Regulations under the Health and Social Care Act 2012: Market entry by means of Pharmaceutical Needs Assessments

Information for NHS England
Executive Summary and Chapters 1-4

Introduction, background/overview of the regulatory system, governance arrangements and pharmaceutical lists/terms of service
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Executive Summary and Chapters 1-4

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Executive summary

- Following consultation in the autumn 2008, two clauses were proposed for the Health Bill (now Health Act 2009) to:
  - require PCTs to develop and publish pharmaceutical needs assessments (PNAs); and
  - then to use PNAs as the basis for determining market entry to NHS pharmaceutical services provision.

- In July 2009, a regulatory advisory group drawn from interested parties was set up and started its work to translate these proposals into reality (a list of members of the Advisory Group is at Annex A). The regulations - The National Health Service (Pharmaceutical Services) Regulations 2012 and guidance were a result of their work on the first clause to require PCTs to develop and publish PNAs.

- From 1 April 2013, local authority Health and Well-being Boards (HWBs) will be responsible for developing and updating pharmaceutical needs assessments (PNAs). The NHS Commissioning Board, now known as NHS England will be responsible for the administration of the pharmaceutical services regulations as part of the implementation of the Health and Social Care Act 2012. As a consequence, the 2012 Regulations have been revised and include for the first time the Local Pharmaceutical Services Regulations – they are entitled the NHS (Pharmaceutical and Local Pharmaceutical Services) Regulations 2013 (“the 2013 Regulations”).

- This guidance has been produced to help NHS England in the assessment and determination of applications to provide NHS pharmaceutical services under the new market entry test.

Content

- Chapter 1 is a general introduction to the guidance.

- Chapter 2 discusses the provisions in the NHS Act 2006 relating to pharmaceutical services and gives an overview of the Regulations that provide further details of the requirements these provisions place on NHS England.

- Chapter 3 gives details of the governance arrangements that NHS England will need to put in place in order to discharge their statutory duties set out in the 2013 Regulations. It also provides an overview of the process of determining an application.

- Chapter 4 explains the requirement on NHS England to prepare, maintain and publish lists of contractors who undertake to provide pharmaceutical services within the relevant HWB area.
Market entry by means of pharmaceutical needs assessments - introduction, background/overview of the regulatory system, governance arrangements and pharmaceutical lists/terms of service

It also covers the information that applicants must provide, the fee that must accompany the market entry application and the arrangements that NHS England is required to enter into when including a contractor onto one of their lists.

- Chapter 5 deals with *routine applications* submitted to meet current needs identified within the relevant HWB PNA.

- Chapter 6 deals with *routine applications* submitted to meet future needs identified within the relevant HWB PNA.

- Chapter 7 deals with *routine applications* submitted to secure improvements or better access identified within the relevant HWB PNA.

- Chapter 8 deals with *routine applications* submitted to secure improvements or better access where these were not included within the relevant HWB PNA, i.e. they provide “unforeseen benefits”.

- Chapter 9 deals with *routine applications* submitted to secure future improvements or better access specified within the relevant HWB PNA.

- Chapter 10 deals with the procedures for applications which are “excepted” from the market entry test where contractors wish to relocate to different premises either within the relevant HWB’s area or to a neighbouring HWB’s area and the relocation would not result in significant change to *pharmaceutical services* or local pharmaceutical services.

- Chapter 11 deals with the procedures for applications which are “excepted” from the market entry test where pharmacy contractors wish to provide services where “all persons receiving those services do so otherwise than at those premises” (distance-selling).

- Chapter 12 deals with the procedures for applications which are “excepted” from the market entry test where contractors wish to apply for a “change of ownership”.

- Chapter 13 deals with the procedures for applications which are “excepted” from the market entry test where contractors wish to apply for a “change of ownership” and relocate to different premises either within the relevant HWB area or to a neighbouring HWB area and the relocation would not result in significant change to *pharmaceutical services* or local pharmaceutical services.

- Chapter 14 outlines the procedures for dealing with the provision of *pharmaceutical services* in *controlled localities*.

- Chapter 15 deals with the provision of *pharmaceutical services* by *dispensing doctors*. 
Transitional provisions

- **Schedule 9** to the NHS (Pharmaceutical Services) Regulations 2013 (the “2013 Regulations”) sets out the provisions for matters, which are not finally determined on the “appointed day”. For the purposes the 2013 Regulations, the “appointed day” is 1 April 2013 when the 2013 Regulations come into force.

- In summary, for market entry purposes, outstanding applications to PCT pharmaceutical lists under the 2012 Regulations (including those who by virtue of the transitional provisions within the 2012 Regulations but which were applications under the 2005 Regulations) are preserved – therefore, any matters that are ongoing on 1 April 2013 continue.
Chapter 1: Introduction

Status of advice

1. This guidance is a working document and may be subject to change as and when there are amendments to The National Health Service (Pharmaceutical Services and Local Pharmaceutical Services) Regulations 2013 (referred to in this guidance as the “2013 Regulations”) or associated legislation.

