Primary hip replacement prostheses and their evidence base: systematic review of literature

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Abstract

Objective To determine the extent to which prostheses with no readily available evidence to support their use are being implanted in primary total hip arthroplasty.

Design Systematic review of the literature.

Data sources The 9th annual report of the National Joint Registry of England and Wales (NJR) was analysed to identify prostheses with an Orthopaedic Data Evaluation Panel rating of “unclassified” or “pre-entry” used in primary total hip arthroplasty in 2011. A systematic review of those prostheses was carried out using PubMed, Cochrane, Embase, OVID, and Google databases.

Study selection Prostheses used in primary total hip arthroplasty as published in the NJR’s 9th annual report were analysed. Only literature that included the name of the prosthesis was included. Literature yielded in the search results was excluded if it reported animal, non-orthopaedic, non-total hip arthroplasty, or non-device related studies.

Results The systematic review found that 24% (57/235) of all hip replacement implants available to surgeons in the UK have no evidence for their clinical effectiveness. It also shows that 10 617 (7.8%) of the 136 593 components used in primary hip replacements in 2011 were implanted without readily identifiable evidence of clinical effectiveness. These comprised 157 cemented stems (0.5% of 34 655 implanted), 936 (2.8% of 33 367) uncemented stems, 1732 (7.1% of 24 349) cemented cups, and 7577 (17.1% of 44 222) uncemented cups.

Conclusions This study shows that a considerable proportion of prostheses available to orthopaedic surgeons have no readily available evidence of clinical effectiveness to support their use. Concern exists about the current system of device regulation, and the need for a revised process for introducing new orthopaedic devices is highlighted.

Introduction

Medical device regulation has been the subject of recent debate.1-4 Both professional and public confidence in the system is at a low point. This is particularly true in orthopaedics, where the premature failure of some metal-on-metal hip replacements has added considerably to the global burden of hip revision. As a result, the British Orthopaedic Association has been at the forefront of demanding that a good evidence base accompanies each orthopaedic treatment. In addition, the association’s practice strategy document “Restoring your Mobility” has joined others in calling for radical change in the way medical devices are regulated in the European Union.5-6

In the United States, the regulation of medical devices falls under a single agency, the Food and Drug Agency, in much the same way as drugs are regulated by the European Medicines Agency in the EU. However, medical devices in Europe are evaluated for safety and reliability through any one of 76 notified bodies, which then issue a Conformité Européenne mark, allowing that device to be marketed in Europe.7 All hip replacement prostheses are considered class III devices, which means that the application for approval must include some human clinical investigations,8 although this need not be new research specific to the device if the manufacturer is claiming similarity to an existing product.

Although no clear list of which class III devices have been cleared for use in the United Kingdom or Europe is available,
in the case of hip replacement prostheses agencies exist that allow some comparison of available devices. One of these is the National Joint Registry (NJR). This was established in 2001 by the Department of Health and the Welsh Assembly Government to collect information on all hip and knee replacement operations in England and Wales and to monitor the performance of the prostheses used. In April 2011 reporting became mandatory for all National Health Service hospitals, and this now captures data from most of such operations, with compliance rates per hospital varying between 93% and 100% in the NJR’s latest report. A second agency is the Orthopaedic Data Evaluation Panel (ODEP), set up as part of the NHS to assess follow-up data for different primary hip replacement prostheses. Manufacturers are requested, but not required, to submit data on their product by using a pro-forma. Prostheses are then classified by the number of years post-implantation that the evidence spans (3, 5, 7, or 10) and by the quality of the data supplied (level A being strong evidence, level B reasonable evidence, and level C weak evidence); 10A is the highest rating available. To achieve this 10A benchmark, the prosthesis must have a failure rate of 10% or less at 10 years, a cohort study of more than 500 joints at its start, Kaplan-Meier survivorship data at 10 years, and registry data supporting its use. Products with less than three years of evidence are classified as “pre-entry” as long as the manufacturers keep the ODEP informed of the post-marketing surveillance data. Inclusion in this pre-entry category does not require evidence from peer reviewed literature at either the pre-clinical or clinical stages of development. Therefore, any data, regardless of whether it has been peer reviewed or not, can be submitted. Prostheses termed “unclassified” have had no evidence submitted by the manufacturers. How many of these “unclassified” and “pre-entry” implants actually have peer reviewed evidence to support their use is unclear, but many are widely available for implantation by any orthopaedic surgeon.

