Uptake Rates of Risk-Reducing Surgeries for Women at Increased Risk of Hereditary Breast and Ovarian Cancer Applied to Cost-Effectiveness Analyses: A Scoping Systematic Review

Julia Simões Corrêa Galendi *, Sibylle Kautz-Freimuth, Stephanie Stock © and Dirk Müller *

Institute of Health Economics and Clinical Epidemiology, Faculty of Medicine and University Hospital of Cologne, University of Cologne, 50935 Cologne, Germany; sibylle.kautz-freimuth@uk-koeln.de (S.K.-F); stephanie.stock@uk-koeln.de (S.S.)

* Correspondence: julia.simoes-correa-galendi@uk-koeln.de (J.S.C.G.); dirk.mueller@uk-koeln.de (D.M.)

Simple Summary: For women who have tested positive for BRCA mutations, the decision to make use of preventive surgical options, such as risk-reducing mastectomy (RRM) or risk-reducing bilateral salpingo-oophorectomy (RRSO), depends on the women’s personal preferences and the cultural/social context. Among others, the cost-effectiveness of RRM and RRSO can be affected by the uptake rate of these preventive surgical options. Uptake rates of surgery should be given more attention in the conceptualization of health economic modeling studies for RRM and RRSO. Prospective multicenter studies are recommended to reflect regional and national variations in women’s preferences for preventive surgery.

Abstract: The cost-effectiveness of genetic screen-and-treat strategies for women at increased risk for breast and ovarian cancer often depends on the women’s willingness to make use of risk-reducing mastectomy (RRM) or salpingo-oophorectomy (RRSO). To explore the uptake rates of RRM and RRSO applied in health economic modeling studies and the impact of uptake rates on the incremental cost-effectiveness ratios (ICER), we conducted a scoping literature review. In addition, using our own model, we conducted a value of information (VOI) analysis. Among the 19 models included in the review, the uptake rates of RRM ranged from 6% to 47% (RRSO: 10% to 88%). Fifty-seven percent of the models applied retrospective data obtained from registries, hospital records, or questionnaires. According to the models’ deterministic sensitivity analyses, there is a clear trend that a lower uptake rate increased the ICER and vice versa. Our VOI analysis showed high decision uncertainty associated with the uptake rates. In the future, uptake rates should be given more attention in the conceptualization of health economic modeling studies. Prospective studies are recommended to reflect regional and national variations in women’s preferences for preventive surgery.

Keywords: cost-effectiveness; patient-centered care; economic modeling; genetic testing; breast cancer; risk-reducing surgery

1. Introduction

In recent decades, genetic testing and counseling have evolved to become an essential part of hereditary breast cancer (BC) and ovarian cancer (OC) prevention. Women who are carriers of germline BRCA1 and/or BRCA2 mutations can be offered risk management strategies that can significantly reduce the risk of BC/OC and cancer-related mortality. Risk-reducing mastectomy (RRM) has been shown to decrease the risk of BC and to provide an overall survival benefit for BRCA1 mutation carriers [1]. Risk-reducing salpingo-oophorectomy (RRSO) decreases the risk of OC and improves cancer-related and overall survival [2], while there are inconsistent results on the impact of RRSO on BC risk in BRCA
When opting for one or both risk-reducing surgeries, a woman has to weigh the benefit of reducing the cancer risk against potential negative consequences of these procedures, such as the loss of fertility, premature menopause, or psychological and physical suffering [5,6]. Hence, some women might opt either for delaying preventive surgery or for intensive surveillance instead.

The reimbursement of risk management strategies for BRCA mutation carriers depends on their clinical effectiveness for preventing cancer as well as on their economic consequences. To evaluate the lifelong health economic impact of preventive strategies, models are usually applied with several input parameters. These parameters include data on the course of the disease, such as the cancer incidence, the impact of preventive surgeries in reducing cancer risk, costs, and utilities (i.e., health state preference values).

A recent systematic review of health economic modeling studies concluded that targeted screening followed by risk-reducing strategies might be cost-effective. However, the input parameters applied to these models often differed, notably the uptake rates of surgeries (i.e., the women’s choice for RRM and RRSO) [7]. The uptake of surgery among BRCA mutation carriers varies substantially around the world. According to data from an international database from 10 countries, the rate of RRM was highest in the United States (50%) and lowest in Poland (4.5%). The uptake of RRSO was highest in France (83%) and lowest in China (37%) [8]. Thus, to some extent, the heterogeneity of uptake rates could be explained by cultural differences across countries [7]. However, the study designs used for measuring women’s uptake rates or the strength of recommendation for prophylactic surgeries in the clinical management of BRCA carriers might also explain the observed differences. In addition, there is some evidence that uptake rates of surgeries are sensitive parameters, resulting in potential uncertainty for the model outcomes [7].

To better illuminate how uptake rates are depicted in health economic modeling studies for preventing BRCA-induced cancer and to investigate the relevance of uptake rates for model results, we conducted a scoping review. The objectives of this review were (i) to systematically assess the sources of input data and assumptions for applying uptake rates of surgery within cost-effectiveness modeling studies and (ii) to assess the degree of uncertainty in the model outcomes that may result from different uptake rates in different settings. In addition, we conducted a value of information (VOI) analysis, based on one previously published model, to exemplify the decision uncertainty that results from uncertainty in the model outcomes [9].

2. Materials and Methods
2.1. Literature Review

The reporting of this scoping review is in accordance with the preferred reporting items for systematic reviews and the meta-analyses extension for scoping reviews (PRISMA-ScR) checklist [10]. The protocol for this scoping review was not pre-registered. A literature search was conducted in MEDLINE (via PUBMED) on 21 September 2022 and the Centre for Reviews and Dissemination (CRD) database to search for health economic modeling studies that addressed women who were offered RRM and/or RRSO after screening for germline BRCA mutations. In addition, we screened the studies included in the most recent systematic reviews published on the topic [7,11–14]. The search strategy is provided in the Supplementary Materials (File S1). Two reviewers screened the titles/abstracts of studies and selected potential studies for full text reading. The study selection and data extraction were carried out independently, and, in case of disagreement, consensus was achieved by discussion.

We included cost-effectiveness modeling studies that (i) targeted women at high clinical or familial risk for carrying BRCA mutations or known carriers of BRCA mutations and provided genetic testing for inheritable germline mutations including but not limited to BRCA mutations, (ii) evaluated risk management strategies based on RRM and/or RRSO, and (iii) presented the model outcomes as incremental cost-effectiveness ratios. Studies
were excluded if the reporting of the uptake rates was insufficient, or if the uptake rates were based on an assumption of perfect adherence. There was no language restriction.

From the selected models, we extracted the study characteristics (e.g., strategies used for comparison, model population) and the applied rates of the uptake of surgery (and respective age) in the case of a positive gene test result. In addition, we extracted results from deterministic sensitivity analyses to assess the impact of varying uptake rates of surgery on the incremental cost-effectiveness ratio (ICER). In order to assess if the uptake rates were appropriate for the models’ target population, the cited sources were retraced, from which we extracted data with regard to the study design, setting, number of participants, and time of follow-up.