2. The primary purpose of this document is to help all those working in NHS England with the task of determining applications relating to the provision of pharmaceutical services in England from 1 April 2013. This document is also intended to be of assistance to all others who are affected by such decisions.

3. The law on the subject is complex and contained in Acts of Parliament, Regulations and case law from the courts. Additionally over time, decisions made by the NHS Litigation Authority’s Family Health Services Appeal Unit (FHSAU) will need to be taken into account by NHS England when determining applications. This document is designed to provide staff at all levels with information on the relevant legal provisions and interpretations of those provisions. It is also intended to provide practical advice in relation to the operation of the legal provisions.

4. Although this document contains a lot of detailed reference in the footnotes to the legal provisions, the rules themselves are not, in the main, set out word for word in this guidance. In order to make the document easier to read, the detailed rules have, in most cases, been paraphrased. However, all those responsible for administering or applying the law must bear in mind that it is the law that must be applied, not the interpretation that is set out below.

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1 Reference is made in the Regulations to the determination of applications. Effectively, this means making a decision on an application, i.e. approval, conditional approval, deferral or refusal.
5. This document’s intended legal status is that it is non-statutory guidance, designed to assist NHS England in reaching decisions within the framework of the law. It is not an authoritative statement of the law. In practice, there is no substitute for referring to the law itself, or seeking professional advice as to what the law says and how it applies in particular circumstances. It is essential to understand that decisions must be taken in accordance with the law, and not simply based on the analysis and advice contained in this guidance (or indeed any other commentary on the law). Furthermore, although it is hoped that NHS England will find this guidance helpful, the Department’s view is that NHS England is not obliged to take this guidance into consideration when formulating its decisions. NHS England’s own understanding of the law is fundamentally a matter for it and where it is in doubt, it should seek legal advice.

Previous guidance and transitional arrangements

6. This document relates to applications received under the 2013 Regulations from 1 April 2013 onwards and replaces the original control of entry guidance produced by the Department on 17 September 2009 to accompany the NHS (Pharmaceutical Services) Regulations 2005 (referred to in this guidance as the 2005 Regulations) and the more recent guidance to the NHS (Pharmaceutical Services) Regulations 2012 (the “2012 Regulations”). NHS England should retain that previous guidance, however, as there may be applications that by virtue of the transitional provisions within the 2012 Regulations fell to be dealt with under the 2005 Regulations even after the 2013 Regulations have come into force. This includes any appeals that may arise from decisions on such applications.

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2 It should be noted that in a 2008 Court of Appeal decision, Lord Justice Lawrence Collins stated that “if the Secretary of State issues non-statutory guidance for decision-makers, and there is a radical departure from the guidance, then, although not relevant to the construction of the relevant provisions, the guidance may be relevant to a challenge because the decision-maker may be under an obligation to take it into account and to explain why he has taken that radically different approach.” (Assura Pharmacy Ltd and NHS Litigation Authority (Family Health Services Appeal Unit) and E Moss Ltd (trading as Alliance Pharmacy) December 2008 – available on [http://www.bailii.org/ew/cases/EWCA/Civ/2008/1356.html](http://www.bailii.org/ew/cases/EWCA/Civ/2008/1356.html)). In the light of this, the Department has sought to make its own view clear that decision-makers are not bound to take this particular example of non-statutory guidance into account. However, as Lord Justice Sedley notes in his judgment in Assura, it is currently unresolved at appellate level how an independent tribunal should treat departmental guidance given otherwise than under statutory authority, and reserves his view on the matter to a case where the issue is pivotal. It seems likely therefore that this issue will come up for further judicial consideration in the future.

The Regulations

7. The 2013 Regulations, Statutory Instrument (SI) 2013/349 replace the 2012 Regulations with effect from 1 April 2013. NHS England should ensure it has access to these Regulations to ensure they are acting within the law when determining applications.

8. As with the 2012 Regulations, it is possible that the 2013 Regulations will be amended over time and staff at NHS England should ensure they have access to an up-to-date version of the Regulations.

Transitional provisions

9. Schedule 9 to the NHS (Pharmaceutical Services) Regulations 2013 sets out the provisions for matters, which are not finally determined on the “appointed day”. For the purposes the 2013 Regulations, the “appointed day” is 1 April 2013 when the 2013 Regulations come into force.

10. The transitional arrangements are in summary, for market entry purposes, outstanding applications to PCT pharmaceutical lists under the 2012 Regulations (including those who by virtue of the transitional provisions within the 2012 Regulations but which were applications under the 2005 Regulations) are preserved – therefore, any matters that are ongoing on 1 April 2013 continue.

