Clinical benefit of systemic therapies for recurrent ovarian cancer—ESMO-MCBS scores

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Background: Licensed systemic treatment options for platinum-sensitive recurrent ovarian cancer are platinum-based chemotherapy and maintenance treatment with bevacizumab and poly (ADP-ribose) polymerase inhibitors. For platinum-resistant disease, several non-platinum options are available. We aimed to assess the clinical benefit of these treatments according to the European Society of Medical Oncology (ESMO)-Magnitude of Clinical Benefit Scale (MCBS).

Materials and methods: A PubMed search was carried out including all studies evaluating systemic treatment of recurrent epithelial ovarian cancer, from 1990 onwards. Randomised trials with an adequate comparator and design showing a statistically significant benefit of the study arm were independently scored by two blinded observers using the ESMO-MCBS.

Results: A total of 1127 papers were identified, out of which 61 reported results of randomised trials of sufficient quality. Nineteen trials showed statistically significant results and the studied treatments were graded according to ESMO-MCBS. Only three treatments showed substantial benefit (score of 4 on a scale of 1-5) according to the ESMO-MCBS: platinum-based chemotherapy with paclitaxel in the platinum-sensitive setting and the addition of bevacizumab to chemotherapy in the platinum-resistant setting. The WEE1 inhibitor adavosertib (not licensed) also scores a 4, based on a recent small phase II study. Assessment of quality-of-life data and toxicity using the ESMO-MCBS showed to be complex, which should be taken into account in using this score for clinical decision making.

Conclusion: Only a few licensed systemic therapies for recurrent ovarian cancer show substantial clinical benefit based on ESMO-MCBS scores. Trials demonstrating overall survival benefit are sparse.

Key words: clinical benefit, ovarian cancer, ESMO-MCBS, chemotherapy, targeted therapy

INTRODUCTION

First-line therapy for advanced epithelial ovarian cancer (EOC) consists of debulking surgery and platinum-based chemotherapy. EOC is usually diagnosed at an advanced stage, which has a poor prognosis with the majority of women eventually developing recurrent disease. Ovarian cancer-specific survival at 5 years is 40% for stage III disease and only 20% for stage IV disease.1 Treatment options and prognosis differ depending on the interval between first-line treatment and relapse (the platinum-free interval, PFI), with the highest chance of response to reintroduction of platinum-based chemotherapy in patients with a PFI of >12 months. Patients with relapse occurring within 6 months of platinum-based chemotherapy were historically considered platinum-resistant. Recently, the strict definition of platinum-resistant disease was abandoned. It is recognised that the time elapsed since the last platinum chemotherapy represents a continuum of probability of response to further chemotherapy. Treatment options mentioned in guidelines for this setting have limited response rates (RRs) in patients with a short PFI and most patients die of their disease within 1 year.2 Therapies licensed for treatment of recurrent ovarian cancer, besides carboplatin and paclitaxel, are gemcitabine and bevacizumab (in combination with platinum), liposomal doxorubicin with or without trabectedin, treosulfan, melphalan, topotecan, etoposide and the poly (ADP-ribose) polymerase (PARP) inhibitors olaparib, niraparib and rucaparib. Guidelines mention these drugs as treatment options and advise that the treatment regimen should be chosen based on platinum sensitivity, previously received treatments, BRCA mutation status and physician and patients'...
preferences. The European Society of Medical Oncology (ESMO)-Magnitude of Clinical Benefit Scale (MCBS) is an instrument to score the clinical benefit of anticancer therapies taking into account efficacy, quality of life (QoL) and toxicity. We aimed to assess the clinical benefit of systemic therapies in recurrent ovarian cancer according to the ESMO-MCBS.

**MATERIALS AND METHODS**

**Search strategy and selection**

A PubMed search was carried out in 2019 and updated in March 2021 using the following Medical Subject Headings (MeSH) terms: ‘recurrent’ OR ‘refractory’ AND ‘ovarian’ OR ‘ovary’ AND ‘neoplasm’ OR ‘cancer’ OR ‘carcinoma’; the search was limited to trials in humans. Studies evaluating systemic treatment in recurrent EOC patients were screened, from the introduction of taxanes in 1990 onwards.