This study aimed to establish the number of primary hip replacement prostheses that have no readily available evidence of clinical effectiveness to support their use and how many are being implanted in clinical practice.

Methods

We analysed the NJR’s 9th annual report (2012) to group primary hip replacement prostheses according to their ODEP rating (fig 1⇓). For prostheses with an ODEP rating of pre-entry or unclassified, we did a systematic review of the literature by using PubMed, Cochrane, Embase, OVID, and Google databases to look for peer reviewed papers of any evidence level relating to the prosthesis in question. We did not search for custom, revision, or discontinued prostheses.

The search terms and protocol used were “prosthesis name” AND “hip”. The “prosthesis name” was that given in the NJR’s report as the “brand name” and compared with the manufacturer official website. If a discrepancy in the name was found, we used both names individually for the literature search. We did an additional search for each of the prostheses excluding generalised words from the brand name given in the NJR’s report, such as “cementless” or “stem,” to avoid missing potentially relevant articles.

Two researchers (FK-P and TTM) did the literature research and independently reviewed all results to establish the highest level of evidence for each of the prostheses; a third researcher (AMA) resolved any discrepancies. Titles and abstracts were reviewed, and those that were potentially relevant to the device in question were included. We defined evidence as peer reviewed publications in which the clinical effectiveness of a particular device was assessed. We excluded animal, non-orthopaedic, non-primary hip arthroplasty, and non-device specific studies (fig 1⇓).

We then gave the selected papers an evidence level rating according to the simplified evidence level table from the Centre for Evidence-Based Medicine, Oxford (box 1), and we established the highest level evidence available for each device. If no suitable evidence could be identified, we contacted the manufacturers and asked them to provide some data on their prosthesis. Those that responded with papers were rated; if some data were provided (for example, details of tensile strength or principles to support the device’s use), this would earn a level 5 evidence rating (expert opinion without explicit critical appraisal/pre-clinical biomechanical data).

We then analysed the collected evidence to determine the number of brands implanted with no evidence of clinical effectiveness and, additionally, the number implanted with no evidence at any level. Those implants with only level 5 data were excluded from the analysis, as we believed this level to be inadequate for clinical decision making. We then cross referenced this information with the published NJR9 prostheses tables to find the numbers of prostheses actually implanted into patients in each of these two categories.

Statistical methods

Statistical analysis in this study focused on descriptive statistics. We tabulated frequencies for the appropriate implants. We calculated percentages by using the total number of medical devices in the relevant population as the denominator. We report the findings according to the PRISMA guidelines.

Results

Data from 9th annual NJR report (2012)

The NJR’s report shows that 142 different brands of femoral stems (57 cemented, 85 uncemented) and 119 different brands of acetabular cups (48 cemented, 71 cementless) were used in primary total hip replacement procedures in 2011 (table 1⇓). The proportion of components implanted that achieved the optimal ODEP rating of 10A varied between the different types of prosthesis; 85% of cemented stems implanted achieved a 10A rating, dropping to 72% of cementless stems, 40% of cemented cups, and only 3% of cementless cups (table 2⇓). Forty eight per cent (126/261) of prosthesis brands implanted in primary hip replacements in 2011 were categorised as having an ODEP rating of pre-entry or unclassified. On closer inspection of the published NJR prostheses tables, we determined eight of these brands to be revision implants, leaving 118 unrated primary prosthesis brands included for further analysis (table 1⇓).

