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<b>13. SUPPLEMENTARY NOTES</b>				
<b>14. ABSTRACT (ca. 200 words)</b> Our specific aims were to map origins of replication in the breast cancer genome and compare these sites with genomic positions that bind the estrogen receptor (ER) and genomic regions of DNA amplification. Correlations would support our hypothesis that ER adjacent to replication origins may interact with the replication machinery to drive DNA amplification, hallmark of many cancers. We began our experiments with the well-studied breast cancer cell line MCF7. We refined the lambda exonuclease method to obtain nascent DNA strands (up to 20 times enriched) and demonstrated that the myc origin maps to the same position in HeLa and MCF7 cells. We size fractionated nascent DNA (500-1500 nt) for next generation sequencing which gives better resolution and sensitivity than DNA microarrays. The results of Illumina sequencing revealed that our nascent DNA sample from the whole genome included many of the origins previously reported for 1% of the human genome. Development of this technology serve as the basis for our IDEA Extension grant which was funded to extend the work begun in this parent grant.				
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## FINAL PROGRESS REPORT

### INTRODUCTION

Recent data support the hypothesis that DNA amplification plays a role in establishing the malignant cell phenotype in cancer (Nikolsky et al., 2008). However, the basic mechanism underlying DNA amplification has not yet been elucidated, though it may be more common than originally thought (Gomez 2008; Gomez and Antequera 2008). There appears to be a link between the steroid hormone estrogen and many forms of breast cancer, but the detailed mechanism is unknown. Estrogen can turn on gene expression and thus activate the production of the proteins encoded by these genes. Our recent results in a model system indicated that a steroid hormone can induce gene amplification in which re-replication creates extra copies of the gene. This in turn will also increase production of the protein encoded by the amplified gene. Hormonal induction of gene amplification is a new paradigm for how hormones work, and we wish to see if it applies to breast cancer. We wish to examine if a correlation exists between estrogen receptor (ER) binding at novel sites in the breast cancer genome and juxtaposition with replication origins that escape normal cellular controls and re-replicate, leading to DNA amplification. The recent observations that estrogen induces cell proliferation by retention of MCM proteins in the nucleus and by induction of the loading factor Cdt1 (Pan et al., 2006) support our hypothesis, especially since increases in MCM proteins and Cdt1 have been shown to induce DNA amplification in yeast (Gopalakrishnan et al., 2001; Nguyen et al., 2001; Green et al., 2006) and increased Cdt1 results in re-replication in human cells (Dorn et al., 2008). The N-terminus of Cdt1 is important for re-replication, perhaps through interactions with PCNA and/or cyclin (Teer and Dutta, 2008). Cdt1 and its inhibitor geminin are deregulated in human tumors (Petropoulou et al., 2008). Moreover, stalled replication forks and DNA re-replication lead to DNA breakage and rearrangements (Green and Li, 2005; Raveendranathan et al., 2006; Zhu and Dutta, 2006; Dutta, 2007; Hook et al. 2007) which is a hallmark of cancer. Our research may provide a new paradigm for hormonal induction of breast cancer via gene amplification, leading to new methods of diagnosis and treatment.

### BODY

In the research supported by this grant, we proposed to map estrogen receptor binding sites, origins of replication and regions of DNA amplification in surgically derived breast cancer tissue (see Appendix 1: DOD meeting abstract). We report our progress on these three specific aims. Also, we report on relevant recent publications that support our working model. The P.I. (Susan Gerbi) and two co-P.I.s (Alex Brodsky and Ben Raphael) meet together with their lab personnel roughly once a month to review past results and design future experimental strategies.

The text that follows is the revised final report for this grant, organized according to the Statement of Work that was listed in the approved grant application which was organized according to the three Specific Aims. Figures and tables are located at the end of the document.

## **Statement of Work Item 1: Map ER binding in the human genome**

Months 1-6: Work out the methodology (Gerbi and Brodsky labs)

Chromatin from Breast Cancer Tissue - this is the starting material for each of the three specific aims. In our previous progress report we reported the difficulty in obtaining breast cancer tissue samples, as recent changes in clinical protocols now result in the vast majority of patients receiving chemotherapy prior to breast cancer surgery. We cannot use tissue derived from patients with neoadjuvant therapy, and therefore, our potential supply of material was drastically reduced. As reported previously, we expanded our network of clinical collaborators to include other surgeons and pathologists at both R.I. Hospital and Women and Infants Hospital. With this increased outreach, we have now obtained some surgically derived tissue samples and are hoping for more samples (see **Table 1**). To work out the methodology for chromatin isolation from breast cancer tissue, we used material from the R.I. Hospital breast cancer tumor bank that did not meet our criteria above, but was available for these pilot experiments. We determined that tissue that was freshly obtained and frozen from a current surgery was equivalent to tissue that had been stored frozen for a period of time. For chromatin immunoprecipitation (ChIP) procedures, the tissue is subjected to formaldehyde fixation, homogenized and then sheared by sonication. We found that the breast cancer tissue (portions of 1.0-1.5 cm tumor specimens) was very fibrous and hard to break open by standard homogenization, resulting in low yields of chromatin. We purchased a microhomogenizer (the same model used by Dr. Peggy Farnham for her breast cancer chromatin studies) to use for breast cancer tissue disruption. Our initial results were promising. We were able to prepare sonicated chromatin from cell lines and tissues using the microhomogenizer averaging less than 500 bp in size (**Figure 1**) and that works well for ChIP.

Months 7-20: Carry out ER ChIP-chip experiments

(Brodsky, Gerbi and Raphael labs)

ChIP of ER binding sites in the breast cancer genome - these data had already been obtained by co-P.I. Alex Brodsky for MCF7 cultured breast cancer cells (Carroll et al., 2005 and 2006), and we planned to use them as a reference source as we developed the ChIP methodology for breast cancer tissue. In the grant application we proposed to do ChIP-chip (DNA microarrays of chromatin immunoprecipitated samples). However, that method has now been superseded by ChIP-Seq (DNA sequencing of chromatin immunoprecipitated DNA samples). This method has greater sensitivity, in addition to its better resolution than ChIP-chip. ChIP-Seq revealed 10-13,000 ER-alpha binding sites in the genome of MCF7 breast cancer cells (Lin et al., 2007; Fullwood et al., 2009; Welboren et al., 2009; Hurtado et al., (2011). Because of the difficulty in obtaining tumor sample specimens (see preceding section) and the fact that none of the samples we received were FISH positive for HER2 gene amplification (**Table 1**), we decided to carry out our experiments MCF7 cultured breast cancer cells. An added bonus of this cell culture model system is that the ER-alpha binding sites have already been mapped by several groups by the more advanced method of ChIP-Seq (see above), obviating the need for us to map these sites.

Months 21-24: confirm ER binding site candidates (Gerbi lab)

This was no longer necessary since other groups have already mapped ER-alpha binding sites in the MCF7 breast cancer genome (see above). The decreased experimentation needed for statement of work item 1 allowed us to devote more time to develop methodology for statement of work item 2 (see below).

**Statement of Work Item 2: Map binding sites for the Origin Recognition Complex (ORC) across the human genome (Brodsky, Gerbi and Raphael labs)**

Months 7-20: Map replication origins in the breast cancer genome

As stated above, because of the paucity of breast cancer tissue material, we decided to do our initial experiments to map replication origins on the well studied MCF7 cell line where the ER binding sites are already mapped. This also has the advantage of sample homogeneity which would be a concern for tissue from breast cancer tumors. Our plan was to analyze DNA from unsynchronized cells in order to capture all origins regardless of when during S phase they are activated. We reported in previous progress reports that we obtained polyclonal antibodies for human ORC2 and Cdt1 and a monoclonal antibody against human ORC6 from Aloys Schepers, antibody against human ORC1 from Mel DePamphilis and a mammalian expression clone for FLAG-tagged human ORC1 from Dr. Kohji Noguchi (a former post-doc with Dr. Mel DePamphilis). The Brodsky lab checked these antibodies by Western blots, but there was high background with multiple bands. Moreover, two tries of ChIP with ORC antibody were unsuccessful. Discussions that P.I. Susan Gerbi had with Drs. Aloys Schepers and Michael Leffak at the Cold Spring Harbor DNA Replication Meeting revealed that ChIP on mammalian cells with ORC2 antibodies has a high background. Instead, the Brodsky lab considered cloning ORC1 into a FLAG-tag vector for transfection into MCF7 breast cancer cells, reasoning that ChIP with a FLAG antibody should give better results. However, we decided instead to pursue the more promising approach described below.

