Management Guidelines for Metal-on-metal Hip Resurfacing Arthroplasty: A Strategy on Followup

Abstract

Despite the initial promise of metal-on-metal (MoM) implants as the ideal bearing surface for hip replacements and resurfacings, high short term failure rates due to an adverse reaction to metal debris (ARMD) have led to a dramatic reduction in the number of MoM implants used in the modern era. With over one million patients worldwide having undergone hip operations utilizing a MoM bearing surface, the long term outcomes for such patients remains unknown, and there is much debate as to the most effective management of these patients. Although several regulatory bodies have released guidelines on the management of patients with MoM hips, these recommendations remain open to interpretation, and the most effective management for these patients remains unclear. The aim of this review is to compare the current guidelines for managing patients with MoM hips and also to discuss established ARMD risk factors, evidence regarding the optimum management for patients with MoM hips, and the indications for revision surgery. Furthermore, although specialized laboratory tests and imaging can be used to facilitate clinical decision making, over-reliance on any single tool should be avoided in the decision making process, and surgeons should carefully consider all findings when determining the most appropriate course of action.

Keywords: Adverse reaction to metal debris, follow up, guideline, hip replacement, hip resurfacing, metal-on-metal, patient management

MeSH terms: Replacement, arthroplasty, hip, reoperation, prosthesis

Introduction

The increasing incidence of polyethylene associated complications in Metal-on-Polyethylene (MOP) bearings in the early 2000s led to a resurgence of the metal-on-metal (MoM) bearing in total hip arthroplasty (THA) surgery. Initial promise from MoM THAs was rooted in a proposed reduction in failure rates due to diminished volumetric wear, greater component stability, and avoidance of polyethylene wear-induced osteolysis. Parallel to this, MoM bearing was found to have a number of unique advantages in hip resurfacing. These included the ability to preserve bone stock in the younger patients, low dislocation rates, improved proprioception and gait, and the perceived ease of future conversion to a THA. In addition, it has been suggested that it is safe to return to high impact activity following hip resurfacing – something highly valuable to younger individuals undergoing hip resurfacing procedures.

Despite these initial benefits, high short term failure rates due to adverse reaction to metal debris (ARMD) in MoM bearings have been reported, leading to concerns over their use. ARMD is the sequela of metal debris released from MoM bearings due to wear and corrosion. This process results in destructive soft-tissue masses adjacent to the bearing and often necessitates surgical revision. As such, ARMD lesions have been branded “inflammatory pseudotumor.” Although the size of particulate matter generated by a MoM bearing is significantly smaller than that of a MoP bearing, these adverse inflammatory effects are theorized to be the result of a greater volume of particle generation and these particles being more biologically active. In addition, the significant bone loss and muscle damage that results from ARMD means that short-term outcomes following revision surgery are often poor. The poor outcomes associated with MoM bearings, and the subsequent recall of specific implants has led to a

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dramatic reduction in the number of MoM THA and hip resurfacing procedures performed today. However, as over a million patients worldwide have already been implanted with MoM devices, a clear management strategy for patients with MoM hips is essential.\textsuperscript{20} Regulatory authorities have published guidance on the regular followup and management of patients with MoM hips.\textsuperscript{21-26}

**Metal-on-metal management guidelines**

Management guidelines for MoM hips have been published by five authorities in various countries and regions. These include:

1. United States of America Food and Drug Administration (FDA)\textsuperscript{21}
2. United Kingdom Medical and Healthcare Products Regulatory Agency (MHRA)\textsuperscript{22}
3. European Federation of National Associations of Orthopaedics and Traumatology (EFORT)\textsuperscript{23}
4. Health Canada\textsuperscript{26}
5. Therapeutic Goods Administration of Australia (TGA).\textsuperscript{24}

**Surveillance strategy guidance**

Each authority provides guidance on the patient selection for followup, and the frequency at which this should occur [Table 1].