2.2. Value of Information (VOI) Analysis

Based on a model developed and previously published by our institution [9], we conducted a VOI analysis to estimate whether the costs of additional evidence (e.g., conducting a new study) for reducing decision uncertainty associated with model outcomes are worthwhile. The model, on which the VOI analysis was based, assessed the cost-effectiveness of screen-and-treat strategies for German women at risk of hereditary BC and OC versus no testing. The model had a lifelong time horizon and included the health states ‘well’, ‘breast cancer without metastases’, ‘breast cancer with metastases’, ‘ovarian cancer’, ‘death’, and two post (non-metastatic) breast or ovarian cancer states. The perspective of the German statutory health insurance (SHI) was adopted, and input data were predominantly taken from German sources. While the input data concerning uptake rates are reported in Table 1 (i.e., Müller 2018), all input data are reproduced in the Supplementary Materials (Tables S1 and S2) [9].

The expected value of perfect information (EVPI) is computed as the difference in terms of the net monetary benefit (NMB) between the expected value of a decision made with perfect information and the expected value of the decision based on the current evidence [15]. While the EVPI shows the overall uncertainty, the expected value of partial perfect information (EVPPI) determines which parameters are highly related to decision uncertainty and the potential value of reducing that uncertainty by collecting more data on these specific parameters [16].

The 10,000 iterations generated in the probabilistic sensitivity analysis from our model were entered into the Sheffield Accelerated Value of Information (SAVI), which consists of a regression-based method for the EVPI and EVPPI calculations [17]. The value of eliminating parameter uncertainty associated with the uptake rates was quantified in comparison to three sets of other relevant model parameters—utilities (i.e., the quality-adjusted life year values), cancer incidence on BRCA mutation carriers, and risk reduction of preventive surgeries. These parameter sets were chosen due to their relevance in deterministic sensitivity analyses [9]. The NMB, which indicates the value of an intervention in monetary terms, was calculated for a hypothetical willingness to pay EUR 10,000.

3. Results

After the removal of duplicates, the search yielded 1197 references. After reading titles and abstracts, 31 studies were selected for full text reading. Among these, nineteen health economic modeling studies fulfilled the inclusion criteria [9,18–35]. Four studies were excluded due to insufficient reporting of the uptake rates of risk-reducing surgeries [36] because a perfect uptake of surgery was assumed [37,38] or because of an inappropriate presentation of the model result [39]. More information on the excluded studies is provided in the Supplementary Materials (Table S3). Two studies that were included had not yet been considered in any of the screened systematic reviews [22,31]. Figure 1 shows a flowchart of the study selection process.
The health economic models included covered health systems from different countries, including Norway [19], Australia [20,31,35], Brazil [32,33], the United Kingdom [18,21,25,34], the United States [22,23,28], Canada [24,26,27], Spain [30], and Germany [9]. Table 1 provides an overview of the included models and their uptake rates.

Table 1. Characteristics of modeling studies included.

| Author/Year          | Country                  | Strategies Being Compared | Model Population                                                                 | RRM Uptake Rate (Age, in Years) | RRSO Uptake Rate (Age, in Years) | Combined RRM and RRSO |
|----------------------|--------------------------|---------------------------|----------------------------------------------------------------------------------|---------------------------------|----------------------------------|-----------------------|
| Müller 2018 † [9]    | Germany                  | Testing (sequencing of BRCA1/2) vs. no testing | Women at risk for hereditary BC or OC due to family history, entering the model at age 35 | 0.06 (35)                      | 0.42 (35)                        | 0.45 (35)             |
| Simões Correa Galendi 2020 [33] | Brazil                  | Testing (sequencing of BRCA1/2) vs. no testing | First-degree relatives of index patients (BC or OC) with BRCA1/2 mutations, entering the model at age 30 | 0.10 (30–34)                   | 0.11 (35–39)                     | 0.27 (30–34)          |
| Petelin 2020 [31]    | Australia                | Risk management strategy (including risk-reducing surgeries) vs. population-based breast screening program | BRCA1/2 mutation carriers entering the model at age 20 | 0.31 (39)                      | 0.41 (45)                        |                      |
| Manchanda 2020 [29]  | United Kingdom/USA/Netherlands/China/Brazil/India | Testing (sequencing of BRCA1/2) all general population women ≥ 50 years vs. clinical criteria/FH-based testing | Women at risk for having mutations based on clinical and FH, entering the model at age 30 | 0.47                           | 0.55                           | Not considered       |
Table 1. Cont.

| Author/Year   | Country                  | Strategies Being Compared                                                                 | Model Population                                                                 | RRM Uptake Rate (Age, in Years) | RRSO Uptake Rate (Age, in Years) | Combined RRM and RRSO |
|---------------|--------------------------|-------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------|--------------------------------|----------------------------------|-----------------------|
| Hurry 2020    | Canada                   | Testing (sequencing of BRCA1/2) vs. no testing                                            | Index patients aged 50; first- and second-degree relatives (daughters entered the model at age 20; sisters at 50) | 0.21 (44)                     | 0.44 (54)                        | Not considered         |
| Guzauskas 2020| United States            | Population-based testing (sequencing of BRCA1/2) vs. testing based on FH or clinical risk| Women at risk for having mutations based on clinical and FH, entering the model at age 30 or 45 | 0.10 (30–34) | 0.11 (35–39) | 0.07 (40) Not considered |
| Sun 2019      | United States and United Kingdom | Testing (sequencing of BRCA1/2) for all women with BC vs. based on FH or clinical risk | Index patients (BC); first-degree relatives of index patients with BRCA1/2 mutations, entering the model at different ages | 0.47 (30)                     | 0.55 (30)                        | Not considered         |
| Moya-Alarcón 2019 | Spain                  | Testing (sequencing of BRCA1/2) vs. no testing                                           | Index patients at age 51 (OC); first- and second-degree relatives (daughters, nephews and nieces entered the model at age 23) | 0.25 (45–55) | 0.65 (45–55) | Not considered         |
| Kwon 2019     | Canada                   | Testing followed by RRSO (Sequencing of BRCA1/2) vs. no testing vs. RRSO for all (without testing) | First-degree relatives of index patients (OC), entering the model at age 40 | Not considered | 0.54 (40–50) | 0.33 (40–50) |
| Kemp 2019     | United Kingdom           | Testing (sequencing of BRCA1/2) vs. no testing                                           | Index patients aged 50 years (BC); first- and second-degree relatives (daughters entered the model at age 20; sisters at 50) | BRCA1 0.34 (40) | BRCA1 0.88 (40) | BRCA2 0.25 (40) BRCA2 0.87 (40) Not considered |
| Asphaug 2019  | Norway                   | Full sequencing of BRCA1/2 vs. seven-gene panel vs. 14-gene panel                        | Index patients aged 55 years (BC); first-degree relatives (daughters entered the model at age 25 and sisters at 50) | 0.12 (25–34) | 0.11 (35–60) | 0.10 (25–34) 0.28 (35–39) 0.35 (40–60) Not considered |
| Tuffaha 2018  | Australia                | Testing (sequencing of BRCA1/2) vs. no testing                                           | Index patients at age 40 (BC) with 10% probability for BRCA1/2 mutations; first- and second-degree relatives (children entered the model at age 10, siblings at age 40) | 0.3 (40)                     | 0.54 (40)                        | 0.16 (40)             |
| Ramos 2018    | Brazil                   | Testing (sequencing of BRCA1/2) vs. no testing                                           | First-degree female relatives of index patients (OC) with BRCA1/2 mutations, entering the model at age 30 | 0.18 (30)                     | 0.57 (30)                        | Not considered         |
Table 1. Cont.