The goal is to map DNA replication origins in the breast cancer genome. A problem with the approach of mapping ORC binding sites is that ORC also binds to silent origins that are not used, so we would not know which origins are active in the breast cancer cells. Moreover, as described above, there are problems with the antibodies against ORC. We decided that superior results would be obtained with a more successful approach to isolate small nascent DNA to map replication origins directly by sequencing the nascent strands, rather than using ChIP-chip or ChIP-seq to map ORC binding sites. The short nascent strand sequencing approach allows us to identify by this direct method all origins that are active in the breast cancer genome. Nascent strands have been used to map replication origins for a limited portion (1% ENCODE project) of the human genome (Lucas et al., 2007; Cadoret et al., 2008) and has given more reliable results than BrdU labeling of non-lambda exonuclease treated DNA (Birney et al., 2007; Karnani et al., 2007) where results from the latter do not agree with results of mapping replication bubbles trapped in agarose (Mesner et al., 2010).

In order to identify origins of DNA replication throughout the genome of MCF7 cells, nascent DNA was prepared according to our previous protocol (Gerbi and

Bielinsky, 1997). As summarized in **Figure 2**, genomic DNA was prepared from mid-log phase cells using DNAzol (Invitrogen, Calsbad, CA) and resuspended in TE. Replicative Intermediate (RI) DNA was enriched by passing the genomic DNA over a column of BND-cellulose. The ends of the RI DNA were phosphorylated using T4 Polynucleotide Kinase (New England Biolabs, Ipswich, MA). Next the DNA was digested with lambda exonuclease to enrich nascent strands which are resistant to lambda-exonuclease digestion due to the presence of an RNA primer at their 5' end. Finally, the nascent strands were size fractionated (500 – 1500 bp) on low melting point agarose to eliminate background from Okazaki fragments which occur throughout the genome. Enrichment of nascent strands was confirmed by real-time PCR assaying for enrichment of the c-myc origin of replication (Tao et al., 2000). Primers were designed to be specific to locus 11 (the c-myc origin) and a non-origin sequence about 6 kb upstream at locus 1. The origin mapping experiments are being carried out by Dr. Michael Foulk, a talented postdoc in the Gerbi lab. Due to the fact that the c-myc origin of replication was discovered in HeLa cells, we first compared the enrichment of nascent strands between HeLa cells and MCF7 cells in order to confirm that the replication origin maps to the same position at the c-myc locus in MCF7 cells. Our results confirmed that this was indeed the case: In HeLa cells, the c-myc origin of replication was enriched about 12 fold, while in the MCF7 cells, it was enriched about 11 fold when we used the DNA nascent strand isolation protocol above, suggesting a replication origin exists at the same locus in MCF7 cells (**Figure 2**). These experiments also demonstrated the feasibility of isolating nascent strands from MCF7 cells for further analysis. Subsequently, we were able to reliably obtain ~ 100-150 ng nascent strands from 100 ug starting genomic DNA from asynchronous MCF7 cells. Real-time PCR showed that the preps were enriched for the c-myc origin between 11.0 and 19.6 fold (**Figure 2**).

Several preparations of nascent DNA from MCF7 cells were pooled and submitted for next generation sequencing. Ben Raphael aligned the resulting Illumina reads to the human genome using MAQ resulting in 5.6 million mapped reads. We counted the number of reads that align to genomic intervals defined by 283 replication origins identified in 1% of the human genome (Cadoret et al., 2008; ENCODE project). For each of these 283 intervals, we compared the read count of the interval to the expected read count under a uniform distribution of reads to intervals. We found that 78 of the 283 origins were enriched ( $P < 10^{-3}$ ) for nascent strands (**Figure 3**; **Table 2**). These initial results were encouraging, and suggest that our nascent DNA preparation is enriching for replication origins. However, our sequencing coverage was not high enough to robustly detect new replication origins and there was some contaminating bacterial DNA.

However, these experiments proved difficult to reproduce. We determined that the cause of this variability was in the poor quality of the preparation of lambda exonuclease we were using (our previous source from Invitrogen had been discontinued so we had switched to enzyme from New England BioLabs). By discussion with Drs. Mechali and Prioleau whose labs are in France, P.I. Susan Gerbi learned that the company Fermentas could prepare high quality lambda exonuclease by special order.

Therefore, we contracted with Fermentas to obtain a high quality, high concentration preparation of lambda exonuclease. The original nascent strand protocol was modified so that the phosphorylated R1 DNA was digested with 240 units of lambda exonuclease (versus 15 units previously) overnight following a protocol developed by Cadoret et al. (2008) for mapping replication origins in the ENCODE subset of the human genome. Using this protocol we achieved about 20 fold enrichment of the c-myc origin in MCF 7 cells, which is excellent. We then, however, noticed some variability in nascent strand enrichment from preparation to preparation. We did controls that revealed that the pH optimum for the Fermentas recombinant lambda exonuclease is broader than for the previous Invitrogen purified enzyme. At the previously used higher pH of 8.8 we found that there was degradation of RNA; this would compromise the integrity of the RNA primers on the nascent DNA and render the nascent DNA susceptible to lambda exonuclease digestion (**Figure 4**). We found that the Fermentas preparation of lambda exonuclease is still active at pH 8.0 and that RNA degradation does not occur at that pH (**Figure 4**). Additionally, we found that heating the samples resulted in degrading the RNA so we have modified the original protocol to eliminate heating steps where possible (**Figure 4**).

For the Illumina sequencing, we isolated about 50-150 ng nascent strands from 100 ug starting genomic DNA. Several nascent strand preparations were pooled (about 500 ng total) and subsequently amplified at the Yale University sequencing facility for Illumina sequencing. However, it required a lot of effort to obtain this amount of nascent DNA. Also, PCR artifacts can be introduced during the amplification step. To overcome these problems, the DOD IDEA Extension grant will allow us to try Helicos rather than Illumina sequencing. The first report of Helicos sequencing appeared just a year ago (Harris et al. 2008) and holds much promise (Gupta 2008). This true single molecule sequencing (tSMS) approach omits the necessity for DNA amplification, significantly reducing the amount of nascent DNA starting material required, about 10 ng. In comparison, the Illumina platform for complete genome coverage requires 500-1000 ng nascent DNA. Moreover, since the nascent DNA preparation enriches for the single stranded leading strands near an origin of replication the sequences should map to opposite strands on either side of the origin. This data will provide a signature for authentic origins of replication reducing the potential for calling false positives. There are only a few Helicos machines in operation world-wide. We have been given access on a fee-for-service basis to the Helicos sequencing machine at the Dana Farber Cancer Institute where one of us (Alex Brodsky) was previously a postdoc prior to joining the faculty at Brown University. We intend to also sequence the nascent DNA using Illumina and compare the results between the two platforms. We anticipate that there will be ~25,000-30,000 replication origins in the human genome. The replication origins, once mapped, will be compared to estrogen receptor binding sites (the ENCODE data showed a correlation for c-JUN and c-FOS as potential regulators of origins; Cadoret et al., 2008) and to regions of DNA amplification in breast cancer cells.

To sum up, we have accomplished much more for Statement of Work Item 2 than originally presented in the grant proposal. Instead of using ChIP-chip or ChIP-Seq to map ORC binding sites, we have refined the methodology to allow direct mapping of all



active origins in the breast cancer genome by isolation of short nascent strands of DNA and sequencing them. Our pilot run on the Illumina platform was successful. Based on this progress, we are grateful to have received a DOD IDEA Extension award.

### **Statement of Work Item 3: Mapping DNA amplification sites in the breast cancer genome**

One of us (Ben Raphael, co-P.I.) and others have identified chromosomal changes in the genome of MCF7 breast cancer cells (Volik et al. 2003 and 2006; Raphael et al., 2008; Hampton et al., 2008), including sites of DNA amplification. Therefore, once we have mapped the replication origins in MCF7 cells (Statement of Work Item 2), we can directly compare their locations with the already mapped locations of ER binding sites (Statement of Work Item 1) and DNA amplification (Statement of Work Item 3), as per our specific aims for this grant. If we extend the study to breast cancer tissue, array comparative genome hybridization (aCGH) will be used to determine the sites of DNA amplification in this tissue.

