Mainstreaming Genetic Testing for Epithelial Ovarian Cancer by Oncology Providers: A Survey of Current Practice

by

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Abstract

With significant deficits in early detection and poor treatment response, ovarian cancer is a devastating diagnosis for many women. Up to 25% of epithelial ovarian cancer (EOC) is due to a hereditary predisposition, most commonly in \textit{BRCA1} and \textit{BRCA2}. Pathogenic variants in \textit{BRCA1}, for example, confer a 45% lifetime risk of EOC; whereas, the general population risk is only 1-2%. Knowledge of an affected individual’s genetic status can have significant implications for treatment and prognosis. Women with pathogenic variants in \textit{BRCA1/2} have an enhanced response to platinum-based chemotherapy and poly ADP-ribose polymerase (PARP) inhibitors, leading to an improved prognosis. Identifying individuals who harbor pathogenic variants in ovarian cancer predisposition genes is therefore of critical importance.

Current National Comprehensive Cancer Network (NCCN) guidelines recommend that all individuals diagnosed with EOC be offered germline genetic testing. While this would ideally be performed by genetics professionals, a shortage of genetic counselors precludes timely access to these services. This study sought to investigate the current genetic testing practices of oncology providers in order to determine the feasibility of oncologist-led genetic testing for patients with EOC. A survey was distributed to members of the Society of Gynecologic Oncologists with questions regarding timing, frequency, and type of genetic testing, referrals to genetics professionals, confidence with aspects of genetics services, and any barriers that currently hinder these processes. Results of the study were encouraging, with the majority of providers always
ordering genetic testing for patients with EOC; testing was most commonly ordered at diagnosis and was typically multi-gene panel testing that included *BRCA1/2*, consistent with current recommendations. Provider confidence with the genetic testing process was generally high, especially for deciding which patients to refer to genetics professionals. Patient disinterest and concerns for insurance coverage were commonly cited barriers to testing and referrals. Thus, oncologist-led genetic testing for patients with EOC, with referrals to genetics professionals when appropriate, has the potential to be a viable alternative service delivery model and warrants additional investigation.
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Preface

I would like to extend my gratitude to my committee members for their hard work and dedication to this study. Rachelle Huziak, my committee chair, Dr. Andrea Durst, and Dr. Phuong Mai assisted me on numerous occasions with their expertise in clinical cancer genetics. I would also like to offer thanks to Dr. Sarah Taylor, who was instrumental in the distribution of the survey. Finally, I would like to thank my family, friends, and colleagues for their support and encouragement.
1.0 Introduction

Ovarian cancer is the 5th leading cause of cancer death for women in the United States, as difficulties in early detection and treatment response often lead to a poor prognosis. Most ovarian cancers are not diagnosed until advanced stages, where the 5-year survival rate is only 29%.\(^1\) Importantly, 20-25% of epithelial ovarian cancer (EOC) is related to a hereditary predisposition.\(^2\) The majority of pathogenic variants occur in \textit{BRCA1} and \textit{BRCA2}. Mismatch repair genes related to Lynch syndrome such as \textit{MLH1}, \textit{PMS2}, \textit{MSH2}, \textit{MSH6}, and \textit{EPCAM} also contribute to ovarian cancer cases, as well as other moderate-risk genes.\(^3\) Due to the high rate of germline pathogenic variants, current NCCN, ASCO, and SGO guidelines recommend that all women diagnosed with EOC be offered genetic testing.\(^4-6\)

For affected individuals, standard-of-care treatment for late-stage ovarian cancer is surgery and platinum-based chemotherapy. Women with pathogenic variants in \textit{BRCA1/2} or other genes that disrupt homologous recombination, whether germline or somatic, have favorable response to poly ADP-ribose polymerase (PARP) inhibitors.\(^7\) These women therefore often have an improved prognosis.\(^8\) Similarly, patients with pathogenic variants in mismatch repair genes have targeted immunotherapy treatment options and often have improved survival compared to most patients.\(^9\) Knowledge of genetic status is therefore critical in tailoring treatments and informing prognosis.

Current screening methods for ovarian cancer are ineffective and do not improve survival of patients.\(^10,11\) For women at a significantly increased risk of developing EOC, risk-reducing surgery is the most effective preventative measure.\(^12\) Germline genetic testing is therefore essential in order to identify at-risk individuals; tumor testing alone is not sufficient. Many of the genes implicated in ovarian cancer predispositions have additional cancer risks where additional
screening and risk-reducing interventions could be considered.\textsuperscript{4,13} Pathogenic variants in the \textit{BRCA1}/2 genes substantially increase the risk for breast cancer and have additional risks for male breast, prostate, pancreatic, and melanoma cancers.\textsuperscript{14} In the mismatch repair genes, pathogenic variants result in significantly increased risks for colorectal and endometrial cancers and slightly increased risks for skin, gastric, bile duct, small bowel, and pancreatic cancers.\textsuperscript{13} Not only does this knowledge help to inform care for the patient, but at-risk family members can also be evaluated to clarify if their risks are elevated or at the general population level. Germline testing for ovarian cancer predisposition genes is therefore useful in informing patients and their families of potential inherited cancer risks.

Unfortunately, there is limited capacity for patients to be seen by a genetics healthcare professional, and long wait times can seriously hinder appropriate testing and counseling for individuals.\textsuperscript{15} Certain patient characteristics have also been shown to be associated with less genetic testing, including Black race, greater poverty, and less insurance.\textsuperscript{16} A recent study found that only 35\% of women with ovarian cancer have received genetic testing despite current guidelines.\textsuperscript{17} Comprehensive knowledge of the factors that are limiting genetic testing of patients with EOC are still not fully understood. In light of this issue, there has been a call to mainstream germline genetic testing by oncologists for patients with EOC. Studies in the United Kingdom have shown favorable outcomes with this service delivery model including high patient and provider satisfaction, rapid turnaround time, and appropriate referrals.\textsuperscript{18,19} It is not only critical that oncologists order appropriate testing and utilize the results to guide treatment, but they must also refer patients with positive or unclear results to genetics services to ensure comprehensive care.

This study sought to investigate the current practices for genetic testing in patients diagnosed with EOC by oncology providers. A survey was sent to members of the Society of
Gynecologic Oncologists, including oncologists, physician assistants, and nurses. Information collected included the type, frequency, and timing of genetic testing as well as referrals to genetics providers and questions to elucidate barriers to testing and referrals.

The specific aims of this study were as follows:

a) To assess the current practices of oncology providers regarding the ordering and use of genetic testing in patients with epithelial ovarian cancer

b) To identify the frequency with which oncology providers make referrals to genetics professionals for their patients

c) To evaluate the prevalence and types of genetics education that oncology providers have received

d) To identify the comfort level of oncology providers with different aspects of cancer genetics services

The results of this study will help to identify successes and barriers that currently exist in the United States regarding the mainstreaming of genetic testing for patients with EOC. Future studies can further investigate ways to streamline this pathway, address any issues that exist, and improve the patient experience.
2.0 Literature Review

2.1 Ovarian Cancer

In 2018, the World Health Organization (WHO) estimated the incidence of ovarian cancer diagnoses to be 295,414 globally; at the same point in time, there were 184,799 new ovarian cancer deaths.\textsuperscript{20} In the United States alone, the incidence of ovarian cancer is 14.8 in 100,000 women.\textsuperscript{21} Most women are diagnosed after menopause; it is rare for women under the age of 40 to have ovarian cancer.\textsuperscript{22} As the 5\textsuperscript{th} leading cause of cancer death in women, the 5-year survival rate for invasive ovarian cancer at all stages is 47%. For metastatic stages—which compromises 59% of diagnoses—the 5-year survival rate drops to 28%.\textsuperscript{1}

2.1.1 Screening and Surveillance

A deficit in the ability to detect ovarian cancer early results in most women being diagnosed at later stages of the disease, contributing to a poor prognosis. When still localized, the 5-year survival rate of ovarian cancer is 92%.\textsuperscript{1} Vague and non-specific symptoms such as frequent urination, feeling full quickly, abdominal pain, and bloating can arise, but these often do not present until advanced stages of the disease, if at all.\textsuperscript{10} Attempts to detect ovarian cancer earlier through screening measures have been studied. A randomized control trial with 78,216 average-risk women either receiving normal gynecologic care or an annual transvaginal ultrasound combined with cancer antigen 125 (CA-125) screening did not result in a decrease in mortality nor a stage shift. Additionally, the screening program had a false positive rate of 5%, and the invasive
diagnostic follow-up procedure resulted in serious medical complications in 15% of the study participants.\textsuperscript{10}

Some screening studies have found evidence of a slight increase in early detection of ovarian cancer. A randomized control trial with average-risk women in Japan found that a higher proportion of women were diagnosed at Stage I in the screening group (63%) compared to the control group (38%), but this difference was not statistically significant.\textsuperscript{23} Additional studies have examined serial CA-125 screening with scores using the Risk of Ovarian Cancer Algorithm (ROCA) in either average-risk women or women with a significant family history or a BRCA1/2 pathogenic variant.\textsuperscript{11,24-26} ROCA utilizes CA-125 data from thousands of women to determine the risk of having ovarian cancer based on CA-125 fluctuations. The researchers obtained a baseline CA-125 level for each woman, and ROCA was recalculated with every serial CA-125 value. If ROCA demonstrated an elevated risk for ovarian cancer, then a transvaginal ultrasound was performed. There was no reduction in mortality for average-risk women, and the program was not cost-effective.\textsuperscript{11,25,26} For high-risk women, there was a significant increase in the detection of early-stage ovarian cancer compared to historical controls as well as a low false positive rate, but the data are not sufficient to replace the current recommendation for high-risk women to undergo a risk-reducing salpingo-oophorectomy (RRSO).\textsuperscript{24}

\textbf{2.1.2 Histological Subtypes}

Ovarian cancer is a heterogenous disease with different histologic subtypes. First, ovarian cancer types can be broken down by the cell from which it originates: epithelial (90% of cases), sex-cord stromal, germ cell, and mixed-cell type.\textsuperscript{27} Epithelial ovarian cancer can be further broken down into multiple histological subtypes (Table 1).\textsuperscript{27-31} Information on the histological subtype is
pivotal to understand relevant risk factors and best treatment approaches. The WHO classifies ovarian tumors into one of two types. Type I tumors notably include low-grade serous, endometrioid, clear cell, and mucinous carcinomas; these tumors are typically low grade and have a slow progression. Rarely, type I tumors can be malignant Brenner or seromucinous. Somatic variants in these tumors typically involve $KRAS$ and $BRAF$.\textsuperscript{27} The type II category encompasses high-grade serous carcinomas, carcinosarcomas, and undifferentiated carcinomas. These tumors often progress rapidly and aggressively, and they are likely to carry somatic variants in $TP53$ as well as somatic or germline variants in $BRCA1/2$.\textsuperscript{2,27,32}

<table>
<thead>
<tr>
<th>Type</th>
<th>Histological Subtype</th>
<th>Estimated Prevalence</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Low-grade serous</td>
<td>3.5%</td>
</tr>
<tr>
<td></td>
<td>Endometroid</td>
<td>11%</td>
</tr>
<tr>
<td></td>
<td>Clear cell</td>
<td>12%</td>
</tr>
<tr>
<td></td>
<td>Mucinous</td>
<td>3.5%</td>
</tr>
<tr>
<td></td>
<td>Other</td>
<td>2%</td>
</tr>
<tr>
<td>II</td>
<td>High-grade serous</td>
<td>68%</td>
</tr>
<tr>
<td></td>
<td>Other</td>
<td>Rare</td>
</tr>
</tbody>
</table>

2.1.3 Risk and protective factors for epithelial ovarian cancer

For a woman living in the United States, the average lifetime risk of developing epithelial ovarian cancer is 1-2%.\textsuperscript{33,34} However, there are several factors that are known to increase or decrease the risk of EOC. Li et al. (2015) created an epidemiologic risk prediction model using data from 202,206 Western European women. Although this model has never been validated, it does provide insight into potential risk and protective factors, many of which have been supported by other studies.\textsuperscript{35} Certain factors have only been shown to have an association with particular
histological subtypes, which emphasizes the importance of understanding the etiology of ovarian cancer subtypes (Table 2).