Pharmaceutical Needs Assessments

11. This document does not cover those regulations (regulations 3-9) and Schedule 1 that relate to HWB pharmaceutical needs assessments (PNAs). Separate guidance has been produced to assist HWBs in understanding the requirements of these.

Other guidance documents

12. Other guidance has been produced to assist NHS England in understanding the requirements of the 2013 Regulations. These include fitness to practise, performance related sanctions including market exit and the charging of fees for applications (see Annex F).
Structure of the document

13. The document is structured so that each type of application that NHS England may receive has its own chapter. NHS England will then be able to work through the correct process for each type of application without having to cross-reference to other chapters. This does mean that the document as a whole is repetitive in parts but it is hoped that the repetition of certain elements within each chapter provides a clearer process which is easy to follow.

14. Throughout the document, where reference is made to another document, the web address will be given. Where documents are Department of Health publications, the Gateway reference will also be given.

15. Within Regulation 2 of the 2013 Regulations, there is a list of definitions. Where one of these terms is used in this document, it will appear in italics so that readers are able to understand each term in the correct context. Annex A contains a glossary of such terms.
Chapter 2: Background and overview of the Regulatory system

1. This chapter discusses the provisions within the National Health Service Act 2006\(^4\) (the 2006 Act) relating to pharmaceutical services and gives an overview of the Regulations that provide further details of the requirements these provisions place on NHS England.

2. This document provides guidance on the following Parts of the Regulations:

- Part 3 – general matters relating to pharmaceutical lists and applications in respect of them;
- Part 4 – excepted applications;
- Part 5 – specific grounds for refusal or deferral of applications that are not linked to fitness to practise grounds;
- Part 6 – refusal, deferral and conditional inclusion in a pharmaceutical list on fitness to practise grounds;
- Part 7 – controlled localities, reserved locations and pharmacies in these areas;
- Part 8 – dispensing doctors; and
- Schedules 2 (procedures) and 3 (appeals).

Arrangements for pharmaceutical services

3. Sections 126 to 133 inclusive of the 2006 Act cover the provision of pharmaceutical services.

4. Section 126 of the 2006 Act places an obligation on NHS England to put arrangements in place so that drugs, medicines and listed appliances ordered via NHS prescriptions can be supplied to persons. This section makes provision for the types of healthcare professional whom the Regulations may allow to order drugs, medicines and listed appliances on an NHS prescription.

5. Appliances must be listed in the Drug Tariff in order to be prescribed on NHS prescriptions and are therefore referred to as listed appliances\(^5\).

6. This duty on NHS England to make arrangements set out in Section 126 of the 2006 Act is alongside a duty on the Secretary of State to make Regulations to facilitate NHS England to discharge this duty.

7. Section 126 requires NHS England to ensure that they have arrangements in place for:

- the provision of proper and sufficient drugs, medicines and listed appliances which are ordered on NHS prescriptions by doctors;
- the provision of proper and sufficient drugs, medicines which are ordered on NHS prescriptions by dentists;
- the provision of proper and sufficient drugs, medicines and listed appliances which are ordered on NHS prescriptions by other specified descriptions of healthcare professionals; and
- such other services that may be prescribed.

8. It is this duty that underpins the provision of those services set out in Schedules to the Regulations by pharmacy contractors and dispensing appliance contractors. Section 132 makes provision for regulations that allow for arrangements to provide dispensing services by GPs. Their terms of service are set out in Schedule 6 to the Regulations.

9. Section 127 of the 2006 Act allows the Secretary of State to make provision for other services in Directions. It is from this Section that advanced and enhanced services are derived. The requirements for these services are set out in Directions and are referred to as directed services in the 2013 Regulations.

10. The 2006 Act allows for arrangements to be made for the provision of three types of pharmaceutical services, depending on the type of contractor providing the services:

- essential\(^6\) services, other mandatory arrangements, advanced services and enhanced services provided by pharmacy contractors;

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\(^5\) The exhaustive list of appliances that may be supplied under the Regulations is set out in Part IX of the Drug Tariff. Any drug/medicine may be supplied unless it is included in the selected list set out in Schedule 1 to the NHS (General Medical Services Contracts)(Prescription of Drugs etc) Regulations 2004, which is reproduced in Part XVIIIA of the Drug Tariff. There is also a “grey list” of drugs that GPs should only prescribe on the NHS under restrictive conditions.

\(^6\) Essential services are defined as those set out in paragraph 3 in Schedule 4 and does not include other mandatory arrangements that apply to all pharmacy contractors, such as clinical governance arrangements. The term “essential services” is not used to describe those services provided by dispensing appliance contractors as set out in Schedule 5.
• services set out in Schedule 5 and advanced services as provided by dispensing appliance contractors; and
• dispensing services provided by dispensing GPs as set out in Schedule 6.

