ANNEX I

LIST OF THE NAMES, PHARMACEUTICAL FORM, STRENGTH OF THE MEDICINAL PRODUCTS, ROUTES OF ADMINISTRATION AND MARKETING AUTHORISATION HOLDERS IN THE MEMBER STATES

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|---|---|---|---------------------------------|-------------------------|
| AT - Austria | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | Dotarem - Injektionslösung | 279,32 mg/ml | solution for injection | intravenous use |
| AT - Austria | Insight Agents GmbH Ringstrasse 19 B 69115 Heidelberg Germany | Gadopentetsäure Insight 500 Mikromol/ml Injektionslösung | 469 mg/ml Gadopentetat- Dimeglumin (78,63 mg/ml Gadolinium) | solution for injection | intravenous use |
| AT - Austria | Bayer Austria GmbH Herbststrasse 6-10 1160 Wien Austria | Gadovist 1,0 mmol/ml Injektionslösung in Fertigspritzen/Patronen | 604,72 mg/ml Gadobutrol (157,25 mg/ml Gadolinium) | solution for injection | intravenous use |
| AT - Austria | Bayer Austria GmbH, Herbststrasse 6-10 1160 Wien Austria | Gadovist 1,0 mmol/ml - Injektionslösung | 604,72 mg/ml Gadobutrol (157,25 mg/ml Gadolinium) | solution for injection | intravenous use |
| AT - Austria | Insight Agents GmbH Ringstrasse 19 B 69115 Heidelberg Germany | Magnegita 500 Mikromol/ml Injektionslösung | 469 mg/ml Gadopentetat- Dimeglumin (78,63 mg/ml Gadolinium) | solution for injection | intravenous use |
| AT - Austria | Bayer Austria GmbH, Herbststrasse 6-10 1160 Wien Austria | Magnevist 0,5 mmol/ml - Injektionslösung | 469 mg/ml Gadopentetsäure Dimeglumin | solution for injection/infusion | intravenous use |
| AT - Austria | Bracco S.p.A. Via Egidio Folli, 50 20134 Milano Italy | MultiHance 0,5 M - Injektionslösung | 334 mg/ml Gadobensäure Dimeglumin | solution for injection | intravenous use |
| AT - Austria | Bracco S.p.A. Via Egidio Folli, 50 20134 Milano Italy | MultiHance 0,5 mmol/ml Injektionslösung in einer Fertigspritze | 334 mg/ml Gadobensäure Dimeglumin | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|---|---|------------------------|----------------------------|
| AT - Austria | GE Healthcare Handels GmbH Europlaza Gebäude E Technologiestr. 10 1120 Wien | Omniscan 0,5 mmol/ml - parenterale Kontrastmittellösung | 287 mg/ml Gadodiamid | solution for injection | intravenous use |
| | Austria | 7: ::025 | 101.10 | | • |
| AT - Austria | Bayer Austria GmbH, Herbststrasse 6-10 1160 Wien Austria | Primovist 0,25 mmol/m Injektionslösung in einer Fertigspritze | 181,43 mg/ml Gadoxetsäure-Dinatrium | solution for injection | intravenous use |
| AT - Austria | Bayer Austria GmbH, Herbststrasse 6-10 1160 Wien Austria | Primovist 0,25 mmol/ml Injektionslösung in einer Durchstechflasche | 181,43 mg/ml Gadoxetsäure-Dinatrium | solution for injection | intravenous use |
| AT - Austria | Bracco S.p.A. Via Egidio Folli, 50 20134 Milano Italy | Prohance - Injektionslösung | 279,3 mg/ml Gadoteridol (78,61 mg/ml Gadolinium) | solution for injection | intravenous use |
| BE - Belgium | Codali S.A. Avenue Henri Dunant 31 1140 Bruxelles Belgium | DOTAREM | 0,5 mmol/ml | Solution for injection | intravenous use |
| BE - Belgium | Insight Agents GmbH Ringstrasse 19 B 69115 Heidelberg Germany | GADOPENTETATE CURAGITA 500MICROMOL/ML | 500 micromol/ml | Solution for injection | intravenous use |
| BE - Belgium | Bayer SA-NV J.E. Mommaertslaan 14 1831 Diegem Belgium | GADOVIST | 1,0 mmol-ml | Solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|---------------|--|--------------|-----------------|------------------------|-------------------------|
| BE - Belgium | Insight Agents GmbH Ringstraat 19 B 69115 Heidelberg Germany | MAGNEGITA | 500 micromol-ml | Solution for injection | intravenous use |
| BE - Belgium | Bayer SA-NV J.E. Mommaertslaan 14 1831 Diegem Belgium | MAGNEVIST | 0,5 mmol-ml | Solution for injection | intravenous use |
| BE - Belgium | BRACCO Imaging Deutschland GmbH Max-Stromeyer-Strasse 116 D - 78467 Konstanz Germany | MULTIHANCE | 0,5 M | Solution for injection | intravenous use |
| BE - Belgium | GE HEALTHCARE BVBA Kouterveldstraat 20 1831 DIEGEM Belgium | OMNISCAN | 0,5 mmol-ml | Solution for injection | intravenous use |
| BE - Belgium | Bayer SA-NV J.E. Mommaertslaan 14 1831 Diegem Belgium | PRIMOVIST | 0,25 mmol-ml | Solution for injection | intravenous use |
| BE - Belgium | BRACCO Imaging Deutschland GmbH Max-Stromeyer-Strasse 116 D - 78467 Konstanz Germany | PROHANCE | 279,3 mg-ml | Solution for injection | intravenous use |
| BG - Bulgaria | Bayer Schering Pharma AG Muellerstrasse 178 13353 Berlin Germany | Gadovist | 604.72 mg/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|------------------------|--|--------------------------|-----------------|--|-------------------------|
| BG - Bulgaria | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Primovist | 181,43 mg/ml | solution for injection | intravenous use |
| BG - Bulgaria | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Magnevist | 469 mg/ml | solution for injection | intravenous use |
| BG - Bulgaria | GE Healthcare AS Nycoveien 1-2 P.O.Box 4220 Nydalen N-0401 Oslo Norway | Omniscan | 287 mg/ml | solution for injection | intravenous use |
| BG - Bulgaria | Insight Agents GmbH Ringstrasse 19 B D-69115, Heidellberg Germany | Magnegita | 500 micromol/ml | Solution for injection | Intravenous use |
| CY - Cyprus | PHADISCO LTD 185 YIANNOU GRANIDIOTI AVE, 2235 LATSIA CYPRUS | OMNISCAN | 0.5MMOL/ML | solution for injection | intravenous use |
| CY - Cyprus | BAYER HELLAS ABEE SOROU 18-20 151 25 MAROUSI, ATHENS, GREECE | PRIMOVIST PFS | 0.25MMOL/ML | solution for injection in prefilled syringes | intravenous use |
| CY - Cyprus | BAYER HELLAS ABEE SOROU 18-20 151 25 MAROUSI, ATHENS, GREECE | PRIMOVIST | 0.25MMOL/ML | solution for injection | intravenous use |
| CZ - Czech Republic | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding | GADOVIST 1,0 m mol/ml | 1 mmol/l | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|------------------------|--|---|--------------|------------------------|----------------------------|
| | Germany | | | | |
| CZ - Czech Republic | Insight Agents GmbH Ringstrasse 19 B 69115 Heidelberg Germany | Magnegita 500 mikromol/ml injekční roztok | 0,5 mmo/ml | solution for injection | intravenous use |
| | Bracco Imaging Deutschland GmbH | ProHance | 279,3 mg-ml | solution for injection | intravenous use |
| CZ - Czech Republic | Max-Stromeyer-Strasse 116 D-78467 Konstanz Germany | | | | |
| CZ - Czech Republic | Bracco Imaging Deutschland GmbH Max-Stromeyer-Strasse 116 D-78467 Konstanz Germany | MultiHance | 529 mg/ml | solution for injection | intravenous use |
| CZ - Czech Republic | GE Healthcare AS Nycoveien 1-2, P.O.Box 4220 Nydalen N-0401 Oslo Norway | Omniscan 0,5mmol/l | 287 mg/ml | solution for injection | intravenous use |
| CZ - Czech Republic | Guerbet BP 57400 95943 Roissy CdG cedex France | Dotarem | 279.32 mg/ml | solution for injection | intravenous use |
| CZ - Czech Republic | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Primovist 0,25 mmol/ml | 0.25 mmol/ml | solution for injection | intravenous use |
| DE - Germany | BRACCO Imaging Deutschland GmbH Max-Stromeyer-Strasse 116 D - 78467 Konstanz Germany | MultiHance | 529 mg/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|---|--------------|---|-------------------------|
| DE - Germany | BRACCO Imaging Deutschland GmbH Max-Stromeyer-Strasse 116 D - 78467 Konstanz Germany | MultiHance | 0,5 mmol/ml | solution for injection, prefilled syringe | intravenous use |
| DE - Germany | BRACCO Imaging Deutschland GmbH Max-Stromeyer-Strasse 116 D - 78467 Konstanz Germany | MultiHance XL | 529 mg/ml | solution for injection | intravenous use |
| DE - Germany | Bayer Vital GmbH D-51368 Leverkusen Germany | Gadovist 1,0 mmol/ml Injektioslösung | 604.72 mg/ml | solution for injection | intravenous use |
| DE - Germany | Bayer Vital GmbH D-51368 Leverkusen Germany | Gadovist 1.0 mmol/ml Injektioslösung in Fertigspritzen/Patronen | 604.72 mg/ml | solution for injection | intravenous use |
| DE - Germany | Marotrast GmbH Otto-Schott-Str. 15 D-07745 Jena Germany | Gadograf 1,0 mmol/ml Injektionslösung in Fertigspritzen | 604.72 mg/ml | solution for injection | intravenous use |
| DE - Germany | Marotrast GmbH Otto-Schott-Str. 15 D-07745 Jena Germany | Gadograf 1,0 mmol/ml | 604.72 mg/ml | solution for injection | intravenous use |
| DE - Germany | GE Healthcare Buchler GmbH & Co.KG Gieselweg 1 D-38110 Braunschweig Germany | Omniscan 0,5 mmol/ml Injektioslösung | 287 mg/ml | solution for injection | intravenous use |
| DE - Germany | GE Healthcare Buchler GmbH & Co.KG Gieselweg 1 | Omniscan 0,5 mmol/ml Injektioslösung in Fertigspritzen | 287 mg/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|--|--------------|------------------------|----------------------------|
| | D-38110 Braunschweig Germany | | | | |
| DE - Germany | Guerbet 15 rue des Vanesses Zone Paris Nord II F-83420 VILLEPINTE France | Dotarem 0,5 mmol/ml Injektionslösung in Fertigspritzen | 279.32 mg/ml | solution for injection | intravenous use |
| DE - Germany | Guerbet 15 rue des Vanesses Zone Paris Nord II F-83420 VILLEPINTE France | Dotarem 0,5 mmol/ml Injektionslösung in Durchstechflaschen (für Mehrfachentnahme) | 279.32 mg/ml | solution for injection | intravenous use |
| DE - Germany | Guerbet 15 rue des Vanesses Zone Paris Nord II F-83420 VILLEPINTE France | Dotarem 0,5 mmol/ml Injektionslösung in Durchstechflaschen | 279.32 mg/ml | solution for injection | intravenous use |
| DE - Germany | Bayer Vital GmbH D-51368 Leverkusen Germany | Magnevist 0,5 mmol/ml, Injektionslösung | 469.01 mg/ml | solution for injection | intravenous use |
| DE - Germany | Bayer Vital GmbH D-51368 Leverkusen Germany | Gadopentetat- Dimeglumin 0,5 mmol/ml, Injektionslösung | 469.01 mg/ml | solution for injection | intravenous use |
| DE - Germany | be imaging GmbH DrRudolf-Eberle-Str. 8-10 D-76534 Baden-Baden Germany | Magnevision 0,5 mmol/ml Injektionslösung | 469 mg/ml | solution for injection | intravenous use |
| DE - Germany | Covidien Deutschland GmbH Gewerbepark 1 D-93333 Neustadt Germany | Marktiv 500 Mikromol/ml Injektionslösung | 469 mg/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|---|---|--------------|------------------------|-------------------------|
| DE - Germany | be imaging GmbH DrRudolf-Eberle-Str. 8-10 D-76534 Baden-Baden Germany | Magnevision b.e. 0,5 mmol/ml Injektionslösung | 469 mg/ml | solution for injection | intravenous use |
| DE - Germany | Helm AG Nordkanalstr. 28 D-20097 Hamburg Germany | Gadopentetat Dimeglumin Helm AG Injektionslösung | 469 mg/ml | solution for injection | intravenous use |
| DE - Germany | Insight Agents GmbH Ringstr. 19 B D-69115 Heidelberg Germany | Magnegita 500 Mikromol/ml Injektionslösung | 469.01 mg/ml | solution for injection | intravenous use |
| DE - Germany | Insight Agents GmbH Ringstr. 19 B D-69115 Heidelberg Germany | Gadopentetat Insight 500 Mikromol/ml Injektionslösung | 469.01 mg/ml | solution for injection | intravenous use |
| DE - Germany | Marotrast GmbH Otto-Schott-Str. 15 D-07745 Jena Germany | Magnograf 0,5 mmol/ml, Injektionslösung | 469.01 mg/ml | solution for injection | intravenous use |
| DE - Germany | ratiopharm GmbH Graf-Arco-Str. 3 D-89079 Ulm Germany | Gadopentetat-MRT- ratiopharm | 469 mg/ml | solution for injection | intravenous use |
| DE - Germany | Sanochemia Diagnostics Deutschland GmbH Stresemannallee 4 c D-41460 Neuss Germany | MR-Lux | 469 mg/ml | solution for injection | intravenous use |
| DE - Germany | Bracco IMAGING Deutschland GmbH Max-Stromexer-Str. 116 | ProHance | 279.3 mg/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|---|--------------|---|-------------------------|
| | D-78467 Konstanz Germany | | | | |
| DE - Germany | Bayer Vital GmbH D-51368 Leverkusen Germany | Primovist 0,25 mml/ml Injektionslösung, Fertigspritze | 181.43 mg/ml | solution for injection | intravenous use |
| DE - Germany | Bayer Vital GmbH D-51368 Leverkusen Germany | Primovist 0,25 mml/ml Injektionslösung, Durchstechflasche | 181.43 mg/ml | solution for injection | intravenous use |
| DK - Denmark | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | Dotarem | 279,3 mg/mL | solution for injection | intravenous use |
| DK - Denmark | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | Dotarem | 279,3 mg/mL | Solution for injection, prefilled syringe | intravenous use |
| DK - Denmark | GE Healthcare AS Nycoveien 1-2, 4220 Nydalen, Oslo N-0401 Norway | Omniscan | 0,5 mmol/mL | solution for injection | intravenous use |
| DK - Denmark | Insight Agents GmbH Ringstr. 19 B D-69115 Heidelberg Germany | Magnegita | 0,5 mmol/mL | solution for injection | intravenous use |
| DK - Denmark | Insight Agents GmbH Ringstr. 19 B D-69115 Heidelberg Germany | Gadopentetat "Insight" | 0,5 mmol/mL | solution for injection | intravenous use |
| DK - Denmark | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Gadovist | 1 mmol/mL | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|--------------|-------------|---|-------------------------|