### Table 2 Risk and protective factors for invasive epithelial ovarian cancer

<table>
<thead>
<tr>
<th>Evidence Level</th>
<th>Risk Factor</th>
<th>Protective Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strong</td>
<td>Family history of ovarian cancer(^{34,36})</td>
<td>Parity(^{35,40})</td>
</tr>
<tr>
<td></td>
<td>Hormone replacement therapy(^{35,37})</td>
<td>Oral contraceptive use(^{35,40,41})</td>
</tr>
<tr>
<td></td>
<td>Endometriosis (clear-cell, low-grade serous, endometroid)(^{38})</td>
<td>Tubal ligation(^{42})</td>
</tr>
<tr>
<td></td>
<td>Increasing height(^{39})</td>
<td>Bilateral salpingo-oophorectomy(^{43,44})</td>
</tr>
<tr>
<td></td>
<td>Higher BMI(^{35,39})</td>
<td></td>
</tr>
<tr>
<td>Moderate</td>
<td>Older age at menopause(^{35,45})</td>
<td>Older age at last birth(^{49})</td>
</tr>
<tr>
<td></td>
<td>Younger age at menarche(^{45})</td>
<td>Breastfeeding(^{45,50})</td>
</tr>
<tr>
<td></td>
<td>Diabetes mellitus(^{46})</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Smoking (mucinous)(^{47})</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Genital powder(^{48})</td>
<td></td>
</tr>
<tr>
<td>Insufficient</td>
<td>Incomplete pregnancy(^{51-53}), infertility drugs(^{54,55}), polycystic ovary syndrome (PCOS)(^{56}), pelvic inflammatory disease (PID)(^{57}), alcohol(^{58})</td>
<td>Multiple births(^{59}), hysterectomy(^{43,45})</td>
</tr>
</tbody>
</table>

For family history, the increase in risk for EOC depends on the degree of relation to the affected family member, as well as the number of family members affected. A woman with an affected first degree relative is about three times more likely to develop ovarian cancer than a woman without this family history.\(^{34,36}\) If a woman has more than one affected first-degree relative, she is 10 times more likely to develop ovarian cancer.\(^{60}\) BRCA1/2 pathogenic variants are estimated to account for a quarter of the familial relative risk for first-degree relatives. Other factors that increase familial relative risk include having affected relatives with EOC diagnosed under 50 years old and affected relatives with serous histology.\(^{36,60}\)
Surgical removal of the ovaries is currently recommended for women at an increased risk of ovarian cancer because it is the strongest protective factor.\textsuperscript{4} RSSO has been shown to dramatically reduce the risk of ovarian cancer in multiple studies, with risk reduction estimates around 80%.\textsuperscript{43,44} Oral contraceptive use is also an established protective factor. A meta-analysis that combined results from 45 epidemiological studies found that the use of oral contraceptives reduces a woman’s risk for ovarian cancer by 27% compared to women who reported never using them. Moreover, the longer the duration of use, the lower her risk; taking oral contraceptives for at least five years reduces the risk of developing ovarian cancer by approximately 50%.\textsuperscript{41} Further research is still needed to clarify risk estimates with knowledge of genetic predispositions and establish more evidence for factors that are currently not well supported.

2.2 Genetic Predispositions to Ovarian Cancer

Several genes have been identified to be associated with an increased susceptibility to EOC.\textsuperscript{61-65} It is currently estimated that about 20-25% of ovarian cancer is due to a hereditary predisposition.\textsuperscript{2,66} \textit{BRCA1} and \textit{BRCA2} pathogenic variants account for the majority of hereditary ovarian carcinomas, followed by pathogenic variants related to Lynch syndrome.\textsuperscript{3,9,61,62,66} Other genes have been shown to contribute moderately to EOC risk such as \textit{RAD51D}, \textit{RAD51C}, \textit{BRIP1}, \textit{BARD1}, and \textit{PALB2}, among others.\textsuperscript{63,64} Germline genetic testing can identify unaffected individuals who would benefit from preventative strategies; it can also help to direct treatment, inform prognosis, and elucidate additional beneficial screening for affected individuals. Current NCCN guidelines recommend that any woman with a diagnosis of epithelial ovarian, fallopian tube, and/or peritoneal cancers be offered germline genetic testing.\textsuperscript{7}
2.2.1 \textit{BRCA1} and \textit{BRCA2}

Both \textit{BRCA1} and \textit{BRCA2} are associated with the autosomal dominant Hereditary Breast and Ovarian Cancer (HBOC) syndrome. These genes are necessary for double-stranded DNA breaks to be repaired by homologous recombination.\textsuperscript{61} Several studies have aimed to quantify the frequency of \textit{BRCA1/2} germline pathogenic variants in women with ovarian cancer. A case-control study in Australia with 1,001 women found germline \textit{BRCA1/2} pathogenic variants in 14.1\% of patients with non-mucinous ovarian carcinomas. When only looking at patients with high-grade serous histology, this prevalence increased to 22.6\%.\textsuperscript{61} A prospective cohort study of 104 patients with EOC found that 21.15\% of patients carried \textit{BRCA1/2} pathogenic variants, and patients with high-grade serous histology had a higher prevalence (25.7\%) of these variants.\textsuperscript{67} One study specifically sought to estimate the prevalence of \textit{BRCA1/2} variants in an ethnically diverse sample of 585 patients with EOC and found that 22.5\% carried a pathogenic variant, again with a higher prevalence (27.6\%) in those with a serous histology.\textsuperscript{68} Importantly, not all patients identified to have a pathogenic variant in \textit{BRCA1/2} had a reported family history of breast or ovarian cancer. 44\% of patients in the Australia study had no suggestive family history at all.\textsuperscript{61} This study, and many others, are consistently finding that family history is not sufficient to predict pathogenic variant status; this supports current NCCN guidelines to offer genetic testing to any woman with epithelial ovarian cancer, regardless of family history.\textsuperscript{3,7,61,62,66-69}

For the general population, the lifetime risk of ovarian cancer is only 1-2\% with an average age of diagnosis of 63 years old.\textsuperscript{22,33,34} In the largest prospective cohort study to date, researchers estimated age-specific risks for breast, ovarian, and contralateral breast cancer in \textit{BRCA1/2} pathogenic variant carriers. The cumulative lifetime risk (to age 80) of ovarian cancer for a \textit{BRCA1} carrier was 44\%, while for \textit{BRCA2} it is around 17\%.\textsuperscript{70} The average age of diagnosis for ovarian
cancer also differs between the two genes, with BRCA1 pathogenic variant carriers typically being diagnosed at younger ages. BRCA1 carriers have a median age of diagnosis of ovarian cancer of 54 years compared to approximately 60 years for BRCA2. Importantly, other cancer risks exist for BRCA1/2 pathogenic variant carriers. Perhaps most notable is the lifetime breast cancer risk, which can be as high as 87%; other associated cancers include male breast, prostate, pancreatic, and melanoma. The benefit of identifying BRCA1/2 germline pathogenic variants is thus apparent, as surveillance and risk-reducing strategies could detect cancer at earlier stages or decrease the risk of developing many of these cancers.

2.2.2 Mismatch repair genes

Germline pathogenic variants in certain mismatch repair (MMR) genes—MLH1, PMS2, MSH2, and MSH6—and EPCAM are associated with Lynch syndrome, another autosomal dominant cancer predisposition syndrome. While the main cancer risks are colorectal and endometrial, there is an increased risk of ovarian cancer as well, along with skin, gastric, bile duct, small bowel, and pancreatic cancers. One study that sequenced germline DNA from 1,915 women with ovarian cancer found that 0.4% of ovarian cancer patients had pathogenic variants in MMR genes. The lifetime risk of ovarian cancer depends on the specific MMR gene that is mutated. MLH1 and MSH2 have been shown to have higher lifetime risks than MSH6 and PMS2. In a recent multicenter prospective observational study, the cumulative incidence of ovarian cancer was 11% for MLH1 pathogenic variant carriers, 17.4% for MSH2, 10.8% for MSH6, and 3% for PMS2.