**Symptomatic and asymptomatic metal-on-metal hip investigations and imaging**

Each of the five authorities also provide guidance regarding the investigations and imaging patients with MoM hips should receive [Table 2], and further discusses which metal ions should be sampled and the acceptable values for each [Table 3].

**Which patients are at risk of adverse reaction to metal debris?**

Patients can be stratified into high- and low-risk groups depending on implant specific and patient specific factors. Implant specific factors that increase the risk of ARMD are THAs with large diameter femoral head components or the DePuy ASR implants. In addition, several implant specific factors correlate with a lower risk of ARMD, including THAs with small diameter femoral head components, and hip resurfacing implants.\textsuperscript{22} Several studies have described patient factors associated with increased failure rates in MoM hips. In a systematic review, an increased risk of developing ARMD, and higher rates of dislocation, aseptic loosening and revision after MoM hip resurfacings were found in female patients.\textsuperscript{27} Although the reasons for this finding are unclear, it has been theorized that the higher prevalence of developmental dysplasia in females may affect the accuracy of acetabular component positioning, resulting in higher rates of impingement, edge loading, and wear.\textsuperscript{28} In addition, metal hypersensitivity has been reported as an important factor in the pathology of ARMD following histological examination of peri prosthetic tissue.\textsuperscript{29,30} Another theorized risk factor for ARMD is that of low body mass index (BMI). A recent study demonstrated a negative correlation between BMI and elevated Cr level.\textsuperscript{28} Although the authors suggested that differences in activity level could explain this relationship, further studies

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**Table 1: Surveillance strategy guidance by authorities**

<table>
<thead>
<tr>
<th>Authority</th>
<th>Recommended followup schedule for asymptomatic patients</th>
<th>Recommended followup schedule for symptomatic patients</th>
<th>Additional guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDA*</td>
<td>Every 1-2 years</td>
<td>Every 6 months</td>
<td>Closer followup should be considered for those with increased risk factors of increased device wear. These include female, those who received hip resurfacing with a small femoral head component (≤44 mm), those with bilateral hip implants, and those with suboptimal component alignment</td>
</tr>
<tr>
<td>MHRA</td>
<td>Annual followup for THA cases with large femoral head component (≥36 mm)</td>
<td>Annual followup for hip resurfacing cases</td>
<td>No guidelines are given for patients that do not meet these criteria</td>
</tr>
<tr>
<td>EFORT</td>
<td>Annual followup for THA cases with a large femoral head component (≥36 mm) and hip resurfacing cases with defined risk factors for ARMD</td>
<td>No specific guidelines provided</td>
<td>Defined risk factors are described as female gender, low coverage arc, and small femoral head size</td>
</tr>
<tr>
<td>Health Canada*</td>
<td>Annual followup for first 5 years for asymptomatic patients with MoM hips</td>
<td>No specific guidelines provided</td>
<td>N/A</td>
</tr>
<tr>
<td>TGA</td>
<td>Annual follow-up for cases with large (≥45) or small (≤36) femoral head components</td>
<td>TGA Australia recommends annual followup for symptomatic patients with MoM hip cases</td>
<td>No guidelines given for patients other than those described</td>
</tr>
</tbody>
</table>

*Unlike other authorities, the FDA and Health Canada make no distinction between the followup of hip replacement and hip resurfacing. FDA=Food and Drug Administration, MHRA=Medical and Healthcare Products Regulatory Agency, EFORT=European Federation of National Associations of Orthopedics and Traumatology, TGA=Therapeutic Goods Administration, THA=Total hip arthroplasty, ARMD=Adverse reaction to metal debris, MoM=Metal-on-metal, N/A=Not available
assessing activity level in isolation have shown this not to be the case.31

**What is the evidence base for the frequency of followup?**

As asymptomatic ARMD lesions have been observed in up to 61% of MoM hips, followup strategies according to symptoms will result in undetected ARMD in a large number of patients.32-34 This necessitates regular investigations to identify asymptomatic ARMD.

