

**Subject: Specialised Services Circular (SSC)**

**Sent on behalf of: Chair of the SW Specialised Service Circular Group**

Dear Colleagues,

Please find attached the following Specialised Services Circular(s):

<b>SSC Number</b>	<b>SSC Title</b>	<b>Trusts approved to prescribe in accordance with the SSC, providing appropriate internal governance arrangements are in place</b>
2733	<b>NICE Technology Appraisal Final Draft Guidance: Crizotinib for treating ROS1-positive advanced non-small-cell lung cancer</b>	All South West Acute Trusts

**Is an implementation plan required from all SW trusts (regardless of commissioned status) for this SSC? No**

**For all other South West region trusts this is for information only.**

Trusts should ensure that use is registered on the Blueteq system (if appropriate).

Treatment will only be funded where the drugs minimum dataset is fully and accurately populated.

Please direct any queries to: [england.speccomm-southwest@nhs.net](mailto:england.speccomm-southwest@nhs.net)

**All Chief Executives**

**All Medical Directors**

**All Chief Pharmacists**

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12 November 2024

Dear Colleagues,

**Re: NICE Technology Appraisal Final Draft Guidance: Crizotinib for treating ROS1-positive advanced non-small-cell lung cancer**

I am writing to advise you regarding the funding position on the recently published NICE Technology Appraisal Final Draft Guidance (FDG) for crizotinib for treating ROS1-positive advanced non-small-cell lung cancer.

The FDG can be found at: <https://www.nice.org.uk/guidance/indevelopment/gid-ta11450>.

NICE in their FDG published on 31<sup>st</sup> October 2024 has stated that:

Crizotinib is recommended as an option for treating ROS1-positive advanced non-small-cell lung cancer in adults, only if they have not had ROS1 inhibitors

Crizotinib has been available in this indication via the Cancer Drugs Fund (CDF) since 31<sup>st</sup> May 2018 (TA529). NICE has reviewed the evidence collected as part of the CDF managed access agreement and has recommended crizotinib for use in routine commissioning. Crizotinib will receive interim funding via the Cancer Drugs Fund (CDF) from 31<sup>st</sup> October 2024 in line with these recommendations and according to a set of treatment criteria which translates the NICE recommendation into a clinical guide as to use in practice. These treatment criteria can be found on the national CDF list at <https://www.england.nhs.uk/cancer/cdf/cancer-drugs-fund-list/> or on the application form(s) on the Blueteq site.

NHS England will then routinely commission crizotinib in patients with ROS1-positive non-small-cell lung cancer, incorporating these treatment criteria, including those contained within this letter from 30 days after the day of publication of the final guidance.

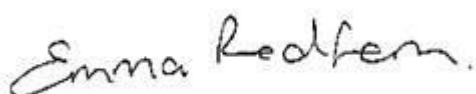
In addition:

- Trusts must ensure that they are purchasing crizotinib at the agreed proposed patient access scheme (PAS) discounted price. This discounted price will be applied automatically at point of invoice and applies to all indications. Trusts should refer to the CAP portal for further information on the PAS price. The CAP portal is available at <https://nhsengland.sharefile.eu/Authentication/Login>

- Trusts must ensure that, until 30 days after publication of the final guidance from NICE, only invoices for the drug procurement costs of crizotinib in this indication are directed to the CDF and that they are also submitting complete and accurate information via the CDF minimum dataset (MDS).
- In line with the terms and conditions included in the NHS Standard Contract and as per the agreement that Cancer Services are commissioned with Trusts, Schedule 6a Reporting Requirements for drugs will apply. Payment of Trust invoices will be contingent on the completion of the MDS record and this information being made available in a timely way.
- Trusts must ensure they are registering crizotinib use on SACT. The SACT dataset is a mandated dataset as part of the Health and Social Care Information Standards. This is listed as a Schedule 6 national information requirement within the NHS Standard Contract.
- Patients must be registered via Blueteq (CRI3) and meet the clinical criteria on the registration form during the interim funding period. Trusts do not need to submit a new form for patients who have started treatment since 31<sup>st</sup> May 2018.
- **Payment of Trust invoices will be contingent on Blueteq registration, the full SACT and CDF MDS record applicable to the drug being completed and this information being made available in timely way.**
- Trusts must ensure that local governance aspects (e.g. technical issues, education & training, patient information) have been identified and addressed for all staff groups (as appropriate) in order to permit the safe delivery of this therapy.

I would be grateful if you could cascade this information to relevant clinical teams within your organisation to support the consistent adoption of the policy nationally.

With best wishes,



Emma Redfern  
Medical Director NHS England South West  
Region



Tracey Williams  
Principal Pharmacist