Ovarian cancer is typically diagnosed between ages 42 and 49 for patients with Lynch syndrome. One study that examined data on more than 800 women from a Lynch syndrome
registry and cohort study calculated a median age of onset of 46 years for ovarian cancer; the range of age of onset was wide, with the youngest woman being 20 years old and the oldest being 75.\textsuperscript{9} Tumor histology is most often endometroid, although serous carcinomas are also associated with Lynch syndrome.\textsuperscript{9,73,74} The survival rate of ovarian cancers related to Lynch syndrome is high—likely in part due to the difference in tumor histology characteristics—with studies finding a 10-year survival of 84-87\%.\textsuperscript{13,73}

2.2.3 Other genes

Pathogenic variants in other genes involved in homologous recombination have been identified to increase a woman’s risk of EOC. Recently identified genes include RAD51C, RAD51D, BRIP1, and PALB2.\textsuperscript{63,64,75-77} One study examined pedigrees from 480 families with a history of breast and ovarian cancer that had negative BRCA1/2 testing results. The researchers found that 1.3\% of families had a pathogenic variant in RAD51C segregating with disease.\textsuperscript{76} Another study examined 911 BRCA1/2-negative families with breast and ovarian cancer and found that 0.9\% carried pathogenic variants in RAD51D. The relative risk of ovarian cancer for pathogenic variant carriers was estimated to be 6.3 times population risk, while the relative risk for breast cancer was 1.3 and not statistically significant.\textsuperscript{75} Pathogenic variants in BRIP1 increase risk mainly for high-grade serous EOC, although the specific risk is not currently well-understood. Older ages of diagnosis are also associated with BRIP1 pathogenic variants.\textsuperscript{63} One study found that certain frameshift BRIP1 pathogenic variants significantly increased the risk for ovarian cancer, while more common missense variants were not associated with increased risk.\textsuperscript{77} Finally, PALB2 pathogenic variants are known to confer breast cancer susceptibility, but more studies are
finding that ovarian cancer risk may be increased as well. Further research is needed to better
determine the estimated ovarian cancer risk associated with pathogenic variants in this gene.63,65

Massively parallel sequencing has identified other potential ovarian cancer susceptibility
genes, but more studies are needed to fully understand their penetrance and risks. One study
utilized this sequencing technology with 360 women with ovarian, fallopian tube, or peritoneal
cancers and found germline pathogenic variants in 23% of them. 18% of pathogenic variants were
in BRCA1/2, and the other 6% were in MSH6, RAD51C, BRIP1, PALB2, TP53, BARD1, CHEK2,
MRE11A, NBN, and RAD50. At the time of this study, RAD51D had not been implicated in ovarian
cancer predisposition and was therefore not included. All genes identified, with the exception of
MSH6 and TP53, are involved in the homologous recombination pathway. Importantly, the women
with MSH6 and TP53 germline pathogenic variants did not meet criteria for Lynch syndrome or
Li-Fraumeni syndrome, respectively. 30% of women in the study had no family history of breast
or ovarian cancer, again pointing to the need for testing without such a history.3

2.3 Ovarian Cancer Genetic Testing

In the field of cancer, genetic testing can be ordered for a variety of reasons. Regardless of
whether an individual is affected, genetic testing can increase knowledge about personal and
family cancer risks. It can provide an explanation for a personal or family history of cancer if a
genetic pathogenic variant is found. For affected individuals, genetic testing can result in
personalized treatment. Unaffected persons may benefit from enhanced surveillance or options for
risk reduction. Sometimes, genetic testing can provide reassurance that there is likely not a single-
gene predisposition to cancer in the family.
2.3.1 Utility of ovarian cancer genetic testing

For patients with ovarian cancer, genetic testing can inform prognosis and guide treatment. Tumor testing alone can provide this information, but given the high frequency of germline pathogenic variants in patients with EOC, it does not provide all needed information. Germline results must be delineated in addition to somatic results to provide the patient and their family with the most comprehensive medical information. BRCA1/2 testing was historically more commonly used, but with the advent of next-generation sequencing, multi-gene panel testing has become both feasible and necessary and has been utilized more frequently in recent years.

2.3.1.1 BRCA1/2: Management and Screening

Primary treatment of ovarian cancer is typically platinum chemotherapy and surgery to remove the tumor, followed by maintenance therapy. A study comparing ovarian cancer outcomes in carriers of BRCA1/2 pathogenic variants with non-carriers found an improved prognosis and a better response to platinum-based chemotherapy. The presence or absence of a BRCA1/2 pathogenic variant guides the selection of and provides information on the magnitude of benefit of maintenance therapy as well. Maintenance therapy is typically achieved through poly ADP-ribose polymerase (PARP) inhibitors, which prevent cancer cells from repairing their damaged DNA and can help to trigger apoptosis. Until recently, PARP inhibitors were only recommended for metastatic treatment of homologous recombination (HR) deficient tumors—such as a BRCA1/2 mutated tumor—after at least three other lines of therapeutic therapy had been utilized. The PARP inhibitor Niraparib is now FDA-approved as first-line maintenance therapy after response to platinum-based chemotherapy for late-stage ovarian cancer, regardless of the HR status of the tumor. It is important to note, however, that the response to Niraparib was significantly better for
HR deficient tumors. For example, in the clinical trial, the median progression-free survival was 21.9 months for HR deficient tumors compared to 13.8 months in the overall population; this value for the placebo group was 10.4 months.\textsuperscript{8} Knowledge of \textit{BRCA1/2} tumor status is therefore essential for treatment and prognosis purposes.

As mentioned previously, tumor testing alone is not sufficient due to the high prevalence of germline pathogenic variants in patients with ovarian cancer.\textsuperscript{67} Identifying somatic mutations in the tumor provides information for treatment of the tumor only; identifying germline mutations present in the entire body provides information for screening and risk reduction of all cancers associated with the mutation. Unaffected relatives also greatly benefit from genetic testing for ovarian cancer susceptibility genes. If a pathogenic variant is identified in an affected relative, cascade testing permits the testing of blood relatives for the same variant to identify those at increased risk for EOC. As there are currently no effective screening methods, the main benefit of genetic testing lies in risk reduction.\textsuperscript{12,24} For women with pathogenic variants in \textit{BRCA1/2} who are unaffected with ovarian cancer, it is recommended that they undergo a risk-reducing salpingo-oophorectomy once childbearing is completed, ideally between the ages of 35-40 for \textit{BRCA1} and between the ages of 40-45 for \textit{BRCA2}. If there is an ovarian cancer diagnosis in the family prior to the recommended age for RRSO, then earlier prophylactic surgery should be discussed.\textsuperscript{4} Although this procedure significantly reduces the risk of ovarian cancer, and possibly breast cancer, it does incur some risks associated with early menopause, such as osteoporosis and heart disease.\textsuperscript{82,83}

There are guidelines in place for the additional cancer risks associated with \textit{BRCA1/2} pathogenic variants as well. Note that while these recommendations are standard, they may or may not be indicated in patients with EOC, depending on their clinical picture. Breast cancer screening
begins at age 25 with an annual MRI. Beginning at age 30, a mammogram should occur annually
with the MRI. The option of a risk-reducing bilateral mastectomy should be discussed. Pancreatic
screening may also be justified, although data for its efficacy are limited.\textsuperscript{4} The American
Gastroenterological Association currently recommends screening in individuals with pathogenic
variants in \textit{BRCA1/2} only if there is a first degree relative with pancreatic cancer.\textsuperscript{84}

\textbf{2.3.1.2 Mismatch repair genes: Management and Screening}

Women diagnosed ovarian cancer who have mismatch repair deficient (dMMR) tumors
may have tailored therapy options. While there are not as many targeted therapy options for non-
\textit{BRCA1/2} pathogenic variants in ovarian cancer tumors, a growing body of evidence is suggesting
that targeted therapy is warranted for dMMR tumors.\textsuperscript{85} Pembrolizumab is a specific
immunotherapy, for example, that has been shown to have increased efficacy in dMMR ovarian
cancer tumors. There are numerous clinical trials available for patients with dMMR tumors, and
thus targeted therapies will likely become more widely available in the future. Patients with dMMR
tumors have also been shown to have improved survival, making it a useful marker for prognosis.\textsuperscript{9}
For unaffected women with pathogenic variants in \textit{MLH1} and \textit{MSH2}, risk-reducing RR\textit{S}SO is
recommended; for individuals with pathogenic variants in \textit{PMS2}, \textit{EPCAM}, or \textit{MSH6}, there is
currently insufficient evidence to recommend RR\textit{S}SO.\textsuperscript{86}

There are additional screening and management guidelines for germline MMR pathogenic
variants, which may or may not be applicable to individuals affected with ovarian cancer. The two
largest risks and most well-defined screening guidelines are for colon and endometrial cancer.\textsuperscript{13}
For men and women, colonoscopies should begin between the ages of 20 and 25 (or 2-5 years prior
to the earliest colon cancer in the family, if earlier) and should be repeated every 1-2 years.\textsuperscript{86} Daily
aspirin use has been shown to reduce colorectal cancer risk, although more data is needed in order
to determine an optimal dosage and duration of use. Pancreatic cancer screening is recommended only if an individual has an affected first-degree relative. Many women affected with ovarian cancer have a total abdominal hysterectomy in addition to oophorectomy, and thus have reduced their endometrial cancer risk by as much as possible. Unaffected women should be educated on symptoms of endometrial cancer. Screening via endometrial biopsy every 1-2 years can be considered starting at age 30-35. A risk-reducing hysterectomy can be considered as well.

2.3.1.3 Other genes: Management and Screening

Other moderate-risk pathogenic variants as previously described warrant additional screening, although guidelines are generally not as well-established. Affected women with pathogenic variants that result in homologous recombination deficiency have an improved response to maintenance therapy with PARP inhibitors. Clinical trials or other targeted therapies may become available for other genes and should be investigated when genetic test results are available. For unaffected women, RRSO should be considered between age 40-45 for women with a pathogenic variant in BRIP1 and 45-50 for RAD51C/D, or earlier if family history of early-onset ovarian cancer is present. Further studies are needed to establish whether or not RRSO is warranted for other moderate-risk genes. Women with a pathogenic variant in PALB2 can consider RRSO after menopause if there is a family history of ovarian cancer. They should begin annual breast mammograms with MRI at age 30 and should discuss the option of a bilateral risk-reducing mastectomy. For other genes, breast cancer surveillance should be based on family history.
2.3.2 Factors that Influence Genetic Testing Decisions

There are several factors that may predict whether a person decides to undergo genetic testing, but there is inconsistency from study to study regarding which factors are significant.\textsuperscript{88} Some of these factors have changed as genetic testing methods and anti-discrimination laws have been updated. In older studies, predictors of \textit{BRCA1/2} genetic testing uptake included being Caucasian, older, and wealthier as well as having children, a family history of breast or ovarian cancer, and a higher level of knowledge about genetic testing.\textsuperscript{89-91} One study from 2011 noted that the ability to cure or reduce the risk of a disorder also appeared to be a strong predictor of genetic testing; 77\% of individuals were theoretically willing to undergo testing for a curable condition, while only 50\% of individuals were interested in testing for an incurable disorder.\textsuperscript{92} Studies within the past ten years have conflicting evidence as to whether or not age, race, and education level are predictors of genetic testing. However, relatively consistent predictors of testing uptake include having a personal or family history of cancer, having a higher income, being more knowledgeable about genetics, and perceiving more benefits to the test than risks.\textsuperscript{88,93-95} A recent study also found that having discussed genetic testing with a surgeon was a strong predictor of genetic testing in newly diagnosed breast cancer patients.\textsuperscript{96}

Several studies have attempted to elucidate specific reasons why unaffected individuals decline genetic testing. One study looking only at genetic testing for \textit{BRCA1} in unaffected individuals from HBOC families noted that individuals were more likely to request testing if they had a higher socioeconomic status and adequate health insurance. This study took place in 1996, prior to the passing of the Genetic Information Non-Discrimination Act (GINA), so many individuals cited a fear of discrimination from employers or health insurance as the reason they declined testing.\textsuperscript{91} Another study from 2003 with 13 women at risk of carrying a pathogenic variant
in *BRCA1/2* found that being satisfied with participating in a surveillance program and being emotionally unprepared to cope with testing results were also reasons that testing was declined. Additionally, women who had a higher reluctance toward undergoing prophylactic surgery were less likely to move forward with genetic testing compared to women who were more comfortable with surgical intervention.\(^97\) Newer studies highlight additional reasons why patients opt out of genetic testing, including distrust in genetic information, perception of more risks than benefits, and significant financial barriers.\(^98,99\) One retrospective chart review from 2018 found that 267 of 1082 individuals who met NCCN criteria for *BRCA1/2* analysis did not receive genetic testing. However, only 22% of these individuals were disinterested in testing. 40% were advised to gather additional information from relatives before testing, and 38% desired testing but were prohibited by the cost.\(^98\)

In all aspects of medical care, health literacy plays an important role; this is the ability of a person to get the health-related information they need, understand it, and utilize it appropriately to make medical decisions. All health care professionals have a vital role in understanding how health literacy influences the care that patients receive. When it comes to genetic testing, limited health literacy has been consistently shown to be associated with lower genetic health knowledge. One study examined how genetic health literacy plays a role in understanding genetic testing; the results indicated that patients’ understanding of the utility of genetic testing is positively correlated with genetic health literacy.\(^100\) Another study interestingly noted that patients with low health literacy had a greater perceived importance of genetic information but a lower perceived importance of family health history information, suggesting a disconnect in their understanding of genetics and genetic testing.\(^101\) It is therefore essential that health care providers be aware of their
patient’s level of health literacy in order to appropriately communicate information and create equitable health care for all patients.