Available evidence for pre-entry and unclassified components

A literature search was conducted for these 118 brands of prosthesis. In total, 8111 papers were reviewed by title and abstract. After application of exclusion criteria, 368 papers were reviewed in full and a further 211 were then discarded, leaving 157 relevant papers. These were classified according to the level of evidence they contained (fig 2⇓). Four of 157 papers had a discrepancy in the level of evidence between the two initial researchers, which were resolved by a third researcher. The inter-observer correlation between the researchers who did the
systematic review was 97%. Unpublished data subsequently provided by manufacturers changed the evidence base rating on one cemented stem, two uncemented stem, and three uncemented cup brands.

We could identify no evidence of clinical effectiveness for 48% (57/118) of pre-entry and unclassified prostheses brands. Excluding custom, revision, and discontinued implants, these accounted for 24% (57/235) of the total number of primary hip replacement prostheses brands listed in the NJR’s report (box 2; fig 3). If we counted level 5 evidence (for example, biomechanical data showing equivalence to pre-existing devices), this still left 42% (49/118) of the pre-entry and unclassified prostheses brands and 21% (49/235) of all brands used bereft of any identifiable evidence to support their use.

Use of component with no available evidence

Applying the results of our systematic review to the NJR’s 9th report data showed that 10 617 prostheses without available evidence of clinical effectiveness to support their use were implanted in patients in 2011 (7.8% of the 136 593 prostheses included in the report) (table 2). This number comprised 157 cemented stems, 936 uncemented stems, 1732 cemented cups, and 7 577 uncemented cups.

Cemented stems represented the group with the lowest proportion of devices implanted without available evidence of clinical effectiveness (0.5%; 157/34 655), followed by uncemented stems (2.8%; 936/33 367). Higher numbers of cups were implanted without such evidence; 7.1% (1 732/24 349) of cemented cups, and 17.1% (7 577/44 222) of cementless cups had no available evidence of clinical effectiveness to support their use (table 3; fig 3).

Discussion

This systematic review of the literature shows that 8% of all primary hip replacement prostheses implanted in 2011 and recorded by the National Joint Registry (NJR) had no readily available evidence relating to their safety or effectiveness. This is likely to be an underestimation of the true problem, as much of the evidence that does exist for the other unrated prostheses is of low quality or relates to short term outcomes only. This is of great concern, particularly in light of the widespread publicity surrounding recent safety problems with regard to some resurfacing and other large diameter metal-on-metal joint replacements.

Evidence ratings

The ODEP system of grading primary hip components offers clinicians a simplified, device specific rating for clear comparison of devices’ performance on relevant clinical criteria.
### Box 2 List of prostheses implanted in 2011 with no readily available evidence of clinical effectiveness

**Unclassified devices**

**Cemented stem**
- Excia Cemented (B Braun/Aesculap)
- Kinectiv Cemented (Zimmer)
- Edinburgh* (Implants International)
- Trilliance† (B Braun/Aesculap)
- Answer (Biomet)
- CTI Cemented Stem (Corin); no longer available
- Response (DePuy)
- Wroblewski Resection (DePuy)
- Kinectiv Cemented (Zimmer)

**Uncemented stem**
- H-Max (Lima)
- FH Modular (FH Orthopaedics)
- MiniMax (Medacta UK Ltd)
- Harmony (Symbios SA)
- Lima SL (Lima)
- LPS (DePuy)
- Euros Cementless (Euros)

**Cemented cup**
- Exeter RimFit (Stryker)
- CMK Cemented Cup‡ (Biomet)
- Exceed ABT Cemented (Biomet)
- CCB (Implants International)
- Edinburgh Cup* (Implants International)
- Alfa Cemented Cup (B Braun/Aesculap); discontinued 2008
- PE (Symbios SA)
- Luna (Amplitude)
- Charnley KS (DePuy)
- Cone Cup (Medacta UK Ltd)
- Snap Fit (Waldemar Link)

**Uncemented cup**
- April (Symbios SA)
- Gyros (DePuy)
- Regenerex Ringloc + (Biomet)
- Novation (Exatech)
- Beta Cup (Waldemar Link)
- Split Cup (Surplas)
- Restoration ADM Cup (Stryker)
- MMC Cup (Zimmer)
- Maxera (Zimmer)
- Dynasty (Wright Medical UK Ltd)
- Sirius Cementless Cup (Euros)

