

# OVARIAN CANCER TREATMENT IS INCREASINGLY INDIVIDUALISED

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Up to 70% of patients with advanced Stage III–IV high-grade OC will **relapse**,<sup>1</sup> and >80% of patients with late-stage OC will become **platinum resistant**.<sup>2</sup>

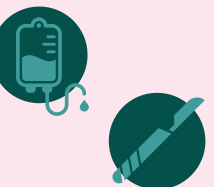
## IN THE PAST



### Patient

A deeper understanding of OC tumour biology has gradually led to subclassifications and the identification of some molecular markers.<sup>2</sup>

In the past, patient characterisation and treatment decisions were based on disease staging, treatment response, and adverse event profiles.<sup>3</sup>



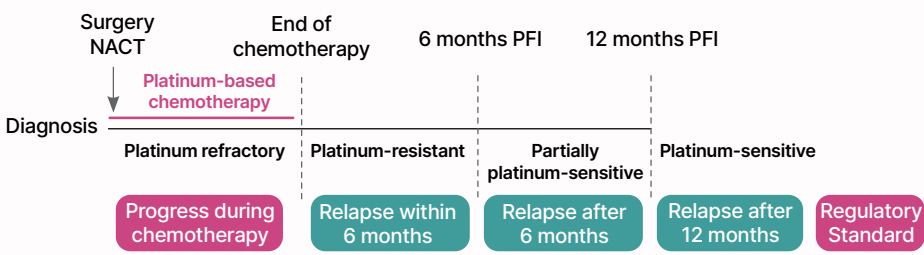
### Treatment

Prior to the development of targeted therapies, treatment options in OC were restricted to surgical intervention and PbC.<sup>4</sup>

A 'regulatory standard' was developed to define PbC eligibility (**Figure 1**), and PROC was defined as a disease that relapses within 6 months of PbC.<sup>5</sup>

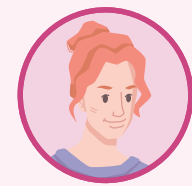
- However, this definition may not be sufficient to predict potential benefit from subsequent PbC or non-platinum-based therapy.<sup>6</sup>

Figure 1<sup>5</sup>



Adverse events should be reported. Reporting forms and information can be found at <https://yellowcard.mhra.gov.uk> or via the MHRA Yellow Card app, available in the Google Play or Apple App Stores. Adverse events should also be reported to AbbVie on [GBPv@abbvie.com](mailto:GBPv@abbvie.com).

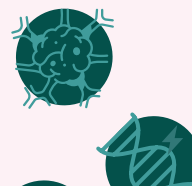
## IN THE PRESENT



### Patient

Due to the identification and validation of **clinically actionable biomarkers**, **individualised treatment** paths for patients are now possible.<sup>7</sup>

Currently, patient characterisation and treatment decisions should also consider tumour characteristics, survival outcomes, tumour burden, adverse event management, quality of life, treatment-free interval, histology, and mutation/ biomarker status.<sup>1,7</sup>

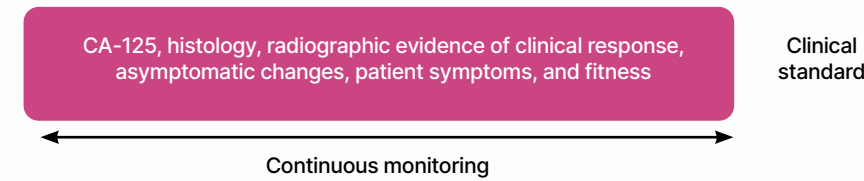


### Treatment

**Therapeutic advancements** in OC, such as bevacizumab or PARP inhibitors, provide more opportunities to treat when PbC is not the best option.<sup>1</sup>