11. Within the 2013 Regulations, the definition of pharmaceutical services differs between the Parts and Schedules:

• Part 2 and Schedule 1 – provisions regarding PNAs – as per the definition in paragraph 10 above, plus local pharmaceutical services (LPS);
• Parts 7 and 8 and Schedule 6 – provisions regarding controlled localities, reserved locations and dispensing doctors, and the terms of service for dispensing doctors – in the context of service provision by services provided by dispensing doctors, pharmaceutical services means the dispensing services provided by dispensing GPs as set out in Schedule 6. In the context of service provision by pharmacy contractors or appliance contractors, it means the services mentioned in relation to those types of contractor in paragraph 10 above, but not LPS; and
• Parts 3 to 6 and 9 to 13 – as per the definition in paragraph 10 above, although in practice, services provided by dispensing GPs will generally not be relevant where pharmaceutical services is used in these Parts, as these Parts relate almost exclusively to market entry and performance sanctions for pharmacy and dispensing appliance contractors.

12. Originally, the underlying policy objective of the legislation was that there should be a distinction between those who prescribe drugs and those who dispense drugs. This principle has, however, always been subject to exceptions, most notably in rural areas – which are known in the Regulations as controlled localities. It is increasingly less distinct as more health professionals, including pharmacists, are able to qualify as prescribers in order to improve access to, and choice of, services for patients.

Regulations for the provision of pharmaceutical services

13. Detailed Regulations relating to the arrangements for securing the provision of these services are made under the authority principally of Section 129 of the 2006 Act. Section 129 (now amended by the Health and Social Care Act 2012) was amended by the Health Act 20097 to reflect the move towards using PNAs as the basis for determining applications for inclusion in a pharmaceutical list and is reproduced in Annex B for ease of reference. In summary, Section 129 makes provisions for Regulations that will govern the provision of pharmaceutical services to assist NHS England in discharging its duty set out in Section 126.

7 http://www.opsi.gov.uk/acts/acts2009/ukpga_20090021_en_1
14. Section 129 also includes provision for Regulations to deal with the process of creating a pharmaceutical list, removing persons from such a list and setting the entry requirements of those entitled to be entered on a list.

15. The drugs, appliances and chemical reagents to be supplied within the NHS and payments for such are listed in the Drug Tariff. The Drug Tariff, which is a statutory requirement under Section 164 is available from The Stationery Office and is available on the NHS Business Service Authority website\(^8\).

16. There are two modes of provision of pharmaceutical services in the Regulations:

(a) by chemists (and in the context of the 2013 Regulations this covers individual pharmacists, partnerships, bodies corporate and suppliers of appliances); and

(b) by doctors (see Chapter 15 of this guidance).

**Market entry test**

17. The market entry test describes the system whereby NHS England assesses an application that offers to:

- meet an identified current or future need or needs;
- meet identified current or future improvements or better access to pharmaceutical services; or
- provide unforeseen benefits, i.e. applications that offer to meet a need that is not identified in a PNA but which NHS England is satisfied would lead to significant benefits to people living in the relevant HWB area.

\(^8\) [http://www.ppa.org.uk/ppa/edt_intro.htm](http://www.ppa.org.uk/ppa/edt_intro.htm)
18. **Section 129 (2A)** sets out the market entry test as follows:

<table>
<thead>
<tr>
<th>Provision within the Act</th>
<th>Explanation</th>
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<tr>
<td>(2A) The Board is satisfied as mentioned in this subsection if, having regard to the needs statement for the relevant area and to any matters prescribed by the Secretary of State in the regulations, it is satisfied that to grant the application would meet a need in that area for the services or some of the services specified in the application, or secure improvements, or better access, to pharmaceutical services in that area.</td>
<td>Section 129(2A) requires Regulations to make provision for NHS England to approve applications where they are satisfied that to grant the application would meet a need identified within in the HWB PNA. The Regulations may provide overriding reasons for NHS England not to approve such an application, for example, where there are concerns relating to the applicant’s fitness to practise.</td>
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**Types of applications**

There are two types of application that can be made by pharmacy or dispensing appliance contractors within the 2013 Regulations:

- ***routine applications***; and
- ***excepted applications***.

19. **Regulation 12** defines the types of applications that are termed *routine application*. These are applications that are submitted under Part 3 of the Regulations, namely applications:

- to be included in a pharmaceutical list by persons not already included in it; and

applications by persons already included on a pharmaceutical list to:

- open, within the relevant HWB area, additional premises from which to provide the same or different *pharmaceutical services*;
- relocate to different premises and to provide the same or different *pharmaceutical services* from those new premises; and

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9 When PCTs were responsible for developing and updating PNAs, they were required to grant applications where the PNA had identified a need or improvement. After 1 April 2013, as HWBs will be responsible for updating and developing PNAs, NHS England has a discretion as to whether or not they grant the application if a need is identified in a HWB PNA.
• provide services in addition to those that are already provided at the listed chemist premises.

