**National respiratory audit programme (NRAP)**

**NRAP: National data opt-out (NDO) statement**

Version 6: March 2024

All patients have the right to decide whether information about them and their healthcare is used for research and planning. The national data opt-out allows information held on the NHS Spine to be checked against an individual’s NHS number indicating whether they are happy to for their data to be used. This screen ensures that data of a patient who has opted-out is removed from data collections before submission to non-exempt national clinical audits, registries, confidential enquiries etc.

**What happens if a patient decides to opt-out of NRAP audits?**

* It is the responsibility of each hospital or NHS Trust in England to ensure that information is not entered into the web tool for patients that have a national data opt-out in place.
* Patient details should be screened before entering data into the web tool, to see whether they have chosen to opt out of confidential patient information being used for research and planning purposes.
* If a patient decides during their treatment that they would like to opt-out, their data should not be entered into the NRAP web tool, and they should be provided with the necessary National data opt-out information to formally log their opt-out preference (see below for this information).
* If a patient wishes to opt out but their treatment has already taken place, and the record for that episode of care has already been entered onto the NRAP web tool, we recommend that they contact the hospital that treated them or the audit team to discuss this.

*Please note that national data opt-out preferences do not apply retrospectively (to data already submitted and used for research and planning purposes).*

* If a patient thinks that their information is being used inappropriately, they have the right to complain to the [**Information Commissioners Office**](https://ico.org.uk/) (ICO).

**What information should I be providing to patients who wish to opt-out of their data being used for planning and research purposes?**

More information on the National data opt-out is available at [**https://digital.nhs.uk/services/national-data-opt-out**](https://digital.nhs.uk/services/national-data-opt-out) or [**www.nhs.uk/your-nhs-data-matters**](http://www.nhs.uk/your-nhs-data-matters).

These web pages will provide the patient with:

* information on what is meant by confidential patient information
* finding examples of when confidential patient information is used for individual care and examples of when it is used for purposes beyond individual care
* finding out more about the benefits of sharing data
* understand more about who uses the data
* finding out how their data is protected
* being able to access the system to view, set or change their opt-out setting
* finding the contact telephone number if you want to know any more or to set/change your opt-out by phone
* knowledge of the situations where the opt-out does not apply

Patients can also find out more about how patient information is used at:

[**https://www.hra.nhs.uk/information-about-patients/**](https://www.hra.nhs.uk/information-about-patients/) (which covers health and care research); and

[**https://understandingpatientdata.org.uk/what-you-need-know**](https://understandingpatientdata.org.uk/what-you-need-know) (which covers how and why patient information is used, the safeguards and how decisions are made)

More information about how NRAP uses patient data, including ‘how to say no’ to being included, can be found at the following audit workstream links:

COPD: [**www.rcp.ac.uk/projects/outputs/support-service-teams-copd**](http://www.rcp.ac.uk/projects/outputs/support-service-teams-copd)

Adult Asthma: [**www.rcp.ac.uk/projects/outputs/support-service-teams-adult-asthma**](http://www.rcp.ac.uk/projects/outputs/support-service-teams-adult-asthma)

Children and young people’s asthma: [**www.rcp.ac.uk/projects/outputs/support-service-teams-children-and-young-people-s-asthma**](http://www.rcp.ac.uk/projects/outputs/support-service-teams-children-and-young-people-s-asthma)

Pulmonary rehabilitation: **www.rcp.ac.uk/projects/outputs/support-service-teams-pulmonary-rehabilitation**

Patients can change their mind about their choice at any time. For patients wishing to opt back in, ie to have their data included in the audit, it can take up to 21 days for this change to take effect (fair processing period).

To help support services ensure that they are compliant with applying the national data opt-out, we also have available a set of frequently asked questions (NDO FAQs).



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