| Author/Year | Country   | Strategies Being Compared | Model Population                                                                 | RRM Uptake Rate (Age, in Years) | RRSO Uptake Rate (Age, in Years) | Combined RRM and RRSO |
|-------------|-----------|---------------------------|----------------------------------------------------------------------------------|---------------------------------|----------------------------------|-----------------------|
| Li 2017 [28]| United States | Full sequencing of BRCA1/2 vs. five-gene panel | Women at risk for hereditary BC or OC due to family history or other hereditary syndromes, entering the model at age 40 or 50 | 0.42 (50) | Not considered | Not considered |
| Eccleston 2017 [21] | United Kingdom | Testing (sequencing of BRCA1/2) vs. no testing | Index patients age 50 years (OC) First- and second-degree relatives (daughters entered the model at age 20, sisters at 50) | BRCA1 0.34 (40) | BRCA2 0.25 (40) | BRCA1 0.88 (40) |
| NICE 2013 [18] | United Kingdom | Testing (sequencing of BRCA1/2) vs. no testing | First-degree female relatives of index patients (BC or OC) with BRCA1/2 mutations, entering the model at different ages 20–70 | 0.42 (30) | 0.54 (35) | 0.15 |
| Kwon 2010 [26] | Canada | Testing (different criteria for sequencing of BRCA1/2) vs. no testing | Subgroups of women with BC before age 40 or 50, regardless of ethnicity of family history | 0.20 (50–55) | § | 0.55 (50–55) | Not considered |
| Holland 2009 [23] | United States | Testing (sequencing of BRCA1/2) vs. no testing | Women with 10% pre-test probability of having a mutation, † who entered the model at age 35 | 0.15 (35) | 0.25 (35) | Not considered |
| Breheny 2005 [20] | Australia | Testing (sequencing of BRCA1/2) vs. no testing | First-degree relatives of individuals with BRCA1/2 mutations, entering the model at age 25 | 0.30 (38) | - | Not considered |

Abbreviations: BC: breast cancer, OC: ovarian cancer, RRM: mastectomy, RRSO: salpingo-oophorectomy, FH: family history. † Model used for value of information analysis; ‡ implies some familial history, but not necessarily a known mutation in the family; § in this population, RRM referred to contralateral mastectomy, assuming unilateral mastectomy as first-line BC treatment; ¥ individual simulation with clinical trial data.

3.1. Strategies Being Compared

A screen-and-treat intervention comprising BRCA genetic testing (i.e., full sequencing of BRCA genes) followed by RRM and/or RRSO was compared with a no prevention strategy by 10 studies [9,20,21,23–25,30,32,33,35]. A reference model developed by NICE compared testing vs. no testing; in this model, a proportion of women received risk-reducing surgery independent of the provision or outcome of testing [18]. In addition, in seven models, risk-reducing surgery was offered to both intervention and controls with differences between the compared strategies: Two models compared two testing strategies, namely full sequencing of BRCA genes versus a 7- or 14-gene panel [19,28], while five studies compared testing women based on familial/clinical risk versus different populational criteria [22,26,27,29,34]. While most studies provided immediate surgery for women who had tested positive, five studies modeled a woman’s option to delay surgery [19,22,26,30,33].

3.2. Study Population

In eight models, the model population was composed of index patients (i.e., the first person in the family diagnosed with a BRCA mutation after a diagnosis of either BC or OC), followed by cascade testing of first- and second-degree healthy relatives [18,19,21,24,25,30,34,35]. Nine
models addressed healthy women at increased risk for BRCA mutations due to familial risk [18,19,21,24,25,30,34,35]. Whereas some studies limited the population to first- or second-degree relatives of women affected by cancer with BRCA mutations [20,27,32,33], others defined the population by an established familial risk (with or without a known mutation in the family) [9,22,23,28,29]. Kwon et al. included only index patients diagnosed with BC at different ages [26].

Women entered the models at different ages, varying from 10 years [35] for siblings and children to 55 years [19] for healthy women and from 40 [26] to 55 years [19] for index patients. In most models, risk-reducing surgery was offered immediately after entering the model, while in five studies, the possibility of delaying surgery was accounted for [19,22,26,30,33].

3.3. Uptake Rates Applied to the Models

The uptake rate of RRM applied to the included health economic models ranged from 6% [9] to 47% [34], while those of RRSO varied between 10% [19] and 88% [21]. Figure 2A,B illustrates the variability in the uptake rates of RRM and RRSO for two different age groups, whereas the actual rates are described in Table 1.

Figure 2. Variability of the uptake rates applied to the models in two age groups. (A) Uptake rates of risk-reducing mastectomy; (B) uptake rates of risk-reducing salpingo-oophorectomy.
Whereas in some studies, the uptake rates were obtained from local centers or smaller departments [9,18,21,25,28,30,34], in others, the information was obtained from multicentric studies or registries reflecting larger regions of a country [19,24,27,32,33,35]. The rates were obtained from a single study [19,23,25,28,30,32,33,35] or from multiple studies [18,24,26,29,34]. Uptake rates without providing a reference or based on unpublished data were found in three studies [20,21]. Most studies considered country-specific evidence, with the exception of four [25,29,32,33]. Considerations with regard to the appropriateness of the selected uptake rates for the models’ target population were missing in all studies.

3.4. Sources of Uptake Rates

The sources of uptake rates were published between 2000 [40] and 2014 [41–43]. Most studies were based on retrospective data obtained from registries, hospital records, or questionnaires (n = 10). In seven models, the rates were based on prospective studies with women followed from 1 to 11 years [44] or based on a systematic review [45]. In these studies, women were recruited from the United States [42,44,46–50], the United Kingdom [42,46,50–53], Australia [54], Spain [41], Canada [43,45,48,49,55,56], and the Netherlands [40]. Two studies included women from several countries (i.e., Austria, Canada, France, Israel, Italy, Norway, Poland, and the United States) [48,49]. Table 2 details the methodological characteristics of the sources of data regarding uptake rates (as cited in each model study included).

Table 2. Sources of uptake rates cited by the included health economic models.

| Author/Year       | Source of Uptake Rate (Year) | Study Design                                    | Country                        | Number of Participants | Follow-Up |
|-------------------|------------------------------|-------------------------------------------------|--------------------------------|------------------------|-----------|
| Müller 2018 † [9] | Unpublished                  | Cross-sectional (single-center, hypothetical responses of women in a counseling situation) | Germany                        | 136 women at different ages following individual genetic counseling | -         |
| Simões Correa Galendi 2020 [33] | Chai (2014) [42] | Prospective, multi-center (post-testing counseling) | United States, United Kingdom | 1499 healthy women with inherited BRCA1/2 mutations | At least 0.5 years |
| Petelin 2020 [31] | Petelin (2019) [57] | Prospective and retrospective collected clinical data from a single specialized cancer center | Australia                      | 983 women with BRCA1/2 mutations (302 had BC at diagnosis) | 6.5 years |
| Manchanda 2020 [29] | Evans (2009) [58] | Matched controls (regional cancer registries) | United Kingdom                  | 221 healthy women with known BRCA1/2 mutations | 7 years |
| Hurry 2020 [24] | RRM: Metcalfe (2007) [56] Retrospective databases of mutation carriers Hospital discharges (RRSO) | Canada                          | RRM: 342 women with BRCA mutations, healthy and previous BC RRSO: 2119 who underwent hysterectomy (with or without BSO) or sterilization RRM: 4 years RRSO: | |
| Guzauskas 2020 [22] | Chai (2014) [42] | Prospective, multi-center (post-testing counseling) | United States, United Kingdom | 1499 healthy women with inherited BRCA1/2 mutations | At least 6 months |
| Sun 2019 [34] | RRM: Evans (2009) [58] RRSO: Manchanda (2012) [52] | Matched controls (regional cancer registries) Prospective observational cohort | United Kingdom | RRM: 105 women with BRCA mutations (healthy and BC) RRSO: 1133 women at high risk, less than 50% had BRCA4 mutations | 7 years |
| Moya-Alarcón 2019 [30] | Esteban (2015) [41] | Retrospective (hospital data) | Spain                           | 969 women from 682 families | 6 years |
### Table 2. Cont.