#### Months 1-6: Refine methods to analyze DNA amplification (Raphael lab)

DNA double strand breaks have been shown to play a role in DNA amplification. As stated in our previous progress report, co-P.I. Ben Raphael developed a novel method called Neighborhood Breakpoint Correlation (NBC) to identify correlated rearrangement breakpoints from CGH data (*BMC Bioinformatics*, in revision). Unlike previous methods for aCGH analysis that focus on finding common genomic intervals of amplification or deletion that might harbor oncogenes or tumor suppressor genes, respectively, NBC focuses on the precise localization of the boundaries (breakpoints) of these intervals. We hypothesize that pairs of such highly conserved breakpoints might indicate fusion genes or other common rearrangements. The algorithm employs a statistical model derived from the binomial distribution to assess the statistical significance of breakpoints that shared by multiple patients. The algorithm also identifies genes or pairs of genes that each contains one or more breakpoints in a statistically significant number of patients.

In preliminary analysis, Ben Raphael examined a collection of 36 primary prostate tumors for breakpoints in the well-known TMPRSS2-ERG fusion gene (Tomlins et al. 2005). He applied NBC to identify changes in copy number (breakpoints) in each patient and then identified common breakpoints that appear in a statistically significant number of patients. Specifically, he identified 12 statistically significant rearrangements, one of which is the TMPRSS2-ERG fusion gene. It is detected in 5 patients with a p-value of  $2.7 \times 10^{-10}$  (**Figure 5**). In a larger analysis, he examined a collection of data from 233 patients with glioblastoma, including 227 primary tumor samples and 107 matched blood samples from The Cancer Genome Atlas. He predicted 93 statistically significant rearrangements that are further classified as gene truncations, germline structural variants, and fusion genes. The power of his method to detect correlated breakpoints increases with larger sets of patients. We will apply these methods to the aCGH data that will be generated in the present research project on DNA amplification in breast cancer cells. This will allow us to uncover additional candidate fusion genes or regulatory fusions, particularly fusions near ER binding sites.

We now report here some new data derived by computational analysis by co-P.I. Ben Raphael. He combined aCGH data and ESP data from estrogen receptor (ER) binding data in MCF7 breast cancer cells determined initially by co-P.I. Alex Brodsky using chromatin immunoprecipitation (ChIP-chip) (Carroll et al. 2006). The original ChIP-chip study (Carroll et al., 2006) interpreted the data in the context of the reference human genome, even though it is well known that MCF7 exhibits extensive genomic aberrations including copy number changes and structural rearrangements. We examined how knowledge of these genomic changes affects the interpretation of the ChIP-chip data. Using scan statistics (Glaz et al., 2001), we identified regions of the reference genome that contained significantly few or significantly many ER binding sites. We found 38 gaps, defined as sequenced genomic regions > 6.9 Mb with no ER binding sites. Under the null hypothesis of ER sites distributed uniformly on the genome, the probability of finding one such gap is  $< 10^{-4}$ . The copy numbers of the probes in these gaps determined using Agilent 44K array CGH had mean  $\log_2$ -ratio of -0.28, significantly below the mean value of -0.05 over all probes (p-value by T-test  $< 10^{-100}$ ) and implying that the gaps in ER binding sites are a result of deletions in the MCF-7 genome.

We next identified unusual clusters of ER binding sites in the ChIP-chip data, where a cluster was defined as 12 or more ER sites in a 1 Mb region of the genome. Under the null hypothesis, the probability of such a cluster is  $< 6 \times 10^{-5}$ , but we identified 11 clusters in the data. The mean  $\log_2$ -ratio in these segments is 1.3, suggesting that clusters of ER binding sites are preferentially found in amplified regions. This could be due to the fact that the ChIP-chip assay has higher sensitivity in amplified regions, or due to the fact that the statistical model used to call binding sites is imprecise in amplified regions. To assess whether there might be amplification of the regions harboring ER clusters, we examined BAC array CGH data on 51 breast cancer cell lines from (Neve et al., 2006). We found that one of the 11 ER clusters is preferentially amplified in ER-positive cell lines and preferentially deleted in ER-negative breast cancer cell lines ( **Figure 6** ). Moreover, we found differential expression of one of the four genes in this region using OncoPrint (Rhodes et al., 2007). TLE3 is significantly under-expressed in ER-negative breast tumors compared to ER-positive breast tumors (Sotiriou et al., 2003; Minn et al., 2005; Wang et al., 2005; Hess et al., 2006) (p-values  $10^{-12}$ ,  $2 \times 10^{-11}$ ,  $2 \times 10^{-10}$ ,  $8 \times 10^{-9}$  based on t-test). This result is consistent with the amplification of this region as determined by array CGH, and also suggests estrogen dependent regulation of this gene.

We also looked for structural rearrangements that might yield regulatory fusions between bound ER sites in the ChIP-chip data and genes identified as differentially regulated in response to estrogen. Using ESP data to identify rearrangements (Volik et al., 2006; Raphael et al., 2008), we found 27 examples of such candidate fusions ( $P < 2 \times 10^{-11}$  by a permutation test.). Moreover, in 11/27 cases, the genes involved in these putative fusions have no bound ER site within 100 kb of the transcription start site. Several of these genes are implicated in breast cancer including BRCC3 (a subunit in the BRCA1/2 containing protein complex), PTK6, and STK6.

### Months 21-24: Whole genome SNP arrays (Brodsky and Gerbi labs)

Having shifted from tumor specimens to MCF7 breast cancer cells, SNP arrays were no longer needed.

To sum up, we have done more work in the Statement of Work Item 3 than originally described in the grant application, having developed many tools for computational analysis.

Concluding remarks – The recent finding that the transcription factor c-Myc interacts with the pre-replication complex to control DNA replication (Dominguez-Sola et al., 2007; Lebofsky and Walter, 2007) and that the androgen receptor interacts with MCM7 of the pre-replication complex (Shi, 2008) provides precedence for our hypothesis that the ligand-bound estrogen receptor may play a direct role in regulating replication origins beyond its traditional role as a transcription factor. We are grateful for the DOD funding that allowed us to initiate experiments to test our hypothesis and look forward eagerly to results from the DOD funded IDEA Extension award.

### **KEY RESEARCH ACCOMPLISHMENTS**

- Further refinement in the method to isolate nascent (newly replicated) DNA, lowering the pH to prevent RNA degradation during lambda exonuclease digestion, thereby reducing the prep-to-prep variability in origin enrichment.
- PCR mapping of the myc replication origin showed that it is located in the same position in HeLa and MCF7 cells.
- A trial run of Illumina sequencing of nascent strands included many of the replication origins previously reported for 1% of the human genome
- Improvement of the methodology for analysis of aCGH data to identify common aberrations and common breakpoints.
- Computational analysis suggested that clusters of ER binding sites are preferentially found in amplified regions.

### **REPORTABLE OUTCOMES**

- Method to isolate nascent (newly replicated) DNA
- Preliminary data mapping replication origins in the breast cancer genome
- Methodology for analysis of aCGH data to identify common aberrations and common breakpoints.
- Computational results suggesting that clusters of ER binding sites are preferentially found in amplified regions.

We anticipate writing a paper for publication in a high profile journal describing the map of replication origins in the entire human genome (from breast cancer cells) once our data are complete.

Based on our successful upward trajectory with these experiments, we have received a DOD Idea Expansion award. This award would allow us to complete and expand our promising experiments.

## **CONCLUSION**

Recent publications cited in this progress report support our hypothesis that the estrogen receptor may interact with the replication machinery and promote DNA amplification in breast cancer cells. We have improved the experimental protocol from what was initially approved in this grant. Instead of identifying origins by ORC ChIP, we are isolating size-fractionated nascent strands to use them for next generation sequencing. Our results will be the first to map replication origins on the entire human genome. The data will be compared to map positions of ER binding sites in the genome and regions of DNA amplification. A positive correlation will directly support our hypothesis and will provide a new way of thinking about the role of steroid hormones in cancer. The results will begin to elucidate the mechanism of induction of DNA amplification and could provide a platform for new methods of diagnosis and treatment of breast cancer.

