National Cancer Medicines Advisory Group (NCMAG) Programme

NCMAG Proposal Form Guidance

Background

The purpose of the NCMAG Programme is to provide advice on the safe implementation of new uses of cancer medicines where a “Once for Scotland” approach would be of benefit to improve consistency, equity of access, avoid duplication and support recovery from the pandemic. This would include medicines in the following categories which are out with the remit of Scottish Medicines Consortium (SMC):

- Off-label uses of licensed cancer medicines, branded, generic or biosimilar and
- On-label uses of licensed generic medicines, known as off-patent use

The programme is intended to work alongside SMC and existing local systems to support access to cancer medicines. The work programme is driven by clinical need in consultation with Scottish Government Health Department (SGHD) Pharmacy and Medicines Division and Cancer Policy Teams, National Cancer Recovery Group, and stakeholders across NHSScotland.

Decisions on treatment need to be made on an individual patient basis by clinicians in discussion with the patient and, where appropriate, the multidisciplinary team. NCMAG Council will make decisions on proposals which will apply to groups of patients, notwithstanding that individual documented patient assessment and discussions will still be required.

Proposals to the group

The Programme will appraise and issue advice on the off-label and off-patent use of cancer medicines, including:

- Off-label uses of licensed cancer medicines (branded, generic or biosimilar):
  - for an illness or patient population not specified within the marketing authorisation
  - for administration by a different route, dose or frequency
- On-label uses of licensed generic medicines, known as off-patent use. This category is anticipated to include medicines which are not recommended by SMC, however the patent has expired since SMC advice was published, with the medicine now available at lower cost and current cost-effectiveness is unknown.

The programme will not consider:

- medicines without any marketing authorisation (unlicensed medicines) in the UK
- situations where a marketing authorisation is likely to be sought for the proposed medicine in the off-label use within 24 months

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Visit NCMAG Programme Webpage for more information
• in situations where a regulatory decision on marketing authorisation is pending for a comparator product, in the same off-label use as a proposal received by NCMAG, the suitability of the proposal for NCMAG review will be considered on a case-by-case basis
• established off-label uses which have already become standard of care nationally
• paediatric indications
• treatments that do not impact on disease behaviour, for example analgesics for cancer pain
• medicines and uses within SMC remit.

Process

Who can make a proposal?

Individual consultants wishing to implement a change should initially seek support from specialist colleagues within their team/Managed Clinical Network and from specialist colleagues across Scotland. Submissions to NCMAG for changes to practice should be led by a tumour site specialist working in collaboration with specialist consultant colleagues across NHSScotland. Pharmacy input and support should be sought to work up proposal submissions. The nominated lead will submit the proposal for consideration.

What does the proposal entail?

It requires the completion of a proposal form with key information which will support the NCMAG Council to appraise the evidence and decide whether to support the routine use of the proposed treatment.

It is important that prior to submission you check your proposal against the following criteria:

• The medicine meets off-label or off-patent criteria defined above.
• The proposal is supported by at least one published research article.
• There is evidence of national consensus with support from all three cancer networks. The submitting clinician is required to provide contact details of the supporting applicants included in the proposal. This will allow the supporting applicants to be sent an acknowledgement email on receipt of the proposal form.
• Clearly defined eligible patient population for whom the proposed use is intended.

How should a proposal form and supporting documents be submitted?

• The nominated lead will submit the proposal via email to the NCMAG mailbox: his.ncmag@nhs.scot.
• In addition to this document, further advice on the process/proposal prior to submission can be sought via the NCMAG email address.
• The NCMAG team may be in contact with the nominated lead to request further information in advance of the screening and prioritisation step or in advance of consideration at the NCMAG Council.
• Council meetings to review proposals will be held at least every 3 months and, depending on demand, possibly more frequently.
• Proposers are required to submit proposals by set submission deadlines which can be found here.
• The aim is for proposals to be scheduled for review within 4 months of receipt of submission.
Proposers will be notified via email confirming the date of the NCMAG Council meeting at which their proposal will be considered.

What happens to submitted proposals?

Screening and prioritisation

- Once a proposal is received it will be reviewed against pre-determined criteria by the NCMAG Programme Team to assess its suitability for consideration by the NCMAG Council.
- The proposer will be notified via email confirming if the proposal has or has not been considered appropriate for review.
- In the event that the proposal is likely to have a significant service implication, we may require additional input from service managers to explore this further.

Depending on the volume of workload, there may be a need to prioritise proposals received. Prioritisation will follow pre-determined criteria and be conducted by the NCMAG Executive.

Prior to the meeting

- For proposals considered appropriate for consideration by the Council, the NCMAG team will confirm the meeting date to the proposer.
- Proposals suitable for consideration by the NCMAG Council will go through an internal evidence review process, including a systematic literature review, appraisal and quality assessment of the evidence relevant to the proposal.
- The proposer may be asked to provide additional information prior to the meeting to support the review.
- Ahead of the meeting the proposer will be issued with a presentation template to support contribution at the council meeting. If the proposer is unable to attend, we would request the proposer nominates a deputy who would be able to present on their behalf.
- The NCMAG team will liaise with National Procurement to consider matters related to procurement and supply of the proposed medicine, where relevant.

At the meeting

- The NCMAG Council includes a wide range of stakeholder representatives, who will consider proposals at the meeting.
- The proposer or a deputy is invited to join the NCMAG council meeting to provide a brief presentation on the proposal, using the template provided in advance.
- The NCMAG team will present the results of the internal clinical and economic evidence review.
- The NCMAG Council members are invited to ask any questions of the proposer and NCMAG team.
• The proposer is asked to temporarily leave the meeting to allow the NCMAG Council members to consider the proposal and make a decision on whether the proposal will be supported or not supported by NCMAG.

• Decisions are based on the best available information provided on the day of the meeting.

• Once the NCMAG Council members have reached a decision, the proposer will be invited to rejoin the meeting for the confidential sharing of the decision outcome and rationale for the decision.

After the meeting

• The NCMAG team will complete the draft advice document, taking into consideration the discussion and decision made by the Council.

• The draft advice document is shared with the proposer in confidence for comment on accuracy only.

If the proposal is supported the proposer will be requested to work with the supporting pharmacist to complete the National Systemic Anticancer Therapy (SACT) protocol to share nationally along with the advice.

The final advice document, along with the SACT protocol, will be issued to Health boards and Regional Cancer Networks and added to the NCMAG Programme webpage.

Boards will still need to consider local governance issues and the service/budget impact of changes. All cancer networks and Boards are expected to facilitate access to medicines supported by the group.

How long will the advice be valid?

All advice will be reviewed every three years, or when new information or evidence becomes available. Reviews may result in advice being extended, changed or withdrawn.