Identification and Characterization of Splice Variants in BRCA Genes Using Reverse Transciptase Polymerase Chain Reaction and Multiplex Ligation-Dependent Probe Amplification

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**Objectives:** A large subset of the unclassified variants in the BRCA1/2 genes are in intronic sequences or close to exon–intron boundaries and may affect splicing. Putative BRCA1 and BRCA2 splice variants, as predicted in silico, were studied experimentally using reverse transcriptase polymerase chain reaction (RT-PCR) and a new approach based on multiplex ligation-dependent probe amplification (MLPA) technology.

**Methods:** Short-term lymphocyte cultures from controls and unclassified sequence variant carriers were established with PHA and IL-2. Analysis by RT-PCR was performed for the following BRCA2 putative splice variants: c.425G>T, c.6935A>T, c.6842_6937del (exon 12 skipping) and c.476_631del (exon 6 and 7 skipping). Because these in-frame transcripts are also expressed in healthy controls and have an unknown function, their clinical relevance remains unclear. This has led to 46 bp of intron retention. These variants were considered to be pathogenic. Additionally, alternative splicing events were observed in controls, such as c.6842_6937del (exon 12 skipping) and c.476_631del (exon 6 and 7 skipping). Because these in-frame transcripts are also expressed in healthy controls and have an unknown function, their clinical relevance remains unclear. This has consequences for the classification of c.6935A>T, which leads to increased expression of c.6842_6937del compared with controls. We found that, with MLPA, it is also possible to detect alternative splicing events.

**Conclusions:** We characterized several new pathogenic BRCA2 splice variants and demonstrated that, with MLPA technology, it is possible to set up a multiplex screen for semi-quantitative detection of alternative splice events that is also widely applicable for genes other than BRCA1/2.
Implementation of High Throughput Parallel Sequencing in a Diagnostic Setting: Multiplexed Amplicon Sequencing of the Breast Cancer Genes BRCA1 and 2

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“Next generation” sequencing opens new perspectives for mutation analysis of large genes. We optimized a multi-sample barcode amplicon sequencing strategy for mutation detection in BRCA1 and 2 on the GS-FLX instrument (Roche/454 Life Sciences).

For the first two runs (POP1 and 2), 112 amplicons covering the complete coding sequence of BRCA1 and 2 were pooled in equimolar amounts. Per run, 11 patients were sequenced in both directions for all amplicons. For a third run (POP3), all 112 amplicons were generated in 15 multiplex polymerase chain reaction (PCR) reactions. Hereby, all PCR reactions were optimized to obtain a final equimolar mix for all fragments (CV < 30%). The sequence coverage was much more uniform in POP3 than in POP1 and POP2, indicating a successful optimization and showing that the bias introduced by multiplexing and pooling was limited. In POP3, less than 40% coverage was obtained only for 6 amplicons. All sequence reads for the 22 patients of POP1 and POP2 were analyzed with our in-house variant identification pipeline (Deschrijver et al., submitted) using the empirically determined filters (minimum coverage: 40; maximum homopolymer length: 7; minimum variant frequency: 33%). These results were compared with data obtained by Sanger sequencing, producing 0 false negatives and 5 false positives. The filter settings applied for POP1 and POP2 are currently validated for POP3.

This approach is not yet cost-competitive with the high-resolution melting curve analysis (De Leeneer et al., 2008, 2009) currently applied in our laboratory, but it outperforms traditional Sanger sequencing. Further optimization of sequence coverage uniformity, pooling of patients, and Titanium throughput will allow further cost reduction. For the optimization of next-generation amplicon sequencing, a strong collaboration is required between people involved in molecular diagnostics and bio-informaticians to obtain clinically reliable results.

Prevalence of BRCA1/2 Mutations in Unselected and Selected French Canadian Breast Cancer Cases Diagnosed at the CRCHUM

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Objectives: The main purpose of this study was to estimate the prevalence of BRCA gene mutations and the yield of genetic testing among unselected breast cancer patients less than 50 years of age compared with selected patients between 51 and 65 years of age (with a family history of 2 or more breast or ovarian cancers).

Methods: In an open-cohort clinic at the CHUM in which French-Canadian breast cancer patients from 2005–2009 were recruited, we studied 932 subjects (687 < 50 years of age and 245 > 50 years of age). All subjects were tested for BRCA gene mutations.

Results: Of the unselected women with breast cancer (<50 years of age), 5.0% were BRCA mutation carriers (1.2% BRCA1, 3.8% BRCA2). In those with a family history (selected subjects or >50 years of age), 4.9% displayed BRCA mutations (0.8% BRCA1, 4.1% BRCA2). Among unselected patients, the refusal rates for the genetic tests were 10.2% and 14.9% among selected breast cancer patients. Of the study subjects, 99% agreed to be told the results of their genetic tests; the remaining 1% refused to be told their results.

Conclusions: The results of this open-cohort study of the genetic epidemiology of breast cancer showed that the prevalence of BRCA mutations among French-Canadian breast cancer patients is around 5.0% (5.0% in unselected or <50-year-old patients, 4.9% in selected or older subjects). This type of study is feasible for genetic screening and the refusal rate for participation in a genetic study of breast cancer is very low.

The Establishment of a Cohort of Girls in Ontario Breast Cancer Families

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Background: Emerging evidence indicates that childhood and adolescence may be periods sensitive to factors associated with breast cancer initiation or progression. We initiated a study to determine predictive and protective factors against breast cancer development. To investigate factors related to breast cancer risk, we are planning a study of girls.

Methods: To determine the interest of parents and daughters for participating in the Legacy—Youth study plans to examine several intermediate phenotypes associated with breast cancer, including pubertal breast development, breast tissue characteristics, age at menarche, sex and growth exposures, as well as genetic determinants.

Objectives: To determine the interest of parents and daughters for participating in the Legacy—Youth study and to aid in its design, we are conducting a pilot study to evaluate the feasibility of recruitment, biopspecimen, and data collection in 100 girls; to determine whether, as compared with Tanner stage, optical spectroscopy can be used as an efficient measure of breast development in the girls; and to define parental and offspring levels of acceptance or dissatisfaction across data collection measures.

Methods: Parents and their 6- to 17-year-old daughters from two high-risk cancer genetics clinics in the Greater Toronto Area and those already enrolled in the Ontario Familial Breast Cancer Registry, a U.S. National Cancer Institute–supported population-based registry, will be invited to participate. The pilot involves the completion of self- and interview-administered epidemiologic questionnaires for daughter and parent, clinic-administered measures of height and weight; blood, buccal, and urine sampling; measures of pubertal breast development using self-Tanner staging and experimental optical spectroscopy modalities; and qualitative exit interviews for thorough exploration of participant likes and dislikes.

Results: We will present recruitment outcomes from our first group of girls.

Conclusions: Completion of the pilot study will allow us to demonstrate feasibility, to estimate response rates, and to assess the interest of families in participating in Legacy—Youth.

Mutations in Belgian Families with Breast and/or Ovarian Cancers

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Germline mutations in BRCA1 and BRCA2 genes cause predisposition to breast and ovarian cancers. Here, we describe the experience of our Human Genetics Center on a total of approximately 700 patients living in the southern part of Belgium. These patients, apparently from independent families, were referred by oncologists or by geneticists. All of them had a personal and/or familial history suggestive of genetic predisposition to breast and/or ovarian cancer.

BRCA1 and BRCA2 genes were screened by standard scanning methods (denaturing gradient gel electrophoresis, denaturing high-pressure liquid chromatography, or high resolution melt), followed by sequencing of abnormal profiles. Large rearrangements of the two genes were investigated by MLPA (multiplex ligation-dependent probe amplification from MRC-Holland).

A total of 186 variants (excluding known polymorphisms) were identified (106 in BRCA1, 80 in BRCA2). This represents a detection rate of 26.5%.

In BRCA1, 26 (24.5%) were variants of uncertain significance, and 80 (75.5%) were clear mutations, of which 13 were large rearrangements of the gene. Four recurrent mutations (deletion from exons 8 to 13, c.2772C>T, c.4391_4393delinsTT, and c.5137delc) account for 27.5% (respectively 7.5%, 7.5%, 6.25%, and 6.25%) of the mutations. In BRCA2, 37 (46%) were variants of uncertain significance, and 43 (54%) were clear mutations, of which only 1 was a large rearrangement of the gene. Two recurrent mutations (c.9117G>A and c.4936_4939delc) account for 30% (respectively 16.3% and 14%) of the mutations.

A causal mutation has hence been identified in a total of 17.5% of the patients (11.4% in BRCA1, 6.1% in BRCA2). These results underline the role of large rearrangements of BRCA1 (16.25% of the mutations identified in this gene) in Belgium and point to the need for collaborative efforts to classify precisely variants of uncertain significance, because they represent 33.8% of all variants detected.
Breast and Ovarian Cancer Risk Among Heterozygotes for Xeroderma Pigmentosum Nucleotide Excision Repair Gene Mutations

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**Background:** Xeroderma pigmentosum (XP) is a rare autosomal recessive disorder with an estimated frequency of 10–6 in North America. The condition is characterized by severe photosensitivity, abnormal pigmentation, and an increase, by a factor of 1000 or more, in the risk of cancers in sun-exposed tissues (skin and eye) in affected individuals. The disorder is caused by mutations in a number of genes, all (XP-A–XP-G) but one (XP-V) variant involved in the nucleotide excision repair (NER) pathway. The heterozygote (clinically normal-appearing carrier) frequency of XP gene mutations is estimated at about 1 in 300. Cancer risk of heterozygotes is not known. We report on the frequency of hereditary breast cancer among XP families.

**Methods:** The study population consists of 22 XP families with detailed multi-generation pedigrees and epidemiology information ascertained under a protocol to study the natural history of the disorder. Hereditary breast cancer was defined as the occurrence of at least 2 cases of premenopausal breast cancer (diagnosed <50 years of age) or 1 case of premenopausal breast cancer and 1 case of ovarian cancer in one side of the family.

**Results:** The family histories of 5 of the 22 XP patients were compatible with hereditary breast cancer. In 3 families with XP, in both affected individuals with XP, there were 4 cases (including the heterozygote mother of the proband) of hereditary breast cancer, 1 case of ovarian cancer, and 1 case of prostate cancer in the maternal side. Comprehensive sequencing failed to identify a mutation in the BRCA1 and BRCA2 genes in this family. Overall, a family history compatible with hereditary breast cancer was found in approximately 23% of XP families reported here.

**Conclusions:** Heterozygote carriers of XP gene mutations may be at increased risk for breast and ovarian cancer. A proportion of breast and ovarian cancer cases in the population may be attributed to mutations and polymorphisms in the NER genes.

Improvement in the Detection Rate of the BRCA1/BRCA2 French-Canadian Mutation Panel at the Service de Médecine Génétique, CHUM S. Nolet, K. Latour, S. Côté, N. Dumas, M.C. Binet, M. Breuguet, L. Gabor, P. Hamet, N. Larouche, G. Proulx, D. Provencher, J. Richard, C. Wilmar, C. Maugard. Centre hospitalier de l’Université de Montréal (CHUM), Montreal, Quebec, Canada.

**Introduction:** The genetics of the French-Canadian (FC) population of Quebec is shaped by well-documented founder effects that have allowed for the identification of common mutations in specific genes. In BRCA1 and BRCA2, hereditary breast and ovarian cancer, several mutations are part of a FC panel which is used as a first line of testing.

**Objectives:** To evaluate the detection rate of the FC panel currently used in the CHUM diagnostic laboratory. To determine the change in detection rate when mutations, reported in at least two distinct FC families, are added to the panel.

**Methods:** A retrospective review of results obtained for FC families tested through the Hereditary Cancer Clinic using the laboratory and clinical records. Included in our analysis were index cases for whom the affected branch of the family was of FC descent, for whom a whole blood sample could be obtained, and who had both panel testing and sequencing if panel testing was negative (n = 301).

**Results:** From January 2005 to December 2008, 408 cases of FC descent were screened for mutations (R1443X, 2953del3+C, 3768insA, W3216insa, 6503delTT for BRCA1; 8735delaG, E1953X, 3398del5, 2816insa, 6503delTT for BRCA2). Of the 301 index cases corresponding to our inclusion criteria, 69 were identified as mutation carriers. Of these 69 carriers, 57 were found to carry a mutation from the FC panel (82.6%). The remaining 12 had a mutation identified by sequencing (17.4%). 5 of these mutations (2080insA, 2244insA, 2800del1A for BRCA1; 3773delTT, R2336H for BRCA2) were reported in at least 2 distinct families. By adding these mutations to our in-house panel, our detection rate will reach 97.1% (67/69).

**Conclusions:** This retrospective review of our laboratory data indicates that the detection rate of our in-house FC panel of 9 mutations is 82.6%. By adding 5 mutations, this detection rate will reach 97.1%. These preliminary data indicate that testing using the FC panel in appropriate families continues to be a cost-effective initial screening. The data are insufficient to preclude further testing by sequencing in panel-negative families. A periodic review of mutations identified by sequencing in FC families is important to improve and validate the efficiency of the FC panel.

