Out of licence cancer drug or regimen

Funding request form



This form is to be used to request Bupa funding for your patient to receive cancer drugs that are not licensed in the UK for the condition being treated.

For us to accurately assess your request, you need to:

- 1. Complete this form in **FULL** including all relevant elements of the patient's current medical condition and medical history
- 2. Attach MDT notes (section 5)
- 3. Attach accompanying evidence (section 6)

at least three working days before the treatment is due to take place.

We're unable to assess funding based on incomplete forms or evidence. If we need to ask you for more information, this is likely to delay our funding decision and risks delaying your patient's treatment.

1. Patient's details Miss Mrs Ms Mr Dr Other (please state) Title (please tick) Patient's name Date of birth Bupa membership number 2. Clinician's details Name Bupa provider number Hospital name Phone number **Email address**

3. Patient's medical information Primary diagnosis Stage of disease Are there any metastases? Yes, please specify site below No Are you treating? Primary tumour Metastases Side effects, please give details below Current performance status (ECOG) 4. Drug(s) requested (including supporting drugs) Drug 2 Drug 1 Drug 3 Drug 4 Drug name Cycle length Number of cycles or until progression/ toxicity? Number of treatments per cycle Route of administration Dosage Setting Relevant gene expression/ hormone status Proposed start date Is this part of a compassionate

use scheme?

| Bupa will only fund experimental drug treatment following use of an in treatment is proposed? | itial licenced treatment where | available. What | line of |
|---|---|---|----------------|
| | | | |
| What is this treatment's intent? | -adjuvant Adjuvant | | |
| Outline previous therapies tried and the response | | | |
| | | | |
| Explain what standard treatments are available to other patients (included why these are not appropriate for this patient. | ding on the NHS) with this co | ndition or stage | of disease and |
| | | | |
| Will the patient be receiving any other SACT / radiotherapy alongside | any of the drugs above? | | |
| | | | |
| | | | |
| 5. Multidisciplinary Team (MDT) | | | |
| Has this regime been agreed by an MDT? | Yes. Please provide notes and supportive evidence as described in section 6. | No. We're unable to treatment for y without an MD | our patient |
| Name of hospital where MDT was carried out | | | |
| MDT speciality area | Date of MDT | D M M | YYY |
| If an MDT is not suitable to be carried out, please explain why | | | |
| | | | |
| 6. Evidence base for the use of this treat | ment at the give | n indicat | ion* |
| Select 1 option at minimum | | | |
| Does the proposed treatment have an EMA licence for the given indication?* | Yes, please attach a co evidence along with th | | No |
| Does the proposed treatment have a positive outcome from a Health Technology Assessment for the given indication*? | Yes, please attach a co | | ☐ No |
| Is there evidence of benefit through a predictive genetic test for the given indication*? | Yes, please attach a co | | ☐ No |
| Is there an existing NHS guideline recommending the proposed treatment for the given indication*? | Yes, please attach a co | | ☐ No |
| Is there an existing ESMO or NCCN guideline recommending the proposed treatment for the given indication* that is supported by a published phase III study? | Yes, please attach a co | | ☐ No |
| Are there any published Phase III clinical trial results (not interim findings) confirming clinical efficacy? | Yes, please attach a co | | No |

 $^{^{\}ast}\mbox{including tumour type, stage, phase of treatment and pharmacogenetic markers.}$

| 7. Trials and compassionate use | | |
|---|--|-----------|
| Is there a registered clinical trial investigating the proposed treatment for which the patient would meet the inclusion criteria? | Yes, please give details below | ☐ No |
| | | |
| 8. Consultant's declaration | | |
| I understand that the clinical information I've supplied may be conside that my patient (or their legal representative) has given their permission review this information, they've been given an opportunity to do so be | on for me to share this information and, w | |
| I confirm that the information that I have supplied is the full clinical pic will submit all relevant evidence correctly in accordance with Good Me | | |
| Consultant's name | Date D M | 1 Y Y Y Y |
| General Medical Council number | | |
| | | |

9. Further information

Our customers' health insurance schemes may cover the cost of some cancer drugs that aren't licensed or drugs that we don't routinely fund in the UK. When assessing funding requests, we look at the strength and quality of the evidence of clinical effectiveness and the anticipated measurable outcomes. These outcomes may include improvements in overall survival, progression-free survival, clinical response, and adverse effects.

Please return this form to us by secure email to OncologyTeam@bupa.com

Please be aware that information you send to this email address may not be secure unless you send us your email through Egress Switch. For more information and to sign up for a free Egress Switch account, go to https://switch.egress.com/ui/learn. You won't be charged for sending secure emails to a Bupa email address using the Switch service.