Titles and abstracts were screened independently by two observers (KEB and MvK) and categorised into three categories: category I contained large phase III randomised controlled trials (RCTs) in the target population with an adequate comparator and a primary endpoint of overall survival (OS) or progression-free survival (PFS); category II for other studies that could potentially be scored on the ESMO-MCBS; and category III for studies that did not meet the inclusion criteria for scoring (e.g. wrong population, surgical interventions only, review articles, retrospective studies and case series). After all articles had been assigned to a category, the assessments of the two independent observers were compared and conflicting results were resolved at a consensus meeting. Studies in category I and II were analysed in more detail by reading the full paper, assessing study methodology, patient numbers, study design, endpoints and outcome. From category II, only randomised studies comparing a new agent or combination to a clinically relevant comparator were included, and thus single-arm studies and studies evaluating different dose regimens of the same drug were excluded from the analysis. Non-platinum-based chemotherapy was inferior to platinum-based chemotherapy in the MITO-8 study, and platinum-based chemotherapy was therefore considered the preferred comparator treatment for the included trials in the platinum-sensitive setting. For the platinum-resistant setting, any treatment option mentioned in the ESMO or American Society of Clinical Oncology guidelines was considered an adequate comparator.

Finally, references of included studies and relevant guidelines were checked to identify relevant publications not retrieved by the search strategy. All included studies showing a statistically significant benefit of the study arm were graded according to the ESMO-MCBS.

**Grading according to ESMO-Magnitude of Clinical Benefit Scale**

The ESMO-MCBS can be used to rank the value of systemic therapies based on reported relative and absolute benefits in terms of improved survival (PFS, disease-free survival, OS) and better survival (e.g. QoL, toxicity). For palliative treatments, as is the case in recurrent ovarian cancer, there are several MCBS evaluation forms available and the correct form is chosen based on the primary endpoint used in the study (OS, PFS, QoL, toxicity or RR) and the duration of survival in the control group. Palliative treatments are graded 1-5, where grades 4 and 5 represent a substantial clinical benefit. The preliminary score for palliative treatments is upgraded when the study treatment shows an improvement in QoL or a reduction in grade 3-4 toxicities impacting daily well-being. For treatments that only show a benefit in PFS, but not OS, the preliminary score is down-graded if the study treatment has increased toxicity or does not demonstrate improvement in QoL. The MCBS allows for scoring of clinical benefit in a maximum of three pre-specified subgroups, provided adjustments for multiple comparisons are taken into account.

**RESULTS**

**Search results and trial selection**

Our search retrieved 1127 studies in ovarian cancer published since 1990. There was an excellent overall agreement between the two observers (98%) when categorising the studies into the three specified groups. A flow chart indicating the selection procedure is shown in Figure 1. Sixty-five studies were directly included based on the abstract (listed as category I by both observers). Nine hundred and fifty-two studies were directly excluded based on the abstract (category III). The remaining 110 studies (category II) were reviewed in more detail and discussed at a consensus meeting. Sixteen studies from category II matched criteria for inclusion in the analysis. After detailed assessment of full-text articles, 29 RCTs from category I and II were excluded because of an inadequate comparator, premature termination of the trial or the availability of a higher-quality trial studying the same treatment (Supplementary Table S1, available at https://doi.org/10.1016/j.esmoop.2021.100229).

Finally, nine additional studies were included after checking relevant guidelines and references.

In total, 61 publications reporting the results of 46 original clinical trials were assessed. Fourteen phase III studies and five randomised phase II studies with statistically significant results were graded for clinical benefit using the ESMO-MCBS (Tables 1 and 2). The remaining 27 studies did not report a statistically significant result and were therefore not graded on the ESMO-MCBS (Table 3).

**Clinical benefit of treatment in platinum-sensitive recurrent ovarian cancer**

**Platinum-based chemotherapy.** Nine randomised phase III studies evaluating chemotherapy in platinum-sensitive disease in a total of 5470 patients were assessed for clinical benefit. Of these nine trials, five (55%) showed significant improvement of either PFS (3/9), OS (1/9) or QoL (1/9) (Table 1).
With regard to the platinum-based options, the highest clinical benefit (ESMO-MCBS grade 4) was obtained for the combination of paclitaxel plus platinum-based chemotherapy. In the ICON4 trial, comparing the combination of platinum-based chemotherapy with paclitaxel to carboplatin or cisplatin monotherapy, the combination resulted in a 5-month improvement of OS (median OS 29 versus 24 months; hazard ratio (HR) 0.82 [95% confidence interval (CI) 0.69-0.97]). Carboplatin/paclitaxel was used as the comparator treatment in trials carried out after ICON4. Treatment with carboplatin plus pegylated liposomal doxorubicin (PLD) has a lower ESMO-MCBS grade (grade 3) than carboplatin/paclitaxel, but the comparator arm in these two studies was different. Namely, while carboplatin/paclitaxel was compared to platinum monotherapy in the ICON4 trial, carboplatin plus PLD was compared to carboplatin/paclitaxel in the CALYPSO study. In this non-inferiority study, PFS was longer for carboplatin/PLD as compared to carboplatin/paclitaxel [median PFS 11.3 versus 9.4 months; HR 0.82 (0.72-0.94)]. Global QoL scores were equal, but there was an improvement in various symptom subscales with carboplatin/PLD, including peripheral neuropathy, global chemotherapy side-effects and impact on body image resulting in a grade 3 score (form 2c). Early treatment discontinuation was also less frequent for carboplatin/PLD (6%) than for carboplatin/paclitaxel (15%).