**Pre-entry devices**

**Cemented stem**
- Corail Cemented (DePuy)

**Uncemented stem**
- Metafix Stem* (Corin)
- MiniHip* (Corin)
- Silent (DePuy)
- Amoda (Comis Orthopaedics)
- Trabecular Cementless Stem (Zimmer)
- Novation Element Stem (Exatech)
- Novation Stem (Exactech)

**Cemented cup**
- Apollo‡ (Biomet)
- Polarcup Cemented (Endo Pius (UK) Ltd)

**Uncemented cup**
- CSF plus* (Joint Replacement Instrumentation Ltd)
- Continuum (Zimmer)
- Trinity* (Corin)
Regulatory process

The regulation process also seems to be entirely inadequate. The award of a Conformité Européenne mark is conditional on a device meeting a series of laboratory based standards that may not equate to the safety or effectiveness of an implant in patients. A Conformité Européenne mark may also be awarded in cases of “existing similarity,” where the new device closely resembles an existing design. However, as discussed above, small changes in design have been shown to have major deleterious effects on an implant’s clinical effectiveness or lifespan.  

The NJR is effective in auditing current practice but was designed to monitor the success of implants in relation to when and how often they need to be replaced or revised. Thus, although patient reported outcome measures are due to be introduced as part of the annual report, in its current form the NJR’s report does not recognise problems with implants until they are revised and so may not be the best tool for evaluating new prostheses. Most implants are typically introduced in small numbers initially, which makes outliers difficult or impossible to detect. This is shown by the recent problems with some metal-on-metal joint replacements, which were identified in single centre cohort studies more than three years before the NJR identified them as outliers. Arguably, well designed controlled studies would have identified the problem even more quickly with fewer patients experiencing the adverse consequences of a substandard design. In addition, the NJR lists only implants that are in open use and for which reports are submitted. Consequently, devices that are available to some surgeons but have not yet been made available on the open market do not appear in the NJR’s report. These factors can lead to a delay in the identification of failing implants. The National Institute for Health and Care Excellence (NICE) suggests that the use of more refined outcome measures such as radiostereometric analysis, which aims to identify early loosening of implants by using bi-planar radiographs, may help to detect early problems. The phased introduction of devices, combined with the use of surrogate outcome measures, has been called for to provide early identification of poorly performing implants. This correlates with the IDEAL framework ensuring that the introduction of new devices is controlled and regulated in phases. The future of medical device regulation needs to be a careful balance between the requirement to facilitate innovation and the imperative to safeguard patients. Following the example of pharmaceutical regulation by implementing a phased introduction of new orthopaedic implants would seem prudent. It has been proposed that innovative technologies should be made available in a few specialist units, where the evidence can be objectively gathered and any problems with implants’ resilience identified quickly using surrogate outcome measures. The counter argument debates the damage over-regulation can do to innovation and development in the field of medicine. The cost of medical implants and devices may also rise if they are required to undergo lengthy pre-clinical and clinical testing. However, significant cost savings may accrue if the number of devices on the market were limited to only those devices with a solid evidence base for their use. Tackling this question requires a delicate balance.

Many of the concerns in this process relate to the evidence required to market an implant or device. Public availability of a list of current medical implants and devices, including pre-marketing data and peer reviewed publications, is needed. This may improve transparency in the early stages of implant introduction, allowing surgeons and patients to make more informed decisions.

Limitations of study

A major limitation to our study relates to the requirement that a prosthesis be specifically named in a publication to meet our inclusion criteria. Relevant published evidence may therefore not have been identified, for instance, if a specific implant was simply referred to as “an uncemented acetabular cup” or as a “proximal loading femoral stem.” However, we felt that the explicit naming of a prosthesis is necessary if surgeons or commissioners are to locate evidence that can inform their decision making as to the use of a particular prosthesis. Nevertheless, evidence for some implants may have been missed if the device had undergone a change of brand name since evidence was published. However, in all cases in which no evidence could be found, we contacted the manufacturers directly to request supportive data for the use of their implant, which gave the opportunity for missed evidence to be brought to our attention.

