

Care Information Choices, England Experimental Statistics, April 2016

A patient may opt out, or withdraw the decision to opt out of information that identifies them being shared outside of the HSCIC for purposes beyond direct care.

This publication reports on the number of patient opt-outs as a percentage of patients registered at Clinical Commissioning Groups (CCG) in England where patient opt-out information has been received from GP practices, as of April 2016.

Percentage of patient opt-outs received by the HSCIC (of over 56 million patients in England):

2.2%
Opted out

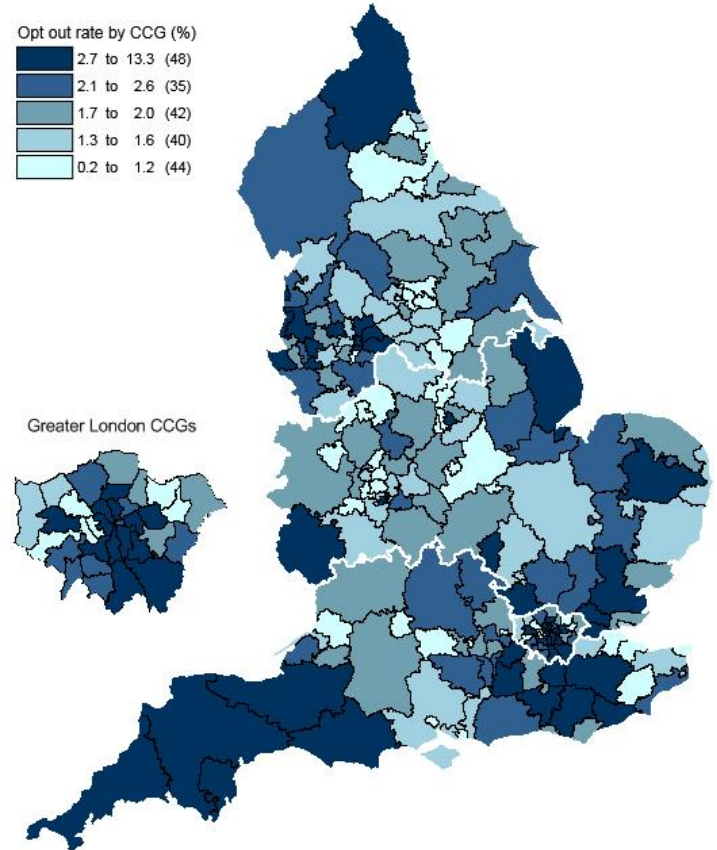
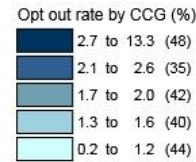
Around 1 in 45 patients

Opting out of information that identifies you being shared outside of the HSCIC for purposes beyond direct care.

Opt-out levels vary by CCG and over time. However 95.7 per cent of CCGs have between a 0 per cent and 5 per cent opt-out rate as of April 2016.

www.hscic.gov.uk/pubs/careinfochoicesapr16
www.hscic.gov.uk/yourinfo

Percentage of opt-outs mapped by CCG



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