The remaining three trials with significant results investigated carboplatin doublet chemotherapy plus bevacizumab (versus carboplatin doublet chemotherapy), carboplatin plus gemcitabine (versus carboplatin monotherapy) and carboplatin plus PLD plus bevacizumab (versus carboplatin/gemcitabine/bevacizumab), which all showed an improved PFS over the comparator, but did not result in OS benefit and showed no improved QoL. This qualifies as only limited clinical benefit with ESMO-MCBS grades of 3, 2 and 1, respectively. An overview of the relative benefit between the different arms and comparators studied and graded for the platinum-sensitive setting is provided in Figure 2A.
| Treatment | Study name | Subgroups | n | Control | Primary endpoint | PFS control | PFS gain | HR control | OS gain | OS HR | ORR | Toxicity/QoL adjustment | MCBS score (form) | Ref. |
|-----------|------------|-----------|---|---------|------------------|-------------|----------|------------|---------|-------|-----|--------------------------|-----------------|------|
| Platinum-sensitive recurrence | Carboplatin/PLD | CALYPSO | 976 | Carboplatin plus paclitaxel | PFS (non-inferiority) | 9.4 | 1.9 | 0.82 (0.72-0.94) | 30.7 | 2.3 | 0.99 (0.85-1.16) | Global QoL equal but better scores on subscales for carboplatin/PLD (less neurotoxicity, better body image; early discontinuation in 6% versus 15%) | 3 (2c) | 8,9 |
| | Paclitaxel plus platinum-based chemotherapy | ICON4/AGO-OVAR 2.2 | 802 | Platinum monotherapy (carboplatin or cisplatin) | OS | 24 | 5 | 0.82 (0.69-0.97) | 24.7 | 2.3 | 0.99 (0.85-1.16) | | 4 (2a) | 7 |
| | Carboplatin plus gemcitabine | | 356 | Carboplatin | PFS | 5.8 | 2.8 | 0.72 (0.58-0.90) | 17.3 | 0.7 | 0.96 (0.75-1.23) | –1 | No QoL benefit and no OS benefit | 2 (2b) | 10 |
| | Carboplatin/PLD/ bevacizumab | | 682 | Carboplatin/ gemcitabine/ bevacizumab | PFS | 11.7 | 1.7 | 0.81 (0.68-0.91) | 27.8 | 4.1 | 0.81 (0.67-0.98) | 1 | New QoL benefit and no OS benefit | 1 (2b) | 11 |
| | Carboplatin doublet plus bevacizumab | MITO16b/ MANGO-OV2/ ENGOT-ov17 | 406 | Carboplatin doublet | PFS | 8.8 | 3 | 0.51 (0.41-0.65) | | | | 3 (2b) | 12 |
| Maintenance therapy after response to second-line platinum-based chemotherapy | Cediranib | ICON6 - ITT unselected | 282 | Placebo | PFS | 8.7 | 2.3 | 0.56 (0.44-0.72) | 0.86 (0.67-1.11) | –1 | No QoL benefit and no OS benefit | 1 (2b) | 21,28 |
| | Niraparib | | 553 | Placebo | PFS | 5.5 | 14.5 | 0.27 (0.17-0.41) | 0.45 (0.34-0.61) | 0.38 (0.24-0.59) | | | 3 (2b) | 3 (2b) | 13 |
| | Rucaparib (ITT) | ARIEL3 | | Placebo | PFS | 5.4 | 11.2 | 0.36 (0.30-0.45) | 0.23 (0.16-0.34) | 0.32 (0.24-0.42) | | | 3 (2b) | 3 (2b) | 18 |
| | Olaparib (tablets) | SOLO2 - ITT All had gBRCA1/2 mutation | 295 | Placebo | PFS | 5.5 | 13.6 | 0.30 (0.22-0.41) | 38.8 | 12.9 | 0.74 (0.54-1.00) | –1 for no OS benefit | 2 (2b) | 20,27 |
| | Addition of bevacizumab to carboplatin plus gemcitabine | OCEANS | 484 | Carboplatin plus gemcitabine | PFS | 8.4 | 4 | 0.48 (0.39-0.61) | 32.9 | 0.7 | 0.95 (0.77-1.18) | | | 2 (2b) | 22,23 |

Continued
Studies with negative results included the addition of the folate receptor antibody farletuzumab to carboplatin plus taxane,17 carboplatin plus topotecan14 and carboplatin plus micellar paclitaxel.15

In conclusion, for platinum-sensitive recurrence, the first-choice treatment is a carboplatin-based combination regimen. Direct comparison in the CALYPSO trial demonstrated that carboplatin plus PLD has a slightly better toxicity profile than carboplatin/paclitaxel. Carboplatin/paclitaxel renders comparable OS16 but may result in more toxicity and early discontinuation.

