD51 and the five hRA mplex, and weaker inter paralogs increased in may indicate a regulat of to purify the paraloc cance of these interac	we focused on the D51 paralogs. We eractions. Some of the presence of ory role for ader	2 characterizatic detected strong of these weaker i ATP and decrease hosine nucleotide	on of interaction nteraction ed in the es. We also		
Approved for Public Re 13. ABSTRACT (Maximum 200 W) During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor the functional signifi currently in progress.	of the funded period, NRAD51 and the five hRA Domplex, and weaker inter a paralogs increased in may indicate a regulat of to purify the paraloc cance of these interact	we focused on the D51 paralogs. We eractions. Some of the presence of ory role for ader	2 characterizatic detected strong of these weaker i ATP and decrease hosine nucleotide	on of interaction nteraction ed in the es. We also		
Approved for Public Re 13. ABSTRACT (Maximum 200 W) During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor the functional signifi currently in progress.	of the funded period, NRAD51 and the five hRA Domplex, and weaker inter a paralogs increased in may indicate a regulat of to purify the paraloc cance of these interact	we focused on the D51 paralogs. We eractions. Some of the presence of ory role for ader	2 characterizatic detected strong of these weaker i ATP and decrease hosine nucleotide	on of interaction nteraction ed in the es. We also		
Approved for Public Re 13. ABSTRACT (Maximum 200 W) During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor the functional signifi currently in progress.	of the funded period, NRAD51 and the five hRA Domplex, and weaker inter a paralogs increased in may indicate a regulat of to purify the paraloc cance of these interact	we focused on the D51 paralogs. We eractions. Some of the presence of ory role for ader	2 characterizatic detected strong of these weaker i ATP and decrease hosine nucleotide	on of interaction ed in the es. We also		
Approved for Public Re 13. ABSTRACT (Maximum 200 WA During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor the functional signifi currently in progress. 14. SUBJECT TERMS:	of the funded period, nRAD51 and the five hRA omplex, and weaker inte paralogs increased in may indicate a regulat of the purify the paralo cance of these interact	we focused on the aD51 paralogs. We eractions. Some of the presence of ory role for ader ogs and heterodime etions and the rol	2 characterizatic detected strong of these weaker i ATP and decrease nosine nucleotide ers thereof. Exam le of adenosine n	on of interaction interaction ed in the es. We also mination of nucleotides		
Approved for Public Re 13. ABSTRACT (Maximum 200 WA During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor the functional signifi currently in progress. 14. SUBJECT TERMS:	of the funded period, nRAD51 and the five hRA omplex, and weaker inte paralogs increased in may indicate a regulat of the purify the paralo cance of these interact	we focused on the aD51 paralogs. We eractions. Some of the presence of ory role for ader ogs and heterodime etions and the rol	2 characterizatic detected strong of these weaker i ATP and decrease nosine nucleotide ers thereof. Exam le of adenosine n	on of interaction interaction ed in the es. We also ination of sucleotides =		
Approved for Public Re 13. ABSTRACT (Maximum 200 WA During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor the functional signifi currently in progress. 14. SUBJECT TERMS:	of the funded period, NRAD51 and the five hRA Domplex, and weaker inter a paralogs increased in may indicate a regulat of to purify the paraloc cance of these interact	we focused on the aD51 paralogs. We eractions. Some of the presence of ory role for ader ogs and heterodime etions and the rol	2 characterizatic detected strong of these weaker i ATP and decrease nosine nucleotide ers thereof. Exam le of adenosine n	TRIBUTION CODE on of interaction anteraction ed in the es. We also mination of mucleotides f ER OF PAGES 8		
Approved for Public Re 13. ABSTRACT (Maximum 200 WA During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor the functional signifi currently in progress. 14. SUBJECT TERMS: breast cancer, hRAD51 17. SECURITY CLASSIFICATION	of the funded period, nRAD51 and the five hRA omplex, and weaker inte paralogs increased in may indicate a regulat to purify the paralo cance of these interact	we focused on the D51 paralogs. We eractions. Some of ory role for ader ogs and heterodime ctions and the rol	2A2 12b. DIS 12b. DIS 14b. DIS 14b. DIS 15. NUMBE 16. PRICE	TRIBUTION CODE on of interaction interaction ed in the es. We also mination of sucleotides in the state of the state of the state state of the state		
Approved for Public Re 13. ABSTRACT (Maximum 200 WA During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor the functional signifi currently in progress. 14. SUBJECT TERMS: breast cancer, hRAD51 17. SECURITY CLASSIFICATION OF REPORT	of the funded period, iRAD51 and the five hRA omplex, and weaker inte paralogs increased in may indicate a regulat t to purify the paralo cance of these interac homologs, DNA recombin 18. SECURITY CLASSIFICATION OF THIS PAGE	we focused on the aD51 paralogs. We eractions. Some of the presence of ory role for ader ogs and heterodime etions and the rol	2A2 12b. DIS 12b. DIS 14b. DIS 14b. DIS 15. NUMBE 16. PRICE	TRIBUTION CODE on of interaction anteraction ed in the es. We also mination of mucleotides f ER OF PAGES 8		
Approved for Public Re 13. ABSTRACT (Maximum 200 WA During the third year interactions between h suggesting a stable co signals between hRAD51 presence of ADP which continued in our effor the functional signifi currently in progress. 14. SUBJECT TERMS: breast cancer, hRAD51 17. SECURITY CLASSIFICATION	of the funded period, nRAD51 and the five hRA omplex, and weaker inte paralogs increased in may indicate a regulat ct to purify the paralo cance of these interact homologs, DNA recombin 18. SECURITY CLASSIFICATION	we focused on the D51 paralogs. We eractions. Some of ory role for ader ogs and heterodime etions and the rol	2A2 CATION 2 Characterizatic detected strong 12b. DIS 12b. DI	TRIBUTION CODE on of interaction interaction ed in the es. We also mination of sucleotides ER OF PAGES 8 CODE		