2.3.3 Psychological Impact of Genetic Testing Results

Previous studies have generally shown favorable psychosocial outcomes after genetic testing, with a slight increase in anxiety immediately after results are disclosed, followed by a return to pre-testing anxiety levels.\textsuperscript{102,103} A comprehensive review by Hirschberg \textit{et al.} found that both men and women who receive genetic testing for hereditary cancer syndromes experience no clinically significant long-term distress. Any psychological distress that does arise during the testing process appears to decrease over the course of the first year after testing. Individuals tested positive for a pathogenic variant in a cancer predisposition gene who continue to have higher levels of distress after testing tend to have a higher level of baseline distress, a history of depression and/or psychotropic drug use, an elevated risk perception, complicated grief or unresolved loss, especially if they lost a relative due to hereditary cancer. Additionally, individuals with children were more likely to have long-term distress.\textsuperscript{104} Being aware of these risk factors can facilitate the counseling process and can better prepare patients for their testing results.

2.4 Alternative Service Delivery Models

The growing demand for genetics services, combined with the deficit in genetics health care professionals, has led to a call for alternative service delivery models.\textsuperscript{15,105} The most recent practice analysis conducted by the American Board of Genetic Counselors (ABGC) found that
only 77.3% of certified genetic counselors are currently working in clinic, and 40.6% are working in cancer. More studies are necessary to clarify how many cancer genetic counselors are needed to meet current demand. Geographical barriers are a major access concern; a study in California found that genetics providers are concentrated in major metropolitan areas, and on average patients travel approximately 80 miles to access these services. To increase patient access, many cancer genetics providers have incorporated alternative service delivery models into their practice such as telephone, telemedicine, pre-recorded videos, or group genetic counseling. These models mitigate issues with travel distance and are generally more convenient for both the counselor and the patients, but issues with billing or reimbursement, equipment setup, and internet access, as well as the inability to physically see the patient hinder some efforts.

Studies have shown that alternative care delivery models result in an increase in genetics knowledge and access to services, although the effects on risk perception and possible lingering misconceptions warrant further study. The Cancer Risk Education Intervention Tool (CREdIT) was designed to facilitate pre-counseling education for low literacy women who are at an increased risk for breast and ovarian cancer. Participants’ general genetics knowledge improved after viewing the CREdIT slides, but changes in risk perception varied between participants—sometimes inappropriately increasing or decreasing. Additionally, some misconceptions remained after viewing the content; one woman still thought that she inherited her cancer predisposition from her niece, for example.

2.4.1 Cancer Genetic Testing by Non-Genetic Health Professionals

With the increase in interest in genetics services, many non-genetic health professionals (NGHPs) are ordering genetic testing themselves instead of referring patients to a genetics
provider. A recent survey investigating the attitudes, knowledge, and skills of NGHPs with regards to hereditary cancer testing found that these providers generally have positive views about their communication skills and are confident in their ability to take a family history and order genetic testing. The majority of providers responded that they discussed the benefits and limitations of close cancer surveillance (95%) and prophylactic surgery (89%), discussing results with family members (91%), and confidentiality (89%). Importantly, however, only 71% of providers thought it was their responsibility to manage emotions; 68% felt confident in their ability to interpret variants, and 55% had received formal training on communicating hereditary information. Other studies have noted that an increase in knowledge corresponds to an increase in the confidence of the provider to deliver genetics services, and many providers asked for more formal training in genetics. A study by the American Society of Breast Surgeons found that the majority of breast surgeons provided genetic testing and counseling, created 3-generation pedigrees and provided pre- and post-test counseling. Still, 11.8% of respondents did not feel confident in their ability to provide counseling and desired more educational support in genetics. It thus appears that NGHPs are willing and eager to provide genetics services, but more educational opportunities to increase their knowledge of genetics may be necessary to improve confidence in all aspects of the genetic testing process.

2.4.1.1 Knowledge Gaps and Educational Needs

One method to mitigate this issue is to provide more formal education in topics related to hereditary cancer genetics and genetic testing so that providers can better understand this process and identify which patients require a referral. A plethora of studies have examined the current level of genetics knowledge in NGHPs and the effects of educational programs. Genomic literacy and confidence appear to differ greatly between practice groups, provider specialty, and years of
practice. For example, OB-GYNs and specialists (e.g. surgeons and oncologists) generally have greater genetics knowledge and confidence compared to primary care physicians. Common knowledge gaps include the inability to recognize paternal inheritance of HBOC, the need for comprehensive rearrangement testing in high-risk women, and appropriate testing options for a VUS result. One study surveyed providers in Florida who order BRCA1/2 testing and assessed the educational needs and preferences among this group; responses from mostly physicians and nurse practitioners indicated that in-person training was most strongly preferred. Minimal time off work and continuing education credits were also strong motivators to participate in formal education training. Other commonly reported preferred educational methods include multi-modal genetics courses that combine distance learning with interdisciplinary in-person training, seminars detailing clinical genetics services referral pathways, electronic referral guidelines, and example case scenarios. A study conducted by the City of Hope demonstrated that a targeted outreach program among community-based clinicians resulted in a 40% increase in cancer genetics knowledge on average. 77% of participants felt that they could use the course information and materials to better counsel and refer patients for hereditary cancer risk assessment. Various professional societies and academic institutions currently offer seminars, workshops, and web-based curriculum and resources to help educate providers.

2.4.1.2 Motivations for and barriers to cancer genetics referrals

Although education programs aim to increase genetic literacy and knowledge, they are also intended to increase appropriate referrals to hereditary cancer genetics services. A survey of NGHPs indicated that common reasons for referral include eligibility based on the patient’s personal or family history, the need for enhanced risk assessment, improved medical management, concern for family members, and patient request. Major barriers to referral included a lack of
knowledge about a cancer genetics program and limited awareness of improved insurance coverage and anti-discrimination legislation.\textsuperscript{125} Another study that surveyed gastroenterologists and colorectal surgeons found that 82.7\% had referred a patient to cancer genetic counseling. The majority of those surveyed were able to correctly identify which patients were at a “much higher” risk to develop colorectal cancer than the general population; fewer were able to identify those at “somewhat higher” risk than the general population. Risk categories were based off the American Gastroenterological Association’s criteria for high, moderate, and average risk to develop colorectal cancer. The survey identified barriers to referral including insurance coverage issues and discrimination fears, as well as a lack of clear guidelines from professional organizations. Similar to previously mentioned studies, physicians were more confident ordering testing than interpreting the results or providing emotional support.\textsuperscript{126}

\textbf{2.4.1.3 Adverse effects of incorrect testing and non-referrals}

Many studies illustrate the negative effects that occur when testing is ordered incorrectly, or when appropriate referrals are not made. A national case series examined patterns from 21 cases submitted to the National Society of Genetic Counselors Cancer Special Interest Group that exhibited adverse outcomes of testing by NGHPs. Major patterns noted were: the wrong genetic test was ordered (e.g. \textit{BRCA1/2} analysis instead of MMR genes or full-sequencing when a familial pathogenic variant was known), the results were mis-interpreted (e.g. a VUS was counseled as if pathogenic or incorrectly considering a patient to have a “true negative” result), and inadequate genetic counseling was given.\textsuperscript{127} Negative outcomes of errors in genetic testing and result interpretation include unnecessary prophylactic surgery, unnecessary testing, psychosocial distress, false reassurance, and increased cost to insurance companies and patients.\textsuperscript{127,128} Provider education could lead to improved availability of services and more appropriate care.
2.4.2 Oncologist-Led Genetic Testing for Epithelial Ovarian Cancer Patients

Data from the National Health Interview Survey (NHIS) and a recent study indicated that only a third of individuals with a history of ovarian cancer have undergone genetic testing, despite current NCCN guidelines that all these women should receive genetic testing.\(^7\),\(^{17,93}\) Only 15% of patients with ovarian cancer had even discussed genetic testing with a healthcare provider.\(^93\) An increase in public awareness, targeted at providers and women with ovarian cancer, may help to increase utilization of genetics services.\(^90,93\) Professional organizations like the American College of Obstetricians and Gynecologists (ACOG), American Society of Clinical Oncology (ASCO), and Society of Gynecologic Oncologists (SGO) have published recommendations and guidelines that support germline and somatic genetic testing for epithelial ovarian cancer patients at the time of diagnosis; they state informed consent is required, including a discussion of the benefits, limitations, and implications of genetic testing results.\(^5,6,129,130\) SGO specifically recommends that patients begin with genetic counseling to discuss testing options, but they also acknowledge that if these services are not available from a genetic counselor, it is appropriate to receive counseling from a provider trained in cancer genetic counseling.\(^6\)

A streamlined \textit{BRCA1/2} genetic testing pathway for ovarian cancer patients has been proposed and studied in recent years. Colombo \textit{et al.} published the results of a study entitled Evaluating a Streamlined Onco-genetic \textit{BRCA} Testing and Counseling Model Among Patients with Ovarian Cancer (ENGAGE). In this study, a clinical team of oncologists and oncology nurses received training on \textit{BRCA1/2} testing and genetic counseling techniques. Pre-test counseling was provided by this team, although patients could elect to have counseling from a genetic counselor or geneticist prior to testing if desired. Results were disclosed by the clinical team, and patients with a positive result were recommended to undergo further genetic counseling with a genetic
counselor or geneticist. The results of the study were favorable, with a median turnaround time of 4.1 weeks in the United States. Patient satisfaction was 99%, and 80% of oncologists agreed that the process worked well and was efficient. However, only 50% of genetic counselors and geneticists agreed that patients received accurate \textit{BRCA1/2} testing information in the pre-test counseling session, suggesting that the process still requires some fine-tuning.\textsuperscript{19}