Maintenace therapy after platinum-based chemotherapy. Nine randomised studies in 3394 patients evaluated maintenance therapy after response to platinum-based chemotherapy for recurrent disease. Seven of nine trials (78%) reported significant results: three phase III RCTs evaluating maintenance therapy with PARP inhibitors,17-20 one phase III RCT assessing maintenance with cediranib, a tyrosine kinase inhibitor targeting the vascular endothelial growth factor receptor,21 one phase III RCT studying bevacizumab maintenance22,23 and two phase II RCTs with the PARP inhibitor olaparib (capsules).24,25 In six studies, maintenance therapy was initiated only after response to platinum-based chemotherapy for recurrent disease. Three studies were conceptually different because the investigational agent was started concomitantly with chemotherapy and continued thereafter (one study with olaparib25 and two studies with bevacizumab22,23).

The phase II STUDY19 compared olaparib capsules with placebo: there was no significant improvement in OS or QoL for the intention-to-treat (ITT) population or for the prespecified BRCA-mutated subgroup (germline or somatic). This resulted in the preliminary score of 3 for the observed PFS benefit which was downgraded to a grade 2 in both populations.24 Treatment with olaparib tablets was assessed in the phase III SOLO2 trial, in which all included patients had a germline BRCA (gBRCA) mutation.20 The mature OS results were recently published.27 The observed OS benefit in the ITT population (n = 295) did not reach statistical significance with an HR of 0.74 (0.54-1.00; P = 0.054), which results in a downgrade to grade 2 for the ITT gBRCA-mutated population. Remarkably, a pre-planned sensitivity analysis in patients with Myriad-confirmed gBRCA mutation (n = 286 of the total 295 patients) did show a statistically significant OS benefit of 15 months [HR 0.71 (0.52-0.97); P = 0.0306] compared to placebo (OS 37.4 months). However, because there was no correction for multiple testing, this analysis was considered exploratory and the results were not scored on the ESMO-MCBS.

Treatment with PARP inhibitors niraparib and rucaparib resulted in moderate benefit according to ESMO-MCBS (grade 3) when compared to placebo, all based on PFS as primary endpoint. This score was the same for the benefit in the unselected ITT population as well as the prespecified subgroup analyses based on BRCA status, although the included study populations differed somewhat between studies. The absolute benefit was largest for
Table 2. Phase II studies scored for clinical benefit (n = 5)

| Treatment | Study name | Subgroups | n | Control | Primary endpoint | PFS control | PFS gain | HR OS | OS gain | OS HR | Toxicity/QoL adjustment | MCBS score | Ref. |
|-----------|------------|-----------|---|---------|-----------------|-------------|----------|-------|---------|-------|---------------------------|------------|------|
| Maintenance therapy after response to second line platinum-based chemotherapy | Olaparib (capsules) | STUDY19 | - ITT | 265 Placebo | PFS | 4.8 | 3.6 | 0.35 | (0.25-0.49) | 27.8 | 2 | 0.73* | (0.55-0.95) | 6 | 2 (2b) |
| | | | - BRCA-mut | 136 | 4.3 | 6.9 | 0.18 | (0.10-0.31) | 30.2 | 4.7 | 0.62 | (0.42-0.93) | 8 | 2 (2b) |
| | | | - BRCA-wt | 118 | 5.5 | 1.9 | 0.54 | (0.34-0.85) | 26.6 | -2.1 | 0.84 | (0.57-1.25) | 2 | 2 (2b) |
| Olaparib (capsules) during carboplatin/paclitaxel followed by olaparib maintenance | - ITT unselected | 173 | Carboptatin/paclitaxel followed by no further treatment | PFS | 9.6 | 2.6 | 0.51 | (0.34-0.77) | 2 | (2b) |
| Platinum-resistant recurrence | Sorafenib plus topotecan | TRIAS | 185 | Topecan plus placebo | PFS | 4.4 | 2.3 | 0.60 | (0.43-0.83) | 3 | (2b) |
| | Berzosertib plus gemcitabine | 70 | Gemcitabine | PFS | 3.4 | 1.9 | 0.57 | (0.33-0.98) | 9.9 | 3.8 | 0.84 | (0.53-1.32) | 2 | (2b) |
| | Adavosertib plus gemcitabine | 99 | Gemcitabine | PFS | 3 | 1.6 | 0.55 | (0.35-0.90) | 7.2 | 4.2 | 0.56 | (0.35-0.91) | 4 | (2a) |

HR, hazard ratio; ITT, intention to treat; MCBS, Magnitude of Clinical Benefit Scale; OS, overall survival; PFS, progression-free survival; QoL, quality of life; w/t, wild type.