The Characterization of Genomic Instability in Lymphoblastoid Cell Lines Derived from Heterozygous PALB2 Mutation Carriers

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**Objectives:** PALB2, upon interaction with its nuclear partner, the breast cancer gene, BRCA2, is known to promote localization and stability. PALB2 is estimated to confer a moderately increased risk for breast cancer, likely because of a disruption of genomic regulation. To date, the underlying pathogenesis of PALB2 mutations has yet to be illuminated by common molecular mechanisms. It has previously been suggested that Fanconi anemia proteins, a group of which PALB2 is a member, may play a critical role in the telomere maintenance pathway, because of their core function in genetic recombination. In the current study, we attempt to establish both a molecular and cellular link between PALB2 heterozygous mutations and genomic instability, an important precursor to carcinogenesis.

**Methods:** Six lymphoblastoid cell lines (l.l.s) were analyzed in the current study, two expressing wild-type PALB2, and four carrying different mutation PALB2 truncating mutations: 2521delA, 3323delA, Q775X, and 229delIT. Genomic integrity was assessed by telomeric quantitative fluorescence in situ hybridization (q-fish). Further investigations of genomic abnormalities were conducted through the use of centromeric ratio and spectral karyotyping. The response to cellular cytotoxicity was examined in the presence of mitomycin C, bleomycin, and cisplatin. Cell survival was measured via photo spectrophotometry through the metabolism of the tetrazolium salt WST-1.

**Results:** No statistical difference was observed in telomere number or median telomeric intensity between the two PALB2 control lines (C1 and C2; p = 0.287). Similarly no significant differences were observed in telomere number or intensity between C1/C2 and the 3323delA carrier (p = 0.386 vs. C1 and p = 0.786 vs. C2). Probe intensity was markedly reduced in the 2521delA carrier as compared with both C1 and C2. The Q775X carrier was observed with a reduced telomere count and differed significantly from either control with respect to mean probe intensity (p = 2.22 × 10^-3 vs. C1 and p = 8.52 × 10^-3 vs. C2). Similarly, the 229delIT carrier was observed with an increased telomere count and significantly differed in mean probe intensity (p = 6.05 × 10^-3 vs. C1 and p = 1.77 × 10^-2 vs. C2). No significant differences were observed in centromere number throughout all six l.l.s. No duplications, translocations, deletions, or rearrangements were observed to be consistent with the pathogenicity of the PALB2 variants. No l.c.s showed any significantly altered response to any of the cytotoxic agents.

**Conclusions:** The current study may suggest a possible association between PALB2 heterozygous truncating mutations and impaired genomic regulation. However, a new model system for analysis, such as a PALB2 tumour cell line, is required to further elucidate this potential association.
Mutations in PALB2 have been reported to be associated with a doubled to tripled increased risk of breast cancer. Women with mutations in PALB2 who also have a strong family history of breast cancer are predicted to be at a high absolute risk for the disease comparable to that in women who carry BRCA2 mutations. A current paucity of information about PALB2 mutation carrier risks prevents testing for mutations in this gene being incorporated into evidenced-based clinical genetics services.

**Objectives:** To conduct a large case–control family study using population-based and clinic-based resources, to characterize PALB2 mutations in terms of breast cancer risk, and to devise criteria to identify women most likely to carry PALB2 mutations.

**Methods:** High-resolution melt curve analysis and Sanger sequencing.

**Results:** To date, we have screened 736 early-onset population-based cases for PALB2 mutations and identified 2 carriers, both in women with extremely strong family histories of breast cancer. We have also screened 400 women from multiple-case breast cancer families and identified 4 families that carry PALB2 mutations. Testing for these mutations in relatives has identified an additional 20 PALB2 mutation carriers. Breast cancers arising in affected female carriers are commonly grade 3 (7/12 reviewed to date) and estrogen receptor–positive (7/19 reviewed to date), 9 are infiltrating ductal carcinomas, and 3 are lobular or pleomorphic lobular carcinomas.

**Conclusions:** We and others have produced preliminary and circumstantial evidence that women who carry a mutation in PALB2 are at an elevated risk of breast cancer, and that depending on family history, these risks could be of clinical relevance comparable to that for women with BRCA2 mutations. PALB2 mutations are rare, but for the women who carry these mutations and their families, it is potentially important that mutations be identified so that the women can be offered appropriate prevention, screening, and clinical management.
Identification of a BRCA2 Founder Mutation in the Manitoba Mennonite Population

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The Mennonites are a distinct ethnic and religious group characterized by their Anabaptist beliefs and pacifism. Over the last few centuries, persecution forced this group to migrate throughout Northern and Eastern Europe, and then to Southern Manitoba (Canada), the American Midwest, Mexico, and Central and South America. There are about 66,000 Mennonites residing in the Canadian province of Manitoba, accounting for approximately 6% of its total population. Founder mutations have been identified in the Mennonite population for a number of genetic diseases, including hypophosphatasia, X-linked congenital stationary night blindness, and familial hypercholesterolemia. We have identified a mutation, c.5238dupT (p.Asn1746Tyr), in the BRCA2 gene in 4 unrelated Mennonite families. The index patients were referred for genetic testing of BRCA1 and BRCA2 genes because of personal and family histories of breast and ovarian cancer. One family also had a history of pancreatic cancer. Of the 4 families, 3 were long-time residents of Manitoba; 1 family had recently emigrated from Bolivia. Further investigations into the pedigrees did not reveal any common relatives between the families. These mutations were independently identified by us sequencing following the observation of a truncated band using the protein truncation test targeted to exon 11 of BRCA2. The c.5238dupT mutation has been reported only a limited number of times in the Breast Cancer Information Core database and in the literature, and it has been restricted to patients of Dutch, German, Mennonite, and Western and Eastern European descent. The finding of 4 large pedigrees of Mennonite descent indicates the presence of a founder mutation in the Manitoba Mennonite population. The population frequency of this BRCA2 c.5238dupT mutation among Manitoba Mennonites is not known, but it may be of value to offer targeted mutation analysis to individuals of Mennonite descent who have a personal and/or family history of breast and ovarian cancer. However, it is important to note that the mutation, as one of the founders, is not identified in Manitoba Mennonites with strong family histories of cancer. Such families that meet testing criteria and do not carry the c.5238dupT mutation should be offered full screening for BRCA1 and BRCA2.

Missense Variant D1739V Supported by Multiple Lines of Evidence

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Objective: The clinical relevance of most non-truncating mutations in the BRCA1 and BRCA2 genes, including amino acid substitutions, is unknown. The assessment of these unclassified variants (UVs), which is crucial for appropriate risk management, requires a multimodal approach.

Methods: By integrating classical genetic, tumour histopathologic, functional, evolutionary conservation, and computational approaches, we investigated the pathogenic nature of a missense mutation, D1739V (c.5216A>T, p.Asp1739Val), of BRCA1.

Results: By integrating classical genetic, tumour histopatholog, functional, evolutionary conservation, and computational approaches, we investigated the pathogenic nature of a missense mutation, D1739V (c.5216A>T, p.Asp1739Val), of BRCA1.

Conclusions: All types of data generated in this study support a pathogenic nature of the variant D1739V, and we therefore propose to classify this amino acid substitution as a deleterious BRCA1 mutation.

The Breast Cancer Genes International Agency for Research on Cancer Database: A Tool to Improve Evaluation of BRCA1 and BRCA2 Unclassified Sequence Variants

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BRCA1 and BRCA2 are the two major high-risk breast cancer susceptibility genes, and the scale of clinical mutation screening of these genes probably exceeds that of any other set of cancer susceptibility genes. One complication resulting from genetic testing is that between 5% and 15% of patients screened are found to carry an unclassified sequence variant (UV). Observations of UVs create problems for the clinical lab staff who initially detect the UVs through to the patients who receive the ambiguous information.

The Breast Cancer Information Core has developed a Bayesian integrated evaluation of UVs based on co-segregation, personal and family history, tumour histopathology, and co-occurrence of variants in trans. Integrated evaluation includes a prior probability based on these in silico analyses of the substitutions, both from the perspective of potential effects on mRNA splicing and potential effects on protein function. To facilitate the evaluation of BRCA1 and BRCA2 UVs, we created the BRCA International Agency for Research on Cancer (iARC) database. This database contains the set of all possible single nucleotide substitutions (which constitute a sizeable preponderance of all UVs) to the open reading frames of BRCA1 and BRCA2. In silico analysis programs can be run on this finite set. The BRCA iARC database summarizes predicted potential to interfere with mRNA splicing (MaxEnt and NNsplice) and predicted missense substitution severity (position and Align-GVGD, using curated protein multiple sequence alignments) on the entire set. These scores are integrated to give a prior probability of pathogenicity for each substitution. Thus the database, and the Web site through which we present it (breca.iarc.fr), provides a starting point for the integrated evaluation of most BRCA1 and BRCA2 UVs.

Pathogenicity of the BRCA1 Missense Variant D1739V Supported by Multiple Lines of Evidence

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Objectives: Breast cancer is the most common cancer to affect South African women, with a lifetime risk of 1 in 13 for the African. Discovery of the BRCA1 genes introduced a new era for genetic testing, for not only could affected patients be informed regarding their risk for other cancer types, but other at-risk relatives could also be identified. The discovery of these genes urged many countries, including South Africa, to launch investigations into the prevalence of mutations in these genes.

Methods: A total of 157 African breast and/or ovarian cancer families (23 cases) were screened using conventional polymerase chain reaction analyses. Families carrying identical mutations were genotyped to determine whether the mutations resulted from independent events or a common ancestor. A genetic counselling protocol, based on classical clinical counselling strategies, was developed for patients requesting testing.

Results: BRCA1 mutations accounted for 19% (30/157) and BRCA2 mutations for 47% (73/157) of mutation-positive families. Three recurrent mutations were identified, two in BRCA1, 1493delC and E881X, and one in BRCA2, 8162delG. Genotype and genealogical analyses indicated common ancestors, with a specific European founding couple identified for each. Conversely, the 3 African founder mutations accounted for 94% (97/103) of BRCA2 mutation-positive families. A diagnostic testing protocol has since been implemented. Because of incomplete penetrance, preliminary investigations into potential single-nucleotide polymorphisms revealed that homoygosity for the variant allele of WA1 IF172–16C→G proved to be associated with an increased breast cancer risk among BRCA2 mutation carriers affected with breast cancer specifically.

Conclusions: The 3 founder mutations are unique and limited to the South African African population. The uniqueness could possibly be attributable to both mutations being de novo events; or the particular mutations might be extremely rare in the European populations from which the forefathers originally came; or lastly, both mutations were initially present in the European populations, but have become extinct.
MEETING ABSTRACTS

APPLIED RESEARCH

P018

Frequency of the c.1100delC CHEK2 Mutation in a Cohort of Belgian Familial Breast Cancer Patients: The Leuven Experience

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Nowadays genetic testing for hereditary breast cancer is offered worldwide. In about 20% of familial cases, the disease-causing mutations can be identified; the rest remain unexplained. Many of these mutations occur in the high-penetrance breast cancer genes BRCA1 and BRCA2. A limited percentage are attributable to pathogenic mutations in moderate-penetrance genes such as CHEK2.

Objectives: To determine the frequency of CHEK2 mutations in Belgian patients with breast cancer.

Methods: The frequency of CHEK2 mutations was assessed by multiplex ligation-dependent probe amplification (PLIA) in 867 consecutive breast cancer patients and carriers, representing 1100 Belgian families with breast cancer.

Results: The frequency of CHEK2 mutations in Belgian patients with breast cancer was 3.4%—very close to the frequency found in the Netherlands. Compared with the frequency in non-carriers, the CHEK2 mutation was more frequent in bilateral breast cancer patients and in those patients with an age of onset over 46 years. Our data are consistent with the observation that the CHEK2 gene is a multi-organ gene.

Conclusions: The frequency of CHEK2 mutation in Belgium warrants its implementation in the diagnostic service.

P020

Clinical, Pathologic, and Genetic Characteristics of Chinese Patients with BRCA1-Related Breast Cancer

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Background: Breast cancers attributable to underlying germline BRCA1 and BRCA2 mutations are associated with particular pathologic features that may differ from those of sporadic breast cancers. We report clinical and pathologic characteristics of breast cancer in Chinese women with and without mutations and of carriers of BRCA1 mutations as compared with BRCA2 mutations.

Methods: Based on age and family history, 226 high-risk women were recruited from March 2007 to November 2008. Medical information was prospectively collected from the patients and medical records. BRCA1 and BRCA2 mutations were detected using full gene sequencing and multiplex ligation-dependent probe amplification.

Results: Of the 226 female probands tested, 28 women (12.4%) were BRCA1 mutation carriers, and among these carriers, 11 (39.3%) had BRCA1 and 17 (60.7%) had BRCA2 mutations. The BRCA1 mutation carriers were more likely to have a familial history of breast and ovarian cancer, high-grade cancers, and triple negative (TNBC) cancers. Prevalence of TNBC was 48.3% in BRCA1 carriers and 25.6% in non-carriers; it was 67.7% in BRCA1 and 35.3% in BRCA2 carriers. The frequency of CHEK2 mutation was 4.9% in BRCA1 mutation carriers and 3.6% in BRCA2 mutation carriers.

Conclusions: BRCA1-related breast cancer in this Chinese population is associated with family history and adverse pathologic and prognostic features, with BRCA2 mutations being more prevalent, but BRCA1 carriers having more aggressive and TNBC cancers. Compared with Caucasian populations, the prevalences of BRCA2 mutations and TNBC are elevated.