| DK - Denmark | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Gadovist PFS | 1 mmol/mL | solution for injection, prefilled syringe | intravenous use |
| DK - Denmark | Bracco International B.V Strawinskylaan 3051 NL-1077 ZX Amsterdam Netherlands | Prohance | 279,3 mg/mL | solution for injection | intravenous use |
| DK - Denmark | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Magnevist | 469 mg/mL | solution for injection | intravenous use |
| DK - Denmark | Bracco S.p.A. Via Emilio Folli, 50 20134 Milano Italy | Multihance | 334 mg/ml | solution for injection | intravenous use |
| DK - Denmark | Bracco S.p.A. Via Emilio Folli, 50 20134 Milano Italy | Multihance | 334 mg | Solution for injection, prefilled syringe | intravenous use |
| EE - Estonia | Bayer Schering Pharma AG DE-13342 Berlin- Germany | MAGNEVIST | 469 mg/ml | solution for injection | intravenous use |
| EE - Estonia | GE Healthcare AS PO 4220, Nycoveien 1-2 NO-0401 Nydalen Norway | OMNISCAN | 0,5 mmol/ml | solution for injection | intravenous use |
| EE - Estonia | Bayer Schering Pharma AG DE-13342 Berlin- Germany | GADOVIST | 1 mmol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|--------------|--------------|--|----------------------------|
| EE - Estonia | Bayer Schering Pharma AG DE-13342 Berlin- Germany | PRIMOVIST | 0.25 mmol/ml | solution for injection | intravenous use |
| EE - Estonia | Bayer Schering Pharma AG DE-13342 Berlin- Germany | PRIMOVIST | 0.25 mmol/ml | solution for injection in pre-filled syringe | intravenous use |
| EE - Estonia | Insight Agents GmbH Ringstr. 19 B D-69115 Heidelberg Germany | MAGNEGITA | 469 mg/ml | solution for injection | intravenous use |
| ES - Spain | GE HEALTHCARE BIO-SCIENCES S.A. Avda. de Europa 22 Alcobendas 28108 Madrid Spain | OMNISCAN | 0,5 mmol/ml | solution for injection syringe | intravenous use |
| ES - Spain | GE HEALTHCARE BIO-SCIENCES S.A. Avda. de Europa 22 Alcobendas 28108 Madrid Spain | OMNISCAN | 0,5 mmol/ml | solution for injection | intravenous use |
| ES - Spain | Bracco S.p.A. Via Egido Folli, 50 20134 Milano Italy | MULTIHANCE | 0,5 M | solution for injection | intravenous use |
| ES - Spain | QUIMICA FARMACEUTICA BAYER S.L. Av. Baix Llobregat 3-5 Sant Joan Despi Barcelona 08970 Spain | PRIMOVIST | 0,25 mmol/ml | solution for injection | intravenous use |
| ES - Spain | QUIMICA FARMACEUTICA BAYER S.L. Av. Baix Llobregat 3-5 Sant Joan Despi | PRIMOVIST | 0,25 mmol/ml | solution for injection (syringe) | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|---|--------------|-------------|--|----------------------------|
| | Barcelona 08970 Spain | | | | |
| ES - Spain | Bracco International B.V Strawinskylaan 3051 NL-1077 ZX Amsterdam Netherlands | PROHANCE | (0.5 M) | solution for injection (syringe) | intravenous use |
| ES - Spain | Bracco International B.V Strawinskylaan 3051 NL-1077 ZX Amsterdam Netherlands | PROHANCE | (0.5 M) | solution for injection (vial) | intravenous use |
| ES - Spain | QUIMICA FARMACEUTICA BAYER S.L. Av. Baix Llobregat 3-5 Sant Joan Despi Barcelona 08970 Spain | GADOVIST | 1,0 mmol/ml | solution for injection (vial) | intravenous use |
| ES - Spain | QUIMICA FARMACEUTICA BAYER S.L. Av. Baix Llobregat 3-5 Sant Joan Despi Barcelona 08970 Spain | GADOVIST | 1,0 mmol/ml | solution for injection (syringe and cartridge) | intravenous use |
| ES - Spain | QUIMICA FARMACEUTICA BAYER S.L. Av. Baix Llobregat 3-5 Sant Joan Despi Barcelona 08970 Spain | MAGNEVIST | 0,5 mmol/ml | solution for injection | intravenous use |
| ES - Spain | QUIMICA FARMACEUTICA BAYER S.L. Av. Baix Llobregat 3-5 Sant Joan Despi | MAGNEVIST | 0,5 mmol/ml | solution for injection (syringe) | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|--------------|-------------|----------------------------------|----------------------------|
| | Barcelona 08970 Spain | | | | |
| ES - Spain | QUIMICA FARMACEUTICA BAYER S.L. Av. Baix Llobregat 3-5 Sant Joan Despi Barcelona 08970 Spain | MAGNOGRAF | 0,5 mmol/ml | solution for injection | intravenous use |
| ES - Spain | QUIMICA FARMACEUTICA BAYER S.L. Av. Baix Llobregat 3-5 Sant Joan Despi Barcelona 08970 Spain | MAGNOGRAF | 0,5 mmol/ml | solution for injection (syringe) | intravenous use |
| ES - Spain | Guerbet BP 57400 95493 Roissy CdG cedex France | Dotarem | 0,5 mmol/ml | solution for injection | intravenous use |
| ES - Spain | Guerbet BP 57400 95493 Roissy CdG cedex France | Dotarem | 0,5 mmol/ml | solution for injection (syringe) | intravenous use |
| FI - Finland | GE Healthcare AS Nycoveien 1-2 4220 Nydalen, Oslo N-0401 Norway | OMNISCAN | 0.5 mmol/ml | Solution for injection | intravenous use |
| FI - Finland | Bayer Schering Pharma Oy Pansiontie 47 20210 Turku Finland | MAGNEVIST | 0.5 mmol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|---|--------------------------|-----------------|--|-------------------------|
| FI - Finland | Insight Agents GmbH Ringstr. 19 B, 69115 HEIDELBERG Germany | MAGNEGITA | 500 micromol/ml | solution for injection | intravenous use |
| FI - Finland | Insight Agents GmbH Ringstr. 19 B, 69115 HEIDELBERG Germany | GADOPENTETATE INSIGHT | 500 micromol/ml | solution for injection | intravenous use |
| FI - Finland | Bracco S.p.A. Via Emilio Folli, 50 20134 Milano Italy | MULTIHANCE | 334 mg/ml | solution for injection | intravenous use |
| FI - Finland | Bracco S.p.A. Via Emilio Folli, 50 20134 Milano Italy | MULTIHANCE | 334 mg/ml | Solution for injection, pre-filled syringe | intravenous use |
| FI - Finland | Bayer Schering Pharma Oy Pansiontie 47, 20210 Turku Finland | PRIMOVIST | 0.25 mmol/ml | solution for injection | intravenous use |
| FI - Finland | Bayer Schering Pharma Oy Pansiontie 47, 20210 Turku Finland | PRIMOVIST | 0.25 mmol/ml | Solution for injection, pre-filled syringe | intravenous use |
| FI - Finland | Bracco International B.V Strawinskylaan 3051, NL-1077 ZX Amsterdam Netherlands | PROHANCE | 279.3 mg/ml | solution for injection | intravenous use |
| FI - Finland | Bayer Schering Pharma Oy Pansiontie 47, 20210 Turku Finland | GADOVIST | 1 mmol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|---|--|-------------------|---|----------------------------|
| FI - Finland | Bayer Schering Pharma Oy Pansiontie 47, 20210 Turku Finland | GADOVIST | 1 mmol/ml | Solution for injection (syringe) | intravenous use |
| FI - Finland | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | DOTAREM | 279,3 mg/ml | solution for injection | intravenous use |
| FR - France | GE HEALTHCARE 11, avenue Morane Saulnier 78140 Vélizy Villacoublay France | OMNISCAN 0,5 mmol/ml, solution injectable | 28,7 g / 100 ml | solution for injection | intravenous use |
| FR - France | GE HEALTHCARE 11, avenue Morane Saulnier 78140 Vélizy Villacoublay France | OMNISCAN 0,5 mmol/ml, solution injectable en seringue pré-remplie | 287 mg / 1 ml | Solution for injection (pre-filled syringe) | intravenous use |
| FR - France | BRACCO IMAGING France 7, place Copernic 91080 Courcouronnes France | MULTIHANCE 0,5 mmol/ml, solution injectable (IV) | 529 mg / 1 ml | solution for injection | intravenous use |
| FR - France | BRACCO IMAGING France 7, place Copernic 91080 Courcouronnes France | MULTIHANCE 0,5 mmol/ml, solution injectable en seringue pré-remplie | 529 mg / 1 ml | solution for injection | intravenous use |
| FR - France | BAYER SANTE 220, avenue de la Recherche 59120 Loos France | MAGNEVIST, solution injectable (IV) | 46,901 g / 100 ml | solution for injection | intravenous use |
| FR - France | BAYER SANTE 220, avenue de la Recherche 59120 Loos France | MAGNEVIST, solution injectable en seringue pré-remplie (IV) | 46,901 g / 100 ml | Solution for injection (pre-filled syringe) | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|--|--------------------|---|----------------------------|
| FR - France | BRACCO IMAGING France 7, place Copernic 91080 Courcouronnes France | PROHANCE 1396,5 mg/5 ml, solution injectable | 1396,50 mg / 5 ml | solution for injection | intravenous use |
| FR - France | BRACCO IMAGING France 7, place Copernic 91080 Courcouronnes France | PROHANCE 1396,5 mg/5 ml, solution injectable en seringue pré-remplie | 1396,5 mg / 5ml | Solution for injection (pre-filled syringe) | intravenous use |
| FR - France | BRACCO IMAGING France 7, place Copernic 91080 Courcouronnes France | PROHANCE 2793 mg/10 ml, solution injectable | 2793 mg / 10 ml | solution for injection | intravenous use |
| FR - France | BRACCO IMAGING France 7, place Copernic 91080 Courcouronnes France | PROHANCE 2793 mg/10 ml, solution injectable en seringue pré-remplie | 2793 mg / 10 ml | Solution for injection (pre-filled syringe) | intravenous use |
| FR - France | BRACCO IMAGING France 7, place Copernic 91080 Courcouronnes France | PROHANCE 4189,5 mg/15 ml, solution injectable | 4189,50 mg / 15 ml | solution for injection | intravenous use |
| FR - France | BRACCO IMAGING France 7, place Copernic 91080 Courcouronnes France | PROHANCE 4189,5 mg/15 ml, solution injectable en seringue pré-remplie | 4189,50 mg / 15 ml | Solution for injection (pre-filled syringe) | intravenous use |
| FR - France | PRACCO IMAGING France 7, place Copernic 91080 Courcouronnes France | PROHANCE 4748,10 mg/17 ml, solution injectable en seringue pré-remplie | 4748,1 mg / 17 ml | Solution for injection (pre-filled syringe) | intravenous use |
| FR - France | BRACCO IMAGING France 7, place Copernic 91080 Courcouronnes France | PROHANCE 5586 mg/20 ml, solution injectable | 5586 mg / 20 ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|---|-------------------|---|----------------------------|
| FR - France | BAYER SANTE 220, avenue de la Recherche 59120 Loos France | GADOVIST 1,0 mmol/mL, solution injectable | 604,72 mg / 1 ml | solution for injection | intravenous use |
| FR - France | BAYER SANTE 220, avenue de la Recherche 59120 Loos France | GADOVIST 1,0 mmol/mL, solution injectable en seringue préremplie | 604,72 mg / 1 ml | Solution for injection (pre-filled syringe) | intravenous use |
| FR - France | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | DOTAREM 0,5 mmol/ml, solution injectable en flacon | 27,932 g / 100 ml | solution for injection | intravenous use |
| FR - France | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | DOTAREM 0,5 mmol/ml, solution injectable en seringue pré-remplie | 27,932 g / 100 ml | Solution for injection (pre-filled syringe) | intravenous use |
| GR - Greece | GE HEALTHCARE PLAPOUTA 139 & LAMIAS ST NEO IRAKLEIO 14121 GREECE | OMNISCAN | 287mg/ml | solution for injection | intravenous use |
| GR - Greece | BAYER HELLAS ABEE SOROU 18-20, 151 25 MAROUSI, ATHENS, GREECE | MAGNEVIST | 469.01mg/ml | solution for injection | intravenous use |
| GR - Greece | GEROLYMATOS P.G.N. AEBE ASKLIPIOU ST KRYONERI ATTICA GREECE | MULTIHANCE | 529mg/mlm | solution for injection | intravenous use |
| GR - Greece | GEROLYMATOS P.G.N. AEBE ASKLIPIOU ST KRYONERI ATTICA GREECE | MULTIHANCE | 529mg/mlm | Solution for injection prefilled syringe | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|---------------------------------------|--------------|--|-------------------------|
| GR - Greece | BAYER HELLAS ABEE SOROU 18-20, 151 25 MAROUSI, ATHENS, GREECE | PRIMOVIST | 0.25 mmol/ml | Solution for injection | intravenous use |
| GR - Greece | BAYER HELLAS ABEE SOROU 18-20, 151 25 MAROUSI, ATHENS, GREECE | PRIMOVIST "PFS" | 0.25 mmol/ml | Solution for injection prefilled syringe | intravenous use |
| GR-Greece | BAYER HELLAS ABEE SOROU 18-20, 151 25 MAROUSI, ATHENS, GREECE | GADOVIST | 1 mmol/ml | solution for injection | intravenous use |
| GR-Greece | BAYER HELLAS ABEE SOROU 18-20, 151 25 MAROUSI, ATHENS, GREECE | GADOVIST PFS | 1 mmol/ml | Solution for injection prefilled syringe | intravenous use |
| GR-Greece | Hospital Line SA K. Palama 36 GR-143 43, N. Chalkidona, Athens Greece | Dotarem | 1,4 mg/ml | solution for injection | intravenous use |
| GR-Greece | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | VASOVIST | 0.25mmol/ml | solution for injection | intravenous use |
| HU - Hungary | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | DOTAREM 0,5 mmol/ml injection 15ml | 0,5 mmol/ml | injection | intravenous use |
| HU - Hungary | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | DOTAREM 0,5 mmol/ml injection 20ml | 0,5 mmol/ml | injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|---|-------------|---|-------------------------|
| HU - Hungary | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | DOTAREM 0,5 mmol/ml injection 60ml | 0,5 mmol/ml | injection | intravenous use |
| HU - Hungary | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | DOTAREM 0,5 mmol/ml injection 100ml | 0,5 mmol/ml | injection | intravenous use |
| HU - Hungary | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | GADOVIST 1,0 mmol/ml solution for injection | 1 mmol/ml | solution for injection | intravenous use |
| HU - Hungary | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | GADOVIST | 1 mmol/ml | solution for injection, prefilled syringe | intravenous use |
