Partner Organisations:

Health Research Authority, England NHS Research Scotland

NIHR Clinical Research Network, England NISCHR Permissions Co-ordinating Unit, Wales

HSC Research & Development, Public Health Agency, Northern Ireland

Notification of Non-Substantial/Minor Amendments(s) for NHS Studies

This template **must only** be used to notify NHS/HSC R&D office(s) of amendments, which are **NOT** categorised as Substantial Amendments.

If you need to notify a Substantial Amendment to your study then you MUST use the appropriate Substantial Amendment form in IRAS.

Instructions for using this template

- For guidance on amendments refer to http://www.hra.nhs.uk/research-community/during-your-research-project/amendments/
- This template should be completed by the CI and optionally authorised by Sponsor, if required by sponsor guidelines.
- This form should be submitted according to the instructions provided for NHS/HSC R&D at http://www.hra.nhs.uk/research-community/during-your-research-project/amendments/which-review-bodies-need-to-approve-or-be-notified-of-which-types-of-amendments/. If you do not submit your notification in accordance with these instructions then processing of your submission may be significantly delayed.

1. Study Information

Full title of study:	Improving perioperative care through the use of quality data: Patient Study of the Perioperative Quality							
	Improvement Programme							
IRAS Project ID:	215928							
Sponsor Amendment Notification number:								
Sponsor Amendment Notification date:	11/08/2017							
Details of Chief Investigator:								
Name [first name and surname]	Dr Suneetha Ramani Moonesinghe							
Address:	Anaesthetics Department, Podium 3, Maple Link corridor, University College Hospital 235 Euston Road							
Postcode:	NW1 2BU							
Contact telephone number:	07956620717							
Email address:	ramani.moonesinghe@nhs.net							
Details of Lead Sponsor:								
Name:	Suzanne Emerton							
Contact email address:	randd@uclh.nhs.uk							
Details of Lead Nation:								
Name of lead nation delete as appropriate	England							
If England led is the study going through CSP? delete as appropriate	Yes							
Name of lead R&D office:	Joint Research Office, UCL, London, WC1 E6BT							

Partner Organisations:

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NHS Research Scotland
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2. Summary of amendment(s)

This template must only be used to notify NHS/HSC R&D office(s) of amendments, which are NOT categorised as Substantial Amendments. If you need to notify a Substantial Amendment to your study then you MUST use the appropriate Substantial Amendment form in IRAS.

	ced supporting For office use only	With this form)		6.0v	16.06.2017	_																
List relevant supporting	numbers (please ensure all referenced supporting	Documents are submitted with this form. Document		Patient Information	Sheet		Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form	Patient Consent Form
idment applies to		Sites	All sites or list	affected sites	All sites or list	affected sites		All sites or list	All sites or list affected sites	All sites or list affected sites All sites or list	All sites or list affected sites All sites or list affected sites	All sites or list affected sites All sites or list affected sites Listed sites	All sites or list affected sites All sites or list affected sites Listed sites	All sites or list affected sites All sites or list affected sites Listed sites	All sites or list affected sites All sites or list affected sites Listed sites	All sites or list affected sites All sites or list affected sites Listed sites	All sites or list affected sites All sites or list affected sites Listed sites	All sites or list affected sites All sites or list affected sites Listed sites	All sites or list affected sites All sites or list affected sites Listed sites	All sites or list affected sites All sites or list affected sites Listed sites	All sites or list affected sites All sites or list affected sites Listed sites	All sites or list affected sites All sites or list affected sites Listed sites
Amendment applies to (delete/ list as appropriate)		Nation			Northern	Ireland	Landle C	Scotland	SCOTIAND	Scotland	Wales	Wales Scotland	Wales Scotland	Wales	Wales	Wales	Wales	Wales	Wales	Wales Scotland	Wales	Wales Scotland
Brief description of amendment (please enter each separate amendment in a new row)			Minor amendment to Patient Information Sheet and	patient consent form to further clarify the data	linkage which will be taking place.								Addition of new sites: University Hospitals Of North Midlands NHS Trust	Addition of new sites: University Hospitals Of North Midlands NHS Trust	Addition of new sites: University Hospitals Of North Midlands NHS Trust	Addition of new sites: University Hospitals Of North Midlands NHS Trust	Addition of new sites: University Hospitals Of North Midlands NHS Trust	Addition of new sites: University Hospitals Of North Midlands NHS Trust	Addition of new sites: University Hospitals Of North Midlands NHS Trust	Addition of new sites: University Hospitals Of North Midlands NHS Trust	Addition of new sites: University Hospitals Of North Midlands NHS Trust	Addition of new sites: University Hospitals Of North Midlands NHS Trust
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Partner Organisations:

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NIHR Clinical Research NHS Research Scotland

NISCHR Permission

HSC Research & Development, Public Health Agency, Northern Ireland

NIHR Clinical Research Network, England NISCHR Permissions Co-ordinating Unit, Wales

3. Declaration(s)

Declaration by Chief Investi	(D\	niet inves	tig	gai	lOI
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- I confirm that the information in this form is accurate to the best of my knowledge and I take full responsibility for it.
- I consider that it would be reasonable for the proposed amendment(s) to be implemented.

Signature of Chief Investigator:

Print name:

SR MOONESINGHE

Date:

11.08.2017

Optional Declaration by the Sponsor's Representative (as per Sponsor Guidelines)

The sponsor of an approved study is responsible for all amendments made during its conduct.

The person authorising the declaration should be authorised to do so. There is no requirement for a particular level of seniority; the sponsor's rules on delegated authority should be adhered to.

• I confirm the sponsor's support for the amendment(s) in this notification.

Signature of sponsor's representative

Print name DAN . WILSOL

Post: ULL SPOASOR REPRESENTATIVE

Organisation: U 4

Date 15/08 /2017