BASIC RESEARCH

P030

Genetic Risk Modifiers in Hereditary Breast and/or Ovarian Cancer Families

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Objectives: To study the effect of 6 polymorphisms, recently described as being risk modifiers of sporadic breast cancer (BC) or ovarian cancer (OC) in BRCA1 families.

Methods: We recorded the cancer history (tumour site, age of diagnosis) of 548 women (293 carriers, 255 non-carriers) from 125 BC families (72 BRCA1, 53 BRCA2). The polymorphisms genotyped were: rs3351G/A and rs254110Pro/Arg3, localized in the progesterone receptor gene, CASP9-D02H, CASP8-652 6Nins/del, FGFR2 (rs2081582), and TNRC9 (rs3803662). Familial clustering was taken into account in the statistical analyses.

Results: Two polymorphisms modified the risk of BC in BRCA1 and BRCA2 families: rs3351 G/A heterozygous genotype increased the risk (odds ratio; OR: 2.41; 95% confidence interval (CI): 0.98 to 5.96; p = 0.056), and FGFR2 had a protective effect (OR: 0.52; 95% CI: 0.28 to 0.96; p = 0.037). FGFR2 was also significantly associated with increased risk of bilateral BC (OR: 2.67; 95% CI: 1.45 to 4.92; p = 0.002), whereas CASP9-652 6Nins/del had a trend toward a protective effect, restricted to BRCA1 mutation carriers (OR: 0.71; 95% CI: 0.48 to 1.05; p = 0.084). Furthermore, rs254110 was significantly associated with OC among the BRCA2 non-carriers (phenocopies—OR: 8.17; 95% CI: 2.17 to 30.76; p = 0.002).

Conclusions: We found evidence that certain polymorphisms involved in hormone-mediated cell proliferation and apoptosis modify the BC and OC risks in BC families. This may be relevant for an individual assessment of the most suitable preventive option among those currently available for each of those patients.

P032

Genome-Wide Linkage Analysis of High-Risk Breast Cancer Families from the International BRCA1/2 Consortium

M. S. Smith, et al.*

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Introduction: Variability observed in penetrance, age of onset, and site of the tumour, both among and within BRCA1 families, suggests that other low-penetrance genetic variants modify the cancer risk.

Objectives: To study the effect of 6 polymorphisms, recently described as being risk modifiers of sporadic breast cancer (bc) or ovarian cancer (oc) in BRCA1 families.

Methods: We recorded the cancer history (tumour site, age of diagnosis) of 548 women (293 carriers, 255 non-carriers) from 125 BC families (72 BRCA1, 53 BRCA2). The polymorphisms genotyped were: rs3351G/A and rs254110Pro/Arg3, localized in the progesterone receptor gene, CASP9-D02H, CASP8-652 6Nins/del, FGFR2 (rs2081582), and TNRC9 (rs3803662). Familial clustering was taken into account in the statistical analyses.

Results: Two polymorphisms modified the risk of BC in BRCA1 and BRCA2 families: rs3351 G/A heterozygous genotype increased the risk (odds ratio; OR: 2.41; 95% confidence interval (CI): 0.98 to 5.96; p = 0.056), and FGFR2 had a protective effect (OR: 0.52; 95% CI: 0.28 to 0.96; p = 0.037). FGFR2 was also significantly associated with increased risk of bilateral BC (OR: 2.67; 95% CI: 1.45 to 4.92; p = 0.002), whereas CASP9-652 6Nins/del had a trend toward a protective effect, restricted to BRCA1 mutation carriers (OR: 0.71; 95% CI: 0.48 to 1.05; p = 0.084). Furthermore, rs254110 was significantly associated with OC among the BRCA2 non-carriers (phenocopies—OR: 8.17; 95% CI: 2.17 to 30.76; p = 0.002).

Conclusions: We found evidence that certain polymorphisms involved in hormone-mediated cell proliferation and apoptosis modify the BC and OC risks in BC families. This may be relevant for an individual assessment of the most suitable preventive option among those currently available for each of those patients.

Although studies have shown that germline mutations in BRCA1 and BRCA2 account only for approximately half of all site-specific familial breast cancer (bc), to date there has been little success in using a linkage approach to identify further bc susceptibility loci. Thus, it seems likely that there are additional high-penetrance loci to be discovered. In 2006, we published the results of a linkage analysis in 149 multiple-case bc pedigrees. Although several regions of interest were identified, no definitive evidence of linkage was found, likely because of genetic heterogeneity. To address this, we have now analyzed a further set of 125 families that were part of an international consortium in familial bc including families from North America (n = 53), Europe (n = 57), and Australia (n = 14). All families had at least 3 cases of bc diagnosed at or before the age of 60 and were previously screened for mutations in BRCA1 and BRCA2. The primary analysis used MERLIN under a dominant mode of inheritance using the same model as in Smith et al., 2006.

Results: The linkage analyses identified one region with a conditional heterogeneity logarithm of odds (cLOD) of 2.2, and three others with LOD > 1.0. In a combined analysis of the new and existing data, suggestive evidence of linkage was identified on chromosome 4 (cLOD = 2.4, α = 0.15). When families were stratified by the presence of bilateral bc or of 5 or more cases of bc, the evidence (or the linkage) was increased (cLOD = 3.3, α = 0.30 and LOD = 3.6, α = 0.26 respectively). To further refine the hypothesized locus on chromosome 4, we genotyped a further 12 tandem repeat markers spanning the linked region. Fine mapping of likely-linked families has narrowed the region likely containing a bc susceptibility gene to 14.6 Mb covering approximately 18 Mb of dna. We are currently sequencing the most plausible candidate genes within this region.
A Mouse Model Reveals a Possible Function for BRCA1 in Human Uterine Leiomyosarcoma

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Uterine leiomyosarcoma (ULMS) is a rare gynecologic malignancy with a low survival rate. Currently, there is no effective treatment for ULMS. Infrequent occurrences of human ULMS hamper the understanding of the initiation and progression of the disease, thereby limiting the ability to develop efficient therapies. To elucidate the roles of the p53 and BRCA1 tumour suppressor genes in gynecologic malignancies, we generated mice in which both p53 and BRCA1 were conditionally deleted using anti-Müllerian hormone type 1 and 2 receptor (Amhr2)-driven Cre recombinase. Mice with a conditional deletion of p53 developed uterine tumours that resembled human ULMS. A concurrent deletion of p53 and BRCA1 accelerated the progression of these tumours. Consistent with the hypothesis that BRCA1 plays a role in ULMS, we have shown that the BRCA1 protein is absent in 29% of human ULMS. We have demonstrated that BRCA1 promoter methylation is the likely mechanism of BRCA1 downregulation in human ULMS. Our findings provide a rationale for investigating therapies that target BRCA1 deficiency in ULMS.

Morphologic and Immunohistochemical Correlation of Ovarian Epithelial Dysplasia in Prophylactic Oophorectomies from BRCA1/2 Mutation Carriers

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Objectives: Histopathologic examination of material from prophylactic salpingo-oophorectomies performed in patients at genetic risk has revealed frequent abnormalities interpreted as possible pre-cancerous “ovarian dysplasia” lesions. We sought to study the morphologic features and immunohistochemical expression patterns of neoplasia-associated markers in prophylactically removed ovaries.

Methods: Morphologic features and immunohistochemical expression patterns of Ki-67, p53, and ALCAM (an enzyme significantly associated with early-stage ovarian cancer) were evaluated in 35 prophylactic oophorectomies from BRCA1/2 carriers. Several cases were followed from formalin-fixed, paraffin-embedded tissue blocks were all read blindly by two gynecologic pathologists. Immunohistochemical staining results were correlated with morphologic findings.

Results: Mean dysplasia score was significantly higher in the genetic risk group than in controls (9.55 vs. 3.62, p < 0.0001). Increased ALCAM expression was observed in prophylactically removed ovaries compared with normal ovaries, and expression patterns of Ki-67 and p53 were low in both groups.

Conclusions: The increased dysplasia score and ALCAM expression in ovaries from BRCA1/2 carriers might be consistent with progression towards neoplastic transformation and could justify the use of the term “dysplasia” or intraepithelial ovarian neoplasia. Ovarian dysplasia may be a pre-malignant, noninvasive pathologic entity that could be an important step in early neoplasia, especially in ovaries from BRCA1/2 carriers.
Screening and Prevention Utilization in Unaffected BRCA Carriers

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Objectives: The purpose of this research is to carefully quantify the uptake of screening breast magnetic resonance imaging (MRI), chemoprevention, risk-reducing mastectomy (RRM), and risk-reducing salpingo-oophorectomy (RRSO) in unaffected BRCA carriers. We also aim to compare predictors of each of these options.

Methods: We thoroughly surveyed all women who received BRCA counselling and testing at University of California–San Francisco in the last 15 years. Using the 2008 Cancer Risk Program survey, BRCA testers were queried regarding screening and preventive options, risk perception, cancer worry, knowledge, and opinions regarding screening and prevention.

Results: The survey response rate was more than 80%, with 1137 returns. Approximately 20% of this population tested positive for a known deleterious BRCA mutation, and about half of the BRCA positives were unaffected with breast or ovarian cancer. Of the 102 unaffected BRCA carriers, 66 had BRCA1 mutations, and 36 had BRCA2 mutations. Their median age was 38 years (range: 20–64 years). Risk-reducing mastectomy was chosen by 30% of these women, and RRSO was chosen by 58%. The median time between BRCA testing and undergoing a risk-reducing surgery was 5 months. Of women with breast tissue at risk, 73% received screening MRI, and 7% chose chemoprevention (3.5% with tamoxifen, 3.3% with raloxifene). Cancer screening knowledge did not differ between women choosing use, chemoprevention, or surgeries. Breast cancer risk perception was significantly lower in women who chose MRI as compared with other options (p < 0.05), but cancer worry did not differ between option choices. As women aged, they tended to choose the following options, from younger to older age: MRI, RRM, RRSO, chemoprevention (p for trend < 0.05).

Conclusions: The uptake of screening breast MRI in unaffected BRCA carriers in this cohort was higher by a factor of approximately 10 than was the uptake of chemoprevention. The median time to MRI and RRSO, for women who chose those options, was quite short, at approximately 5 months. It is important to present unaffected BRCA carriers with a range of options for screening and prevention, understanding that there is a wide variation of uptake for each option.

Review of BRCA1 and BRCA2 Mutation Screening at Guy’s Hospital, London, U.K.

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In response to the 2004 U.K. National Institute for Health and Clinical Excellence Guidelines for Familial Breast Cancer, our screening strategy for the BRCA1 and BRCA2 genes was changed from screening approximately two thirds of each gene using a protein-truncation test and fluorescent heteroduplex analysis to a complete screen of both genes using sequencing and multiplex ligation-dependent probe amplification (MLPA).

Since May 2006, we have screened 564 new referrals and have found BRCA1 or BRCA2 pathogenic mutations in 99 individuals (18%). A variant of unknown significance (VUS) has been identified in a further 83 individuals (15%). We have also retrospectively completed a full screen on 738 individuals who previously had a negative two-thirds screen. This involved a significant collaboration with the Clinical Genetics Team who reviewed all individuals seen before January 2000 to ensure that they met current testing guidelines. In this group, BRCA1 or BRCA2 pathogenic mutations were identified in 45 individuals (6%), and a VUS was identified in a further 48 individuals (7%). Of the 144 (65 BRCA1, 79 BRCA2) pathogenic mutations identified, 127 (51 BRCA1, 76 BRCA2) were detected by sequencing, and 17 (14 BRCA1, 3 BRCA2) by MLPA. A total of 40 BRCA1 mutations and 44 BRCA2 mutations have been seen in only 1 individual. Of the 11 BRCA1 and 12 BRCA2 mutations that have been identified more than once, most have been seen in only 2 unrelated individuals. However BRCA1 c4065_4066delTCAAA has been seen in 4 unrelated individuals, duplication of BRCA1 exon 13 in 3 unrelated individuals, BRCA2 c.755_7556delACAG in 6 unrelated individuals, and BRCA2 c.7988A>T in 5 unrelated individuals.

In total, 126 different VUS have been identified, of which 101 have been seen in only 1 individual. These comprise 68 missense variants (23 BRCA1, 45 BRCA2), 30 synonymous variants (13,17), and 28 in intronic variants (8,20).

A Rare Case of Anal Melanoma in a BRCA1 Mutation Carrier

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Objectives: Mutations in BRCA1 and BRCA2 are associated with an increased risk of breast, ovarian, prostate, and pancreatic cancer. There have been reports of a mild association of BRCA1 and BRCA2 mutations with cutaneous and ocular melanoma. We present a rare case of an anal melanoma in a BRCA1 Ashkenazi Jewish founder mutation carrier.

Methods: A 62-year-old male of Ashkenazi Jewish descent presented with symptoms of anal pain and hematochezia. Sigmoidoscopy revealed an anal mass, and biopsy demonstrated invasive malignant melanoma of the anal canal. Staging work-up revealed liver metastases. Family history was significant for a sister who died at age 30 from breast cancer, and a 40-year-old daughter diagnosed with ovarian cancer at age 36 who tested positive for a BRCA1 mutation, 187delAG.