| Author/Year | Source of Uptake Rate (Year) | Study Design | Country | Number of Participants | Follow-Up |
|-------------|-------------------------------|--------------|---------|------------------------|-----------|
| Kwon 2019 [27] | Retrospective (multicenter study, questionnaire after receiving genetic test) | United States | RRSO: 703 women, healthy and with previous BC with BRCA mutations | 3.9 years |
| Kemp 2019 [25] | Retrospective (unpublished single hospital data) | United Kingdom | 858 women with BRCA mutations (unclear if previous cancer diagnosis) | - |
| Asphaug 2019 [19] | Retrospective (multi-center, questionnaire after receiving genetic test) | Austria, Canada, France, Israel, Italy, Norway, Poland, United States | RRM: 1290 RRSO: 177 women, healthy and with previous BC with BRCA mutations | 3.9 years |
| Tuffaha 2018 [35] | Prospective (multicenter, interviewer-administered questionnaire, surgery confirmed from pathology and medical records) | Australia | 325 healthy women with inherited BRCA mutations | 3 years |
| Ramos 2018 [32] | Retrospective (multicenter, questionnaire after receiving genetic test) | Various, Canada | RRM: 766/RRSO: 1383 women, healthy and with previous BC, with BRCA mutations | 3.9 years |
| Li 2017 [28] | Retrospective (registry data) | United States | 136 women with inherited BRCA mutations without previous cancer diagnosis | 1–11 years |
| Eccleston 2017 [21] | Retrospective (unpublished single hospital data) | United Kingdom | 858 women with BRCA mutations (unclear if previous cancer diagnosis) | - |
| NICE 2013 [18] | Matched controls (regional cancer registries) Retrospective (regional cancer registries) Retrospective (medical records) | United Kingdom | RRM: 105 RRSO: 314 RRM/RRSO: 554 All women with BRCA mutations, healthy or with BC | 7 years 5 years 6 years |
| Kwon 2010 [26] | Retrospective (medical records) Prospective (questionnaire, medical records) Prospective (single-center, hospital data) Retrospective (multicenter, questionnaire after receiving genetic test) Prospective (multicenter, questionnaires) | United States, the Netherlands | Metcalfe (2004): 390 women with early-stage BC, who are known carriers or are likely to carry BRCA1/2 mutations and were treated with unilateral mastectomy Other studies: healthy women with BRCA mutations and diagnosis of BC | 9 years |
| Holland 2009 [23] | Meta-analysis (five studies for uptake of BC, six studies for uptake of OC) | Various | 354 healthy, pre-symptomatic women who knew their mutation status and who had no prior history of BC or OC | - |
| Breheny 2005 [20] | Provided abbreviation not identifiable | - | - | - |

Abbreviations: BC: breast cancer, OC: ovarian cancer, RRM: risk-reducing mastectomy, RRSO: risk-reducing salpingo-oophorectomy. † Model used for value of information analysis.
The lowest uptake rate of RRM among healthy *BRCA* mutation carriers was reported by Metcalfe et al. in a retrospective cohort of 177 Norwegian women [48]. In that study, only 5% opted for RRM during the study follow-up [48]. In contrast, 51% of the 257 women retrospectively followed by Meijers-Heijboer et al. opted for RRM as the preferred strategy [40].

With regard to RRSO, the highest uptake was reported by Chai et al. (i.e., 86% of *BRCA1* and 70% of *BRCA2* mutation carriers under 50 years) [42]. In this study, all women were unaffected by cancer. In contrast, the lowest uptake of RRSO (26%) was reported by a retrospective single-center study that addressed women with previous BC to prevent a recurrent or contralateral BC [51].

### 3.5. Impact of Varying Uptake Rates in Sensitivity Analyses

Most studies provided information about the impact of varying the uptake of surgery in a deterministic sensitivity analysis, except for three [25,27,32]. In all but one of these studies [21], higher uptake rates improved the incremental cost-effectiveness ratio (ICER). Table 3 summarizes the results of the deterministic sensitivity analysis reported by the models.

| Author/Year | Strategies Being Compared | ICER | Deterministic Sensitivity Analysis (Impact on the ICER by Varying the Uptake Rates) |
|-------------|---------------------------|------|-------------------------------------------------------------------|
| Müller 2018 *†* [9] | Testing vs. no testing | EUR 17,027/QALY | 5% lower uptake of RRSO and RRSO combined with RRM increased ICER by 70%. |
| Simões Correa Galendi 2020 [33] | Testing vs. no testing | BRL 24,264/QALY (USD 11,726/QALY) | At a 75% reduced uptake of RRSO, the ICER increased by 25% and 15% for *BRCA1* and *BRCA2* mutation carriers, respectively. At a 75% reduced uptake of RRM, the ICER decreased by 1% and 17% for *BRCA1* and *BRCA2* mutation carriers, respectively. |
| Petelin 2020 [31] | Risk management strategy vs. population-based breast screening program | AUD 32,359/QALY (*BRCA1*) AUD 48,263/QALY (*BRCA2*) | Half the uptake rate for RRM or RRSO increased the ICER by about 5%. |
| Manchanda 2020 [29] | Populational testing vs. clinical criteria/FH-based testing | UK: USD 21,191/QALY USA: USD 16,552/QALY NL: USD 25,215/QALY China: USD 23,485/QALY Brazil: USD 20,995/QALY India: USD 32,217/QALY | Considering an uptake rate of RRSO or RRM of 50% lower (or 50% higher) increased (or reduced) the ICER by 10%. |
| Hurry 2020 [24] | Testing vs. no testing | CAD 14,294/QALY (USD 10,555/QALY) | 50% increase in RRS uptake rates (RRSO 0.66 and RRM 0.32), and mean age of RRSO 50 years reduced the ICER 85%. |
| Guzauskas 2020 [22] | Population-based testing vs. testing based on FH or clinical risk | USD 87,700/QALY | Considering an uptake rate of RRSO or RRM of 50% lower (or 50% higher) increased (or reduced) the ICER by 10%. |
| Sun 2019 [34] | Testing for all women with BC vs. based on FH or clinical risk | UK: GBP 10,464/QALY USA: USD 65,661/QALY | 10% higher uptake of RRSO reduced the ICER by 10%, and 10% lower uptake increase the ICER by 10% (for the UK payer perspective). 10% higher uptake of RRSO increased the ICER by 5%, 10% lower uptake decreased the ICER by 40% (for the US payer perspective). |
| Moya-Alarcón 2019 [30] | Testing vs. no testing | EUR 31,621/QALY | Considering an uptake rate of RRSO or RRM 25% lower (or 25% higher) increased (or reduced) the ICER by 5%. |
| Kwon 2019 [27] | Testing followed by RRSO vs. no testing | USD 7888 per QALY | Not reported |
| Kemp 2019 [25] | Testing vs. no testing | USD 1330/QALY | Not reported |
| Asphaug 2019 [19] | Full sequencing of *BRCA1/2* vs. seven-gene panel vs. 14-gene panel | USD 53,310/QALY | Considered negligible by the author. |
### Table 3. Cont.