Another limitation is that the evidence for some early phase implants will not have been detected if they are part of ongoing prospective cohort studies or randomised controlled trials that have yet to report. We have included, where identified, “pre-entry” devices in our analysis, and these prostheses are likely to be undergoing prospective studies and so would not yet have published evidence available in the literature. However, the number of prostheses implanted as part of prospective and unreported clinical studies is likely to represent only a small proportion of the 10 617 devices implanted with no available evidence.

The evidence presented in this study relates to prostheses implanted in the UK. However, we believe the results of this study can be applied to other healthcare settings, given that most of the prostheses in our systematic review are available in most other developed countries, in Europe, Australasia, and the United States.

Conclusion

NICE has set a clear benchmark of an ODEP rating of 10C as the minimum clinical standard it recommends for general implantation. Our review has determined that only 49%
(118/261) of prostheses implanted overall achieved a 10A/B/C rating and that almost one in four prosthesis brands available to surgeons have no evidence to support their use. Although these brands are used relatively infrequently, at least 7% of all prostheses implanted lack evidence of clinical effectiveness or longevity. This study shows that the need still exists for an improved and more rigorous approach to regulation of devices to avoid devices with no available evidence being used in a widespread and uncontrolled manner.

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Ethical approval: Not needed.

Data sharing: No additional data available

Declaration of transparency: The lead author affirms that this manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as planned (and, if relevant, registered) have been explained.

1. Wilmshurst P. The regulation of medical devices. BMJ 2011;342:d2822.
2. European Commission. Exploring innovative healthcare—the role of medical technology innovation and regulation. European Commission, 2011.
3. Somel S. Medical device development: U.S. and EU differences. 2006. http://appliedclinicaltrialsonline.fda.gov/appliedclinicaltrials/article/articleDetail.jsp?id=386480.
4. Cohen D. Out of joint: the story of the ASR. BMJ 2011;342:d2905.
5. McCullough P. The EU’s system for regulating medical devices. BMJ 2012;345:e7126.
6. Dias J, Kay P, Porter M, Briggs T. Restoring your mobility. British Orthopaedic Association, 2012.
7. European Commission Enterprise and Industry. Notified bodies. http://ec.europa.eu/enterprise/newapproach/vindas/index.cfm?useaction=directive.notifiedbody&dir_id=13&type_dir=NCBPDI&proc_id=99999&proc_id=99999&com_id=99999&proc_id=99999&_lang=en.
8. Fraser AG, Daubert JC, Van de Werf F, Estes NA 3rd, Smith SC Jr, Krucoff MW, et al. Clinical evaluation of cardiovascular devices: principles, problems, and proposals for European regulatory reform. Report of a policy conference of the European Society of Cardiology. Eur Heart J 2011;32:1673-86.
9. National Joint Registry for England and Wales. 9th annual report. National Joint Registry, 2012 (available from www.hqip.org.uk/national-joint-registry-9th-annual-report-2012/).
10. NHS Supply Chain. ODEP criteria. www.supplychain.nhs.uk/odep/.
11. OCEBM Levels of Evidence Working Group. The Oxford 2011 levels of evidence: Oxford Centre for Evidence-Based Medicine, www.cebm.net/index.aspx?o=5663.
12. Foster RL. Reporting guidelines: CONSORT, PRISMA, and SQUIRE. J Spec Pediatr Nurs 2012;17(1):1-2.
13. De Staiger RN, Hang JR, Miller LN, Graves SE, Davidson DC. Five-year results of the ASR XL Acetabular System and the ASR Hip Resurfacing System: an analysis from the Australian Orthopaedic Association National Joint Replacement Registry. J Bone Joint Surg Am 2011;93:2297-93.
14. Hauptfleisch J, Glyn-Jones S, Beard DJ, Gill HS, Murray DW. The premature failure of the Charney Elite-Plus stem: a confirmation of RSA predictions. J Bone Joint Surg Br 2006;88:179-83.
15. Roy N, Hossan S, Ayeo C, McGee HM, Jacobs LG. 3M Capital hip arthroplasty: 3-8-year follow-up of 208 primary hip replacements. Acta Orthop Scand 2002;73:400-2.
16. Murray DW, Carr AJ, Bulstrode CJ. Which primary total hip replacement? J Bone Joint Surg Br 1995;77:520-7.
17. McCullough P, Altman DG, Campbell WB, Plum DR, Glassizou P, Marshall JC, et al. No surgical innovation without evaluation: the IDEAL recommendations. Lancet 2009;374:1105-12.
18. Nielssen RG, Pilie BG, Karrholm J, Malchau H, Niewenhuize MJ, Valstar ER. RSA and registries: the quest for phased introduction of new implants. J Bone Joint Surg Am 2011;93(suppl 3):1-82.5.
19. Glynn-Jones S, Pandit H, Kwon YM, Doll H, Gill HS, Murray DW. Risk factors for inflammatory pseudotumour formation following hip resurfacing. J Bone Joint Surg Br 2009;91:1566-74.
20. Smith AJ, Dickie P, Howard PW, Bjom AW. For the National Joint Registry for England and Wales. Failure rates of metal-on-metal hip resurfacings: analysis of data from the National Joint Registry for England and Wales. Lancet 2012;380:1759-66.
21. Di Mario C, James S, Dukie D, Sabale M, Dergekhian M. Commentary: The risk of over regulation. BMJ 2011;342:d3020.
22. National Institute for Health and Clinical Excellence. Guidance on the selection of prostheses for primary total hip replacement. NICE, 2000.