Barriers to implementing this type of streamlined pathway have been reported, including a limited number of providers that were appropriately trained, time constraints, insufficient health insurance, fear of discrimination, and issues with family communication.\textsuperscript{124,131} Issues that arise once the pathway has been implemented include difficulty interpreting a VUS result and lack of patient awareness of result implications for their own treatment.\textsuperscript{19,132} Pre-test counseling requires improvements to better educate patients, and referrals to a genetics provider are likely warranted for not only positive results, but also VUS results and a negative result for a patient with a family history.\textsuperscript{18} Still, other studies have shown comparable results with quick turnaround time, high patient satisfaction, high rates of genetic counseling appointment attendance, and appropriate changes in treatment based on genetic testing results.\textsuperscript{18,131,133,134} Patients were satisfied with the timing of testing and did not appear to be overwhelmed with the additional testing at the time of diagnosis.\textsuperscript{132} Thus, oncologist-led genetic testing for patients with EOC appears to be a feasible service model.
3.0 Manuscript

3.1 Background

Every year, nearly 300,000 women receive a diagnosis of ovarian cancer across the world. As the 5th leading cause of cancer death in women, the diagnosis is often devastating. Genetic predispositions to ovarian cancer have been identified and are thought to contribute to 20-25% of cases. Hereditary Breast and Ovarian Cancer syndrome (HBOC), Lynch syndrome, and pathogenic variants in moderate-risk genes have all been demonstrated to increase a woman’s lifetime risk of developing ovarian cancer.

Due to the high frequency of germline pathogenic variants, current NCCN guidelines advise that all women diagnosed with epithelial ovarian cancer (EOC) be offered germline genetic testing. In spite of this, only about 35% of women with this diagnosis have undergone genetic testing. Knowledge of a germline pathogenic variant in an affected individual can inform prognosis, guide treatment, and clarify the need for management of other cancer risk. Studies show that women with BRCA1/2 pathogenic variants have an improved prognosis and a better response to platinum-based chemotherapy and PARP inhibitors. These women may also benefit from additional cancer screening due to an increased risk of breast cancer, depending on their clinical status. Similarly, women with germline pathogenic variants in a Lynch-syndrome related gene have a higher survival rate, and some treatment options such as Pembrolizumab are targeted for mismatch-repair deficient tumors. Surveillance with frequent colonoscopies and endometrial biopsies might also be indicated for colon and endometrial cancer risk management.
Tumor testing alone can provide information to determine indications for targeted treatment options, but it cannot differentiate between germline and somatic variants. Identification of a germline pathogenic variant provides information on additional cancer risks, and it helps facilitate testing for at-risk family members. As such, tumor testing alone is not sufficient for women diagnosed with ovarian cancer. Germline genetic testing, or tumor testing followed by germline testing if there are results suspicious for a germline pathogenic variant, is necessary to provide comprehensive risk assessments to the patient and their families.

While the demand for genetic services has increased, there is a deficit in genetics health professionals.\textsuperscript{15,105} This has resulted in a call for alternative genetics service delivery models. Many non-genetics health professionals (NGHPs) are increasingly ordering genetic testing themselves, especially within the realm of hereditary cancer testing. Studies of oncologist-led testing pathways have been favorable so far, with quick turnaround times and high levels of patient and provider satisfaction.\textsuperscript{19,133,134} However, knowledge gaps and barriers to cancer genetics referrals after testing is completed persist and must be addressed for this service model to provide appropriate care. A recent study found that most NGHPs’ self-reported proficiency discussing the benefits and limitations to close surveillance and risk-reducing surgery, importance of delineating information to family members, and confidentiality of results. However, only 68\% felt confident in interpreting all variants, and 55\% had never received formal training in communicating this information.\textsuperscript{112} This theme has been replicated in other studies, as the American Society of Breast Surgeons reported that most providers were ordering genetic testing and were counseling patients on results, but 11.8\% of respondents did not feel confident in their counseling abilities and desired more educational support in genetics.\textsuperscript{114} Targeted outreach programs have been successful in the past in improving genetics knowledge, appropriate referrals, and counseling abilities.\textsuperscript{123} Errors in genetics
service delivery can have severe negative outcomes including unnecessary prophylactic surgery, inappropriate testing, psychological distress, false reassurance, and increased cost to insurance companies and patients.127,128

The goal of this study was to evaluate the current genetic testing practices of oncology providers for patients with epithelial ovarian cancer. Data surrounding the genetic testing practices of gynecologic oncologists are limited, and these individuals are central healthcare providers for women diagnosed with ovarian cancer. This study will help to elucidate how and when gynecologic oncologists are ordering genetic testing and referring patients to genetics services, and it could help highlight any barriers that currently exist. It will also assess provider confidence with different aspects of cancer genetics services to identify areas that may require educational outreach. Addressing these issues will facilitate access to genetic testing services for all patients with ovarian cancer and their families.

3.2 Methods

3.2.1 Study Population

A survey was sent to full and associate members of the Society of Gynecologic Oncology, a professional society comprised of gynecologic oncologists, researchers, medical oncologists, physician assistants, nurse practitioners, registered nurses, fellows, residents, students, patients, and caregivers. As healthcare systems differ vastly by country, only members who practice in the United States were recruited. Participants were deemed eligible if they are currently in clinical practice or have been in clinical practice within the past year.
3.2.2 Survey Development

Prior to survey distribution, exempt IRB approval was obtained from the University of Pittsburgh (Appendix A). The survey was developed utilizing the Qualtrics Survey System, which was accessible through the University of Pittsburgh license. The survey collected information on participant demographics, genetic testing and referral practices, and confidence with different aspects of the genetic testing process (Appendix B). It was estimated to take approximately 15 minutes to complete. Demographic information recorded included current job title, primary area of medical practice, location (urban/suburban/rural), practice setting, years of experience, age, gender, and ethnicity. Genetic testing questions inquired about ordering frequency, timing, type of test (e.g., multi-gene panel vs. \textit{BRCA}1/2 analysis only), barriers to testing, and how often results guide treatment decisions. Certain survey questions permitted multiple responses to be selected. Questions about referrals to genetics services similarly asked about frequency, timing, and barriers.

Several potential barriers to ordering genetic testing were asked in the survey, including patient disinterest, insurance coverage or discrimination concerns, lack of effect on medical management, lack of necessity, and confusion. A list of barriers to making cancer genetics referrals was provided for the respondents to select from, including patient disinterest, services already provided, insurance coverage or discrimination concerns, burden of additional appointments, lack of services in region, and confusion or lack of knowledge about services. Finally, a group of questions asked about education or training in genetics and confidence in: taking a 3-generation family history, ordering genetic testing, interpreting all possible results, counseling a patient on the meaning of the result, and deciding which patients should be referred to cancer genetics services. Respondents completed the survey online. Due to time constraints, the survey was not piloted.
3.2.3 Recruitment and Survey Distribution

An e-mail was sent out to invite members of SGO to participate in the research survey on December 14th, 2020. The recruitment e-mail explained the goals of the study, pointed out potential risks and benefits to taking the survey, provided researcher contact information, and had a link to the survey (Appendix C). The survey was open for a total of six weeks and closed on January 22nd, 2021. Two reminder e-mails were sent out with two weeks and one week remaining. An informed consent paragraph appeared before any survey questions could be viewed and required the participant to acknowledge to continue (Appendix B). Survey completion was used as proof of informed consent. Respondents that completed less than half the questions were not included, as not all questions necessary for analysis were answered (i.e., missing information on frequency of referrals, provider confidence, and demographic information).

3.2.4 Data Analysis

Descriptive statistics were performed and bar graph data visualizations were created using Qualtrics for demographic information, testing and referral practices and barriers, and provider confidence. Cross tabulations were calculated between provider testing practice and both use of genetic testing results to guide treatment and frequency of referrals made. Statistical analysis was performed using STATA statistical software. Fisher’s exact test was utilized to determine if there was a significant association between provider testing practice and confidence levels with different aspects of cancer genetics services and referrals. P-values were considered significant if they were less than 0.05. All outputs from statistical analysis can be found in Appendix D.
3.3 Results

3.3.1 Demographic Information

The survey was sent to 1,444 members of SGO who are full or associate members currently residing in the United States. A total of 186 responses were received, therefore giving the study a 12.9% response rate. Twelve responses were excluded because less than half of the questions were answered. Four additional responses from individuals who are not currently in clinical practice were also excluded. Hence, 170 responses were included in the analysis.

Almost 82% of respondents were gynecologist oncologists (Table 3), and nurse practitioners comprised the next largest group (10.6%). Thirty-three percent of respondents had 10-19 years of experience in patient care. Approximately 25% of respondents had 0-9 years of experience, and another 25% had 20-29 years. 17% of respondents had 30 or more years of experience. The majority (95.3%) of respondents worked primarily in gynecology/obstetrics or gynecologic oncology. Most respondents were white (84.7%), non-Hispanic (94.1%), and between the ages 30-69 (97.1%), and slightly more respondents were female (63.5%) than male. Individuals were asked two questions about practice type and location. Note that total percentages can be higher than 100 due to the ability to select multiple responses. The majority of respondents (53.5%) worked at an academic center at least some of the time and 36.5% of respondents were hospital based. More individuals reported to be part of a group practice (20%) than a private practice (8.8%). Most respondents worked in an urban setting (66.5%) at least some of the time, followed by suburban (40.6%) and then rural (7.1%). Individuals that selected “Other” specified working at referral centers and in the military (Table 3). Patients with epithelial ovarian cancer made up less than half of all patients seen in clinical practice for 90% of providers.
<table>
<thead>
<tr>
<th>Current job title</th>
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<tr>
<td>Gynecologic oncologist</td>
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<td>Other</td>
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<td>10 - 19</td>
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<td>20 - 29</td>
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<td>30 or more</td>
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<tr>
<td>Hospital based</td>
<td>36.5% (62)</td>
</tr>
<tr>
<td>Academic center</td>
<td>53.5% (91)</td>
</tr>
</tbody>
</table>
3.3.2 Ordering, Treatment, and Referral Practices

The majority (85%) of respondents always ordered genetic testing for patients with EOC (Table 4). The frequency that providers used genetic testing results to guide treatment and referred patients to cancer genetics was examined separately based on how often they ordered genetic testing. Of providers who always ordered genetic testing, about 30% also always used genetic testing results to guide treatment. 60% of these providers always referred patients to cancer genetics services. Of providers who often or sometimes ordered genetic testing, 23.8% always used the results to guide treatment, and a third always referred patients to cancer genetics. Only four providers responded that they never ordered their own genetic testing; all of these individuals always referred patients to cancer genetics.