| Author/Year       | Strategies Being Compared | ICER             | Deterministic Sensitivity Analysis (Impact on the ICER by Varying the Uptake Rates) |
|-------------------|----------------------------|------------------|-----------------------------------------------------------------------------------|
| Tuffaha 2018 [35] | Testing vs. no testing    | AUD 18,900       | Significant reducing the uptake rates by 10%, the ICER increased 40–50%.          |
| Ramos 2018 [32]   | Testing vs. no testing    | BRL 908/case of cancer avoided | Not reported                                                                       |
| Li 2017 [26]      | Full sequencing of BRCA1/2 vs. five-gene panel | USD 69,920/QALY  | Considering an uptake rate of RRM 50% lower (or 50% higher) increased the ICER by 50% (or reduced the ICER by 40%). |
| Eccleston 2017 [21] | Testing vs. no testing | GBP 4339/QALY   | Considering an uptake rate of RRSO 75% lower increased the ICER by 40%.           |
| NICE 2013 [18]    | Testing vs. no testing    | GBP 18,114/QALY § | The ICER increased about 30% when applying a realistic scenario (40% choose no procedure) over an ideal scenario (100% uptake). |
| Kwon 2010 [26]    | Testing vs. no testing    | USD 9084/QALY    | The ICER decreased as the rate of RRM increased and dominated above an 80% RRM rate. Higher rates (until 60%) of RRSO also decreased the ICER, and higher than 60%, the incremental benefits decreased faster than the incremental costs, increasing the ICER. |
| Holland 2009 [23] | Testing vs. no testing    | USD 9000/QALY    | Varying the uptake rate of RRM from 0% to 50%, the latter reduced the ICER by 10%. |
| Breheny 2005 [20] | Testing (sequencing of BRCA1/2) vs. no testing | USD 477/cancer-free year gained (BRCA1) USD 2150/cancer-free year gained (BRCA2) | Varying the uptake rate of RRM from 0% to 50%, the latter reduced the ICER by 10%. |

Abbreviations: ICER: incremental cost-effectiveness ratio, RRM: risk-reducing mastectomy, RRSO: risk-reducing salpingo-oophorectomy. † Model used for value of information analysis. § women aged 40–49 at 10% pre-test probability.

In seven studies, the impact of varying uptake rates on the cost-effectiveness was remarkable [9,21,23,24,26,33,35]. For instance, varying uptake rates changed the cost-effectiveness ratio from 20% to 40% [21] to more than 70% [9]. A common aspect of these studies is that the strategies being compared comprise a screen-and-treat intervention versus a no-testing strategy (i.e., no surgery in the comparator arm). In contrast, in six of the modeling studies, the authors considered the impact of varying the uptake rates to be slight (≤10%) or negligible [18–20,22,29,34]. In all of these studies, a risk-reducing surgery was offered to both the intervention and the comparator arm [18,19,22,29,34].

### 3.6. VOI Analysis

The overall EVPI per person is estimated at EUR 1680, which is the value of acquiring perfect information (i.e., eliminating all uncertainty) about all parameters applied to the model (detailed in Tables S1 and S2). The EVPPI per person for the predefined parameter sets is shown in Figure 3. The EVPPI value indicates to what extent more information on these sets of parameters would reduce the decision uncertainty (i.e., the chance that the decision-maker incorrectly opts for the strategy with lower payoffs, which, in our model, was the no-testing strategy). The maximum return in terms of the net monetary benefit from removing uncertainty around the uptake rates was EUR 239 (standard error (SE): EUR 24), corresponding to 14% of the total EVPI. The second set of parameters with the highest EVPPI was cancer incidence in the BRCA mutation carriers (EUR 207, SE: 25), followed by the risk reduction of preventive surgeries (EUR 188, SE: 25), and the lowest were the utilities (EUR 154, SE: 27).
4. Discussion

According to this comparison, the uptake rates of risk-reducing surgeries applied in cost-effectiveness models are sensitive parameters. In the models’ deterministic sensitivity analyses, there was a clear trend that a lower uptake rate increased the ICER and vice versa. Considering the vast potential of both RRM and RRSO for reducing the risk of cancer and cancer-related mortality, this is a little surprising. However, in one analysis, the authors reported a slightly higher cost-effectiveness ratio compared to the base case when the uptake of RRM was increased in a sensitivity analysis. The authors explained this counterintuitive effect with high costs for preventive treatment, which were not offset by survival gains (because of the high survival rates in women who do not undergo RRM) [21].

While deterministic analyses demonstrate the model’s sensitivity to a single input parameter, the VOI analysis evaluates the uncertainty of multiple parameters simultaneously. By sampling each parameter several times from a given range at each iteration, a more reliable estimation of the uncertainty can be provided, especially in models with parameters that correlate to each other [15]. The VOI analyses indicate the potential NMB forgone by making the decision between two treatment alternatives with current (i.e., uncertain) parameters, in comparison to making the decision with perfect information. As a decision rule for VOI analyses, the cost of future studies to gather more information about uncertain model parameters should not exceed the NMB elicited in the VOI analysis [15].

The high EVPPI of uptake rates indicates that gathering more information about the uptake rates would have a slightly higher impact on reducing decision uncertainty than additional information about other parameter sets (i.e., cancer incidence on BRCA mutation carriers, risk reduction of preventive surgeries, or utilities). Although the VOI calculation reflects the uncertainty in the German model, this finding is likely to be replicated in similar models. The VOI analysis can be easily replicated using regression-based methods based on the iterations generated in the probabilistic sensitivity analysis [17].

The uptake rates applied to modeling studies varied substantially. To reflect the attitudes and preferences of the different target populations, different sources of input data have been chosen for models. This variability might be explained by several factors, such as (i) cultural differences, (ii) individual-related factors, (iii) age-dependent factors, and (iv) an improved acceptance of preventive surgeries over time.

(i) Cultural differences (e.g., perception of health and disease, femininity, autonomy) and the risk of financial and social discrimination might influence the preference for genetic testing and risk-reducing surgery [40]. In a previous systematic review, it was suggested that cultural differences between countries could explain the variability in uptake rates to
a large degree [7]. However, according to our results, most models used country-specific
data, and there was even considerable variability within countries. For instance, among
studies conducted in the UK, the rates of RRM varied from 0.21 [46] to 0.43 [42], while
those for RRSO varied from 0.26 [58] to 0.86 [42]. Similarly, studies conducted in the United
States showed that the uptake rates of RRM varied from 0.36 [45] to 0.42 [42,50], and those
for RRSO from 0.33 to 0.71 [48].