### **Personnel paid from this grant:**

(nb – these are personnel in the three different groups of PI and co-PIs; some of the lab personnel only worked briefly on this project)

PI: Susan A. Gerbi

Co-P.I.s: Alexander Brodsky, Benjamin Raphael

Postdoctorals: Michael Foulk, Yutaka Yamamoto

Graduate Students: Crystal Kahn, Anna Ritz

Research Assistants: Jacob Bliss, Megan Frayne, Mark Grabiner, Sara Hillenmyer, Ingrid Mercer, Shellee Morehead, Hannah Sanford, Heidi Smith

Undergraduate Dishwashers: Carolyn Crisp, Sydney Ember, Emily Hartman, Theeradej Thaweerattanasin

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## APPENDICES

### Appendix 1: Meeting Abstracts

Meeting abstract: DOD 2008 Era of Hope meeting (Baltimore, MD)

#### Hormonal Involvement in Breast Cancer Gene Amplification

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Genetic instability and rearrangements, including gene amplification, is a hallmark of cancer. Amplification of the HER2 (ErbB2/Neu) gene occurs in invasive breast cancer (~25%) and in ductal carcinoma *in situ* (50-60%). HER2 amplification and concomitant over-expression of this growth factor promotes cancer cell growth, acting as a metastasis-promoting factor. It would be desirable to prevent HER2 gene amplification, thereby moderating the aggressive growth of breast cancer cells. The problem is that no one knows what triggers gene amplification. Our recent research suggested that the trigger may be the steroid hormone estrogen. Do genetic or epigenetic changes produce novel binding site(s) for the estrogen receptor (ER) near the HER2 replication origin, inducing gene amplification? Our hypothesis is that ER interacts with the replication machinery to drive re-replication of the HER2 locus, resulting in DNA amplification.

Our specific aims and the study design are:

**(1)** Map ER binding sites in surgically derived HER2 amplified breast cancer tissue, using chromatin immunoprecipitation (ChIP) with an antibody against ER. The immunoprecipitated DNA will be used as a probe for DNA microarray chips ("ChIP-chip") to screen the human genome for hormone receptor binding sites. We will look for differences in ER binding sites between cancer cells and non-cancer cells from the same patient. The positive candidates will be confirmed by quantitative PCR following ChIP.

**(2)** Map replication origins using short nascent strands as probes for DNA microarray chips. Data analysis will identify replication origins that are near ER binding sites, with special attention given to novel ER sites in the cancer genome. An alternate and/or confirmatory approach is sequential ChIP ("re-ChIP") on chip experiments where DNA is immunoprecipitated by antibodies against ER and against Origin Recognition Complex polypeptide 2 (ORC2), thereby pulling down DNA fragments bound by both antigens.

**(3)** Quantify level of HER2 amplification and identify sites of co-amplification in the genome. DNA will be isolated from the same tissue samples used for specific aims (1) and (2) for use as probes for whole genome SNP arrays to quantify gene copy

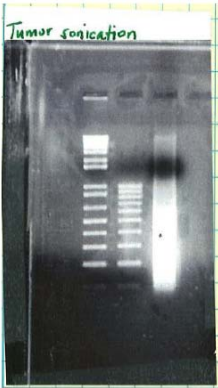
numbers, thereby identifying regions of amplification. The level of HER2 gene amplification will be quantified and any sites of co-amplification will be determined. This study will examine if a correlation exists between ER binding at novel sites in the breast cancer genome and juxtaposition with putative replication origins that escape normal cellular controls and re-replicate, leading to DNA amplification. This may provide a new paradigm for hormonal induction of breast cancer via gene amplification, leading to new methods of diagnosis and treatment. Our results will indicate if there are other regions that co-amplify with the HER2 locus in the ER positive, HER2 amplified breast cancer patient samples. Other co-amplified genes, within the HER2 amplicon and/or at other regions, could serve as additional novel targets for therapies similar to the approach of using Herceptin to target HER2.

## SUPPORTING DATA

Figures 1-6  
Tables 1-2

### Figures

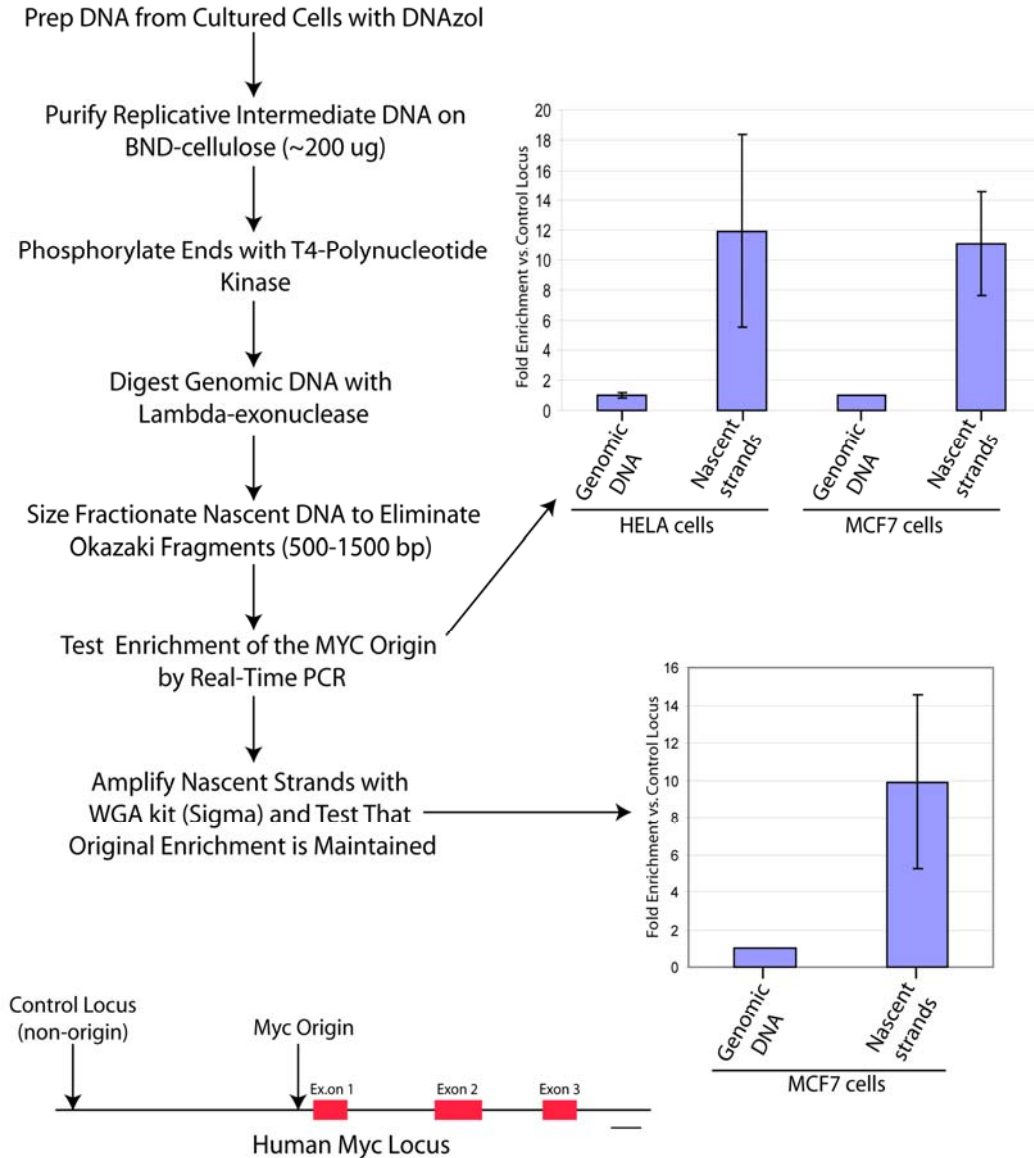
Figure 1:



**Figure 1** : Representative gel showing good shearing of genomic DNA from an ER+ breast tumor sample using the Bioruptor. The largest signal observed is in the range of 500 bp.

Figure 2:

### Flow Chart for Preparing Nascent DNA to Hybridize to Microarrays



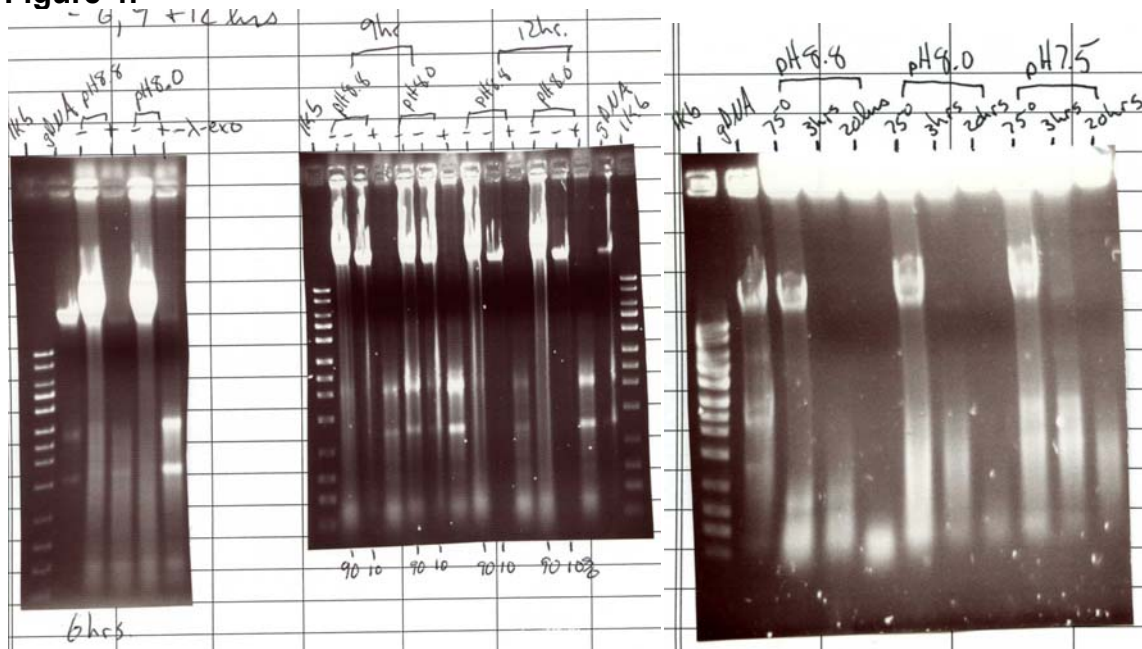
Adapted from Tao et al., *JCB* 78: 442-57 (2000).