Table 4. Treatment and Referral Tendencies by Testing Practice

<table>
<thead>
<tr>
<th>Used results to guide treatment</th>
<th>I always ordered genetic testing 85.3% (145)</th>
<th>I often/sometimes ordered genetic testing 12.4% (21)</th>
<th>I never ordered genetic testing 2.3% (4)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Always</em></td>
<td>29.7% (43)</td>
<td>23.8% (5)</td>
<td>50.0% (2)</td>
</tr>
<tr>
<td><em>Often/Sometimes</em></td>
<td>69.7% (101)</td>
<td>76.2% (16)</td>
<td>50.0% (2)</td>
</tr>
<tr>
<td><em>Never/Unsure</em></td>
<td>0.6% (1)</td>
<td>0% (0)</td>
<td>0% (0)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Referred patients to cancer genetics</th>
<th>I always ordered genetic testing 85.3% (145)</th>
<th>I often/sometimes ordered genetic testing 12.4% (21)</th>
<th>I never ordered genetic testing 2.3% (4)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Always</em></td>
<td>60.7% (88)</td>
<td>33.3% (7)</td>
<td>100% (4)</td>
</tr>
<tr>
<td><em>Often/Sometimes</em></td>
<td>33.8% (49)</td>
<td>61.9% (13)</td>
<td>0% (0)</td>
</tr>
<tr>
<td><em>Never/Unsure</em></td>
<td>5.5% (8)</td>
<td>4.8% (1)</td>
<td>0% (0)</td>
</tr>
</tbody>
</table>
Regarding the timing of testing, 64.5% of respondents indicated that they ordered testing at diagnosis (Figure 1). 37.3% of providers ordered before starting chemotherapy, and 34.3% ordered after beginning chemotherapy. Other responses included when a patient relapsed, after completion of chemotherapy, or after surgery has been performed.

![Figure 1. Percentage of providers who ordered genetic testing at indicated times in patient treatment](image)

Respondents were also asked to indicate the types of testing that they had ordered for patients with EOC in the past year and were permitted to select multiple types of testing. Panel testing was performed by 95.8% of providers (Figure 2). The second most frequently selected type of testing (44.0%) was somatic testing with reported germline mutations, followed by somatic testing without reported germline mutations (30.1%). Only 13.9% of providers indicated that they performed BRCA1/2 analysis alone in the past year. Other responses specified that test selection is deferred to genetic counselors.
The majority of respondents (53.6%) felt that none of the potential barriers listed prevented or hindered them from ordering genetic testing. The most commonly cited barriers to testing were patient disinterest (30.4%) and insurance coverage concerns (13.7%). Other responses (9.5%) specified long wait times for genetic counseling appointments, the financial confusion that testing causes patients, and a lack of patient interest to share information with family. Three of the four providers who never ordered genetic testing clarified that genetic counselors are consulted to order testing based on the patient’s personal history and their family history. No providers felt that genetic testing was unnecessary or were confused by genetic testing.

Providers were asked to select the timing at which they referred patients with EOC to cancer genetics services in the past year. About half of providers referred patients before genetic testing for pre-test counseling or at diagnosis (Figure 3). Almost 25% of providers referred patients
after ordering testing if there was a positive result; 14.3% referred patients after ordering testing if the result was unclear. 3.1% of providers refer after testing if there is a negative result. Other responses included after somatic testing if a germline mutation is suspected, when a result is in a gene the provider is not familiar with, when family history warrants further evaluation, or upon patient request.

Figure 3. Percentage of providers who referred patients to cancer genetics services at indicated times in patient treatment

40% of providers did not feel that any of the listed barriers prevented or hindered them from referring a patient to cancer genetics services. A third of providers felt that patient disinterest was a barrier. 17.3% of providers felt that they provided the necessary services to the patient. Insurance coverage and burdening patients with additional appointments were less commonly
indicated as barriers. Lack of cancer genetics services in the area was selected by 5% of providers. No respondents cited confusion about cancer genetics services as a barrier to referrals.

3.3.3 Genetics Education

Providers were asked to indicate if they had attended educational courses, events, and programs. The most commonly reported form of genetics education (71.2%) was talks at conferences focused on genetics. Continuing education events, genetics courses in medical school, and webinars were the next most commonly selected responses. About 10% of respondents did not have experience with any of the genetics education forms listed. Other forms of genetics education specified included a Masters or PhD in Genetics, research, on-the-job training in genetics, years of experience, and the City of Hope course.

3.3.4 Provider Confidence

Five questions were posed to assess providers’ confidence level with various aspects of cancer genetics services: taking a detailed family history, ordering genetic testing, interpreting results, counseling a patient on the meaning of results, and making a referral to cancer genetics services. Provider confidence levels were broken down and analyzed by testing practices. The majority of those who always ordered genetic testing were very confident in each part of the testing process; however, only 43.4% of these providers were very confident in their ability to take a detailed family history, and 13.1% described themselves as not confident. Similarly, almost 50% of those who often or sometimes ordered testing were only somewhat confident with this process. The majority of these providers were somewhat or not confident with aspects of the testing process,
except when it comes to knowing which patients to refer. 86% of providers who sometimes or often order testing are very confident in their ability to identify patients that warrant a referral. Only four providers never ordered testing, and their confidence levels varied. Three of the four providers were not confident with their ability to order testing, and all four were very confident in their ability to appropriately make referrals.

Fisher’s exact test determined there was no statistically significant difference between providers with different testing practices in regard to their confidence taking a detailed family history and deciding which patients to refer to cancer genetics. However, there was a significant difference between providers in confidence ordering testing as well as interpreting and counseling a patient on the meaning of genetic test results. Providers who always ordered genetic testing tended to describe themselves as very confident, while many providers who often or sometimes ordered testing tended to be somewhat confident (Table 5).
Table 5. Provider Confidence with Aspects of Cancer Genetics Services by Testing Practices

<table>
<thead>
<tr>
<th>Confidence Level</th>
<th>I always ordered genetic testing</th>
<th>I often/sometimes ordered genetic testing</th>
<th>I never ordered genetic testing</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Taking a detailed, 3-generation family history</strong></td>
<td></td>
<td></td>
<td></td>
<td>0.968</td>
</tr>
<tr>
<td>Very confident</td>
<td>43.8% (63)</td>
<td>38.1% (8)</td>
<td>50.0% (2)</td>
<td></td>
</tr>
<tr>
<td>Somewhat confident</td>
<td>43.0% (62)</td>
<td>47.6% (10)</td>
<td>50.0% (2)</td>
<td></td>
</tr>
<tr>
<td>Not confident</td>
<td>13.2% (19)</td>
<td>14.3% (3)</td>
<td>0.0% (0)</td>
<td></td>
</tr>
<tr>
<td><strong>Ordering genetic testing</strong></td>
<td></td>
<td></td>
<td></td>
<td>0.008*</td>
</tr>
<tr>
<td>Very confident</td>
<td>61.0% (86)</td>
<td>45.0% (9)</td>
<td>25.0% (1)</td>
<td></td>
</tr>
<tr>
<td>Somewhat confident</td>
<td>30.5% (43)</td>
<td>45.0% (9)</td>
<td>0.0% (0)</td>
<td></td>
</tr>
<tr>
<td>Not confident</td>
<td>8.5% (12)</td>
<td>10.0% (2)</td>
<td>75.0% (3)</td>
<td></td>
</tr>
<tr>
<td><strong>Interpreting all possible results</strong></td>
<td></td>
<td></td>
<td></td>
<td>0.005*</td>
</tr>
<tr>
<td>Very confident</td>
<td>60.0% (87)</td>
<td>28.6% (6)</td>
<td>50.0% (2)</td>
<td></td>
</tr>
<tr>
<td>Somewhat confident</td>
<td>37.2% (54)</td>
<td>57.1% (12)</td>
<td>25.0% (1)</td>
<td></td>
</tr>
<tr>
<td>Not confident</td>
<td>2.8% (4)</td>
<td>14.3% (3)</td>
<td>25.0% (1)</td>
<td></td>
</tr>
<tr>
<td><strong>Counseling a patient on meaning</strong></td>
<td></td>
<td></td>
<td></td>
<td>0.002*</td>
</tr>
<tr>
<td>Very confident</td>
<td>62.8% (91)</td>
<td>19.0% (4)</td>
<td>50.0% (2)</td>
<td></td>
</tr>
<tr>
<td>Somewhat confident</td>
<td>35.2% (51)</td>
<td>76.2% (16)</td>
<td>50.0% (2)</td>
<td></td>
</tr>
<tr>
<td>Not confident</td>
<td>2.1% (3)</td>
<td>4.8% (1)</td>
<td>0.0% (0)</td>
<td></td>
</tr>
<tr>
<td><strong>Deciding which patients to refer to cancer genetics</strong></td>
<td></td>
<td></td>
<td></td>
<td>0.999</td>
</tr>
<tr>
<td>Very confident</td>
<td>84.8% (123)</td>
<td>85.7% (18)</td>
<td>100% (4)</td>
<td></td>
</tr>
<tr>
<td>Somewhat confident</td>
<td>15.2% (22)</td>
<td>14.3% (3)</td>
<td>0.0% (0)</td>
<td></td>
</tr>
<tr>
<td>Not confident</td>
<td>0.0% (0)</td>
<td>0.0% (0)</td>
<td>0.0% (0)</td>
<td></td>
</tr>
</tbody>
</table>
3.4 Discussion

3.4.1 Current Genetic Testing and Referral Practices by Oncology Providers

Current NCCN guidelines recommend that all patients with epithelial ovarian cancer (EOC) be offered germline genetic testing. The first goal of this study was to gain an understanding of how oncology providers are currently ordering and utilizing genetic testing results in their practice for patients with EOC. Results from this study are encouraging, demonstrating that the overwhelming majority (85%) of oncology providers always order genetic testing for patients with EOC. Providers most often order genetic testing at diagnosis and typically order multi-gene panel or somatic testing. Many providers did not feel that listed barriers to testing and referrals applied to them, although patient disinterest and insurance coverage issues were sometimes indicated.

Recent literature has found that only about 35% of patients with ovarian cancer have received genetic testing in spite of guidelines recommending testing for all of these patients. However, this study found that 85% of gynecologic oncologists always ordered testing for patients with EOC in the past year. With such a high rate of providers always ordering testing, one would expect a higher percentage of patients to have received testing. There are several possible explanations for this discrepancy. First, this study focused on provider testing practice within the past year across the country. The study that estimated 35% of ovarian cancer patients to have received genetic testing collected data for women in California and Georgia only from 2012-2019. Additionally, the recent emergence of companion testing may also be increasing the proportion of patients with ovarian cancer who have received testing; Myriad Genetics launched their version of companion testing in 2014, for example. Therefore, better awareness of guidelines and improved
options for testing may have increased testing rates in very recent years. Additionally, providers might be ordering genetic testing for their patients with EOC, but genetic testing may not always be completed. Disease-related burden or patient disinterest could be preventing patients from going through with the testing, despite the order being in place. About 30% of providers listed patient disinterest as a barrier to testing, so it is possible that this is contributing to the discrepancy in ordering versus testing rates.