| HU - Hungary | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | MAGNEVIST | 0,5 mmol/ml | solution for injection | intravenous use |
| HU - Hungary | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | MAGNEVIST | 0,5 mmol/ml | injection in a pre- filled syringe | intravenous use |
| HU - Hungary | Bracco S.p.A. Via Emilio Folli, 50 20134 Milano Italy | MULTIHANCE | 0,5M | solution for injection | intravenous use |
| HU - Hungary | GE Healthcare AS Nycoveien 1-2 4220 Nydalen, Oslo N-0401 | OMNISCAN 0,5 mmol/ml injection | 0,5 mmol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|---|--------------|--|-------------------------|
| | Norway | | | | |
| HU - Hungary | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | PRIMOVIST | 0,25 mmol/ml | solution for injection | intravenous use |
| HU - Hungary | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | PRIMOVIST | 0,25 mmol/ml | solution for injection, pre-filled syringe | intravenous use |
| IE - Ireland | GE Healthcare AS Nycoveien 1-2 4220 Nydalen, Oslo N-0401 Norway | Omniscan 0.5mmol/ml solution for injection, glass vial/bottle | 0.5 mmol/ml | solution for injection | intravenous use |
| IE - Ireland | GE Healthcare AS Nycoveien 1-2 4220 Nydalen, Oslo N-0401 Norway | Omniscan 0.5mmol/ml solution for injection, polypropylene bottles | 0.5 mmol/ml | solution for injection | intravenous use |
| IE - Ireland | GE Healthcare AS Nycoveien 1-2 4220 Nydalen, Oslo N-0401 Norway | Omniscan 0.5mmol/ml solution for injection, prefilled syringe | 0.5 mmol/ml | solution for injection, prefilled syringe | intravenous use |
| IE - Ireland | Bayer Limited The Atrium Blackthorn Road Dublin 18 Ireland | Primovist 0.25 mmol/ml Sol for Injection | 0.25 mmol/ml | solution for injection | intravenous use |
| IE - Ireland | Bayer Limited The Atrium Blackthorn Road | Primovist 0.25 mmol/ml Sol for Injection, prefilled syringe | 0.25 mmol/ml | solution for injection, prefilled syringe | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--------------------------------|---------------------------|-------------|-------------------------|-------------------------|
| | Dublin 18 Ireland | | | | |
| | Bayer Limited | Gadovist 1.0 mmol/ml | 1.0 mmol/ml | solution for injection | intravenous use |
| | The Atrium | Solution for Injection | | | |
| IE - Ireland | Blackthorn Road | J | | | |
| | Dublin 18 | | | | |
| | Ireland | | | | |
| | Bayer Limited | Gadovist 1.0 mmol/ml | 1.0 mmol/ml | solution for injection, | intravenous use |
| | The Atrium | Solution for Injection in | | in prefilled syringe | |
| IE - Ireland | Blackthorn Road | prefilled syringe | | | |
| | Dublin 18 | | | | |
| | Ireland | | | | |
| | Bayer Limited | Gadovist 1.0 mmol/ml | 1.0 mmol/ml | solution for injection, | intravenous use |
| | The Atrium | Solution for Injection in | | in prefilled cartridge | |
| IE - Ireland | Blackthorn Road | prefilled cartridge | | | |
| | Dublin 18 | | | | |
| | Ireland | | | | |
| | Bracco International B.V | Prohance 279.3 mg/ml | 279.3 mg/ml | solution for injection | intravenous use |
| IE - Ireland | Strawinskylaan 3051 | Solution for Injection, 5 | | | |
| TE - Heland | NL-1077 ZX Amsterdam | ml vial | | | |
| | Netherlands | | | | |
| | Bracco International B.V | Prohance 279.3 mg/ml | 279.3 mg/ml | solution for injection | intravenous use |
| IE - Ireland | Strawinskylaan 3051 | Solution for Injection, | | | |
| 1E - Heland | NL-1077 ZX Amsterdam | 10 ml vial | | | |
| | Netherlands | | | | |
| | Bracco International B.V | Prohance 279.3 mg/ml | 279.3 mg/ml | solution for injection | intravenous use |
| IE - Ireland | Strawinskylaan 3051 | Solution for Injection, | | | |
| 1E - Heland | NL-1077 ZX Amsterdam | 15 ml vial | | | |
| | Netherlands | | | | |
| | Bracco International B.V | Prohance 279.3 mg/ml | 279.3 mg/ml | solution for injection | intravenous use |
| IE - Ireland | Strawinskylaan 3051 | Solution for Injection, | | | |
| | NL-1077 ZX Amsterdam | 20 ml vial | | | |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|----------------|--------------------------------------|---------------------------|--------------|------------------------|-------------------------|
| | Netherlands | | | | |
| | GUERBET | Dotarem 279.32mg/ml | 279.32 mg/ml | solution for injection | intravenous use |
| IE - Ireland | BP 57400 | Solution for Injection, | | | |
| IL - II cianu | 95943 Roissy Charles de Gaulle Cedex | glass pre-filled syringes | | | |
| | France | | | | |
| | GUERBET | Dotarem 279.32mg/ml | 279.32 mg/ml | solution for injection | intravenous use |
| IE - Ireland | BP 57400 | Solution for Injection, | | | |
| IL - IICiana | 95943 Roissy Charles de Gaulle Cedex | glass vials | | | |
| | France | | | | |
| | Bracco SpA | Multihance 0.5 M | 529 mg/ml | solution for injection | intravenous use |
| IE - Ireland | Via Egidio Folli 50 | solution for injection | | | |
| IL - II cland | I-20134 Milan | | | | |
| | Italy | | | | |
| | Bracco SpA | Multihance 529 mg/ml | 529 mg/ml | solution for injection | intravenous use |
| IE - Ireland | Via Egidio Folli 50 | solution for injection in | | in prefilled syringe | |
| IL - Il cland | I-20134 Milan | prefiled syringe | | | |
| | Italy | | | | |
| | Insights Agents GmbH | Magnegita 500 | 500 mmol/ml | solution for injection | intravenous use |
| IE - Ireland | Ringstrasse 19B | micromol/ml solution | | | |
| IL - Il claird | D-69115 Heidelberg | for injection | | | |
| | Germany | | | | |
| | Bayer Limited | Magnevist 0.5 mmol/ml | 0.5 mmol/ml | solution for injection | intravenous use |
| | The Atrium | Solution for Injection | | | |
| IE - Ireland | Blackthorn Road | | | | |
| | Dublin 18 | | | | |
| | Ireland | | | | |
| | Bayer Limited | Magnevist 0.5mmol/ml | 0.5 mmol/ml | solution for Injection | intravenous use |
| | The Atrium | Solution for Injection in | | in pre-filled syringe | |
| IE - Ireland | Blackthorn Road | pre-filled syringe. | | | |
| | Dublin 18 | | | | |
| | Ireland | | | | |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|---|--------------|-------------------------|---|-------------------------|
| IS - Iceland | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Magnevist | 469 mg/ml | solution for injection | intravenous use |
| IS - Iceland | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Magnevist | 469 mg/ml | solution for injection, prefilled syringe | intravenous use |
| IS - Iceland | GE Healthcare AS Nycoveien 1-2 4220 Nydalen, Oslo N-0401 Norway | Omniscan | 0,5 mmol/ml | solution for injection | intravenous use |
| IS – Iceland | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Gadovist | 1,0 mmol/ml | Solution for injection | Intravenous use |
| IT - Italy | Bayer Schering Pharma AG Müllerstrasse 170-178, DE-13342 Berlin-Wedding Germany | Magnevist | 469 mg/ml | solution for injection | intravenous use |
| IT - Italy | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | Dotarem | 0,5 mmol/ml | solution for injection | intravenous use |
| IT - Italy | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | Dotarem | 0,0025 mmol/ml | solution for injection | intravenous use |
| IT - Italy | GE Healthcare Via Galeno 36, 20126 Milano | Omniscan | 287 mg/ml (0,5 mmol/ml) | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|----------------|---|--------------|---------------------|---|-------------------------|
| | Italy | | | | |
| IT - Italy | Bracco Imaging Italia Via Egidio Folli, 50 20134 Milano Italy | ProHance | 279,3 mg/ml (0.5 M) | solution for infusion | intravenous use |
| | <i>y</i> | M-14:TT | 224 /1 (0.5 M) | 14: 6:4: | :4 |
| IT - Italy | Bracco Imaging Italia Via Egidio Folli, 50 20134 Milano Italy | MultiHance | 334 mg/ml (0,5 M) | solution for injection | intravenous use |
| IT - Italy | Bayer SpA Viale Certosa, 130 20156 Milano Italy | Gadovist | 604.72 mg/ml | solution for injection | intravenous use |
| IT - Italy | Bayer SpA Viale Certosa, 130 20156 Milano Italy | Primovist | 0,25 mmol/ml | solution for injection | intravenous use |
| LT - Lithuania | Bayer Schering Pharma AG Müllerstrasse 178 DE-13353 Berlin Germany | Gadovist | 1 mmol/ml | solution for injection (pre- filled syringe) | intravenous use |
| LT - Lithuania | Bayer Schering Pharma AG Müllerstrasse 178 DE-13353 Berlin Germany | Primovist | 0,25 mmol/ml | solution for injection | intravenous use |
| LT - Lithuania | Bayer Schering Pharma AG Müllerstrasse 178 DE-13353 Berlin Germany | Primovist | 0,25 mmol/ml | solution for injection (pre-filled syringe) | intravenous use |
| LT - Lithuania | Bayer Schering Pharma AG Müllerstrasse 178 DE-13353 Berlin | Magnevist | 0,5 mmol/ml | solution for injection/infusion | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|-----------------|--------------------------------|----------------|-----------------|------------------------|-------------------------|
| | Germany | | | | |
| | Insight Agents GmbH | Magnegita | 500 micromol/ml | solution for injection | intravenous use |
| LT - Lithuania | Ringstr. 19 B | | | | |
| LT - Litiluania | D-69115 Heidelberg | | | | |
| | Germany | | | | |
| | GE Healthcare AS | Omniscan | 0,5 mmol/ml | solution for injection | intravenous use |
| | Nycoveien 1-2 | | | | |
| LT - Lithuania | P.O. Box 4220 Nydalen, | | | | |
| | Oslo N-0401 | | | | |
| | Norway | | | | |
| | Codali S.A. | DOTAREM | 37,7G | solution for injection | intravenous use |
| LU - | Avenue Henri Dunant 31 | | | | |
| Luxembourg | 1140 Bruxelles | | | | |
| | Belgium | | | | |
| | Insight Agents GmbH | GADOPENTETATE | 469MG/ML | solution for injection | intravenous use |
| LU - | Ringstr. 19 B | INSIGHT | | | |
| Luxembourg | D-69115 Heidelberg | | | | |
| | Germany | | | | |
| | Bayer SA-NV | GADOVIST 1 | 604,72mg | solution for injection | intravenous use |
| LU - | J.E. Mommaertslaan 14 | | | | |
| Luxembourg | 1831 Diegem | | | | |
| | Belgium | | | | |
| | Bayer SA-NV | GADOVIST PFS-1 | 604,72mg /1ml | solution for injection | intravenous use |
| LU - | J.E. Mommaertslaan 14 | | | | |
| Luxembourg | 1831 Diegem | | | | |
| | Belgium | | | | |
| | Insight Agents GmbH | MAGNEGITA | 78,63 MG/1ML | solution for injection | intravenous use |
| LU - | Ringstr. 19 B | | | | |
| Luxembourg | D-69115 Heidelberg | | | | |
| | Germany | | | | |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------------|---|--|-----------------|------------------------|-------------------------|
| LU - Luxembourg | Bayer SA-NV J.E. Mommaertslaan 14 1831 Diegem Belgium | MAGNEVIST | 4,69G/10 ML | solution for injection | intravenous use |
| LU - Luxembourg | Bracco IMAGING Deutschland GmbH Max-Stromeyer-Str. 116 D-78467 Konstanz Germany | MULTIHANCE | 529mg/1 ml | solution for injection | intravenous use |
| LU - Luxembourg | GE HEALTHCARE BVBA Kouterveldstraat 20 1831 DIEGEM Belgium | OMNISCAN | 287MG/1 ML | solution for injection | intravenous use |
| LU - Luxembourg | Bracco IMAGING Deutschland GmbH Max-Stromeyer-Str. 116 D-78467 Konstanz Germany | PROHANCE | 279,3MG/1 ML | solution for injection | intravenous use |
| LV - Latvia | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Gadovist 1,0 mmol/ml solution for injections | 1,0 mmol/ml | solution for injection | intravenous use |
| LV - Latvia | Insight Agents GmbH Ringstr. 19 B D-69115 Heidelberg Germany | Magnegita 500 micromol/ml solution for injection | 500 micromol/ml | solution for injection | intravenous use |
| LV - Latvia | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Magnevist 0,5 mmol/ml solution for injection | 0,5 mmol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|---------------------|--|---|--|--|-------------------------|
| LV - Latvia | GE Healthcare AS Nycoveien 1-2, 4220 Nydalen, Oslo N-0401 Norway | Omniscan 0,5 mmmol/ml solution for injection | 0,5 mmol/ml | solution for injection | intravenous use |
| LV - Latvia | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Primovist 0,25 mmol/ ml solution for injection | 0,25 mmol/ml | solution for injection | intravenous use |
| LV - Latvia | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Primovist 0,25 mmol/ ml solution for injection pre-filled syringe | 0,25 mmol/ml | Solution for injection, pre-filled syringe | intravenous use |
| MT - Malta | GE Healthcare AS Nycoveien 1-2, 4220 Nydalen, Oslo N-0401 Norway | Omniscan | 0.5mMol/ml (287 mg equiv. 0.5 mmol) | solution for injection | intravenous use |
| MT - Malta | Bayer plc Bayer House, Strawberry Hill, Newbury, Berkshire, RG14 1JA UK | Magnevist | 469.01 mg | Solution for injection | intravenous use |
| MT - Malta | Bayer plc Bayer House, Strawberry Hill, Newbury, Berkshire, RG14 1JA UK | Primovist | PFS 0.25 mmol/ml | Solution for injection, prefilled syringe | intravenous use |
| NL - Netherlands | Guerbet Nederland B.V. Avelingen-West 3A 4202 MS GORINCHEM Netherlands | Dotarem | 0,5 mmol/ml | solution for injection | intravenous use |
| NL - Netherlands | Bayer B.V. Energieweg 1 3641 RT MIJDRECHT Netherlands | Gadovisit | 1,0 mmol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|---------------------|---|-----------------------|--------------|------------------------|-------------------------|