The necessity of blood metal ion concentration testing has been repeatedly demonstrated, with higher complication rates shown in patients with elevated blood metal ion concentrations.7,35-37 However, the optimum frequency of these blood tests remains unclear. One study found that blood metal ion levels in patients with MoM THAs increased significantly over an 8-month period and that similar increases were not shown in patients who received hip resurfacing.38 This led the researchers to suggest that annual blood metal ion measurements are of values in THA followup but not in hip resurfacing. An additional study found that repeating blood metal ion measurements at a mean postoperative time of 27 months in MoM hip resurfacing cases showed no significant increases in Co when compared to the concentration after surgery.36 Although these findings highlight the ambiguity with regards to optimum followup frequency, elevated blood levels require further investigation. One study has suggested that if metal ion levels are elevated, radiographs should be evaluated for osteolysis and inappropriate component position. If either is found, then revision surgery should be considered.39

In addition to the unclear evidence regarding metal ion blood tests, the benefits of short-term imaging are also questionable. Two previous studies have suggested that patients with normal initial imaging do not need repeat imaging in the short-term, as no significant change was found when repeat imaging was performed at various time intervals.40,41 However, the usefulness of magnetic resonance imaging (MRI) scans in surgical decision making decreases drastically if not regularly updated. One study demonstrated that MRI scans performed over 12 months previously had a sensitivity of 29%. This was compared to a sensitivity of 88% for scans that were 3-month-old.42 From these results, it was suggested that repeat imaging should be performed at least annually, and the usefulness of results over a year old severely questioned. Furthermore, as cyst progression has not been observed earlier than 6 months postoperatively, annual followup has been recommended in other studies.43 In addition, although some work has begun to assess the time-course of growth of the pseudotumors, there is a large discrepancy between results. Growth of pseudotumors, remission of small masses, and static pseudotumors have all been documented, highlighting the need for a greater

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**Table 2: Guidance regarding investigation and imaging of MoM hips by authorities**

<table>
<thead>
<tr>
<th>Authority</th>
<th>Guidance regarding investigation and imaging of asymptomatic MoM hips</th>
<th>Guidance regarding investigation and imaging of symptomatic MoM hips</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDA</td>
<td>No guidance provided</td>
<td>Radiograph of hips, imaging (MARS MRI, CT or ultrasound) and metal ion sampling</td>
</tr>
<tr>
<td>MHRA</td>
<td>Metal ion sampling for asymptomatic THA cases, and imaging is recommended if blood metal ion levels rise</td>
<td>Metal ion sampling and imaging (MARS MRI or ultrasound)</td>
</tr>
<tr>
<td>EFORT</td>
<td>Radiographs and metal ion sampling for all asymptomatic hips. Further imaging for asymptomatic hips with an abnormality in radiographs, or with Co concentrations between 2-7 µg/L</td>
<td>Radiograph of hips, imaging (MARS MRI, CT or ultrasound) and metal ion sampling</td>
</tr>
<tr>
<td>Health Canada</td>
<td>No guidance provided</td>
<td>Radiographs, imaging (MARS MRI or ultrasound) and metal ion sampling</td>
</tr>
<tr>
<td>TGA</td>
<td>Radiographs, imaging and metal ion sampling for cases with femoral heads ≤36 mm or ≥45 mm</td>
<td>Radiographs, imaging (MARS MRI or ultrasound) and metal ion sampling</td>
</tr>
</tbody>
</table>

FDA=Food and Drug Administration, MHRA=Medical and Healthcare Products Regulatory Agency, EFORT=European Federation of National Associations of Orthopedics and Traumatology, TGA=Therapeutic Goods Administration, MARS=Metal artifact reduction sequence, MRI=Magnetic resonance imaging, CT=Computed tomography, MoM=Metal-on-metal, THA=Total hip arthroplasty

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**Table 3: Metal ion sampling recommendations and advised thresholds by authorities**