(ii) Individual-related factors are also prone to affecting the preferences of women
towards RR surgeries. Individual factors that increase the uptake of RRSO include a per-
sonal history of BC [49,50,52], parity [44,46,52], and a woman’s postmenopausal status [52],
while the uptake of RRM tends to be higher among both parous women and those who
have a first-degree relative with BC [44,46]. In addition, women who had a family history
of OC were more likely to undergo any surgical option [50].

Moreover, many of these individual factors are (iii) age-dependent. While women
who have tested positive for BRCA1/2 should consider an RRSO by the age of 35 or right
after completion of childbearing [59,60], a prospective study shows that the usage of RRM
and RRSO occurs later than recommended [42]. The proportion of women that opt for
a risk-reducing surgery increases after age 40 probably because fertility is no longer a
concern, and the cumulative risk of cancer is more paramount [42]. Accordingly, the uptake
rates applied by the models were, in general, lower for women younger than 35 years, with
the lowest uptake rate applied for RRSO (10%) in women younger than 35 [19].

Finally, recent evidence indicates (iv) an improved acceptance of preventive surgeries
over time. A reason for this trend could be the improvement in genetic counseling protocols
and the cross-center knowledge transfer [61]. Increased uptake rates of risk-reducing
surgeries over time due to improved adherence have been observed for the uptake of
RRM in women with BRCA mutations, while the uptake of RRSO remained stable [58].
Nevertheless, the trend for RRM was not confirmed by the modeling studies included in
our review.

As a limitation of this literature review, it should be acknowledged that there was no
protocol registration, and a critical appraisal within sources of evidence was not conducted.
Furthermore, because sources of uptake rates were identified only if used for a cost-
effectiveness model, it is not possible to draw firm conclusions on temporal, regional, or
 cultural trends or individual factors. To evaluate these relationships more precisely, a
comprehensive literature review of observational studies has to be performed. However,
our review could demonstrate how sensitive models were when depicting the complexity
inherent to the uptake rates.

The usage of outdated sources of evidence for decision-making carries substantial
uncertainty regarding the payer’s outcomes. The improved counseling for BRCA mutation
carriers in recent years might have gradually reduced women’s reluctance in opting for
risk-reducing surgery, resulting in higher uptake rates. Hence, country-specific, prospective,
multi-center studies including post-testing counseling with respect to age and subgroups
should be performed to reflect the current status of women’s preferences for or against
surgical prevention. However, as long as updated evidence is not available, modelers—
at least in some countries—have to rely on data obtained from retrospective surveys,
cross-sectional studies, or medical records without accounting for follow-up. In this case,
assessing uncertainty associated with the uptake rates applied to the models is of utmost
importance to provide the decision-maker with a realistic assessment of the economic
consequences when adopting a screen-and-treat strategy for women with BRCA mutations.

5. Conclusions

The uptake rates of risk-reducing surgeries applied to modeling studies assessing the
cost-effectiveness of screen-and-treat strategies vary considerably. Uptake rates of surgery
are associated with high uncertainty, especially in modeling studies comparing a screen-
and-treat intervention versus a no-testing strategy. Country-specific and prospective studies
including non-directive counseling should be performed to reflect women’s preferences for
or against surgical prevention and would provide a stronger evidence base for economic modeling studies.

**Supplementary Materials:** The following supporting information can be downloaded at: [https://www.mdpi.com/article/10.3390/cancers14071786/s1](https://www.mdpi.com/article/10.3390/cancers14071786/s1), File S1: Search strategy; Table S1: Input parameters for the German model included in the value of information analysis [9]; Table S2: Input parameters for the German model not included in the value of information analysis [9]; Table S3: Excluded studies [36–38,62–70].

**Author Contributions:** Conceptualization, J.S.C.G., D.M. and S.S.; methodology, J.S.C.G. and D.M.; software, J.S.C.G.; validation, D.M. and S.K.-F.; writing—original draft preparation, J.S.C.G., D.M. and S.K.-F.; writing—review and editing, J.S.C.G., D.M., S.K.-F. and S.S.; supervision, S.S. All authors have read and agreed to the published version of the manuscript.

**Funding:** This research received no external funding.

**Conflicts of Interest:** The authors declare no conflict of interest.

**References**

1. Heemskerk-Gerritsen, B.A.M.; Jager, A.; Koppert, L.B.; Obdeijn, A.I.; Collée, M.; Meijers-Heijboer, H.E.J.; Jennen, D.J.; Oldenburg, H.S.A.; van Engelen, K.; de Vries, J.; et al. Survival after bilateral risk-reducing mastectomy in healthy BRCA1 and BRCA2 mutation carriers. *Breast Cancer Res. Treat.* 2019, 177, 723–733. [CrossRef] [PubMed]

2. Domchek, S.M.; Friebel, T.M.; Singer, C.F.; Evans, D.G.; Lynch, H.T.; Isaacs, C.; Garber, J.E.; Neuhusen, S.L.; Matloff, E.; Eeles, R.; et al. Association of risk-reducing surgery in BRCA1 or BRCA2 mutation carriers with cancer risk and mortality. *JAMA* 2010, 304, 967–975. [CrossRef] [PubMed]

3. Kotsopoulos, J.; Huzarski, T.; Gronwald, J.; Singer, C.F.; Moller, P.; Lynch, H.T.; Armel, S.; Karlan, B.; Foulkes, W.D.; Neuhusen, S.L.; et al. Bilateral Oophorectomy and Breast Cancer Risk in BRCA1 and BRCA2 Mutation Carriers. *J. Natl. Cancer Inst.* 2017, 109, djw177. [CrossRef] [PubMed]

4. Mavaddat, N.; Antoniou, A.C.; Mooij, T.M.; Hooning, M.J.; Heemskerk-Gerritsen, B.A.; Nougès, C.; Gauthier-Villars, M.; Caron, O.; Gesta, P.; Pujol, P.; et al. Risk-reducing salpingo-oophorectomy, natural menopause, and breast cancer risk: An international prospective cohort of BRCA1 and BRCA2 mutation carriers. *Breast Cancer Res.* 2020, 22, 8. [CrossRef]

5. Manchanda, R.; Abdelraheim, A.; Johnson, M.; Rosenthal, A.N.; Benjamin, E.; Brunell, C.; Burnell, M.; Side, L.; Gessler, S.; Saridogan, E.; et al. Outcome of risk-reducing salpingo-oophorectomy in BRCA carriers and women of unknown mutation status. *BJOG. Int. J. Obstet. Gynaecol.* 2011, 118, 814–824. [CrossRef] [PubMed]

6. Nelson, H.D.; Pappas, M.; Zakher, B.; Mitchell, J.P.; Okinaka-Hu, L.; Fu, R. Risk assessment, genetic counseling, and genetic testing for BRCA-related cancer in women: A systematic review to update the U.S. Preventive Services Task Force recommendation. *Ann. Intern. Med.* 2014, 160, 255–266. [CrossRef]

7. Koldenhoff, A.; Danner, M.; Civello, D.; Rhiem, K.; Stock, S.; Muller, D. Cost-Effectiveness of Targeted Genetic Testing for Breast and Ovarian Cancer: A Systematic Review. *Value Health* 2021, 24, 303–312. [CrossRef]