Table of Contents

Cover1
SF 2982
Table of Contents
Introduction4
Body4
Key Research Accomplishments6
Reportable Outcomes7
Conclusions7
References7
Appendices

(4) INTRODUCTION:

DNA repair is essential for genomic integrity, and failure of repair pathways may lead to a mutator phenotype and to tumorigenesis (1, 2). Homologous DNA recombination (HR) is a prominent pathway for the repair of double-strand break (DSB) and other DNA lesions and is dependent on human RAD52 epistasis group proteins including hRAD51 and its five paralogs (3-6). Evidence for the involvement of both BRCA1 and BRCA2 in hRAD51-mediated repair processes is accumulating (7-9). The purpose of this study is to characterize physical and functional interactions among hRAD51 paralogs and BRCA1/BRCA2 in order to better understand HR.

(5) BODY

Brief summary of previously reported work

<u>Aim I:</u> done, results have been reported in Cancer Res. (Ref.11). The coding region of the hRAD51 gene has been examined for mutations in tumor tissues with high frequencies of 15q15 deletions, and the promoter region has been tested for hypermethylation. No changes have been found in the tumors compared to normal tissues.

<u>Aim II:</u> partially done, results have been reported. All known human RAD51 homologs involved in mitotic recombination have been cloned into appropriate vectors and overexpressed in bacteria or in a baculovirus system. Polyclonal antibodies against these proteins have been generated and characterized. Monoclonal antibodies have become available. Previously unknown chromosomal locations of RAD51 homologs have been determined. Purification of XRCC2 has been reported.

Aim III: partially done. Basic interactions between hRAD51 and its human paralogs have been reported. We have cloned members of the human RAD52 epistasis group into appropriate expression vectors: hRAD51, hRAD51B (a.k.a. hRAD51L1 or hRAD51-H2), hRAD51C (a.k.a. hRAD51L2), hRAD51D (a.k.a. hRAD51-H3, hRAD51L3), hXRCC2, hXRCC3 as well as hRAD52 and hRAD54. This collection allowed us to clearly identify static interactions between these six human mitotic RecA homologs, hRAD52, hRAD54 (Aim IIIa). Interaction studies were done using an in vitro GST-fusion-IVTT method as described (10, 11). In this assay, hRAD51 gave a strong signal with itself as expected. We also found (weaker) interactions with hRAD51B, hRAD51D, hXRCC3, and hRAD52. hRAD51B showed strong interactions with hRAD51C and hRAD51D and weaker signals with hXRCC3. hRAD51C interacted strongly with hRAD51B and hRAD51D and less strongly with XRCC3. hRAD51D interactions were detected with all other hRAD51 homologs. The strongest signal was seen with hXRCC2, and hXRCC2 seems to interact only with hRAD51D. hXRCC3 also seems to be able to interact with all hRAD51 homologs, but seems to bind best to hRAD51D. In addition, we detected signals in the hRAD52 and hRAD54 lanes. hRAD52 bound to hRAD51, hRAD51B, hRAD51D, hXRCC3 and itself.