20. In practice, contractors and NHS England may rarely use the 2013 Regulations to submit and consider applications to provide additional services to those that they already provide (Regulation 12(b)(iii)). It is more likely that NHS England will use other processes when commissioning enhanced services, but they should be aware of this route and ensure they have processes to determine such applications. If an application to provide directed services is received, NHS England has a duty to determine it unless it is withdrawn by the applicant.

21. Section 129(2A) does not apply to some types of application and these are known as excepted applications.

22. Provision for these types of application are included in Part 4 of the Regulations and include:

• applications to provide directed services (Regulation 23);
• relocations that do not result in significant change to pharmaceutical services provision (Regulation 24);
• change of ownership applications (Regulation 26);
• applications for temporary listings arising out of suspensions (Regulation 27);
• applications from persons exercising a right of return to a pharmaceutical list (Regulation 28); and
• applications relating to emergencies requiring the flexible provision of pharmaceutical services (Regulation 29).

Applications to relocate to new premises

23. NHS England should note that there are two different types of relocation applications.

24. The first type is a routine application that falls under Regulation 12(b)(ii). This is an application to relocate to new premises in order to meet a need identified within the PNA, and which would result in a significant change to pharmaceutical services provision in the relevant HWB area.
Example

The lease on the premises of a contractor is due to expire and the new rent would make the pharmacy financially unviable. The premises are on the High Street of a town. Within the HWB PNA, there is an identified current need for a pharmacy within a new housing development on the other side of the town. The contractor could therefore submit an application under Regulation 12(b)(ii) to relocate from their current premises to new premises within the housing development and offer to provide the services that the PNA identifies are needed at that new location.

25. Where NHS England receives an application to relocate under Regulation 12(b)(ii), then apart from not needing further fitness to practise information from the applicant, they should treat it as though it was an application to join a pharmaceutical list to meet a need identified within the HWB PNA. Chapters 5 to 9 of this guidance set out how to process and determine such an application.

26. The second type is an excepted application that falls under Regulation 24. In order to meet the requirements of Regulation 24, the relocation must not result in a significant change to pharmaceutical services provision.

Example

The lease on the premises of a contractor is due to expire and the building is due to be demolished. The contractor identifies suitable premises 200 m down the road and therefore submits an application under Regulation 24 to relocate to these new premises. Further information on this type of application can be found in Chapter 10.

Rural dispensing

27. Where NHS England has determined that an area is “controlled” (i.e. rural in character - see Chapter 14), provided certain conditions are met, doctors as well as pharmacists can dispense NHS medicines. GPs, may, in general, dispense NHS prescriptions only with NHS England approval and only to their own patients who live in such controlled localities and live more than 1.6 km (as the crow flies) from a pharmacy. The main purpose of this is to ensure patients in rural areas who might have difficulty getting to their nearest pharmacy can access the dispensed medicines they need.

28. A GP who wishes to apply to dispense to patients need only show that to do so would not prejudice the proper provision of relevant NHS services locally (known as the “prejudice test”). Relevant NHS services for these purposes are medical, pharmaceutical or local pharmaceutical services provided by other contractors.
29. The “prejudice test” will also apply to applications from pharmacy contractors in rural areas and further information can be found in Chapter 14.

30. NHS England may determine reserved locations in controlled localities (see Chapter 14 for further information). If an application is made for new pharmacy premises in a reserved location, only the market entry test is applied to the application, not the prejudice test.

31. Further information on rural dispensing can be found in Chapter 14 and 15 of this document.

Appeals against NHS England decisions on pharmaceutical applications

32. Under the Regulations, most NHS England decisions on market entry applications are appealable to the Secretary of State for Health who has delegated this responsibility to the NHS Litigation Authority (NHSLA). The appellate function is undertaken by the NHSLA’s Family Health Services Appeal Unit (FHSAU). More information about their work is available on their website10.

33. **Schedule 3** of the 2013 Regulations sets out the actions that the FHSAU may take. In summary, the FHSAU may generally:

- confirm the decision or determination of NHS England;
- quash the decision or determination of NHS England and re-determine the application;
- substitute its decision or determination for any decision or determination NHS England could have made; or
- quash NHS England’s decision and remit the matter to NHS England for it to re-determine the application.

34. Where the FHSAU remits the matter back to NHS England, this is generally where there have been procedural concerns of an administrative nature (i.e. NHS England has not followed a procedure set down in the 2013 Regulations).

10 [http://www.nhsla.com/FHSAU/Decisions/Pharmacy+2005+Regulations/]
35. For the purposes of the 2013 Regulations, the FHSAU’s decision becomes NHS England’s decision on the matter. The FHSAU’s decision may only be overruled by a court.