The high prevalence of multi-gene testing found in this study is in concordance with other recent studies. The utility of multi-gene panel testing is well-established, as it can detect more pathogenic variants in clinically actionable genes than BRCA1/2 testing alone. Interestingly, a third of providers ordered somatic testing without reported germline mutations despite the high prevalence of germline pathogenic variants in patients with EOC. It is possible that patients were not interested in germline testing. Providers may perform follow-up germline testing if there is a somatic result that is suspicious for being germline or refer individuals with suspicious tumor test results to cancer genetics professionals. Additionally, current ASCO guidelines recommend that women with EOC with negative germline testing should have somatic testing. It is therefore also possible that providers ordered somatic testing as a follow-up test to negative germline testing.

Because this survey only asked providers to list all types of testing they have ordered within the last year, further research is needed to clarify the context in which somatic testing is being ordered.

In this study, testing at diagnosis was the most commonly reported timing of testing, in compliance with current guidelines. ASCO guidelines recommend germline genetic testing at the diagnosis of EOC in order to determine whether therapy with PARP inhibitors is indicated and to make decisions regarding neoadjuvant chemotherapy. Interestingly, the majority of providers—regardless of their testing practice—indicated that they often or sometimes use test results to guide
treatment. However, both positive and negative genetic test results are informative for treatment and prognosis. Negative genetic test results can predict response to therapies and rule out the use of certain tailored therapies.\textsuperscript{7,8,78} Perhaps providers do not consider negative genetic test results to be informative for treatment purposes as no tailored therapies are available to wild-type patients. Further research is needed to investigate the situations in which providers do not use genetic test results to guide treatment. Testing was also ordered before or after beginning chemotherapy by about a third of survey respondents. The survey did not probe further into why testing was ordered at the indicated times, but there are many possible explanations. The patient may not have been seen by the provider until this stage in their treatment, or there could have been delays in being able to proceed with testing. A frequently mentioned “other” response to testing barriers was long wait times for genetic counseling appointments. Only four respondents indicated that genetic counselors handle all genetic testing for patients EOC. This further supports the notion that an alternative service pathway for patients with EOC to undergo genetic testing is needed to improve access, as many other studies have found that there are not enough cancer genetics providers to meet patient needs.\textsuperscript{15,105,106}

The majority of providers that always ordered genetic testing for patients with EOC also always referred these patients to cancer genetics services, while those who ordered testing less often tended to refer patients less frequently. One possible interpretation of this trend is that providers who order their own testing have a greater knowledge of genetics and are more likely to recognize their limitations in providing all necessary services surrounding genetic testing, such as coordinating testing for family members and addressing patient emotions. This would be in line with other studies which have found that providers who have a greater familiarly and knowledge in genetics have higher rates of referrals.\textsuperscript{117,122,126} This study did not examine the reasons why these
providers were making referrals to genetics services. Apart from patient request or eligibility, commonly described motivations for referrals in current literature are to enhance risk assessment, improve medical management, and address concern for family members. Additionally, recent studies have found that many non-genetics healthcare professionals experience difficulty managing patient emotions and providing support. When asked about barriers to referrals, only 20% of providers surveyed responded that they provided all the needed services to the patients. Therefore, one explanation for the high rate of referrals among providers who order their own testing is that they recognize the benefits of genetic counseling and the additional services that can be provided by genetics professionals.

3.4.2 Types of Genetics Education

The most common forms of genetics education that respondents reported to have received were talks at conferences about genetics, Continuing Education events, and webinars. These platforms, while not formal training in genetics, have been frequently utilized by providers in the past and have been shown to increase genetics knowledge in previous studies. Some providers specified that multiple years of experience in their career supplied their genetics training. Providers with multiple years of experience should still be encouraged to improve their knowledge in genetics due to the rapidly evolving nature of the field. Educational sessions should continue to be offered in a variety of platforms to ensure accessibility.
3.4.3 Confidence Levels of Oncology Providers with Genetic Testing Process

Overall, providers were most confident in deciding which patients to refer to cancer genetics services and were least confident in taking a detailed, 3-generation family history. Confidence levels with these aspects of genetics did not significantly differ between providers with different testing practices. Previous studies have suggested that non-genetics healthcare providers are proficient at identifying high-risk patient that require referrals, but they tend to be less adept at identifying moderate or low risk patients.\textsuperscript{118,126} While high confidence levels for referring patients is reassuring, this study did not assess provider knowledge with or appropriateness of referrals being made. It is important that future studies ensure that appropriate referrals are being made for all patients. Confidence levels for obtaining a family history were slightly lower than expected; less than half of respondents felt very confident performing this task. One recent study found that 91\% of non-genetics healthcare providers felt confident taking a cancer genetics family history, and a survey of breast surgeons found that 90\% of respondents were willing and able to take 3-generation pedigrees for patients.\textsuperscript{112,114} Further research can determine what providers are struggling with when taking a family history.

Finally, provider confidence with ordering testing, interpreting all possible results, and counseling the patients on the meaning of the results significantly differed between providers with different testing practices. Providers who always ordered their own genetic tests tended to label themselves as “very confident” in these tasks, while providers who ordered their own tests only often or sometimes labeled themselves “somewhat confident.” Many studies have demonstrated that the more confident a provider is with aspects of genetics services, the more likely they are to deliver those services themselves.\textsuperscript{112-114,125,126} These results are therefore not surprising and are in agreement with current literature. Recent studies have also shown that many providers struggle
with appropriately interpreting a variant of uncertain significance, negative results in the context of a strong family history, and results in less well-studied genes.\textsuperscript{18,19,62,112,127} These issues may be playing a role in provider confidence, but ultimately future studies are required to tease out why providers who order their own genetic testing less frequently tend to be less confident with interpreting results and counseling patients.

### 3.4.4 Study Limitations and Future Directions

This study provided new information on oncology providers’ genetic testing practices for patients with EOC, but several limitations exist. The sample may not be truly representative of all oncology providers, as there was a low response rate. The results may also be subject to response bias, as providers who are more interested or more confident in cancer genetics may have been more inclined to respond to the survey. Finally, this survey was descriptive in nature, and so underlying causes were not able to be examined.

In spite of the limitations, the results of this study are encouraging and suggest that mainstreaming genetic testing for patients with EOC by oncology providers is an achievable service delivery model. Future studies should continue to delve into this pathway by examining providers’ genetics knowledge and the appropriateness of testing and referrals. More specific questions, perhaps with case scenarios or general knowledge questions, could help to elicit whether providers are accurately ordering, interpreting, and acting on genetic information. It would also be of interest to learn more about why the trends observed in this study exist. For example, with such a high percentage of providers always ordering testing, one would expect a higher percentage of patients to have received testing. Future studies can help to determine if testing rates have dramatically increased in recent years or if there is a significant absence of follow-through with
genetic test orders. Additionally, it would be of interest to determine the context that somatic testing is being ordered. Are providers ordering it alone, or prior to or after germline testing? Is a particular method more cost effective? Examining answers to questions such as these can help to improve this oncologist-led alternative service delivery pathway and increase access to genetics services.

3.5 Conclusion

As up to 25% of epithelial ovarian cancer diagnoses have an underlying genetic predisposition, accessible genetic testing is crucial to appropriately guide treatment for these patients. This study adds to the growing body of evidence that oncologist-led genetic testing for patients with EOC is a potentially feasible alternative service delivery model. The majority of oncology providers are ordering multi-gene testing for this patient population and most commonly are doing so at diagnosis, in concordance with current NCCN and ASCO guidelines. Additionally, most providers who order testing independently are still referring patients to cancer genetics services, which can help to address patient emotions and risk to family members in more detail when necessary. Genetics educational support has been consistently shown to be highly desired, and results of this study suggest that genetics talks at conferences and continuing education events are popular modalities. Overall, oncologist-led genetic testing for patients with EOC with appropriate referrals to genetics services appears to be an achievable way to increase access to genetics services, but further research is necessary to tease out differences in provider confidence and investigate reasons for trends observed in this study.
The implications of this study’s results extend into both genetic counseling and public health. Assessing and monitoring community needs is the first essential service of public health. It is well-established that patients with epithelial ovarian cancer require genetics services for comprehensive care. The Center for Disease Control considers both Hereditary Breast and Ovarian Cancer syndrome and Lynch syndrome to be Tier 1 genomic conditions; there is significant evidence that identifying individuals with these conditions can have a significant public health impact. The shortage of genetics healthcare professionals necessitates alternative service delivery models in order to increase access to genetics services. Some service models can still incorporate genetics professionals themselves, such as telephone, telemedicine, pre-recorded videos, or group genetic counseling. However, these models alone still cannot meet the current demand. If every patient diagnosed with EOC should be offered genetic testing, then genetic testing cannot be solely completed by genetics professionals.

This study sought to investigate a potential solution to the low percentage of genetic testing among patients with EOC, which reflects the second essential service of public health. The low rate of genetic testing is a hazard affecting these patients, as tailored treatment opportunities could be missed and at-risk family members may not be appropriately identified. The results of this study suggest that oncologist-led genetic testing for patients with EOC is a feasible alternative service delivery model, with 85% of gynecologic oncology providers always ordering testing for these patients. Supporting this service pathway has the potential to improve testing accessibility and increased testing rates, which in turn could lead to improved health outcomes for patients and their families.
Within public health, there are three core functions: assessment, policy development, and assurance. This study assessed how oncology providers are using genetic testing in their practice and how confident they feel about different aspects of cancer genetics. This data is critical information that can be used to identify aspects of this service model that are working well and aspects that require targeted outreach to improve the process. This survey demonstrated that providers are generally ordering multi-gene panel testing at diagnosis for these patients, which is in compliance with current guidelines. Additionally, providers were very confident in their ability to know which patients require a referral to cancer genetics services. Providers were generally less confident in taking a detailed family history, so educational outreaches can strive to highlight the importance of family history and the familial implications of genetic test results. Addressing the aspects of genetics services that providers struggle with helps to assure that oncologist-led genetic testing delivers appropriate care.