The patient presented for genetic counselling and testing. The patient’s peripheral blood was sent for analysis of the three Ashkenazi Jewish founder mutations: BRCA1 187delAG, BRCA1 5383insC, and BRCA2 6174delT.

Results: Molecular analysis identified a deleterious mutation, 187delAG, in the BRCA1 gene. This is a frameshift mutation that results in premature truncation of the BRCA1 protein at amino acid position 39. This confirms the diagnosis of hereditary breast and ovarian cancer syndrome (BRCA1/2).

Conclusions: This case represents the first report of an anal melanoma in a BRCA1 carrier and sheds light on an unusual cancer presentation in patients with BRCA1/2. Earlier studies have shown an association between melanoma and BRCA2 mutations, but there has been little to-no evidence of this association in BRCA1 carriers, especially carriers with Ashkenazi Jewish founder mutations. Detection of the BRCA1 187delAG mutation in this patient leads to promising therapeutic options using novel systemic agents for this highly aggressive, potentially lethal malignancy.
A Rare Case of Ocular Melanoma in a BRCA1 Mutation Carrier
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Objectives: Hereditary breast and ovarian cancer syndrome (HBOC) is associated with a mildly increased risk of cutaneous and ocular melanoma. Earlier studies have demonstrated ocular melanoma in association with BRCA2, but not BRCA1. We present a rare case of an ocular melanoma and primary peritoneal carcinoma in a BRCA1 mutation carrier.

Methods: A 62-year-old Caucasian female presented with an abnormal Papanicolaou smear and underwent a laparotomy, which revealed primary peritoneal carcinoma. Her past medical history was significant for decreased night vision, leading to the diagnosis of a left ocular melanoma (spindle-cell, type B) at age 57 years. She underwent a therapeutic enucleation at that time. Family history is remarkable for early-onset breast cancer in her sister and two nieces, all diagnosed in their 40s. The patient presented for genetic counselling and testing following the diagnosis of primary peritoneal carcinoma. The patient’s peripheral blood was sent for full sequence analysis of the BRCA1 and BRCA2 genes. Direct DNA sequence analysis was performed on the polymerase chain reaction products corresponding to the entire BRCA1 and BRCA2 coding region.

Results: The molecular analysis identified a deleterious mutation, c.267_268delAG, in the BRCA1 gene. This is a frameshift mutation that results in premature truncation of the BRCA1 protein at amino acid position 808, confirming the diagnosis of HBOC.

Conclusions: The patient described above developed two rare malignancies, ocular melanoma and primary peritoneal carcinoma. Earlier studies have identified a 2%-3% prevalence of cancers in individuals with HBOC.

Risk Factors for Carcinoma of the Fallopian Tube in Women with a Germline BRCA Mutation and in Sporadic Cases
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Objectives: To evaluate risk factors that may relate to the development of fallopian tube cancer in women with and without a BRCA mutation.

Methods: Subjects with fallopian tube cancer were identified from a large international registry of women that carry a BRCA1 or BRCA2 mutation (n = 56) and from a population-based study of ovarian and fallopian tube cancer in Ontario, Canada (n = 67). BRCA mutation status was established for all subjects. All subjects completed a questionnaire about medical history and lifestyle factors. Subjects were matched with controls for date of birth within 4 years, for all subjects completed a questionnaire about medical history and lifestyle factors. Increasing parity was associated with a decreased risk of death from prostate cancer (hazard ratio (HR): 0.90; 95% confidence interval (CI): 0.45 to 1.80). Results: We studied 105 women with fallopian tube cancer (48 with a BRCA1 mutation, 10 with a BRCA2 mutation, and 47 with no identified BRCA mutation) and 958 matched controls. Increasing parity was associated with a decreased risk of fallopian cancer in non-carriers (trend per birth—odds ratio (OR): 0.93; 95% CI: 0.87 to 1.00; p = 0.02), in BRCA1 carriers (OR: 0.98; 95% CI: 0.87 to 1.12; p = 0.07), and in BRCA2 carriers (OR: 0.94; 95% CI: 0.85 to 1.05; p = 0.06), but was significant only for non-carriers. Oral contraceptive use was protective for BRCA1 carriers (trend per year of use—OR: 0.90; 95% CI: 0.82 to 1.00; p = 0.03), but was not protective for non-carriers (OR: 0.99; 95% CI: 0.88 to 1.10; p = 0.80) or BRCA2 carriers (OR: 0.95; 95% CI: 0.82 to 1.11; p = 0.53). No significant effects were observed for body mass index, hormone replacement therapy, or tubal ligation.

Conclusions: Parity and oral contraceptive use were associated with a decreased risk of fallopian tube cancer, but these effects were not observed consistently across all groups stratified by mutation. The risk factors for fallopian tube cancer appear to be similar to risk factors for ovarian cancer.

Detection and Prevention of Hereditary Breast Cancer
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Objectives: There is a paucity of data looking at how breast cancers are diagnosed in BRCA carriers and the stage in which these women are diagnosed. Our hypothesis is that BRCA1/2 carriers will more often present with a palpable mass than be identified through imaging and that they will be diagnosed with advanced-stage disease that requires traditional chemotherapy. We also wanted to examine the relationship between mutation status and pathology. We conducted a retrospective evaluation of female breast cancer patients who carried BRCA1/2 mutations to determine how women find their cancers and how those cancers are treated. Our study included 66 women who were diagnosed with breast cancer and who carried a BRCA1/2 mutation. Almost half the women presented with palpable masses, but only 1/2 were diagnosed by mammographic findings. Surprisingly, of the 66 cancers, 50 were diagnosed at stage 0-ii, and only 5 were diagnosed at stage iii or higher. BRCA1 cancers were more often triple-negative than were BRCA2 cancers (42% and 21% respectively). Although these cancers were diagnosed at early stages, 92% of BRCA1 carriers and 68% of BRCA2 carriers still were treated with traditional chemotherapy. We feel that, if a woman is presented with this data showing that an impending cancer will more often than not require chemotherapy, she may choose more aggressive management and opt for prophylactic mastectomy.

Conclusions: BRCA1 cancers are triple-negative and may also change recommendations regarding the use of anti-hormonal agents and leave women with fewer prevention options. All of this information should be presented to women to help them make an informed decision about prevention when they receive results.

Detection and Treatment of Hereditary Breast Cancer
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Objectives: To determine clinicopathologic characteristics of BRCA-associated prostate cancer in men of Ashkenazi Jewish (AJ) ancestry.

Methods: We determined BRCA mutation prevalence in 832 AJ men with localized prostate cancer and 435 AJ controls, and compared age, stage, prostate-specific antigen, Gleason score (GS), overall survival, and other clinical outcome measures among 26 BRCA mutation carriers and 906 non-carriers. Kruskal–Wallis tests were used to compare age of diagnosis and OS, and logistic regression models were used to determine associations between carrier status, prostate cancer risk, and OS. Hazard ratios were estimated using Cox proportional hazards models.

Results: BRCA2 mutations were associated with triple risk for prostate cancer (odds ratio (OR): 3.18; 95% confidence interval (CI): 1.52 to 6.66; p = 0.002). BRCA1 mutation conferred no increased risk, BRCA2 carriers presented at a younger age (median: 62 years vs. 68.2 years; p = 0.05). Adjusting for age, BRCA1 mutation carriers were more likely than non-carriers to have a Gleason score of 7 or higher (hazard ratio (HR): 2.37; 95% CI: 1.14 to 4.94; p = 0.021), but BRCA1 mutation carriers were not (HR: 0.62; 95% CI: 0.12 to 3.10; p = 0.56). After 7254 person–years of follow-up, BRCA1 and BRCA2 mutation carriers, as compared with non-carriers, had an increased risk of developing biochemical recurrence (HR: 4.32; 95% CI: 1.31 to 13.62; p = 0.016 and HR: 2.41; 95% CI: 1.23 to 4.75; p = 0.011 respectively) and castrate metastases (HR: 2.15; 95% CI: 0.28 to 16.31; p = 0.46 and HR: 3.01; 95% CI: 1.26 to 7.14; p = 0.013). There was a greater risk of death from prostate cancer (HR: 5.16; 95% CI: 1.09 to 24.53; p = 0.039 and HR: 5.48; 95% CI: 2.03 to 14.79; p = 0.001) for BRCA1 and BRCA2 carriers respectively.

Conclusions: BRCA2 mutation carriers had an increased risk of prostate cancer, a younger age of onset, and a more aggressive phenotype. Prostate cancer cases with either BRCA1 or BRCA2 mutations were characterized by an adverse clinical outcome. These results may affect clinical management of this subset of hereditary prostate cancer.
**P058**

**Single-Staged Breast Cancer Reconstruction Using the Combined Spectrum Expander/Implant**

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**Introduction:** Breast reconstruction is an important procedure that requires attention to detail for optimal outcome. Spectrum implants ( Mentor Corporation, Santa Barbara, CA, U.S.A.) have provided us with the tools to optimally adjust for achieving symmetry and patient satisfaction. Single-staged breast reconstruction with saline implants was first introduced by Becker in 1982. Despite advances in our surgical techniques and prosthesis, the reported complication rate remains high at 28%.

**Purpose:** We present the senior author’s experience over a period of 7 years for immediate and delayed breast reconstruction as it relates to implant reconstruction. This is the first reported study of breast reconstruction using Spectrum Expander/Implant with the use of the same retropectoral technique and no alloderm. All implants were saline-filled.

**Methods:** A retrospective review analysis of patients undergoing Spectrum implants by the senior author from 2001 to 2008 is presented. Specific attention is paid to factors associated with major complications.

**Results:** A total of 44 patients underwent 67 Spectrum breast implant reconstructions. The mean age of the study patients was 42.7 years (standard deviation: ±11.6 years; range: 17–67 years). There was no skin necrosis (0%). Complications requiring reoperation and general anesthesia were capsular contracture (n = 10; 14.9%), perioperative infection (n = 3; 4.5%); and delayed sudden deflation (n = 2; 3.0%). Only 1 patient was subsequently judged not suitable for reoperation/reconstruction because of severe neuropenia. The subcutaneous valves were removed under local anesthesia during nipple reconstruction and no alloderm. All implants were saline-filled.

**Conclusions:** Use of a versatile integrated all-in-one expander/implant (Spectrum) can achieve great results with minimal complication and reoperation rates.

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**P059**

**Early Experiences of the Calgary Breast Health Program**

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*Tom Baker Cancer Centre and Centre for Health Services—Calgary, Breast Health Program, Calgary, Alberta, Canada; and Queen’s University, Kingston, Ontario, Canada.

**Background:** In 2007, funding was obtained to formalize and evaluate the Calgary Breast Health Program high-risk breast cancer clinic, which commenced February 2008. The team included a physician, nurse, psychologist, and clerical assistant. The target patient population included individuals at high genetic risk (per medical genetics), women diagnosed with lobular carcinoma in situ or atypical hyperplasia, and women who received mantle radiotherapy in childhood or teenage years. The initial appointment involved self-completion of a needs assessment questionnaire, a brief meeting with the psychologist, breast (<1 ovarian) cancer risk assessment, overview of screening and risk-reduction options, and a nurse-led discussion of lifestyle strategies and signs or symptoms of breast (<1 ovarian) cancer. The pilot period was completed in March 2009, and the clinic continues to operate under the same model.

**Objectives:** To describe the demographic profile of clients; the perceived pre-consultation needs of clients; the recommendations made by the clinic; and the referrals generated.

**Methods:** All clients seen from February 2008 through May 2009 who agreed to be contacted regarding further research were sent a risk assessment questionnaire. Pertinent information on consenting clients will be abstracted from charts. Descriptive statistics will be applied to the data obtained.

**Results:** By May 31, 2009, the clinic had received 116 referrals, and 73 consultations had been completed. Referrals came from medical genetics (n = 57), primary care (n = 38), surgery (n = 20), gynecology (n = 1), and psychology (n = 1). Of the referrals not seen, most were referred directly to medical genetics (n = 22) or were waiting on a pending appointment (n = 6). For consenting clients seen by the clinic, aggregate data describing demographic profile, perceived pre-consultation needs, recommendations made, and referrals made are presented.

**Conclusions:** The introduction of a formal high-risk breast cancer clinic in Calgary has been a feasible undertaking. Assessment and revision of the current clinic model is ongoing.

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**P060**

**Hereditary Breast/Ovarian Cancer High-Risk Clinic: Our First 12 Years**

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**Objectives:** To describe the clinic’s first 12 years of experience with high-risk screening for women with confirmed BRCA1 or BRCA2 gene mutations.

**Methods:** To date, 262 women with BRCA1/2 mutations have been seen by the clinic physician and nurse specialist for consultation. Prophylactic mastectomy with reconstruction is an option discussed with all women, and high-risk breast screening is offered as the alternative. Screening includes clinical breast exam and imaging every 6 months, with an alternating schedule of breast screen (from age 25) and mammography (from age 30). Prophylactic bilateral salpingo-oophorectomy (pso) is recommended for all mutation carriers. Ovarian cancer screening is not currently recommended. Women are discharged to their family physician’s care after prophylactic surgeries, and those with a new cancer diagnosis or recurrence are referred to an oncologist.