* P value did not meet prespecified study criteria for significant OS result.

Clinical benefit of treatment in the platinum-resistant setting

A total of 8137 patients with platinum-resistant ovarian cancer were treated in 28 randomised studies included in our analysis, out of which only 6 trials (21%) resulted in significant improvement of either PFS (5/6) or OS (1/6). Two trials showed substantial benefit (ESMO-MCBS grade 4) in this setting. In the AURELIA study, with either PIDD, a PARP inhibitor showed a median OS benefit of 4.2 months compared to placebo (HR 0.64 (95% CI 0.48-0.88)). Although OS was similar in the physician-choice chemotherapy arm (HR 0.70 (95% CI 0.50-0.98)).

HR 0.48 (95% CI 0.38-0.66). Although OS was similar in HR 0.48 (95% CI 0.38-0.66). Although OS was similar in HR 0.48 (95% CI 0.38-0.66) and HR 0.48 (95% CI 0.38-0.66), the benefit of olaparib maintenance therapy after response to second line platinum-based chemotherapy was not seen in the Phase III study.

In conclusion, despite the promising results, PARP inhibitors do not provide substantial benefit in patients with platinum-sensitive recurrence. The benefit of olaparib maintenance therapy after response to platinum-based chemotherapy was not seen in the Phase III study.
| Treatment                                      | Study name              | n   | Control Design | Primary endpoint | PFS control | PFS gain (HR, CI) | OS control | OS gain (HR, CI) | OS HR | ORR | Toxicity/ QoL | ESMO grading | Ref. |
|------------------------------------------------|-------------------------|-----|----------------|-----------------|-------------|------------------|-------------|------------------|--------|-----|---------------|--------------|------|
| Platinum-sensitive recurrence                  | Farletuzumab with carboplatin/taxane | 1100| Placebo with carboplatin/taxane | Phase III RCT | PFS | 9.0 | 0.7 | 0.99 (0.81-1.21) | Not significant | 13   |
|                                                | Carboplatin/ topotecan   | 550 | Carboplatin plus paclitaxel, gemcitabine or PLD | Phase III | PFS | 10 | 0 | Not provided | Not significant | 14   |
| Non-platinum-based CT                           | MITO-8                  | 215 | Platinum-based CT | Phase III RCT | OS | 9 | −4 | Not provided | 24.5 | −2.7 | Not provided | PFS control arm significantly better | 5    |
|                                                | Carboplatin/ paclitaxel micellar | 789 | Carboplatin/paclitaxel (conventional) | Phase III non-inferiority RCT | Non-inferiority for PFS | 10.1 | 0.2 | 0.86 (0.72-1.03) | Increased neutropenia, no difference in neurotoxicity | 15   |
| Maintenance therapy after response to second-line platinum-based chemotherapy | Vismodegib maintenance after complete response to second- or third-line chemotherapy | 104 | Placebo | Phase II RCT | PFS | 5.8 | 1.7 | 0.79 (0.46-1.35) | Not significant | 29   |
|                                                | Carboplatin/ paclitaxel plus bevacizum | 674 | Carboplatin/paclitaxel plus placebo | Phase III RCT | OS | 37.3 | 4.9 | 0.83 (0.68-1.01) | Not significant | 26   |
| Platinum-resistant recurrence                  | PLD³                    | 474 | Topotecan | RCT | PFS | 3.9 | −0.2 | NS | 13.8 | 0.69 | 0.82 (0.68-1.00) | Not significant | 46,47 |
|                                                | Epidoxorubicin plus paclitaxel¹ | 234 | Paclitaxel | Phase III RCT | OS | 14.0 | −2.0 | NS | Not significant | 48   |
|                                                | PLD                     | 195 | Gemcitabine | Phase III RCT | PFS | 3.1 | 0.5 | NS | Not significant | 49   |
|                                                | PLD³                    | 153 | Gemcitabine | Phase III RCT | TTP | 3.7 | 0.9 | NS | Not significant | 50   |
| Topotecan/etoposide (TE) or topotecan/ gemcitabine | PLD                    | 502 | Topotecan | Phase III RCT | OS | 17.2 | 0.6 (TE) | 1.18 | 0.90-1.53 | Not significant | 51   |
|                                                | Canfosfamide"           | 461 | PLD or topotecan | Phase III RCT | OS | 13.5 | −5.0 | 1.71 | Control arm significantly better | 52   |
|                                                | Pertuzumab plus gemcitabine | 130 | Gemcitabine plus placebo | Phase II RCT | PFS | 2.9 | 0 | 0.66 (0.43-1.03) | Not significant | 53   |
|                                                | Pertuzumab plus carboplatin and either paclitaxel or gemcitabine | 149 | Carboplatin and either paclitaxel or gemcitabine | Phase II RCT | PFS | 8.6 | −0.7 | 1.16 (0.90-1.49) | Not significant | 54   |
|                                                | Paclitaxel/carboplatin OR paclitaxel/ topotecan | 165 | Paclitaxel | Phase II RCT | PFS | 3.7 | 1.1 | 0.92, NS | 1.7 | 0.95, NS | Not significant | 55   |
|                                                | Olaparib 200 mg or 400 mg " | 97  | PLD | Phase II RCT | PFS | 7.1 | NA | 0.88 (0.62-1.28) | Not significant | 56   |
|                                                | Patupilone              | 829 | PLD | Phase III RCT | OS | 12.7 | 0.5 | 0.93 (0.79-1.09) | Not significant | 57   |
|                                                | Docetaxel plus vandetanib | 129 | Docetaxel | RCT | PFS | 3.5 | −0.5 | 1.01 (0.79-1.27) | Not significant | 58   |