| NL - Netherlands | Bayer B.V. Energieweg 1 3641 RT MIJDRECHT Netherlands | Magnevist | 0,5 mmol/ml | solution for injection | intravenous use |
| NL - Netherlands | Bracco IMAGING Deutschland GmbH Max-Stromeyer-Str. 116 D-78467 Konstanz Germany | Multihance | 0,5 mmol/ml | solution for injection | intravenous use |
| NL - Netherlands | GE Healthcare B.V. Cygne Centre De Rondom 8 5612 AP EINDHOVEN Netherlands | Omniscan | 0,5 mmol/ml | solution for injection | intravenous use |
| NL - Netherlands | Bayer B.V. Energieweg 1 3641 RT MIJDRECHT Netherlands | Primovist | 0,25 mmol/ml | solution for injection | intravenous use |
| NL - Netherlands | Bracco IMAGING Deutschland GmbH Max-Stromeyer-Str. 116 D-78467 Konstanz Germany | Prohance | 0,5 mmol/ml | solution for injection | intravenous use |
| NL - Netherlands | Insight Agents GmbH Ringstr. 19 B D-69115 Heidelberg Germany | Magnegita | 0,5 mmol/ml | solution for injection | intravenous use |
| NL - Netherlands | Insight Agents GmbH Ringstr. 19 B D-69115 Heidelberg Germany | Gadopentetate Insight | 0,5 mmol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|--------------|--------------|------------------------|----------------------------|
| NO - Norway | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | Dotarem | 279,3 mg/ml | solution for injection | intravenous use |
| NO - Norway | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Magnevist | 469 mg/ml | solution for injection | intravenous use |
| NO - Norway | Bracco S.p.A. Via Emilio Folli, 50 20134 Milano Italy | MultiHance | 334 mg/ml | solution for injection | intravenous use |
| NO - Norway | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Primovist | 0,25 mmol/ml | solution for injection | intravenous use |
| NO - Norway | Bracco International B.V Strawinskylaan 3051 NL-1077 ZX Amsterdam Netherlands | Prohance | 279,3 mg/ml | solution for injection | intravenous use |
| NO - Norway | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Gadovist | 1 mmol/ml | solution for injection | intravenous use |
| NO - Norway | GE Healthcare AS Nycoveien 1-2 4220 Nydalen, Oslo N-0401 Norway | Omniscan | 0,5 mmol/ml | solution for injection | intravenous use |
| PL - Poland | GE Healthcare AS Nycoveien 1-2 P.O. Box 4220 Nydalen, | Omniscan | 0,5 mmol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|---------------|---|--------------|--------------|---|-------------------------|
| | Oslo NO-0401 Norway | | | | |
| PL - Poland | Bracco Imaging Deutschland GmbH Max-Stromeyer-Str. 116 D-78467 Konstanz Germany | ProHance | 279,3 mg/ml | solution for injection | intravenous use |
| PL - Poland | Bayer Schering Pharma AG D-13342 Berlin Germany | Gadovist 1,0 | 604,72 mg/ml | solution for injection | intravenous use |
| PL - Poland | Bayer Schering Pharma AG D-13342 Berlin Germany | Magnevist | 469 mg/ml | solution for injection | intravenous use |
| PL - Poland | Bracco ALTANA Pharma GmbH Max-Stromeyer-Str. 116 78467 Konstanz Germany | Multihance | 529 mg/ml | solution for injection | intravenous use |
| PL - Poland | Bayer Schering Pharma AG D-13342 Berlin Germany | Primovist | 0,25 mmol/ml | solution for injection | intravenous use |
| PT - Portugal | Bayer Portugal S.A. Rua Quinta do Pinheiro, 5 2794-003 Carnaxide Portugal | Primovist | 0.25 mmol/ml | solution for injection | intravenous use |
| PT - Portugal | Bayer Portugal S.A. Rua Quinta Pinheiro, 5, 2794-003 Carnaxide Portugal | Primovist | 0.25 mmol/ml | solution for injection (pre-filled syringe) | intravenous use |
| PT - Portugal | Lusal - Produção Químico- Farmacêutica Luso-Alemã Lda. Rua Quinta Pinheiro, 5, Outurela 2794- 003 Carnaxide | Gadovist | 1 mmol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|---------------|---|--|-----------------------------------|---|-------------------------|
| | Portugal | | | | |
| DT Destroyal | Lusal - Produção Químico- Farmacêutica Luso-Alemã Lda. | Gadovist | 1 mmol/ml | solution for injection (pre-filled syringe) | intravenous use |
| PT - Portugal | Rua Quinta Pinheiro, 5, Outurela 2794- 003 Carnaxide Portugal | | | | |
| PT - Portugal | A. Martins & Fernandes S.A. Rua Raúl Mesnier du Ponsard, 4 B 1750-243 Lisboa Portugal | Dotarem | 377 mg/ml | solution for injection | intravenous use |
| PT - Portugal | Satis-Radioisótopos e Protecções Contra Sobretensões Eléctricas Unipessoal Lda. Edificio Ramazzotti, Av. do Forte, n.º 6 - 6A, 2790-502 Carnaxide Portugal | Omniscan | 287 mg/ml | solution for injection | intravenous use |
| RO - Romania | Bracco S.p.A. Via Egidio Folli, 50 20134 Milano Italy | MULTIHANCE 0,5M | 0.529 g (0.334 g +0.195g)/ml | solution for injection | intravenous use |
| RO - Romania | Bayer Schering Pharma AG Müllerstrasse 178 DE-13342 Berlin- Germany | MAGNEVIST | 469,01 mg/ml | solution for injection | intravenous use |
| RO-Romania | INSIGHT AGENTS GmbH Ringstrasse. 19B 69115 Heidelberg Germany | MAGNEGITA 500 micromol/ml, soluție injectabilă | 500 micromol/ml (469,01 mg/ml) | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--|--|---------------|--|----------------------------|
| RO - Romania | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | DOTAREM 0,5 mmol/ml soluţie injectabilă în seringă preumplută | 27.932 g/ml | solution for injection, pre-filled syringe | intravenous use |
| RO - Romania | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | DOTAREM 0,5 mmol/ml soluție injectabilă unidoză | 27.932 g/ml | solution for injection | intravenous use |
| RO - Romania | GUERBET BP 57400 95943 Roissy Charles de Gaulle Cedex France | DOTAREM 0,5 mmol/ml soluție injectabilă multidoză | 27.932 g/ml | solution for injection | intravenous use |
| RO - Romania | Bayer Schering Pharma AG Müllerstrasse 178 DE-13342 Berlin- Germany | PRIMOVIST 0,25 mmol/ml, soluție injectabilă în seringă preumplută | 181,430 mg/ml | solution for injection, pre-filled syringe | intravenous use |
| RO - Romania | Bayer Schering Pharma AG Müllerstrasse 178 DE-13342 Berlin- Germany | GADOVIST 1,0 mmol/ml,soluție injectabilă | 604,720 mg/ml | solution for injection | intravenous use |
| RO - Romania | GE Healthcare AS Nycoveien 1-2 P.O. Box 4220 Nydalen, N-0401 Oslo Norway | OMNISCAN, soluție injectabilă | 287,000 mg/ml | solution for injection | intravenous use |
| SE - Sweden | Bayer Schering Pharma AG Müllerstrasse 178 DE-13342 Berlin-Wedding Germany | Magnevist | 0,5 mmol/m | solution for injection | intravenous use |
| SE - Sweden | Bracco SpA Via Egidio Folli 50 I-20134 Milan | Multihance | 334 mg/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--------------------------------------|--------------|-------------------|-------------------------|-------------------------|
| | Italy | | | | |
| | Bracco SpA | Multihance | 334 mg/ml | solution for injection, | intravenous use |
| SE - Sweden | Via Egidio. Folli 50 | | | pre-filled syringe | |
| SE Sweden | I-20134 Milan | | | | |
| | Italy | | | | |
| | Bracco International BV | Prohance | 279,3 mg/ml | solution for injection | intravenous use |
| SE - Sweden | Stravinskylaan 3051 | | | | |
| Z_ Z C Z | NL-1077 ZX Amsterdam | | | | |
| | The Netherlands | | | | |
| | GE Healthcare AS | Omniscan | 0,5 mmol/ml | solution for injection | intravenous use |
| GE G 1 | P.O.Box 4220 | | | | |
| SE - Sweden | Nydalen | | | | |
| | N-0401 Oslo | | | | |
| | Norway GE Healthcare AS | Omniscan | 0,5 mmol/ml | solution for injection, | introven our use |
| | P.O.Box 4220 | Ommscan | 0,3 1111101/1111 | pre-filled syringe | intravenous use |
| SE - Sweden | Nydalen | | | pre-filled syringe | |
| SE - Sweden | N-0401 Oslo | | | | |
| | Norway | | | | |
| | Bayer Schering Pharma AG, | Primovist | 0,25 mmol/ml | solution for injection | intravenous use |
| | Müllerstrasse 178 | Timovist | 0,23 1111101/1111 | solution for injection | mitavenous use |
| SE - Sweden | DE-133 42 Berlin | | | | |
| | Germany | | | | |
| | Bayer Schering Pharma AG, | Primovist | 0,25 mmol/ml | solution for injection, | intravenous use |
| SE - Sweden | Müllerstrasse 178 | | | pre-filled syringe | |
| SE - Sweden | DE-133 42 Berlin | | | | |
| | Germany | | | | |
| | GUERBET | Dotarem | 279,3 mg/ml | solution for injection | intravenous use |
| SE - Sweden | BP 57400 | | | | |
| SL - Sweden | 95943 Roissy Charles de Gaulle Cedex | | | | |
| | France | | | | |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|---------------|---|--|-----------------|--|----------------------------|
| SE - Sweden | Bayer Schering Pharma AG Müllerstrasse 178, DE-13342 Berlin-Wedding Germany | Gadovist | 1,0 mmol/ml | solution for injection | intravenous use |
| SE - Sweden | Bayer Schering Pharma AG Müllerstrasse 178, DE-13342 Berlin-Wedding Germany | Gadovist | 1,0 mmol/ml | solution for injection, pre-filled syringe | intravenous use |
| SE - Sweden | Insight Agents GmbH Ringstr.19 B D-69115 Heidelberg Germany | Magnegita | 500 mikromol/ml | solution for injection | intravenous use |
| SE - Sweden | Insight Agents GmbH Ringstr.19 B D-69115 Heidelberg Germany | Gadopentetsyrad imegluminat Insight | 500 mikromol/ml | solution for injection | intravenous use |
| SE - Sweden | Bayer Schering Pharma AG Müllerstrasse 178 DE-13342 Berlin-Wedding Germany | Magnograf | 0,5 mmol/ml | solution for injection | intravenous use |
| SE - Sweden | Bayer Schering Pharma AG Müllerstrasse 178 DE-13342 Berlin-Wedding Germany | Magnograf | 0,5 mmol/ml | solution for injection pre-filled syringe | intravenous use |
| SI - Slovenia | Higieia d.o.o. Zastopstva in trgovina, Blatnica 10, 1236 Trzin, Slovenia | Omniscan 0,5 mmol/ml raztopina za injiciranje | 0,5 mmol/ml | solution for injection | intravenous use |
| SI - Slovenia | Insight Agents GmbH Ringstr. 19 B D-69115 Heidelberg Germany | Magnetita 500 mikromolov/ml raztopina za injiciranje | 500 micromol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|---------------|--|--|--------------|------------------------|----------------------------|
| SI - Slovenia | Bayer Schering Pharma AG Müllerstrasse 170-178 DE-13342 Berlin-Wedding Germany | Magnevist | 0,5 mmol/ml | solution for injection | intravenous use |
| SI - Slovenia | Auremiana izvozno uvozno trgovsko podjetje, d.o.o., Sežana, Partizanska 109, 6210 Sežana | Multihance 0,5 mmol/ml raztopina za injiciranje | 334 mg/ml | solution for injection | intravenous use |
| SI - Slovenia | Emporio Medical d.o.o., Prešernova 5, 1000 Ljubljana, Slovenia | Dotarem 0,5mmol/ml raztopina za injiciranje | 27,93 g/ml | solution for injection | intravenous use |
| SI - Slovenia | Bayer Schering Pharma AG Müllerstrasse 170-178, DE-13342 Berlin-Wedding Germany | Gadovist 1,0 mmol/ml raztopina za injiciranje | 1,0 mmol/ml | solution for injection | intravenous use |
| SI - Slovenia | Bayer Schering Pharma AG Müllerstrasse 170-178, DE-13342 Berlin-Wedding Germany | Primovist 0,25 mmol/ml raztopina za injiciranje v napolnjeni injekcijski brizgi | 0,25 mmol/ml | solution for injection | intravenous use |
| SK - Slovakia | Bayer Schering Pharma AG Müllerstrasse 170-178, DE-13342 Berlin-Wedding Germany | Gadovist 1,0 mmol/ ml | 604,72 mg/ml | solution for injection | intravenous use |
| SK - Slovakia | GE Healthcare AS Nycoveien 1-2, 4220 Nydalen, Oslo N-0401 Norway | OMNISCAN | 0,5 mmol/ml | solution for injection | intravenous use |
| SK - Slovakia | Bayer Schering Pharma AG Müllerstrasse 170-178, DE-13342 Berlin-Wedding Germany | Primovist 0,25 mmol/ ml, injekčný roztok | 0,25 mmol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|------------------------|---|----------------------------------|--------------------------------|------------------------|-------------------------|
| SK - Slovakia | Bracco IMAGING Deutschland GmbH Max-Stromexer-Str. 116 D-78467 Konstanz Germany | PROHANCE | 279,3 mg/ml | solution for injection | intravenous use |
| SK - Slovakia | Bayer Schering Pharma AG Müllerstrasse 170-178, DE-13342 Berlin-Wedding Germany | Magnevist | 469 mg/ml | solution for injection | intravenous use |
| UK - United Kingdom | GE Healthcare AS Nycoveien 1-2, 4220 Nydalen, Oslo N-0401 Norway | Omniscan injection | 0.5 mmol/ml and 0.5 mmol/litre | solution for injection | intravenous use |
| UK - United Kingdom | Bayer plc Bayer House, Strawberry Hill, Newbury, Berkshire, RG14 1JA UK | Magnevist | 0.5 mmol/ml | solution for injection | intravenous use |
| UK - United Kingdom | Bracco S.p.A. Via Emilio Folli, 50 20134 Milano Italy | Multihance | 0.5 M/ml | solution for injection | intravenous use |
| UK - United Kingdom | Bayer plc Bayer House, Strawberry Hill, Newbury, Berkshire, RG14 1JA UK | Primovist solution for injection | 0.25 mmol/ml | solution for injection | intravenous use |
| UK - United Kingdom | Bracco International B.V., Strawinskylaan 3051 Amsterdam 107 zx Netherlands | Prohance | 0.5 M/ml | solution for injection | intravenous use |
| UK - United Kingdom | Bayer plc Bayer House, Strawberry Hill, Newbury, | Gadovist | 1.0 mmol/ml | solution for injection | intravenous use |