- Despite all of these advancements, treating patients with PROC remains a clinical challenge.<sup>8</sup>
- Today, a '**clinical standard**' has emerged whereby suitability for PbC is determined through the assessment of many patient factors (**Figure 2**).<sup>1</sup>
- For some patients with recurrent OC, **PbC rechallenge may not be clinically appropriate**.<sup>1</sup>
- The concept of a '**platinum-ineligible**' status allows for more **individualisation** of therapy.<sup>1</sup>

Figure 2<sup>1</sup>



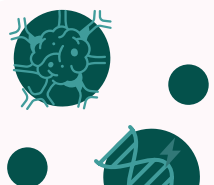
## IN THE FUTURE



### Patient

An **increasing understanding** of tumour biology in OC is informing novel therapeutic development.<sup>4</sup>

In the future, patient characterisation and treatment decisions may also consider tumour cell **subtype burden**, cumulative **adverse event burden**, patient **choice**, **sequencing**, and **evolving biomarker/ mutation** status. (Moore, personal communication)



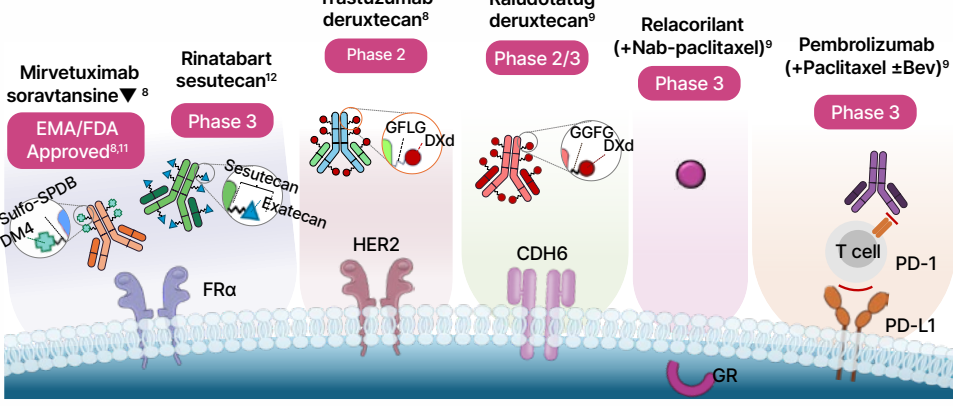
### Treatment

Advancements in **targeted therapies** and **health technologies** may usher in a future of **personalised** care.<sup>7-9</sup>

With perspectives on platinum sensitivity changing, novel approaches to treatment are needed to fill the gap.<sup>2,4</sup>

Several **antibody–drug conjugates** and **targeted therapies** are in late-stage development, which may expand therapeutic options in PROC (**Figure 3**).<sup>7,10</sup>

Figure 3



## CONCLUSION

- Evolving understanding of patient and tumour biology is leading to a **changing tide in PROC diagnosis and treatment**.<sup>1,7</sup>
- The **emerging 'clinical standard'** determines suitability for PbC through the assessment of many patient factors.<sup>1</sup>
- The future of **tailored therapy** will **harmonise** considerations for **tumour biology** and what is right for the **patient**.<sup>2,8,9</sup>



Prescribing information and indication for UK HCPs can be found [here](#). December 2025 UK-ELAH-250156

**Abbreviations:**  
Bev: bevacizumab; CDH6, cadherin 6; FRα: folate receptor alpha; GR: glucocorticoid receptor; HER2: human epidermal growth factor receptor 2; Nab: nanoparticle albumin-bound; NACT: neoadjuvant chemotherapy; OC: ovarian cancer; PARP: poly-ADP ribose polymerase; PbC: platinum-based chemotherapy; PD-1: programmed cell death protein 1; PD-L1: programmed death-ligand 1; PFI: platinum-free interval; PROC: platinum-resistant cancer; SPDB: N-Succinimidyl 4-(2-pyridyldithio)butyrate.

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Clinicaltrials.gov. NCT06619236. <https://clinicaltrials.gov/study/NCT06619236>.