Continued
### Table 3. Continued

| Treatment | Design | Primary end point | Control arm | OS control gain | OS HR | OS gain | ESRO grading | Ref. |  |
|-----------|--------|-------------------|-------------|-----------------|-------|---------|--------------|------|---|
| Seribantumab plus paclitaxel RCT PFS | Phase II RCT PFS | 5.2 | 0.4 | 1.1 | 18.9 | 0.8 | 1.22 | Not significant | 59 |
| Motolimod plus PLD 297 PLD Phase II RCT PFS 5.2 | 0.4 | 1.21 | 18.9 | 0.8 | 1.22 | Not significant | 60 |
| Oncolytic reovirus plus paclitaxel Phase II RCT PFS 4.3 | 0.1 | 1.11 | 18.9 | 0.8 | 1.22 | Not significant | 61 |
| Pazopanib plus paclitaxel Phase II RCT PFS 6.2 | 1.3 | 0.84 | 18.9 | 0.8 | 1.22 | Not significant | 62 |
| Cabozantinib plus paclitaxel Phase II RCT PFS 5.5 | 1.1 | 0.84 | 18.9 | 0.8 | 1.22 | Not significant | 63 |
| Linsitinib (two different regimens) plus paclitaxel Phase II RCT PFS 5.6 | 1.4 | 0.77 | 18.9 | 0.8 | 1.22 | Not significant | 64 |
| Olaratumab plus PLD 123 PLD Phase II RCT PFS 4.0 | 0.2 | 1.04 | 18.9 | 0.8 | 1.22 | Not significant | 65 |
| Alisertib plus paclitaxel Phase I/II RCT PFS 4.7 | 2.0 | 0.75 | 18.9 | 0.8 | 1.22 | Not significant | 66 |

**Endpoints:**
- **OS:** Overall survival
- **PFS:** Progression-free survival
- **HR:** Hazard ratio
- **ORR:** Objective response rate
- **Toxicity/QoL:** Quality of life

**Grading:**
- ESMO grading: 0-4
- Control arm: 0-1
- OS control gain: 0-3
- OS HR: 0-4
- OS gain: 0-5

**Ref.:** Not significant

**Note:**
- The addition of bevacizumab did result in improvement in a predefined analysis of a subscale of abdominal/gastrointestinal (GI) symptoms. The preliminary score of 3 was upgraded to an ESMO-MCBS grade 4 based on this QoL benefit. The combination of the WEE1 inhibitor adavosertib with gemcitabine also improved both PFS and OS compared to gemcitabine monotherapy [median OS 7.2 versus 11.4 months; HR 0.56 (95% CI 0.35-0.91)]. However, this was assessed in a phase II study and a difference in prognostic baseline factors between groups might bias the results of this small trial. Therefore, the ESMO-MCBS score of 4 should be considered as preliminary.