**Results:** Of BRCA1/2 mutation carriers attending the clinic, 16% have had at least 1 new cancer diagnosis; 46 new cancers have been detected (26 in BRCA1-positive, 20 in BRCA2-positive), with 4 women having 2 new diagnoses. Cancers diagnosed include 24 invasive breast cancers, 7 ovarian cancers, 4 cases of ductal carcinoma in situ (dcis), 2 cases of lobular carcinoma in situ (lcis), 2 fallopian tube (ft) cancers, 2 peritoneal cancers (both after pso), 2 pancreatic cancers, 1 malignant melanoma, 1 gastric cancer, and 1 colorectal cancer. Of the breast cancers, 10 were identified on MRI screening. Six cancers were diagnosed by pathology review at the time of prophylactic surgery, including 3 ovarian cancers, 1 ft cancer, 1 breast cancer, and 1 lcis, with an additional lcis diagnosis on contralateral mastectomy.

**Conclusions:** Over the past 12 years, the clinic has provided a valuable service to high-risk women in our province. Experienced clinicians offer information, support, referral to expert surgeons, breast imaging, and clinical follow-up. This high-risk clinic will provide ongoing opportunities to evaluate the efficacy of specific cancer risk management strategies.

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**P061**

**Somatic Complaints in Women with Risk-Reducing Salpingo-Oophorectomy to Prevent Hereditary Breast/Ovarian Cancer**

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**Background:** Risk-reducing salpingo-oophorectomy (rrso) is effective for preventing ovarian cancer in women from hereditary breast/ovarian cancer families, but the procedure may be associated with somatic morbidity and complaints. In this cross-sectional study, we aimed to study self-reported somatic complaints in women who had undergone rrso.

**Methods:** Based on surgical records, we identified and invited 503 women who had undergone rrso after genetic counselling at a median of 6 years earlier. Among these, 338 (67%) completed questionnaires on demographic and medical issues, somatic complaints, and mental distress. We randomly allocated 5 age-matched controls per case (n = 1690) among participants with intact ovaries from the population-based Norwegian Nord-Trøndelag Health Study.

**Results:** The rrso group had significantly more palpitations (p < 0.02), constipation (p = 0.01), pain and stiffness (p = 0.02), musculoskeletal disease (p = 0.01), and osteoporosis (p = 0.02) than the control group. In a multivariate analyses of the total sample, membership in the rrso group was significantly associated with osteoporosis [odds ratio (or) 2.4; 95% confidence interval (ci) 1.3 to 4.5], musculoskeletal disease (or 1.7; 95% ci 1.0 to 2.7), osteoporosis (or 2.4; 95% ci 1.5 to 3.5), and constipation (or 1.4; 95% ci 1.1 to 2.0). Current use of hormonal replacement therapy (hrt) was significantly associated with musculoskeletal disease (or 1.5; 95% ci 1.3 to 2.3), osteoporosis (or 2.0; 95% ci 1.2 to 3.3), pain and stiffness (or 1.4; 95% ci 1.1 to 1.8), fibromyalgia (or 1.6; 95% ci 1.1 to 2.5), nausea (or 1.5; 95% ci 1.1 to 2.1) and constipation (or 1.5; 95% ci 1.2 to 1.9). In multivariate analyses of the rrso group only, no significant association was observed between the use of hrt and somatic complaints, but time since rrso was significantly associated with osteoporosis (or 1.2; 95% ci 1.1 to 1.3), nausea (or 0.9; 95% ci 0.8 to 1.0), and constipation (or 0.9; 95% ci 0.9 to 1.0).

**Conclusions:** The rrso group had more palpitations, constipation, and musculoskeletal disease than did the control group. Use of hrt was associated with musculoskeletal pain conditions and gastrointestinal symptoms, which may indicate that women with more symptoms choose hrt. Follow-up examinations after rrso should also focus on somatic complaints.
**Effect of Reminder Telephone Calls on Mammography Compliance in High-Risk Women**

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The effect of a reminder telephone call intervention designed to encourage mammography has not been systematically studied in a high-risk population in which compliance is crucial. We hypothesize that a simple reminder telephone call would significantly increase mammography uptake in high-risk women as compared with a control group. In 447 high-risk women, consent to participate in the study was obtained. Interestingly, 346 (77%) of these women self-reported being compliant in obtaining annual mammography for the preceding 2 years. Verification was established in a subset of these women by obtaining their mammogram reports. Subjects who were noncompliant by self-report (n = 32) were randomized to the intervention or the control group. Reminder and follow-up telephone calls were completed on 16 women in the experimental group, and follow-up calls were completed on 15 women in the control group. One woman was excluded from the study after being diagnosed with breast cancer upon obtaining a mammogram. A statistical difference (p = 0.0017) was observed between the two groups in support of the hypothesis that mammography compliance in high-risk women can be increased if an intervention such as a simple reminder call is implemented. Future studies should identify barriers in obtaining mammograms in these high-risk individuals.

**Results and Uptake of Risk-Reducing Salpingo-Oophorectomy in Women Over the Age of 40 Years at High Familial Risk of Ovarian Cancer**

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**Objectives:** To evaluate outcomes and factors affecting uptake of risk-reducing salpingo-oophorectomy (RRSO) in women at high risk for ovarian cancer over the age of 40 years.

**Methods:** Prospectively collected data of women attending a Familial Cancer Clinic between 2004 and 2009 were analyzed. The preferred management option recommended to women over the age of 40 years with a 10% or greater estimated lifetime risk of ovarian cancer was RRSO with peritoneal washings. Those declining surgery opted for screening under the United Kingdom Familial Ovarian Cancer Screening Study. A strict histopathologic protocol with serial slicing was used for RRSO specimens.

**Results:** Of 1518 high-risk women from breast/ovarian cancer families seen from 2004 to 2009, 517 were below 40 years of age. Of the 1001 others, 181 were BRCA1/2 carriers, and 820 had unknown mutation status (UMS). Of the 1001 women, 263 (of whom 33.5% were carriers) chose RRSO, and 738 (of whom 12.6% were carriers) preferred screening.

The median ages (and interquartile ranges) for the RRSO and screening groups were 50.9 (45.3, 57) years and 47.5 (43.2, 54.9) years respectively (p = 0.001). Women undergoing RRSO were more likely to have had breast cancer (p < 0.0005), to have a relative below the age of 50 years with ovarian cancer (p = 0.012), to have a parity of P2 or greater (p = 0.039), to be postmenopausal (p < 0.0005), and to carry BRCA gene mutations (p < 0.0005). The median age of women undergoing RRSO was lower in BRCA carriers than in UMS women (p < 0.000001).

The occult cancer rate in the whole cohort was 3.8% [95% confidence interval (ci): 1.84% to 6.88%]. Of the 7 tubal and 3 primary ovarian cancers, 6 were in mutation carriers, for a 6.8% incidence (95% ci: 2.5% to 14.25%), and 4 were in UMS women for a 2.3% incidence (95% ci: 0.63% to 5.75%; p = 0.09).

**Conclusions:** Various factors affect decision-making in high-risk women. BRCA carriers and UMS women both have a high occult malignancy rate, and the age at detection of lesions is similar in both groups. It is important that women are made aware of these facts, because UMS women tend to delay surgery till becoming postmenopausal.

* Equal contribution.

**Results of a Phase II Open-Label Nonrandomized Trial of Cisplatin Chemotherapy in Patients with BRCA1-Positive Metastatic Breast Cancer**

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**Purpose:** To evaluate the efficacy of cisplatin chemotherapy in BRCA1 mutation carriers with metastatic breast cancer.

**Design:** In a phase II open-label study, 20 patients with metastatic breast cancer who carried a mutation in BRCA1 were treated with cisplatin 75 mg/m2 intravenously every 3 weeks as part of a 21-day cycle for 6 cycles. Restaging studies to assess response were performed after cycles 2 and 6, and every 3 months thereafter.

**Results:** Between July 2007 and January 2009, 20 patients were enrolled. At baseline, 65% had received prior adjuvant chemotherapy, and 55%, prior chemotherapy for metastatic breast cancer. Mean age in the group was 48 years (range: 32-70 years); 30% were estrogen (e) or progesterone receptor (pR)-positive; 70% were e/pe or nene-negative; and 0% were nene-positive. Overall response rate was 80%; 9 patients experienced a complete clinical response (45%), and 7 experienced a partial response (35%). The 1-year survival was 93%. Cisplatin-related adverse events, including nausea (50%), anemia (5%), and neutropenia (35%), were mostly mild to moderate in severity. One patient discontinued therapy because of grade 4 neutropenia.

**Conclusions:** This phase II study demonstrates that cisplatin chemotherapy has high activity in women with a BRCA1 mutation and metastatic breast cancer and is generally well tolerated.
**MEETING ABSTRACTS**

### EDUCATION AND COMMUNICATION

**P070**

### Increasing Medical Student Awareness of Inherited Susceptibility to Breast Cancer

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In 2007, it was reported that North American medical school curricula are lacking in the area of clinical genetics (Thurston VC et al. Acad Med 2007;82:441–5). Implementation of an online self-directed learning module (SLM) provides students with an interactive case-based approach that simulates real-world uses of genetic testing and highlights the implications of such tests on patient decision-making.

**Objectives:** To design and implement a SLM for medical students to improve their knowledge of hereditary cancers using breast cancer as the model.

**Methods:** The SLM will be designed by a medical student working together with a clinical geneticist and a medical oncologist. The objectives for the module are focused on providing students with knowledge in three areas:

- How to interpret a patient’s personal and family history to assess level of risk
- Understanding the implications of genetic testing as they relate to hereditary cancers
- Increasing awareness of the diversity of psychosocial issues facing women and their families who are found to have a BRCA mutation

**Results:** Assessment of the efficacy of this SLM as a teaching tool will be carried out using a focus group of medical students. The focus group will examine the effectiveness of the SLM in increasing student knowledge and in addressing the limitations of the modality so that those limitations can be overcome.

**Conclusions:** SLMs present an excellent opportunity to provide early impact in medical school curricula. By integrating knowledge from several key disciplines (genetics, oncology, and gynecology) with appropriate emphasis on the psychosocial aspects of risk assignment, medical students will have an increased understanding of adult-onset genetic conditions.

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### GENETIC COUNSELLING

**P080**

### Secondary Use of Data and Samples from Deceased Participants: What Are the Legal and Ethical Constraints?

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**Context:** Biologic samples and data used in research may, in some cases, be of great use for another research project. This is called “secondary use.” The secondary use of data and samples is current practice among health care researchers, even more so since the increasing use of biobanks. However, particular issues arise when the participant to whom the samples and data are linked is deceased. Although ethical and legal framing of secondary use of samples and data of live participants is in place and is consistent across various normative texts, the framework concerning deceased participants is often less developed and sometimes inexistential. Although the value of these samples is often inestimable, using them may sometimes prove to be complex.

**Methods:** Our research consists of an analysis of the legal and ethical framework in place in Quebec and Canada concerning the secondary use of data and samples from deceased individuals. We analysed how the various norms address some specific questions such as the scope of consent for use after death and considerations concerning the wishes of living relatives related to the use of data and samples of a deceased individual, when there is no prior indication of individual choice.

**Results:** We provide a synthesis of the elements that need to be taken into account before considering the secondary use of such material. We will examine, by way of example, the procedure set in place to allow secondary use in the case of the continuation of the INHERET breast cancer project, funded by the Canadian Institutes of Health Research Team in Familial Risks of Breast Cancer. Finally, we will conclude by presenting lessons learned, which could be helpful for the elaboration of a uniform ethical and legal framework in Canada.

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**P081**

### Breast Cancer at Less Than 36 Years of Age Is a Strong Predictor of BRCA1/2 Status in French-Canadian Cancer Families

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The presence of a breast cancer case diagnosed before 36 years of age is a strong indicator for the presence of a BRCA1/2 mutation in French-Canadian (1/0) hereditary breast and/or ovarian cancer (HBOC) families. Earlier studies were limited by screening for the mutations that recur in this founder population. We have assessed the contribution of BRCA1/2 to 75 HBOC and 51 HBOC (p53) families who have at least 3 cancer cases per family and who have undergone Myriad Genetics mutation screening and multiplex ligation-dependent probe amplification analysis for rearrangements. About 50% of the HBOC families were mutation-negative. About 32% were 3-case families, in which 71% were mutation-negative (p = 0.014). The distribution of cases relative to an index tested case in the HBOC mutation-negative group differed from that in the mutation-positive group (p = 0.004), where more first-degree relatives with breast cancer were observed in the mutation-negative families (56%) than in the mutation-positive families (35%, p = 0.0009). The frequency of breast cancer diagnosed before the age of 36 years in the HBOC mutation-negative families (6%) differed significantly from that in the HBOC (23%, p = 0.0002) and HBOC (23%, p < 10−10) mutation-positive families. Thus, although 50% of the HBOC families harboured a mutation, about 80% of families with at least 1 breast cancer case diagnosed below the age of 36 years were mutation-positive. About 69% of the HBOC families were mutation-positive, and as compared with the 19% of mutation-negative families, 54% of those mutation-positive families contained at least 1 breast cancer case diagnosed below the age of 36 years (p = 0.017). Of the 18 HBOC families with fewer than 3 cases of breast cancer, half of the 10 mutation-positive families contained at least 1 breast cancer case diagnosed below the age of 36 years as compared with no cases diagnosed in mutation-negative HBOC families in this sub-group. These observations reaffirm the significance of the presence of very young age of diagnosis in HBOC and HBOC families and should facilitate decisions aimed at genetic testing for BRCA mutations.
**P084**

**A Study of Preventive Practices Among Canadian Women with a BRCA1 or BRCA2 Genetic Mutation**

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**Methods:** Eligible patients were identified from a database of 307 probands who had previously been selected for genetic screening.