8. Metcalfe, K.; Eisen, A.; Senter, L.; Armel, S.; Bordeleau, L.; Meschino, W.S.; Pal, T.; Lynch, H.T.; Tung, N.M.; Kwong, A.; et al. International trends in the uptake of cancer risk reduction strategies in women with a BRCA1 or BRCA2 mutation. *Br. J. Cancer* 2019, 121, 15–21. [CrossRef]

9. Muller, D.; Danner, M.; Schmutzler, R.; Engel, C.; Wassermann, K.; Stollenwerk, B.; Stock, S.; Rhiem, K. Economic modeling of risk-adapted screen-and-treat strategies in women at high risk for breast or ovarian cancer. *Eur. J. Health Econ.* 2019, 20, 739–750. [CrossRef]

10. Trico, A.C.; Lillie, E.; Zarin, W.; O’Brien, K.K.; Colquhoun, H.; Levac, D.; Moher, D.; Peters, M.D.J.; Horsley, T.; Weeks, L.; et al. PRISMA Extension for Scoping Reviews (PRISMA-ScR): Checklist and Explanation. *Ann. Intern. Med.* 2018, 169, 467–473. [CrossRef]

11. Ficarazzi, F.; Vecchi, M.; Ferrari, M.; Pierotti, M.A. Towards population-based genetic screenings for breast and ovarian cancer: A comprehensive review from economic evaluations to patient perspectives. *Breast* 2021, 58, 121–129. [CrossRef]

12. Jayasekera, J.; Mandellblatt, J.S. Systematic Review of the Cost Effectiveness of Breast Cancer Prevention, Screening, and Treatment Interventions. *J. Clin. Oncol.* 2020, 38, 332–350. [CrossRef] [PubMed]

13. Meshkani, Z.; Aboutorabi, A.; Moradi, N.; Langarizadeh, M.; Mottaghi, A.G. Population or family history based BRCA gene tests of breast cancer? A systematic review of economic evaluations. *Hered. Cancer Clin. Pract.* 2021, 19, 35. [CrossRef] [PubMed]

14. Sroczynski, G.; Gogollari, A.; Kuehne, F.; Hallsson, L.R.; Widschwendter, M.; Pashayan, N.; Siebert, U. A Systematic Review on Cost-effectiveness Studies Evaluating Ovarian Cancer Early Detection and Prevention Strategies. *Cancer Prev. Res. 2020*, 13, 429–442. [CrossRef] [PubMed]

15. Fenwick, E.; Steuten, L.; Knies, S.; Ghabri, S.; Basu, A.; Murray, J.F.; Koffijberg, H.E.; Strong, M.; Sanders Schmider, G.D.; Rothery, C. Value of Information Analysis for Research Decisions—An Introduction. Report 1 of the ISPOR Value of Information Analysis Emerging Good Practices Task Force. *Value Health* 2020, 23, 139–150. [CrossRef]
16. Rothery, C.; Strong, M.; Koffijberg, H.E.; Basu, A.; Ghabri, S.; Knies, S.; Murray, J.F.; Sanders Schmidler, G.D.; Steuten, L.; Fenwick, E. Value of Information Analytical Methods: Report 2 of the ISPOR Value of Information Analysis Emerging Good Practices Task Force. Value Health 2020, 23, 277–286. [CrossRef]

17. Strong, M.; Oakley, J.E.; Brennan, A. Estimating multiparameter partial expected value of perfect information from a probabilistic sensitivity analysis sample: A nonparametric regression approach. Med. Decis. Mak. 2014, 34, 311–326. [CrossRef]

18. Developed for NICE by the National Collaborating Centre for Cancer. Familial Breast Cancer: Full Cost Effectiveness Evidence Review & Reports. Available online: https://www.nice.org.uk/guidance/cg164/update/CG164/documents/familial-breast-cancer-update-full-evidence-review-reports-for-health-economics2 (accessed on 1 June 2021).

19. Asphaug, L.; Melberg, H.O. The Cost-Effectiveness of Multigene Panel Testing for Hereditary Breast and Ovarian Cancer in Norway. MDM Policy Pract. 2019, 4, 2381468318821103. [CrossRef]

20. Breheny, N.; Geelhood, E.; Goldblatt, J.; O’Leary, P. Cost-Effectiveness of Predictive Genetic Tests for Familial Breast and Ovarian Cancer. Available online: https://isspjournal.biomedcentral.com/track/pdf/10.1186/1746-5996-5-67.pdf (accessed on 1 June 2021).

21. Eccleston, A.; Bentley, A.; Dyer, M.; Strydom, A.; Vereecken, W.; George, A.; Rahman, N. A Cost-Effectiveness Evaluation of Germline BRCA1 and BRCA2 Testing in UK Women with Ovarian Cancer. Value Health 2017, 20, 567–576. [CrossRef]

22. Guzautkas, G.F.; Garbett, S.; Zhou, Z.; Spencer, S.J.; Smith, H.S.; Hao, J.; Hassen, D.; Snyder, S.R.; Graves, J.A.; Peterson, J.F.; et al. Cost-effectiveness of Population-Wide Genomic Screening for Hereditary Breast and Ovarian Cancer in the United States. JAMA Netw. Open 2020, 3, e2022874. [CrossRef]

23. Holland, M.L.; Huston A Fau-Noyes, K.; Noyes, K. Cost-effectiveness of testing for breast cancer susceptibility genes. Value Health 2009, 12, 207–216. [CrossRef] [PubMed]

24. Hurry, M.; Eccleston, A.; Dyer, M.; Hoskins, P. Canadian cost-effectiveness model of BRCA-driven surgical prevention of breast/ovarian cancers compared to treatment if cancer develops. Int. J. Technol. Assess. Health Care 2020, 36, 104–112. [CrossRef] [PubMed]

25. Kemp, Z.; Turnbull, A.; Yost, S.; Seals, S.; Mahamadlallie, S.; Poyastro-Pearson, E.; Warren-Perry, M.; Eccleston, A.; Tan, M.M.; Teo, S.H.; et al. Evaluation of Cancer-Based Criteria for Use in Mainstream BRCA1 and BRCA2 Genetic Testing in Patients with Breast Cancer. JAMA Netw. Open 2019, 2, e194428. [CrossRef] [PubMed]

26. Kwon, J.S.; Gutierrez-Barrera, A.M.; Young, D.; Sun, C.C.; Daniels, M.S.; Lu, K.H.; Arun, B. Expanding the criteria for BRCA mutation testing in breast cancer survivors. J. Clin. Oncol. 2010, 28, 4214–4220. [CrossRef]

27. Kwon, J.S.; Tinker, A.V.; Hanley, G.E.; Pansegrau, G.; Sun, S.; Carey, M.S.; Schrader, I. BRCA mutation testing for first-degree relatives of women with high-grade serous ovarian cancer. Gynecol. Oncol. 2019, 152, 459–464. [CrossRef]

28. Li, Y.; Arellano, A.R.; Bare, L.A.; Bender, R.A.; Strom, C.M.; Devlin, J.J. A Multigene Test Could Cost-Effectively Help Extend Life Expectancy for Women at Risk of Hereditary Breast Cancer. Value Health 2017, 20, 547–555. [CrossRef] [PubMed]

29. Manchanda, R.; Sun, L.; Patel, S.; Evans, O.; Wilchust, J.; de Freitas Lopes, A.C.; Gaba, F.; Brentnall, A.; Duffy, S.; Cui, B.; et al. Economic Evaluation of Population-Based BRCA1/BRCA2 Mutation Testing across Multiple Countries and Health Systems. Cancers 2020, 12, 1929. [CrossRef]