<table>
<thead>
<tr>
<th>Authority</th>
<th>Metal ion sampling recommendations</th>
<th>Thresholds advised</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDA</td>
<td>Sampling of whole blood cobalt and chromium</td>
<td>Concentrations above 7 µg/L should be a cause for concern</td>
</tr>
<tr>
<td>MHRA</td>
<td>Sampling of whole blood cobalt and chromium</td>
<td>Asymptomatic patients with concentrations between 2 µg/L and 7 µg/L require cross-sectional imaging</td>
</tr>
<tr>
<td>EFORT</td>
<td>Sampling of whole blood cobalt only</td>
<td>Concentrations above 7 µg/L should be a cause for concern</td>
</tr>
<tr>
<td>Health Canada</td>
<td>Sampling of whole blood and serum cobalt and chromium</td>
<td>Not provided</td>
</tr>
<tr>
<td>TGA</td>
<td>Sampling of whole blood or serum cobalt and chromium</td>
<td>Not provided</td>
</tr>
</tbody>
</table>

FDA=Food and Drug Administration, MHRA=Medical and Healthcare Products Regulatory Agency, EFORT=European Federation of National Associations of Orthopaedics and Traumatology, TGA=Therapeutic Goods Administration
Thresholds for concern with metal ion levels

Metal ion levels have been extensively used as a surrogate marker for patients with MoM bearings related wear.\(^4\) The population background level of Co in blood has been shown to be 0.5 µg/L, and the mean Co blood levels in patients with well functioning MoM hip implants ranges between 0.2 and 4 µg/L.\(^4\) There is correlation seen with wear rates of the MoM joint, where blood levels of 2 µg/L can be expected with wear rates of 2 mm\(^2\)/years.\(^3\)\(^,\)\(^7\) In addition, Co and Cr ion levels are influenced by the type of MoM implant, head size, implant design, and component positioning.\(^4\)\(^,\)\(^5\) Although one study has shown that Co is more reliable than Cr, and that whole blood testing is quicker and more accurate than serum, either whole blood or serum, and Co or Cr, can be used for monitoring as long as the testing modality is consistent.\(^4\)\(^,\)\(^6\)\(^,\)\(^9\) Furthermore, there should be standardized protocols in place for the collection of serum or blood samples, as needle type and collection techniques can influence reported ion concentrations.\(^2\)\(^,\)\(^5\)\(^,\)\(^6\)

A retrospective study evaluating the use of Co and Cr blood sampling established that cut-off value of 7 µg/L had a specificity of 89% and a sensitivity of 52% in detecting failed MoM articulations in patients with normal imaging results, and has been extensively adopted.\(^5\) Similarly, Hart et al. demonstrated 93% specificity with a threshold set at 7 µg/L.\(^3\) Although no single metal ion threshold will reliably identify all ARMD patients, EFORT suggests that 7 µg/L may be too high a threshold, instead recommending a threshold of between 2 µg/L and 7 µg/L, with the exact level still to be determined. In addition, the interpretation of metal ion levels can be difficult in patients with a systemic renal disease or additional metallic implants, such as in patients with bilateral MoM implants. Therefore, metal ions levels should be used to guide and should not be used in isolation to determine the need for revision surgery.\(^3\)\(^,\)\(^6\)\(^,\)\(^5\)\(^2\) Surgeons must carefully consider findings from clinical examination, blood test results, and radiologic investigations when determining the likelihood of a failing MoM THA or hip resurfacing.

The essence of clinical examination

Physical examination is essential in the assessment of painful hips in patients with MoM implants. Patients with ARMD commonly present with early recurrence of preoperative symptoms, or with persistent pain. In addition, radiation of pain to the greater trochanter and down the thigh is a common feature and can result in patients displaying an antalgic gait.\(^5\) Patients may also present with a palpable mass or fluid collection near the hip.\(^4\) Over time, this may progress to instability, with or without dislocation, and patients may complain of a “clicking” or “clunking” sensation in the hip. Patients may also experience other symptoms, such as stiffness or reduced range of movement often most evident in the abduction. It is, therefore, imperative to assess the active and passive range of motion of the hip joint. Reproduction of pain on passive extension and active flexion may indicate concomitant iliopsoas tendinopathy. Abductor weakness may indicate peri-prosthetic soft-tissue involvement and is occasionally accompanied by a rash indicating a reaction to the metal ions.\(^5\)