36. NHS England may find it useful periodically to view the FHSAU’s decisions for learning and training purposes.
Chapter 3: Governance arrangements

1. This chapter gives details of the governance arrangements that need to be put in place in order for NHS England to discharge its statutory duties set out in the 2013 Regulations. It also provides an overview of the process of determining an application. Further detailed information on this process can be found in the subsequent chapters.

Decision-making process

2. It is imperative that NHS England has robust decision-making processes for market entry decisions including an assurance that there are no conflicts of interest. It is good practice that terms of reference are developed for the panel that makes the key decisions and that these are made open and transparent with all stakeholders involved in the process. It is especially important for the panel involved in making the key decisions to be clear on their remit and responsibilities.

Decision-making group

3. Those persons who are involved in the determination or deferral of an application must have access to the 2013 Regulations and any amending Regulations. NHS England should also consider the training that such persons receive and should ensure that members of panels are advised of subsequent amendments and receive training where this is appropriate.

4. Paragraph 26, Schedule 2 sets out a list of persons who must take no part in determining or deferring any routine or excepted application. They are:

- a contractor included in a pharmaceutical list or any employee of such a contractor;
- a person who assists in the provision of pharmaceutical services under Part 1 of Chapter 7 of the 2006 Act;
- Local Pharmaceutical Services (LPS) contractors, or someone who provides or assists in the provision of LPS;
- providers of primary medical services;
- members of a provider of primary medical services that is a partnership, or shareholders in a provider of primary medical services if that provider is a company limited by share;
- persons employed or engaged by a primary medical services provider; or
- persons employed or engaged by an (Alternative Provider Medical Services) APMS contractor in any capacity relating to the provision of primary medical services.
5. These persons must take no part in decisions, whether their involvement would give rise to a reasonable suspicion of bias or not.

6. Additionally, no person with an interest or association in the application may take part in determining or deferring a particular application where their involvement would give rise to a reasonable suspicion of bias. For example, a lay member who sits on the panel should be excluded from taking part in the determination of an application which has been submitted for the village they live in.

7. Furthermore, any person who may be subject to pressure about a particular application must not take part in the determining or deferral of it (paragraph 26(2) of Schedule 2). NHS England should therefore check the membership of the panel that considers applications to ensure it complies with this paragraph. Additionally, NHS England may wish to seek declarations of conflict of interest from members at the beginning of the meeting and to record any such declarations formally.

8. Panels may wish to take professional advice from persons who are excluded by virtue of paragraph 26 of Schedule 2 to aid the determination or deferral of an application. Such persons are able to give advice to the panel, but must not be seen to be part of the determination or deferral of an application. Hence, if any persons are present to give professional advice to the panel, they should withdraw before any vote on a decision is made. It is important to take account of the provisions of Article 6 of the Human Rights Act 1998 (right to a fair trial), as well as general public law considerations of fairness, when deciding whether to include such people.

9. Where professional advice is sought, it is good practice to declare at the start of any panel meeting that these individuals are excluded from any determination or deferral of an application. It is also good practice to make a note in the minutes that any professional advisers have retired from the room before decisions are finalised and voting takes place.

10. The importance of these points is that if this is not followed, an appeal or judicial review could be based on an argument that due process has not been followed and that inappropriate persons have been party to the determination of deferral of an application. Equally, however, the panel has a duty to make a sufficient enquiry into the matter before it, so that it has a sufficiently full picture for a fair decision to be made. Therefore, sometimes, obtaining the advice of professional advisers will be unavoidable.

11. Appeals against NHS England decisions to the NHSLA are difficult to avoid, but with appropriate and robust decision-making processes, their likelihood of success can be minimised. Appeals procedures can be difficult and require a lot of time and effort.

12. NHS England need to be sure that data protection and confidentiality is monitored throughout the whole process and sound administrative and audit processes for applications are important.
13. Some decisions may be delegated to an officer of NHS England and need not go to a full panel meeting for approval, for example, where a change of ownership application has been received. If it meets the requirements of the Regulations and no issues have been identified in any fitness to practise checks, NHS England may decide it is appropriate for an officer to determine the application instead of the panel.

**Oral hearings**

14. Oral hearings are not required to be held for all decisions and NHS England should make a judgement on when and for what type of decisions an oral hearing is necessary. This is likely to be based on the complexity of the application, previous applications in the locality, the number and type of representations made in respect of the application from those notified of it, and decisions of the NHSLA.

15. Where NHS England decides to hold an oral hearing, the procedure to be followed is set out in paragraph 25 of Schedule 2.

**Notification of decision**

16. NHS England must make a decision on the application as soon as is practicable and no later than:

- four months after receiving the application and all supporting information or documentation, if it is one that is *notifiable* (see later chapters); or
- within 30 days if the application is not a *notifiable* one (see later chapters).

(Paragraph 27 of Schedule 2)

17. Once the decision is made, NHS England must *notify* certain persons of that decision (paragraph 28 of Schedule 2). The decision letter must include the reasons for the decision (paragraph 28(6) of Schedule 2), and should not simply say that the application has been granted or refused.