30. Moya-Alarcon, C.; Gonzalez-Dominguez, A.; Simon, S.; Perez-Roman, I.; Gonzalez-Martin, A.; Bayo-Lozano, E.; Sanchez-Heras, A.B. Cost-utility analysis of germline BRCA1/2 testing in women with high-grade epithelial ovarian cancer in Spain. Clin. Transl. Oncol. 2019, 21, 1076–1084. [CrossRef]

31. Petelin, L.; Hossack, L.; Shanahan, M.; Mitchell, G.; Liew, D.; James, P.A.; Trainer, A.H. Cost-effectiveness of long-term clinical management of BRCA pathogenic variant carriers. Genet. Med. 2020, 22, 831–839. [CrossRef]

32. Ramos, M.C.A.; Folgueira, M.; Maistro, S.; Campolina, A.G.; Soarez, P.C.; Bock, G.H.; Novaes, H.M.D.; Diz, M. Cost effectiveness of the cancer prevention program for carriers of the BRCA1/2 mutation. Rev. Saude Publica 2018, 52, 94. [CrossRef]

33. Simes Correa-Galendi, J.; del Pilar Estevez Diz, M.; Stock, S.; Müller, D. Economic Modelling of Screen-and-Treat Strategies for Brazilian Women at Risk of Hereditary Breast and Ovarian Cancer. Appl. Health Econ. Health Policy 2021, 19, 97–109. [CrossRef] [PubMed]

34. Sun, L.; Brentnall, A.; Patel, S.; Buist, D.S.M.; Bowles, E.J.A.; Evans, D.G.R.; Eccles, D.; Hopper, J.; Li, S.; Southey, M.; et al. A Cost-effectiveness Analysis of Multigene Testing for All Patients with Breast Cancer. JAMA Oncol. 2019, 5, 1718–1730. [CrossRef] [PubMed]

35. Tuffaha, H.W.; Mitchell, A.; Ward, R.L.; Connelly, L.; Butler, J.R.G.; Norris, S.; Scuiffham, P.A. Cost-effectiveness analysis of germ-line BRCA testing in women with breast cancer and cascade testing in family members of mutation carriers. Genet. Med. 2018, 20, 985–994. [CrossRef] [PubMed]

36. Norum, J.; Grindedal, E.M.; Heramb, C.; Karsrud, I.; Ariansen, S.L.; Undlien, D.E.; Schlichting, E.; Maehle, L. BRCA mutation carrier detection. A model-based cost-effectiveness analysis comparing the traditional family history approach and the testing of all patients with breast cancer. ESMO Open 2018, 3, e000328. [CrossRef] [PubMed]

37. Hoskins, P.; Eccleston, A.; Hurry, M.; Dyer, M. Targeted surgical prevention of epithelial ovarian cancer is cost effective and saves money in BRCA mutation carrying families of women with epithelial ovarian cancer. A Canadian model. Gynecol. Oncol. 2019, 153, 87–91. [CrossRef] [PubMed]

38. Tengs, T.O.; Berry, D.A. The Cost Effectiveness of Testing for the BRCA1 and BRCA2 Breast-Ovarian Cancer Susceptibility Genes. Available online: http://www.scopus.com/inward/record.url?scp=0033997009&partnerID=8YFLogXK (accessed on 2 June 2021).

39. Carbonara, N.; de Jong, D.; Pellegrino, R.; Ressa, C.; Tommasi, S. A Cost Decision Model Supporting Treatment Strategy Selection in BRCA1/2 Mutation Carriers in Breast Cancer. J. Pers. Med. 2021, 11, 847. [CrossRef] [PubMed]
62. Balmana, J.; Sanz, J.; Bonfill, X.; Casado, A.; Rue, M.; Gich, I.; Diez, O.; Sabate, J.M.; Baiget, M.; Alonso, M.C. Genetic counseling program in familial breast cancer: Analysis of its effectiveness, cost and cost-effectiveness ratio. *Int. J. Cancer* 2004, 112, 647–652. [CrossRef]

63. Gamble, C.; Havrilesky, L.J.; Myers, E.R.; Chino, J.P.; Hollenbeck, S.; Plichta, J.K.; Kelly Marcom, P.; Shelley Hwang, E.; Kauff, N.D.; Greenup, R.A. Cost Effectiveness of Risk-Reducing Mastectomy versus Surveillance in BRCA Mutation Carriers with a History of Ovarian Cancer. *Ann. Surg. Oncol.* 2017, 24, 3116–3123. [CrossRef]

64. Patel, S.; Legood, R.; Evans, D.G.; Turnbull, C.; Antoniou, A.C.; Menon, U.; Jacobs, I.; Manchanda, R. Cost-effectiveness of population based BRCA1 founder mutation testing in Sephardi Jewish women. *Am. J. Obstet. Gynecol.* 2018, 218, e1–e12. [CrossRef]

65. Rubinstein, W.S.; Jiang, H.; Dellefave, L.; Rademaker, A.W. Cost-effectiveness of population-based BRCA1/2 testing and ovarian cancer prevention for Ashkenazi Jews: A call for dialogue. *Genet. Med.* 2009, 11, 629–639. [CrossRef] [PubMed]

66. Manchanda, R.; Legood, R.; Pearce, L.; Menon, U. Defining the risk threshold for risk reducing salpingo-oophorectomy for ovarian cancer prevention in low risk postmenopausal women. *Gynecol. Oncol.* 2015, 139, 487–494. [CrossRef] [PubMed]

67. Manchanda, R.; Legood, R.; Antoniou, A.C.; Gordeev, V.S.; Menon, U. Specifying the ovarian cancer risk threshold of ‘pre-menopausal risk-reducing salpingo-oophorectomy’ for ovarian cancer prevention: A cost-effectiveness analysis. *J. Med. Genet.* 2016, 53, 591–599. [CrossRef] [PubMed]

68. Manchanda, R.; Patel, S.; Antoniou, A.C.; Levy-Lahad, E.; Turnbull, C.; Evans, D.G.; Hopper, J.L.; Macinnis, R.J.; Menon, U.; Jacobs, I.; et al. Cost-effectiveness of population based BRCA testing with varying Ashkenazi Jewish ancestry. *Am. J. Obstet. Gynecol.* 2017, 217, 578.e1–578.e12. [CrossRef]

69. Manchanda, R.; Patel, S.; Gordeev, V.S.; Antoniou, A.C.; Smith, S.; Lee, A.; Hopper, J.L.; MacInnis, R.J.; Turnbull, C.; Ramus, S.; et al. Cost-effectiveness of Population-Based BRCA1, BRCA2, RAD51C, RAD51D, BRIP1, PALB2 Mutation Testing in Unselected General Population Women. *J. Natl. Cancer Inst.* 2018, 110, 714–725. [CrossRef]

70. Zhang, L.; Bao, Y.; Riaz, M.; Tiller, J.; Liew, D.; Zhuang, X.; Amor, D.J.; Huq, A.; Petelin, L.; Nelson, M.; et al. Population genomic screening of all young adults in a health-care system: A cost-effectiveness analysis. *Genet. Med.* 2019, 21, 1958–1968. [CrossRef]