It is vital to obtain a complete history, perform a thorough clinical examination, and conduct relevant investigations to exclude other potential diagnoses. As joint sepsis must be suspected in patients with a history of complicated wound healing, physical examination of the skin should be performed, looking for signs of infection and previous scars. A comprehensive spine and neurovascular examination should also be performed to evaluate possible neurogenic and vascular pain generators.\(^4\) Due to the destructive nature of ARMD, appropriate assessment in a timely manner is essential. Due to the high rate of asymptomatic pseudotumors, it is also very important for general practitioners to remain vigilant when examining any patient who has previously undergone a MoM THA or a hip resurfacing.

When should revision surgery be considered and conducted?

All the authorities recommend revision surgery in patients with abnormal cross-sectional imaging and/or rising blood metal ions. In addition, the FDA recommends deciding on revision surgery in the context of the overall clinical picture, examination findings and test results, and further suggests early revision surgery in patients with progressive lesions. Both MHRA and EFORT recommend revision surgery if there are abnormal imaging findings and/or if increasing blood metal ion levels are found. EFORT further suggests that a blood Co level above 20 µg/L is an indication for revision. Conversely, Health Canada recommends revision surgery if patients are asymptomatic and if a positive MRI appearance (soft tissue mass) is found. They also suggest revision surgery in asymptomatic cases if a soft-tissue mass, or masses, found on MRI are increasing in size. TGA recommends revision surgery if symptoms are persistent with imaging abnormalities and/or increasing of blood metal ions.

Poor short term outcomes have been reported following ARMD revision, with only a few studies assessing prognostic factors of post revision outcome.\(^5\)\(^,\)\(^6\)\(^,\)\(^1\)\(^8\)\(^,\)\(^5\)\(^6\)\(^,\)\(^7\) One study attributed improved outcomes following ARMD revision to early surgery and increased experience.\(^3\) In the latest internationally recognized guidelines published by Scientific Committee on Emerging and Newly Identified Health Risks (SCENIHR), there was no consensus on the appropriate timing of revision surgery for ARMD in MoM THA or hip resurfacings.\(^2\) SCENIHR do
however recommended that revision should be considered when symptoms become persistent, progressive, and unmanageable. Furthermore, they suggest that any patient exhibiting progressive osteolysis, expanding or large pseudotumors, progressive neck thinning or very high metal ion concentrations should also be considered for revision surgery. Although these recommendations are useful until prognostic factors of adverse outcomes are identified, the need for defined thresholds for revision surgery is paramount. Surgeons should keep in mind that a decision to revise should not be based on a single investigation. Instead, the decision should take the implant type, patient symptoms, imaging findings, metal ion levels, and the activity levels of the patient into account. When revising MoM implants, it is recommended to utilize a non-MoM articulation such as ceramic-on-ceramic or ceramic-on-polyethylene to reduce local metal ion release to decrease the chance of recurrence of ARMD, because poorer outcomes have been reported in patients revised using MoM articulations.

**Conclusion**

Patients with MoM bearings should be monitored closely with physical examination, radiographs, metal ion measurement, and imaging. Evidence supporting the management of MoM hips is still lacking in many scenarios, and regulatory guidelines are open to interpretation. When considering revision surgery, no single investigation or aspect of the history should be taken in isolation. Decisions should be taken on a case-by-case basis, with consideration given to all aspects of investigation results and the patient’s clinical condition. Until supporting evidence is available, an evidence-based multidisciplinary approach is considered a safe method to help surgeons make decisions and potentially improve patient outcomes.

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**Conflicts of interest**

There are no conflicts of interest.

**References**


38. Amanatullah DF, Sucher MG, Bonadurer GF. Metal-on-metal hip prostheses: Correlation between debris in the synovial fluid and levels of cobalt and chromium ions in the bloodstream. Int Orthop 2014;38:469-75.