**Results:** Of the 625 eligible women contacted, 464 (74%) women returned completed questionnaires. Of whom 77% (356) had undergone no preventive options. The women’s mean age was 51.6 years (range: 25–70 years), and they received their genetic test results a mean of 79.2 months (range: 18–239 months) before completing the questionnaire. Of the 218 women without breast cancer, 147 (67%) opted against prophylactic mastectomy, because of body image and sexuality concerns, and a preference for other preventive options; 49 (22%) opted against prophylactic oophorectomy because of concerns about medical side effects, menstrual symptoms, and effects on sexuality; 195 (89%) opted against preventive tamoxifen because of concerns about side effects and lack of recommendation; and 49 (22%) opted against preventive magnetic resonance imaging because of lack of access, availability, and recommendation.

**Conclusions:** Health care providers should ensure that women with a BRCA1 or BRCA2 mutation are presented with all preventive options, should be more proactive in addressing the concerns of these women, and should encourage informed decision-making about suitable preventive options.

**P085**

**Efficiency of BRCApro and Myriad Models and the Use of Mutation Probability Thresholds for BRCA1/2 Mutation Detection: A Retrospective Evaluation**

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Considerable differences exist between countries in the mutation probability methods and thresholds used to select patients for BRCA1 and BRCA2 genetic testing.

**Objectives:** To assess the efficiency of the mutation probability methods BRCApro and Myriad, as compared with criteria based on cancer history in the family, to select patients eligible for BRCA1/2 mutation detection.

**Methods:** We retrospectively calculated the BRCApro and Myriad probabilities in 307 probands who had previously been selected for DNA analysis according to a list of eligibility criteria based on number of first- and second-degree relatives affected with breast cancer (bc) and/or ovarian cancer (oc), age of diagnosis, presence of bilateral bc, and oc in men. Until the year 2000, DNA analysis was performed using the protein truncation test (PTT). The DNA from patients in whom no mutation was initially found was resequenced using the PTT later on under new complete genetic screening.

**Results:** The PTT detected 21 mutations (6.5%); complete genetic screening identified an additional 32 mutations (10.4%) and 11 unclassified variants (3.6%). Compared with cancer history, a threshold of 10% with BRCApro or Myriad excluded about 40% of the patients from analysis, including 4 patients with a mutation and probabilities under 10% with both programs. All 4 probands had a BRCA2 mutation. The BRCApro and Myriad methods showed similar specificity at 10% threshold, and overall, BRCApro was more sensitive than Myriad. Only 2 of the probands with a uv had probabilities above 20% with BRCApro and Myriad.

**Conclusions:** The mutation probability models BRCApro and Myriad are economically more than efficient than cancer history criteria alone. For the detection of BRCA1 mutations, BRCApro, at a 10% threshold, is equally sensitive; however, for BRCA2 mutations, the method is most sensitive. Most of the patients with a uv have low probabilities with both methods, which may be an indicator of the lack of pathogenicity of those uv.

**P082**

**Transitional-Cell Ovarian Carcinoma Is Associated with Germline BRCA1/2 Mutations**

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**Objectives:** The population prevalence of BRCA1 and BRCA2 mutations for women with epithelial ovarian carcinoma is estimated to be around 11%–15%, with the predominant histology of tumours being serous. Our study is designed to estimate the strength of the association between transitional-cell ovarian carcinomas and germline mutations in BRCA1 and BRCA2.

**Methods:** We searched our pathology department’s database for cases of epithelial ovarian carcinoma with transitional-cell histology. Patients (or their surviving first-degree relatives) were offered genetic counselling and testing for germline mutations in BRCA1 and BRCA2. Pathology slides were reviewed and confirmed by a gynecologic pathologist.

**Results:** Between 1996 and 2008, 25 patients with a mean age of 58 years (range: 39–75 years) were included. In all patients and one patient’s son underwent genetic testing. The other patients could not be tested because they were either deceased or lost to follow-up. We identified 6 BRCA1 and 2 BRCA2 mutation carriers with a mean age of 57 years (range: 51–63 years). A conservative estimate of the prevalence of BRCA1 and BRCA2 mutations, including all 25 transitional-cell histology patients as the denominator, identified a frequency of 32% (95% confidence interval: 14%–50%). If we had been able to offer testing to all patients in our transitional-cell cohort, the prevalence of BRCA1/2 mutations may have been even greater. All patients who tested positive had a family history of cancer, including 5 families with breast cancer, 3 with ovarian cancer, and 2 with colon cancer.

**Conclusions:** Transitional-cell histology apparently has a higher prevalence of BRCA1/2 mutation carriers than is found in the general population. Further molecular characterization may elucidate whether the transitional-cell and serous histologies represent distinct phenotypes.

**P083**

**Long-Term Stability of Knowledge Acquired During Genetic Counselling for Breast/Ovarian Cancer Susceptibility**

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**Purpose:** Studies have shown that standard genetic counselling increases knowledge of breast and ovarian cancer genetics. However, little is known about the retention of such information long after test result disclosure. This study assesses the stability of knowledge acquired during pre-test genetic counselling up to 3 years following BRCA1/2 test result disclosure.

**Methods:** Participants were a consecutive series of 429 French-Canadian women (89 carriers, 120 non-carriers, and 220 with an inconclusive test result) from 187 families, who were tested for BRCA1/2 mutations as part of the InHerit BCaR multidisciplinary research program. Knowledge about BRCA1/2 testing was assessed by a 19-item true–false measure at the pre-test genetic counselling session and 3 years after result disclosure. Knowledge scores are expressed as percentages of correct answers.

**Results:** Three years post-disclosure, mean knowledge scores were moderate (56.7 ± 18.2) and lower than those at pre-test genetic counselling (58.9 ± 18.7; paired t-test p = 0.0059 [intra-class correlation coefficient: 0.58]). An average of 70% of participants who answered correctly to a given item at pre-test also answered correctly at 3 years. In multivariate models, higher education (p = 0.02) and older age (p = 0.04) were associated with a lower retention of information acquired during pre-test genetic counselling. However, women’s retention of information was not affected by test result, cancer status, psychological distress, or elapsed time between pre- and post-measurements.

**Conclusions:** Although knowledge acquired during a standard BRCA1/2 genetic counselling session could be improved, information correctly learned was remembered for an extended period of time by most participants.
P087 Is It Time for Population Genetic Testing for BRCA1 and BRCA2 in the Jewish Population? 
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Introduction: There are 2 founder mutations in BRCA1 and 1 founder mutation in BRCA2 that are present in up to 2.5% of Ashkenazi Jewish women. To date, it has not been proposed that the entire female Jewish population be eligible for screening. However, given the high frequency of mutations in the Jewish population at large, a full-population approach may be rational. Current guidelines for genetic testing in Ontario, Canada, stipulate the family history of cancer that must be present to confer eligibility for testing. Little is known about the appropriateness of these guidelines in the Jewish population and the acceptance of genetic testing in this group of women.

Methods: Eligible subjects were women who self-identified as (Ashkenazi or Sephardic) Jewish, who were between the ages of 25 and 80 years, and who resided in Ontario. Study subjects were recruited through a description of the study that was published in a national newspaper on a single occasion in May 2008. Women were asked to complete a study questionnaire and a family history questionnaire, and to provide a blood or saliva sample. Risks of mutation were estimated for each woman using in-house software.

Results: The overall mutation prevalence in the 2080 women who enrolled in the study was 1.1% (0.5% in BRCA1 and 0.6% in BRCA2). Of the 22 families, 10 (45%) would have met the criteria for genetic testing for BRCA1 and BRCA2 under current guidelines from the Ontario Ministry of Health and Long-Term Care. There were no differences in risk of mutation for those with and without a BRCA1 or BRCA2 mutation (3.9% vs. 1.2%, p = 0.23). However, estimates of mutation risk for women with a BRCA1 mutation were higher than for those without a mutation (7.6% vs. 1.2%, p ≤ 0.01–0.04). This was not observed for women with a BRCA2 mutation compared with those with no BRCA1 mutation (0.8% vs. 1.2%, p = 0.32).

Conclusions: The prevalence of BRCA mutations in unscreened Jewish women from Ontario, Canada, was lower than previously reported. However, many of the women would not have met the criteria for testing based on family history of cancer. Population screening for BRCA mutations should be considered for the Jewish population.

P088 Recruitment of a Population-Based Sample of Black Women for a Study of Inherited Breast Cancer Using a Cancer Registry-Based Approach 
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Objectives: We investigated BRCA1 and BRCA2 (BRCA) mutations in a population-based sample of black women diagnosed with breast cancer below the age of 50 years, because of the disproportionate numbers affected with early-onset breast cancer. Furthermore, because black people are typically underrepresented in genetic research studies, we sought to understand the factors associated with recruitment activities and study participation.

Methods: Following intutional research board approvals, the State Cancer Registry released demographic, clinical, and contact information on all eligible participants. State-mandated recruitment methods included 2 mailings, 3 weeks apart, with a telephone response card for patients who did not wish to be contacted by telephone. If no patient response was received within 3 weeks of the second mailing, a member of the study team contacted the patient by telephone. Participation involved a genetic counselling session over the telephone, and collection of blood or saliva for BRCA testing.

Results: Of the 209 eligible patients identified by the cancer registry, contact was established with 87, of whom 82 were eligible for study participation. The overall rate of interest in study participation was 80% (including 93% for passive follow-up and 68% for active follow-up), of whom 48 have been consented to date. There were no differences in the clinical and demographic characteristics of participants and nonparticipants.

Conclusions: This is the first study conducted through a State Cancer Registry in which the primary goal was to recruit participants for genetic counselling and testing for inherited breast cancer. Use of active follow-up methods was important in study recruitment. In contrast with many earlier studies, our results suggest that black women with early-onset breast cancer are interested in participating in studies of genetics and breast cancer, indicating that participation rates in black women may be enhanced by carefully considering methods to facilitate participation.

P089 Critical Evaluation of BRCA1/2 Mutation Detection Rates in Southwestern Ontario 
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Deleterious mutations in the BRCA1/2 genes have not been identified in high-risk families as often as originally expected. In clinical practice, predictions of BRCA1/2 mutations probability are sometimes made using various statistical models. When such models are applied to the same person, a wide range of probabilities is revealed. Although trying to provide best practice to patients, economic challenges also exist in the health care environment. For these reasons, the Cancer Genetics team at the London Health Sciences Centre critically evaluated our experiences regarding cancer risk assessment and our associated BRCA1/2 mutation detection rate. In Ontario in 2000, a provincial committee established 13 eligibility criteria for hereditary breast/ovarian cancer (HBOC) screening based on the medical literature at that time, offering gene analysis to individuals with a perceived 10% or greater risk of having a BRCA1/2 gene mutation. The methodology of this study included a review of all pedigrees, of eligibility criteria, and of genetic test results of individuals (n = 1270) who participated in index BRCA1/2 testing from 2000 to 2008. A consistent approach to determining which eligibility criteria were used in each case was pertinent to the analysis. The results revealed that one of the criteria (3 or more cases of breast and/or ovarian cancer, any age), representing 48% (608/1270) of all individuals tested, reported a 12.3% (75/608) overall mutation detection rate. However, a subset of this group (3 or more cases of breast cancer, age over 50 years), representing 31% (189/608) of this category, showed only a 2.6% (5/189) positivity rate. Using a conservatively priced in-house screening test ($1500), it takes approximately $56,000 to detect 1 mutation-positive individual in this subset. This finding could represent a significant cost savings to the current service testing program in Ontario. Our next goal is to determine if other cancer genetics programs in Ontario experience similar results in their own region, to provide further evidence to influence provincial decision-making.

P090 We Are On the Manchester Bandwagon—So How Are We Doing? 
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Background: The American Society of Clinical Oncology recommends offering BRCA1/2 mutation searching to cancer-affected individuals when the probability of identifying a mutation is 10% or higher (J Clin Oncol 1996;14:1730–6). In September 2007, our unit began prospectively triaging familial breast and/or ovarian cancer families to genetic testing when the probability of identifying a mutation is 10% or higher (J Med Genet 2005;42:439). This method was chosen because of its simplicity; the score can be calculated manually without the need for a computer, and the score can be easily recalculated in an outpatient setting after clarification of a family history.

Objectives: To retrospectively audit our unit’s BRCA1/2 mutation detection rate, based on the combined-MS.

Patients and Methods: Between early 1995 and April 30, 2009, our service initiated BRCA1 and BRCA2 genetic testing in 939 breast and/or ovarian cancer–affected probands. Using in-house clinical criteria, 827 of the 939 had initially been triaged to BRCA1/2 testing, and a retrospective combined-MS was calculated for these probands; 112 had been triaged to testing using a prospectively calculated combined-MS.