**Figure 2:** Flow chart of the nascent strand preparation protocol. At the bottom is a cartoon of the c-myc locus. The arrows indicate the targets of the real-time PCR primers. At right is real-time PCR data showing enrichment of the c-myc origin in both HeLa and MCF7 cells (top) and the maintenance of enrichment after WGA amplification of ht nascent strands (bottom).

**Figure 3:**

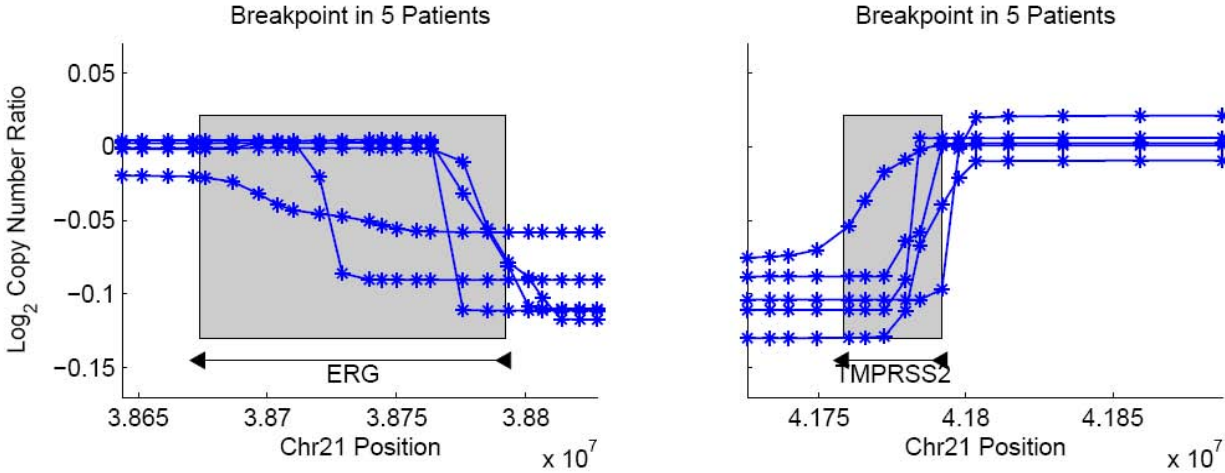
**Figure 3:** Chart of Illumina reads from MCF7 cell nascent strands mapped to the 283 origins identified by Cadoret et al., 2008. The red line indicates the cut off for statistical significance.

**Figure 4:**



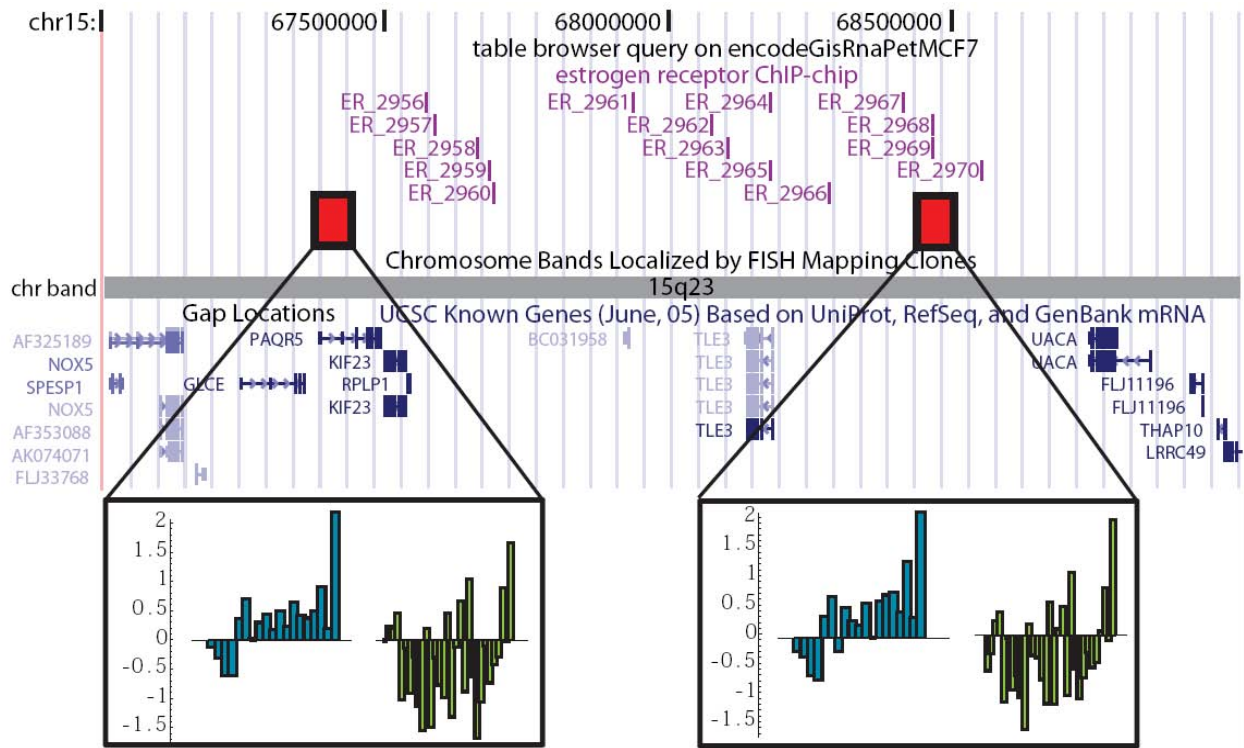
**Figure 4:** Gels from the notebook showing that the Fermentas lambda exonuclease is active at pH 8.0 and that the rRNA is more stable at this pH over the course of the digestion. 5 ug of DNA/RNA was digested at the indicated pH and for the time indicated. In the middle gel, the undigested sample was loaded in two adjacent lanes with one receiving 90% of the sample and the other the remaining 10%. The gel on the right shows that heating the sample to 75° C degrades the rRNA despite the pH (75° lanes).

**Figure 5:**



**Figure 5:** We identify the TMPRSS2-ERG fusion gene in 5 prostate cancer patients. The segmentations for each patient are shown in blue, and the asterisks denote probe locations. The deletion fuses the 5' end of TMPRSS2 to the 3' end of ERG, and the relative copy number at these breakpoints is conserved across the deletion.

**Figure 6:**



**Figure 6:** A statistically significant cluster of 15 ER binding sites (purple lines, top) identified on Chromosome 15.



**TABLE 1: Log of Breast Cancer Tumor Samples Received from R.I. Hospital**

All tumor specimens were provided by Dr. Shamlal Mangray (Pathology Department, RI Hospital) and were frozen at -80 degrees. All samples were from female patients (identity unknown – coded by the Pathology Department) without neoadjuvant chemotherapy. Most were patients of Dr. Thersa Graves. The samples were 1.0-1.5 cm.