| Member State | Marketing Authorisation Holder | Product name | Strength | Pharmaceutical Form | Route of Administration |
|--------------|--------------------------------------|----------------------|-------------|------------------------|----------------------------|
| | Berkshire, RG14 1JA | | | | |
| | UK | | | | |
| | GUERBET | Dotarem solution for | 0.5 mmol/ml | solution for injection | intravenous use |
| UK - United | BP 57400 | injection | | | |
| Kingdom | 95943 Roissy Charles de Gaulle Cedex | | | | |
| | France | | | | |

ANNEX II

SCIENTIFIC CONCLUSIONS AND GROUNDS FOR AMENDMENT OF THE SUMMARY OF PRODUCT CHARACTERISTICS AND PACKAGE LEAFLETS PRESENTED BY THE EUROPEAN MEDICINES AGENCY

SCIENTIFIC CONCLUSIONS

OVERALL SUMMARY OF THE SCIENTIFIC EVALUATION OF GADOLINIUM-CONTAINING CONTRAST AGENTS (see Annex I)

The gadolinium-containing contrast agents (GdCAs) – gadoversetamide, gadodiamide, gadopentetic acid, gadobenic acid, gadofosveset, gadoxetic acid, gadoteridol, gadobutrol and gadoteric acid - are intravenous agents used for contrast enhancement with magnetic resonance imaging (MRI) and with magnetic resonance angiography (MRA). The GdCAs are available for different types of MR scan varying from product to product, including liver, brain and whole body scan.

GdCAs have been associated with nephrogenic systemic fibrosis (NSF), a rare, serious and life-threatening syndrome involving fibrosis of the skin, joints and internal organs in patients with severe renal impairment. GdCAs were first associated with nephrogenic systemic fibrosis (NSF) in January 2006 when five end-stage renal failure patients undergoing MRA developed signs of NSF two to four weeks after GdCAs administration. This followed a cluster of 25 cases of NSF (20 in Denmark and 5 in Austria) in patients with severe renal impairment, to whom gadodiamide had been administered. Since June 2006 there have been reports of NSF associated with other GdCAs and this issue has been subject to close regulatory reviews leading to risk minimisation measures at the national level.

On 6 November 2008 Denmark asked the CHMP, under Article 31 of Directive 2001/83/EC, to give its opinion on whether the marketing authorisations for GdCAs should be varied in relation to its use in special patient's population more at risk to develop nephrogenic systemic fibrosis (NSF). On 19 November 2008, the European Commission triggered the corresponding procedure under Article 20 of Council Regulation (EC) 726/2004, for GdCAs which are centrally authorised (gadoversetamide and gadofosveset).

The CHMP reviewed all the information made available by the Marketing Authorisation Holders.

The estimated relative risk for NSF calculated based on the number of unconfounded cases and GdCA is higher for gadodiamide (100%), gadoversetamide (94%), and gadopentetic acid (10%) and <1% for gadoteridol and gadoteric acid. No relative risk was estimated for the other GdCAs as their usage is too low.

All GdCAs are chelate complexes containing Gd³⁺, the highly toxic gadolinium ion, which potentially may be released through transmetallation *in vivo*. The extent of transmetallation differs significantly between the complexes with the linear chelates more likely to release Gd³⁺ than the cyclical chelates where the gadolinium ion is caged in a cavity. Other factors such as renal impairment would likely increase the toxicity of the complexes by slowing the clearance of Gd³⁺.

Based on the above the CHMP recognised that there are different categories of NSF-risk for GdCAs:

High risk:

- a) Linear non-ionic chelates including gadoversetamide (OptiMARK) and gadodiamide (Omniscan).
- b) Linear ionic chelate: gadopentetic acid (Magnevist, Gado-MRT-ratiopharm, Magnegita).

Medium risk:

Linear ionic chelates including gadofosveset (Vasovist), gadoxetic acid (Primovist) and gadobenic acid (MultiHance).

Low risk:

Macrocyclic chelates including gadoteric acid (Dotarem), gadoteridol (ProHance) and gadobutrol (Gadovist).

The CHMP recognises that within the high risk group the risk of NSF with gadodiamide and gadoversetamide appears higher than with gadopentetic acid based on physicochemical properties, studies in animals and the number of cases of NSF reported. However as the risk with gadopentetic acid remains substantially higher than the NSF risk with the other lower risk contrast agents, the CHMP recommended that gadopentetic acid should be retained in the high risk group and be subject to the same risk minimisation measures.

In order to minimise the recognised risk associated with GdCAs and the development of NSF, the CHMP agreed on the following measures for the following at risk patient groups:

Use during pregnancy and lactation

Use during pregnancy is not recommended for any GdCA due to the possibility of gadolinium accumulation in human tissues. Although only small amounts of gadolinium are excreted into human breast milk, the immaturity of foetal kidneys could delay the excretion of gadolinium leading to the possibility of long-term accumulation of gadolinium in tissues. Discontinuation of breast feeding for at least 24 h is therefore recommended for all patients receiving high NSF-risk GdCAs. For all other GdCAs the continuation or suspension of breast feeding is left to the discretion of the mother in consultation with the doctor.

Renally impaired patients and haemodialysis

The use of high risk GdCAs is contraindicated in patients with severe renal impairment. Strong warnings are included in the GdCAs of medium and low NSF risk as regards use in patients with severe renal impairment but subject to dose restriction to a minimum during a scan and with a minimum 7 day interval between administrations.

For patients with moderate renal impairment, since the risk is unknown for the high risk category of GdCAs it was agreed that use should only be considered after careful consideration of the benefit-risk, subject to dose restriction to not more than one injection of the minimum dose during a scan with a minimum 7 day interval between administrations.

There is no evidence that supports the use of haemodialysis for preventing or treating NSF in patients not already undergoing haemodialysis, but this may be useful at removing GdCAs in patients already on haemodialysis. This information is reflected in all GdCAs' product information.

Liver transplant patients

Patients undergoing liver transplantation are at particular risk of NSF if exposed to GdCAs particularly to the high-risk GdCAs. Therefore the use of high-risk GdCAs is contraindicated in this population. Strong warnings are included for medium and low NSF risk GdCAs as regards use in this particular special population. However, if use is necessary then dose restrictions to a minimum dose during one scan with a minimum 7 day interval between administrations are recommended.

