PUBLIC PETITIONS COMMITTEE CONSIDERATION OF PE1517
QUESTIONS / ISSUES ARISING FROM COMMITTEE MEETINGS

THURSDAY 29 SEPTEMBER 2016

Scottish Government—

- Please provide an update on the work of the expert group.

TUESDAY 8 MARCH 2016

UK Government—

- How can the Public Petitions Committee best influence the debate on the regulation of mesh implants at a UK level?

TUESDAY 26 JANUARY 2016

The Scottish Government—

- Please ensure that the expert group’s work is published and that it operates in the most open and transparent way possible.
- Please provide an update on the SIMS trial in the light of the conclusions of the Independent Review’s interim report
- Please comment on reports that some mesh manufacturers have used plastic that was not for internal human use.
- Please comment on the petitioners’ calls for a Scottish medical watchdog to replace the MHRA.

TUESDAY 12 MAY 2015

British Society of Urogynaecology—
Royal College of Obstetricians and Gynaecologists—

- What are your views on the petition?

European Commission, Scientific Committee on Emerging and Newly Identified Health Risks—

- Please provide an update on when the SCENIHR will be publishing its opinion on TVM devices.

TUESDAY 24 FEBRUARY 2015

The Scottish Government—

- Please ensure that the evidence received by the Committee at its meeting on 24 February 2015 is taken into account by the Scottish Government’s Independent Review of Transvaginal Mesh Implants.
TUESDAY 11 NOVEMBER 2014

Department of Health (UK Government) —

- What is the view of the Department of Health on the use of polypropylene transvaginal mesh devices?
- Please provide more information on the working group considering the use of polypropylene transvaginal mesh devices which is being chaired by NHS England.

Cabinet Secretary for Health—

- When will the report from the review of the use of polypropylene transvaginal mesh devices be published?
- What is the Scottish Government’s current view on the liability risks for NHS Scotland with regard to the use of these devices?
- Please provide clarification on why mesh devices continue to be used.
- Please provide reassurance that women undergoing these procedures are able to give their informed consent and that the risks are being clearly explained.
- The Committee would also welcome assurance that any adverse effects arising from the use of mesh medical devices are being consistently and accurately recorded and monitored.

TUESDAY 3 JUNE 2014

Medicines and Healthcare products Regulatory Agency (MHRA)—
NHS National Services Scotland—
European Commission—
Royal College of Surgeons of Edinburgh—
British Medical Association Scotland—
Scottish Regional NHS Boards—

- What are your views on what the petition seeks and the discussions that took place at the meeting on 3 June?