<u># code</u>	<u>Date</u>	<u>ER</u>	<u>PR</u>	<u>HER2</u>	<u>Age</u>	<u>Comments</u>	
33	1	10/25/07	pos	used		for H4 test1	
34	2	11/13/07		also		normal tissue	
35	3	12/17/07	pos	pos	neg	also	normal tissue
36	4	1/18/08			post-menopausal(PM)	normal tissue (tube J)	
Path # 213A/J							
37 5							
38	6	1/18/08			45	used:C=H4 test (1 g)	
Path # 375C/F							
			also			normal tissue (tube F)	
39 7							
40	8	1/28/08	pos	pos	neg	60(PM)	also normal tissue
41 9							
42	10 (SG5)	1/28/08	pos	pos	neg	49	used: H4/ER/mock(0.2 g)
							also normal tissue
43 11							
44	12 (SG6)	1/28/08	pos	pos	neg	41	also normal tissue
3	SG8	9/09	3+	2+	Neg	73	8 cm tumor, lymph node mets
5	SG10 (5-10%)	9/09	3+	1+	Neg	28	4 cm tumor, sentinel lymph node (SLN) micromets
6	SG11	9/09	3+	3+	Neg	54	2.5 cm tumor, no mets to SLN
8	SG13	9/09	Neg	Neg	Neg	53	1.7 cm tumor,

								no mets to SLN
9	SG14	9/09	3+	3+	2+(FISH neg)	33		4 cm tumor no mets to SLN
10	SG15 No	9/09	3+	3+	2+(FISH neg)	66		2.4 cm tumor, SLN sampling, FNA of node negative
11	SG16	9/09	Neg	Neg	2+(FISH neg)	80		1.1 cm tumor, no mets to SLN
12	SG17 Axillary	9/09	3+	Neg	2+(FISH neg)	69		2.5 cm tumor, lymph node (ALN) negative
13	SG18	9/09	3+	3+	FISH neg	59		1.3 cm tumor, no mets to SLN
4	19	11/8/10	pos	pos	neg	also		normal tissue
5	20	11/8/10	pos	pos	neg	also		normal tissue
6	21	11/8/10	pos	pos	neg	also		normal tissue
7	22	11/8/10	pos	pos	neg	also		normal tissue
8	25	11/8/10	pos	pos	neg	also		normal tissue
9	26	11/8/10	pos	pos	neg	also		normal tissue
10	27	11/8/10	pos	pos	neg	also		normal tissue
11	28	11/8/10	pos	pos	neg	also		normal tissue
12	29	11/8/10	pos	pos	neg	also		normal tissue
13	30	11/8/10	pos	pos	neg	also		normal tissue
14	31	11/8/10	pos	pos	neg	also		normal tissue

**TABLE 2: Spreadsheet of MCF7 nascent DNA Illumina reads mapped to the ENCODE data set.**

	A	B	C	D	E	F	G	H	I
1	Origin - Chrom	Origin Start	Origin End	Num Reads	Reads/kb	pval (Poisson)	Corrected pval	Mapped	Mapped/kb
2	chr1 1	64131110	64132987 37		19.712310		0	2651049 0.94	680
3	chr1 1	64289674	64293005 75		22.51576	0	0		
4	chr1 1	64326897	64328689 35		19.53125	0	0		
5	chr1 1	64367486	64368648 34		29.25990	0	0		
6	chr1 3	112596565	112597931 27		19.76574	0	0		
7	chr1 6	154879	156172 22		17.01469	0	0		
8	chr1 6	271684	272766 25		23.10536	0	0		
9	chr1 6	343083	344285 27		22.46256	0	0		
10	chr1 6	353862	355393 26		16.98236	0	0		
11	chr2	220057705	220059150 27		18.68512	0	0		
12	chr2	220081216	220083396 37		16.97248	0	0		
13	chr2	220084625	220085841 29		23.84868	0	0		
14	chr2	220111158	220113002 39		21.14967	0	0		
15	chr2	220124753	220128235 81		23.26249	0	0		
16	chr2	220133214	220134642 25		17.50700	0	0		
17	chr2 1	39607655	39608789 24		21.16402	0	0		
18	chr5	131620957	131622317 53		38.97059	0	0		
19	chr6	4162251	4162406 46		29.7150	0	0		

		3 1		76				
20	chr9	1308139 41	1308161 02 37	17.121 70	0	0		
21	chr9	1308303 74	1308311 23 19	25.367 16	0	0		
22	chrX	1528895 95	1528907 89 38	31.825 80	0	0		
23	chrX	1533394 31	1533408 44 33	23.354 56	0	0		
24	chr1 6	265752	267214 23	15.731 87	0.00000000 0001	0.00000000 004		
25	chr2	2199899 90	2199915 33 24	15.554 12	0.00000000 0001	0.00000000 004		
26	chr2	2200485 83	2200516 56 47	15.294 50	0.00000000 0001	0.00000000 004		
27	chr2 0	3365297 3	3365411 6 18	15.748 03	0.00000000 0001	0.00000000 004		
28	chr6	1083854 66	1083867 97 20	15.026 30	0.00000000 0001	0.00000000 004		
29	chr7	2711991 5	2712143 9 24	15.748 03	0.00000000 0001	0.00000000 004		
30	chr9	1308904 21	1308918 63 22	15.256 59	0.00000000 0001	0.00000000 004		
31	chr9	1309104 97	1309150 36 70	15.421 90	0.00000000 0001	0.00000000 004		
32	chrX	1528535 10	1528547 17 19	15.741 51	0.00000000 0001	0.00000000 004		
33	chr1 1	6416729 6	6416897 1 24	14.328 36	0.00000000 002	0.00000000 06		
34	chr1 4	9876731 6	9876915 8 27	14.657 98	0.00000000 002	0.00000000 06		
35	chr1 6	47415 48	530 16	14.349 78	0.00000000 002	0.00000000 06		
36	chr1 9	5917491 3	5917871 8 55	14.454 66	0.00000000 002	0.00000000 06		
37	chr9	1309804 91	1309816 77 17	14.333 90	0.00000000 002	0.00000000 06		
38	chr1 4	9880831 6	9880954 3 17	13.854 93	0.00000000 03	0.00000000 9		
39	chr1 6	371926 3	73475 21	13.557 13	0.00000000 03	0.00000000 9		

40	chr2	2200183 96	2200200 76 22		13.095 24	0.00000000 03	0.00000000 9		
41	chr5	1318600 38	1318615 73 20		13.029 32	0.00000000 03	0.00000000 9		
42	chr1 1	6415478 2	6415672 6 24		12.345 68	0.00000000 5	0.0000001		
43	chr1 4	9878226 9	9878357 3 16		12.269 94	0.00000000 5	0.0000001		
44	chr1 8	2401172 5	2401289 1 14		12.006 86	0.00000000 5	0.0000001		
45	chr1 9	5906297 3	5906418 6 15		12.366 03	0.00000000 5	0.0000001		
46	chr6	4185459 6	4185569 7 14		12.715 71	0.00000000 5	0.0000001		
47	chr7	2711405 5	2711597 9 24		12.474 01	0.00000000 5	0.0000001		
48	chr7	2713506 8	2713651 4 18		12.448 13	0.00000000 5	0.0000001		
49	chr9	1311805 25	1311817 12 15		12.636 90	0.00000000 5	0.0000001		
50	chr1 1	6430203 5	6430380 8 20		11.280 32	0.00000006	0.000002		
51	chr7	2722534 7	2722705 0 20		11.743 98	0.00000006	0.000002		
52	chr7	9006354 2	9006491 6 16		11.644 83	0.00000006	0.000002		
53	chr9	1311387 58	1311402 99 18		11.680 73	0.00000006	0.000002		
54	chrX	1533260 81	1533272 18 13		11.433 60	0.00000006	0.000002		
55	chrX	1534167 20	1534181 74 16		11.004 13	0.00000006	0.000002		
56	chr1 1	6424950 6	6425146 7 20		10.198 88	0.00000007	0.000002		
57	chr2 0	3366722 6	3366896 2 19		10.944 70	0.00000007	0.000002		
58	chr7	2716040 4	2716151 0 12		10.849 91	0.00000007	0.000002		
59	chrX	1534280 85	1534294 58 14		10.196 65	0.00000007	0.000002		
60	chr1	1496978 53	1496998 82 19		9.3642 2	0.0000007	0.00020		