18. Where an application is approved, NHS England is also required to send a template *notice of commencement* with the decision letter (paragraph 29 of Schedule 2). NHS England may choose to pre-fill the template with the relevant data, but in any case, the applicant is required to complete and return the template to NHS England fourteen days prior to their intention to commence the provision of *pharmaceutical services* (paragraph 34(2) of Schedule 2). Commencement must begin within the period for which an application is approved.
Appeals against NHS England decisions on pharmaceutical applications

19. The main procedures regarding appeals are set out in **Schedule 3**.

20. An appeal must be made in writing to the NHSLA within 30 days from the date on which NHS England decision letter is sent. Appeals are heard by the Family Health Services Appeal Unit (FHSAU) on behalf of the Secretary of State for Health.

21. In order to be valid, the appeal must contain a concise and reasoned statement of the grounds for appeal. Appeals may be sent by e-mail attachment initially, but should always be followed up by hard copies by letter or by fax.

22. An appeal under Schedule 3 can generally only be made by the applicant or by a contractor who has been notified of the decision by NHS England. There are some exceptions to this relating to **controlled localities** and **reserved locations** (see Chapters 14 and 15 for further information).

23. Apart from some **controlled locality** and **reserved location** appeals, an appeal cannot be made, for example, by a Local Pharmaceutical Committee (LPC) or Local Medical Committee (LMC). Where more than one appeal is received in relation to a decision, the FHSAU can determine them at the same time.

24. Market entry appeals are determined by the Pharmacy Appeals Committee of the FHSAU. Some other types of appeal are determined by a senior officer of the FHSAU.

25. The FHSAU’s deliberations are not limited to simply reviewing NHS England’s decision. It may determine the appeal by reconsidering the application *de novo* (i.e. from the beginning).

26. The majority of cases are decided based on correspondence with the FHSAU and other documentation related to the original decision. Occasionally, for example, if there are material differences in the facts presented by the parties, the FHSAU will convene a panel to hold an oral hearing.

27. The FHSAU must give at least 14 days’ notice of the hearing to the appellant and a list of other persons who are entitled to make oral representations at the hearing. This list includes:

- the applicant (if different from the appellant);
- NHS England;
- the relevant local representative committees; and
any “additional presenters” – that is, any person to whom a copy of the notice of appeal is sent who made written representations on the appeal in the course of which they indicated they would wish to make oral representations if there were a hearing, provided that the FHSAU is satisfied that it would be desirable to hear further evidence from them at the hearing.

28. The appellant and any person who whom a notice of the hearing is sent can attend with any representatives they wish to accompany them.

29. The FHSAU’s target is to ensure all appeals are processed within 26 weeks. For those appeals determined without an oral hearing, the target is to process appeals within 15 weeks.

30. Whilst there are standard complaints procedures that apply to any NHS body, there are no further powers for review of an appeal decision once it has been issued. The FHSAU’s decision can only be set aside by the High Court.
Chapter 4: Pharmaceutical lists and terms of service

1. This chapter explains the requirement on NHS England to prepare, maintain and publish lists of contractors who undertake to provide pharmaceutical services within the relevant HWB area. It also covers the information that applicants must provide, the fee that must accompany the market entry application and the arrangements that NHS England are required to enter into when including a contractor on a list.

Pharmaceutical lists

2. NHS England required to prepare, maintain and publish lists of persons (other than GPs and dentists) who undertake to provide pharmaceutical services from premises in the relevant HWB area (Regulation 10(1)). It should be noted that although the regulation makes reference to persons being included on a list, the word “persons” refers to the contractor. In that respect, a pharmaceutical list differs from the “performer lists” that NHS England hold for individual GPs, dentists and optometrists. Currently, there is no pharmaceutical equivalent of the performer list.

3. Regulation 10(1) requires NHS England to prepare, maintain and publish:

- a list of persons who undertake to provide pharmaceutical services in the relevant HWB area, in particular by way of the provision of drugs and the appliances that they supply in their normal course of business, i.e. community pharmacies run by sole traders, partnerships or bodies corporate (Regulation 10(2)(a)); and

- a list of persons who undertake to provide pharmaceutical services only by way of the provision of appliances, i.e. dispensing appliance contractors (Regulation 10(2)(b)).

4. It should be noted that community pharmacies that also provide a range of appliances should only be included on the first list; they do not also need to be included in the list of persons providing appliances.

5. A list must contain as a minimum, the following information:

- the contractor’s name (this should not be the trading name, rather the name of the organisation that is running the pharmacy);

- the address of the premises from where the contractor has undertaken to provide pharmaceutical services – the listed chemist premises (Regulation 10(3)(a)); and

- the days and times at which services are provided, during both core opening hours (Regulation 10(3)(b)(i)) and supplementary opening hours at those premises (Regulation 10(3)(b)).
6. **Regulation 10 (3)** sets out the minimum information requirements of a list and NHS England is free to add additional information to a list and may wish to include the following:

- the trading name of the pharmacy; and
- any *advanced* and *enhanced services* that are provided.