This study provides valuable information about the current genetic testing practice of gynecologic oncology providers. Future studies can continue to explore this practice by investigating provider knowledge and the appropriateness of referrals. Cancer genetic professionals have a key role to play in supporting this service pathway. They can help to educate and assist providers with questions that arise during the testing process. Furthermore, as referrals for positive or unclear results are a pivotal aspect of this service model, cancer genetics professionals are essential for comprehensive care of patients and their family members. The public health impact that this service model could provide is significant and warrants additional investigation.
Appendix A IRB Approval

EXEMPT DETERMINATION

<table>
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<th>August 14, 2020</th>
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<tbody>
<tr>
<td>IRB:</td>
<td>STUDY20070321</td>
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<tr>
<td>PI:</td>
<td>Megan Czekalski</td>
</tr>
<tr>
<td>Title:</td>
<td>Mainstreaming Genetic Testing for Epithelial Ovarian Cancer by Oncology Providers: A Survey of Current Practice</td>
</tr>
<tr>
<td>Funding:</td>
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The Institutional Review Board reviewed and determined the above referenced study meets the regulatory requirements for exempt research under 45 CFR 46.104.

**Determination Documentation**

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<tr>
<td>Exempt Category:</td>
<td>(2)(ii) Tests, surveys, interviews, or observation (low risk)</td>
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<tr>
<td>Approved Documents:</td>
<td>• Czekalski Survey, Category: Data Collection;</td>
</tr>
<tr>
<td></td>
<td>• Czekalski HRP-721, Category: IRB Protocol;</td>
</tr>
<tr>
<td></td>
<td>• Czekalski Informed Consent, Category: Consent Form;</td>
</tr>
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</table>

If you have any questions, please contact the University of Pittsburgh IRB Coordinator, Larry Ivanco.

Please take a moment to complete our Satisfaction Survey as we appreciate your feedback.
Appendix B Survey Content

This study is being conducted by Megan Czekalski, a genetic counseling student at the University of Pittsburgh, for the completion of her thesis project. The purpose of this research study is to investigate current practices by oncology providers regarding cancer genetic testing for patients with epithelial ovarian cancer. We are surveying all members of the Society of Gynecologic Oncologists with this brief questionnaire, which will take approximately 15 minutes to complete. The survey collects information on your background (e.g. specialty, years of experience, etc.) and how you incorporate genetic testing in your practice. There are no foreseeable risks associated with this project, nor are there any direct benefits to you. You will not receive any payment for participation. Your responses will not be identifiable in any way, and all responses are confidential. Your participation is voluntary, and you may discontinue answering the survey questions at any time. Should you have any questions, you can contact Megan Czekalski at mac547@pitt.edu.

Q1 Are you in clinical practice currently or have you been in clinical practice within the past year?

☐ Yes

☐ No

☐ Decline to answer

Q2 Over the past year, approximately what percentage of your total practice involved patients diagnosed with epithelial ovarian cancer?

☐ I did not see any patients diagnosed with epithelial ovarian cancer in the past year

☐ 1-25%

☐ 26-50%

☐ 51-75%

☐ 76-100%

☐ Unsure
Many of the following questions will ask about genetic testing in your practice. For the purposes of this survey, we are interested in germline (hereditary) genetic testing only. Unless otherwise specified, only include somatic (tumor) genetic testing if germline results are also reported.

Q3 How often have you ordered genetic testing for patients with epithelial ovarian cancer in the past year?

☐ Never
☐ Sometimes
☐ Often
☐ Always
☐ Unsure

Q4 When did you typically order genetic testing for patients with epithelial ovarian cancer in the past year? **Select all that apply.**

☐ At diagnosis
☐ Before beginning chemotherapy
☐ After beginning chemotherapy
☐ Other (please specify): ___________________________________________________
☐ Unsure
Q5 What kind of genetic testing did you typically order for patients with epithelial ovarian cancer in the past year? **Select all that apply.**

- [ ] BRCA1/2 testing
- [ ] Mismatch repair gene testing (e.g. MLH1, PMS2, MSH2, MSH6; typically associated with Lynch syndrome)
- [ ] Panel testing (e.g. multi-gene testing that examines BRCA1/2 and other genes at the same time)
- [ ] Somatic testing (with reported germline mutations)
- [ ] Somatic testing (without reported germline mutations)
- [ ] Other (please specify): ____________________________________________
- [ ] Unsure
Q6 What barriers hinder or prevent you from ordering genetic testing for patients with epithelial ovarian cancer? **Select all that apply.**

☐ I do not think genetic testing is necessary

☐ It will not change medical management for the patient

☐ I am concerned about insurance discrimination against patients

☐ I am concerned about insurance not covering genetic testing

☐ My patient was not interested

☐ I am confused by genetic testing

☐ Other (please specify): __________________________________________

☐ None of these apply

Q7 How often have you used genetic testing results to guide the treatment (e.g. chemotherapy) of patients with epithelial ovarian cancer in the past year?

☐ Never

☐ Sometimes

☐ Often

☐ Always

☐ Unsure
Q8 How often have you referred patients with epithelial ovarian cancer to cancer genetics services in the past year?

- Never
- Sometimes
- Often
- Always
- Unsure

Q9 When did you typically refer patients with epithelial ovarian cancer to cancer genetics services in the past year? Select all that apply.

- At diagnosis
- Before genetic testing for pre-test counseling
- After ordering genetic testing if there is a positive result
- After ordering genetic testing if there is an uncertain result
- After ordering genetic testing if there is a negative result
- Other (please specify): ________________________________________________
- Unsure
Q10 What barriers hinder or prevent you from referring patients with epithelial ovarian cancer to cancer genetics services? **Select all that apply.**

☐ I provide the needed services to patients

☐ I am concerned about insurance discrimination against patients

☐ I am concerned about insurance not covering genetic counseling or testing

☐ I do not want to burden my patients with additional appointments

☐ My patient was not interested

☐ There are no cancer genetics services in the area

☐ I do not know who to refer my patients to

☐ I am confused about these services

☐ Other (please specify): ______________________________________________________

☐ None of these apply
Q11 Have you completed any training or continuing education in genetics? **Select all that apply.**

- [ ] Completed course(s) in genetics in medical school
- [ ] Completed a residency or fellowship in genetics
- [ ] Watched webinar(s)
- [ ] Attended talk(s) at conference(s) about genetics
- [ ] Attended Continuing Education event(s)
- [ ] Taken formal course(s) in genetics after medical school
- [ ] Obtained certification(s) in genetics
- [ ] Other (please specify): ________________________________
- [ ] None of these apply

Q12 How confident are you in taking a detailed, 3-generation family history?

- [ ] Not confident
- [ ] Somewhat confident
- [ ] Very confident
- [ ] Decline to answer
Q13 How confident are you in ordering genetic testing?

- Not confident
- Somewhat confident
- Very confident
- Decline to answer

Q14 How confident are you in interpreting all possible genetic testing results (e.g. positive, negative, or a VUS)?

- Not confident
- Somewhat confident
- Very confident
- Decline to answer

Q15 How confident are you in counseling a patient on the meaning of their genetic testing results?

- Not confident
- Somewhat confident
- Very confident
- Decline to answer
Q16 How confident are you in deciding which patients should be referred to cancer genetics?

- Not confident
- Somewhat confident
- Very confident
- Decline to answer

Q17 What is your current job title?

- Gynecologic oncologist
- Medical oncologist
- Surgical oncologist
- Nurse practitioner
- Physician assistant
- Registered nurse
- Other (please specify): ________________________________
- Decline to answer
Q18 What is your primary area of medical practice?

- Gynecology/obstetrics
- Internal medicine
- Family practice
- Other (please specify): 
- Decline to answer

Q19 Which of the following best describes the location where you work? Select all that apply.

- Rural
- Suburban
- Urban
- Other (please specify): 
- Decline to answer
Q20 Which of the following best describes your practice setting? Select all that apply.

- [ ] Private practice
- [ ] Group practice
- [ ] Hospital based
- [ ] Academic center
- [ ] Other (please specify): ________________________________________________
- [ ] Decline to answer

Q21 Approximately how many years of experience do you have in patient care?

- [ ] 0-9
- [ ] 10-19
- [ ] 20-29
- [ ] 30 or more
- [ ] Decline to answer
Q22 Which category below includes your age?

- 29 or younger
- 30-49
- 50-69
- 70 years or older
- Decline to answer

Q23 What gender do you identify as?

- Female
- Male
- Nonbinary
- Other (please specify): __________________________________
- Decline to answer
Q24 What ethnicity do you identify as? Select all that may apply.

☐ American Indian or Alaskan Native

☐ Asian

☐ Black or African-American

☐ Native Hawaiian or other Pacific islander

☐ White

☐ Other (please specify): ________________________________________________

☐ Decline to answer

Q25 Are you of Hispanic, LatinX, or Spanish origin?

☐ Yes

☐ No

☐ Decline to answer
Appendix C Recruitment E-mail

Dear SGO member,

I’m reaching out to you today to invite you to participate in a research study. This study is being conducted by Megan Czekalski, a second-year genetic counseling student at the University of Pittsburgh.

She is conducting an online, anonymous survey as part of her master’s thesis project entitled “Mainstreaming Genetic Testing for Epithelial Ovarian Cancer by Oncology Providers: A Survey of Current Practice.” This research project seeks to investigate current practices by oncology providers regarding cancer genetic testing for patients with epithelial ovarian cancer. We are surveying all members of the Society of Gynecologic Oncologists with this brief questionnaire, which will take approximately 15 minutes to complete.

The survey collects information on your background (e.g., specialty, years of experience, etc.) and how you incorporate genetic testing in your practice. There are no foreseeable risks associated with this project, nor are there any direct benefits to you. You will not receive any payment for participation. Your responses will not be identifiable in any way, and all responses are confidential. Your participation is voluntary, and you may discontinue answering the survey questions at any time.

To participate in the survey, please use this link: https://pitt.co1.qualtrics.com/jfe/form/SV_8cEm74MY0W2mMJv

Disclaimer: This survey is not in connection to the Society of Gynecologic Oncology.

This project is being conducted in partial fulfillment of a Master of Science in Genetic Counseling at the University of Pittsburgh. This study has been approved by the University of Pittsburgh Institutional Review Board, STUDY20070321.

Thank you for your time and participation. Should you have any comments or questions, please feel free to contact Megan at mac547@pitt.edu. You may also reach out to other members of the thesis committee:

Rachelle C. Huziak, MS, CGC: huziakr@upmc.edu
Phuong Mai, MD, MS: maip@upmc.edu
Andrea L. Durst, MS, DrPH, CGC: adurst@pitt.edu
CONFIDENCE TAKING FAMILY HISTORY

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. tab fhx order_group, exact exp

Key

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Fisher’s exact = 0.968

CONFIDENCE ORDERING GENETIC TESTING

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CONFIDENCE INTERPRETING TESTING

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CONFIDENCE COUNSELING A PATIENT ON MEANING

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CONFIDENCE DECIDING WHICH PATIENTS TO REFER

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Bibliography

27. Kurman RJ, Carcangiu ML, Herrington CS, Carcangiu ML. WHO Classification of Tumours of Female Reproductive Organs. Lyon, FRANCE: International Agency for Research on Cancer (I A R C) (UN); 2014.


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