Paediatric patients

The use of the high risk category of GdCAs in neonates up to 4 weeks of age is contra-indicated. The use of medium and low risk GdCAs in neonates should only be considered after careful consideration subject to dose and interval administration restrictions.

Due to the immature renal function of infants below 1 year of age the use of all GdCAs should be subject to careful consideration and to dose and interval administration restrictions to not more than one injection of the minimum dose during a scan with a minimum 7 day interval between dose administrations.

Elderly patients

No dose adjustments are recommended but screening of 65 years and older patients for renal dysfunction is of particular importance prior to the administration of GdCAs.

Other precautionary measures

Screening for renal dysfunction

For all patients to whom high NSF risk GdCAs will be administered, mandatory screening for renal dysfunction by laboratory tests is required. This screening is recommended for all patients who will receive medium and low NSF-risk GdCAs. Laboratory tests are more effective to assess the renal function of all at-risk patients, since changes in renal function are often not reflected symptomatically or clinically.

In addition to the minimisation measures included in the product information, the CHMP having considered the evidence that toxic free gadolinium ions are retained in human tissues concluded that studies evaluating the potential for long-term retention of gadolinium in bone are needed. Therefore, the MAHs are requested to submit to the CHMP protocols and timelines for the studies of gadolinium accumulation in human bone within 3 months of the decision on this referral procedure. The testing of bone samples from patients undergoing hip and knee replacement surgery is recommended. Co-factors that may increase the risk of NSF such as serum calcium and phosphate levels at the time of administration of a GdCA should be studied and biomarkers evaluated.

In addition, the MAHs should submit a cumulative review on NSF cases annually for 3 consecutive years commencing one year after the decision on this referral procedure.

The need to have a harmonised traceability method across Europe for effective monitoring of the use of GdCAs was agreed. The use of "sticky labels" detachable from the vials and syringes are considered an appropriate method to be implemented for all GdCAs.

The MAH for Omniscan (gadodiamide) disagreed with the proposed labelling warnings with respect to screening patients for renal dysfunction and requested a re-examination of the opinion.

The MAH supported the CHMP proposed risk minimisation for all patients to be screened for renal dysfunction irrespectively of the GdCAs. However, screening should only require laboratory testing following evaluation of the patient's medical history and the minimisation measure should be the same for all GdCAs.

Having considered the detailed grounds for re-examination provided by the MAH in writing, the CHMP agrees that medical history could identify some patients with possible renal dysfunction. However, medical history alone could not be relied on, as it will not be sufficient to identify all at-risk patients. Laboratory tests are more effective to assess the renal function of all at-risk patients, since changes in renal function are often not reflected symptomatically or clinically. Encouraging appropriate testing of renal function should ensure identification of patients at risk and ensure use of appropriate diagnostic agents.

This risk minimisation was applied in accordance with the three different categories of NSF-risk for GdCAs recognised by the CHMP based on their thermodynamic and kinetic properties. Therefore and considering the overall benefit/risk, the CHMP agreed that for all patients to be administered with high NSF-risk GdCAs mandatory screening by laboratory tests should be performed.

Based on the above, the CHMP concluded that its Opinion of 19 November 2009 should be maintained with the recommended amendments to relevant sections of the Summary of Product Characteristics and Package Leaflet as set out in Annex III to the opinion.

GROUNDS FOR AMENDMENT OF THE SUMMARY OF PRODUCT CHARACTERISTICS AND PACKAGE LEAFLETS

Whereas

- The Committee considered the referral made under article 31 of Directive 2001/83/EC, as amended, for gadolinium containing contrast agents (GdCAs) initiated by Denmark.
- The Committee assessed the grounds for re-examination submitted by the MAH of Omniscan (gadodiamide) on 25 January 2010 and the scientific discussion within the Committee;
- The Committee considered all the available data submitted on the safety of the gadolinium containing contrast agents, in relation to the risk of NSF.
- The Committee, concluded that gadolinium contrast agents are associated with NSF and that the
 risk is increased in renal impaired patients, liver transplant patients, the paediatric population,
 during pregnancy and lactation and in the elderly.
 The CHMP also recognised that according to their risk for NSF, GdCAs can be classified in 3
 risk categories: high, medium and low risk.
- The CHMP concluded that the Product Information of all GdCAs should include safety
 information to minimise the risk of NSF and therefore recommended the amendments to the
 relevant sections of the Summaries of Product Characteristics and Package Leaflets in
 accordance to the risk category.
 Furthermore, Risk Minimisation Measures on the traceability as well as the long-term effects of
 these products in Europe are recommended.

As a consequence, the CHMP has recommended the maintenance of the Marketing Authorisations for the medicinal products referred to in Annex I for which the amendments to the relevant sections of the Summary of Product Characteristics and Package Leaflet are set out in Annex III and in accordance to the conditions set out in Annex IV.

ANNEX III

AMENDMENTS TO THE SUMMARY OF PRODUCT CHARACTERISTICS AND PACKAGE LEAFLETS

AMENDMENTS TO BE INCLUDED IN THE RELEVANT SECTIONS OF THE SUMMARY OF PRODUCT CHARACTERISTICS FOR HIGH RISK GADOLINIUM-CONTAINING CONTRAST AGENTS

(gadodiamide, gadopentetic acid)

4.2 Posology and method of administration

[Use currently approved text with the addition of guidance for special populations (patients with renal impairment, neonates, infants and the elderly)]

Special Populations

Renal impairment

{Invented name} is contraindicated in patients with severe renal impairment (GFR < 30 ml/min/1.73m²) and in patients in the perioperative liver transplantation period (see section 4.3). {Invented name} should only be used after careful risk/benefit evaluation in patients with moderate renal impairment (GFR 30-59 ml/min/1.73m²) at a dose not exceeding $\{x\}$ mmol/kg body weight (see section 4.4). More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

[If use is permitted in neonates and infants, add a statement on use in neonates and infants]

Neonates up to 4 weeks of age and infants up to 1 year of age

{Invented name} is contraindicated in neonates up to 4 weeks of age (see section 4.3). Due to immature renal function in infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration at a dose not exceeding {x} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

[If use is restricted by age for a particular indication add a statement on use in neonates and infants and specify the indication and age at which use is not recommended]

Neonates up to 4 weeks of age, infants up to 1 year of age and children

{Invented name} is contraindicated in neonates up to 4 weeks of age (see section 4.3).

Due to immature renal function in infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration at a dose not exceeding {x} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

Use for whole body MRI is not recommended in children less than 6 months of age.

[If use is restricted for use in children greater than 1 year of age, specify the indication and age at which use is not recommended]

Neonates up to 4 weeks of age, infants up to 1 year of age and children

{Invented name} is contraindicated in neonates up to 4 weeks of age (see section 4.3).Use of {Invented name} is not recommended in children less than 2 years of age.

Elderly (aged 65 years and above)

No dosage adjustment is considered necessary. Caution should be exercised in elderly patients (see section 4.4).

4.3 Contraindications

[Use currently approved text with addition of contraindication below]

{Invented name} is contraindicated in patients with severe renal impairment (GFR <30ml/min/1.73m²), in patients in the perioperative liver transplantation period and in neonates up to 4 weeks of age (see section 4.4).

4.4 Special warnings and precautions for use

[With the exception of information on impaired renal function, use in neonates and infants and use in the elderly, use currently approved text]

Impaired renal function

Prior to administration of {Invented name}, all patients should be screened for renal dysfunction by obtaining laboratory tests.

There have been reports of nephrogenic systemic fibrosis (NSF) associated with use of {Invented name} and some other gadolinium-containing contrast agents in patients with acute or chronic severe renal impairment (GFR $< 30 \text{ ml/min}/1.73\text{m}^2$). Patients undergoing liver transplantation are at particular risk since the incidence of acute renal failure is high in this group. Therefore {Invented name} must not be used in patients with severe renal impairment, in patients in the perioperative liver transplantation period and in neonates (see section 4.3).

The risk for development of NSF in patients with moderate renal impairment (GFR 30–59 ml/min/1.73 m²) is unknown, therefore, {Invented name} should be only used after careful risk-benefit evaluation in patients with moderate renal impairment.

Haemodialysis shortly after {Invented name} administration may be useful at removing {Invented name} from the body. There is no evidence to support the initiation of haemodialysis for prevention or treatment of NSF in patients not already undergoing haemodialysis.

[If use is not authorised for infants below the age of 1 year a statement in section 4.4 is not necessary. If use is authorised in neonates and infants add the following statement]

Neonates and infants

{Invented name} is contraindicated in neonates up to 4 weeks of age (see section 4.3). Due to immature renal function in infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration.

[If use is only authorised in infants above the age of 6 months add the following statement]

<u>Infants</u>

Due to immature renal function in infants up to 1 year of age, {Invented name} should only be used in patients 6 to 12 months of age after careful consideration.

Elderly

As the renal clearance of {active substance} may be impaired in the elderly, it is particularly important to screen patients aged 65 years and older for renal dysfunction.

4.6 Pregnancy and lactation

Pregnancy

[Amend currently approved text for data in pregnant women and animal studies in line with the CHMP Guideline on Risk Assessment of Medicinal Products on Human Reproduction and Lactation: From Data to Labelling (24 July 2008)]

There are no data from the use of {active substance} in pregnant women. Animal studies do not indicate direct or indirect harmful effects with respect to reproductive toxicity (see section 5.3). {Invented name} should not be used during pregnancy unless the clinical condition of the woman requires use of {active substance}.

or

There are no data from the use of {active substance} in pregnant women. Animal studies have shown reproductive toxicity at repeated high doses (see section 5.3). {Invented name} should not be used during pregnancy unless the clinical condition of the woman requires use of {active substance}.

Lactation

[Amend currently approved text for data in lactating women and animal studies in line with the CHMP Guideline on Risk Assessment of Medicinal Products on Human Reproduction and Lactation: From Data to Labelling (24 July 2008)]

It is unknown whether {active substance} is excreted in human milk. There is insufficient information on the excretion of {active substance} in animal milk. A risk to the suckling child cannot be excluded. Breast-feeding should be discontinued for at least 24 hours after the administration of {Invented name}.

or

It is unknown whether {active substance} is excreted in human milk. Available data in animals have shown excretion of {active substance} in milk (for details see section 5.3). A risk to the suckling child cannot be excluded. Breast-feeding should be discontinued for at least 24 hours after the administration of {Invented name}.

4.8 Undesirable effects

Cases of nephrogenic systemic fibrosis (NSF) have been reported with {Invented name} (see section 4.4).

4.9 Overdose

[Use currently approved text with the addition of haemodialysis wording]

{Invented name} can be removed by haemodialysis. However there is no evidence that haemodialysis is suitable for prevention of nephrogenic systemic fibrosis (NSF).

6.6 Special precautions for disposal and other handling

[*Use currently approved text for information on disposal*]

The peel-off tracking label on the {vials/syringes/bottles} should be stuck onto the patient record to enable accurate recording of the gadolinium contrast agent used. The dose used should also be

recorded.

AMENDMENTS TO BE INCLUDED IN THE RELEVANT SECTIONS OF THE SECTIONS OF THE PACKAGE LEAFLET FOR HIGH RISK GADOLINIUM-CONTAINING CONTRAST AGENTS

(gadodiamide, gadopentetic acid)

2. BEFORE YOU ARE GIVEN {Invented name}

Take special care with {Invented name}:

[Use currently approved text with addition of the information on the NSF]

You should not be given {Invented name} if you suffer from severe kidney problems, or if you are a patient who is about to have or has recently had a liver transplant, as use of {Invented name} in patients with these conditions has been associated with a disease called nephrogenic systemic fibrosis (NSF). NSF is a disease involving thickening of the skin and connective tissues. NSF may result in severe joint immobility, muscle weakness or may affect the normal working of internal organs which may potentially be life threatening.

{Invented name} should also not be given to newborn babies up to age of 4 weeks.

Tell your doctor if:

[*Use currently approved text*]

- your kidneys do not work properly
- you have recently had, or soon expect to have, a liver transplant

[Use currently approved text with the addition of information on impaired renal function]

Before you receive {Invented name}, you will need to have a blood test to check how well your kidneys are working.

[Use currently approved text with the addition of information on use in neonates and infants]

[If use is authorised in infants add the following statement]

{Invented name} should not be used in newborn babies up to the age of 4 weeks. As kidney function is immature in infants up to 1 year of age, {Invented name} will only be used in infants after careful consideration by the doctor.

[If use is only authorised in infants above the age of 6 months add the following statement]

{Invented name} should not be used in newborn babies up to the age of 4 weeks. As kidney function is immature in infants up to 1 year of age, {Invented name} will only be used in patients 6 to 12 months of age after careful consideration by the doctor.

[*If use is not authorised in children under 2 years add the following statement*]

{Invented name} should not be used in newborn babies up to the age of 4 weeks and is not recommended in children under 2 years.

Pregnancy and breast-feeding

Ask your doctor for advice before taking any medicine.

Pregnancy

You must tell your doctor if you think you are or might become pregnant as {Invented name} should not be used during pregnancy unless strictly necessary.