7. **Regulation 10(4)** also requires NHS England to prepare, maintain and publish, a list of all *NHS chemists* in the HWB area who participate in the electronic prescription service (EPS) service. Included in this list is the address of any premises at which the EPS service is provided (**Regulation 10(5)**).

8. In addition to these two lists, NHS England is also required to prepare and publish a list of any *dispensing doctors* in the HWB area (**Regulation 46(1)**). Further information on this list can be found in Chapter 15.

NHS England should ensure that each HWB has access to these lists, which is sufficient to enable the HWB to carry out its functions under the 2013 Regulations (**Regulation 10(6)**).

### Inclusion in or amendment to a pharmaceutical list – information to be provided

9. **Regulation 10(7)** refers to Schedule 2 of the Regulations. **Part 1 of Schedule 2** contains the information to be supplied by a person who is:

- seeking inclusion in a pharmaceutical list that they are not already included in; or
- included in a pharmaceutical list and is seeking to:
  - open additional premises from which to provide the same or different *pharmaceutical services*;
  - to relocate to different premises to provide the same or different *pharmaceutical services*; or
  - to provide additional services to those they are already listed to provide at their *listed chemist premises*.

10. Where the applicant is not already included in a pharmaceutical list, they are required to submit information on their fitness to practise to NHS England (for further details, please see guidance on fitness to practise). NHS England must ensure that where they need to come to a decision as to whether the applicant is suitable to be included on a pharmaceutical list or not, this decision must be made before considering the market entry application.
11. Schedule 2 also sets out the procedure to be followed by applicants when they make an application and other related matters such as notification and determination of applications by NHS England.

12. The Regulations do not set out a template form to use for the purposes of providing this information. However, applications should include such information as NHS England considers necessary for it to be able to determine the application. Paragraph 11 of Schedule 2 sets out the procedure to be followed where not all the relevant information and documentation is provided to NHS England.

13. It should be noted, however, that applicants would not be obliged to use NHS England and NHS England cannot refuse to determine an application simply because the information has been provided in a different format.

Terms of service: general

14. When including a contractor on a pharmaceutical list, Regulation 11 requires NHS England to ensure that the terms of their inclusion include the following:

- their terms of service as set out in Schedules 4 (community pharmacy contractors) or Schedule 5 (dispensing appliance contractors);
- any obligation in the 2013 Regulations that is only applicable if the contractor is a person to whom the obligation is applicable, for example, the 100 hours per week condition that is applicable to some pharmacies that relied on that exemption under the 2005 Regulations;
- the conditions set out in the Drug Tariff, where applicable;
- the terms of the arrangement for the provision of any directed services; and
- Regulation 3 of the Local Involvement Networks (Duty of Services Providers to Allow Entry) Regulations 2008\(^\text{11}\) as it applies to pharmacy and dispensing appliance contractors.

15. Additionally, where NHS England has placed fitness to practise conditions on the inclusion of that contractor under Regulation 35 or Chapter 6 of Part 7 of the 2006 Act, the arrangements must make reference to those conditions.

16. NHS England should, therefore, ensure that when writing to notify applicants that their application has been approved, reference is made to these requirements.

17. This requirement applies equally to both pharmacy and appliance contractors. Regulation 47 makes similar provision for dispensing doctors (see chapter 14).

Charges for applications

18. Since April 2008, NHS England has been required to charge for certain types of application. Details on this are set out in Directions and guidance produced to assist NHS England.

19. Applicants should, therefore, ensure they submit the information set out in Part 1 of Schedule 2 where applicable and also the appropriate fee. Until all these elements are received, the application is deficient and they should not begin to process the application.

20. The fees are non-refundable as they are a contribution towards NHS England’s costs of processing and determining the application.

21. During the passage of the Health Act 2006 a general review, after 18 months of implementation, of charging for NHS pharmaceutical services applications was promised (September 2009). The review looked at progress in implementing charging as well as the impact of charging on the NHS, pharmacy and appliance contractors and applicants. It also sought to find out whether the current fee levels were fair and reasonable.

22. The Government published the outcome of the review in October 2011. The Government noted that the experience of the effect of charging by the NHS is variable. However, chemist contractors did not feel significantly affected by charging for applications. Thus, the fees for applications were retained. The Government also noted that most respondents felt that the current fee levels were fair and reasonable. Therefore, it did not feel there was sufficient evidence to change current fee levels.

23. Fees will continue to be paid for applications under the 2013 Regulations. Further details of this are set out in The Pharmaceutical Services (Fees for Applications) Directions 2013 and guidance to these Directions is included in Annex E.