One trial showed moderate benefit (grade 3) of sorafenib plus topotecan compared to topotecan alone, with a statistically significant 2.3-month PFS gain, evaluated in a phase II study. Other treatments, such as PLD plus trabectedin compared to PLD monotherapy, trebananib plus paclitaxel compared to paclitaxel monotherapy and the ataxia telangiectasia and Rad3-related protein (ATR) inhibitor berzosertib plus gemcitabine compared to gemcitabine monotherapy, had only limited benefit (grade 2).

An overview of the relative benefit between the different arms and comparators studied and graded for the platinum-resistant setting is provided in Figure 2B.

Therapies studied in the platinum-resistant setting that were unable to improve PFS or OS are listed in Table 3. This includes studies that compared the different agents used in the comparator arm of the AURELIA trial (PLD, paclitaxel or topotecan), making it difficult to recommend one of these agents over the other. Therefore, due to lack of convincing evidence, the choice of therapy will largely be based on previously received treatments and physician and patients’ preferences regarding scheduling and side-effects.

In conclusion, topotecan, PLD and paclitaxel are used for treatment of platinum-refractory ovarian cancer but there are no studies comparing these therapies to best supportive care. Addition of bevacizumab to these agents improved PFS and some symptoms; however, no OS benefit was observed. The WEE1 inhibitor adavosertib combined with gemcitabine has a preliminary score of 4 based on an OS benefit observed in a phase II study; this drug has not yet been licensed. Other treatments or combinations have shown only limited benefit compared to monotherapy with topotecan, PLD or paclitaxel.

### DISCUSSION

Analysis of the clinical benefit of systemic therapies for recurrent ovarian cancer using the ESMO-MCBS shows substantial benefit of only a few licensed treatments: carboplatin with either paclitaxel or PLD in the platinum-sensitive setting and the addition of bevacizumab to topotecan, paclitaxel or PLD in the platinum-resistant setting. Considering that over 18,000 patients were enrolled in randomised clinical trials in ovarian cancer over the last three decades, this result is disappointing. This is in line with...
Figure 2. Relative difference in MCBS between comparators in platinum-sensitive (A) and platinum-resistant (B) ovarian cancer.

An overview of the relative benefit between the different arms and comparators studied and graded for the platinum-sensitive and -resistant setting is provided. The distance between two comparators is representative for the MCBS grading with larger distance representing more benefit according to MCBS for the intervention versus comparator.

*In the MITO16b study, carboplatin doublets studied were carboplatin with either paclitaxel, gemcitabine or liposomal doxorubicin.

MCBS, Magnitude of Clinical Benefit Scale; OS, overall survival; PFS, progression-free survival; PLD, liposomal doxorubicin; PRO, patient-reported outcomes; QoL, quality of life.
the moderate improvement of 5-year survival rate in advanced ovarian cancer of only 2% from 1983 to 2012. Most of the studied treatments were tested in large RCTs and did not improve PFS, OS or QoL. Therefore, the lack of benefit is more likely due to the inefficacy of the drugs rather than due to badly designed trials. Treosulfan, melphalan, paclitaxel and etoposide received a broad label for treatment of ovarian cancer decades ago based on registration trials that included patients with newly diagnosed EOC. These drugs were not tested specifically in the recurrent setting in large trials; therefore, these were not included in the current analysis and not scored on the ESMO-MCBS.