Results: Pathogenic BRCA1/2 mutations were detected in 152 of 939 probands tested (16.1%). When the combined-MS was considered, pathogenic BRCA1/2 mutations were detected in almost one quarter of probands with a combined-MS above 15 (136/566, 24.0%), but in less than 5% of probands with a combined-MS of 15 or lower (16/373, 4.3%). The probability of detecting a pathogenic mutation increased as the combined-MS increased, with the 10% mutation detection threshold reached at a combined-MS of 16 (based on a best-fit trend line, where R2 = 0.92).

Conclusions: Our experience suggests that it is appropriate to triage familial breast and/or ovarian cancer families to BRCA1/2 genetic testing when the combined Manchester score is more than 15.
Non-Acceptance of an Appointment at the Familial Cancer Clinic: A Preliminary Telephone Survey

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Purpose and Methods: A telephone survey was used to determine why some referred clients do not complete and return pre-clinic forms (PCFs—includes family history form, personal medical record consent, individual medical history form, and clinic location preference form), despite being informed that an appointment will be arranged only after the PCFs are returned.

Results: A group of 47 clients who failed to return PCFs more than 5 months after receiving them were contacted by telephone (7 men, 40 women). Of the 47, 29 (61%) were referred for assessment of familial breast/ovarian cancer. The non-responders gave 57 reasons for not returning PCFs. Most reasons related to difficulties completing the family history form (n = 35 (61%)). Other reasons included appointment not a priority because of ongoing cancer treatment (n = 4), concerns about genetic information (n = 4), not understanding why the referral was made (n = 2), recent death of a close relative (n = 2), and concerns about family confidentiality (n = 1).

Two thirds (n = 30 (63%)) indicated they wanted an appointment “now” and would return their PCFs (2 men, 28 women; 18 familial breast/ovarian). However, only 13 (43%) returned PCFs in the 6-month period after contact. Almost all reported difficulties completing the family history form (n = 28 (93%)).

One third (n = 17 (36%)) indicated that they would not return PCFs; 11 did not want an appointment, and 6 did not want an appointment, but would consider re-referral at a later date (5 men, 12 women; 11 familial breast/ovarian). Of the 17, 7 (41%) reported difficulties filling out the family history forms.

Conclusions: Almost 2 in 3 non-responders want an appointment, but have not returned their PCFs, primarily because of difficulties with the family history form. We need to explore ways to assist completion of family history forms while minimizing the impact on genetic counsellor workload and showing respect for passive indications by non-responders that they do not want an appointment.

To Test or Not to Test: The Influence of Ashkenazi Jewish Ethnicity

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Cancer genetic counsellors are well aware of the increased risk for individuals of Ashkenazi Jewish background carrying a BRCA1 mutation and of the fact that breast/ovarian cancer is more likely to have a genetic basis in Jewish than in non-Jewish families. What they may not be as familiar with is the existence of a community of mainly religious Jewish women who are resistant to genetic counselling/testing because of unique issues surrounding their religious beliefs.

Some religious women may find breast/ovarian cancer treatments and/or genetic testing present religious dilemmas above and beyond the emotional dilemmas that all women confront with cancer. When faced with a diagnosis of breast or ovarian cancer, these women have questions and specific psychosocial concerns that may not be understood by the general cancer community, including genetic counsellors and cancer support groups. Some of their concerns surround issues of community, the Mikvah (ritual bath), and cancer as a stigma affecting marriage prospects. These questions need to be addressed by sources knowledgeable in Jewish lore and practices. A national, not-for-profit organization was formed several years ago to address these unique psychosocial, genetic, and practical issues pertinent to Jewish women living with or at high risk of breast cancer. Acquaintance with this organization and an in-depth description of these issues will be the substance of this poster, so that professionals who meet with resistance on the part of their Jewish patients will have a better understanding of how to give them the support they need within the parameters of Jewish religious customs.

Pre-implantation Genetic Diagnosis for BRCA1/2: The Patients’ Perspective

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Objectives: BRCA1/2 carriers face a 50% chance that mutations will be transmitted to their offspring, and concern about passing on BRCA1/2 is a common reason for genetic testing. Pre-implantation genetic diagnosis (PGD) is a promising technology for BRCA carriers. Pre-implantation genetic diagnosis is a type of assisted reproductive technology that involves genetic analysis of embryos obtained through in vitro fertilization. Couples can transfer only mutation-free embryos to the mother’s uterus to initiate pregnancy. However, use of PGD for adult-onset cancer susceptibility is also controversial. As health care professionals increasingly become involved in discussions of family planning and reproductive technologies in their work with hereditary breast/ovarian cancer, the viewpoints of patients may provide insight for genetic counsellors and physicians attempting to navigate this ethically complex terrain. We report findings of a qualitative investigation of the views of reproductive-age BRCA1/2 mutation carriers regarding PGD, focusing on suggestions for integrating PGD into medical practice.

Methods: Reproductive-age patients (n = 22; 21 women, 1 man; 91% European American), recruited from the clinical genetics service of a national comprehensive cancer centre, participated. Participants view a brief presentation regarding PGD for BRCA1/2 mutations, followed by an in-depth interview about their attitudes toward using PGD to prevent transmission of BRCA1/2 mutations, including their suggestions for discussing PGD in the context of cancer-risk counselling. Interviews are recorded and transcribed verbatim, and analyzed using methods of grounded theory.

Results: Themes regarding information management—including timing, delivery, and depth—were discussed. Although most participants want some information about PGD, suggesting it is “at least glossed over” in the initial counselling session, many emphasized that information should be limited, maintaining the focus on the patient undergoing testing/counselling.

Conclusions: BRCA1/2 mutation carriers are interested in learning about PGD, physicians and genetic counsellors must be prepared to discuss this technology. Suggestions are provided for tailoring information to the needs and values of patients.

Ethical Aspects of Pre-implantation Genetic Diagnosis for BRCA1/2

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Objectives: Individuals with inherited breast and ovarian cancer face difficult ethical choices when making reproductive decisions. In pre-implantation genetic diagnosis (PGD), embryos created by in vitro fertilization are genetically tested and unaffected ones are implanted. This study aimed to determine if couples attempting PGD to exclude BRCA1/2 mutations faced ethical and clinical challenges different from those for other cancer predispositions.

Methods: Couples referred for PGD for cancer predisposition had a preliminary consultation to outline the process. Applications for treatment licenses were made to the Human Fertilization and Embryology Authority (HFEA) for each new disorder and for individual patients with BRCA1/2 mutations. Applications were also made for National Health Service funding.

Results: Between June 1, 1999, and May 31, 2009, 100 referrals were made for PGD for 15 different cancer-predisposing genes. Among 61 couples who opted for PGD, work-up was started for 41 couples who received funding. Overall, 18 couples of PGD were completed for 12 couples, resulting in the birth of 5 healthy singleton and 4 ongoing pregnancies (pregnancy rate of 50% (9/18) per PGD cycle started). The largest referral group was for breast and ovarian cancer attributable to BRCA1 (25 couples) and BRCA2 (6 couples). However, only 55% (11/13) from this group opted for PGD as compared with 87% (13/15) for NF1, 71% (12/17) for RB1, and 65% (11/17) for APC mutations. Patients with BRCA1/2 mutations refused PGD, because the time taken by the sequential process of funding, protocol development, and HFEA licensing interfered with their planned prophylactic surgery. Couples with children rarely chose to have PGD.

Conclusions: The uptake of PGD for BRCA1/2 is clearly much less than for other cancer predispositions. Our study shows that this finding has as much to do with treatment of the female partner as the fact that BRCA1/2 have incomplete penetrance as compared with other cancer predispositions.
Objectives: Most large genomic rearrangements (LGRs) in BRCA1 and BRCA2 are missed using conventional sequencing technologies. In the United States, LGRs can be found using the BRACAnalysis Large Rearrangement Test (BART) offered by Myriad Genetics. The BART is performed in parallel with DNA sequencing on individuals who meet clinical criteria set by Myriad Genetics. The BART can be ordered separately for individuals who do not meet the established criteria. We were interested in determining how many of our patients who carried an LGR in BRCA1 or BRCA2 met the criteria.

Methods: We performed a retrospective chart review of individuals seen at our center for BRCA1/2 testing after the introduction of this new technology. The family histories of individuals with an LGR were assessed and determined to meet or not meet the clinical criteria set by Myriad for inclusion of BART.

Results: In 5 individuals, an LGR was detected on BART. All LGRs were in the BRCA1 gene and included del(exon 5), del(exons 1–2), del (exons 19–20), and exon 14–20del26 kb (2 individuals). In 1 of the 5, cancer history clearly fulfilled the Myriad criteria. In 2 of the 5, cancer histories fulfilled the Myriad criteria only because the genetic counsellor obtained cancer histories on the individual’s great aunts, great uncles, and great grandparents. In 2 of the 5, the individual’s personal and family cancer histories did not meet the clinical criteria set by Myriad.

Conclusions: Although LGRs are expected to account for a small minority of mutations in BRCA1/2, the phenotypes of these LGRs are not well understood. Families harbouring an LGR do not necessarily meet Myriad’s stringent criteria. Testing using BART should be considered in all families who are offered BRCA1/2 testing.

Objectives: Bi-allelic mutations of the MYH gene, implicated in base excision repair, are known to cause adenomatous colorectal polyps and are associated with very high colorectal cancer risk. The extracolonic manifestations of MYH-associated polyposis coli have not been fully characterized. One previous report suggested a substantially increased breast cancer (bc) risk in female MYH mutation carriers (Nielsen et al. J Med Genet 2005;42:e54).

Methods: We describe an extended Canadian kindred in which a wide spectrum of tumour types occurred, including myeloma and colorectal, breast, bladder, and uterine cancers.

Results: Two previously recognized MYH mutations, G382X (1145G>A) and E182X (544G>T), were identified in the male proband who presented with bilateral bc and in whom BRCA1/2 testing was inconclusive. Of the remaining 2 sisters unaffected by bc, one died at age 47 with colon cancer. The MYH test results for the woman with bilateral bc and her sister who developed breast cancer at age 68, are pending and will be presented.

Conclusions: Our findings will contribute to the scant literature concerning bc risk in MYH mutation carriers. Further studies of the frequency of bc in mono-allelic and bi-allelic mutation carriers are warranted.

Objectives: The purpose of this study is to determine the prevalence of BRCA1 and BRCA2 mutations among women with carcinoma of the fallopian tube.

Methods: Two series of women diagnosed with carcinoma of the fallopian tube were studied. The first series of subjects was identified from the Ontario Cancer Registry. Those women were diagnosed between 1990 and 1998 or between 2002 and 2004. The second series was identified from Cedars Sinai Medical Centre, Los Angeles, California, U.S.A. They were diagnosed between 1991 and 2007. Each subject was approached to provide a family history and a blood sample for genetic testing for mutations in the BRCA1 and BRCA2 genes.

Results: We recruited 116 patients with fallopian tube cancer (70 from Ontario, 46 from Los Angeles). A deleterious mutation was found in 36 patients (31.0%), 26 in BRCA1 (22.4%) and 10 in BRCA2 (8.6%). A family history of ovarian or breast cancer was positive for 26 women; of these, 15 had a mutation (57.5%). In 19 of the patients with a previous history of breast cancer, 15 (79.0%) had a mutation. Of the women who were diagnosed with fallopian tube cancer before the age of 60 years, 40.0% had a mutation, as compared with 19.2% of the women diagnosed after the age of 60 years.

Conclusions: Approximately 31% of women with fallopian tube cancer have a mutation in BRCA1 or BRCA2. All patients diagnosed with invasive fallopian tube cancer should be considered candidates for genetic testing.
A Randomized Controlled Trial of Population Testing for BRCA1/2

**Objectives:** To compare a systematic population-based approach to genetic testing for cancer-predisposing genes with the current family history-based approach.

**Methods:** A randomized controlled trial for BRCA1/2 genes (disease model) in the Ashkenazi Jewish (AS) community (population model), with two arms: population screening (study arm) and family history (control arm). Primary outcomes include BRCA founder mutations (FMs) detected; acceptability; psychological impact; uptake of screening or preventive strategies; and health economics. Inclusion criteria are age over 18 years and AS ethnicity. Exclusion criteria are first-degree relationship to BRCA carrier and past BRCA testing. Recruitment depends on self-referral, and a variety of population-based strategies are used to increase awareness. All participants undergo genetic counselling (group or individual) before making a decision regarding testing. Consenting volunteers provide a blood sample and are subsequently followed at 7 days, 3 months, 1 year, 2 years, and 3 years after the test result to ascertain the benefits and disadvantages of testing. Follow-up questionnaires evaluate attitude, knowledge, satisfaction, psychological impact, quality-of-life, uptake of screening or preventive strategies, lifestyle behaviours, and health economics.

**Results:** We present data on our experience of initial participants from the pilot phase of 1000 volunteers. Of the 450 people recruited to date, 31% are men, and 69% are women (median age: 59 years; interquartile range: 45 to 66 years). The post-counselling participation rate exceeds 85%. The founder mutation rate is 2.2% (95% confidence interval: 0.55% to 5.04%). Of these participants, 350 have been randomized, and 3 carriers were detected in the systematic screening arm, and 1 was detected in the family history arm.