Summary of current work

Aim IIIa: Interaction studies. Since all RAD51 homologs have highly conserved ATP binding domains (Walker boxes) and ATP binding/hydrolysis seems to have a crucial role during homologous recombination (12), we further investigated whether interactions between the hRAD51 homologs could be modified by adenosinenucleotide. We found that GSThRAD51 precipitates **IVTT**hRAD51D in the presence of ADP plus sodium aluminum tetrafluoride (NaAlF4; Figure 1, Lanes 4 and 5). The function of NaAlF₄ is unknown. However, it is generally regarded that NaAlF₄ NDP-bound Walker stabilizes A/B motif proteins in a pseudotransition state.



Figure 1. GST/IVTT precipitation analysis. A GST fusion construct of hRAD51 (GST-hRAD51) was bound to glutathione beads and exposed to 35 S-labeled *in vitro* transcribed translated hRAD51D (IVTT-hRAD51D) in the presence of 1mM ADP, ATP, ATP γ S and/or ADP-NaAlF₄ as indicated. Precipitated proteins were separated by PAGE and visualized using a PhosphoImager system. No significant IVTT material was precipitated by GST alone (data not shown).

As mentioned above, hRAD51D forms a very stable heterodimer with hXRCC2. We have purified the hRAD51D/hXRCC2 heterodimer using a baculovirus expression system. Immuno-precipitation experiments (Figure 2) support the conclusion that hRAD51D/hXRCC2 interacts with the hRAD51-ADP- NaAlF₄ transition state.

	purified proteins			IP with hRAD51 antibody						
Mg-ATP							+			
Mg-ADP, NaAlF ₄								+		+
hRAD51	+			+	+	+	+	+	+	+
hXRCC2		+			+		+	+		
hXRCC2/hRAD51D			+			+			+	+
hXRCC2 antibody			Ó		1					
hRAD51 antibody	-									

Figure 2. Immunoprecipitation of hXRCC2 and hXRCC2-hRAD51D with hRAD51. Purified hRAD51 was bound to Protein A beads preincubated with hRAD51 antibody and subsequently exposed to purified hXRCC2 or hXRCC2-hRAD51D in the absence of adenosine nucleotide or in the presence of 1mM ADP or ADP-NaAlF₄ as indicated. Precipitated proteins were separated on SDS-PAGE and probed with a monoclonal antibody to hXRCC2 (hXRCC2; upper panel) or the hRAD51 antibody (hRAD51; lower panel) Lanes 1-3 contain purified proteins which had not been subjected to immunoprecipitation.

Preliminary data have also shown that binding of hRAD51 to the hXRCC3-GST fusion protein is influenced by adenosine nucleotides. Binding of hRAD51 to hXRCC3 is increased in the presence of ADP and ATP in the presence of NaAlF₄ (Figure 8).

These data indicate a regulatory role of ATP hydrolysis and/or binding. We currently perform experiments using monoclonal



Figure 3. GST/IVTT precipitation analysis (see legend to Figure 1).

antibodies to XRCC3 and hRAD51C, respectively, to further confirm these findings.

In addition, co-immunoprecipitation experiments using specific antibodies and HeLa cell extracts are currently in progress.

<u>Aim III b) Purification and characterization of hRAD51 derivatives.</u> Purification of hXRCC2, hRAD51D/hXRCC2 heterodimer, hRAD51C/hXRCC3 heterodimer have been completed. Purification of hRAD51B/hRAD51C heterodimer and biochemical studies to characterize the human RAD51 paralogs are currently in progress.