Based on their mechanism of action, PARP inhibitors are expected to especially benefit patients with homologous recombination-deficient tumours, such as BRCA-mutated ovarian cancer. HRD can also be caused by other alterations in the homologous recombination repair pathway. Based on the recently published mature OS results, maintenance treatment with olaparib (tablets) did not yield a substantial clinical benefit (grade 2) for patients with a gBRCA mutation. Surprisingly, a sensitivity analysis in the population with a Myriad-confirmed gBRCA mutation, which included only nine additional patients, did result in a statistically significant OS benefit. However, because correction for multiple testing was not applied, this sensitivity analysis is considered exploratory and its results were not scored on the ESMO-MCBS. Grading of the other licensed PARP inhibitor therapies (niraparib and rucaparib) on the ESMO-MCBS labels their benefit as only ‘moderate’ with a score of 3. This score was not higher in the pre-specified subgroup of patients with a BRCA mutation compared to the unselected ITT population. The PFS gain in the ITT population for rucaparib, and in the non-gBRCA-mutated patients for niraparib was, however, numerically lower than in the other subgroups. It is, therefore, essential to understand the benefit of PARP inhibitors in patients with BRCA wild-type disease and absence of HRD, since the gain observed for the ITT population is likely to be largely driven by the inclusion of patients with either HRD tumours or BRCA-mutated tumours. This supports restricting use of olaparib tablets to this group of patients. Niraparib and rucaparib received a broad label, which includes treatment of patients irrespective of BRCA mutation status. The registration trial for olaparib capsules (STUDY19) also included BRCA-negative patients, but it received a restricted licensed indication only in patients with a BRCA mutation. Later, an updated OS analysis indeed only showed a numerical OS gain in patients with a BRCA mutation supporting this. This OS benefit did not meet the prespecified criterion for statistical significance; therefore, the preliminary ESMO-MCBS score of 2 was not upgraded. For olaparib tablets (SOLO2), the ITT population only included patients with a confirmed BRCA mutation. Remarkably, registration was partly based on data from the ITT analysis of STUDY19, leading to a broader licensed label for olaparib tablets, including treatment of patients without a BRCA mutation. Despite the use of debulking surgery for selected patients with recurrent disease, given the low chance of cure for these patients, we considered chemotherapy after debulking surgery for recurrent ovarian cancer a palliative treatment and therefore used ESMO-MCBS forms 2 or 3. For treatments that only show a PFS benefit, without an OS benefit, the ESMO-MCBS score is downgraded if toxicity is increased or QoL analysis does not show an improvement. Obviously, if mature OS data or QoL results are not published, as is yet still the case for most PARP inhibitor trials, this downgrade will not be applied. ESMO-MCBS forms do not currently allow downgrading when OS and QoL data are not published, but this could be relevant in cases in which sufficient time has passed for data to become mature and publication of OS data is no longer expected to follow. In ovarian cancer trials, a PFS benefit often does not predict a benefit in OS. However, mature OS data can be difficult to interpret because of crossover. Furthermore, a difference in sensitivity to reintroduction of some chemotherapies in responders versus non-responders to PARP inhibitors due to a difference in DNA damage repair capability further complicates ascribing an observed difference in OS to a single treatment regimen. An improvement of PFS accompanied by an improvement in QoL could be of clinical value for patients, even in the case that no OS benefit is established.

Downgrading for toxicity can be a matter of debate. For example, in the AURELIA trial, global QoL was comparable but there was an improvement in a subscale of QoL measuring GI symptoms. This subscale analysis was predefined and therefore the preliminary score of 3 was upgraded to a 4. The additional toxicity of bevacizumab, such as an extra 2% GI perforation (grade 2 or higher), arterial thromboembolic events and fistula/abscess formation should be taken into account in the benefit/risk evaluation of this treatment, although this does not meet the criteria for a downgrade of the ESMO-MCBS score.

There are some other points to consider in interpreting the current analysis. First of all, studies from 1990 onwards were included. Surgical treatment and supportive care have changed during this time period, complicating the comparison of study results within these three decades. Furthermore, the ESMO-MCBS was developed to grade the results of well-designed registration studies, and the standards for performing and reporting of clinical trials have become more stringent during the years. However, even in recent times, only a minority of studies testing anticancer drugs were adequately designed to be able to meet ESMO-MCBS thresholds of substantial clinical benefit. When head-to-head comparisons testing superiority for different treatments are not available, and the studies testing those treatments used different comparator arms, it is complicated to compare ESMO-MCBS scores of these treatments. An example is the use of carboplatin/paclitaxel (grade 4, compared to carboplatin) versus carboplatin/PLD (grade 3, compared to carboplatin/paclitaxel in a non-inferiority study) in the platinum-sensitive setting. Despite the lower ESMO-MCBS grade, one could argue that carboplatin/PLD is the better combination given the stronger comparator arm.
in this study. Therefore, when taking into account ESMO-MCBS grading for treatment decision making, it is crucial to critically assess differences in comparators used between studies. We attempted to give insight into the relative benefit of various head-to-head comparisons tested in the platinum-sensitive (Figure 2A) and platinum-resistant (Figure 2B) setting, but this should be interpreted with caution given the heterogeneity between studies. Finally, more detailed consideration of toxicity, impact on QoL and the optimal sequences of therapies on an individual patient level should be taken into account to make treatment decisions.

Conclusion

Only a few licensed systemic therapies for recurrent ovarian cancer show substantial clinical benefit based on ESMO-MCBS scores. Carboplatin + paclitaxel scores a grade 4 indicating substantial clinical benefit in the platinum-sensitive setting. Addition of bevacizumab to either PLD, paclitaxel or topotecan in platinum-resistant disease also has substantial benefit (grade 4), but the QoL data supporting this score are subject to interpretation. The unlicensed WEE1 inhibitor adavosertib combined with gemcitabine has a preliminary score of 4 based on an OS benefit observed in a phase II study. For PARP inhibitors, only moderate clinical benefit can be confirmed, based on OS results for olaparib tablets and PFS results for the other drugs.

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