

## SCHEDULE

### AMENDMENTS TO OTHER ENACTMENTS

**18.** In the General Product Safety Regulations 2005<sup>(1)</sup> in regulation 33 (duty to notify), in paragraph (10)(b), after “human use”, insert “as amended by Directive [2002/98/EC](#) of the European Parliament and of the Council setting standards of quality and safety for the collection, testing, processing, storage and distribution of human blood and blood components, Commission Directive [2003/63/EC](#) amending Directive [2001/83/EC](#) on the Community code relating to medicinal products for human use, Directive [2004/24/EC](#) of the European Parliament and of the Council amending, as regards traditional herbal medicinal products, Directive [2001/83/EC](#) on the Community code relating to medicinal products for human use and Directive [2004/27/EC](#) of the European Parliament and of the Council amending Directive [2001/83/EC](#) on the Community code relating to medicinal products for human use”.

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<sup>(1)</sup> S.I. 2005/1803.