CANDIDATE GENE IN PREDICTING IN VIVO OVARIAN CANCER RESPONSE TO COMBINATION THERAPY WITH PARAPLATIN AND PACLITAXEL

Ramin Mirhashemi¹, J. Fernando Arena¹, Tony Frudakis², Nicholas Lambrou¹, Jane Arboleda¹, Marsha Hunt¹, Maria Medranda¹, Hervy Averette¹, and Manuel Penalver¹

¹Department of Obstetric & Gynecology, University of Miami School of Medicine, P.O. Box 016960, Miami, FL 33136; ²DNAPrint Genomics, Inc., 900 Cocoanut Ave., Sarasota, FL 34236
*rmirhashemi@med.miami.edu

INTRODUCTION. Adverse drug reactions, due at least in part to interindividual variability in drug response, rank between the 4th and 6th leading causes of death in the U.S. The field of “pharmacogenetics” (the study of variability in drug response due to heredity), should help in reducing drug-caused morbidity and mortality[1]. The underlying genetics for the efficacy of the chemotherapeutic treatment of ovarian cancer is still unknown and subject to a high degree of uncertainty and interindividual variability[2]. In the present project, using a pharmacogenomic approach, we evaluate the patient response rates to primary chemotherapeutic treatment of ovarian cancer with combination of paclitaxel and the carboplatin agents. In an attempt to define the genetic determinants for this variability, we are conducting a pharmacogenomics study to predict response based on patient genomics[3].

Our results demonstrate that single nucleotide polymorphic (SNPs) in candidate genes being identified could potentially be used pre-treatment to individualize chemotherapy for patients with ovarian cancer.

METHOD. Our approach is a systematic haplotype-based candidate gene approach, where we genotype each patient at each SNP site for each gene known to be relevant for xenobiotic disposition, then computationally infer haplotypes, and geometrically model the data in order to identify associations between haplotypes and response. We have chosen this novel approach to objectively define an optimal pharmacogenomics solution.

RESULTS. We have constructed SNP maps for 50 genes known to be involved in xenobiotic metabolism and/or the disposition of paclitaxel and carboplatin and have discovered an average of 30 candidate SNPs per gene, which, upon polymorphism screening, validated at a rate of about 18 SNPs per gene. Having obtained genotypes for 57 patients at each of the SNPs in several genes, preliminary results have revealed a statistically significant association (Fishers Exact $p = 0.00332 \pm 0.00043$) between haplotypes at the CYP3A4 gene and a lack of response to paclitaxel/carboplatin as measured by the CA-125 tumor marker. In particular, the CGC and CGT CYP3A4 haplotypes were only present in the nonresponder group ($n = 22$ individuals, 106 readings), when a 50% decrease in CA-125 was used to define responders ($n = 9$ individuals, 37 readings). When a 20% decrease in CA-125 was used, a similar result was obtained (Fishers Exact...
\( p = 0.016 \pm 0.001 \); CGC was found in the nonresponder group 75% of the time, and the CGT haplotype was found in the nonresponder group 100% of the time.

**DISCUSSION.** In the present study, we have investigated SNPs in 50 candidate genes involved in the xenobiotic metabolism/disposition of paclitaxel and carboplatin[4]. We identified and validated SNPs in a reference population, and screened them for the ability to resolve between paclitaxel and carboplatin responders and nonresponders. We have observed 50% reductions in CA-125 tumor marker values are associated with two haplotypes (CGC and CGT) in the CYP3A4 gene. Even when a 20% decrease in CA-125 was used as the criteria for response grouping, rather than a 50% decrease, the two haplotypes were associated with nonresponse. The same results were obtained whether average response per patient was used or whether individual readings were grouped. Patient accrual is still in process and no similar data was found in the literature for comparison. This *in vivo* assay could represent a component of a classifier that could help classify ovarian cancer patients as potential responders or nonresponders to our conventional chemotherapeutic agents.

**REFERENCES**

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