Breast-feeding

Tell your doctor if you are breast-feeding or about to start breast-feeding. Breast-feeding should be discontinued for at least 24 hours after you receive {Invented name}.

3. HOW TO USE {Invented name}

Dosage in special patient groups

[Use currently approved text with the addition of information on impaired renal function, use in neonates and infants and use in the elderly]

You should not be given {Invented name} if you suffer from severe kidney problems or if you are a patient who is about to have or has recently had a liver transplant. {Invented name} should also not be used in newborn babies up to the age of 4 weeks.

If you have moderate kidney problems, you should only receive one dose of {Invented name} during a scan and you should not receive a second injection for at least 7 days.

[If use is permitted in infants, add a statement on use in infants]

As kidney function is immature in infants up to 1 year of age, infants should only receive one dose of {Invented name} during a scan and should not receive a second injection for at least 7 days.

It is not necessary to adjust your dose if you are 65 years of age or older but you will have a blood test to check how well your kidneys are working.

4. POSSIBLE SIDE EFFECTS

[*Use currently approved text with the addition of information on NSF*]

There have been reports of nephrogenic systemic fibrosis (which causes hardening of the skin and may affect also soft tissue and internal organs).

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|---|------|------|
| | | |

The following information is intended for medical or healthcare professionals only:

[*Use currently approved text with the addition of information on NSF*]

Prior to administration of {Invented name}, all patients should be screened for renal dysfunction by obtaining laboratory tests.

There have been reports of nephrogenic systemic fibrosis (NSF) associated with use of {Invented name} and some other gadolinium-containing contrast agents in patients with acute or chronic severe renal impairment (GFR $< 30 \text{ ml/min}/1.73\text{m}^2$). Patients undergoing liver transplantation are at particular risk since the incidence of acute renal failure is high in this group. Therefore {Invented name} must not be used in patients with severe renal impairment, in patients in the perioperative liver transplantation period.

{Invented name} should also not be given to newborn babies up to the age of 4 weeks.

The risk for development of NSF in patients with moderate renal impairment (GFR 30–59 ml/min/1.73 m²) is unknown, therefore, {Invented name} should be only used after careful risk-benefit evaluation in patients with moderate renal impairment at a dose not exceeding {x} mmol/kg body weight. More

than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

Due to immature renal function in infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration at a dose not exceeding {x} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days. {Invented name} should not be given to newborn babies up to age of 4 weeks.

As the renal clearance of {active substance} may be impaired in the elderly, it is particularly important to screen patients aged 65 years and older for renal dysfunction.

Haemodialysis shortly after {Invented name} administration may be useful at removing {Invented name} from the body. There is no evidence to support the initiation of haemodialysis for prevention or treatment of NSF in patients not already undergoing haemodialysis.

{Invented name} should not be used during pregnancy unless the clinical condition of the woman requires use of {active substance}.

Breast-feeding should be discontinued for at least 24 hours after the administration of {Invented name}.

The peel-off tracking label on the {vials/syringes/bottles} should be stuck onto the patient record to enable accurate recording of the gadolinium contrast agent used. The dose used should also be recorded.

AMENDMENTS TO BE INCLUDED IN THE RELEVANT SECTIONS OF THE SUMMARY OF PRODUCT CHARACTERISTICS FOR MEDIUM RISK GADOLINIUM-CONTAINING CONTRAST AGENTS

(gadoxetic acid, gadobenic acid)

4.2 Posology and method of administration

[Use currently approved text with the addition of guidance for special populations (patients with renal impairment, neonates, infants and the elderly)]

Special Populations

<u>Impaired renal function</u>

Use of {Invented name} should be avoided in patients with severe renal impairment (GFR < $30 \, \text{ml/min}/1.73\,\text{m}^2$) and in patients in the perioperative liver transplantation period unless the diagnostic information is essential and not available with non-contrast enhanced MRI (see section 4.4). If use of {Invented name} cannot be avoided, the dose should not exceed {x} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

[If use is permitted in neonates and infants add a statement on use in neonates and infants]

Neonates up to 4 weeks of age and infants up to 1 year of age

Due to immature renal function in neonates up to 4 weeks of age and infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration at a dose not exceeding {x} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

[If use is restricted by age for a particular indication, add a statement on use in neonates and infants and specify the indication and age at which use is not recommended]

Neonates up to 4 weeks of age, infants up to 1 year of age and children

Due to immature renal function in neonates up to 4 weeks of age and infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration at a dose not exceeding {x} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

Use for whole body MRI is not recommended in children less than 6 months of age.

[If use is restricted for use in children greater than 1 year of age, specify the indication and age for which use is not recommended]

Neonates up to 4 weeks of age, infants up to 1 year of age and children

Use of {Invented name} is not recommended in children less than 2 years of age.

Elderly (aged 65 years and above)

No dosage adjustment is considered necessary. Caution should be exercised in elderly patients (see section 4.4).

4.4 Special warnings and precautions for use

[With the exception of information on impaired renal function, use in neonates and infants and use in the elderly, use currently approved text]

<u>Impaired renal function</u>

Prior to administration of {Invented name}, it is recommended that all patients are screened for renal dysfunction by obtaining laboratory tests.

There have been reports of nephrogenic systemic fibrosis (NSF) associated with use of some gadolinium-containing contrast agents in patients with acute or chronic severe renal impairment (GFR < 30 ml/min/1.73m²). Patients undergoing liver transplantation are at particular risk since the incidence of acute renal failure is high in this group. As there is a possibility that NSF may occur with {Invented name}, it should therefore be avoided in patients with severe renal impairment and in patients in the perioperative liver transplantation period unless the diagnostic information is essential and not available with non-contrast enhanced MRI.

Haemodialysis shortly after {Invented name} administration may be useful at removing {Invented name} from the body. There is no evidence to support the initiation of haemodialysis for prevention or treatment of NSF in patients not already undergoing haemodialysis.

[If use is not authorised for infants below the age of 1 year a statement in section 4.4 is not necessary. If use is authorised in neonates and infants add the following statement]

Neonates and infants

Due to immature renal function in neonates up to 4 weeks of age and infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration.

[If use is only authorised in infants above the age of 6 months add the following statement]

Infants

Due to immature renal function in infants up to 1 year of age, {Invented name} should only be used in patients 6 to 12 months of age after careful consideration. Elderly

As the renal clearance of {active substance} may be impaired in the elderly, it is particularly important to screen patients aged 65 years and older for renal dysfunction.

4.6 Pregnancy and lactation

Pregnancy

[Amend currently approved text for data in pregnant women and animal studies in line with the CHMP Guideline on Risk Assessment of Medicinal Products on Human Reproduction and Lactation: From Data to Labelling (24 July 2008)]

There are no data from the use of {active substance} in pregnant women. Animal studies do not indicate direct or indirect harmful effects with respect to reproductive toxicity (see section 5.3). {Invented name} should not be used during pregnancy unless the clinical condition of the woman requires use of {active substance}.

There are no data from the use of {active substance} in pregnant women. Animal studies have shown reproductive toxicity at repeated high doses (see section 5.3). {Invented name} should not be used during pregnancy unless the clinical condition of the woman requires use of {active substance}.

Lactation

Gadolinium containing contrast agents are excreted into breast milk in very small amounts (see section 5.3). At clinical doses, no effects on the infant are anticipated due to the small amount excreted in milk and poor absorption from the gut. Continuing or discontinuing breast feeding for a period of 24 hours after administration of {Invented name}, should be at the discretion of the doctor and lactating mother.

4.8 Undesirable effects

[Use currently approved text with the addition of information on NSF]

Isolated cases of nephrogenic systemic fibrosis (NSF) have been reported with {Invented name} (see section 4.4).

or

Isolated cases of nephrogenic systemic fibrosis (NSF) have been reported with {Invented name}, most of which were in patients co-administered other gadolinium-containing contrast agents (see section 4.4).

or

Isolated cases of nephrogenic systemic fibrosis (NSF) have been reported with {Invented name} in patients co-administered other gadolinium-containing contrast agents (see section 4.4).

or

Cases of nephrogenic systemic fibrosis (NSF) have been reported with other gadolinium-containing contrast agents (see section 4.4).

4.9 Overdose

[Use currently approved text with the addition of haemodialysis wording]

{Invented name} can be removed by haemodialysis. However there is no evidence that haemodialysis is suitable for prevention of nephrogenic systemic fibrosis (NSF).

6.6 Special precautions for disposal and other handling

[Use currently approved text for information on disposal]

The peel-off tracking label on the {vials/syringes/bottles} should be stuck onto the patient record to enable accurate recording of the gadolinium contrast agent used. The dose used should also be recorded.

AMENDMENTS TO BE INCLUDED IN THE RELEVANT SECTIONS OF THE SECTIONS OF THE PACKAGE LEAFLET FOR MEDIUM RISK GADOLINIUM-CONTAINING CONTRAST AGENTS

(gadoxetic acid, gadobenic acid)

2. BEFORE YOU ARE GIVEN {Invented name}

Take special care with {Invented name}:

[Use currently approved text with the addition of information on impaired renal function, use in neonates and infants and use in the elderly]

Tell your doctor if:

- your kidneys do not work properly
- you have recently had, or soon expect to have, a liver transplant

[The following statement should be added]

Your doctor may decide to take a blood test to check how well your kidneys are working before making the decision to use {Invented name}, especially if you are 65 years of age or older.

[If use is not authorised in neonates and infants add the following statement]

The safety of {Invented name} in persons under 18 years has not yet been tested.

[*If use is authorised in neonates and infants add the following statement*]

Neonates and infants

As kidney function is immature in babies up to 4 weeks of age and infants up to 1 year of age, {Invented name} will only be used in these patients after careful consideration by the doctor.

[If use is only authorised in infants above the age of 6 months add the following statement]

Infants

As kidney function is immature in infants up to 1 year of age, {Invented name} will only be used in patients 6 to 12 months of age after careful consideration by the doctor.

Pregnancy and breast-feeding

Ask your doctor for advice before taking any medicine.

Pregnancy

You must tell your doctor if you think you are or might become pregnant as {Invented name} should not be used during pregnancy unless strictly necessary.

Breast-feeding

Tell your doctor if you are breast-feeding or about to start breast-feeding. Your doctor will discuss whether you should continue breast-feeding or interrupt breast-feeding for a period of 24 hours after you receive {Invented name}.

3. HOW TO USE {Invented name}

Dosage in special patient groups

[Use currently approved text with the addition of information on impaired renal function, use in neonates and infants and use in the elderly]

The use of {Invented name} is not recommended in patients with severe kidney problems and patients who have recently had, or soon expect to have, a liver transplant. However if use is required you should only receive one dose of {Invented name} during a scan and you should not receive a second injection for at least 7 days.

Neonates, infants, children and adolescents

[If use is permitted in neonates and infants add a statement on use in neonates and infants]

As kidney function is immature in babies up to 4 weeks of age and infants up to 1 year of age, {Invented name} will only be used in these patients after careful consideration by the doctor. Neonates and infants should only receive one dose of {Invented name} during a scan and should not receive a second injection for at least 7 days.

[If use is restricted for use in children greater than 1 year of age, specify the indication and age for which use is not recommended]

Use of {Invented name} is not recommended in children less than 2 years of age.

[If use is restricted by age for a particular indication, add a statement on use in neonates and infants and specify the indication and age at which use is not recommended]

Use for whole body MRI is not recommended in children less than 6 months of age.

Elderly

It is not necessary to adjust your dose if you are 65 years of age or older but you may have a blood test to check how well your kidneys are working.

4. POSSIBLE SIDE EFFECTS

[*Use currently approved text with the addition of information on NSF*]

There have been reports of nephrogenic systemic fibrosis (which causes hardening of the skin and may affect also soft tissue and internal organs).

or

There have been reports of nephrogenic systemic fibrosis (which causes hardening of the skin and may affect also soft tissue and internal organs) most of which were in patients who received {Invented name} together with other gadolinium-containing contrast agents.

or

There have been reports of nephrogenic systemic fibrosis (which causes hardening of the skin and may affect also soft tissue and internal organs) in patients who received {Invented name} together with other gadolinium-containing contrast agents.

or

There have been reports of nephrogenic systemic fibrosis (which causes hardening of the skin and may affect also soft tissue and internal organs) associated with use of other gadolinium-containing contrast agents.

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The following information is intended for medical or healthcare professionals only:

[Use currently approved text with the addition of information on NSF]

Prior to administration of {Invented name}, it is recommended that all patients are screened for renal dysfunction by obtaining laboratory tests.

There have been reports of nephrogenic systemic fibrosis (NSF) associated with use of some gadolinium-containing contrast agents in patients with acute or chronic severe renal impairment (GFR< 30ml/min /1.73 m²). Patients undergoing liver transplantation are at particular risk since the incidence of acute renal failure is high in this group. As there is a possibility that NSF may occur with {Invented name}, it should therefore be avoided in patients with severe renal impairment and in patients in the perioperative liver transplantation period unless the diagnostic information is essential and not available with non-contrast enhanced MRI. If use of {Invented name} cannot be avoided, the dose should not exceed {x} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

Due to immature renal function in neonates up to 4 weeks of age and infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration at a dose not exceeding {x} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

As the renal clearance of {active substance} may be impaired in the elderly, it is particularly important to screen patients aged 65 years and older for renal dysfunction.