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What is already known on this topic
The high failure rate of some metal-on-metal hip replacements has highlighted the need for an adequate evidence base for orthopaedic implants.

Many implants are available to orthopaedic surgeons, but how many of these have evidence of clinical effectiveness to support their use is not known.

What this study adds
A quarter of prostheses available to the surgeon for use in primary total hip arthroplasty in the UK have no available evidence of clinical effectiveness to support their use.

Almost 8% of all primary hip replacement prostheses implanted in 2011 had no readily available evidence relating to their safety or effectiveness.

Tables

Table 1 Breakdown of Orthopaedic Data Evaluation Panel’s rating for each device category from 9th National Joint Registry of England and Wales report. Values are numbers (percentages)

| ODEP rated | Cemented stem (n=57) | Uncemented stem (n=85) | Cemented cup (n=48) | Uncemented cup (n=71) | Total (n=261) |
|------------|----------------------|------------------------|---------------------|-----------------------|---------------|
| 10A        | 14                   | 16                     | 11                  | 9                     | 50            |
| 10B        | 2                    | 1                      | 1                   | 2                     | 6             |
| 10C        | 2                    | 2                      | 0                   | 1                     | 5             |
| 7A         | 3                    | 3                      | 1                   | 4                     | 11            |
| 7B         | 2                    | 1                      | 0                   | 1                     | 4             |
| 5A         | 5                    | 4                      | 2                   | 12                    | 23            |
| 5B         | 0                    | 1                      | 2                   | 0                     | 3             |
| 3A         | 4                    | 7                      | 1                   | 2                     | 14            |
| 3B         | 0                    | 0                      | 0                   | 1                     | 1             |
| Total      | 32                   | 35                     | 18                  | 32                    | 117 (45)      |

Unrated

|               | Cemented stem (n=57) | Uncemented stem (n=85) | Cemented cup (n=48) | Uncemented cup (n=71) | Total (n=261) |
|---------------|----------------------|------------------------|---------------------|-----------------------|---------------|
| Unclassified  | 14                   | 33                     | 23                  | 24                    | 94            |
| Pre-entry     | 3                    | 13                     | 3                   | 13                    | 32            |
| Custom/revision/discontinued | 8                    | 4                      | 4                   | 2                     | 18            |
| Total         | 25                   | 50                     | 30                  | 39                    | 144 (55)      |
Table 2 | Numbers (percentages) of prostheses implanted, 2011