**Conclusions:** Initial experience suggests that it is feasible to undertake a randomized controlled trial for population-based BRCA1/2 gene testing. Preliminary results show the presence of carriers in individuals without a strong family history of cancer.
**Genetic Counselling: What Is Important to Spouses of BRCA1/2-Positive Women?**

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**Introduction:** A limited number of studies have examined the interests and preferences of non-affected spouses in the process of genetic testing and counselling for genetic disorders that have sex-limited expression. Our hypothesis was that non-affected spouses of BRCA1- or BRCA2-positive women would have specific preferences and needs that were unaddressed. This pilot study ascertained those items.

**Subjects and Methods:** A questionnaire was distributed to attendees of the Men’s Facilitated Networking Session, FORCE (Facing Our Risk of Cancer Empowered) meeting. All participants, but 1 male, were Caucasian (11/12); ages ranged from 27 years to 75 years (median age: 48 years). The sample was not socioeconomically representative of the general population: 92% were in professional occupations or retired; 100 had a bachelor’s degree; 42% had advanced graduate degrees.

**Results:** In the study group, 100% prioritized genetic counselling before testing and did not differentiate between determining a son’s or daughter’s risk; 92% desired discussion of insurance discrimination issues; 83% preferred receiving results with family members present; 75% felt that determining the cancer risk for self and offspring was very important, wanted a summary letter or additional counselling sessions without the spouse present, and felt that the sex of the health care provider was important; 41% felt that determining the cancer risk for self and offspring was very important, wanted a summary letter or additional counselling sessions without the spouse present, and felt that the sex of the health care provider was important; 25% desired an annual visit with a genetic counsellor.

**Conclusions:** Non-affected spouses have unmet preferences and needs. Important needs were insurance discrimination issues, receiving results in person, without the proband present, and felt that the sex of the health care provider was important; 25% desired an annual visit with a genetic counsellor.

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**The Lived Experiences of Women Possessing a Genetic Predisposition to Developing Breast Cancer: A Grounded Theory Study**

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**Objectives:** Genetic testing to determine whether a woman carries a BRCA1 or BRCA2 mutation, which may entail an up to 80% lifetime risk of developing breast cancer, is relatively recent. There is an apparent void in the literature investigating the lived experiences of these women, and qualitative studies of these individuals are virtually nonexistent. The purpose of this ongoing study is to better comprehend the lived experiences of young women who have been identified as carrying a BRCA1/2 mutation. In our study, concepts such as the nature of these women’s received support, the impact of the experience, and the presence of stigma are being examined.

**Methods:** Six women between the ages of 20 and 40 years who were identified as BRCA1/2 mutation carriers will be invited to participate in a semi-structured interview. The interview will be digitally recorded, transcribed, and analyzed through the use of constant comparison analysis (Glaser and Strauss, 1967).

**Results:** This preliminary report is based on the first completed interview. Three major dimensions emerged from 16 raw data themes. More specifically, the themes of Confirmation of Fears, Concern, and Coping were revealed. Generally, this woman was found to experience various negative emotions. The discovery of a positive test result was perceived to be a confirmation of previous fears, which then led to fears for her daughters and about judgments from others. Finally, the woman was able to engage in coping behaviours such as altering her way of life and seeking social support. Results for the remaining 5 planned participants will be presented.

**Conclusions:** The results of this qualitative study may lead to recommendations regarding the additional support and psychological assistance that should be considered in young women undergoing genetic testing for inherited predisposition to breast cancer.

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**Impact of a Hospital-Based Focused, Counsellor-Led Support Group Among BRCA Carriers**

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**Objectives:** Individuals carrying a mutation in the breast and ovarian cancer susceptibility genes BRCA1 and BRCA2 often face significant psychological and emotional challenges. To address the unique medical and psychological needs of BRCA carriers, we implemented a hospital-based, focused support group for BRCA mutation carriers led by genetic counsellors. This study evaluates the impact of such a support group on the knowledge, emotional well-being, and family dynamics of BRCA mutation carriers.

**Methods:** We invited 115 individuals carrying a BRCA mutation to participate in a voluntary electronic questionnaire about their experience in the support group. Participants were asked to rate the impact of the support group on medical knowledge, emotional needs, family dynamics, and overall benefit.

**Results:** The online questionnaire was completed by 42 individuals, 62% of whom were BRCA1-positive, 30% were BRCA2-positive, and 8% were BRCA-negative family members. Most of the respondents were between the ages of 40 and 55 years. Of the respondents, 85% felt that attending the support group significantly improved their medical knowledge; 70% felt that the group provided significant emotional support; and 96% felt that they were better able to broach the subject of their BRCA status with family members. Slightly more than half of respondents (56%) experienced significantly lowered anxiety levels regarding their BRCA status, and a preponderance (78%) reported diminished feelings of isolation since attending the support group.

**Conclusions:** Involvement in a focused, interactive BRCA-positive support group resulted in significant improvement in medical knowledge, family dynamics, and anxiety in the participants. Genetic counsellors with a specialty in cancer genetics proved to be effective in leading the sessions and achieving its goals. This study demonstrates that ongoing involvement in similar support groups provides BRCA carriers with important benefits in dealing with the many challenges faced by this unique group of individuals.

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**Decision-Making About Inherited Breast/Ovarian Cancer Risk: Exploring Dimensions of Genetic Responsibility**

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**Background and Objectives:** Because genetic information has implications for family members, some choices about genetic risk may be influenced by perceptions of responsibility to relatives. Drawing upon 25 semi-structured interviews with test recipients in Canada, this study explored decisions about inherited breast/ovarian cancer.

**Methods:** Semi-structured interviews.

**Results:** Qualitative data analysis revealed the pervasive significance of genetic responsibility in test decisions. We highlight three dimensions of genetic responsibility:

- To know about the self for self
- To know about the self for others
- To know about the self to oblige others to know

**Conclusions:** These dimensions of genetic responsibility have implications for test decisions, family relationships, and other family members’ desire to know (or not know) and to act (or not act) with respect to their own genetic risk. In particular, genetic responsibility may play out as a framing of a relative’s moral obligation to know their risk that could obviate any interest they might have in not knowing. We conclude that perceptions of responsibility to, and of, other family members be thoroughly explored in genetic counselling sessions.
Women who are notified that they carry a BRCA1/2 mutation are presented with surgical options to reduce their risk of breast and ovarian cancer, including risk-reducing mastectomy (RRM) and risk-reducing oophorectomy (RRO). There is growing evidence that a subgroup of women do not make decisions about RRM and RRO immediately following genetic testing; rather, they consider these decisions years later. Women’s perspectives on the timing of these decisions are not well understood. Accordingly, the purpose of this study was to describe how women construct the “right time” to consider decisions about RRM and RRO. In-depth interviews were conducted with 21 women who were BRCA1/2 carriers and were analyzed using qualitative, constant comparative methods. The study findings revealed how important it was for the women to make decisions about RRM and RRO one at a time. Moreover, the women constructed the “right time” to consider decisions about risk-reducing surgery to be when:

- decisions fit into their lives,
- they had enough time to think about decisions,
- they were ready emotionally to deal with decisions,
- all of the issues and conflicts were sorted out,
- there were better options available, and
- the health care system was ready for them.

These findings offer novel insights relevant to health care professionals who provide decision support to women considering RRM and RRO decisions.

**Objective:** To examine the challenges facing health care providers in deciding to test adult children (individuals 17 years of age and older) for a mutation in BRCA1/2.

**Methods:** A focus group was conducted consisting of 1 medical geneticist and 3 genetic counsellors, who all specialize in cancer genetics and who all have encountered a case in which a young individual requested testing for a mutation in BRCA1/2 known to run in the family. The session was audiorecorded. Themes were extracted and examined.

**Results:** Recognizing the different arguments in favour of and against testing young people based on theoretical benefits and harms, the group discussed their clinical experience and their case-based decisions. The adult children were mainly female and came with only 1 parent who was not necessarily the carrier. The prominent theme encountered was to elicit the extent to which the young person’s demand for the test was “voluntary,” that is, being “influenced” but not “coerced” by the parents. This assessment became more difficult by the value of autonomy based on a rational individual, traditional bias on psychological assessment, and the limitations of professional training for the changing demands in this select population. Sex differences in the parent’s perspectives on the daughter’s desire to seek testing and contradictory behaviour in relation to being tested or to receiving the result were also relevant themes for the group. An illustrative case in which the resource of a second opinion was used is also presented.

**Conclusions:** In the face of the “new generation” seeking predictive testing for BRCA1/2, this study illustrates the need for more research and training for the professionals who are more often involved with a complex family dynamic rather than an individual choice. Case-by-case decision based on second opinion or psychological assessment in more difficult family interactions can be adequate resources in some cases.
Factors Associated with Choosing Risk-Reducing Mastectomy in BRCA Mutation Carriers
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Objectives: Lifetime risk of breast cancer among BRCA mutation carriers is markedly reduced with prophylactic mastectomy (PM), yet only a subset elect to undergo the procedure. This study investigates factors that might influence patients’ decisions regarding risk-reducing surgery.

Methods: We conducted a prospective observational study of BRCA mutation carriers with no personal history of cancer presenting to Cedars Sinai Medical Center between 1999 and 2009. Charts were analyzed for factors that might have influenced choices regarding prophylactic surgery, including age, family history, history of breast biopsy, and history of previous cosmetic surgery.

Results: Of 59 BRCA carriers, 22 (37.3%) elected PM. Average time from results disclosure to PM was 1.4 years. In the PM group, as compared with the no-PM group, history of mother dying of breast cancer was 63.6% versus 5.4%, \( p < 0.001 \), and history of a mother dying of ovarian cancer was 13.6% versus 43.2%, \( p = 0.02 \). Of those who had PM, 96% as compared with 40.5% in the no-PM group had bilateral salpingo-oophorectomy (BSO), \( p < 0.001 \). Interestingly, 14 of 15 patients (93.3%) who had BSO in the no-PM group had a history of mother dying of ovarian cancer. No significant differences in mean age at the time of genetic testing, number of affected first-degree relatives, history of breast biopsy, or history of cosmetic surgery were detected between the groups.

Conclusions: BRCA carriers with family history of a mother dying of breast cancer were most likely to choose PM. Among those who did not undergo PM, uptake of BSO was highest among those whose mother died of ovarian cancer. The impact of having a mother die of cancer may be of under-recognized importance.

High Satisfaction Rates in Women after Deep Inferior Epi gastric Artery Perforator Flap Breast Reconstruction
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Objectives: Breast reconstruction (BR) is aimed at improving quality of life (QoL) after mastectomy. Patient satisfaction is an important indicator in evaluating the success of BR. This study explored patient satisfaction and its determinants in women undergoing deep inferior epigastric artery perforator ( DIEP ) flap BR and the impact of the procedure on body image, sexuality, and QoL.

Methods: Patient satisfaction and QoL was studied in 72 women who underwent DIEP flap BR, using a study-specific questionnaire and the Short Form 36.

Results: Patient satisfaction was very high. Approximately 90% of patients reported that they had been sufficiently informed about the procedure and its consequences, that their preoperative expectations had been met, that the reconstructed breast felt like their own, that they would choose the same procedure again, and that they would recommend this procedure to a friend. Patient satisfaction was positively and significantly related to the reconstructed breast or breasts feeling like their own. Women with secondary reconstructions were more positive about changes in sexuality and femininity than were women with primary BR. There were no clinically relevant differences in QoL between our study population and a random sample of Dutch females.

Conclusions: Women with DIEP flap BR reported high satisfaction rates. However, to compare these satisfaction rates with other forms of BR, a prospective study in comparable groups is currently ongoing.

Opening the Psychological Black Box in Genetic Counselling: Recollections and Interpretations of Counselees
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Objectives: It is often assumed that the communication of DNA test results has direct psychological and medical consequences for the counselee. However, subjective factors may mediate between DNA test result and impact. The present study assessed mediation effects of the counselee recollections and interpretations regarding cancer risks and the likelihood that the cancer in their family is heritable.

Methods: In this retrospective study, women tested for BRCA1 or BRCA2 completed questionnaires 5 years after disclosure. Participants had received an unclassified-variant test result ( \( n = 76 \) ), uninformative-negative DNA test results ( \( n = 76 \) ), or pathogenic mutations ( \( n = 51 \) ) in BRCA1/2.

Results: The counselees, 10%–42% correctly recalled and interpreted what the counsellors actually communicated about cancer risks and heredity likelihood. Cancer risks were recalled and interpreted as lower, heredity likelihood as higher. Moderate correlations between recalling and interpreting suggested distinctive processes. Cancer risks were recalled more correctly when measured in percentages instead of in categories. Unclassified DNA variants were recalled and interpreted as giving very high cancer risks and heredity likelihood, uninformative results as very low. Psychological and medical outcomes were predicted by cancer risks and heredity likelihood as actually communicated by counsellors. However, these predictions were lowered or became insignificant when the recollections and interpretations of the counselees were included in analyses and path analyses, suggesting mediation. Overestimations predicted life changes, breast/ovarian removal in all counselees, and low quality of life in unclassified variants.

Conclusions: The impact of test results is strongly mediated by the recollections and interpretations of communicated risks by counsellors. Subjective processes should be addressed during counselling, especially if unclassified-variants are disclosed.