61	chr1 1	2148231	2149404	11	9.3776 6	0.0000007	0.00020		
62	chr1 1	6443852 7	6443994 8 14		9.8522 2	0.0000007	0.00020		
63	chr2 1	3929917 0	3930034 3 11		9.3776 6	0.0000007	0.00020		
64	chr6	4157844 7	4157971 9 12		9.4339 6	0.0000007	0.00020		
65	chrX	1528220 36	1528241 75 21		9.8176 7	0.0000007	0.00020		
66	chr1	1497502 98	1497516 52 12		8.8626 3	0.0000007	0.00196		
67	chr1 1	6416007 2	6416118 7 10		8.9686 1	0.0000007	0.00196		
68	chr1 9	5938544 5	5938723 4 16		8.9435 4	0.0000007	0.00196		
69	chr2	2201444 95	2201459 16 12		8.4447 6	0.0000007	0.00196		
70	chr2 1	3331464 9	3331691 0 19		8.4033 6	0.0000007	0.00196		
71	chr5	1320257 55	1320269 55 10		8.3333 3	0.0000007	0.00196		
72	chr6	4151882 6	4152040 9 14		8.8439 7	0.0000007	0.00196		
73	chr6	4185607 5	4185773 9 14		8.4134 6	0.0000007	0.00196		
74	chr7	2711614 6	2711801 9 15		8.0085 4	0.0000007	0.00196		
75	chr7	2724076 6	2724219 5 12		8.3974 8	0.0000007	0.00196		
76	chr9	1308385 87	1308402 06 13		8.0296 5	0.0000007	0.00196		
77	chrX	1529488 33	1529507 48 17		8.8772 8	0.0000007	0.00196		
78	chrX	1530161 22	1530176 75 13		8.3709 0	0.0000007	0.00196		
79	chrX	1532560 57	1532571 54 9		8.2041 9	0.0000007	0.00196		
80	chr1 1	6436863 1	6436988 4 10		7.9808 5	0.000006	0.01676		
81	chr1 1	6441233 7	6441399 9 13		7.8219 0	0.000006	0.01676		

82	chr1 3	1127529 96	1127542 03 9		7.4565 0	0.00006	0.01676		
83	chr1 4	9848627 8	9848775 7 11		7.4374 6	0.00006	0.01676		
84	chr1 6	176977	177951 7		7.1868 6	0.00006	0.01676		
85	chr1 6	391235	392389 9		7.7989 6	0.00006	0.01676		
86	chr2 2	3035513 5	3035625 5 8		7.1428 6	0.00006	0.01676		
87	chr6	4165412 0	4165575 0 12		7.3619 6	0.00006	0.01676		
88	chr7	1159514 28	1159529 38 12		7.9470 2	0.00006	0.01676		
89	chr9	1309425 15	1309456 15 22		7.0967 7	0.00006	0.01676		
90	chr9	1312141 69	1312164 97 17		7.3024 1	0.00006	0.01676		
91	chr1 3	1128112 87	1128125 98 9		6.8649 9	0.00045	0.12623		
92	chr1 4	9888185 9	9888320 2 9		6.7014 1	0.00045	0.12623		
93	chr1 6	144166	145603 9		6.2630 5	0.00045	0.12623		
94	chr1 9	5909347 0	5909564 7 15		6.8902 2	0.00045	0.12623		
95	chr1 9	5918659 0	5918779 0 8		6.6666 7	0.00045	0.12623		
96	chr1 9	5963361 4	5963465 6 7		6.7178 5	0.00045	0.12623		
97	chr2 1	3332165 8	3332345 7 11		6.1145 1	0.00045	0.12623		
98	chr2 1	3344575 2	3344685 9 7		6.3234 0	0.00045	0.12623		
99	chr2 2	3084905 1	3085026 5 8		6.5897 9	0.00045	0.12623		
100	chr2 2	3090151 6	3090271 5 8		6.6722 3	0.00045	0.12623		
101	chr6	7416071 3	7416199 2 8		6.2548 9	0.00045	0.12623		
102	chr6	1085474 74	1085487 73 9		6.9284 1	0.00045	0.12623		

103	chr7	27176097	271772728		6.80851	0.00045	0.12623		
104	chr9	131086093	13108805712		6.10998	0.00045	0.12623		
105	chrX	152794389	1527956608		6.29426	0.00045	0.12623		
106	chr1	149712058	1497137859		5.21135	0.00291	0.81993		
107	chr1	21272250	21282886	6	5.64440	0.00291	0.81993		
108	chr1	112469591	1124711448		5.15132	0.00291	0.81993		
109	chr1	112703946	11270584511		5.79252	0.00291	0.81993		
110	chr1	98724314	987256457		5.25920	0.00291	0.81993		
111	chr1	59206956	59208156	6	5	0.00291	0.81993		
112	chr5	56241199	562426258		5.61010	0.00291	0.81993		
113	chr7	27250014	272511766		5.16351	0.00291	0.81993		
114	chr7	114544682	1145457456		5.64440	0.00291	0.81993		
115	chr9	131153533	13115594914		5.79470	0.00291	0.81993		
116	chrX	122821773	1228228616		5.51471	0.00291	0.81993		
117	chrX	153213703	1532149597		5.57325	0.00291	0.81993		
118	chr1	130932556	1309336635		4.51671	0.01590	4.48334		
119	chr1	131047767	1310492307		4.78469	0.01590	4.48334		
120	chr1	112428205	1124296226		4.23430	0.01590	4.48334		
121	chr1	112530134	1125318638		4.62695	0.01590	4.48334		
122	chr1	112645963	1126471846		4.91400	0.01590	4.48334		
123	chr1	112668983	1126706268		4.86914	0.01590	4.48334		



124	chr19	59138068	591393376	4.72813	0.01590	4.48334		
125	chr19	59357776	593591076	4.50789	0.01590	4.48334		
126	chr266	2204241	2204258678	4.70312	0.01590	4.48334		
127	chr218	2343693	2343706876	4.38276	0.01590	4.48334		
128	chr210	3369762	336993228	4.70035	0.01590	4.48334		
129	chr221	3080796	308097107	4.00229	0.01590	4.48334		
130	chr539	1316362	1316374485	4.13565	0.01590	4.48334		
131	chr549	1420782	14208117614	4.78305	0.01590	4.48334		
132	chr64	4159980	416011726	4.38596	0.01590	4.48334		
133	chr75	8971253	897137446	4.96278	0.01590	4.48334		
134	chr787	1159267	1159287809	4.51581	0.01590	4.48334		
135	chr773	1159533	1159546466	4.71328	0.01590	4.48334		
136	chr768	1167502	1167515106	4.83092	0.01590	4.48334		
137	chr719	1267744	1267757756	4.42478	0.01590	4.48334		
138	chr752	1268198	1268213447	4.69169	0.01590	4.48334		
139	chrX89	1528365	1528377355	4.36300	0.01590	4.48334		
140	chrX02	1532027	1532038485	4.36300	0.01590	4.48334		
141	chr184	1498136	1498149154	3.24939	0.07078	19.96030		
142	chr1395	1127512	1127527165	3.51865	0.07078	19.96030		
143	chr195	5910427	591066338	3.39271	0.07078	19.96030		
144	chr216	3377390	337754266	3.94737	0.07078	19.96030		

14	chr2	3393714	3393873		3.1348				
5	1	1	6 5		0	0.07078	19.96030		
14	chr2	3034187	3034283		3.1348				
6	2	5	2 3		0	0.07078	19.96030		
14	chr2	3120077	3120281		3.9177				
7	2	5	7 8		3	0.07078	19.96030		
14		5614752	5614906		3.2573				
8	chr5	5	0 5		3	0.07078	19.96030		
14		1421762	1421781		3.5696				
9	chr5	13	74 7		1	0.07078	19.96030		
15		4154306	4154434		3.9001				
0	chr6	0	2 5		6	0.07078	19.96030		
15		4154494	4154602		3.7037				
1	chr6	2	2 4		0	0.07078	19.96030		
15		4158509	4158673		3.6674				
2	chr6	8	4 6		8	0.07078	19.96030		
15		4171917	4172014		3.0959				
3	chr6	8	7 3		8	0.07078	19.96030		
15		7428752	7428863		3.6003				
4	chr6	5	6 4		6	0.07078	19.96030		
15		2722201	2722331		3.0792				
5	chr7	6	5 4		9	0.07078	19.96030		
15		1162900	1162913		3.8197				
6	chr7	06	15 5		1	0.07078	19.96030		
15		1163810	1163824		3.6791				
7	chr7	43	02 5		8	0.07078	19.96030		
15		1270160	1270172		3.4275				
8	chr7	91	58 4		9	0.07078	19.96030		
15		1309828	1309843		3.1545				
9	chr9	08	93 5		7	0.07078	19.96030		
16	chr1	1123952	1123970		2.2296				
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16	chr1	1125614	1125627		2.2338				
1	3	43	86 3		0	0.24468	68.99993		
16	chr1	1126063	1126075		2.4650				
2	3	07	24 3		8	0.24468	68.99993		
16	chr1	1126845	1126863		2.2123				
3	3	21	29 4		9	0.24468	68.99993		
16	chr1	9869625	9869798		2.9019				
4	4	8	1 5		2	0.24468	68.99993		
16	chr1	5907789	5908036		2.4340				
5	9	7	2 6		8	0.24468	68.99993		