Haemodialysis shortly after {Invented name} administration may be useful at removing {Invented name} from the body. There is no evidence to support the initiation of haemodialysis for prevention or treatment of NSF in patients not already undergoing haemodialysis.

{Invented name} should not be used during pregnancy unless the clinical condition of the woman requires use of {active substance}.

Continuing breast feeding or discontinuing {Invented name} for a period of 24 hours after administration, should be at the discretion of the doctor and lactating mother.

The peel-off tracking label on the {vials/syringes/bottles} should be stuck onto the patient record to enable accurate recording of the gadolinium contrast agent used. The dose used should also be recorded.

AMENDMENTS TO BE INCLUDED IN THE RELEVANT SECTIONS OF THE SUMMARY OF PRODUCT CHARACTERISTICS FOR LOW RISK GADOLINIUM-CONTAINING CONTRAST AGENTS

(gadoteric acid, gadoteridol, gadobutrol)

4.2 Posology and method of administration

[Use currently approved text with the addition of guidance for special populations (patients with renal impairment, neonates, infants and the elderly)]

Special Populations

<u>Impaired renal function</u>

{Invented name} should only be used in patients with severe renal impairment (GFR < 30 ml/min/1.73m²) and in patients in the perioperative liver transplantation period after careful risk/benefit assessment and if the diagnostic information is essential and not available with noncontrast enhanced MRI (see section 4.4). If it is necessary to use {Invented name}, the dose should not exceed {x} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

[If use is permitted in neonates and infants add a statement on use in neonates and infants]

Neonates up to 4 weeks of age and infants up to 1 year of age

Due to immature renal function in neonates up to 4 weeks of age and infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration at a dose not exceeding {x} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

[If use is restricted by age for a particular indication add a statement on use in neonates and infants and specify the indication and age at which use is not recommended]

Neonates up to 4 weeks of age, infants up to 1 year of age and children

Due to immature renal function in neonates up to 4 weeks of age and infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration at a dose not exceeding {X} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

Use for whole body MRI is not recommended in children less than 6 months of age.

[If use is restricted for use in children greater than 1 year of age specify the indication and age at which use is not recommended]

Neonates up to 4 weeks of age, infants up to 1 year of age and children

Use of {Invented name} is not recommended in children less than 2 years of age.

Elderly (aged 65 years and above)

No dosage adjustment is considered necessary. Caution should be exercised in elderly patients (see section 4.4).

4.4 Special warnings and precautions for use

[With the exception of information on impaired renal function, use in neonates and infants and use in the elderly, use currently approved text]

Impaired renal function

Prior to administration of {Invented name}, it is recommended that all patients are screened for renal dysfunction by obtaining laboratory tests.

There have been reports of nephrogenic systemic fibrosis (NSF) associated with use of some gadolinium-containing contrast agents in patients with acute or chronic severe renal impairment (GFR < 30 ml/min/1.73m²). Patients undergoing liver transplantation are at particular risk since the incidence of acute renal failure is high in this group. As there is a possibility that NSF may occur with {Invented name}, it should therefore only be used in patients with severe renal impairment and in patients in the perioperative liver transplantation period after careful risk/benefit assessment and if the diagnostic information is essential and not available with non-contrast enhanced MRI.

Haemodialysis shortly after {Invented name} administration may be useful at removing {Invented name} from the body. There is no evidence to support the initiation of haemodialysis for prevention or treatment of NSF in patients not already undergoing haemodialysis.

[If use is not authorised for infants below the age of 1 year a statement in section 4.4 is not necessary. If use is authorised in neonates and infants add the following statement]

Neonates and infants

Due to immature renal function in neonates up to 4 weeks of age and infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration.

[If use is only authorised in infants above the age of 6 months add the following statement]

Infants

Due to immature renal function in infants up to 1 year of age, {Invented name} should only be used in patients 6 to 12 months of age after careful consideration.

Elderly

As the renal clearance of {active substance} may be impaired in the elderly, it is particularly important to screen patients aged 65 years and older for renal dysfunction.

4.6 Pregnancy and lactation

Pregnancy

[Amend currently approved text for data in pregnant women and animal studies in line with the CHMP Guideline on Risk Assessment of Medicinal Products on Human Reproduction and Lactation: From Data to Labelling (24 July 2008)]

There are no data from the use of {active substance} in pregnant women. Animal studies do not indicate direct or indirect harmful effects with respect to reproductive toxicity (see section 5.3). {Invented name} should not be used during pregnancy unless the clinical condition of the woman requires use of {active substance}.

or

There are no data from the use of {active substance} in pregnant women. Animal studies have shown reproductive toxicity at repeated high doses (see section 5.3). {Invented name} should not be used during pregnancy unless the clinical condition of the woman requires use of {active substance}.

Lactation

Gadolinium containing contrast agents are excreted into breast milk in very small amounts (see section 5.3). At clinical doses, no effects on the infant are anticipated due to the small amount excreted in milk and poor absorption from the gut. Continuing or discontinuing breast feeding for a period of 24 hours after administration of {Invented name}, should be at the discretion of the doctor and lactating mother.

4.8 Undesirable effects

[*Use currently approved text with the addition of information on NSF*]

Isolated cases of nephrogenic systemic fibrosis (NSF) have been reported with {Invented name} (see section 4.4).

or

Isolated cases of nephrogenic systemic fibrosis (NSF) have been reported with {Invented name}, most of which were in patients co-administered other gadolinium-containing contrast agents (see section 4.4).

or

Isolated cases of nephrogenic systemic fibrosis (NSF) have been reported with {Invented name} in patients co-administered other gadolinium-containing contrast agents (see section 4.4).

or

Cases of nephrogenic systemic fibrosis (NSF) have been reported with other gadolinium-containing contrast agents (see section 4.4).

4.9 Overdose

[Use currently approved text with the addition of haemodialysis wording]

{Invented name} can be removed by haemodialysis. However there is no evidence that haemodialysis is suitable for prevention of nephrogenic systemic fibrosis (NSF).

6.6 Special precautions for disposal and other handling

[Use currently approved text for information on disposal]

The peel-off tracking label on the {vials/syringes/bottles} should be stuck onto the patient record to enable accurate recording of the gadolinium contrast agent used. The dose used should also be recorded.

AMENDMENTS TO BE INCLUDED IN THE RELEVANT SECTIONS OF THE SECTIONS OF THE PACKAGE LEAFLET FOR LOW RISK GADOLINIUM-CONTAINING CONTRAST AGENTS

(gadoteric acid, gadoteridol, gadobutrol)

2. BEFORE YOU ARE GIVEN {Invented name}

Take special care with {Invented name}:

[Use currently approved text with the addition of information on impaired renal function, use in neonates and infants and use in the elderly]

Tell your doctor if:

- your kidneys do not work properly
- you have recently had, or soon expect to have, a liver transplant

[The following statement should be added]

Your doctor may decide to take a blood test to check how well your kidneys are working before making the decision to use {Invented name}, especially if you are 65 years of age or older.

[If use is not authorised in neonates and infants add the following statement]

The safety of {Invented name} in persons under 18 has not yet been tested.

[*If use is authorised in neonates and infants add the following statement*]

Neonates and infants

As kidney function is immature in babies up to 4 weeks of age and infants up to 1 year of age, {Invented name} will only be used in these patients after careful consideration by the doctor.

[If use is only authorised in infants above the age of 6 months add the following statement]

Infants

As kidney function is immature in infants up to 1 year of age, {Invented name} will only be used in patients 6 to 12 months of age after careful consideration by the doctor.

Pregnancy and breast-feeding

Ask your doctor for advice before taking any medicine.

Pregnancy

You must tell your doctor if you think you are or might become pregnant as {Invented name} should not be used during pregnancy unless strictly necessary.

Breast-feeding

Tell your doctor if you are breast-feeding or about to start breast-feeding. Your doctor will discuss whether you should continue breast-feeding or interrupt breast-feeding for a period of 24 hours after you receive {Invented name}.

3. HOW TO USE {Invented name}

Dosage in special patient groups

[Use currently approved text with the addition of information on impaired renal function, use in neonates and infants and use in the elderly]

The use of {Invented name} is not recommended in patients with severe kidney problems and patients who have recently had, or soon expect to have, a liver transplant. However if use is required you should only receive one dose of {Invented name} during a scan and you should not receive a second injection for at least 7 days.

Neonates, infants, children and adolescents

[If use is permitted in neonates and infants add a statement on use in neonates and infants]

As kidney function is immature in babies up to 4 weeks of age and infants up to 1 year of age, {Invented name} will only be used in these patients after careful consideration by the doctor. Neonates and infants should only receive one dose of {Invented name} during a scan and should not receive a second injection for at least 7 days.

[If use is restricted for use in children greater than 1 year of age, specify the indication and age for which use is not recommended]

Use of {Invented name} is not recommended in children less than 2 years of age.

[If use is restricted by age for a particular indication, add a statement on use in neonates and infants and specify the indication and age at which use is not recommended]

Use for whole body MRI is not recommended in children less than 6 months of age.

Elderly

It is not necessary to adjust your dose if you are 65 years of age or older but you may have a blood test to check how well your kidneys are working.

4. POSSIBLE SIDE EFFECTS

[Use currently approved text with the addition of information on NSF]

There have been reports of nephrogenic systemic fibrosis (which causes hardening of the skin and may affect also soft tissue and internal organs).

or

or

There have been reports of nephrogenic systemic fibrosis (which causes hardening of the skin and may affect also soft tissue and internal organs) most of which were in patients who received {Invented name} together with other gadolinium-containing contrast agents.

There have been reports of nephrogenic systemic fibrosis (which causes hardening of the skin and may affect also soft tissue and internal organs) in patients who received {Invented name} together with other gadolinium-containing contrast agents.

or

There have been reports of nephrogenic systemic fibrosis (which causes hardening of the skin and may affect also soft tissue and internal organs) associated with use of other gadolinium-containing contrast agents.

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The following information is intended for medical or healthcare professionals only:

[Use currently approved text with the addition of information on NSF]

Prior to administration of {Invented name}, it is recommended that all patients are screened for renal dysfunction by obtaining laboratory tests.

There have been reports of nephrogenic systemic fibrosis (NSF) associated with use of some gadolinium-containing contrast agents in patients with acute or chronic severe renal impairment (GFR< 30ml/min /1.73 m²). Patients undergoing liver transplantation are at particular risk since the incidence of acute renal failure is high in this group. As there is a possibility that NSF may occur with {Invented name}, it should therefore only be used in patients with severe renal impairment and in patients in the perioperative liver transplantation period after careful risk/benefit assessment and if the diagnostic information is essential and not available with non-contrast enhanced MRI. If it is necessary to use {Invented name}, the dose should not exceed {x} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

Due to immature renal function in neonates up to 4 weeks of age and infants up to 1 year of age, {Invented name} should only be used in these patients after careful consideration at a dose not exceeding {X} mmol/kg body weight. More than one dose should not be used during a scan. Because of the lack of information on repeated administration, {Invented name} injections should not be repeated unless the interval between injections is at least 7 days.

As the renal clearance of {active substance} may be impaired in the elderly, it is particularly important to screen patients aged 65 years and older for renal dysfunction.

Haemodialysis shortly after {Invented name} administration may be useful at removing {Invented name} from the body. There is no evidence to support the initiation of haemodialysis for prevention or treatment of NSF in patients not already undergoing haemodialysis.

{Invented name} should not be used during pregnancy unless the clinical condition of the woman requires use of {active substance}.

Continuing breast feeding or discontinuing {Invented name} for a period of 24 hours after administration, should be at the discretion of the doctor and lactating mother.

The peel-off tracking label on the {vials/syringes/bottles} should be stuck onto the patient record to enable accurate recording of the gadolinium contrast agent used. The dose used should also be recorded.

ANNEX IV CONDITIONS OF THE MARKETING AUTHORISATIONS

CONDITIONS OF THE MARKETING AUTHORISATIONS

Annual cumulative safety review

The MAHs should provide to the CHMP an annual cumulative review on nephrogenic systemic fibrosis (NSF) cases commencing one year after Commission Decision and for 3 consecutive years.

Long-term effects study

The MAHs should submit to the CHMP protocols and timelines for studies evaluating the potential for long-term accumulation of gadolinium in human bone. Co-factors that may increase the risk of NSF such as serum calcium and phosphate levels at the time of administration of GdCA should be studies and biomarkers evaluated. The testing of bone samples from patients undergoing hip and knee replacement surgery is recommended. This should be submitted to the CHMP within 3 months of the Commission Decision on this Referral procedure.

Communication

National Competent Authorities should ensure that prescribers will be informed of the measures agreed by CHMP to minimise the risk of NSF. The communication should be based on the "key message document" agreed by CHMP.

Other Minimisation Measures

In order to have a harmonised traceability method across Europe for the effective monitoring of the use of GdCAs the National Competent Authorities, coordinated by the Reference Member State (where applicable) should ensure the implementation by the MAHs of detachable ("sticky") labels on the vials and syringes of the GdCAs.