| Rating          | Cemented stem | Uncemented stem | Cemented cup | Uncemented cup | Total  |
|-----------------|---------------|-----------------|--------------|----------------|--------|
| 10A             | 30 689 (88.6) | 23 920 (71.7)   | 9751 (40.0)  | 1531 (3.5)     | 65 891 |
| 10B             | 520 (1.5)     | 147 (0.4)       | 41 (0.2)     | 35 (0.1)       | 743 (0.6)|
| 10C             | 118 (0.3)     | 142 (0.4)       | 0 (0)        | 69 (0.2)       | 329 (0.2)|
| 7A              | 260 (0.8)     | 222 (0.7)       | 223 (0.9)    | 18 639 (42.1)  | 19 944 (14.2)|
| 7B              | 164 (0.5)     | 3514 (10.5)     | 0 (0)        | 6 (<0.1)       | 3684 (2.7)|
| 5A              | 2172 (6.3)    | 1870 (5.6)      | 8436 (34.6)  | 10 530 (23.8)  | 23 008 (16.9)|
| 5B              | 0 (0)         | 49 (0.1)        | 1171 (4.8)   | 0 (0)          | 1220 (0.9)|
| 3A              | 419 (1.2)     | 639 (1.9)       | 1735 (7.1)   | 3405 (7.7)     | 6198 (4.5)|
| 3B              | 0 (0)         | 0 (0)           | 0 (0)        | 7 (<0.1)       | 7 (<0.1)|
| Unclassified    | 204 (0.6)     | 1061 (3.2)      | 2746 (11.3)  | 1434 (3.2)     | 5445 (4.0)|
| Pre-entry       | 84 (0.2)      | 1775 (5.3)      | 206 (0.8)    | 8552 (19.3)    | 10 617 (7.6)|
| Custom/revision/discontinued | 25 (0.1) | 28 (0.1)       | 40 (0.2)     | 14 (<0.1)      | 107 (0.1)|
| Total           | 34 655 (100)  | 33 367 (100)    | 24 349 (100) | 44 222 (100)   | 136 593 (100)|
| Evidence level | Cemented stem | Uncemented stem | Cemented cup | Uncemented cup | Total |
|----------------|---------------|-----------------|--------------|----------------|-------|
| 1A             | —             | —               | —            | —              | —     |
| 1B             | 1 (20)        | 2 (218)         | —            | 3 (47)         | 6 (285) |
| 1C             | —             | —               | —            | —              | —     |
| 2A             | —             | —               | —            | —              | —     |
| 2B             | —             | 3 (552)         | —            | 2 (1,213)      | 5 (1,765) |
| 2C             | —             | —               | 1 (1)        | —              | 1 (1) |
| 3A             | —             | —               | —            | —              | —     |
| 3B             | 1 (36)        | 1 (9)           | 1 (149)      | —              | 3 (194) |
| 4              | 6 (190)       | 18 (958)        | 11 (1069)    | 7 (1726)       | 42 (3941) |

| No of prostheses available with no evidence | 9 (157); 9 unclassified | 14 (936); 7 pre-entry; 7 unclassified | 13 (1732); 2 pre-entry; 11 unclassified | 21 (7577); 10 pre-entry; 11 unclassified | 57 (10 617); 20 pre-entry; 37 unclassified |
| No of prostheses implanted with no evidence | 0.5% of those implanted (157 of 34 655) | 2.8% of those implanted (936 of 33 367) | 7.1% of those implanted (1732 of 24 349) | 17.1% of those implanted (7577 of 44 222) | 7.8% of those implanted (10 617 of 136 593) |

Revision devices have been excluded (2 cemented stems, 3 uncemented stems, and 3 uncemented cups).
Figures

**Fig 1** Process used to identify unrated devices and determine evidence levels. NJR=National Joint Registry; ODEP=Orthopaedic Data Evaluation Panel.

**Fig 2** Flow chart of literature search and evidence level classification.
Fig 3 Percentage of available prosthesis brands with no evidence of clinical effectiveness in 2011 (top) and percentage of prostheses implanted with no evidence of clinical effectiveness in 2011 (bottom)