166	chr1 9	5934000 3	5934136 8 3		2.1978 0	0.24468	68.99993		
167	chr2	2200715 68	2200741 68 7		2.6923 1	0.24468	68.99993		
168	chr2	2345541 79	2345555 16 3		2.2438 3	0.24468	68.99993		
169	chr2	3360464 0 7	3360627 9 4		2.4509 8	0.24468	68.99993		
170	chr2	3384729 1 4	3384917 3 4		2.1287 9	0.24468	68.99993		
171	chr5	5608806 6	5608947 8 3		2.1246 5	0.24468	68.99993		
172	chr5	1315718 51	1315736 44 5		2.7886 2	0.24468	68.99993		
173	chr5	1318011 34	1318028 91 5		2.8457 6	0.24468	68.99993		
174	chr5	1320481 84	1320494 26 3		2.4154 6	0.24468	68.99993		
175	chr5	1422362 58	1422372 49 2		2.0181 6	0.24468	68.99993		
176	chr7	2713713 6	2713833 0 3		2.5125 6	0.24468	68.99993		
177	chr8	1192501 80	1192514 04 3		2.4509 8	0.24468	68.99993		
178	chrX	1227262 13	1227275 33 3		2.2727 3	0.24468	68.99993		
179	chrX	1229231 20	1229244 70 4		2.9629 6	0.24468	68.99993		
180	chrX	1532624 07	1532638 80 3		2.0366 6	0.24468	68.99993		
181	chr1	2989390 3 5	2989541 3 3		1.9893 9	0.61202	172.58983		
182	chr1	5298863 4 3	5299017 1 2		1.3003 9	0.61202	172.58983		
183	chr1	128359 6	129221 1		1.1600 9	0.61202	172.58983		
184	chr1	458042 4	459289 2		1.6038 5	0.61202	172.58983		
185	chr1	5906535 9 8	5906716 9 3		1.6565 4	0.61202	172.58983		
186	chr1	5915533 9 0	5915717 3 3		1.6277 8	0.61202	172.58983		

187	chr20	33520581	335216192		1.92678	0.61202	172.58983		
188	chr20	33593121	335947223		1.87383	0.61202	172.58983		
189	chr21	32736476	327381372		1.20409	0.61202	172.58983		
190	chr21	33143912	331456123		1.76471	0.61202	172.58983		
191	chr21	39377239	393785012		1.58479	0.61202	172.58983		
192	chr22	31035855	310368602		1.99005	0.61202	172.58983		
193	chr22	31371562	313731262		1.27877	0.61202	172.58983		
194	chr5	55873281	558743782		1.82315	0.61202	172.58983		
195	chr5	141999954	1420015052		1.28949	0.61202	172.58983		
196	chr5	142361833	1423637802		1.02722	0.61202	172.58983		
197	chr6	74108528	741103973		1.60514	0.61202	172.58983		
198	chr6	108711089	1087122492		1.72414	0.61202	172.58983		
199	chr7	89870825	898724262		1.24922	0.61202	172.58983		
200	chr7	116591693	1165934042		1.16891	0.61202	172.58983		
201	chr7	116906722	1169087864		1.93798	0.61202	172.58983		
202	chr7	117218481	1172196542		1.70503	0.61202	172.58983		
203	chr8	119302424	1193038202		1.43266	0.61202	172.58983		
204	chr9	130824308	1308258492		1.29786	0.61202	172.58983		
205	chr9	131164410	1311653861		1.02459	0.61202	172.58983		
206	chrX	122763000	1227641462		1.74520	0.61202	172.58983		
207	chr1	149438460	149440397	0	0	1	282		

208	chr1	1715184	1716695	1	0.6618	1	1	282		
209	chr1	130855483	130856606	1	0.89047	1	1	282		
210	chr1	38937995	38939318	1	0.75586	1	1	282		
211	chr1	29850448	29851781	1	0.75019	1	1	282		
212	chr1	112433768	112434909	1	0.87642	1	1	282		
213	chr1	112602511	112604575	2	0.96899	1	1	282		
214	chr1	112612988	112614632	1	0.60827	1	1	282		
215	chr1	98771506	98772663	1	0.86430	1	1	282		
216	chr1	98956703	98958083	1	0.72464	1	1	282		
217	chr1	41735285	41736157	0	0	1	1	282		
218	chr1	41850700	41852158	0	0	1	1	282		
219	chr1	42013413	42014920	1	0.66357	1	1	282		
220	chr1	60988052	60989443	0	0	1	1	282		
221	chr1	24064501	24065770	0	0	1	1	282		
222	chr1	59225201	59226338	1	0.87951	1	1	282		
223	chr1	59588883	59590167	1	0.77882	1	1	282		
224	chr2	51885784	51887168	1	0.72254	1	1	282		
225	chr2	118506164	118507311	0	0	1	1	282		
226	chr2	220218828	220219997	1	0.85543	1	1	282		
227	chr2	220221572	220222989	1	0.70572	1	1	282		
228	chr2	234539246	234540367	1	0.89206	1	1	282		

229	chr2	234549286	234550644	0	0	1	282		
230	chr2	234623194	234624340	0	0	1	282		
231	chr2	334493709	334510621		0.59418	1	282		
232	chr2	337635206	337647781		0.79872	1	282		
233	chr2	328258313	328272101		0.72622	1	282		
234	chr2	329248515	32926140	0	0	1	282		
235	chr2	333711916	333725231		0.75358	1	282		
236	chr2	338695114	338708301		0.75988	1	282		
237	chr2	303197329	30320501	0	0	1	282		
238	chr2	31162231	31163655	0	0	1	282		
239	chr2	314429426	31444794	0	0	1	282		
240	chr2	315361529	31537121	0	0	1	282		
241	chr2	315504512	315516471		0.83612	1	282		
242	chr5	55981348	559828311		0.67431	1	282		
243	chr5	56085418	56086517	0	0	1	282		
244	chr5	56119168	56120680	0	0	1	282		
245	chr5	131744261	1317455221		0.79302	1	282		
246	chr5	132151529	1321526591		0.88496	1	282		
247	chr5	132248525	1322506281		0.47551	1	282		
248	chr5	141950090	141951462	0	0	1	282		
249	chr5	142199652	1422013761		0.58005	1	282		

250	chr5	142259552	142261293	0	0	1	282		
251	chr6	74011080	74012121	0	0	1	282		
252	chr6	108552017	108553295	0	0	1	282		
253	chr6	132399844	132401367	1	0.65660	1	282		
254	chr7	27145830	27147560	1	0.57803	1	282		
255	chr7	89671933	89673120	1	0.84246	1	282		
256	chr7	89716391	89718325	0	0	1	282		
257	chr7	90230960	90232415	0	0	1	282		
258	chr7	90283033	90284598	1	0.63898	1	282		
259	chr7	90434126	90435342	1	0.82237	1	282		
260	chr7	114067204	114068337	0	0	1	282		
261	chr7	114309758	114311461	1	0.58720	1	282		
262	chr7	114511851	114512914	1	0.94073	1	282		
263	chr7	115921833	115923372	0	0	1	282		
264	chr7	115932826	115934434	0	0	1	282		
265	chr7	115985825	115987901	2	0.96339	1	282		
266	chr7	116126066	116128112	2	0.97752	1	282		
267	chr7	116223147	116226022	0	0	1	282		
268	chr7	116494825	116495804	0	0	1	282		
269	chr7	116606707	116607969	0	0	1	282		
270	chr7	116618519	116619873	0	0	1	282		

27 1	chr7	1169141 62	1169156 47 1		0.6734 0	1	282		
27 2	chr7	1169925 50	1169937 26	0	0	1	282		
27 3	chr7	1262729 23	1262741 35	0	0	1	282		
27 4	chr7	1263061 77	1263068 34	0	0	1	282		
27 5	chr7	1264407 57	1264421 09	0	0	1	282		
27 6	chr7	1268922 41	1268933 57	0	0	1	282		
27 7	chr8	1190124 20	1190138 04	0	0	1	282		
27 8	chr8	1193743 75	1193754 34 1		0.9442 9	1	282		
27 9	chr9	1310006 00	1310017 95 1		0.8368 2	1	282		
28 0	chrX	1226621 45	1226635 18	0	0	1	282		
28 1	chrX	1226930 12	1226942 69 1		0.7955 4	1	282		
28 2	chrX	1230616 67	1230640 36	0	0	1	282		
28 3	chrX	1539083 51	1539093 90	0	0	1	282		