<u>Aim IV a) Interactions of hRAD51 derivatives with BRCA2, BARD1, and RPA</u>. In order to test BRCA2 in our GST interaction assay, it was necessary to divide the 10.3kb BRCA2 ORF into four overlapping fragments of approximately 3kb each since we could not generate a full length labeled BRCA2 IVTT protein in sufficient quantities. Interaction experiments with these fragments, the hRAD51 paralogs, BARD1, and RPA are currently in progress.

(6) KEY RESEARCH ACCOMPLISHMENTS:

• Interactions between hRAD51 and its homologs have been further characterized (Aim IIIa).

• Interactions between hRAD51 homologs seem to be modified in the presence of ATP or ADP (Figures 1-3). These findings support a regulatory role for adenosine nucleotides during HR.

(7) REPORTABLE OUTCOMES

All findings listed under (6) are reportable. Data describing interactions of hRAD51 and paralogs have been presented at the Era of Hope meeting in Orlando (Sept.25-28, 2002). A manuscript about the function of hXRCC2 (not described here) has been submitted to Molecular Cell for publication.

(8) CONCLUSIONS

Multiple interactions exist between hRAD51 and the human hRAD51 paralogs which seem to be in part regulated by adenosine nucleotides. Purification and biochemical characterization of these proteins is in progress in order to study the functional relevance of these interactions in the process of homologous recombination.

(9) **REFERENCES**

- 1. Loeb, L. A. A Mutator Phenotype in Cancer. Cancer Res, 61: 3230-3239, 2001.
- 2. Fishel, R. The Selection for Mismatch Repair Defects in Hereditary Nonpolyposis Colorectal Cancer. Cancer Research, *61*: 7369-7374, 2001.
- Kowalcsykowski, S. C., Dixon, D. A., Eggleston, A. K., Lauder, S. D., and Rehrauer, W. M. Biochemistry of homologous recombination in Escherichia coli. Microbiol Rev, 58: 401-465, 1994.
- 4. Kogoma, T. Stable DNA replication: interplay between DNA replication, homologous recombination, and transcription. MMBR, *61*: 212-238, 1997.
- 5. Baumann, P. and West, S. C. Role of the human RAD51 protein in homologous recombination and double-strand-break repair. TIBS, 23: 247-251, 1998.
- Cox, M. M. RECOMBINATIONAL DNA REPAIR OF DAMAGED REPLICATION FORKS IN ESCHERICHIA COLI: Questions. Annu. Rev. Genet., 35: 53-82, 2001.
- Chen, C.-F., Chen, P.-L., Zhong, Q., Sharp, Z. D., and Lee, W.-H. Expression of BRC Repeats in Breast Cancer Cells Disrupts the BRCA2-Rad51 Complex and Leads to Radiation Hypersensitivity and Loss of G2/M Checkpoint Control. J. Biol. Chem., 274: 32931-32935, 1999.
- Yuan, S.-S. F., Lee, S.-Y., Chen, G., Song, M., Tomlinson, G. E., and Lee, E. Y.-H. P. BRCA2 Is Required for Ionizing Radiation-induced Assembly of Rad51 Complex in Vivo. Cancer Res, 59: 3547-3551, 1999.
- 9. Moynahan, M. E., Chiu, J. W., Koller, B. H., and Jasin, M. Brca1 Controls Homology-Directed DNA Repair. Mol Cell, 4: 511-518, 1999.
- Guerrette, S., Wilson, T., Gradia, S., and Fishel, R. Interactions of Human hMSH2 with hMSH3 and hMSH2 with hMSH6: Examination of Mutations Found in Hereditary Nonpolyposis Colorectal Cancer. Mol Cell Biol, 18: 6616-6623, 1998.

- 11. Schmutte, C., Marinescu, R. C., Sadoff, M. M., Guerrette, S., Overhauser, J., and Fishel, R. Human exonuclease I interacts with the mismatch repair protein hMSH2. Cancer Res, *58*: 4537-4542, 1998.
- 12. Baumann, P., Benson, F. E., and West, S. C. Human Rad51 Protein Promotes ATP-Dependent Homologous Pairing and Strand Transfer Reactions In Vitro. Cell, 87: 757-766, 1996.

(10) APPENDICES

none.